

THE BIOLOGICAL BULLETIN

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VOLUME 135
JULY TO DECEMBER, 1968

Printed and Issued by
LANCASTER PRESS, Inc.
PRINCE & LEMON STS.
LANCASTER, PA.



THE BIOLOGICAL BULLETIN is issued six times a year at the Lancaster Press, Inc., Prince and Lemon Streets, Lancaster, Pennsylvania.

Subscriptions and similar matter should be addressed to The Biological Bulletin, Marine Biological Laboratory, Woods Hole, Massachusetts. Agent for Great Britain: Wheldon and Wesley, Limited, 2, 3 and 4 Arthur Street, New Oxford Street, London, W. C. 2. Single numbers, \$3.75. Subscription per volume (three issues), \$9.00.

Communications relative to manuscripts should be sent to Dr. W. D. Russell-Hunter, Marine Biological Laboratory, Woods Hole, Massachusetts 02543.

Second-class postage paid at Lancaster, Pa.

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THE BIOLOGICAL BULLETIN

PUBLISHED BY THE MARINE BIOLOGICAL LABORATORY

THE MARINE BIOLOGICAL LABORATORY

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II. ACT OF INCORPORATION

No. 3170

COMMONWEALTH OF MASSACHUSETTS

Be It Known, That whereas Alpheus Hyatt, William Sanford Stevens, William T. Sedgwick, Edward G. Gardiner, Susan Minns, Charles Sedgwick Minot, Samuel Wells, William G. Farlow, Anna D. Phillips, and B. H. Van Vleck have associated themselves with the intention of forming a Corporation under the name of the Marine Biological Laboratory, for the purpose of establishing and maintaining a laboratory or station for scientific study and investigation, and a school for instruction in biology and natural history, and have complied with the provisions of the statutes of this Commonwealth in such case made and provided, as appears from the certificate of the President, Treasurer, and Trustees of said Corporation, duly approved by the Commissioner of Corporations, and recorded in this office;

Now, therefore, I, HENRY B. PIERCE, Secretary of the Commonwealth of Massachusetts, *do hereby certify* that said A. Hyatt, W. S. Stevens, W. T. Sedgwick, E. G. Gardiner, S. Minns, C. S. Minot, S. Wells, W. G. Farlow, A. D. Phillips, and B. H. Van Vleck, their associates and successors, are legally organized and established as, and are hereby made, an existing Corporation, under the name of the MARINE BIOLOGICAL LABORATORY, with the powers, rights, and privileges, and subject to the limitations, duties, and restrictions, which by law appertain thereto.

Witness my official signature hereunto subscribed, and the seal of the Commonwealth of Massachusetts hereunto affixed, this twentieth day of March, in the year of our Lord One Thousand Eight Hundred and Eighty-Eight.

[SEAL]

HENRY B. PIERCE,
Secretary of the Commonwealth.

III. BYLAWS OF THE CORPORATION OF THE MARINE
BIOLOGICAL LABORATORY

(Revised August 12, 1966)

I. The members of the Corporation shall consist of persons elected by the Board of Trustees.

II. The officers of the Corporation shall consist of a Chairman of the Board of Trustees, President, Director, Treasurer and Clerk.

III. The Annual Meeting of the members shall be held on the Friday following the second Tuesday in August in each year at the Laboratory in Woods Hole, Massachusetts, at 9:30 A.M., and at such meeting the members shall choose by ballot a Treasurer and a Clerk to serve one year, and nine Trustees to serve four years, and shall transact such other business as may properly come before the meeting. Special meetings of the members may be called by the Trustees to be held at such time and place as may be designated.

IV. Twenty-five members shall constitute a quorum at any meeting.

V. Any member in good standing may vote at any meeting, either in person or by proxy duly executed.

VI. Inasmuch as the time and place of the Annual Meeting of members are fixed by these bylaws, no notice of the Annual Meeting need be given. Notice of any special meeting of members, however, shall be given by the Clerk by mailing notice of the time and place and purpose of such meeting, at least (15) days before such meeting, to each member at his or her address as shown on the records of the Corporation.

VII. The Annual Meeting of the Trustees shall be held promptly after Annual Meeting of the Corporation at the Laboratory in Woods Hole, Massachusetts. Special meetings of the Trustees shall be called by the Chairman, the President, or by any seven Trustees, to be held at such time and place as may be designated, and the Secretary shall give notice thereof by written or printed notice, mailed to each Trustee at his address as shown on the records of the Corporation, at least one (1) week before the meeting. At such special meeting only matters stated in the notice shall be considered. Seven Trustees of those eligible to vote shall constitute a quorum for the transaction of business at any meeting.

VIII. There shall be three groups of Trustees:

(A) Thirty-six Trustees chosen by the Corporation, divided into four classes, each to serve four years. After having served two consecutive terms of four years each, Trustees are ineligible for re-election until a year has elapsed.

(B) Trustees *ex officio*, who shall be the Chairman, the Director, the Treasurer, and the Clerk.

(C) Trustees *Emeriti*, who shall be elected from present or former Trustees by the Corporation. Any regular Trustee who has attained the age of seventy years shall continue to serve as Trustee until the next Annual Meeting of the Corporation, whereupon his office as regular Trustee shall become vacant and be filled by election by the Corporation and he shall become eligible for election as Trustee *Emeritus* for life. The Trustees *ex officio* and *Emeriti* shall have all the rights of the Trustees, except that *Trustees Emeriti* shall not have the right to vote.

The Trustees and officers shall hold their respective offices until their successors are chosen and have qualified in their stead.

IX. The Trustees shall have the control and management of the affairs of the Corporation. They shall elect a Chairman of the Board of Trustees who shall be elected annually

and shall serve until his successor is selected and qualified and who shall also preside at meetings of the Corporation. They shall elect a President of the Corporation who shall also be the Vice Chairman of the Board of Trustees and Vice Chairman of meetings of the Corporation, and who shall be elected annually and shall serve until his successor is selected and qualified. They shall appoint a Director of the Laboratory for a term not to exceed five years, provided the term shall not exceed one year if the candidate has attained the age of 65 years prior to the date of the appointment. They may choose such other officers and agents as they may think best. They may fix the compensation and define the duties of all the officers and agents; and may remove them, or any of them except those chosen by the members, at any time. They may fill vacancies occurring in any manner in their own number or in any of the offices. The Board of Trustees shall have the power to choose an Executive Committee from their own number, and to delegate to such Committee such of their own powers as they may deem expedient. They shall from time to time elect members to the Corporation upon such terms and conditions as they may think best.

X. The Associates of the Marine Biological Laboratory shall be an unincorporated group of persons (including associations and corporations) interested in the Laboratory and shall be organized and operated under the general supervision and authority of the Trustees.

XI. The consent of every Trustee shall be necessary to dissolution of the Marine Biological Laboratory. In case of dissolution, the property shall be disposed of in such manner and upon such terms as shall be determined by the affirmative vote of two-thirds of the Board of Trustees.

XII. The account of the Treasurer shall be audited annually by a certified public accountant.

XIII. These bylaws may be altered at any meeting of the Trustees, provided that the notice of such meeting shall state that an alteration of the bylaws will be acted upon.

RESOLUTIONS ADOPTED AT TRUSTEES' MEETINGS EXECUTIVE COMMITTEE

I. RESOLVED:

(A) The Executive Committee is hereby designated to consist of not more than ten members including the *ex officio* members who shall be the Chairman of the Board of Trustees, President, Director and Treasurer; six additional Trustees, two of whom shall be elected by the Board of Trustees each year, to serve for a three-year term. (August 11, 1967)

(B) The Chairman of the Board of Trustees shall act as Chairman of the Executive Committee, and the President as Vice President. A majority of the members of the Executive Committee shall constitute a quorum and a majority of those present at any properly held meeting shall determine its action. It shall meet at such times and places and upon such notice and appoint such sub-committees as the Committee shall determine. (August 12, 1966)

(C) The Executive Committee shall have and may exercise all the powers of the Board during the intervals between meetings of the Board of Trustees except those powers specifically withheld from time to time by the Board or law. (August 16, 1963)

(D) The Executive Committee shall keep appropriate minutes of its meetings, and its action shall be reported to the Board of Trustees. (August 16, 1963)

II. RESOLVED:

The elected members of the Executive Committee shall be constituted as a standing "Committee for the Nomination of Officers," responsible for making nominations at the Annual Meeting of the Corporation and of the Board of Trustees, for candidates to fill each office as the respective terms of office expire (Chairman of the Board, President, Director, Treasurer, and Clerk).

IV. REPORT OF THE DIRECTOR

TO: THE TRUSTEES OF THE MARINE BIOLOGICAL LABORATORY

Gentlemen:

I submit herewith the report of the 80th Session of the Marine Biological Laboratory. This was an eventful session. In terms of scientific enthusiasm, the period was marked by accomplishment and innovation. In a more mundane but nevertheless important sense, the final stages of our building program were funded; our endowment was substantially increased and stabilized by the sale of our General Biological Supply House holdings, and a drive has been launched among our numerous friends to complete the refurbishing of our central campus and the equipping of our new buildings.

The Marine Biological Laboratory is owned and operated by scientists. In terms of corporate structure, this may not be a unique design but in terms of actual operations, our structure *is* unique. All of the developments reported for this year, as well as past years, have been the result of cooperative and diligent efforts on the part of many individuals acting in the best interests of a service institution which is theirs in fact as well as in corporate structure. No major decision is made by any single individual or small group; all actions are within the context of policies discussed and acted upon by many. This special aspect of our progress, invaluable and unique, is one of our greatest assets and, at the same time, one of our least understood features. The support we have received during the past years is evidence that our role in science is tangibly appreciated and makes it of critical importance that we, as an institution, continue to serve the best interests of scientists, not of institutions, programs or organizations.

1. Facilities Development

Construction is now under way on the new training building and will be ready for occupancy next season. The dining-dormitory complex is fully planned, but the timing depends upon obtaining the proper building permits. These buildings are designed to satisfy, in large part, recommendations of several of our committees, especially the Buildings and Grounds Committee and the Instruction Committee. The housing of courses in the old wooden buildings provided an historical charm but lacked much in terms of stability and safety for the tools of modern research instruction. While students can survive and even thrive in cramped and antiquated living quarters, both safety and health demanded new buildings. Our mess hall has been uniformly deprecated as a physical structure for years. The new

buildings will provide adequate quarters for all these purposes. In addition, they will allow us to take care of the increasing demand for year-round facilities. The new buildings represent a culmination of our plans and hopes. In this connection I would remind you of the outstanding report given by Dr. Armstrong, Chairman of the Building Committee, last year.

The major sources of funds have been The Ford Foundation and the National Science Foundation. Our gratitude to them can best be expressed by the enhanced service of the MBL to science.

2. The Users of the MBL

Last year approximately 270 established Ph.D. scientists used our facilities, laboratories and library for periods of two months or longer. The peak period of August found 276 in residence, the low period of December, 20 in residence. It is impossible to keep account of the numerous individuals who drop in and out, registered or unregistered, to collect specimens, read in the library or do the occasional critical experiment. The variegated and fluctuating uses of our facilities go smoothly—a remarkable tribute to the genius of our operating personnel.

A word should be added about the composition of our scientist clientele. Over the years, there has been a yearly turnover of about 30%. That is, 30% of those in residence in a given year were not in residence the year before. This is good; MBL must not become a private club. Our facilities must be allocated on the basis of intellectual need. Membership in the corporation signifies support of the institution, not a vested interest in its services.

3. The Woods Hole Scientific Community

Excluding tourists, the largest single population component of Woods Hole is composed of individuals present in the area for scientific reasons. The organizations providing the physical base for the scientific population have a common general objective of forwarding scientific inquiry; the devices used towards furthering this are varied. Scientists cooperate when there is intellectual reason to do so. Organizations may co-exist in a sort of mutualism, promoting scientific cooperation while studiously avoiding interference with scientists by administrative rigidity or exclusiveness. Woods Hole does enjoy a community of scientists and the organizations do enjoy a mutualism. The Woods Hole Oceanographic Institution, with its productive staff and programs, provides a worldwide oceanic oversight second to none and the Bureau of Commercial Fisheries Laboratory, as an integral part of the federal establishment, furnishes the expertise in applied aspects of exploitation of the oceans. The Marine Biological Laboratory, as the largest and best known service laboratory for biology in the world, provides an intellectual framework that could never be achieved by a single programmatic institution such as a university. Our faculty, to use the term in a very broad sense, is truly worldwide and its influence cannot be measured in terms of time, or dollars, or numbers, but only in terms of intellectual achievement.

I am happy to state that scientific cooperation and administrative co-existence in Woods Hole are in a productive if not utopian state. Ideas are exchanged, facilities are shared, and people talk to each other. Any formal inter-institutional ar-

rangements within the Woods Hole community are certainly not to be sought; each institution has its unique way of doing things. I am personally proud to have a multiple set of duties, as an individual but not as an administrative siamese twin.

4. *How Large is Too Large*

In 1937 during August the MBL provided services for 256 senior investigators. This required 73,000 square feet of space, not counting course facilities. In 1967, thirty years later, the peak of 276 senior investigators in residence required space plus supporting facilities of 122,000 square feet. The demand for space increases each year, not rapidly but appreciably. This raises the question: How large should MBL be in terms of peak number of senior investigators? The Executive Committee and Trustees have adopted a very cautious attitude towards the size of MBL. Recognizing that a major contribution of MBL is to provide facilities for scientific work, they see that exchange of ideas and information is equally if not more responsible for the global impact of MBL on the advancement of science. The growth of ideas may well be a more important contribution of scientists working at MBL than is the production of new results. So far, I think we all come out with the idea that MBL is about large enough in terms of numbers of people. The physical size of our plant, however, must continue to grow to serve the stable population size. Our library holdings show a doubling time of less than ten years. This means that ten years from now much more of the Lillie Building will be occupied by the Library. Similar reasoning shows increased needs for physical space for the machines and materials of modern science. I do not think we are too large in terms of number of production scientists but I suspect we are large enough. But unless we anticipate a reduction in our intellectual size, we must continue to grow in terms of our physical plant.

5. *Attendance—1967*

Please refer to the tabular report of attendance.

A precise head count is difficult since there is a considerable number of individuals using our facilities throughout the year for library work, short-term collecting, and consultation. While we know who is here at any one time, no consistent record is maintained for short-term visitors.

From the number of inquiries, it is clear that much more extended use of our facilities over a broader yearly time scale will follow the completion of our new dining-dormitory complex. Figures for attendance for those duly registered are given in the tabular reports.

6. *The Fifth Anniversary of the Systematics-Ecology Program*

Founded in 1962, the Systematics-Ecology Program is a going concern. Under the stewardship of Dr. Carriker, boats, trucks, collections, and other items of logistic support are now provided for investigations into ecology, biogeography and systematics of the Cape Cod area. The program has placed MBL in a position to offer to scientists who wish to study organisms in natural habitats the same fine services as we have for many years offered to those working more specifically with marine forms as experimental material.

In addition, and equally important, have been the direct contributions to science made by individuals working with SEP. Some three dozen senior investigators (*i.e.*, doctoral-plus level) have carried out studies. Ten pre-doctoral students have been trained. Over 150 reports have been made, 28 as major papers in established journals, the rest as reports to scientific meetings and other items existing as abstracts. A manual on preservation of marine forms, a key in invertebrate forms, and a bibliography of marine and estuarine environments have been published and are available at cost from the Supply Department. Also, the Gray Museum is assuming a position of real importance, preserved specimens being available for ready identification of experimental material. A most valuable feature is the collection of colored slides.

These accomplishments in so short a time are impressive: the ground work has been laid to allow MBL to be of service to a community of scientists that might not have been so interested in our establishment in the past. Dr. Carriker has made good progress towards the objectives of SEP. The most cogent way of stating these objectives is to quote from his original plan:

"It should be clear from the foregoing that the explicit intent of the Systematics-Ecology Program is to provide opportunity for individual investigators, or investigators and their graduate students, to pursue research in marine systematics and/or ecology in areas of their research interests within the overall framework of the Program, and under as ideal a research environment as can be devised. Living organisms, chemicals, general supplies, etc., would be available through the service departments of MBL at the expense of the Program. It is to be the function of the Director to provide direction to the Program, facilities, support, program ideas where desired, and to attract visiting investigators who should help decide the course of their researches."

7. The MBL and the Oceans

The Marine Biological Laboratory was founded for the explicit purpose of providing a good place for scientists to work with the advantages of the rich biological resources of the sea. The name and location lend emphasis to the virtues of marine forms but this has never been restrictive. During the past decade worldwide interest has been sharpened in the resources of the sea with strong attention to oceanography and fisheries. Recognition of the importance of studying the sea was early appreciated by MBL scientists. Frank Lillie chaired the National Academy of Sciences Committee, whose recommendations led to the founding of Woods Hole Oceanographic Institution and the enhancement of the Scripps Institute of Oceanography. MBL is fortunate in being located in immediate proximity to oceanographic and fisheries institutions and hopefully will long continue to maintain its pervasive role as a general biological laboratory devoted to the support of sound research directed towards the solution of basic biological problems.

A Sea Grant College Act was enacted by Congress a year ago and a program has been established in NSF. A Marine Sciences Act has established a Commission and a Council charged with recommending National Programs in Marine Sciences and Engineering. Reports have been issued by the council and should be forthcoming soon from the commission. The MBL, as a resource for basic investigations of biological problems, including use of marine forms, is clearly in

a position to exercise leadership in these developments, not as an institution perhaps but because scientists working at MBL have an awareness of the value of work on marine forms that will help guide the more applied developments. I think none of us visualizes MBL as an operating unit of the Sea Grant Complex of the future, but our guidance and enthusiasm are surely needed. Our mission is primarily that of assisting basic biological research.

8. The Fund Drive

At our last meeting the Corporation authorized an effort to obtain additional funds to complete the current building program. To my knowledge this is the first time that the institution has thus formally appealed to its constituents. Records do show that, in the early days, collections were made now and then to pay to have the floors swept, but until rising costs caught up with us, MBL has been able to avoid direct organized solicitation. I am sure all will agree that the request is appropriate and I hope that the response will enable us to finish our program we have planned. After all, as a corporation chartered in the Commonwealth of Massachusetts, we could not, and would not, dip into our meagre capital reserves.

9. A Word on the Care and Feeding of Scientists

We are all aware of the success of our Cottage Colony. This is our major bulwark to insure that use of MBL for research is not limited to those established individuals who own property in the area or who can afford to rent living quarters on the open market. The Cottage Colony exists to take care of the housing needs of young investigators with families and of visitors with families from abroad. Our cottages are nice places to live in and therein lies a danger. Continued occupancy by a single family unit over a period of many years would be contrary to the purposes for which the cottages were provided by foundation and private gifts. It is essential that cottages, apartments, and dormitory rooms never become part of a vested interest.

MBL attempts to provide food services. There has been healthy criticism of these services over many decades. Meanwhile there have been no cases of malnutrition nor epidemics. A highly competent committee is now functioning to make recommendations about our prandial future—nutritional, aesthetic and social.

10. Gifts and Bequests

We continue to take pride in the support afforded by our friends in the community through the MBL Associates. The Laboratory was saddened by the death of Mrs. G. H. A. Clowes, who for many years gave much of herself to us. It was most heartening that her daughter-in-law, Peggy—Mrs. G. H. A. Clowes, Jr.—was able to succeed her as president of the Associates. Our small Supply Department fleet will be graced this year with the addition of the 33' "Dasya" the gift of Mr. Charles L. Morse, Jr. of Penzance Point, his former "Chadroe."

11. Personnel

Finally my thanks and appreciation to our resident staff who provide the important services required throughout the laboratory. It is with sorrow I report the death of Mr. Robert Adams in March, so soon after his retirement at the end of the year.

1. MEMORIALS

ALEXANDER FORBES

BY ALFRED C. REDFIELD

Alexander Forbes died on March 27, 1965, at Milton, Massachusetts, where he had lived since his birth eighty-three years before. He was of the tradition of those nineteenth century men of independent means who in following their curiosity about the phenomena of nature added much to the advancement of science.

His major accomplishments were in neurophysiology, a career he had been encouraged to enter by G. H. Parker and Walter Cannon. Visiting England as a young man he brought to this country first-hand knowledge of the work of Sherrington on reflex action and of Lucas and Adrian on the nature of nervous conduction. Much of his subsequent effort was an attempt to understand the former in terms of the latter. For a period his laboratory at the Harvard Medical School became the leading center in America for neurophysiological investigations. He was the first to apply vacuum tubes to the amplification of bioelectric currents and thus became one of the progenitors of the ubiquitous black boxes of the modern laboratory.

Alex Forbes' energies were too great to be exhausted by a distinguished career in science. His keen sense of social responsibility led him to give unstintingly of his time and means to good causes in which he believed. For many years he was President of the Board of Trustees of the George Junior Republic. He encouraged and supported many young people whose ambitions appealed to him, frequently through channels unknown to them. His avocations were yachting, aviation, and outdoor sports, such as mountain climbing, and skiing—pleasures he could share with others. He was never happier than when having a merry time with a group of friends.

Stimulated by Sir William Grenfell, his combined interests in boating, aviation, and his concern with the welfare of those who follow the sea led him to organize and carry out in 1931 an aerial survey of the coast and mountains of northern Labrador. For this accomplishment he was awarded the Daly Medal of the American Geographical Society. Labrador led to his assignment during the Second World War to lay out an air route across this desolate northern country.

Alexander Forbes became a member of the Corporation of the Marine Biological Laboratory in 1925 and a Trustee of the Woods Hole Oceanographic Institution in 1937. He published two papers on work done at the Laboratory. One was on the avicular organ of *Bugula*, the other on the viscosity of the *Nereis* egg as influenced by radiation. These papers and others on flying fish and the flight of gulls are evidence of the diversity of his interests in biology. His understanding of our purposes and his friendships with many of us made him our natural ambassador to Naushon Island which was his base of operation each summer. Laboratory people have always been generously welcome on the Island and many of us recall with gratitude his personal hospitality, both at Naushon and on his yachts.

The Alexander Forbes Lectureship was established in 1959 by the Grass Foundation, of which the majority of Trustees were at some time his laboratory associates. It provides a lasting memorial to an honored and beloved friend of the Marine Biological Laboratory.

2. THE STAFF

EMBRYOLOGY

I. INSTRUCTORS

MALCOLM S. STEINBERG, Professor of Biology, Princeton University, in charge of course
 JAMES N. CATHER, Associate Professor of Zoology, University of Michigan
 RALPH HINEGARDNER, Assistant Professor of Biology, Columbia University
 IRWIN R. KONIGSBERG, Professor of Biology, University of Virginia
 JAMES W. LASH, Associate Professor of Anatomy, University of Pennsylvania
 HANS LAUFER, Associate Professor of Zoology, University of Connecticut

II. JUNIOR INSTRUCTORS

MAX BURGER, Assistant Professor of Biology, Princeton University
 GARY FREEMAN, Assistant Professor of Zoology, University of Illinois

III. LABORATORY ASSISTANTS

BURR G. ATKINSON, JR., University of Connecticut
 WILLIAM F. SINDELAR, Western Reserve University

IV. LECTURES

M. S. STEINBERG	Introduction to the course
I. R. KONIGSBERG	Teleosts I
I. R. KONIGSBERG	Teleosts II
M. S. STEINBERG	Coelenterates I
M. S. STEINBERG	Coelenterates II
DOROTHY SPANGENBERG	Initiation of metamorphosis (strobilization) in <i>Aurelia</i>
FRED DIEHL	Regeneration in Hydroids
G. FREEMAN	Ctenophores I
G. FREEMAN	Ctenophores II
G. FREEMAN	Sponges
C. R. AUSTIN	Fertilization
MAX BURGER	Biosynthesis of structural elements of cell walls and membranes
D. L. D. CASPER	Symmetry and self-assembly
HUGH HUXLEY	Self-assembly of the contractile units of muscle
M. S. STEINBERG	Self-assembly of multicellular complexes
ROBERT ROSEN	Pattern generation in some model embryological systems
TOM HUMPHREYS	A biochemical approach to species-specific aggregation in marine sponges
LIONEL JAFFE	The early development of <i>Fucus</i> : Prototype of localization
EVERETT ANDERSON	The fine structure of eggs
RAYMOND RAPPAPORT	Cytokinesis
PAUL GREEN	Biochemistry and physics of cell morphogenesis
ROUND TABLE DISCUSSION	What is differentiation?
J. CATHER	Spiralians I
J. CATHER	Spiralians II
J. CATHER	Spiralians III
JACK COLLIER	Biochemistry of the <i>Ilyanassa</i> embryo
JOHN M. ARNOLD	Development of cephalopods

JAMES LASH	Ascidians I
JAMES LASH	Ascidians II
JAMES LASH	Ascidians III
ROGER MILKMAN	The genetics and development of <i>Botryllus</i>
H. LAUFER	Crustaceans I
H. LAUFER	Crustaceans II
PHILIP SIEKEVITZ	The assembly of cellular membranes
H. LAUFER	Crustaceans III
R. HINEGARDNER	Echinoderms I
R. HINEGARDNER	Echinoderms II
JOHN HUGHES	The sex life of the lobster
PAUL WEISZ	The significance of larvae
ROUND TABLE DISCUSSION	Unsolved problems in development
EDWARD KELLENBERGER	Shape-inheritance of a protein polymer (Morphogenesis of phage T4 capsid)
MICHAEL FISCHBERG	Germinal cytoplasm in gall midges and anurans
HELGE STALSBERG	Morphogenetic movements and differential mitotic activity during early heart formation in the chick embryo
JAMES LASH	Chondrogenesis: Genotypic and phenotypic expression
HOWARD GREEN	Cell contact and oncogenic viruses
ALLISON BURNETT	Dedifferentiation, metaplasia, and ions

PHYSIOLOGY

I. CONSULTANTS

- MERKEL H. JACOBS, Professor of Physiology, University of Pennsylvania
 ALBERT SZENT-GYÖRGYI, Director, Institute for Muscle Research, Marine Biological
 Laboratory
 W. D. McELROY, Director, McCollum-Pratt Institute, The Johns Hopkins University
 J. WOODLAND HASTINGS, Professor of Biology, Harvard University

II. INSTRUCTORS

- ANDREW G. SZENT-GYÖRGYI, Professor of Biology, Brandeis University, in charge of
 course
 SYDNEY BRENNER, Medical Research Council, Laboratory of Molecular Biology, Cam-
 bridge, England
 RODERICK K. CLAYTON, Professor of Biophysics, Cornell University
 HARLYN HALVORSON, Professor of Molecular Biology and Bacteriology, University of
 Wisconsin
 HUGH E. HUXLEY, Medical Research Council, Laboratory of Molecular Biology, Cam-
 bridge, England
 MAURICE SUSSMAN, Professor of Biology, Brandeis University
 K. E. VAN HOLDE, Professor of Physical Chemistry, University of Illinois

III. STAFF ASSOCIATES

- HENRY DEPHILLIPS, Trinity College
 GEORGE HOCH, University of Rochester
 EUGENE KATZ, Laboratory of Molecular Biology, Cambridge, England
 DAVID PATTERSON, Brandeis University
 MICHAEL SMOLLEN, University of Rochester

WILLIAM STEINBERG, University of Wisconsin
 RAYMOND E. STEPHENS, Harvard University
 JAMES C. VARY, University of Wisconsin
 ANNE MARIE WEBER, St. Louis University
 BERNARD WEISSBLUM, University of Wisconsin

IV. SPECIAL LECTURERS

SHINYA INOUÉ, Professor of Biology, University of Pennsylvania
 JOHN NICHOLLS, Associate Professor of Physiology, Yale Medical School
 SOL SPIEGELMAN, Professor of Microbiology, University of Illinois

V. ASSISTANTS

ROBERT LEVY, New York University
 PATRICIA S. VARY, University of Wisconsin

VI. LECTURES

ANDREW G. SZENT-GYÖRGYI	Muscle contraction—I Muscle contraction—II
ANNE MARIE WEBER	Control of contraction
K. E. VAN HOLDE	Protein structure—I. Determinants of secondary and tertiary structure Protein structure—II. Quaternary structure Protein structure—III. Control
H. O. HALVORSON	Developmental changes during sporulation: Physiological and genetic control of the formation of the dormant state
WILLIAM STEINBERG	Developmental changes during outgrowth: Protein synthesis and ordered enzyme synthesis
H. O. HALVORSON	Developmental changes during the cell cycle: Effect of gene position on enzyme timing in yeast
MAURICE SUSSMAN	I—Some conceptual and operational aspects of developmental biochemistry II—The cellular slime molds—LCVII III—The cellular slime molds—LCVIII
RODERICK K. CLAYTON	The mechanism of photosynthesis: I—Early formulations and the question of quantum efficiency II—Contemporary formulations and the cooperation of two photochemical processes
GEORGE HOCH	III—Oxido-reduction and phosphate esterification
RODERICK K. CLAYTON	IV—Physical problems and mechanisms
HUGH E. HUXLEY	Muscle structure—I
BERNARD WEISSBLUM	The role of tRNA in the synthesis of various non-proteins
HUGH E. HUXLEY	Muscle structure—II
SYDNEY BRENNER	The genetic aspects of the code—I The genetic aspects of the code—II The genetic aspects of the code—III
SHINYA INOUÉ	Organization of living cell fine structure: I—The mitotic spindle
RAYMOND E. STEPHENS	II—Structural proteins associated with mitosis

R. WOLFE	Methane bacteria and transmethylation
SHINYA INOUÉ	Organization of living cell fine structure: III—Sperm chromosomes
JOHN G. NICHOLLS	Electrical signalling in the nervous system
E. KRAVITZ	A study of synaptic chemistry in single neurons
JOHN G. NICHOLLS	Neural organization in the visual system
F. BECKWITH	Long-term changes in synaptic transmission
B. D. DAVIS	Regulation of gene action
SEYMOUR BEYCHOK	Regulation of RNA and DNA synthesis
CYRUS LEVINTHAL	Optical activity and conformation of ribonuclease
L. LORAND	Pathways of protein foldings
ALBERT SZENT-GYÖRGYI	Enzyme-controlled assembly of protein fibers: Biosynthesis of a blood clot
RUTH HUBBARD	Comments
RICHARD CONE	Chemistry of visual excitation
WILLIAM HAGINS	Early receptor potentials
J. E. DOWLING	Excitatory processes in squid eye
E. F. MACNICHOL, JR.	Visual adaptation and visual pathway anatomy
GEORGE WALD	Retinal mechanisms of color discrimination and spatial contrasts
SOL SPIEGELMAN	Mechanisms of human color vision
ROBERT MARTIN	<i>In vitro</i> RNA synthesis
DAVID YPHANTIS	Electrical and chemical synaptic transmission in ciliary ganglion
A. H. STURTEVANT	Equilibrium centrifugation of non-ideal protein solutions
DOUGLAS R. WILKIE	T. H. Morgan
E. A. ADELBERG	A muscle physiologist's lament: Is thermodynamics useless?
JEAN-PIERRE CHANGEUX	DNA transfer in bacterial conjugation
I. C. GUNSALUS	On the mechanism of allosteric interaction
	Gene transfer in the fluorescent pseudomonads and the biological management of structural diversity

MARINE BOTANY

I. INSTRUCTORS

- WALTER R. HERNDON, Professor of Botany and Assistant Vice-President, Academic Affairs, University of Tennessee, in charge of course
- PHILIP W. COOK, Assistant Professor of Botany, University of Vermont
- MELVIN S. FULLER, Associate Professor of Botany, University of California, Berkeley
- H. WAYNE NICHOLS, Associate Professor of Botany, Washington University
- JANET R. STEIN, Associate Professor of Botany, University of British Columbia, Vancouver

II. SPECIAL LECTURERS

- R. W. WILCE, University of Massachusetts
- L. PROVASOLI, Haskins Laboratories
- R. DRUM, University of Massachusetts
- I. M. LAMB, Harvard University
- F. TRAINOR, University of Connecticut
- F. E. ROUND, University of Bristol, England

III. ASSISTANTS

DAVIS L. FINDLEY, University of Tennessee

DIANE I. FINDLEY, University of Tennessee

THOMAS LEE, University of Connecticut

IV. LECTURES

W. R. HERNDON	Marine environment and local flora
J. R. STEIN	Introduction to algae
M. S. FULLER	Cyanophyta—I
J. R. STEIN	Cyanophyta—II
M. S. FULLER	Myxomycota and Eumycota (Introduction)
P. W. COOK	Isolation and cultivation of algae and fungi
M. S. FULLER	
J. R. STEIN	
P. W. COOK	Chlorophyta (Introduction)
	Work on isolation technique
W. R. HERNDON	Chlorophyta—Chlorococcales
P. W. COOK	Chlorophyceae—Volvocales
W. R. HERNDON	Chlorophyceae—Tetrasporales
M. S. FULLER	Free-living phycomycetes
J. R. STEIN	Prasinophyceae
	Chlorophyceae—Ulotrichales I
	Chlorophyceae—Ulotrichales II
P. W. COOK	Chlorophyceae—Cladophorales, Siphonocladales, Codiales
P. W. COOK	Chlorophyceae—Codiales, Dasycladales, Derbesiales
J. R. STEIN	Chlorophyceae—Oedogoniales, Zygnematales
W. R. HERNDON	Euglenophyceae
LIONEL JAFFE	The early development of <i>Fucus</i>
I. FREIDMANN	Cytology of fertilization in algae
J. R. STEIN	Xanthophyceae
J. R. STEIN	Chrysophyceae, Haptophyceae
M. S. FULLER	Fungal parasites of algae
P. W. COOK	Phaeophyceae—Introduction
J. R. STEIN	Phaeophyceae—Ectocarpales, Sphacelariales
P. W. COOK	Phaeophyceae—Chordariales, Desmarestiales, Sporochnales
W. R. HERNDON	Phaeophyceae—Punctariales, Dictyosiphonales, Dictyotales, Cutleriales
R. W. WILCE	Arctic algal ecology
M. S. FULLER	Physiology of marine fungi
J. R. STEIN	Phaeophyceae—Laminariales
P. W. COOK	Phaeophyceae—Fucales
H. WAYNE NICHOLS	Rhodophyceae—Introduction
L. PROVASOLI	Nutrition and culture of marine algae
H. WAYNE NICHOLS	Rhodophyceae—Bangiophycidae
R. DRUM	Ultrastructure of diatoms
H. WAYNE NICHOLS	Rhodophyceae—Transitional forms, Nemaionales, Gelidiales
I. M. LAMB	Zonation of marine algae of Northeast Atlantic Coast
H. WAYNE NICHOLS	Rhodophyceae—Cryptonemiales, cytology of reproduction
M. D. FULLER	Fungi

H. WAYNE NICHOLS	Rhodophyceae—Gigartinales, Rhodymeniales
H. WAYNE NICHOLS	Rhodophyceae—Ceramiales
F. TRAINOR	Morphogenesis in coenobial algae
H. WAYNE NICHOLS	Rhodophyceae—Summary evolution, experimental aspects of development
W. R. HERNDON	Charophyta
M. S. FULLER	Fungi—marine ascomycetes
M. S. FULLER	Pyrrophyta
F. E. ROUND	Bacillariophyceae I
	Bacillariophyceae II

INVERTEBRATE ZOOLOGY

I. CONSULTANTS

FRANK A. BROWN, JR., Morrison Professor of Biology, Northwestern University
 LIBBIE H. HYMAN, American Museum of Natural History
 CLARK P. READ, Professor of Biology, Rice University
 ALFRED C. REDFIELD, Woods Hole Oceanographic Institution

II. INSTRUCTORS

W. D. RUSSELL-HUNTER, Professor of Zoology, Syracuse University, in charge of course
 GEORGE G. HOLZ, JR., Professor of Microbiology, State University of New York, Upstate
 Medical Center, Syracuse
 NORMAN MILLOTT, Professor of Zoology, Bedford College, University of London, Eng-
 land
 ERIC L. MILLS, Associate Professor of Biology, Institute of Oceanography, Dalhousie
 University, Halifax, Nova Scotia, Canada
 JAMES CASE, Professor of Biology, University of California, Santa Barbara
 FRANK M. FISHER, JR., Assistant Professor of Biology, Rice University
 ROBERT K. JOSEPHSON, Associate Professor of Biology, Western Reserve University
 JONATHAN P. GREEN, Assistant Professor, Division of Biological and Medical Sciences,
 Brown University
 MEREDITH L. JONES, Curator in charge, Division of Worms, United States National Mu-
 seum, Smithsonian Institute

III. JUNIOR INSTRUCTOR

HUGH Y. ELDER, Lecturer in Histology, Institute of Physiology, University of Glasgow,
 Scotland

IV. ASSISTANTS

ALBERT J. BURKY, Syracuse University
 DARRELL R. STOKES, University of Hawaii

V. LECTURES

W. D. RUSSELL-HUNTER	Orientation and functional homologies in invertebrates
ALBERT J. BURKY	Laboratory orientation
DARRELL R. STOKES	

- ROBERT K. JOSEPHSON Cnidaria I—Introduction to the Cnidaria and Ctenophora
Cnidaria II—Nervous system and behavior
Seminar—Physiological mechanisms controlling behavior
in the hydroid, *Tubularia*
- FRANK M. FISHER, JR. Cnidaria III—Function of the nematocysts, feeding
Turbellaria and Trematoda
Cestoda and Rhynchocoela
- W. D. RUSSELL-HUNTER Mollusca I—General molluscan organization: Functioning
of mantle cavity in Gastropoda
Littoral Ecology: Theoretical and physiological
- ERIC L. MILLS Littoral Ecology: Practical and pragmatic
- W. D. RUSSELL-HUNTER Mollusca II—Gastropoda (continued): Mantle cavity and
feeding mechanisms in Bivalvia
Mollusca III—Adaptations in bivalves: Aspects of general
physiology of gastropods and bivalves
Mollusca IV—Functional morphology in Cephalopoda and
minor groups
Short seminar—Some problems of mechanics in molluscs
Mollusca V—Functional morphology in Amphineura, *Neopila*,
archetypes and ancestors
- ERIC L. MILLS The framework of Cape Cod—an introduction to the history
of the Cape and Barnstable Harbor
- MEREDITH L. JONES Annelida introduction: General characteristics, classification,
external morphology
Annelida II—Feeding, respiration and osmoregulation
Polychaetous serendipity: *Magelona* and *Caobangia*
Annelida III—Reproduction, nervous system, locomotion
- W. D. RUSSELL-HUNTER Seminar—Physiological variation and evolution in fresh-
water molluscs
- ERIC L. MILLS Arthropoda I—General features of arthropods: Introduction
to crustacean structure
Arthropoda II—Crustacean structure, physiology, and re-
production
- JAMES CASE Arthropoda III—Neurosecretion and endocrine control
Arthropoda IV—Vegetative physiology
- W. D. RUSSELL-HUNTER Symposium—Invertebrate connective tissues, skeletons and
(Chairman) mechanics
HUGH Y. ELDER The function of connective tissues in invertebrates and the
occurrence of elastic fibers
- ROBERT V. RICE Amino acid composition of bivalve ligaments
- ELAINE A. ROBSON Functions of mesoglea in coelenterates
- PHILIP PERSON Endoskeletal cartilages in invertebrates
- ROBERT K. JOSEPHSON Tentacular rowing by sea anemones
- JAMES CASE Arthropoda V—Neurobiology
- FRANK M. FISHER, JR. Aschelminthes
Entoprocta, Ectoprocta
The biology of asteroids and ophiuroids
- NORMAN MILLOTT Function of hemal system and axial organ in echinoids
- A. FARMANFARMAIAN Photosensitivity in echinoids
- NORMAN MILLOTT The biology of echinoids
The biology of holothurians; development and phylogeny of
echinoderms

ERIC L. MILLS	Short seminar—The biology of an amphipod crustacean sibling species pair
JONATHAN P. GREEN	Protochordata I Protochordata II Porifera
GEORGE G. HOLZ, JR.	The nature of the Protozoa
HUGH Y. ELDER	Body cavities, connective tissues, and invertebrate locomotion
GEORGE G. HOLZ, JR.	Flagellates I Flagellates II—Sarcodines I
JEAN-MARIE BASSOT	Structure and evolution of luminous organs in some marine animals
GEORGE G. HOLZ, JR.	Sarcodines II Ciliates
W. D. RUSSELL-HUNTER	One approach to the zooplankton

MARINE ECOLOGY

I. CONSULTANTS

MELBOURNE R. CARRIKER, Marine Biological Laboratory
 BOSTWICK H. KETCHUM, Woods Hole Oceanographic Institution
 EDWIN T. MOUL, Rutgers University
 JOHN H. RYTHER, Woods Hole Oceanographic Institution

II. INSTRUCTORS

W. ROWLAND TAYLOR, Associate Professor of Oceanography, The Johns Hopkins University, in charge of course
 DENNIS J. CRISP, Director, Marine Science Laboratories, University College of North Wales, U. K.
 HOWARD L. SANDERS, Senior Scientist, Woods Hole Oceanographic Institution
 LAWRENCE B. SLOBODKIN, Professor of Zoology, University of Michigan
 GOTRAM UHLIG, Biologische Anstalt Helgoland, Germany

III. SPECIAL LECTURERS

LUIGI PROVASOLI, Haskins Laboratories, New York
 VICTOR ZULLO, Systematics-Ecology Program, Marine Biological Laboratory

IV. ASSISTANTS

JOHN F. BOYER, University of Chicago
 HERMAN F. BOSCH, The Johns Hopkins University

V. LECTURES

W. ROWLAND TAYLOR
 Introduction to marine ecology
 The marine environment I—Chemistry of sea water
 The marine environment II—Solar radiation through sea water
 Phytoplankton I
 Phytoplankton II

- Primary productivity by phytoplankton
Phytoplankton III
Ecological techniques: Salinity, dissolved oxygen, chlorophyll
Ecological techniques: Plankton sampling gear
- V. ZULLO
Cape Cod as an ecological laboratory
The evolutionary significance of the deep-sea barnacle fauna
- DR. LEADBETTER
Introduction to microbial ecology
Microorganisms and their role in nature
Selective cultures
- H. JANNASCH
Microbial transformations I
Microbial transformations II
Continuous culture in microbial ecology
Bacteriological lab and field techniques
Recent approaches in microbial ecology
- DR. TRUPER
Ecology of photosynthetic bacteria
- D. RHODES
Environmental stress gradients and paleobathymetry
- H. BOSCH
Benthic collecting methods
- R. GREEN
Population dynamics of *Gemma gemma*
- M. JONES
On the spatial distribution of selected benthic invertebrates of San Francisco Bay
- R. HESSLER
Studies of the deep-sea benthos
- J. SIMON
Reproductive ecology of polychaetes
- R. SCHELTEMA
The importance of the larvae of benthic marine invertebrates
- E. UCHUPI
Bathymetry and sediments of the shelf and slope of the Atlantic Coast and Gulf of America
- G. UHLIG
Physiological ecology—a new field of research
Effects of temperature and salinity on marine and brackish water animals
Introduction to benthic microfauna
Methods of research on benthic protozoa
The biology of folliculinids (Ciliata; Heterotrichida), illustrated with film
Life-cycle and ecological observations on *Noctiluca miliaris*, illustrated with film
- R. WILCE
Arctic ecology
- L. SLOBODKIN
Ecology of evolution
Classical population dynamics
Intrinsic rates of increase and reproductive value
Energetics of populations and the theory of predation
Control of abundance in nature
- D. CRISP
The future of ecology in practical affairs
The role of larvae in the ecology of marine invertebrates
Behavior of marine invertebrate larvae during the free swimming period
Settlement behaviour as exemplified by the cyprid and other larval forms
Behavior of the larvae of deposit living invertebrates
Gregarious behavior at settlement
Territorial behavior at settlement

MARINE BIOLOGICAL LABORATORY
 SYSTEMATICS-ECOLOGY PROGRAM

THE STAFF

Director: MELBOURNE R. CARRIKER
 Resident Systematist: VICTOR A. ZULLO
 Resident Ecologist: ROGER H. GREEN
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SEMINARS (WINTER INCLUDED)

P. WEYGOLDT	Comparative reproductive biology in pseudoscorpions
JAMES R. SEARS	Developmental growth patterns and the occurrence of "mitotic waves" in the coenocytes of the marine endophytic green alga, <i>Blastophysa rhizopus</i> Reinke
DAVID K. YOUNG	The food and feeding of Indo-West-Pacific dorid nudibranches
ROGER H. GREEN	Constant <i>versus</i> erratic mortality as a population regulator in the intertidal
KENNETH J. BOSS	Studies in the systematics of the Tellinidae
TORE LEVRING	Submarine light and vertical distribution of marine benthic vegetation
GARRY CHARLTON	Determination of O ₂ in biological tissues
H. PERRY JEFFRIES	Environmental stress on marine communities as revealed by fatty acid and amino acid spectra
ARTHUR C. MATHIESON	Ecological studies of the marine brown alga <i>Phaeostrophion irregulare</i> S. et G. on the Pacific coast of North America
HAROLD H. PLOUGH	The distribution of ascidian species in the Woods Hole region

- SAUL B. SAILA Some aspects of the movement and behavior of the lobster, *Homarus americanus*
- ARTHUR HUMES Concepts and recent trends in work on parasitic copepods
DAVID DEAN Maine's new marine laboratory—The Ira C. Darling Center for Research, Teaching and Service
- KATHARINE HOBSON The feeding and ecology of two sympatric species of lugworm (Arenicolidae Polychaeta)
- ALFRED SENFT Schistosome physiology II—a continuation
JOSEPH L. SIMON The biology of *Histriobdella homari*, a commensal polychaete
- MEREDITH L. JONES Polychaete potpourri
WILLIAM C. SUMMERS Local squid: past, present and future
LOUISE HOBSON Speculation on the transport of phytoplankton carbon into the deep sea
- THOMAS J. M. SCHOPF Is an ectoproct possible? (Oxygen consumption in a small marine invertebrate)
- LAWRENCE PINTER Behavior of *Phidippus* (Salticidae, Arachnida)
DANIEL MERRIMAN The history of oceanography
HOLGER W. JANNASCH Bacterial growth kinetics in microbial ecology
JOHN D. DAVIS Selective attack by the boring polychaete worm, *Polydora websteri*: environment *vs.* distribution
- ROBERT A. CROKER The biology of some closely-related sympatric amphipods
ROBERT T. WILCE Arctic algal ecology
A. LEE MCALESTER Paleobiology of bivalve molluscs
BARRY A. WADE Ecology of New England nudibranchs
HAROLD H. PLOUGH Recollections of Woods Hole, 1913–1923: Problems and personalities
- VICTOR A. ZULLO Barnacles in Europe
RUTH D. TURNER Deep-sea boring bivalves
WILLIAM J. CLENCH Tree snails of the genus *Linguis* in Florida, Hispaniola and Cuba
- GILBERT V. LEVIN The ecology of Mars
MARTYN APLEY On *Melampus bidentatus*, the salt-marsh snail
MICHAEL T. GHISELIN The principles and concepts of systematic biology
DAVID K. YOUNG Interpretations from bulk chemistry of Chesapeake Bay sediments
- GUNTHER STOTZKY Influence of clay minerals and the ecology and population dynamics of microorganisms
- MELBOURNE R. CARRIKER The mechanism of shell penetration by muricid gastropods: chemical or mechanical?
- WILLIAM D. BURBANCK The ecology of *Cyathura*: The development of a problem
PAUL S. GALTISOFF Water quality levels essential for the well-being of aquatic biota (summary of the work of Technical Advisory Committee of FWPCA)
- K. O. EMERY Relict sediments of the continental shelf
RALPH O. BRINKHURST Biology of aquatic oligochaetes
WILLIAM T. LAMMERS Biophysical limnology: The separation of colloidal and suspended particles in natural waters
- FRANK E. ROUND Vertical migration rhythms in benthic algae
DONALD J. ZINN History and biology of Penikese Island: 1872–present

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 STEPHENS, RAYMOND E., Research Fellow in Biology, Harvard University
 STILLMAN, IRVING M. Research Associate, National Institutes of Health
 STRITTMATTER, PHILIPP, Associate Professor of Biochemistry, Washington University
 STUNKARD, HORACE W., Research Associate, American Museum of Natural History
 STURTEVANT, A. H., Thomas Hunt Morgan Professor of Biology, California Institute of Technology

- SURGENOR, DOUGLAS M., Dean, State University of New York School of Medicine, at Buffalo
- SWETT, JOHN E., Associate Professor of Anatomy, State University of New York, Upstate Medical Center, Syracuse
- SZABÓ, GEORGE, Assistant Professor of Anatomy, Harvard Medical School
- SZENT-GYÖRGYI, ALBERT, Director and Chief Investigator, The Institute for Muscle Research, Marine Biological Laboratory
- SZENT-GYÖRGYI, ANDREW G., Professor of Physiology, Brandeis University
- SZENT-GYÖRGYI, EVA M., Research Associate, Brandeis University
- TANZER, MARVIN LAWRENCE, Research Associate, Massachusetts General Hospital and Harvard Medical School
- TASAKI, ICHIKI, Chief, Laboratory of Neurobiology, National Institutes of Health
- TAYLOR, KENNETH M., Professor of Biology, The Institute for Cancer Research, Philadelphia, and San Diego State College
- TAYLOR, ROBERT E., Acting Chief, Biophysics Laboratory, National Institutes of Health
- TAYLOR, W. ROWLAND, Assistant Professor of Oceanography, The Johns Hopkins University
- TAYLOR, WM. RANDOLPH, Professor Emeritus of Botany, University of Michigan
- THALER, M. MICHAEL, Research Fellow, Harvard Medical School
- TIBBS, JACK, University Lecturer, University of St. Andrews, Scotland
- TILNEY, LEWIS G., Harvard University
- TOYODA, JUN-ICHI, Research Associate, The Rockefeller University
- TRINKAUS, J. P., Professor of Biology and Master of Branford College, Yale University
- TROLL, WALTER, Associate Professor, New York University Medical Center
- TWEDELL, KENYON S., Associate Professor of Biology, University of Notre Dame
- UHLIG, GOTRAM, Biologische Anstalt Helgoland, West Germany
- VAN HOLDE, K. E., Professor of Chemistry, University of Illinois
- VARGAS, FERNANDO F., Faculty of Sciences, University of Chile
- VILLEE, CLAUDE A., Andelot Professor of Biochemistry, Harvard University
- VILLEGAS, JORGE, Investigador Asociado, Instituto Venezolano de Investigaciones Cientificas, Caracas, Venezuela
- WALD, GEORGE, Professor of Biology, Harvard University
- WARREN, LEONARD, Professor of Therapeutic Research, University of Pennsylvania School of Medicine
- WATANABE, AKIRA, Consultant, National Institutes of Health
- WATKINS, DUDLEY T., Assistant Professor of Anatomy, University of Connecticut
- WEBB, H. MARGUERITE, Professor of Biological Sciences, Goucher College
- WEBER, ANNEMARIE, Research Associate, St. Louis University
- WEISBLUM, BERNARD, Assistant Professor of Pharmacology, University of Wisconsin
- WIERCINSKI, FLOYD J., Professor of Biology, Northeastern Illinois State College
- WILKIE, DOUGLAS ROBERT, Professor of Experimental Physiology, University College, London, England
- WEISS, LEON, Professor of Anatomy, The Johns Hopkins University
- WYTTENBACH, CHARLES R., Assistant Professor of Zoology, University of Kansas
- ZIGMAN, SEYMOUR, Assistant Professor of Biochemistry, University of Rochester
- ZIMMERMAN, ARTHUR M., Professor of Zoology, University of Toronto, Canada
- ZUSY, REV. DENNIS R., O.P., Research Associate, Northwestern University

Lalor Fellows, 1967

- MICHAEL FISCHBERG, Senior Fellow, Professor of Zoology, University of Geneva, Geneva, Switzerland
- ALLEN W. SCHUETZ, The Johns Hopkins University
- WALTER AUCLAIR, Rensselaer Polytechnic Institute
- JACK TIBBS, University of St. Andrews, Scotland

Lillie Fellow, 1967

- HANS PETER PERLMANN, Wenner-Gren Institute, Stockholm, Sweden

Grass Fellows, 1967

WERNER E. REICHARDT, Senior Fellow, Director, Max-Planck-Institut für Biologie, Germany
 RENE EPSTEIN, Columbia University, College of Physicians & Surgeons
 DONALD T. FRAZIER, University of New Mexico Medical School
 RAYMON M. GLANTZ, New York University Medical School
 EDWARD GRUBERG, University of Illinois
 RONALD R. HOY, Stanford University
 GEORGE J. MPITSOS, University of Virginia
 BERT SHAPIRO, Harvard University
 DENIS A. BAYLOR, Yale University School of Medicine
 STEPHEN R. SHAW, University of St. Andrews, Scotland

Rand Fellow, 1967

G. ADRIAN HORRIDGE, University of St. Andrews, Scotland

Research Assistants, 1967

ABRAHAM, JERROLD L., University of California, San Diego
 ANGEL, RUTH ANN, North Carolina State University
 ANTONELLIS, BLENDA, Western Reserve University
 APLEY, MARTYN L., Syracuse University
 ATKINSON, BURR G., JR., University of Connecticut
 ATKINSON, JAMES W., Emory University
 ATWATER, ILLANI, University of Chile
 BALICKI, PHILIP, Trinity College
 BARNES, STEPHEN, Syracuse University
 BARNHILL, ROBERT, Miami University, Ohio
 BEAUREGARD, LAURENT, JR., Providence College
 BEDWINEK, JOHN, Oberlin College
 BERMAN, MAC, Columbia University
 BETLEM, SUSAN J., University of Rochester
 BICKER, ALVIN A., New York University
 BILLINGS, SUSAN, Harvard Medical School
 BILLETER, LISBETH, New York University School of Medicine
 BOSCH, HERMAN F., The Johns Hopkins University
 BOSSERMAN, BARBARA, Brown University
 BOYER, JOHN F., University of Chicago
 BOYLE, ROBERT, Georgetown University Medical School
 BRADY, ARLENE C., Mellon Institute
 BRANDER, KEITH M., Marine Science Laboratories, UCNW, Bangor
 BRANDHORST, BRUCE P., University of California, San Diego
 BRIDGES, SANDRA H., Washington University
 BRUNER, LYNNEL, The Johns Hopkins University
 BRUZUAL, IRIS B., Instituto Venezolano de Investigaciones Cientificas, Venezuela
 BURKY, ALBERT, Syracuse University
 BURNS, LAWRENCE A., New York University
 CAMPBELL, MARTIN C., University of Chicago
 CHASIS, SARAH, New York University
 CHILDRESS, DIANA, University of Oregon
 COLLINS, SANDRA E., University of Kansas
 COLVILLE, JEAN E., Tulane University
 CONE, MARGARET VIRGINIA, University of Massachusetts
 CONNELL, PENNY MAY, Tulane University
 CONTA, BARBARA A., University of Michigan
 COPSEY, KATHRYN J., University of Miami
 CORDELL, SAUNDRA, Columbia University

DIZON, ANDREW, Yale University
DONAHUE, HELAINE, Washington University School of Medicine
DOYLE, DARRELL J., University of Connecticut
EDDS, KENNETH T., University of Rhode Island
EIGNER, ELIZABETH ANN, University of New Mexico Medical School
EPSTEIN, MILES, Syracuse University
ETIENNE, EARL M., State University of New York, at Albany
FIEL, STANLEY, University of Connecticut
FERNANDEZ, HUGO, Massachusetts Institute of Technology
FINDLEY, DIANE, University of Tennessee
FINDLEY, DAVIS, University of Tennessee
FITZJARRELL, AUSTIN T., Tulane University
FORMAN, DAVID S., The Rockefeller University
FRICK, JANE ALLISON, Columbia University
GELFAN, JANET I., Albert Einstein College of Medicine
GOLDMAN, JAMES E., University of Maryland
GOODENOUGH, DANIEL, Harvard Medical School
GOODYER, PAUL R., Yale University
GREITZER, NAOMI, Barnard College
HALL, TONI JEAN, University of Michigan
HAMILTON, JOHN DAVID, Massachusetts Institute of Technology
HANSON, MUSETTA, College of St. Mary of the Springs
HARRIS, ALBERT, Yale University
HARRIS, EDWARD M., Duke University
HARRIS, ELIZABETH H., Yale University
HART, BARBARA ANN, Single Cell Research Foundation, Inc.
HECHT, RALPH, New York University
HENDRICKSON, WAYNE, The Johns Hopkins University
HEYMANN, PETER W., Washington University
HERMOLIN, JOSEPH, University of Toronto, Canada
HILLMAN, GILBERT, Yale University
HOFMAN, FLORENCE, City College of New York
HOLT, THOMAS, University of Connecticut
HOUSEHOLDER, SALLY, The Rockefeller University
HOWELL, KATHRYN E., Rutgers, The State University
HUEBNER, ERWIN, University of Massachusetts
HUMPHREYS, SUSIE, University of California, San Diego
IRWIN, M. LINDA, Western Reserve University
IVKER, FRANCES S., Indiana University
KARTEN, BARBARA, Columbia University
KATZ, EUGENE R., University of Cambridge, England
KATZ, JAY A., Washington University School of Medicine
KIMBALL, FRANCES, Reed College
KISSIN, EVELYN ANNE, Columbia University
KRATOWICH, NANCY R., Columbia University
KRAWCHENKO, JOHN, State University of New York, Upstate Medical Center, Syracuse
LANG, FRED, University of Illinois
LANGE, MARY T., University of Pennsylvania
LANKTON, JAMES, Harvard University
LAVORGRA, LORENZO, Washington University Medical School
LAWRENCE, LYLE, Marine Biological Laboratory
LEE, THOMAS F., University of Connecticut
LEFFERT, MARK N., New York University School of Medicine
LEHMAN, WILLIAM, Princeton University
LEMON, STANLEY M., Princeton University
LETTIS, PAMELA JEAN, University of Toronto, Canada
LEVY, ROBERT M., Washington University Medical School
LIPSON, ROBERT ALAN, VA Hospital, Brooklyn

LIPSON, STEPHEN JAY, Yale University
LOVERDE, VERNON D., Washington University
LUCKY, PAUL A., Kenyon College
MAHANTI, MAHENDRA, Columbia University
MARSH, RICHARD CHARLES, University of Maryland
MARTINEK, JOHN J., Tulane University
MASON, JOHN MONTGOMERY, JR., Marine Biological Laboratory
McGRATH, RICHARD A., Northwestern University
McLAUGHLIN, JANE A., The Institute for Muscle Research
MELTZER, PAUL S., Dartmouth College
MERRILL, CHARLOTTE, Massachusetts Institute of Technology
MOFFETT, DAVID F., JR., Duke University
MOYER, PATRICIA ELLEN, Radcliffe College
MPITSOS, GEORGE J., University of Virginia
MYHRMAN, ROLF V., Northwestern University
NADOL, JOSEPH B., JR., The Johns Hopkins University School of Medicine
NATALINI, JOHN J., Northwestern University
NICHOLS, T. RICHARD, Brown University
O'BENAR, JOHN O., University of Illinois
OBERPRILLES, JOHN, Tulane University
OSELKA, JANET, Columbia University
PATTERSON, DAVID, Brandeis University
PALMER, JERRY P., State University of New York, Upstate Medical Center, Syracuse
PARMENTIER, JAMES LAWRENCE, University of California, Santa Barbara
PEDERSON, THORU, Syracuse University
PERCY, JONATHAN, Columbia University
PERRY, MAUREEN M., Trinity College, Washington, D. C.
PLATT, TERRY, Harvard University
PRIOR, GWEN, Brandeis University
PRUSCH, ROBERT D., Syracuse University
PYSAR, JOANNE, University of Connecticut
QUATTROPANI, STEVEN, University of Massachusetts
RANSOHOFF, DAVID F., Harvard College
RAVITZ, MELVYN J., Columbia University
RAYNER, ELLEN P., Brandeis University
REA, INA KATHERINE, Washington University
REYNOLDS, GEORGE T., Princeton University
RITCH, ROBERT, Harvard University
ROBBINS, MARCIA, Radcliffe College
ROBERTSON, LOLA E., American Museum of Natural History
ROBINOWITZ, BERNARD, Tulane University Medical School
ROSENBLUM, JUDY LYNN, University of Vermont
ROSSETTO, MICHALANGELO A., The Rockefeller University
RYDER, GARY, State University of New York, Upstate Medical Center, Syracuse
SAGE, JEAN, Indiana University Medical School
SALTZMAN, ORAH, Massachusetts Institute of Technology
SAUL, DAVID, The Johns Hopkins Medical School
SCHOEPP, CLAUDE, Columbia University
SCHWARTZ, SUE, Russell Sage College
SELDIN, EDWARD B., Harvard School of Dental Medicine
SHAPLEY, ROBERT, The Rockefeller University
SHAW, STEPHEN R., University of St. Andrews, Scotland
SIEGEL, BARRY W., Rensselaer Polytechnic Institute
SINDELAR, WILLIAM F., Western Reserve University
SINGER, IRWIN, New York University
SLOANE, MOLLA R., University of Virginia
SMARSH, ANNE, VA Hospital, Boston
SQUIRE, RICHARD D., North Carolina State University

STEINBERG, WILLIAM, University of Wisconsin
 STOKES, DARRELL R., University of Hawaii
 STONER, LARRY C., Syracuse University
 STRUNK, BRIAN L., Princeton University
 SUBERMAN, RICK, University of North Carolina
 SURGENOR, PETER D., Hamilton College
 SURVER, WILLIAM M., University of Notre Dame
 TAM, ANTHONY, State University of New York, Syracuse
 TANNENBAUM, BERNICE, National Institutes of Health
 TANNER, SUSANE J., University of Hawaii
 TARR, MERRILL, Duke University
 THERIAULT, JACQUES, University of Ottawa
 TUTTLE, JOAN P., University of Rochester
 VARY, JAMES C., University of Wisconsin
 WEIANS, PATRICIA, College of St. Mary of the Springs
 WEINER, BEVERLY, Harvard Medical School
 WIKSELL, AUDRE, Washington University School of Medicine
 WILLENBORG, DAVID O., The Johns Hopkins School of Hygiene
 WILSON, FRANK J., University of Pittsburgh
 YOSHIOKA, PAUL M., Tulane University
 YOUNGER, W. B. JERRY, Washington University School of Medicine
 YUTMAN, MARCIA, Columbia University
 ZUClich, JOSEPH A., Columbia University

Library Readers, 1967

ADELBERG, EDWARD, Professor of Microbiology, Yale University
 ALLEN, GARLAND E., Allston Burr Senior Tutor, Professor in the History of Science, Harvard University
 BALL, ERIC G., Professor of Biochemistry, Harvard Medical School
 BATTELLE MEMORIAL INSTITUTE, William F. Clapp Laboratories, Inc.
 BENJAMIN, THOMAS L., Assistant Member, American Medical Association
 BENNETT, MIRIAM F., Professor of Biology, Sweet Briar College
 BERNE, ROBERT M., Chairman and Professor, University of Virginia Medical School
 BERRY, SPENCER J., Assistant Professor of Biology, Wesleyan University
 BERSOHN, RICHARD, Professor of Chemistry, Columbia University
 BODANSKY, OSCAR, Vice President, Chief, Division of Biochemistry, Sloan-Kettering Institute for Cancer Research
 BOETTIGER, EDWARD G., Professor of Physiology, University of Connecticut
 BRIDGMAN, ANNA JOSEPHINE, Chairman and Professor of Biology, Agnes Scott College
 BURBANCK, W. D., Professor of Biology, Emory University
 BUTLER, ELMER G., Osborn Professor of Biology, Princeton University
 CARLSON, FRANCIS D., Professor of Biophysics, The Johns Hopkins University
 CHANGEUX, J. P., Maître-assistant Faculty Science, Institut Pasteur-Paris and Columbia University
 CHASE, AURIN M., Professor of Biology, Princeton University
 CHAUNCEY, HOWARD H., Chief, Research in Oral Diseases, Veterans Administration Central Office, Washington, D. C.
 CLARK, ARNOLD M., Professor of Biology, University of Delaware
 CLEMENT, A. C., Professor of Biology, Emory University
 CLEWE, THOMAS H., Research Associate, Genetic & Developmental Disorders Division, Tulane University
 COHEN, EDWARD P., Associate Professor of Microbiology & Medicine, Rutgers Medical School
 COOPER, JACK ROSS, Associate Professor of Pharmacology, Yale University School of Medicine
 CRANE, ROBERT K., Chairman and Professor of Physiology, Rutgers Medical School
 CROWELL, SEARS, Professor of Zoology, Indiana University
 DE VILLAFRANCA, GEORGE W., Professor, Clark Science Center, Smith College
 EDDS, M. V., JR., Professor of Biology, Brown University

- ELLIOTT, ALFRED M., Professor of Zoology, University of Michigan
 FINGERHUT, SR. MARION ROBERT, Graduate Student, Catholic University of America
 FLAVIN, REV. JOHN W., Associate Professor of Biology, College of the Holy Cross
 FLESCH, PETER, Research Professor of Dermatology, University of Pennsylvania
 GABRIEL, MORDECAI L., Chairman and Professor of Biology, Brooklyn College
 GIBBS, MARTIN, Chairman and Professor of Biology, Brandeis University
 GITLIN, DAVID, Professor of Pediatrics, University of Pittsburgh School of Medicine
 GLADE, RICHARD W., Chairman and Professor of Zoology, University of Vermont
 GOTTSCHALL, GERTRUDE Y., Assistant Professor of Medicine, The Mount Sinai Hospital
 GREIF, ROGER L., Acting Chairman and Professor of Physiology, Cornell University Medical College
 GUREWICH, VLADIMIR, Assistant Professor of Clinical Medicine, New York College of Medicine
 HANDLER, PHILIP, Chairman and Professor of Biochemistry, Duke University
 HURWITZ, CHARLES, Chief, Basic Science Research Laboratory, VA Hospital, Albany
 ILAN, JOSEPH, Assistant Professor of Biology, Temple University
 ISSELBACHER, KURT J., Professor of Medicine, Massachusetts General Hospital
 JACOBS, MERKEL H., Emeritus Professor of Physiology, University of Pennsylvania
 JOHNSON, WILLIAM H., Chairman and Professor of Biology, Rensselaer Polytechnic Institute
 KEMPTON, RUDOLF T., Professor of Biology, Vassar College
 KENNEDY, EUGENE P., Hamilton Kuhn Professor of Biochemistry, Harvard Medical School
 KEOSIAN, JOHN, Professor of Biology, Rutgers—The State University
 KIORTSIS, VASSILI, Director and Professor of Zoological Laboratory & Museum, University of Athens, Greece
 KROPP, ALLEN, Associate Professor of Chemistry, Amherst College
 LINEVINTHAL, CYRUS, Professor of Biophysics, Massachusetts Institute of Technology
 LINEAWEAVER, THOMAS, III, Marine Biological Laboratory
 MARKS, PAUL A., Associate Professor, Columbia University, College of Physicians & Surgeons
 MATEYKO, GLADYS M., Professor of Biology, New York University
 MOSCONA, A. A., Professor of Zoology, University of Chicago
 NACHMANSOHN, DAVID, Professor of Biochemistry, Columbia University, College of Physicians & Surgeons
 NASON, ALVIN, Professor of Biology, Associate Director of McCollum-Pratt Institute, The Johns Hopkins University
 NOVIKOFF, ALEX B., Research Professor, Albert Einstein College of Medicine
 PERLMAN, GERTRUDE E., Associate Professor, The Rockefeller University
 PERLMAN, HANS PETER, Associate Professor of Immunology, Wenner-Gren Institute
 REICHHARDT, WERNER E., Director, Max-Planck-Institut für Biologie, Germany
 ROWLAND, LEWIS P., Professor of Neurology, Columbia University, College of Physicians & Surgeons
 RUGH, ROBERTS, Associate Professor of Radiology, Columbia Medical Center
 SAGER, RUTH, Professor, Hunter College
 SCHLESINGER, R. WALTER, Chairman of Department of Microbiology and Assistant Dean, Rutgers Medical School
 SCHOPF, THOMAS J. M., Systematics-Ecology Program, Marine Biological Laboratory
 SILMAN, ISRAEL, Research Associate, Columbia University, College of Physicians & Surgeons
 SIMON, JOSEPH L., Systematics-Ecology Program, Marine Biological Laboratory
 SPECTOR, ABRAHAM, Assistant Professor of Ophthalmology, Columbia University
 SPERELAKIS, NICK, Professor of Physiology, University of Virginia School of Medicine
 STETTEN, DEWITT, Dean, Rutgers Medical School
 STETTEN, MARJORIE R., Research Professor of Experimental Medicine, Rutgers Medical School
 SUMMERS, WILLIAM C., Systematics-Ecology Program, Marine Biological Laboratory
 TRAVIS, DOROTHY F., Assistant Biologist, Massachusetts General Hospital
 TWAROG, BETTY M., Assistant Professor of Biology, Tufts University
 WADE, BARRY A., Systematics-Ecology Program, Marine Biological Laboratory
 WALNIO, WALTER, Professor of Biochemistry, Rutgers-The State University
 WALLACE, STANLEY L., Associate Director of Medical Service and Associate Clinical Professor of Medicine, Jewish Hospital and State University of New York, Downstate Medical Center

WHEELER, GEORGE E., Associate Professor of Biology, Brooklyn College
 WHITING, ANNA R., Consultant, Oak Ridge National Laboratory
 WICHTERMAN, RALPH, Professor of Biology, Temple University
 WILSON, THOMAS HASTINGS, Associate Professor of Physiology, Harvard Medical School
 YNTEMA, CHESTER L., Professor of Anatomy, State University of New York, at Syracuse
 YOUNG, DAVID K., Systematics-Ecology Program, Marine Biological Laboratory
 ZACKS, SUMNER I., Associate Professor of Pathology, University of Pennsylvania School of
 Medicine & Pennsylvania Hospital
 ZULLO, VICTOR A., Systematics-Ecology Program, Marine Biological Laboratory

Students, 1967

All students listed completed the formal course program, June 12–July 22. Asterisk indicates students completing post-course research program, July 23–September 2.

ECOLOGY

ANDERSON, ROBERT, Syracuse University
 *BEARD, ELIZABETH, University of Wisconsin
 BRODSKY, DIANE, Northwestern University
 FOX, JOHN, The Johns Hopkins University
 *HALL, JOHN, University of North Carolina
 *HALVERSON, ROGER, Syracuse University
 HEITHAUS, EARL, Kenyon College
 *KEELEY, ROSEMARY, Yale University
 KIRCHER, ANN, Duke University
 MCCORMACK, BRIAN, University of Miami
 *MEADE, JAMES, Cornell University
 *MILLER, NANCY, The Johns Hopkins University
 *NYBLADE, CARL, Oberlin College
 SCHNEIDER, DOUGLAS, Amherst College
 *SHERWOOD, MARJORIE, University of California, Los Angeles
 SHULMAN, RICHARD, University of Illinois
 *SPOON, DONALD, Emory University
 WHITE, ALAN, Harvard University
 WURSTER, CHARLES, State University of New York

EMBRYOLOGY

ANDERSON, PAUL N., National Institutes of Health, National Cancer Institute
 *ASTRIN, KENNETH H., University of California, San Diego
 BOETTIGER, DAVID, Western Reserve University
 BROWDER, LEON W., University of Minnesota
 *FRY, BARRY J., Massachusetts Institute of Technology
 *GARFIELD, SANFORD, University of Illinois
 *GRANHOLM, NORMAN A., University of Oregon
 HAKE, JONATHAN C., Washington University
 HELFENSTEIN, MYRNA C., Washington University
 HOUSEHOLDER, SALLY C., The Rockefeller University
 *HUZ, SHULAMITH, New York University, Washington Square College
 *KEDES, LAURENCE H., Massachusetts Institute of Technology
 *KERBY, CATHERINE J., Institute of Molecular Evolution, University of Miami
 *MCMORRIS, F. ARTHUR, Yale University
 *PHILLIPS, DWIGHT E., Tulane University
 RECHSTEINER, MARTIN C., The Johns Hopkins University
 ROZIN, PAUL, University of Pennsylvania
 WATSON, LESLEY F., Yale University

- *WEBSTER, DALE A., Harvard Medical School & Massachusetts General Hospital
 WESLEY, RONALD D., Oregon State University

MARINE BOTANY

- ANDERSON, DARYL J., Washington University
 *ARCHIBALD, PATRICIA ANN, University of Texas
 *CAMPBELL, LOUISE, University of Colorado
 DURKIN, SISTER EILEEN M., RSM, University of California, Berkeley
 *ERLBAUM, PATRICIA A., Fordham University
 GIESMANN, LARRY ALLEN, University of Kentucky
 *HENDRIX, MARY L., Wellesley College
 KAZAMA, FREDERICK, University of California, Berkeley
 LEE, ROBERT EDWARD, University of Massachusetts
 LEE, ROBERT WINGATE, State University of New York, Stony Brook
 McCRACKEN, MICHAEL D., Indiana University
 *MENDEN, TRINA M., Goucher College
 *MULLINS, JOHN Q., University of Tennessee
 *REDEMSKE, JOYCE, University of Illinois
 *ROY, VICTORIA M., College of New Rochelle
 SPOON, JEANETTE A. H., Emory University
 STEWART, DOROTHY J., Drew University
 TUCKER, LYNNE, Oberlin College
 VANDE BERG, WARREN JAMES, Indiana University
 *VILLALARD, MARTINE, University of Rhode Island

PHYSIOLOGY

- *ABZUG, CHARLES, New York Medical College
 *ANDERSON, CARL W., Washington University
 ASBELL, MARY ANN, University of Georgia
 *BELLEMARE, GUY, University of Montreal
 *BLUMENTHAL, THOMAS, The Johns Hopkins University
 CHAN, TEH-SHENG, Yale University
 *CHEN, VICTOR KAI-HWA, State University of New York, at Buffalo
 DEARMAN, HENRY H., University of North Carolina
 *EBERHARD, CAROLYN, Boston University
 ELLIOTT, DONALD A., The Rockefeller University
 FEHER, GEORGE, University of California, San Diego
 GLEICH, GERALD J., Mayo Clinic
 *HILLMAN, PETER, Weizmann Institute, Rehovoth, Israel
 *HORCH, KENNETH W., Yale University
 *HOWELL, WILLIAM H., Columbia University
 *KIRSCHNER, MARC, The Rockefeller University
 KLOCK, PETER A., The Johns Hopkins University
 *KOWIT, JOEL D., Brandeis University
 MUNSE, PATRICIA A., Harvard University
 *NEALSON, KENNETH H., University of Chicago
 *NICKERSON, KENNETH W., University of Cincinnati
 NOSSAL, RALPH J., National Institutes of Health
 OSBORN, MARY, Pennsylvania State University
 PARDUE, MARY LOU, Yale University
 *PIPER, JUDITH A., Marquette University
 REIF, FREDERICK, University of California, Berkeley
 *SEKAER, CHRISTINA M., Western Reserve University
 SHIH, AGNES AN-YA, Yale University
 *SWENARCHUK, LAUREN C., Western Reserve University
 *TABER, ROBERT L., JR., University of Pittsburgh

Auditors : DONALD GRAHAM, Case Western Reserve University
 JEROME S. HARRIS, Duke University
 MARTIN POSNER, Yale University

INVERTEBRATE ZOOLOGY

ASHER, KENNETH D., Ohio Wesleyan University
 BAILEY, BARBARA ANN, University of Oklahoma
 BAILEY, KANIAULONO HELEN, Acadia University
 BALDERSTON, W. LLOYD, Long Island University
 BENNETT, ALBERT F., University of Michigan
 BILLINGS, SUSAN M., Harvard Medical School
 BRUNS, ROMAINE R., Harvard Medical School
 CHENEY, CLARISSA M., Goucher College
 *COLLIER, MARJORIE M., Brooklyn College
 *CRAMER, NANCY MARIE, George Washington University
 CUMBIE, PETER M., Yale University
 EBNER, FORD FRANCIS, Brown University
 ENGELHARDT, CHARLES D., Drew University
 ERICKSON, GREGORY F., University of Illinois
 ETTIENNE, EARL MANUAL, State University of New York, at Albany
 FINGERHUT, SISTER MARION ROBERT, Catholic University of America
 GABE, PHILIP R., University of California, Berkeley
 HENNINGER, ANN LOUISE, Wilson College
 *HUNTER, R. DOUGLAS, Syracuse University
 *JOHNSTON, MICHAEL A., Colorado College
 *KRASILOVSKY, GEORGE H., University of Oregon
 KUCINSKI, HELEN MARY, University of Massachusetts
 KUPRYS, JEANNIE GENOVAITE, Connecticut College
 LUCKY, PAUL ANDREW, Kenyon College
 *McMAHON, ROBERT F., III, Syracuse University
 MYERS, ALLEN COWLES, University of Rhode Island
 NACE, A. GEORGE, III, Cornell University
 NATALINI, JOHN JOSEPH, Northwestern University
 NEWTON, WILLIAM DONALD, University of North Carolina
 OBERDORFER, MICHAEL D., University of Wisconsin
 *PARACER, S. M., University of Massachusetts
 PILLOW, CHRISTINA LOUISE, Biological Abstracts
 RHYS, ANN FERRIER, Oberlin College
 *RICE, HERBERT VIVIAN, JR., Harvard University
 RITZI, EARL MICHAEL, University of Massachusetts
 *SMITH, ELENORA HARRIET, University of Michigan
 TIREY, SANDRA LEE, Rice University
 *WADDELL, JAMES L. F., Glasgow University, Scotland
 WILLIAMS, GABRIEL O., University of California, Berkeley
 WRIGHT, VIRGINIA K., Emory University
 *YOUNG, JANICE E., Goucher College

4. FELLOWSHIPS AND SCHOLARSHIPS, 1967

The Lucretia Crocker Scholarship :

ROGER C. HALVERSON, Marine Ecology Course

MARTINE VILLALARD, Marine Botany Course

The James W. Mavor Scholarship :

JAMES L. F. WADDELL, Invertebrate Zoology Course

The Turtox-Croasdale Scholarship:

PATRICIA ERLBAUM, Marine Botany Course

Gary N. Calkins Memorial Scholarship:

HARBERT VIVIAN RICE, JR.

ELNORA HARRIET SMITH

James Watt Mavor Scholarship:

JAMES L. F. WADDELL

5. TRAINING PROGRAM

FERTILIZATION AND GAMETE PHYSIOLOGY RESEARCH TRAINING PROGRAM

I. INSTRUCTORS

CHARLES B. METZ, University of Miami, Program Chairman

C. R. AUSTIN, Tulane University, Delta Regional Primate Center

DAVID BISHOP, Carnegie Institution of Washington

GEORGE G. BROWN, University of Miami

GIOVANNI GIUDICE, University of Palermo, Italy

GERTRUDE HINSCH, University of Miami

ALBERTO MONROY, University of Palermo, Italy

II. TRAINEES

CLAUDIO BARROS, University of Tulane, Delta Regional Primate Center

GRACIELA CANDELAS, University of Puerto Rico

JOHN CHAMBERLAIN, Princeton University

CHARLES FOURTNER, Carroll College

VITAUTS KALNINS, NATO Science Post-Doctoral, Harvard University

LAWRENCE KOEHLER, Central Michigan University

NORMAN LAZAROFF, Yale University

PATRICIA OLDS, Washington University

RICHARD PELTZ, University of Pennsylvania

HARRY ROY, The Johns Hopkins University

LEONARD SCHECHTMAN, The Johns Hopkins University

W. WINSLOW SCHRANK, University of Maryland

ROSS SHOGER, Associate Professor, Carlton College

PHILIP SKEHAN, Yale University

RICHARD WEISENBERG, University of Chicago

NATILLE HEADRICK, University of Tennessee

III. LECTURES

DAVID BISHOP Testicular sorbitol dehydrogenase in relation to differentiation and modification of germinal epithelium

I. FRIEDMAN Cytology of fertilization in algae

A. CLARK Aberrant fertilization types in the wasp *Habrobracon*

L. BARTH Parallelism between induction and egg activation

P. SOUPART Studies on the hormonal control of rabbit sperm capacitation

D. PATANELLI Physiological control of spermatogenesis

C. VILLEE The role of RNA in mediating the effects of testosterone

H. GRUNDFEST The egg as an electrophysiological system

J. MARSH Egg cell lipids—the lighter side of reproduction?

L. PIKO Some studies of membranes during fertilization

A. MACLAREN Fertilization in the mouse

M. BEDFORD Some electron microscopic studies of fertilization

6. TABULAR VIEW OF ATTENDANCE, 1963-1967

	1963	1964	1965	1966	1967
INVESTIGATORS—TOTAL	490	512	572	555	590
Independent	261	273	284	287	313
Library Readers	51	47	62	77	78
Research Assistants	178	192	227	191	199
STUDENTS—TOTAL	124	126	128	126	132
Invertebrate Zoology	40	40	41	37	41
Embryology	20	20	20	22	20
Physiology	28	30	30	29	31
Botany	20	19	20	18	20
Ecology	16	17	17	20	20
TRAINEES—TOTAL		30	34	29	16
Nerve-Muscle		7	7	7	
Comparative Physiology		7	11	6	
Fertilization & Gamete		16	16	16	16
TOTAL ATTENDANCE	614	668	734	710	738
Less persons represented in two categories	5	7	4	0	4
	<u>609</u>	<u>661</u>	<u>730</u>	<u>710</u>	<u>734</u>
INSTITUTIONS REPRESENTED—TOTAL	120	140	218	198	177
By Investigators	83	117	142	105	114
By Students	73	23	76	76	74
By Library Readers				47	74
By Research Assistants				80	75
FOREIGN INSTITUTIONS REPRESENTED	21	32	27	28	29
By Investigators	15	28	25	20	24
By Students	6	4	2	4	3
By Library Readers				1	3
By Research Assistants				4	7

7. INSTITUTIONS REPRESENTED, 1967

Agnes Scott College	California, University of, at San Diego
Albert Einstein College of Medicine	California, University of, at Santa Barbara
American Medical Association, Chicago	Carlton College
American Museum of Natural History	Carnegie Institution of Washington
Amherst College	Catholic University of America
Argonne National Laboratory	Central Michigan University
Augustana College	Chicago, University of
Barnard College	Cincinnati, University of
Battelle Memorial Institute	Clapp, Wm. F., Laboratories
Biological Information Service, Philadelphia	Colgate University
Boston University	Colorado College
Brandeis University	Colorado, University of, Medical School
Brooklyn College, City University of New York	Columbia Medical Center
Brown University	Columbia University
Bryn Mawr College	Columbia University, College of Physicians & Surgeons
California Institute of Technology	Connecticut College
California, University of, at Berkeley	Connecticut, University of
California, University of, at Los Angeles	Cornell University
California, University of, at Riverside	Cornell University Medical College

- Dartmouth College
 Delaware, University of
 Delta Regional Primate Research Center, Tu-
 lane University
 Drew University
 Duke University
 Emory University
 Fordham University
 Georgetown University
 Georgetown University Medical School
 George Washington University
 Georgia, University of
 Goucher College
 Hamilton College
 Harvard College
 Harvard Medical School
 Harvard School of Dental Medicine
 Harvard University
 Hawaii, University of
 Holy Cross, College of the
 Hunter College, City University of New York
 Illinois Institute of Technology
 Illinois, University of
 Indiana University
 Indiana University Medical Center
 Institute for Cancer Research, Philadelphia
 Institute for Muscle Research, Woods Hole
 Jewish Hospital, Brooklyn
 Johns Hopkins University, The
 Johns Hopkins University School of Hygiene
 & Public Health, The
 Johns Hopkins University School of Medicine,
 The
 Kansas, University of
 Kentucky, University of
 Long Island University
 Marquette University
 Maryland, University of
 Maryland, University of, School of Medicine
 Massachusetts General Hospital
 Massachusetts Institute of Technology
 Massachusetts, University of
 Mayo Clinic
 McLean Hospital Research Laboratory
 Mellon Institute, The
 Miami University at Ohio
 Miami, University of
 Miami, University of, Institute of Molecular
 Evolution
 Michigan State University
 Michigan, University of
 Minnesota, University of
 Mount Holyoke College
 Mount Sinai Hospital
 National Institutes of Health
 New Mexico, University of, Medical School
 New Rochelle, College of
 New York, City College of
 New York Medical College
 New York, State University of, Albany
 New York, State University of, Buffalo
 New York, State University of, Stony Brook
 New York, State University of, College of
 Medicine, Upstate Medical Center, Syracuse
 New York University
 New York University, College of Dentistry
 New York University, Institute of Environ-
 mental Medicine
 New York University Medical School
 New York University, Washington Square
 College
 North Carolina State University at Raleigh
 North Carolina, University of
 Northeastern Illinois State College
 Northwestern University
 Notre Dame, University of
 Oak Ridge National Laboratory
 Oberlin College
 Ohio Wesleyan University
 Oklahoma, University of
 Oregon State University
 Oregon, University of
 Pennsylvania Hospital
 Pennsylvania State University
 Pennsylvania, University of
 Pennsylvania, University of, School of Medi-
 cine
 Pittsburgh, University of
 Pittsburgh, University of, School of Medicine
 Princeton University
 Providence College
 Queens College, City University of New York
 Radcliffe College
 Reed College
 Rensselaer Polytechnic Institute
 Rhode Island, University of
 Rice University
 Rochester, University of
 Rochester, University of, School of Medicine
 & Dentistry
 Rockefeller University, The
 Russell Sage College
 Rutgers, The State University, New Bruns-
 wick
 Rutgers University, Newark
 Rutgers University Medical School
 St. John's College
 St. Louis University
 St. Mary of the Springs, College of
 San Diego State College
 Simmons College
 Single Cell Research Foundation, Inc.
 Sloan-Kettering Institute for Cancer Research
 Smith College
 Smithsonian Institution
 Southern Connecticut State College

Stanford University	Veterans Administration Hospital, Brooklyn
Sweet Briar College	Veterans Administration Hospital, Pittsburgh
Syracuse University	Virginia Polytechnic Institute
Temple University	Virginia, University of
Tennessee, University of	Virginia, University of, Medical School
Texas, University of	Wards Natural Science Company
Toledo, University of	Washington University, St. Louis
Trinity College, Hartford	Washington University School of Medicine
Trinity College, Washington, D. C.	Wellesley College
Tufts University	Wesleyan University
Tulane University	Western Reserve University
Vassar College	Western Reserve University School of Medicine
Vermont, University of	Wilson College
Veterans Administration Center, Los Angeles	Wisconsin, University of
Veterans Administration Central Office, Washington, D. C.	Yale University
Veterans Administration Hospital, Albany	Yale University School of Medicine
Veterans Administration Hospital, Boston	Yeshiva University

FOREIGN INSTITUTIONS REPRESENTED, 1967

Acadia University, Nova Scotia	Marine Science Laboratory, England
Athens, University of, Greece	Max-Planck Institut für Biologie, Germany
Bedford College, University of London	Medical Research Council, England
Biologische Anstalt Helgoland, West Germany	MRC Laboratory of Molecular Biology, England
British Columbia University, Vancouver	Montreal University, Canada
Cambridge, University of, England	Okayama University, Japan
Chile, University of	Ottawa, University of, Canada
Dalhousie University, Nova Scotia	Palermo, University of, Italy
Ehime University, Japan	Puerto Rico, University of
Geneva, University of, Switzerland	St. Andrews, University of, Scotland
Genoa University, Italy	Teheran University, Iran
Glasgow University, Scotland	Tohoku University, Japan
Institut Pasteur, Paris	Toronto, University of, Canada
Instituto Venezolano de Investigaciones Científicas, Venezuela	University College, London, England
International Laboratory of Genetics and Biophysics, Italy	Weizmann Institute, Israel
Ljubljana University, Yugoslavia	Wenner-Gren Institute, Sweden

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The Ford Foundation	Mr. Franklin A. Park
Mr. John A. Gifford	Mrs. Arthur Kemble Parpart
Mr. W. T. Golden	Schering Corporation
The Grass Foundation	Mr. & Mrs. Gerard Swope, Jr.
The Lalor Foundation	Whitehall Foundation, Inc.

8. FRIDAY EVENING LECTURES, 1967

July 7

ALBERTO MONROYRecent Studies on the Molecular Aspects of the
University of Palermo, Italy Activation of the Egg

July 13

WERNER REICHARDT On Structure and Function of Retina and Lamina
 Max-Planck-Institut für Biologie in Diptera
 Alexander Forbes Lecturer at the
 Marine Biological Laboratory

July 14

WERNER REICHARDT Analysis of Optomotor Responses

July 21

ROBERT A. GOOD Development and Involution of Immunologic Ca-
 University of Minnesota Medical pacity
 School

July 28

PHILIP ABELSON Chemical Events on the Primitive Earth
 Editor, Science

August 4

MICHAEL FISCHBERG Information and Development in *Xenopus laevis*
 Université De Geneva, Switzerland
 Senior Lalor Fellow

August 11

G. ADRIAN HORRIDGE The Action of the Retina in Arthropod Compound
 University of St. Andrews, Scotland Eyes of the Apposition Type
 Rand Fellow

August 18

KENNETH S. COLE Membrane Watching
 National Institutes of Health
 (NINDB)

August 25

PETER PERLMANN Aggressor Function of the Vertebrate Lympho-
 University of Stockholm, Sweden cytes: Its Mechanism and Biological Significance
 F. R. Lillie Fellow

9. TUESDAY EVENING SEMINARS, 1967

July 11

K. RANGA RAO Hormones Controlling the White Chromatophores
 MILTON FINGERMAN of the Fiddler Crab, *Uca pugilator*
 C. K. BARTELL Is There a Physiological Limit to the Response of
 Melanophores in *Uca* to the Melanin-dispersing
 Hormone?
 R. B. LOFTFIELD Comparative Biochemistry of RNA and tRNA
 E. A. EIGNER Ligase Reactions

July 18

NORMAN B. RUSHFORTH Behavioral Responses of Isolated Tentacles of
 Hydra
 JOSEPH L. SIMON Behavioral Aspects of *Histriobdella homari*, an
 Annelid Commensal of the American Lobster

- DANIEL S. GROSCH Biosatellite Experiments
 I. General Design of Experiments
 II. *Habrobracon* Investigations
- July 25
- S. EHRENPREIS Modification of Smooth Muscle Receptors by Heat and Urea
- S. LERMAN The Relationship Between Soluble and Insoluble Protein in the Lens
- S. ZIGMAN Demonstration of Phospholipid Splitting as the Factor Responsible for Increased Permeability and Block of Axonal Conduction Induced by Snake Venom—Study on Squid Axons
- P. ROSENBERG Studies on the Penetration of Some Organophosphorus Cholinesterase Inhibitors into the Squid Giant Axon
- F. C. G. HOSKIN Studies on the Penetration of Some Organophosphorus Cholinesterase Inhibitors into the Squid Giant Axon
- August 1
- RICHARD CONE Early Receptor Potential: Photoreversible Charge Displacement in Rhodopsin
- BRUCE GOLDSTEIN Source of the Early Receptor Potential in the Isolated Frog Retina
- JOEL E. BROWN and Photoelectric Potentials in the *Limulus* Eye
 TOM SMITH
- W. A. HAGINS and The Origin of Fast Photoelectric Effects in the
 R. E. MCGAUGHY Squid Retina
- August 8
- W. D. HUMMON Interstitial Marine Gastrotrichs from Woods Hole, Massachusetts
- B. A. WADE Ecological Studies of New England Nudibranchs
- P. PERSON and Water in Biological Systems I. Solubilization of
 H. ZIPPER Yeast and Heart Mitochondrial Cytochrome Oxidase and Cytochrome *b* by Synthetic Zeolites
- J. FELTON Sequential Solubilization of Mitochondrial Cytochrome System Components at Alkaline pH's
- H. ZIPPER and
 P. PERSON
- August 15
- W. W. SCHRANK Electrically Induced Spawning in the Male and
 R. L. SHOGER Female Horseshoe Crab, *Limulus polyphemus*
- L. M. SCHACHTMAN
- D. W. BISHOP
- M. LOCHHEAD and The Development of Oocytes in the Brine Shrimp,
 J. H. LOCHHEAD *Artemia*
- L. C. STONER Temperature Acclimation in *Tetrahymena pyriformis*
- R. H. GREEN Patterns of Invertebrate Distribution in the Barn-
 K. D. HOBSON stable Harbor Intertidal Area
- S. L. SANTOS
- J. A. MILLER, JR. Temperature, pH and Glucose in Experimental Asphyxia of the Newborn

10. MEMBERS OF THE CORPORATION, 1967

Including Action of 1967 Annual Meeting

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WARE, MR. AND MRS. J. LINDSAY	WILSON, DR. MAY G.
WARREN, DR. AND MRS. SHIELDS	WITMER, DR. AND MRS. ENOS
WATT, MR. AND MRS. JOHN B.	WOLFE, DR. CHARLES
WEBSTER, MRS. EDWIN S.	WOLFINSOHN, MRS. WOLFE
WEEDON FOUNDATION, THE	WRINCH, DR. DOROTHY
WESSLING, DR. AND MRS. PHILIP I.	YNTEMA, MRS. CHESTER L.
WHITELEY, MR. AND MRS. GEORGE C., JR.	ZWILLING, MRS. EDGAR

V. REPORT OF THE LIBRARIAN

Photocopying has become a major activity of the Library. During the summer months two Xerox machines were running constantly and an extra employee was hired to operate them. The machines are located in a library room on the third floor of the Lillie building.

The copying service has also doubled the interlibrary loan requests processed over the previous year. During 1967 we received 4,417 separate requests for articles contained here in the library. The requests were sent from universities, government agencies, hospitals, industrial firms and individual students and researchers. We made 342 requests from other libraries for the use of investigators here. Our list of "Serial Publications," containing approximately 4,000 separate titles, has been placed in over 800 libraries throughout the country.

Over 2,000 volumes were sent to the bindery in 1967 and the total holdings are now 133,447. This figure does not include the reprint floor.

Total number of serial titles in library	3,962
Number received currently	2,292
On subscription	921
On exchange	945
On gift basis	345
Number of reference books added in 1966	532
Received from book exhibitors	167
Total number of reprints in collection	240,417
Number added in 1967	2,197

Respectfully submitted,

JANE FESSENDEN,

Librarian

VI. REPORT OF THE TREASURER

The market value of the General Endowment Fund and the Library Fund at December 21, 1967, amounted to \$2,313,618, and the corresponding securities are entered in the books at a value of \$1,397,833. This compares with values of \$2,249,448 and \$1,326,668, respectively, at the end of the preceding year. The average yield on the securities was 3.99% of the market value and 6.6% of the book value. Uninvested principal cash was \$4,728. Classification of the securities held in the Endowment Fund appears in the Auditor's Summary of Investments.

The market value of the Pooled Securities at December 31, 1967, amounted to \$710,152 as compared to book values of \$618,416. These figures compare with values of \$701,945 and \$664,523, respectively, at the close of the preceding year. The average yield on the securities was 3.93% of the market value and 4.5% of the book value. Uninvested principal cash was in the amount of \$387.

In October, the Laboratory sold its holdings in General Biological Supply House for \$2,664,212, realizing a profit of \$2,651,512. The proceeds of this sale are currently invested in short-term commercial notes.

The proportionate interest in the Pool Fund Account of the various Funds, as of December 31, 1967, is as follows:

Pension Funds	19.992%
General Laboratory Investment	24.638%
F. R. Lillie Memorial Fund	2.685%
Anonymous Gift921%
Other:	
Bio Club Scholarship Fund695%
Rev. Arsenius Boyer Scholarship Fund851%
Gary N. Calkins Fund798%
Allen R. Memhard Fund156%
Lucretia Crocker Fund	2.907%
E. G. Conklin Fund491%
Jewett Memorial Fund258%
M. H. Jacobs Scholarship Fund351%
Herbert W. Rand Fellowship	24.501%
Mellon Foundation	11.727%
Mary Rogick Fund	2.571%
Swope Foundation	6.458%

Donations from MBL Associates for 1967 amounted to \$8,010 as compared with \$7,595 for 1966. A gift of a boat was received from Charles L. Morse, Jr., valued at \$12,000. Unrestricted gifts from foundations, societies and companies amounted to \$4,408.

Donations received during the year, donated specifically to the new laboratory under construction, amounted to \$5,900.

During the year we administered the following grants and contracts :

<i>Investigators</i>	<i>Training</i>	<i>MBL Institutional</i>
11 NIH	3 NIH	3 NIH
4 NSF	2 NSF	3 NSF
2 FORD		1 FORD
1 Whitehall		2 ONR
2 ONR		1 AEL
—	—	—
20	5	10

The rate of overhead on continuing grants from Federal agencies is effectively 20% of the amounts expended. For new Federally funded research grants a provisional rate of 25% has been negotiated. However, the Laboratory is required to make an identifiable contribution to the total cost of each research project. The overhead on those grants wherein provision therefore was made in the funds awarded within the year, amounted to \$95,874 as compared with \$85,822 for the preceding year. A proposal to NSF for determining an indirect cost rate for the current year is in course of preparation.

The following is a statement of the auditors :

To the Trustees of Marine Biological Laboratory, Woods Hole, Massachusetts:

We have examined the balance sheet of Marine Biological Laboratory as at December 31, 1967, the related statement of operating expenditures and income and statement of funds for the year then ended. Our examination was made in accordance with generally accepted auditing standards, and accordingly included such tests of the accounting records and such other auditing procedures as we considered necessary in the circumstances. We examined and have reported on financial statements of the Laboratory of the year ended December 31, 1966.

In our opinion, the accompanying financial statements present fairly the assets, liabilities and funds of Marine Biological Laboratory at December 31, 1967 and 1966 and the results of its operations for the years then ended on a consistent basis.

The supplementary schedules included in this report were obtained from the Laboratory's records in the course of our examination and, in our opinion, are fairly stated in all material respects in relation to the financial statements, taken as a whole.

Boston, Massachusetts
March 29, 1968

LYBRAND, ROSS BROS. AND MONTGOMERY

It will be noted from the operating statement that the Laboratory activities for the year under review, amounted to a figure of a little over 1.6 million dollars, which amount is comparable to last years results.

ALEXANDER T. DAIGNAULT,
Treasurer

MARINE BIOLOGICAL LABORATORY

MARINE BIOLOGICAL LABORATORY

BALANCE SHEETS

December 31, 1967 and 1966

Investments

Investments held by Trustee:

	1967	1966
Securities, at cost (approximate market quotation 1967—\$2,313,618)	\$1,397,833	\$1,326,668
Cash	4,728	1,105
	<hr/>	<hr/>
	1,402,561	1,327,773

Investments of other endowment and unrestricted funds:

Pooled investments, at cost (approximate market quotation 1967— \$710,152) less \$5,728 temporary investment of current fund cash	612,688	658,795
Other investments	2,803,850	121,370
Cash	387	50,108
Accounts receivable	54	1,393
Due from current fund	87,542	
	<hr/>	<hr/>
	\$4,907,082	\$2,159,439
	<hr/> <hr/>	<hr/> <hr/>

Plant Assets

Land, buildings, library and equipment (note)	5,695,732	5,649,105
Less allowance for depreciation (note)	1,624,698	1,537,471
	<hr/>	<hr/>
	4,071,034	4,111,634
Construction in progress	110,042	28,453
Cash		6,056
Short-term investments, at cost	50,000	50,000
Due from current funds	15,365	
	<hr/>	<hr/>
	\$4,246,441	\$4,196,143
	<hr/> <hr/>	<hr/> <hr/>

Current Assets

Cash	107,483	48,860
Temporary investment in pooled securities	5,728	5,728
U. S. Treasury bills, at cost		58,219
Accounts receivable (U. S. Government, 1967—\$83,619; 1966—\$55,247) .	139,771	127,817
Inventories of supplies and bulletins	46,372	40,525
Other assets	8,698	31,580
Due to plant funds	(15,365)	
Due to endowment funds	(87,542)	
	<hr/>	<hr/>
	\$ 205,145	\$ 312,729
	<hr/> <hr/>	<hr/> <hr/>

MARINE BIOLOGICAL LABORATORY

BALANCE SHEETS

December 31, 1967 and 1966

	<i>Invested Funds</i>	
	<i>1967</i>	<i>1966</i>
Endowment funds given in trust for benefit of the Marine Biological Laboratory	\$1,402,561	\$1,327,773
Endowment funds for awards and scholarships:		
Principal	427,663	427,663
Unexpended income	32,738	24,297
	460,401	451,960
Unrestricted funds functioning as endowment	2,857,890	206,378
Retirement fund	169,833	150,703
Pooled investments—accumulated gain	16,397	22,625
	<u>\$4,907,082</u>	<u>\$2,159,439</u>
	<i>Plant Funds</i>	
Funds expended for plant, less retirements	5,805,774	5,677,558
Less allowance for depreciation charged thereto	1,624,698	1,537,471
	4,181,076	4,140,087
Unexpended plant funds	65,365	56,056
	<u>\$4,246,441</u>	<u>\$4,196,143</u>
	<i>Current Liabilities and Funds</i>	
Accounts payable and accrued expenses	1,947	69,385
Advance subscriptions	15,210	16,746
Unexpended grants—research	32,353	61,780
Unexpended balances of gifts for designated purposes	23,899	22,281
Current fund	131,736	142,537
	<u>\$ 205,145</u>	<u>\$ 312,729</u>

Note—The Laboratory has since January 1, 1916 provided for reduction of book amounts of plant assets and funds invested in plant at annual rates ranging from 1% to 5% of the original cost of the assets.

MARINE BIOLOGICAL LABORATORY

STATEMENTS OF OPERATING EXPENDITURES AND INCOME

Years Ended December 31, 1967 and 1966

<i>Operating Expenditures</i>		
	<i>1967</i>	<i>1966</i>
Research and accessory services	\$ 323,854	\$ 329,066
Instruction	219,345	179,964
Library and publications (including book purchases—1967, \$34,626; 1966, \$34,082)	96,850	105,857
Direct costs on research grants	451,756	466,824
Direct costs on institution support grants	89,228	110,993
	<hr/>	<hr/>
	1,181,033	1,192,704
Administration and general	182,008	139,888
Plant operation and maintenance	145,900	147,271
Dormitories and dining	203,058	198,668
Additions to plant from current fund		34,528
	<hr/>	<hr/>
	1,711,999	1,713,059
Less depreciation included in plant operation and dormitories and dining above but charged to plant funds	87,226	88,786
	<hr/>	<hr/>
	1,624,773	1,624,273
	<hr/>	<hr/>

<i>Income</i>		
Research fees	99,716	97,186
Accessory services (including sales of biological specimens—1967, \$42,247; 1966, \$34,551)	123,499	133,416
Instruction fees	27,350	28,750
Library fees, bulletins, subscriptions and other	74,131	64,705
Dormitories and dining income	140,257	141,393
Grants for support of institutional activities:		
Instruction and training	220,451	177,825
Support services	89,228	110,993
General	132,097	130,750
Reimbursements and allowances for direct and indirect costs on specific research grants	526,570	534,909
Gifts used for current expenses	24,418	17,125
Investment income used for current expenses	180,955	162,262
	<hr/>	<hr/>
	1,638,672	1,599,314
	<hr/>	<hr/>
Excess current income (expenditures)	\$ 13,899	(\$ 24,959)
	<hr/>	<hr/>

MARINE BIOLOGICAL LABORATORY

STATEMENT OF FUNDS

Year Ended December 31, 1967

	<i>Balance December 31, 1966</i>	<i>Gifts and Other Receipts</i>	<i>Invest- ment Income</i>	<i>Used for Current Expenses</i>	<i>Other Expendi- tures</i>	<i>Balance December 31, 1967</i>
Invested funds	\$2,159,439	\$2,746,799	\$200,383	\$ 179,079	\$ 20,460	\$4,907,082
Unexpended plant funds	<u>\$ 56,056</u>	87,516	3,382		81,589	<u>\$ 65,365</u>
Unexpended research grants	<u>\$ 61,780</u>	938,919		968,346		<u>\$ 32,353</u>
Unexpended gifts for designated purposes	<u>\$ 22,281</u>	28,825		24,418	2,789	<u>\$ 23,899</u>
Current fund	<u>\$ 142,537</u>	13,899(1) (12,000)(2) (12,700)(3)				<u>\$ 131,736</u>
		<u>\$3,791,258</u>	<u>\$203,765</u>	<u>\$1,171,843</u>	<u>\$104,838</u>	
Gifts		34,753				
Grant for facilities construction		81,589				
Grant for research, training and support		938,919				
Appropriated from current income and other		26,726				
Net gain on sale of securities		2,720,072				
(1) Excess of current income over expenditures		13,899				
(2) Gift of boat transferred to plant funds expended ...		(12,000)				
(3) Loss on sale of gift in kind		(12,700)				
		<u>\$3,791,258</u>				
Expended for new Laboratory planning					81,589	
Scholarship awards .					6,530	
Payments to pen- sioners					13,360	
Other					3,359	
					<u>\$104,838</u>	

MARINE BIOLOGICAL LABORATORY

SUMMARY OF INVESTMENTS

December 31, 1967

	<i>Cost</i>	<i>% of Total</i>	<i>Market Quotations</i>	<i>% of Total</i>	<i>Investment Income 1967</i>
Securities held by Trustee:					
General endowment fund:					
U. S. Government securities	\$ 25,065	2.2	\$ 24,430	1.3	\$ 1,250
Corporate bonds	674,563	59.5	555,267	29.2	28,361
Preferred stocks	102,916	9.1	107,375	5.6	2,595
Common stocks	331,875	29.2	1,214,818	63.9	45,455
	<u>1,134,419</u>	<u>100.0</u>	<u>1,901,890</u>	<u>100.0</u>	<u>77,661</u>
General educational board endowment fund:					
U. S. Government securities	53,112	20.2	51,791	12.6	3,202
Other bonds	129,971	49.3	103,843	25.2	4,264
Preferred stocks	31,914	12.1	37,275	9.1	370
Common stocks	48,417	18.4	218,819	53.1	6,826
	<u>263,414</u>	<u>100.0</u>	<u>411,728</u>	<u>100.0</u>	<u>14,662</u>
Total securities held by Trustee	<u>\$1,397,833</u>		<u>\$2,313,618</u>		<u>\$ 92,323</u>
Investments of other endowment and unrestricted funds:					
Pooled investments:					
U. S. Government securities	2,000	.3	1,645	.2	1,229
Corporate bonds	227,194	36.8	181,380	25.6	9,838
Common stocks	389,222	62.9	527,127	74.2	16,845
	<u>618,416</u>	<u>100.0</u>	<u>\$ 710,152</u>	<u>100.0</u>	<u>27,912</u>
Less temporary investment of current fund cash	(5,728)				(266)
	<u>612,688</u>				<u>27,646</u>
Other investments:					
U. S. Government securities	27,938				1,133
Other bonds	15,029				750
Preferred stocks	3,448				135
Common stocks	46,186				57,174
Real estate	17,549				
Short-term commercial notes	2,693,700				30,392
	<u>2,803,850</u>				<u>89,584</u>
Total investments of other endowment and unrestricted funds	<u>\$3,416,538</u>				<u>117,230</u>

REPORT OF THE TREASURER

79

Total		209,553
Custodian's fees charged thereto		(9,170)
		<hr/>
Investment income distributed to invested funds		200,383
Plant investments:		
Federal agency and corporate bonds ..	\$ 50,000	3,382
	<hr/>	<hr/>
Current investments:		
U. S. Treasury bills		1,610
Temporary investment in pooled securities	\$ 5,725	266
	<hr/>	<hr/>
		1,876
		<hr/>
Total investment income		<u>\$205,641</u>

STUDIES ON MEMBRANE TRANSPORT. II. THE ABSORPTION OF
ACETATE AND BUTYRATE BY HYMENOLEPIS
DIMINUTA (CESTODA)¹

C. ARME AND C. P. READ

Department of Biology, Rice University, Houston, Texas 77001

Lipid metabolism in tapeworms has been studied recently by several investigators (Fairbairn *et al.*, 1961; Harrington, 1965; Ginger and Fairbairn, 1966a, b; Jacobsen and Fairbairn, 1967; Lumsden and Harrington, 1966; McMahon, 1961), and there is abundant evidence that *Hymenolepis diminuta* absorbs fatty acids from the external medium and incorporates them into lipid components of the tissues. Von Brand's hypothesis (1966), that the fatty acids in tapeworm tissues are waste products of carbohydrate metabolism, has not been supported by careful *in vivo* and *in vitro* experiments (Ginger and Fairbairn, 1966b; Jacobsen and Fairbairn, 1967), and all direct available evidence suggests that the lipids of *H. diminuta* are synthesized from fatty acids absorbed from the environment. On the other hand, although there is considerable evidence for the mediated absorption of amino acids, sugars, purines, and pyrimidines by this worm (reviewed by Read, 1966), there seem to be no published studies dealing with mechanisms for absorption of fatty acids by *H. diminuta* or other tapeworms. The present investigation was undertaken to characterize systems by which these presumably important nutrients may enter the tissues of some parasitic organisms.

MATERIALS AND METHODS

Read, Rothman and Simmons (1963) described techniques for the maintenance and experimental manipulation of the tapeworm, *Hymenolepis diminuta*. With few exceptions these methods were adopted during the present investigation.

Young male rats (Holtzman Rat Co., Madison, Wisconsin), weighing 80–100 g. at the time of infection, were used as hosts in all experiments; animals received a diet of Purina Laboratory Chow. Ten days following infection with 30 cysticercoids (obtained from previously infected *Tenebrio molitor*), the rats were killed and the intestines removed. Parasites were flushed from the intestine with Krebs-Ringer solution containing 25 mM tris(hydroxymethyl)aminomethane-maleic acid buffer at pH 7.4. This solution (KRT) was used for all subsequent washes and, except as otherwise stated, as a solute in experimental incubations. After removal from the intestine, worms were washed and randomized in groups of five. They were then preincubated in 10 ml. KRT for a period of 30 minutes in a shaker bath at 37° C., after which each five-worm sample was transferred to 4 ml. of incubation solution for a period of one minute. Following incubation worms were washed in KRT, blotted on hard filter paper and dropped into 5 ml. 70% ethanol. After

¹ This work was supported by grants from the National Institutes of Health, U. S. Public Health Service, AI-01384 and 5 TI AI 106.

18–24 hrs. the worms were removed from the ethanol, dried overnight in an oven at 90° C. and weighed. Aliquots of the ethanol extract were mixed with 0.5 ml. of a 5 mM sodium bicarbonate solution prior to drying on aluminum planchettes and counting on a gas flow counter. Uptake data are based on the amount of radioactive material extracted in 70% ethanol and are expressed as $\mu\text{moles/g./min.}$, using the ethanol-extracted dry weight.

Sodium acetate and sodium butyrate labelled with ^{14}C at the C-1 position were obtained from New England Nuclear Co. and Nuclear Chicago Co., respectively. Unlabelled sodium salts of valeric acid (Eastman Organic Chemicals) and octanoic acid (Matheson, Coleman, and Bell) were prepared by titration of the appropriate acid to pH 7.6 with sodium hydroxide. Formate, acetate, propionate and butyrate were purchased as sodium salts of reagent grade.

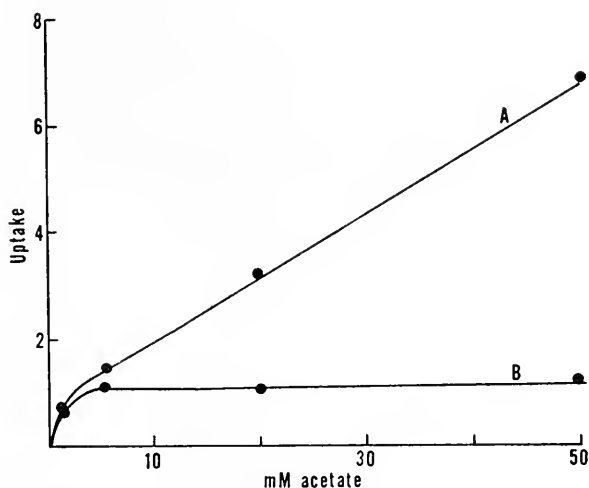


FIGURE 1. The uptake of ^{14}C -acetate as a function of acetate concentration. Curve A is uncorrected uptake. Curve B is uptake corrected by subtracting diffusion, calculated from the linear portion of Curve A. Each point is a mean of 4 determinations.

RESULTS

The relationship between acetate uptake and concentration is shown in Figure 1. The data indicate that the uptake system is not fully saturated even at the highest concentrations tested. At concentrations from 5 to 50 mM the uptake rate increases linearly with increase in concentrations, whereas below approximately 2 mM, acetate uptake is not a linear function of concentration. This suggests that a mediated transport mechanism operates during the uptake of acetate but, at high acetate concentrations, mediated transport is masked by a diffusion component. When it is assumed that at high concentrations of acetate, mediated uptake mechanisms are saturated, a diffusion rate of $0.12 \mu\text{moles/g./min.}$ per unit mM increase in acetate concentration is calculable from the data in Figure 1, curve A. If the data are corrected for a diffusion component of uptake, a typical adsorption isotherm is obtained (Fig. 1, curve B). However, the data do not rule out conclusively the

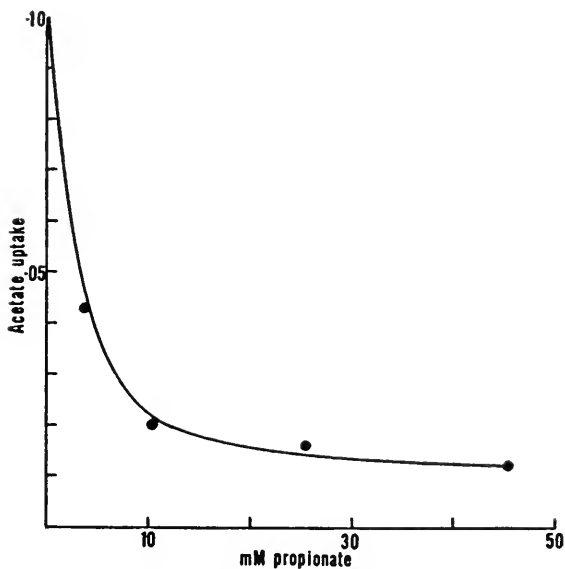


FIGURE 2. The effect of increasing concentrations of propionate on the uptake of acetate (0.1 mM). Each point is a mean of 8 determinations.

possibility that a second system for the mediated transport of acetate is involved, since a second system, saturable at very high concentrations, might appear to yield a linear relationship over the concentration range investigated during the present study. The latter probability is diminished, however, by the observation that propionate does not appear to affect the second component of acetate absorption (Fig. 2); the residual absorption of 0.1 mM acetate in the presence of 50 mM propionate is essentially equal to the diffusion value calculated from the linear portion of the uptake curve shown in Figure 1.

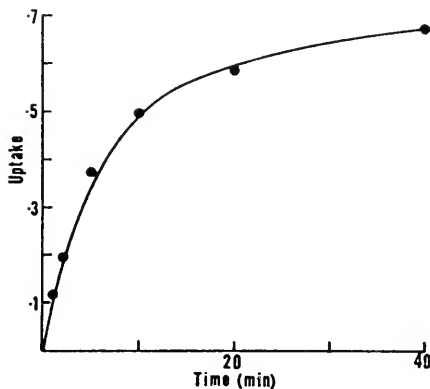


FIGURE 3. Ethanol-extractable ^{14}C , expressed as $\mu\text{moles } ^{14}\text{C-acetate/g}$, after incubation in 0.1 mM $^{14}\text{C-acetate}$. Each point is a mean of 4 determinations.

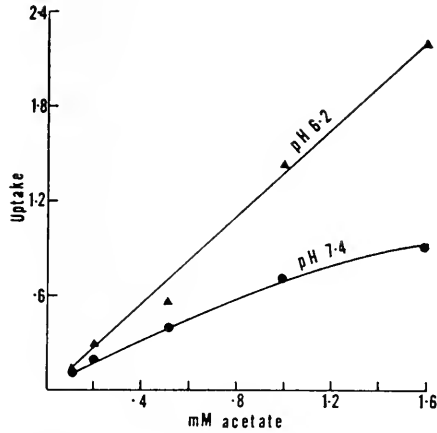


FIGURE 4. The uptake of ^{14}C -acetate at pH 6.2 and 7.4 as a function of acetate concentration. Each point is a mean of 8 determinations.

Analysis of the data on acetate uptake in terms of Michaelis-Menten kinetics allowed an evaluation of K_t (equivalent to Michaelis' constant) and V_{max} (extrapolated maximum velocity). In the concentration range 0.1 to 1.0 mM acetate, K_t and V_{max} were 1.13 mM and 1.25 $\mu\text{moles/g./min.}$, respectively, when the data were not corrected for diffusion. After correction, K_t was 0.92 mM and V_{max} was 0.91.

The uptake of sodium acetate with respect to time is shown in Figure 3. Following 40 minutes incubation in a substrate concentration of 0.1 mM the amount of radioactive material extracted from the worm indicated an internal acetate concentration in the worm water of 0.175 mM. Preliminary chromatography indicated that about 60% of the radioactivity extracted was unaltered acetate, but further study would be required to determine more precisely the proportion of the radioactivity representing metabolically unaltered acetate. However, it may be suggested that mediated acetate uptake is a facilitated diffusion rather than active transport.

TABLE I

Effects of pH on acetate uptake and on propionate inhibition of acetate uptake. Acetate concentration 0.1 mM; propionate concentration 4 mM

pH	Uptake ($\mu\text{moles/g./min.} \pm \text{S.E.}$)		% Inhibition
	Acetate alone	Acetate + propionate	
6.1	161 \pm 10.4	134 \pm 5.0	17
6.4	147 \pm 9.4	102 \pm 5.1	31
6.7	139 \pm 9.0	64 \pm 2.3	54
7.0	135 \pm 14.2	53 \pm 1.4	61
7.5	129 \pm 7.6	39 \pm 1.8	70

Influence of pH on acetate absorption

Worms were pre-incubated for 30 min. in KRT at pH 7.4 and then incubated for 1 min. in 0.1 mM acetate over a pH range of 6.2–8.0. Radioactivity in incubation solutions was assayed at the end of the experiments in order to determine whether any acetate was lost at low pH; no loss was detectable over the time period of the experiment. Acetate uptake increased as a function of decreasing pH and propionate inhibition of acetate decreased with decreasing pH (Table I). The uptake of acetate at pH 6.2 and 7.4 was studied over a substrate concentration

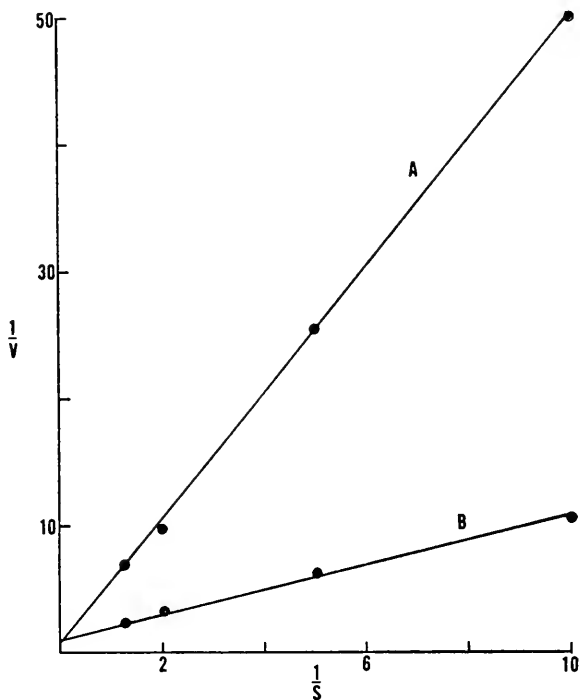


FIGURE 5. The effect of 4 mM propionate on the uptake of ^{14}C -acetate in one-minute incubations. A = inhibited slope; B = uninhibited slope; V = uptake, and S = mM acetate. Each point is a mean of 12 determinations.

range of 0.1–1.6 mM and the results are shown in Figure 4. In contrast to the results obtained at pH 7.4, the uptake of acetate at pH 6.2 was a linear function of acetate concentration.

Effects of other compounds

Effects on acetate uptake of the addition of a variety of compounds were examined. The results of these experiments are shown in Table II. Of the compounds tested, only short chain volatile fatty acids produced a significant effect on acetate uptake.

Acetate uptake at various concentrations in the presence or absence of the individual inhibitory fatty acids of Table II was examined by the Lineweaver-Burke method (1934). A typical double reciprocal plot, that of the effects of propionate on the uptake of acetate, is shown in Figure 5. The approximately common intercept of the slopes of both the inhibited and non-inhibited plots indicates that the inhibition was competitive in nature. Using the K_t and V_{max} values determined experimentally for acetate uptake, the inhibition constants (K_i) for the activity of other fatty acids were determined by application of the equation:

$$\text{Slope} = \frac{K_t}{V_{max}} \left(1 + \frac{[I]}{K_i} \right)$$

TABLE II

*Effects of various compounds on acetate uptake in 1 minute.
Acetate concentration, 0.1 mM; inhibitor concentration, 4.0 mM*

Addition	Uptake (m μ moles/g./min. \pm S.E.)	% Inhibition
none	105 \pm 6.6	—
2,4-dinitrophenol	102 \pm 8.4	0
phlorizin	104 \pm 9.2	0
ouabain	107 \pm 6.3	0
adenine	101 \pm 12.1	0
uracil	102 \pm 9.3	0
galactose	96 \pm 7.2	0
glucose	107 \pm 6.1	0
glycerol	96 \pm 4.5	0
alanine	119 \pm 11.0	0
phenylalanine	120 \pm 10.0	0
arginine	97 \pm 6.9	0
glutamic acid	93 \pm 7.5	0
leucine	104 \pm 6.9	0
lysine	110 \pm 10.7	0
betaine	103 \pm 8.9	0
sarcosine	121 \pm 9.0	0
lactate	91 \pm 14.9	0
succinate	124 \pm 5.9	0
formate	81 \pm 2.3	23
propionate	33 \pm 1.8	69
butyrate	53 \pm 2.9	50
valerate	63 \pm 2.9	40
octanoate	63 \pm 1.4	40

Values for the inhibitor constants of the fatty acids studied are: formate 6.9; propionate 1.25; butyrate 3.6; valerate 5.0; and octanoate 5.1.

To determine whether the inhibitions were fully or partially competitive, the uptake of acetate at a concentration of 0.1 mM was studied in the presence of inhibitor concentration of 1 to 10 mM; the data were then plotted after the method of Dixon (1953). Only formate and acetate were found to be fully competitive inhibitors of acetate uptake.

By an extension of the Michaelis-Menten treatment of enzyme kinetics, Read,

TABLE III

Observed and predicted effects of a mixture of fatty acids on 1-minute acetate uptake. Inhibitors present were sodium propionate (2 mM), sodium butyrate (2 mM), and sodium formate (4 mM)

Acetate concentration (mM)	Uptake $\mu\text{moles/g./min.} \pm \text{S.E.}$	
	Observed	Predicted
0.1	28 \pm 0.6	25
0.2	45 \pm 1.7	50
0.4	110 \pm 10.2	92
0.6	154 \pm 8.6	131

Rothman and Simmons (1963) derived an equation predicting the inhibitory effects of a mixture of amino acids on the uptake of a single amino acid. During the present investigation the effects of a mixture of fatty acids on the uptake of sodium acetate were investigated. Inhibitory effects of a mixture of fatty acids consisting of propionate (2 mM), butyrate (2 mM) and formate (4 mM) on the uptake of acetate at several concentrations are shown in Table III in which observed values are compared to those predicted from the equation:

$$V = \frac{V_{\max}}{\frac{K_t}{S} + 1 + \frac{(K_t)(I)}{(K_i)(S)} + \frac{(K_t)(I')}{(K_i')(S)} + \frac{(K_t)(I'')}{(K_i'')(S)}}$$

where V_{\max} , K_t , and S are values determined for acetate; K_i , K_i' , and K_i'' are independently determined inhibition constants for formate, propionate, and butyrate; and I , I' , and I'' are concentrations of the inhibitors.

Uptake of butyrate 1-¹⁴C

A limited number of observations were made concerning the relationship between butyrate uptake and concentration between 0.1 and 1.6 mM. Over this concentration range the amount of butyrate absorbed by the tapeworm did not increase

TABLE IV

Relative effects of various fatty acids as inhibitors of acetate and butyrate uptake in 1 minute

Inhibitor	Percentage inhibition	
	¹⁴ C-acetate	¹⁴ C-butyrate
Formate	22.9	8.2
Acetate	32.4	16.1
Propionate	69	40.1
Butyrate	50	—
Valerate	40	27.0
Octanoate	40	21.7

linearly with concentration increase and, by methods similar to those outlined above, the K_t and V_{max} values for the butyrate system were 1.5 and 2.5, respectively.

Short chain fatty acids were found to inhibit butyrate uptake and the relative effects of the various inhibitors on both butyrate and acetate uptake are shown in Table IV. The similarity of the relative activity of these several fatty acids as inhibitors of acetate or butyrate uptake suggests similar relative affinities of these inhibitors for the acetate and butyrate transport systems. The uptake of butyrate at several concentrations was determined in the presence and absence of acetate. A Lineweaver-Burke plot of the data so obtained showed the inhibition produced by acetate to be competitive in character and a K_i value of 5.4 for acetate as an inhibitor of butyrate uptake was calculated.

DISCUSSION

It is known that fatty acids enter a number of vertebrate tissues (Fredrickson and Gordon, 1958; Johnston, 1959; Isselbacher, 1965; Quastel, 1965; Hungate, 1966; and others). Quastel (1965) reported that brain tissue accumulated acetate. Transport of acetate, propionate, butyrate, valerate, and hexanoate against a concentration difference has been reported to occur in the small intestine of the rat (Smyth and Taylor, 1958; Barry and Smyth, 1960), although in 60-min. incubations, the concentration ratios reported by the latter workers were always less than 2.0 and the chemical methods used to evaluate the acids were not at all specific (titration after steam distillation).

There is little published information on the kinetics of fatty acid transport. The data of Barry and Smyth (1960) do not permit a conclusion as to whether there is a diffusion component in the absorption of fatty acids by the rat intestine. These workers assumed Michaelis-Menten kinetics and calculated a K_m of 45 mM. This may be a spurious value since the data do not show saturation kinetics and may represent a combination of mechanisms for acetate absorption.

The present experiments have shown that a significant proportion of acetate absorption by *Hymenolepis* occurs by a mediated process. At concentrations above 2 mM, diffusion appears to be a significant component of acetate uptake. In this dual mode of absorption, the uptake of acetate resembles the uptake of uracil by *Hymenolepis* (MacInnis *et al.*, 1965). Since acetate transport is not affected by a variety of organic compounds, other than fatty acids, it may be concluded that acetate transport in *H. diminuta* occurs through a mechanism with fatty acid specificity. The competitive inhibition of acetate uptake by other short chain fatty acids is further evidence for a specific mediated process and suggests that there is a common transport system for these compounds. Indeed, the similarity in the relative inhibition of acetate or butyrate uptake by the other fatty acids is strong evidence for common transport mechanisms for these two compounds, but more extensive experimentation is required to ascertain whether mediated transport of formate, propionate, valerate, heptanoate, and octanoate occurs only through the acetate-butyrate system.

The effects of pH on acetate uptake suggest that the undissociated form enters more rapidly by diffusion, perhaps because of higher solubility in membrane lipid. The data further suggest that mediated transport mainly involves the dissociated

form of acetate. The effect of pH on the propionate inhibition of acetate is also consistent with the above interpretations.

Ginger and Fairbairn (1966b) remarked that during a 2-hour period the rate of absorption of acetate by *H. diminuta* was linear. Estimation of the concentration of acetate present in Ginger and Fairbairn's experiment (using 2.5 μc . acetate, with specific activity of 2.05 $\mu\text{c}/\text{mM}$, in a volume of 10 ml.) indicates a substrate concentration of about 0.121 mM. At this concentration, the absorption of acetate during a 2-hour period should be non-linear with time. However, Ginger and Fairbairn's measurements were of the incorporation of acetate into lipid which may not be a measure of absorption. For example, the incorporation of ^{14}C -labeled amino acids into protein in *Hymenolepis* or *Calliobothrium* is linear with time whereas absorption follows first order kinetics (Harris and Read, in preparation; Fisher *et al.*, in preparation). Care must be taken to differentiate absorption from incorporation into other tissue components. It is apparent that in many cases the rate of absorption does not limit the rate of incorporation of a substance. Or, putting it in different terms, the concentrations required to saturate incorporation systems may be quite low when compared with concentrations required to saturate absorption systems. Similarly, the use of the term "uptake" may create confusion. In recent studies on *Mycoplasma*, Rottem and Razin (1967) reported that acetate "uptake" was decreased by butyrate and propionate. These authors speculated that the effect involved inhibition of acetokinase activity, but it is not possible to determine from the data available whether absorption mechanisms or intracellular enzyme reactions are involved. It would be helpful if the term "uptake" were restricted to denote the movement of solute from an extracellular to an intracellular location.

The competition between short chain fatty acids in their absorption by this worm should be considered in terms of (1) the probability that the quality and relative quantities of fatty acids in the small gut are relatively constant and, to some extent, independent of the fatty acids ingested by the host, and (2) the apparent inability of the worm to carry out *de novo* biosynthesis of higher fatty acids (Ginger and Fairbairn, 1966b; Jacobsen and Fairbairn, 1967).

Harrington (1965) showed that the components of *H. diminuta* lipids are quantitatively altered when the worm is reared in hamsters rather than rats. A clue to the basis of such alterations is found in the evidence that a considerable portion of the fatty acids in the lumen of the small intestine are derived from the host (Ginger and Fairbairn, 1966b).

When this is coupled with the finding that *H. diminuta* appears to be quite limited in its ability to synthesize fatty acids, the worms being limited to reactions resulting in chain lengthening (Ginger and Fairbairn, 1966; Jacobsen and Fairbairn, 1967), specific characteristics of fatty acids of host origin should be reflected in the lipid composition of the worm. Tentative acceptance of this interpretation leads to the conclusion that the transport of fatty acids into the tissues of the worm may be of considerable importance in determining the lipid composition and, in this context, the competitions between fatty acids observed in the present study assume more meaning.

Acetate has been reported to be an end-product of carbohydrate metabolism in *H. diminuta* and to be excreted into the external medium by the worm (Fair-

bairn *et al.*, 1961). It might be argued that acetate absorption is of no physiological significance since the worm produces the compound in considerable quantity. However, it has been shown that labeled acetate in the external medium enters the tissues and is incorporated into tissue lipids (Ginger and Fairbairn, 1966b; Jacobsen and Fairbairn, 1967). The latter authors remarked that the incorporation of ^{14}C -acetate occurred at a much lower rate than the incorporation of higher fatty acids. This might be attributable to dilution of the labeled acetate by the unlabeled acetate originating from carbohydrate metabolism. On the other hand, Ginger and Fairbairn (1966b) observed that only small amounts of ^{14}C previously incorporated into glycogen appeared in lipid during a prolonged incubation in which most of the glycogen was metabolized. This does not appear to be completely consistent with the view that significant portions of the acetate produced endogenously are incorporated into lipid. Further investigation of the relative significance of acetate from metabolic and external sources would be desirable. The present authors can offer no explanation for Daugherty's (1957) report that significant amounts of acetate do not enter the tissues of *H. diminuta* in 15-min. incubations at 37°C .

It is tacitly recognized that the absorption of fatty acids by the worm living in a host may involve important modifications not duplicated in the present experiments. There is evidence that, in the digestive tract, fatty acids, monoglycerides, and conjugated bile salts are present as mixed micelles (Hofmann and Borgström, 1962). This may be of great significance in the absorption of higher fatty acids which are relatively insoluble in water, even at soaps. Thus, while bile may serve as a source of fatty acids (Baxter, 1966), it also contributes bile salts which may be of significance in fatty acid absorption. An investigation of the effects of bile salts on absorption of fatty acids by *Hymenolepis* will be the subject of a subsequent portion of this study. The effects of pH on fatty acid uptake suggest that the antero-posterior pH gradient in the small intestine of the host may be significant in determining the rate of fatty acid uptake and the relative roles of mediated transport and diffusion.

Of a number tested, the only compounds producing competitive or partially competitive inhibition of acetate or butyrate uptake were other volatile fatty acids. This supports the view that the mediated transport of these fatty acids occurs through mechanisms showing some specificity. As a matter of fact, four types of mediated transport showing chemical group specificity are now known in *Hymenolepis diminuta*. Group A includes mechanisms for the mediated transport of amino acids (Read, Rothman and Simmons, 1963); Group B includes mechanisms for the mediated transport of monosaccharide sugars (Phifer, 1960a, b; Read, 1961); Group C includes mechanisms for the transport of purine and pyrimidine bases (MacInnis, Fisher and Read, 1965); and Group D, identified in the present work, includes mechanisms for transport of short chain fatty acids. At higher concentrations, considerable quantities of the compounds of Groups C and D enter the worm tissues by diffusion. Preliminary observations suggest that higher fatty acids, such as palmitic, may be absorbed by *H. diminuta* through mechanisms which are independent of those involved in the absorption of the short chain fatty acids. Transport of higher fatty acids is being examined in some detail.

The technical assistance of Miss Linda J. Rogers is gratefully acknowledged.

SUMMARY

1. At low substrate concentrations (below approximately 2 mM) *Hymenolepis diminuta* absorbs acetate by a mediated process; at high substrate concentrations the main mode of entry is diffusion.

2. Hydrogen ion concentration affects acetate uptake. At low pH the rate is increased and there is a larger diffusion component. Inhibition of acetate uptake by propionate is depressed at low pH.

3. Acetate uptake is inhibited by other short chain fatty acids but is unaffected by a variety of other compounds. The kinetics of the inhibitions were examined and inhibitions were found to be partially competitive in the case of propionate, butyrate, valerate and octanoate and fully competitive in the case of formate. Butyrate uptake is also inhibited by these compounds.

4. The data are discussed in terms of transport of fatty acids, lipid metabolism in cestodes and possible significance in the ecology of the organism.

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THE EFFECTS OF HYDROCORTISONE ON THE BLOOD OF TADPOLES AND FROGS, *RANA CATESBEIANA*

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In his 1957 monograph, Chester Jones stated, "Little information is available from the results of the injection of adrenocortical steroids, used in mammalian work, into Amphibia" (pp. 153-154). Since that time, various observational and experimental attacks on the anterior pituitary-adrenocortical system of the Amphibia have increased our understanding of the adrenal cortex, its role and its control in this group of vertebrates. Aldosterone and corticosterone have been identified as the major cortical steroids of the anurans (Carstensen, Burgers and Li, 1961; Hanke and Weber, 1964; Pesonen and Rapola, 1962; and Phillips and Bellamy, 1963), and recently, Piper and de Roos (1967) have found that both these corticoids have negative feedback influence on the pituitary's secretion of ACTH in *Rana catesbeiana*. Hydrocortisone, an obvious product of the adrenal cortex of mammals, seems to be of lesser significance in the amphibians, although it has been found in cortical extracts or in excretory products of representatives of several genera of this group (Chester Jones, Phillips and Holmes, 1959; Dale, 1962; and Pesonen and Rapola, 1962). Further, studies in our laboratory have established that responses to injections of this hormone do occur in *Rana pipiens* (Bennett and Alspaugh, 1964; and Bennett and Newell, 1965). Changes in the distribution of leucocytes, similar to those seen in mammals and reported originally by Dalton and Selye (1938), were found as early as 12 hours after and as late as 144 hours after the treatment of frogs with hydrocortisone acetate. Of what significance these reactions are in the normal economy of *R. pipiens* is not known. The possible stress imposed by dehydration did not effect lymphopenia and neutrophilia in these frogs as did the injection of hydrocortisone (unpublished observations).

Among the many other questions we have regarding the role of hydrocortisone in Amphibia are these: does this steroid evoke changes in the blood picture of species other than *R. pipiens*; does it have effects on the distribution of white blood cells in immature as well as mature anurans? These are the questions which this study attempted to answer.

MATERIALS AND METHODS

The tadpoles, *R. catesbeiana*, were obtained from a supplier in North Carolina in October, 1967. They were kept in battery jars of pond water at a temperature which ranged between 9.5° and 10.5° C. Fifty tadpoles were studied. The experimental animals, 25, were injected with 0.5 cc. of a 0.2% suspension of hydrocortisone acetate (Nutritional Biochemical Corporation) in distilled water, and the controls, 25, were injected with the same amount of distilled water. The injec-

tions were made into the musculature of the tail in all cases. Forty-eight hours after injection, blood was obtained from the tail; smears were stained with Wright's blood stain. Differential counts of 100 leucocytes were made for each animal.

Frogs, *R. catesbeiana*, were sent from a supplier in Wisconsin from October through December 1967. These animals were kept in individual battery jars containing about 2 cm. of tap water in a room in which the temperature varied between 9.5° and 10.5° C. The 24 experimental animals received 1 cc. of a 0.2% suspension of hydrocortisone acetate in distilled water while 24 controls were injected with 1 cc. of distilled water. The injections were made under the skin of the ventrolateral surface of the posterior trunk. Again, blood samples were withdrawn 48 hours after treatment. Smears were stained with Wright's stain, and differential counts of 100 white cells were made for each frog.

RESULTS AND DISCUSSION

The average per cent distributions of neutrophils and lymphocytes for the control and experimental animals are given in Table I. Also included in this table are

TABLE I

The average per cent distribution of lymphocytes and neutrophils and the Student's "t" values for the differences between counts of the two types of cells for control and experimental tadpoles and frogs

Animals	% Lymphocytes	"t"	% Neutrophils	"t"
Experimental tadpoles	30.08		49.82	
Control tadpoles	58.12	2.07	21.48	2.44
Experimental frogs	17.50		70.71	
Control frogs	57.46	3.10	33.46	2.23

the Student's "t" values for the differences between the two types of blood cells for the controls and experimentals. The difference between the number of lymphocytes in the control and experimental tadpoles was found to be significant ($P < 0.025$), as was the difference between the number of neutrophils in these two groups of animals ($P < 0.01$). The number of lymphocytes in the control frogs was much greater than in the experimentals, and again, the difference was significant ($P < 0.005$). The difference between the counts of neutrophils in these two groups of frogs was also significant ($P < 0.01$).

Thus, it is evident that both the immature and mature forms of *R. catesbeiana* responded to the hydrocortisone as do mature *R. pipiens*. The directions of the responses, *i.e.*, an increase in neutrophils and a decrease in lymphocytes, were also the same in the three groups of animals. However, the average intensity of the reactions varied. In the experimental tadpoles, the neutrophils increased 131% over those of the controls while the lymphocytes decreased 49%. For the experimental frogs, *R. catesbeiana*, the increase in neutrophils was 111%, and the decrease in lymphocytes was 69%. Forty-eight hours after the injection of the same amount of hydrocortisone as was used in the present study, the number of neutrophils was

95% greater and the number of lymphocytes was 41% lesser in mature, experimental *R. pipiens* than in their controls (Bennett and Newell, 1965).

The significant decreases in monocytes, eosinophils and basophils which were observed 72 and 144 hours after administration of comparable doses of hydrocortisone to *R. pipiens* (Bennett and Alspaugh, 1964) were not seen in the blood of *R. catesbeiana* or of adult *R. pipiens* 48 hours after injection of the steroid (Bennett and Newell, 1965). The distribution of these rarer leucocytes may change only after more than 48 hours of exposure to hydrocortisone. However, the distribution of leucocytes in normal frogs varies widely; thus valid generalizations about even the typical white blood cell picture are difficult to state (Schermer, 1967). The differences in intensity of neutrophilia and lymphopenia reported here, as well as the differences between the counts of monocytes, basophils and eosinophils of experimental *R. catesbeiana* and *R. pipiens* at 48 hours and those at 72 and 144 hours after treatment, are in reality only slight and are probably not of biological significance.

The increase in neutrophils and the decrease in lymphocytes in the two species of frogs are certainly significant, and are of interest, especially since they parallel those well known in mammals in which titers of hydrocortisone have been increased by physiological stress or injection. The review of Selye (1950) continues to provide us with details of these changes in mammals. The exact manner in which some of the adrenal steroids effect the responses observed in the blood continues to be problematical. Also not fully understood is the role of the adrenocortical steroids in the immature and/or metamorphosing anuran. Frieden and Naile (1955) reported that hydrocortisone accelerated metamorphosis of thyroxin-treated *R. pipiens* and *R. hechsheri*. The histochemical studies of Rapola (1963) on *Xenopus laevis* also implicate adrenal steroids in metamorphic phenomena. Our work proves only that immature *R. catesbeiana*, as well as mature frogs of this species, do have the system which causes the blood picture of the animals to respond to hydrocortisone as does mature *R. pipiens*. What the mechanism responsible for these changes is; whether it operates in response to intrinsic steroids; whether it is of biological value to these amphibians remain unknown.

SUMMARY AND CONCLUSIONS

1. Differential counts of the leucocytes of tadpoles and frogs, *R. catesbeiana*, were made 48 hours after the injection of hydrocortisone acetate (experimentals) or distilled water (controls).
2. Neutrophilia and lymphopenia were evident in both the immature and mature experimental animals but not in the controls.
3. These changes in the distribution of white blood cells resemble those known in mammals and in *R. pipiens*.

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COLONY GROWTH AND PATTERN IN THE TWO-TENTACLED HYDROID, *PROBOSCIDACTYLA FLAVICIRRATA*

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Proboscidactyla flavicirrata is a species of hydroid whose colony is intimately patterned around its commensal host, a sabellid worm. The hydroid colony covers the distal end of the leathery worm tube, and the polyps interact with the worm during feeding (Uchida and Okuda, 1941; Hand and Hendrickson, 1950). The colony pattern has been described by several workers (Hand, 1954; Brinckmann and Vannucci, 1965) but there is virtually nothing known about the colony development.

The observations presented in this paper indicate that the colony growth of *Proboscidactyla flavicirrata* is dynamic and approaches a steady condition. The colony expands as the worm elongates its tube distally. The colony degenerates in regions distant from the tip. The various axial regions of the stolon mat therefore represent successive ages of tissue. This growth pattern results largely from the unusual ability of the feeding polyps to move along the tube, keeping pace with tube elongation and generating a stolon system behind them as they advance.

METHODS AND MATERIALS

Colonies of *Proboscidactyla flavicirrata* (Brandt) were collected during the summer months from large sabellid tubes (over 5 mm. in diameter) growing on the town floats at Friday Harbor, Washington. During the winter months colonies were kindly collected by Dr. Robert Fernald. Observations on colony and polyp progressions were made on material kept in a running sea water table, with the polyps fed every two days on sabellid eggs.

Polyps were explanted on microscope slides by placing them, with a small portion of attached stolon, upright on the surface. In grafting experiments, polyp segments were isolated with a scalpel and strung together on glass fibers to heal for three hours.

The terms *distal* and *proximal* refer to portions of the worm tube corresponding to the worm's anterior and posterior directions, respectively; the open end of the worm tube is distal. The term *forward* is used in reference to gastrozoid orientation, forward being that direction in which the mouth and tentacles point.

RESULTS

Description of the colony

A *Proboscidactyla* colony is composed of a stolon network and three polyp types: gastrozoid, gonozoid (blastostyle), and dactylozoid. Polyps and stolons are patterned axially along the distal portion of a sabellid worm tube.

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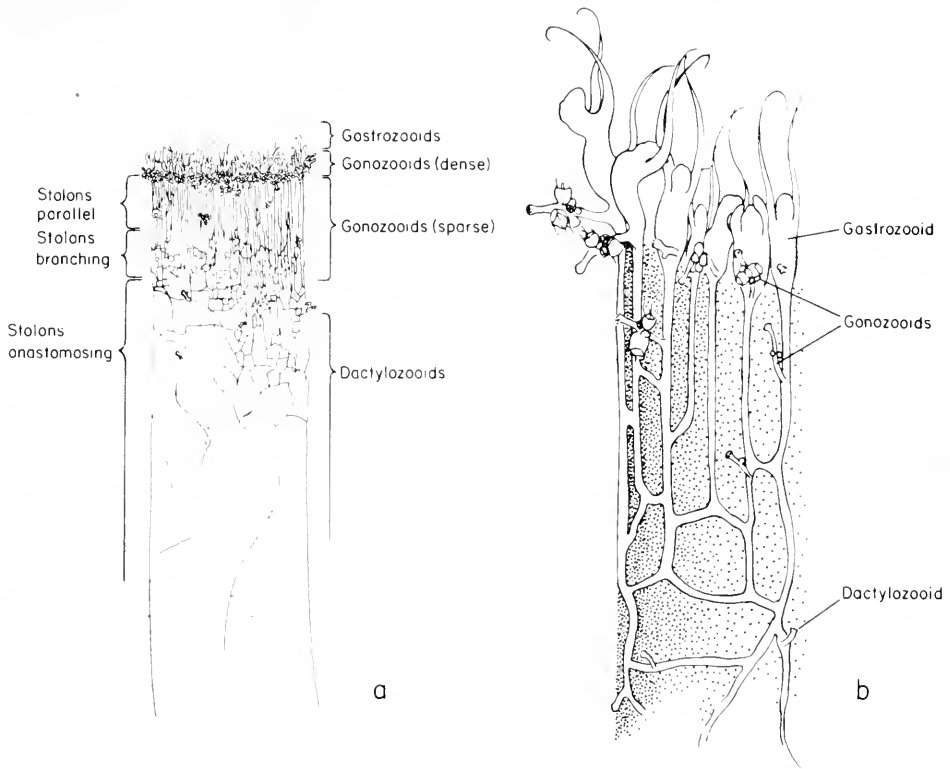


FIGURE 1. *Proboscidactyla flavicirrata* colony pattern. 1a: Axial colonial regions along worm tube. 1b: Polyp insertion and stolon pattern (longitudinal axis greatly shortened).

The gastrozooids are located almost exclusively in a whorl at the rim of the tube (Fig. 1). Gastrozooids are bilateral in structure (Figs. 1b, 3). The curved hypostome inserts vertically on the tip of the arching body column and points in the same direction as the body column arches (Fig. 3), so that the mouth is directed "forward." The two tentacles arising at the base of the hypostome are also directed forward, and a battery of nematocysts is located on the upper side of the hypostome. At the base of the polyp, protruding forward, is a small "foot" (Fig. 3) (see Campbell, 1968a), which may be homologous with the stolon tip of other hydroids. The single stolon connected to each gastrozooid inserts at the base of the polyp "back" (Fig. 3). The stolons (Fig. 1) leading back from the gastrozooids regularly run parallel to the tube axis and alongside adjacent stolons for several millimeters. Further from the tube tip, the stolons become thinner and show increasing branching, thus disrupting the strictly parallel arrangement. In the proximal portions they form an irregular network which becomes indiscernible farther back along the tube.

The gonozooid (Fig. 1), a long column capped by a nematocyst cluster, bears a bouquet of medusa buds midway along its length. Most gonozooids insert on the stolon immediately behind a gastrozooid or on a gastrozooid column. (Almost all

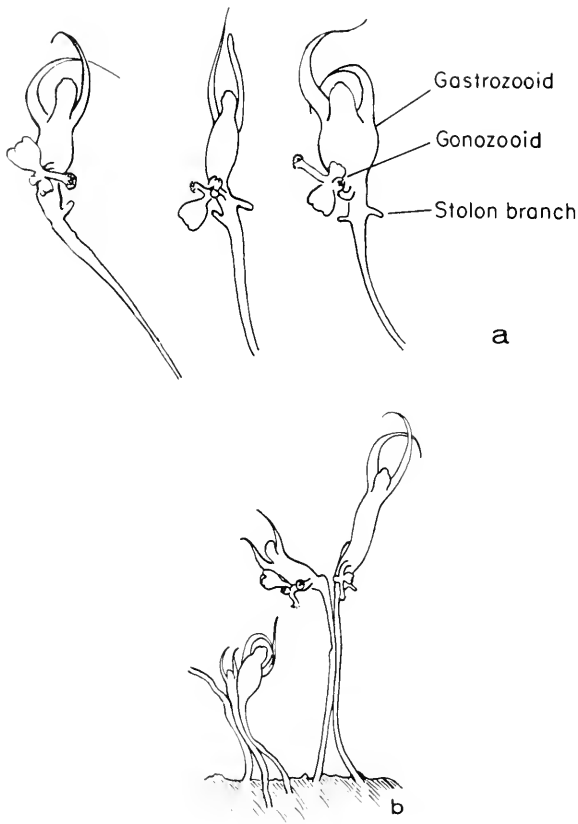


FIGURE 2. Gastrozooids, with gonozooids, migrating along glass tube, 24 hours after colony was stretched over the glass. Tracings from photographs. 2a: Three polyps which have stopped migrating, with stolon branches developing behind. These branches resemble those giving rise to new gastrozooids and to stolon anastomosis. 2b: Several polyps showing tendency to follow stolons. Shaded portion represents the edge of the worm tube.

large gastrozooids have such an associated gonozoid.) However, a few gonozooids are scattered sparsely over the distal half of the colony, inserted at random positions on the stolon network.

The dactylozooids (Hirai, 1960) are short, straight or slightly curved polyps, each crowned by a nematocyst bundle. They are scattered over the proximal region of the colony. Uchida and Okuda (1941) have termed such polyps as minute gonozooids lacking medusa buds. Hand (1954) termed them as young or abortive gonozooids.

Intermediate polyp forms may be found between the territories of gonozooids and dactylozooids. These forms are intermediate in length and often bear one or several minute or abortive medusa buds.

The *Proboscidactyla* colony organization is thus predominantly axiate along the tube, with several regions identifiable in terms of stolon patterns and polyp types.

*Observations on colony growth and expansion**A. Overall pattern*

Some colonies were observed for more than two weeks, during which time they grew in length, accommodating elongation of the worm tubes, but did not change in overall pattern. Furthermore, nearly every one of hundreds of colonies studied throughout the summer and at intervals through the winter showed similar organization regardless of the size of the colony and tube (except for colonies on very small tubes, as discussed below). Apparently the colony is in a steady-state developmental pattern, where continuing growth and expansion do not result in a qualitative change in pattern.



FIGURE 3. Gastrozoid, with gonozoid, migrating on a glass microscope slide. Photograph, taken from the side, was printed from a 16 mm. film. Gastrozoid hypostome is in upper right; two tentacles are at lower right. Arrow points to "foot." The stolon (broken white lines) runs off to the left.

B. Developmental cycle of polyps

Gastrozooids arise through modification of a stolon tip at the rim of the worm tube. This stolon forms as a lateral branch from one of the parallel stolons just behind the tube rim. It does not transform into a polyp until after it has extended slightly beyond the tube rim. At this time there is a slight flattening of the tip in a plane tangential to the tube surface. Two tentacle rudiments first become visible as broad lateral wings on this stolon tip. As these elongate, their positions shift towards the "front," where they eventually become spaced about 40° – 70° from one another. This shifting is probably due to the more rapid growth of the

polyp rudiment on the back surface, rather than to an actual movement of the tentacle rudiments across the column tissue. Tissue above the insertion narrows to form a hypostome. When the polyp rudiment has increased its size by several-fold, it has assumed the typical appearance of an older polyp.

Within a single colony, gastrozooids range from these small young individuals to polyps 4 mm. in length. In some colonies there is evidence of a graded distribution of average polyp size from one side of the tube to the other. No stages of gastrozoid degeneration were consistently found. Thus it appears that gastrozooids arise beside, or intercalated between, existing polyps through transformation of stolon tips, and that the polyps then have a relatively long life during which they grow continuously larger.

Gonozooids arise as small buds on the "backs" of the gastrozooids, about mid-way up the column. Gonozoid buds were found in no other position on the colony. Almost as soon as they are visible the buds are multilobed. As the stalk elongates, the terminal lobe becomes the nematocyst cap, and grows upwards on its own stalk. The other lobes remain in the midstalk region and become medusa buds. The stalk grows relatively rapidly until it is about the same length as the parent gastrozoid. It also changes its position on the gastrozoid, generally moving to the base or just behind the base of the mother polyp.

Gonozooids scattered behind the terminal polyp whorl have few or no medusa buds.

C. *Migratory behavior of gastrozooids*

Gastrozooids are capable of moving across a substratum in a forward direction. This can be demonstrated by extending the polyps' substratum beyond the tube rim, or by explanting polyps onto an appropriate surface under sea water. Under both conditions the gastrozooids will move forward at rates of up to 15 mm. per day.

A simple way to extend the polyps' substratum is to slip the tube over a glass rod. The elasticity of the tube holds the rim tightly against the glass. After 24 hours, many polyps in such colonies will have moved onto the glass (Figs. 2a, b). A single migrating polyp is shown in Figure 3.

The first visible change in a colony after it is stretched over glass is that the "foot" of each gastrozoid swells and elongates. This is apparent after a few hours. Several lines of evidence indicate that this foot directs the subsequent polyp migration. The zooids move forward although not necessarily in a straight line; when their paths curve the feet are also arched. Polyps tend to be guided by physical edges which only the foot may contact; thus, one polyp will follow the stolon of another (Fig. 2b). Finally, if a polyp is cut off from its base, rotated 90° or 180° and grafted back to the base, movement continues in the direction of stolon and foot orientation, not in the polyp's forward direction.

Gonozooids generally move along with the gastrozoid. But occasionally this close relationship is lost and a gonozoid remains isolated on the stolons while the gastrozooids continue advancing.

Mitotic figures (viewed in stained sections of several dozen migrating polyps and their stolons) are absent from the polyp base, but are abundant along the ectoderm and endoderm of stolons behind moving gastrozooids.

Unfed animals move up to several centimeters during the first 36 hours following isolation, but then stop. If fed, polyps migrate for a longer period, but never for more than three days. It is not clear whether food is limiting in these cases; while isolated gastrozooids do diminish greatly in size during movement, some gastrozooids in intact colonies do not.

After migration has stopped, lateral branches are frequently formed from the stolon just behind the gastrozooid (Fig. 2a).

The forward movement of gastrozooids described above is probably involved in the natural growth of the *Proboscidactyla* colony. When worms were grown for five weeks in filtered sea water, the new increments of tube added by the worm were transparent and thus easily distinguished from the previously existing opaque portions. The polyps continued to remain at the rim of the tube, with the entire whorl of gastrozooids eventually located on the transparent tube. Thus movement of polyps along the worm tube occurs during normal growth.

Occasionally, in natural colonies, individual gastrozooids are found away from the tube rim. These are directed in random orientations, their stolons indicating that they had previously been moving parallel to other polyps, but for some reason wandered from the tube rim.

Deduced colonial growth pattern

The observations outlined above suggest a growth pattern by which the entire colony remains in a steady-state condition. As the worm extends its tube at the distal rim, the gastrozooids advance forward, thereby maintaining their terminal positions. The individual polyps thus remain in a strategic ecological position and probably live for a relatively long time.

As gastrozooids advance, the stolon behind the polyps increases in length. This leads to the typical pattern of parallel stolons encircling the distal portion of the worm tube. The stolon system, therefore, is progressively older towards the proximal end of the tube. Apparently the stolons gradually form lateral branches which fuse, leading to progressive anastomosis at greater distances from the tube rim.

Some lateral branches are present in the distal portion. Generally these branches bend forward and extend toward the tube rim between the parent and adjacent stolon. These young stolons are the source of new gastrozooids, their tips being transformed directly into a polyp. Some control process must eventually limit the rate of new gastrozooid formation; possibly high stolon density inhibits lateral branching.

The gonozooids arise on the lower column of the gastrozooid. They remain attached to, or closely associated with, the mother polyp. Occasionally they fall back, are dissociated from the movements of the gastrozooid, and are left behind in the colony. Actually, they still probably continue moving forward to some extent because stolon growth (judging from mitotic figure distributions) occurs along the length of the unbranched stolon portions. These isolated gonozooids gradually lose their medusa buds and diminish in size, becoming dactylozooids.

This model of colonial growth is shown in Figure 4. The colony consists of an active front, which keeps pace with the advancing tube rim, and a stationary

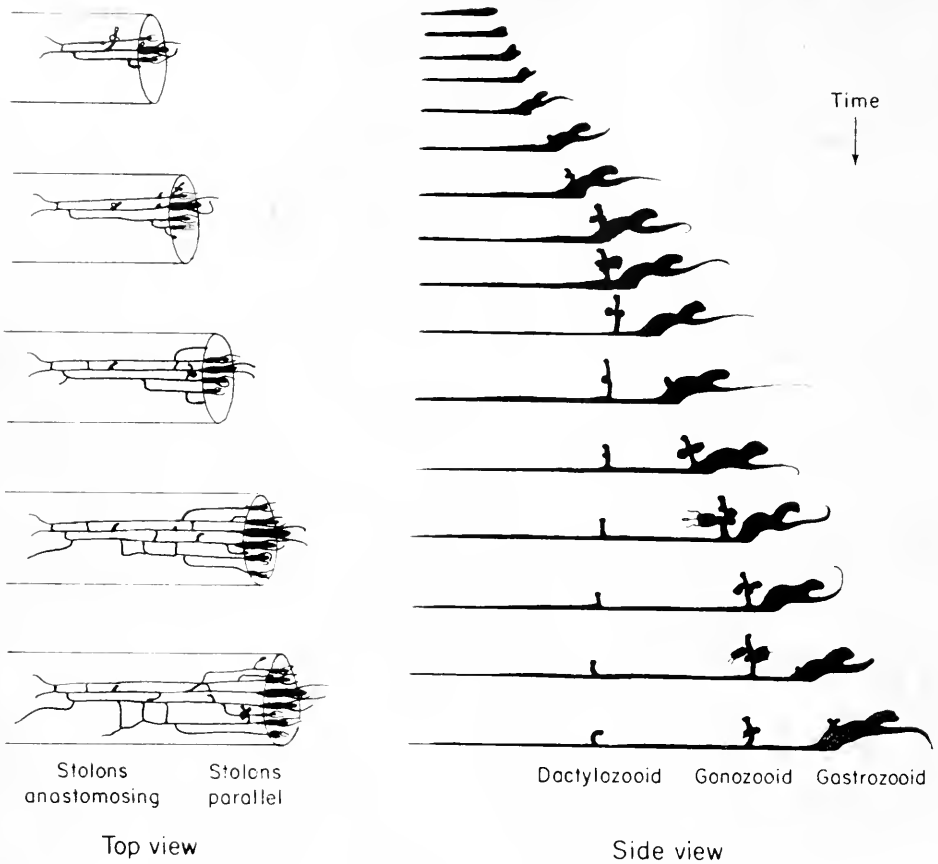


FIGURE 4. Deduced colony progression and development pattern (see text).

network of stolons covering the proximal portion of the worm tube. Individual gastrozooids represent at least relatively permanent components of the active front; they may or may not undergo a gradual replacement cycle. Gonozooids go through a more rapid replacement cycle; individuals occasionally stop advancing with the gastrozoid, stop forming medusa buds, and diminish in size. Each one lost in this manner is replaced by a new one budded from the column of the parent gastrozoid. The stolon system represents the trail of gastrozooids, slowly elaborated by anastomosing.

DISCUSSION

The unique *Proboscidactyla* colonial growth pattern results from the ability of the gastrozooids to move. Colonial succession in other hydroids which have been studied involves extension of the colony boundaries by stolon growth, followed by new polyps arising in the peripheral zones. Older colony regions may die and be

resorbed, giving rise to a net displacement of a colony unchanging in size (such as occurs in *Campanularia*; Crowell, 1953) or the older portions may remain active, in which case the colony expands in size (such as in *Podocoryne*; Braverman, 1964). In neither case is there any movement of any portion of the colony relative to the substratum, except for perhaps the terminal portions of the stolons. In *Proboscidactyla*, on the other hand, individual polyps migrate during colony translocation.

This migration pattern shows two points of similarity to the growth patterns of other hydroids. First, the movement of the gastrozooids appears to be similar and perhaps homologous to the elongation of the stolons in other hydroids. According to this view, the "foot" of the *Proboscidactyla* polyp represents the stolon tip; the gastrozooid would therefore be equivalent to a normal polyp which is situated just behind the stolon tip. Spots of vital dye applied near stolon tips in a number of hydroid genera (*Obelia*, Berrill, 1949; *Cordylophora*, Overton, 1963; *Clytia*, Hale, 1964) do move with the tip as does the *Proboscidactyla* gastrozooid (see also Campbell, 1966, 1968b).

Secondly, *Proboscidactyla* polyp movement resembles that of the upright growth in athecate hydroids bearing terminal polyps, where the individual polyp moves away from the colony center through upright elongation. The main difference between the two growth types is the relation of the polyp and stalk to the substratum.

Some descriptions of *Proboscidactyla* colonies do not indicate the presence of a terminal portion where stolons run parallel. These descriptions are of colonies on small worm tubes. I also observed small colonies to lack regular stolon patterns. Probably larger worm tubes elongate more rapidly than small ones. Since stolons do gradually anastomose, one would expect to find parallel cords of stolons during rapid elongation; at these times polyps can advance forward significant distances before stolon branching occurs. This consideration may explain the differences in colony regularity on large and small worm tubes, and the differences in stolon pattern and polyp distributions among related species (see Hand, 1954).

SUMMARY

1. The colony of the hydroid *Proboscidactyla* covers the terminal portions of a sabellid worm tube. The colony is axially patterned. In distal regions stolons run parallel to the worm tube axis while in proximal regions they form an anastomosing network. Gastrozooids are situated at the tube rim, gonozooids in a whorl behind gastrozooids, and dactylozooids in the proximal colony regions.

2. Gastrozooids will migrate actively forward when they are not at the edge of a substratum. The gonozooid associated with each gastrozooid generally moves with it.

3. The colony appears to be in a steady-state condition. The whorl of gastrozooids, and associated gonozooids, migrates forward as the tube rim advances due to secretion by the worm. The stolon system is generated behind the advancing gastrozooids. The proximal end of the colony progressively loses its regularity and definition. The axiate patterns of the colony at any moment thus trace out the history of the colony.

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RED CELL MORPHOMETRICS AND VOLUME IN THE CICHLID
TELEOST *TILAPIA MOSSAMBICA*, USING A
CHROMIUM-51 LABELING METHOD^{1, 2}

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Investigations of blood and blood cell volumes are limited in fishes, but a cross-section of interpretation of blood cell morphology is offered in the reports of Downey (1909), Jordan and Speidel (1930), Catton (1951) and Weinreb (1963). Blood volume studies have been hampered because the methods used to measure the mass of red cells were inadequate for application to fish. Exsanguination has been frequently attempted with limited success (Ronald, MacNab, Stewart and Beaton, 1964). Derrickson and Amberson (1934) used a perfusion technique on the dogfish, replacing blood with isotonic saline. Indirect dye dilution using Evans Blue was applied by Thorson (1961) on several marine and fresh-water species and by Smith and Bell (1964) on pink and sockeye salmon. Ronald *et al.* (1964) have applied fluorescein dye dilution on the Atlantic cod and Conte, Wagner and Harris (1963) used ⁵¹Cr, Evans Blue and human serum albumin-¹³¹I for blood studies on steelhead trout. The purposes of this investigation were to describe the formed elements and derive the red cell and total blood volumes of *Tilapia mossambica* and to determine the usefulness of ⁵¹Cr as a labeling material in this fish.

METHODS AND MATERIALS

The experimental species, *T. mossambica*, was collected from an estuary of the Enchanted Lakes located near Kailua on the windward side of Oahu, Hawaii. The size range was from 150 to 250 mm. (fork length) with weights ranging from 100 to 250 g. Females were of much greater relative abundance than males. The surface temperature of the estuary averaged 26° C. and the salinity varied from 0 to 25 ppt. Captured fish were held in 55-gallon fiberglass aquaria containing flowing fresh water with a steady temperature of 23.3° C.

Experimental animals were anesthetized with MS 222 (Tricaine methanesulfonate) in a 1:10,000 weight-to-volume solution to a deep stage III level, which is characterized by an absence of swimming movements, respiratory activity or response to external stimulation (Klontz, 1964). They were then placed in a restraining device to irrigate the gills with the MS 222 solution using a pump recycling system. The equipment was similar to that used by Smith and Bell (1964) at the Nanaimo, British Columbia, Station of the Fisheries Research

¹ Part of a thesis submitted to the graduate faculty of the University of Hawaii in partial fulfillment of the requirements for the degree of Master of Science.

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Board of Canada. Because of the unreliability of heart puncture and injection, a technique for cannulation of the ventral aorta was utilized using a variation of a dorsal aorta cannulation employed by Schiffman (1959) and Smith and Bell (1964). The cannula was polyethylene tubing with an inside diameter of 0.025 inch. The tubing was placed over the cut end of a 22-gauge regular bevel needle and filled with a solution of 1% NaCl, 1.2% dextrose and 220 units of heparin per milliliter. The needle was inserted into the skin 4 or 5 mm. to either side of the median sagittal axis of the fish and directly above the ventral aorta. A suitable vessel puncture site was easily selected as the needle and vessel were visible through the membrane of the isthmus. As an attempt to maintain the fish in a constant but an admittedly abnormal environment, the animal was completely immersed in a small blackened aquarium which was supplied with flowing fresh water.

While no tissue erosion was noted around the sutures or puncture sites, hemorrhages developed at the puncture site one to two days after cannulation and after three days, successful sampling could no longer be carried out. It was possible to remove 30 to 50% of the total volume of blood in *Tilapia* without aspiration and no mortalities occurred following repeated injection and blood removal, but it often took the fish 3 or 4 hours to recover to pre-anesthesia levels of activity and appearance.

Smears of fresh, unheparinized blood were used for descriptive study. Smears were fixed in absolute methanol and flooded with Giemsa stain buffered with Sorensen's phosphate buffer to pH 6.8. Red cell dimensions of fixed cells were measured by use of an eyepiece micrometer. Thickness measurements were obtained from living cells in suspension. No differential white cell counts were made. Hematocrits were determined using the micromethod of Smieszko (1960). They were taken from all ventral aorta cannulated fish, serving as controls for blood volume values and were also taken in some fish by severing the tail and exposing the caudal vessels. In each microhematocrit determination three tubes were filled consecutively.

Chromium-51 was supplied as $\text{Na}_2^{51}\text{CrO}_4$ with a specific activity of 20 mc. per milligram. The ^{51}Cr activity of liquid samples, in microhematocrit capillary tubes calibrated to a volume of 0.04 ml., was measured with a crystal scintillation detector. At least two five-minute counts were made on each sample. For labeling, 10 $\mu\text{c.}$ of ^{51}Cr were added to each milliliter of heparinized whole blood in a silicon-coated vial. After this mixture was incubated at room temperature (24°C.) for 75 to 90 minutes, with shaking every 5 minutes, 4 mg. per 10 $\mu\text{c.}$ ^{51}Cr of ascorbic acid were added to reduce free ^{51}Cr . The optimum incubation time was determined from the time labeling activity reached a maximum. At 1 to 10, 20, 30, 45, 50, 60, 75, and 90 minutes after adding ^{51}Cr to a vial of unlabeled blood, 0.01-ml. subsamples of blood were withdrawn and the red cell ^{51}Cr activity was measured. The labeling was inhibited by the addition of an aqueous solution of 0.5 mg. ascorbic acid or by washing the subsample with saline to remove free ^{51}Cr .

The total volume of red cells was calculated from a dilution formula described by Squibb & Co. (1959). About 0.5 to 0.9 ml. of ^{51}Cr -labeled whole blood was injected into the cannula of the donor fish with 0.1 ml. retained as a

TABLE I
The formed elements of T. mossambica blood in circulation

Cell type	Dimensions (microns)					Cell:mc. ratios (by length)
	Whole cell			Nucleus		
	Length	Width	Thickness	Length	Width	
Small lymphocyte	7.5*	—	—	6.3*	—	1.2
Large lymphocyte	12.4*	—	—	5.8*	—	2.1
Macrophage	12.9*	—	—	7.9*	—	1.6
Thrombocyte	9.1	3.7	—	7.0	3.3	1.3
Erythrocyte**	11 ± 0.7	7.5 ± 0.1	3.0	5.0	3.4	2.2

* Recorded as the greatest dimension through the central axis.

** Numbers per mm.³ = 1,549,000 ± 73,000 SD.

reference standard. Serial sampling up to 24 hours post-injection was accomplished by filling capillary tubes directly from the tip of the cannula. Total blood volume was found by dividing the total sample volume of red cells by the respective sample hematocrit and computing an average for all samples.

RESULTS AND DISCUSSION

For comparative purposes the morphological categories of *Tilapia* blood cells were similar to those described by Catton (1951) and Klontz *et al.* (1964).

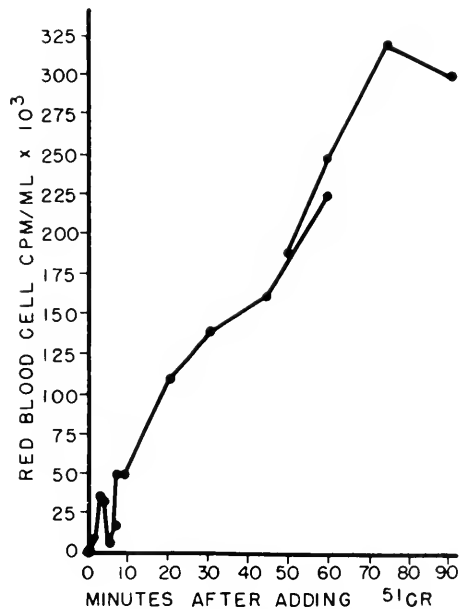


FIGURE 1. Relationship of *in vitro* ⁵¹Cr accumulation by *Tilapia mossambica* RBCs as a function of incubation time.

except cells of the granulocytic series were absent in both living and stained preparations. Morphometric and enumeration data are presented in Table I. The average hematocrits were $25.8 \pm 4.6\%$ ($N = 29$) for caudal vessel samples and $23.0 \pm 3.8\%$ ($N = 9$) for ventral aorta samples obtained prior to blood volume determinations. The nearly 3% difference between the two mean hematocrits was not significant (analysis of variance test; $F = 2.81$, d.f. 1, 36), and probably was a reflection of differences in sampling. Sample hematocrits from ventral aorta cannulations, used for total blood volume determinations, presented a lower average ($20.5 \pm 4.1\%$, $N = 7$) than presample hematocrits. This was a result of the decrease in sample size and not of blood loss. There was also a tendency for hematocrits (caudal vessel samples) to decrease with increasing fish weights. For a wide weight range, this may be correlated with activity or oxygen consumption.

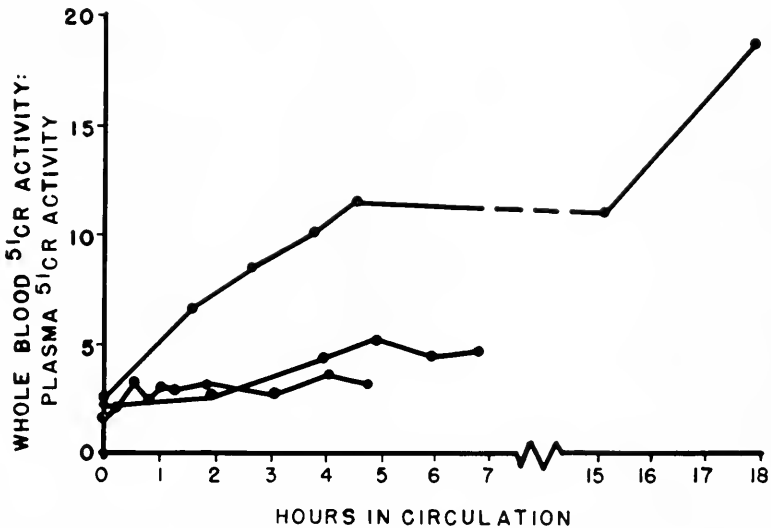


FIGURE 2. Whole blood ^{51}Cr activity:plasma ^{51}Cr activity ratios in three *Tilapia* versus the time of circulation.

The ^{51}Cr uptake curve (Fig. 1) is nearly linear with activity maximizing at 75 to 90 minutes after the addition of ^{51}Cr , although uptake is still rapid at 75 minutes. The dose-dependent effects or possible toxic levels of the ^{51}Cr were not tested. Plasma or unlabeled ^{51}Cr activity levels in *Tilapia* decreased to near background levels 4 to 19 hours after ^{51}Cr infusion in the circulatory system. Initially a large quantity of the radioactivity present in the injection medium does not appear in the red cells. There is a steady decline of plasma activity levels after injection, however, indicating that plasma ^{51}Cr is being rapidly removed from circulation and stored or excreted or both. In all fish samples (serial and single samples from 8 fish) the whole blood activity:plasma activity ratios in the injection medium range from 1.5 to 3.0. After 15 to 19 hours of circulation the values range from 10.2 to 18.8 (Fig. 2). Serial sampling indicates a period

of rapid mixing of injected labeled red cells in circulation during the first 10 to 60 minutes, followed by a long period of slow diffusion where fairly constant total red cell volumes or red cell ^{51}Cr activity levels were obtained. Nevertheless, the picture of dilution and circulation in the restrained or confined fish may not be an actual representation of what occurs while it is free-swimming. Since body movements probably aid peripheral circulation, the weak contractions of the heart while the fish is under anesthesia may supply only a limited flow of blood to body elements other than the head, gills, and viscera.

The mean value for the total blood volume was 3.17 ± 0.48 SD ml./100 g. body weight (range = 2.71 to 4.50). The mean value for the total volume of red cells was 0.65 ± 0.14 SD ml./100 g. body weight (range = 0.43 to 0.84, $N = 7$). A degree of correlation was found among blood volume against weight in seven fish which had moderately constant circulation times (Fig. 3). While the relationship was not significant at the 5% level, there was a definite inverse trend of volume on weight within the weight range tested. This indicates that blood volume in *Tilapia* is not directly proportional with weight as was found in the cod

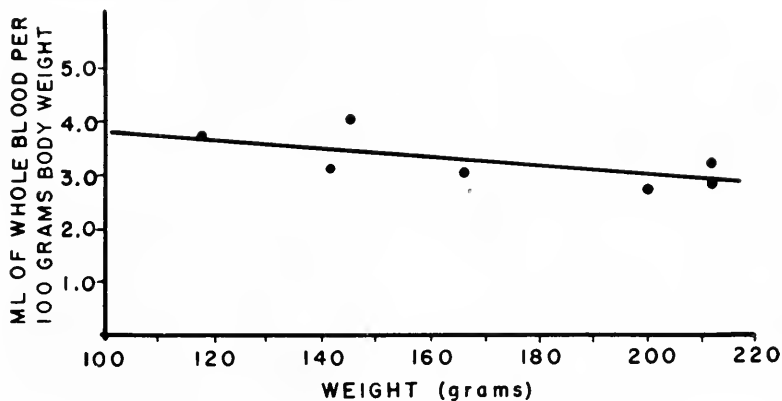


FIGURE 3. Relation of whole blood volume as a percentage of weight (ml. blood/100 g.) on total body weight in *Tilapia mossambica*. Circulation times after the injection of ^{51}Cr labeled RBCs were two to four hours.

(Ronald *et al.*, 1964) but decreases slightly with increasing weight. The total blood volume of *Tilapia* is consistent with those determinations made for other teleosts (Thorson, 1961; Conte *et al.*, 1963; Ronald *et al.*, 1964).

SUMMARY

1. Erythrocytes, small and large lymphocytes, macrophages and thrombocytes were identified. No granulocytes were found in stained or living preparations. Hematocrits averaged $25.8 \pm 4.6\%$ for caudal vessel samples and $23.0 \pm 3.8\%$ for ventral aorta samples. Red cell counts were $1,549,000 \pm 73,000$ SD per mm^3 . Red cells were $11 \pm 0.7 \mu$ long, $7.5 \pm 0.1 \mu$ wide and 3μ thick.

2. A technique is described for cannulation of the ventral aorta to allow repetitive sampling or injection while the fish is restrained or confined.

3. The labeling of erythrocytes with ^{51}Cr followed a linear curve with maximum cell incorporation occurring at approximately 90 minutes post-labeling.

4. A long period of slow mixing was noted for injected ^{51}Cr -RBCs after an initial dilution phase of 10 to 60 minutes. The total average total volume of red cells obtained for *Tilapia* as determined by ^{51}Cr dilution was 0.65 ± 0.14 ml./100 g. body weight and the mean total blood volume was 3.17 ± 0.48 ml./100 g. body weight. There was an allometric relationship suggested between TBV as a percentage of body weight and weight. Higher proportional volumes were associated with lighter animals.

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THE LARVAL DEVELOPMENT OF THE COMMENSAL
CRAB POLYONYX GIBBESI Haig, 1956
(CRUSTACEA: DECAPODA)^{1, 2}

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The larvae of the anomuran crab family Porcellanidae have long been recognized in the plankton by their unique rostral spine (Thompson, 1836). Until recently, however, there was little attempt to rear the larvae in the laboratory under controlled conditions through their complete life cycle. To date, adequate descriptions of the complete larval development are available for species of *Pachycheles* (Kurata, 1964; Knight, 1966; Boschi *et al.*, 1967), *Petrocheles* (Wear, 1965a, 1966), *Petrolisthes* (Gohar and Al Kholi, 1957; Wear, 1964 a, b, 1965b; Greenwood, 1965) *Pisidia* (formerly some *Porcellana* species; Lebour, 1943; Bourdillon-Casanova, 1956), *Porcellana* (Lebour, 1943; Bourdillon-Casanova, 1960) and *Polyonyx* (Knight, 1966).

Polyonyx gibbesi, which is found from Woods Hole, Massachusetts to La Paloma, Uruguay, is a known commensal with the polychaete worm *Chaetopterus variopedatus* (Renier) (see Pearse, 1913; Gray, 1961 and Haig, 1966). Faxon (1879, 1882) briefly described and figured the first and second zoea and the "first stage of the crab" (= megalopa) of *Polyonyx macrocheles* Gibbes (now called *Polyonyx gibbesi* Haig 1956). Faxon's figures and descriptions are unfortunately not sufficiently detailed to allow positive identification of the larvae or comparison with larvae of other species of *Polyonyx*. A second author (MacArthur, 1962, unpublished) also described the larval development of *Polyonyx macrocheles* (sic) but differences between the larvae she studied and mine will require further study before an evaluation can be made.

The purpose of this paper is to illustrate and describe the complete larval development of *Polyonyx gibbesi*. Certain characters are discussed which may enable the larvae of the genus to be identified from the plankton. The relationship between *P. gibbesi* and the Pacific coast form *Polyonyx quadriungulatus* is also discussed.

MATERIALS AND METHODS

Ovigerous female *Polyonyx* were collected from *Chaetopterus* tubes obtained by bucket dredge in the Cape Florida Channel, from an area 3 m. deep and

¹ Contribution No. 939 from the Institute of Marine Sciences, University of Miami, Florida.

² Part of a thesis presented to the Graduate Council of the University of Miami in partial fulfillment of the requirements for the degree of Master of Science. This study was supported in part by Research Grant GM-11244 from the National Institutes of Health, U. S. Public Health Service and a National Science Foundation Institutional Grant through the University of Miami.

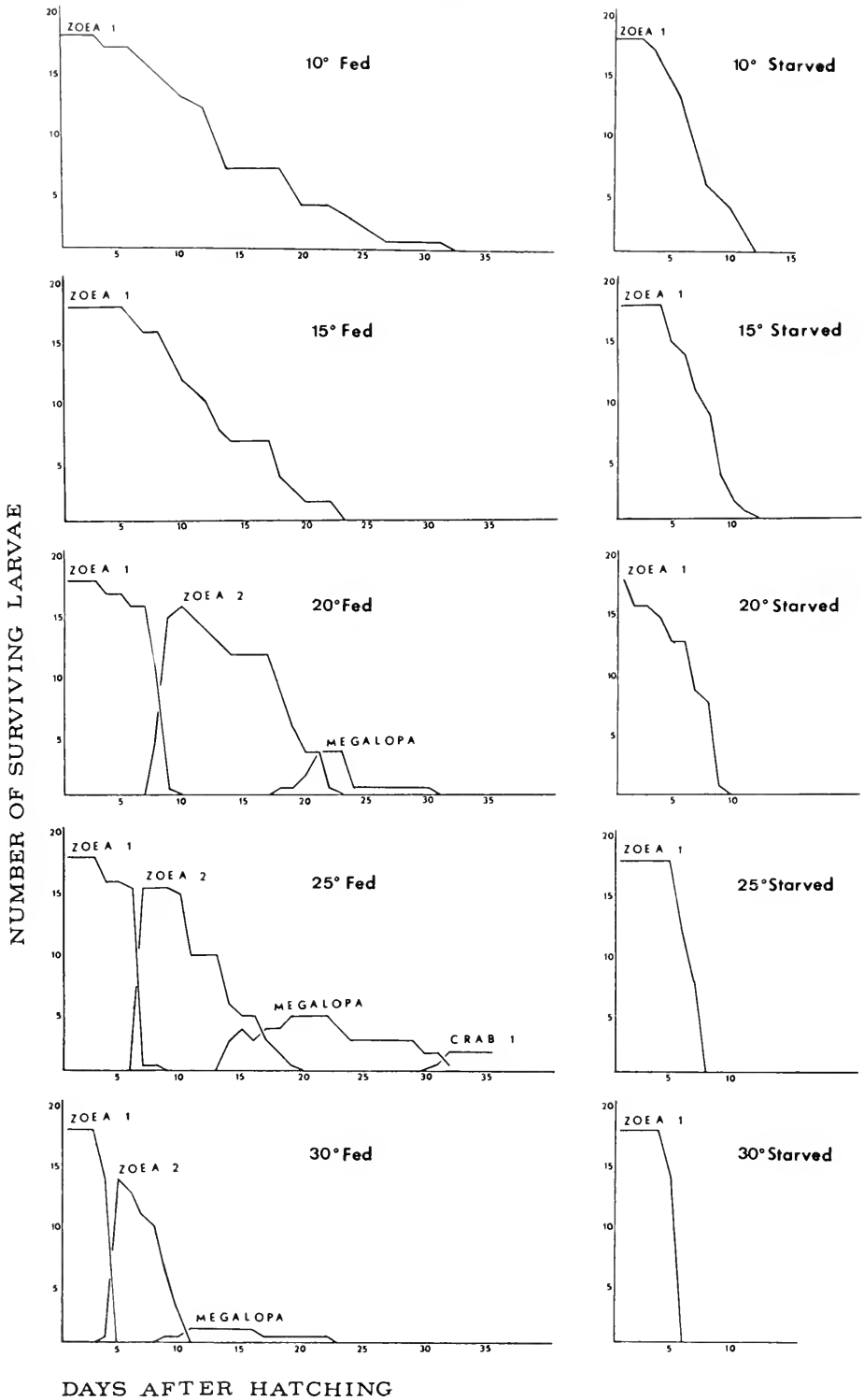


FIGURE 1.

20 m. offshore of Hurricane Harbor, Key Biscayne, Florida. The crabs were isolated in non-flowing sea water in 19-cm. diameter plastic bowls until hatching occurred. Zoeae were placed, one each, in compartmented plastic tackle boxes. Each compartment held about 40 cc. of unfiltered sea water (33–35‰ salinity). Two series of boxes, one with zoeae fed with *Artemia salina* nauplii, and one with starved zoeae, were maintained at 10°, 15°, 20°, 25°, and 30° C. Temperature variation was not more than $\pm 1.5^\circ$ C. Larvae were fed and received a change of water every other day. Illumination was not controlled. A record was kept for all molts and deaths for each larva, with exuviae and dead larvae preserved in 70% ethanol. Larvae obtained from another female were held at 25° C. ($\pm 2^\circ$ C.) and several zoeae were sacrificed every day, and also preserved in 70% ethanol. Appendages were dissected in 40% lactic acid and mounted in Turtox CMC-S. Illustrations were made with a camera lucida attached to a Wild M-20 binocular compound microscope, from slides of individual appendages. Drawings were checked both for accuracy and individual larval variation against appendages dissected from exuviae and sacrificed animals. Measurements were made with a LaFayette objective micrometer. Carapace length was measured from the anterior margin of the zoeal eye to the insertion of the posterior spines on the zoeal carapace, and from the edge of the megalopal frontal region to the posterior margin of the carapace for carapace length, and across the widest part of the carapace for carapace width. The carapace measurements provided are the arithmetic average of 10 specimens measured in each larval stage. One spent female (UMML 32:3581) plus hatched specimens of first and second zoeae and megalopae (UMML 32:3582) were placed in the Museum of the Institute of Marine Sciences.

RESULTS

Polyonyx gibbsii passes through a pre-zoeal stage, two zoeal stages and one megalopal stage. As noted for other porcellanids (*c.g.* Knight, 1966) there is an increase in size of the third maxillipeds, gills and pleopods during the zoeal stages. Some authors (*c.g.* Boschi *et al.*, 1967) consider this increase a substage, though no molt is seen. Others (*c.g.* Boyd and Johnson, 1963) refer to molting that produces additional stages but little or no alteration of form as substages. The term substage needs re-definition. I never observed exuviae other than the two zoeal molts, and I do not use the term substage.

Temperature noticeably affects larval development, altering the duration of each stage or preventing the development to subsequent stages. Figure 1 depicts the length of time the larvae, both fed and starved, spent in each stage at various temperatures. No starved zoeae developed beyond the first stage. Fed zoeae at 10° and 15° C. remained in Stage I, living at 10° C. for a maximum of 32 days, and at 15° C. for a maximum of 22 days, before dying. Animals reared at 20°, 25°, and 30° C. reached the megalopal stage in a minimum of 18, 14 and eight days, respectively. Crab stages were obtained only at 25° C. While the

FIGURE 1. *Polyonyx gibbsii*: Duration of survival in each stage of larval development, fed and starved, at various temperatures. Horizontal scale represents the number of days after hatching. The vertical scale represents the number of surviving larvae. Salinity range = 33–35‰.

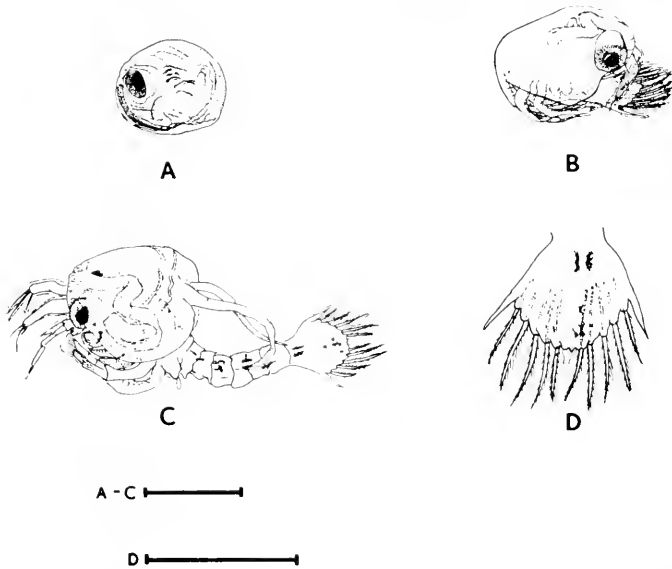


FIGURE 2. *Polyonyx gibbsi*: pre-zoea. A. Larva in egg just before hatching. B. Larva in process of hatching. C. Newly hatched pre-zoea. Remnants of pre-zoeal cuticle surround appendages. Rostral spine and setae on appendages not yet emerged. D. Pre-zoeal telson. Two hairs on central prominence of telson and other fine setae not yet present. Scales equal 0.5 mm.

shortest duration of development occurred at 30° C., no crab stages were obtained. Mortality at this temperature was very high and only three zoeae molted to megalopa and none of these survived longer than 11 days. Thus, 25° C. allowed the best development in the temperature series. At this temperature the first and second zoeal stages lasted six to seven days and the megalopal stage usually lasted 12 to 14 days.

DESCRIPTION OF THE LARVAE

Pre-zoea

The hatching sequence of the entire egg mass lasted about two hours (see Fig. 2, A, B). The pre-zoeal stage also lasts about two hours. The long rostral spine, in the pre-zoea as yet undeveloped, is partially bent outside and under the carapace and partially invaginated into the carapace above the midgut region (Fig. 2, C). The embryonic cuticle in the specimens examined was almost completely fragmented. Setae on the appendages and telson, only partly extruded at the beginning of the stage, become completely extruded toward the end of the stage. The setae on the carapace over the eyes and setae on the central prominence of the telson (Fig. 2, D) are not present. The pre-zoeae swim by rapid abdominal flexion.

In two instances pre-zoeae swam from *Chaetopterus* tubes maintained in the laboratory. If the larvae did not hatch as pre-zoeae they might encounter difficulty in escaping through the narrow neck of the worm tube.

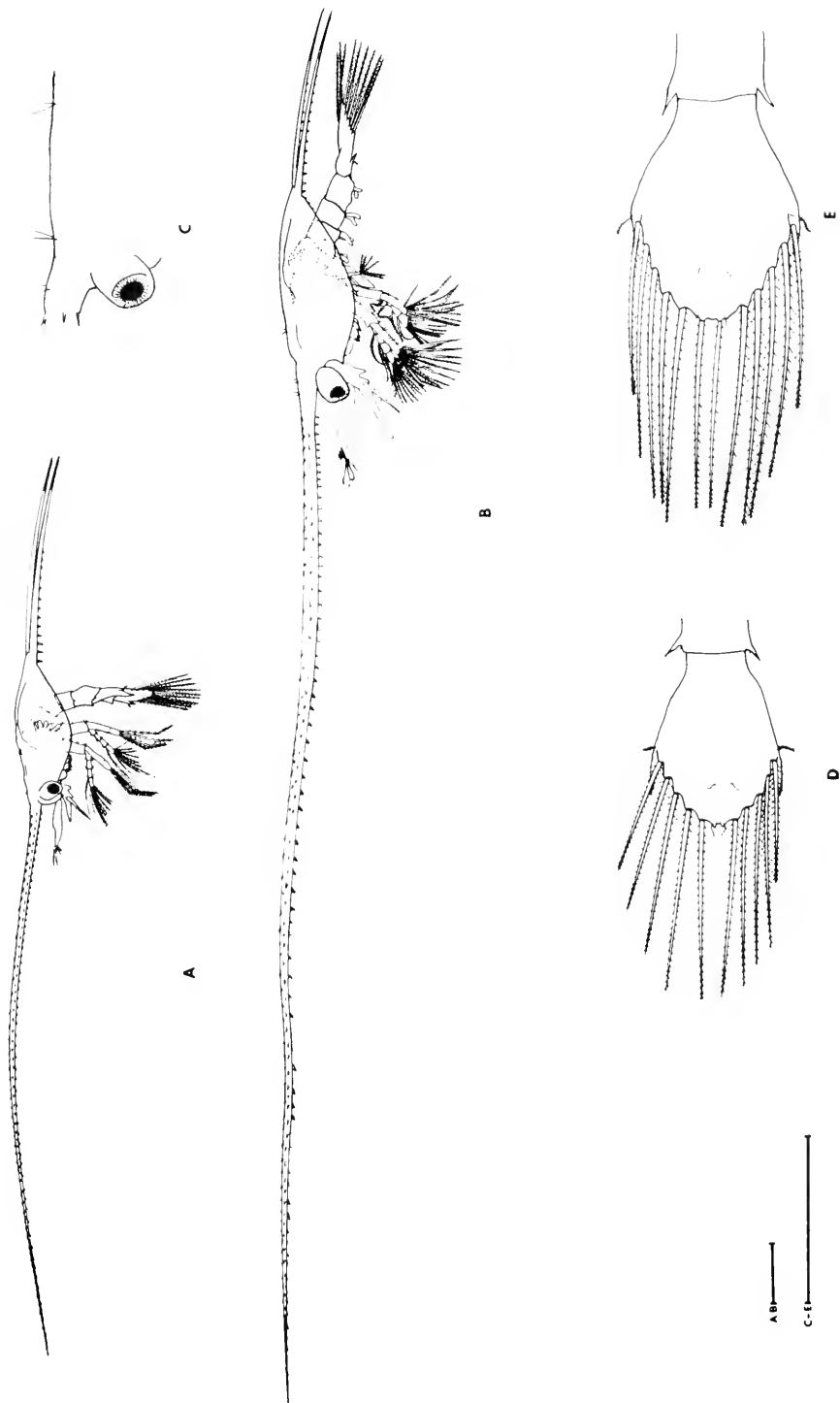


FIGURE 3. *Polyonyx gibbesi*: first and second zoeal stages. A, First zoea. B, Second zoea. C, Caparapace of first zoea showing dorsal setae. D, Telson of first zoea. E, Telson of second zoea. Scales equal 0.5 mm.

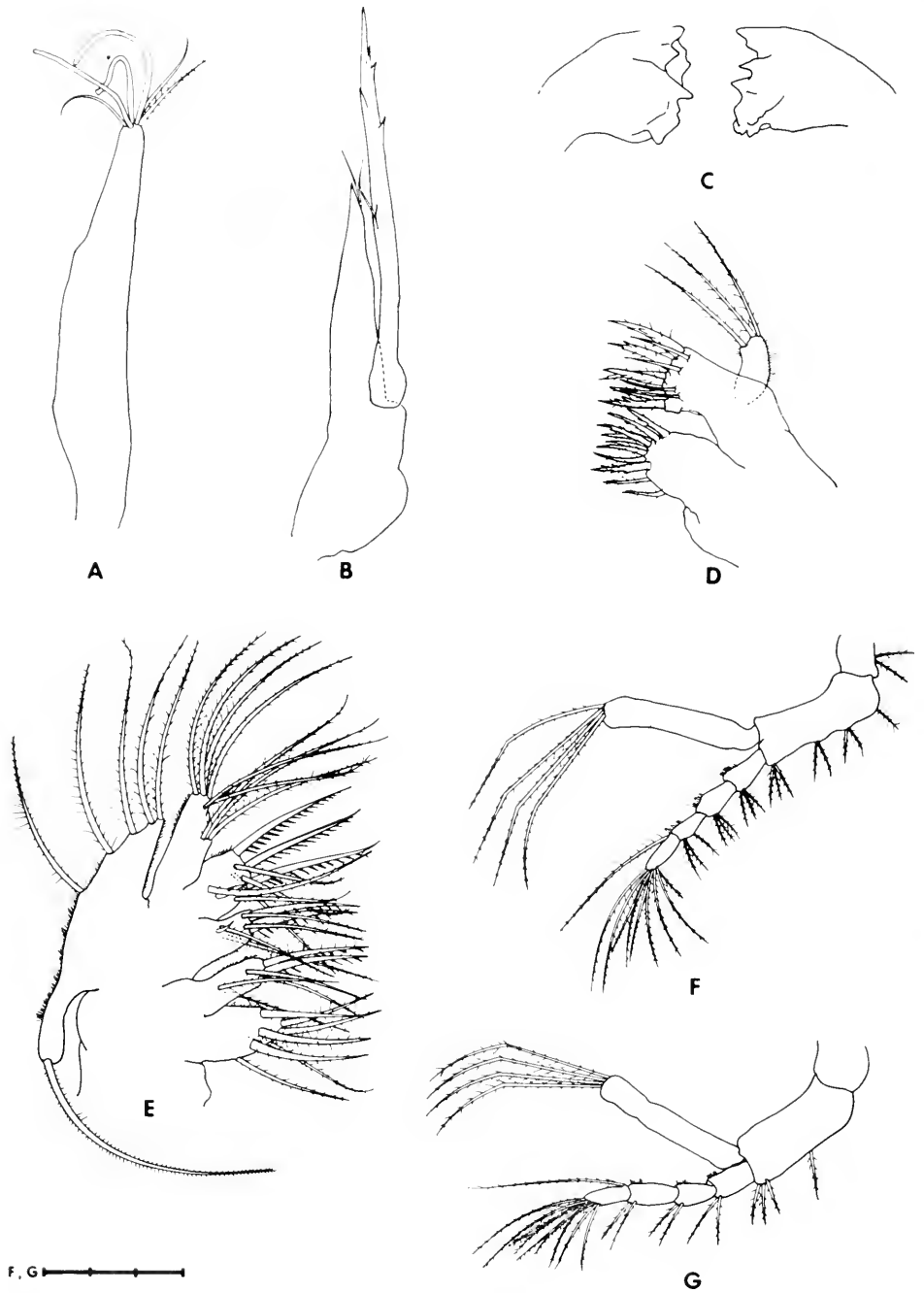


FIGURE 4. *Polyonyx gibbesi*; first zoeal appendages. A. Antennule. B. Antenna. C. Mandibles. D. Maxillule. E. Maxilla. F. Maxilliped 1. G. Maxilliped 2. Scales equal 0.3 mm.

First Zoca

Carapace length: 1.2 mm.

Number of specimens examined: 20

Carapace (Fig. 3, A). Typically porcellanid, produced anteriorly into an extremely long rostral spine up to seven times the length of carapace proper; posteriorly into two straight or divergent posterior carapace spines 1.4 to 1.8 times as long as carapace. Armature of spines as illustrated. Curvature of rostral and posterior carapace spines variable, depending in part on frequency of collision by new zoeae with other objects.

Dorsal surface of carapace with three pairs of fine setae. Placement illustrated in Figure 3, C.

Antennule: (Fig. 4, A). A simple slightly flabelliform rod with two long and one short aesthetasc and two or three setae of variable length, one with fine setules.

Antennae: (Fig. 4, B). Endopodite, fused to protopodite, has a thin subterminal seta. Exopodite a thin spine almost twice as long as endopodite; about six small spinelets distally plus a thin seta halfway down its length.

Mandibles: (Fig. 4, C). Asymmetrical dentate processes without palp.

Maxillule: (Fig. 4, D). Endopodite unsegmented, 3 setae. Coxal and basal endites each with 10 processes as shown.

Maxilla: (Fig. 4, E). Endopodite unsegmented, 4 terminal, two subterminal and three medial setae. Coxal endite with seven processes on proximal lobe, six on distal lobe. Basal endite with seven processes on proximal lobe, nine on distal lobe. Scaphognathite with five setae laterally and one long apical plumose seta. Placement of all setae as illustrated.

Maxilliped 1: (Fig. 4, F). Four and ten setae on terminal segments of exopodite and endopodite, respectively. Basipodite setation progressing distally is 1, 2-3, 2, 3. Small tufts of hair dorsally on endopodite segments one to three.

Maxilliped 2: (Fig. 4, G). Setation on terminal segments similar to maxilliped 1. Basipodite setation 1-2, 3 progressing distally. Tufts of hair dorsally on segments one to three of endopodite as in maxilliped 1.

Maxilliped 3: (Fig. 3, A). A small bifid lobe, with one or two setae.

Percipods: Five buds visible in most specimens; extremely small and distorted in early stage. Both buds and maxilliped 3 increase in size as zoea progresses to Stage II.

Abdomen: (Fig. 3, A). Five somites, each with lateral spine of increasing length on somites nearer the telson. Somites three and four may each have a fine hair dorso-laterally; somite five has two fine setae just above lateral spine on each side.

Pleopods: Absent. Primordia visible in some zoeae from the plankton.

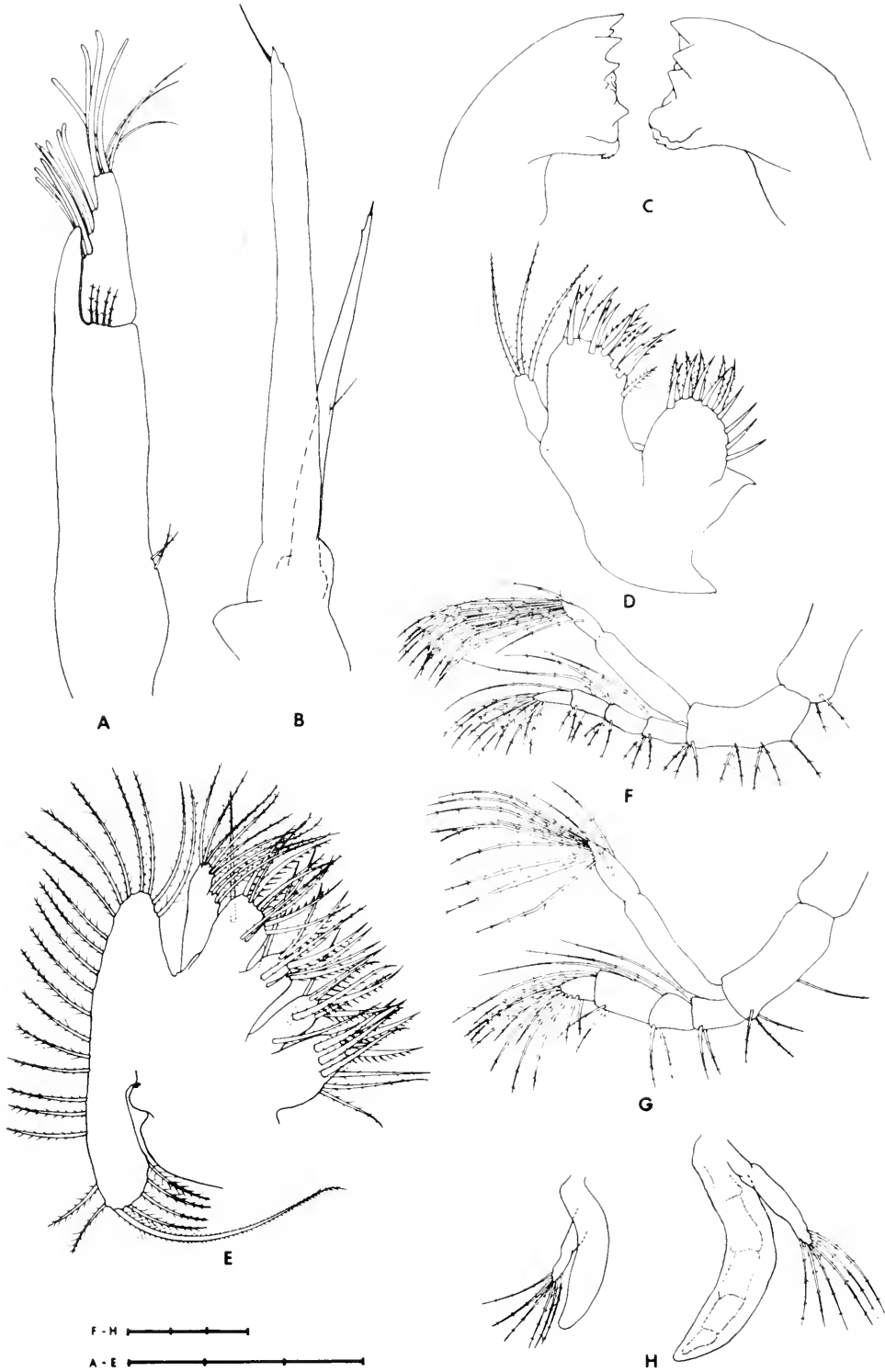


FIGURE 5.

Telson: (Fig. 3, D). Distinctive characters are minutely serrated lateral spines, the thin seta next to each telson spine, two fine hairs on central prominence, minute serrations terminally on articulated plumose setae, and two fine hairs medially on dorsal surface of telson proper. Anal spine present.

Color: Zoea transparent with red-orange chromatophores as follows: dorsally and ventrally surrounding the gut throughout abdomen; dorsally between lateral spines on telson; dorsally on tip of telson; interiorly around mouthparts. Rostral spine diffusely red-orange at tip and intermittently so throughout its length; posterior carapace spines diffusely red-orange only at tips.

Second zoea

Carapace length: 1.7 mm.

Number of specimens examined: 15

Carapace: (Fig. 3, B). Three pairs of dorsal setae persist. Rostral spine about six times carapace length; posterior spines of carapace up to 1.6 times carapace length. Spination on ventral margin of posterior spines may extend onto carapace in some zoeae.

Antennule: (Fig. 5, A). Biramous; exopodite with three or four long aesthetascs and four terminal setae, one seta plumose. Subterminal aesthetascs arranged in three groups as 2, 3, 3 progressing proximally. Endopodite slightly less than half the length of exopodite. At junction of exopodite and endopodite are four small setae; two additional setae on basal medial projection of protopodite.

Antenna: (Fig. 5, B). Similar to Stage I. Exopodite now about $\frac{2}{3}$ as long as endopodite; distal spination almost absent.

Mandibles: (Fig. 5, C). Larger, with three or four teeth and smaller dentate processes. No evidence of palp though Faxon (1879) reported one present and illustrated it as rudiment.

Maxillule: (Fig. 5, D). Endopodite setation same as in Stage I. Coxal and basal endites with 12 processes each, as illustrated. One late stage zoea with one more seta on lateral margin of coxal endite making 13 processes.

Maxilla: (Fig. 5, E). Endopodite setation same as Stage I. Coxal endite with 17-18 setae, 10 on proximal lobe, seven or eight on distal lobe. Basal endite with nine setae on proximal lobe, 11 on distal lobe. Scaphognathite retains apical plumose seta, and now has about 24 setae on margins.

Maxilliped 1, 2: (Fig. 5, F-G). Terminal segments of exopodite and endopodite with about 12 setae. Dorsal tuft of hairs on endopodite segments replaced by single long seta on each segment. Third segment of endopodite of Maxilliped 2 much swollen and nearly twice as long as other segments. Other setation similar to Stage I.

FIGURE 5. *Polyonyx gibbesi*; second zoeal appendages. A. Antennule. B. Antenna. C. Mandibles. D. Maxillule. E. Maxilla. F. Maxilliped 1. G. Maxilliped 2. H. Maxilliped 3, early (l.) and late (r.) stage. Scales equal 0.3 mm.

Maxilliped 3: (Fig. 5, H). Exopodite indistinctly segmented with six terminal setae. Endopodite, the same length or slightly longer than exopodite after molting, increases in length throughout second stage. It measures three times the length of exopodite just before molt to megalopa.

Pereiopods: One and five indistinctly chelate. Gills present. Pereiopods and gills enlarge noticeably throughout duration of Stage II. Toward end of Stage II the almost completely formed pereiopods are tucked under posterior portion of carapace.

Plcopods: (Fig. 3, B). Buds present, of decreasing length on abdominal segments two through five. Buds increase in size as stage progresses.

Telson: (Fig. 3, E). Now with two long articulated plumose setae on central prominence making 8 + 8 processes. Fine hairs below prominence and those adjacent to lateral spines are retained. Setation on dorsal surface unchanged from Stage I.

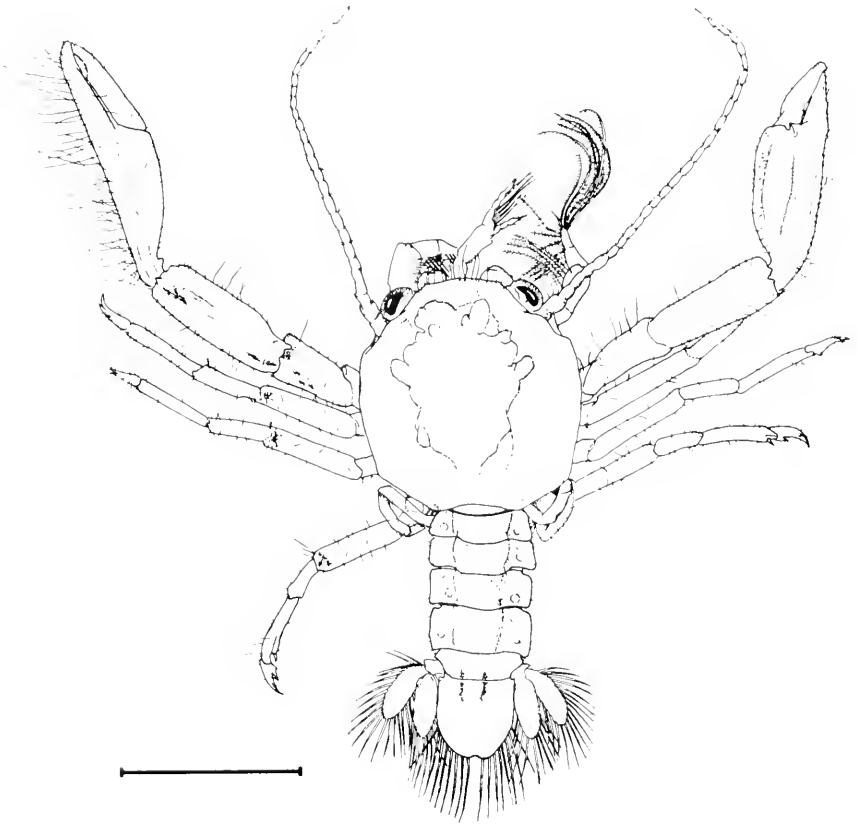


FIGURE 6. *Polyonyx gibbesi*; megalopa. Right antennule and left maxilliped 3 removed for clarity. Scale line equals 1 mm.



FIGURE 7. *Polyonyx gibbsi*; megalopal appendages. A. Antennule (less tips of aesthetascs). B. Antenna (in part). C. Mandible. D. Maxillae. E. Maxilla. F. Maxilliped 1. G. Maxilliped 2. H. Maxilliped 3. Scales equal 0.3 mm.

Color: Chromatophore distribution and color similar to Stage I. As zoea approaches molt to megalopa, rostral spine becomes completely orange. Posterior spines of carapace colored orange but to lesser extent.

Megalopa

The megalopa (Fig. 6) resembles the adult crab closely, enough so that Faxon (1879, 1882) considered it the "first stage of the crab." And, though he acknowledged the presence of biramous pleopods on the abdomen, to substantiate his belief he also cited a lack of persistent zoeal characters expected to appear in a megalopa stage. Because of this, at least one author (*e.g.*, Williams, 1965: 114) has quoted Faxon's error without further evaluation.

Chelae are well developed and fringed with setae on their outer margin. The juvenile and adult crab is always broader than long whereas only in the megalopa are the animals longer than broad. Carapace width-to-length measurements ranged from 1.2 mm. \times 1.2 mm. to 1.4 mm. \times 1.4 mm. First crab measurements were 1.8 mm. wide by 1.6 mm. long.

Carapace: (Fig. 6). Rounded or somewhat quadrate. Frontal region little produced; bears numerous setae. Eyes relatively large compared to first crab stage.

Antennule: (Fig. 7, A). Biramous, with three-segmented peduncle; basal segment enlarged. Lower ramous three-segmented; upper ramous has seven segments with aesthetascs on segments two through five in the following sequence of rows and numbers: one row (10), two rows (10, 3, +2 setae), two rows (3, 2, +1 seta), one row (3). Other setation on both rami as illustrated.

Antenna: (Fig. 7, B). Three-segmented peduncle plus 25 short segments, each bearing several short setae. Terminal segment usually with a long seta.

Mandibles: (Fig. 7, C). Three-segmented palp present, first segment has two setae on distal edge; distal segment with approximately 15–20 strong setae and spines.

Maxillule: (Fig. 7, D). Endopodite two-segmented, with setae as shown. Basal and coxal endites have approximately 29 and 36 processes, respectively, placed as shown.

Maxilla: (Fig. 7, E). Endopodite unsegmented; three or four terminal, two or rarely three subterminal setae. Proximal lobe of coxal endite with at least 10 and up to 13 terminal processes, three subterminal processes and about 20 setae in a ring around middle of lobe. Distal lobe with six or seven terminal and three subterminal processes; seven setae progress down its side. Proximal lobe of basal endite with about 15 processes; distal lobe has about 30. Scaphognathite has 48 or more plumose setae around edge.

Maxilliped 1: (Fig. 7, F). Setation fragile and variable; exopodite with two to ten setae; endopodite with four to six setae; protopodite with 50 or more setae on coxal and basal lobes.

Maxilliped 2: (Fig. 7, G). Exopodite elongate, about 20 setae on its two segments. Setation on four-segmented endopodite progressing distally is 9, 8, 18–20,

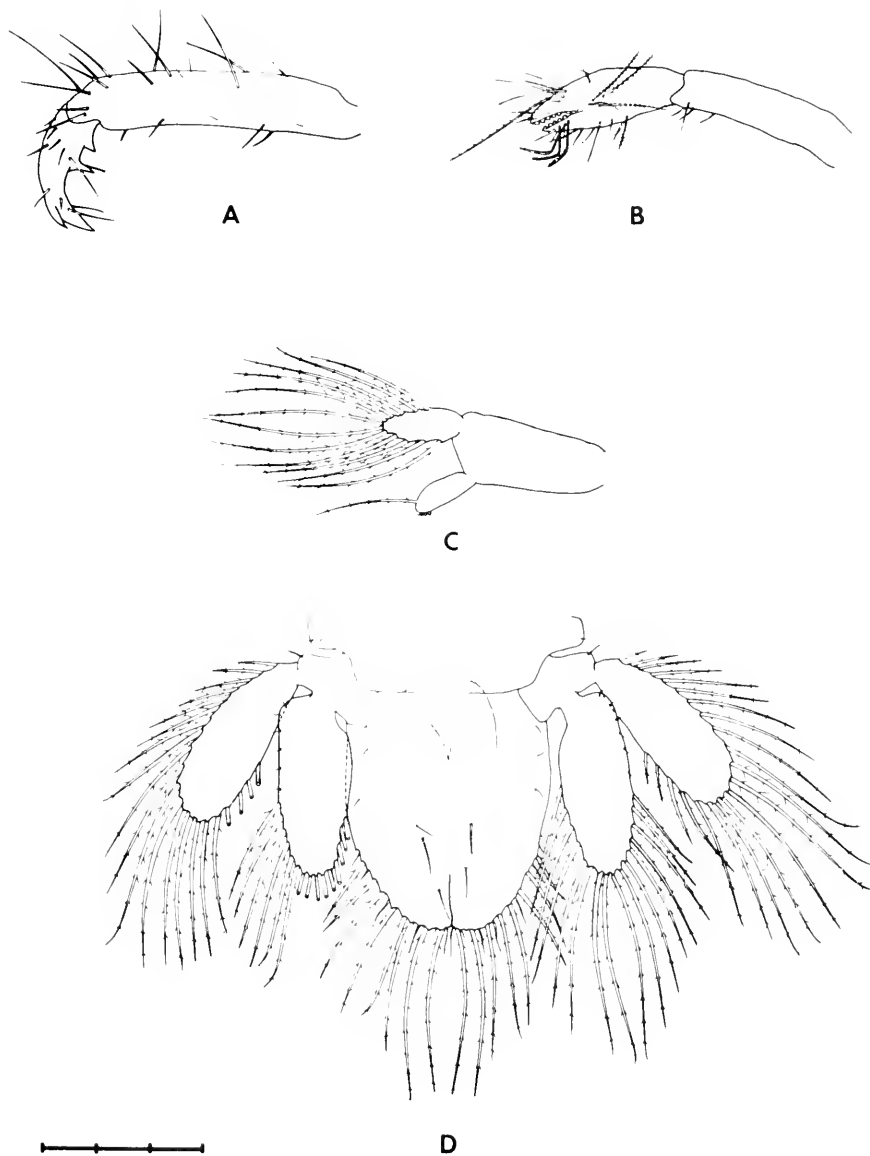


FIGURE 8. *Polyonyx gibbsii*; locomotory appendages of megalopa. A. Pereiopod 2. B. Pereiopod 5. C. Pleopod. D. Tail-Fan. Scale equals 0.3 mm.

about 18. Basipodite and coxopodite with about six and 12 setae, respectively. Placement as illustrated.

Maxilliped 3: (Fig. 7, H). Numerous setae on inner margin of basal lobes; about 14 setae and small spine on coxal lobe, as shown. Processes on five-segmented endopodite as follows: ischium, about 14; merus, about 16; carpus, about 18; pro-

podus, about 23; dactylus, about 15. Ischium, merus and carpus have thin, plate-like extensions. Exopodite has six to eight setae, placed as illustrated.

Pereiopods: (Fig. 6; 8 A, B). Chelipeds large, flattened, subequal, somewhat distorted, covered with setae. Walking legs setose, with two small blunt spines on distal edge of propodus (spines may be poorly developed); dactyl with three or four accessory spines. Pereiopod five chelate, gape appearing dentate in some, approximately 25 setae and three pectinate scythe-like hooks near gape.

Pleopods: (Fig. 8, C). Biramous, of decreasing size on abdominal somites nearer the telson. Exopodites with usually 12 setae; endopodites with one subterminal seta and an appendix interna.

Telson: (Fig. 8, D). 8 + 8 plumose setae plus additional spines and setae as

TABLE I
Comparison of zoeal appendages in two species of *Polyonyx*.
Data for *P. quadriungulatus* from Knight (1966)
and for *P. gibbesi* from the present work.

Appendage	Zoea I		Zoea II	
	<i>P. gibbesi</i>	<i>P. quadriungulatus</i>	<i>P. gibbesi</i>	<i>P. quadriungulatus</i>
ANTENNULE Exopodite Endopodite	Simple rod 3 aesthetascs; 2-3 setae	Simple rod 3 aesthetascs; 3 setae	Biramous 11 aesthetascs; 4 setae 1/2 length of exopodite	Biramous 11 aesthetascs; 4 setae 1/2 length of exopodite
ANTENNA Exopodite Endopodite	Biramous 2 × length of endopodite; spinous	Biramous 3 × length of endopodite; spinous	Similar to Stage I 2/3 length of endopodite Spination reduced As in Stage I	Similar to Stage I About equal to endopodite Spination as in Stage As in Stage I
MANDIBLE	Asymmetrical simple processes	Asymmetrical simple processes	3 large teeth No palp	2 teeth figured No palp
MAXILLULE Endopodite Coxal endite Basal endite	3 apical setae 4 spines; 6 setae 6 spines; 4 setae	3 apical setae 4 spines; 6 setae 6 spines; 4 setae	3 setae 5 spines; 7 setae 7 spines; 5 setae	3 setae 5 spines; 7 setae 7 spines; 5 setae
MAXILLA Endopodite Coxal endite Basal endite Scaphognathite	9 setae 13 processes 16 processes 5 setae; 1 spine	9 setae 13 processes 16 processes 5 setae; 1 spine	9 setae 17-18 processes 20-21 processes 24 (+1 apical) setae	9 setae 17 processes 20 processes 21-26 (+1 apical) setae
MAXILLIPED 1 Coxopodite Basipodite Endopodite Exopodite	2 setae 1,2-3,2,3, setae 3,3,3,10 setae 4 natatory setae	2 setae 1,2,2,3 setae 3,3,3,8 setae 4 natatory setae	2 setae 1,2,2,3 setae 4,4,12 setae 12 setae	2 setae 1,2,2,3 setae 4,4,9 setae 11-12 setae
MAXILLIPED 2 Coxopodite Basipodite Endopodite Exopodite	1 seta or naked 1-2,3 setae 2,2,2,10 setae 4 natatory setae	Naked 1,2 setae 2,2,2,8 setae 4 natatory setae	1 seta or naked 1,3 setae 3,3,3,12 setae 12 setae	Naked 1,2 setae 3,3,3,9 setae 11-14 setae
MAXILLIPED 3 Endopodite Exopodite	Small bifid lobe	Rudimentary	Now functional Increases in length 2 indistinct segments; 6 setae	Now functional Increases in length 2 indistinct segments; 6 setae
PEREIOPODS	Present as buds	Present as buds	Developing Chelation seen	Developing Chelation seen
ABDGMEN	5 somites with lateral spines	5 somites with lateral spines	Pleopod buds present Increase in size seen	Pleopod buds present Increase in size seen
TELSON	7 + 7 processes; 2 hairs on central prominence	7 + 7 processes; 2 hairs on central prominence	8 + 8 processes, hairs on prominence retained	8 + processes, hairs on prominence retained

shown. Uropods biramous, exopodites with 18–24, endopodites 12–16, setae around distal margins.

DISCUSSION

At present the complete larval development is known for only two of the 23 described species of *Polyonyx*, one from the eastern Pacific and one from the western Atlantic. A comparison between the eastern Pacific *Polyonyx quadriungulatus* and *P. gibbesi* shows that the zoeal stages are almost exactly similar in number and placement of setae (see Table I). *P. gibbesi* differs most notably in having more setae on the terminal segments of the endopodites of the maxillipeds in both zoeal stages (10 and 12) than *P. quadriungulatus* (8 and 9). Only detailed examination of each appendage reveals further differences between the zoeae of the two species (*e.g.*, different number of setae and spines on the second maxillae).

More easily observed is the relative length of the antennal exopodite to the endopodite in the two species (see Table I). Knight (1966) considered this a good character for distinguishing *P. quadriungulatus* from *Porcellana* and *Pisidia*, the two other members of Lebour's (1943) triad relationship. It is also possible on this basis to separate *P. gibbesi* from *P. quadriungulatus*. In the latter the antennal exopodite is three times the length of the endopodite in Stage I zoea, becoming about equal to the endopodite in Stage II. In *P. gibbesi* the antennal exopodite is only twice as long as the endopodite in Stage I and becomes $\frac{2}{3}$ the length of the endopodite in Stage II. Thus, the exopodite is always shorter in the zoeal stages of *P. gibbesi* than in *P. quadriungulatus*.

A third character which distinguishes *P. gibbesi* from *P. quadriungulatus* is seen in the dorsal setation of the zoeal carapace. The eastern Pacific form has but two setae in both zoeal stages while *P. gibbesi* has three pairs in both stages. Setae on the dorsal surface of the telson, often difficult to observe, may be an additional feature to separate larvae of the two species.

Differences in the megalopae of the two species are less distinct. The mouthparts are quite similar in both form, and number and placement of setae. A comparison of the megalopae in Table II shows that detailed examination is again necessary to separate the two forms. In general, however, *P. gibbesi* has more setae on the mouthparts than *P. quadriungulatus*. It also lacks both the two small spines at the bases of long setae and the articulated spines on the posterior distal margin of the propodus which *P. quadriungulatus* possesses.

The adult morphology of the two species is quite similar. Haig (1960, p. 239) stated that "Aside from *Polyonyx nitidus* Lockington, *P. quadriungulatus* is most closely related to . . . *P. gibbesi*." Further, both *P. gibbesi* and the two California species just mentioned belong to Johnson's (1958) "*P. sinensis* group" (Haig, 1960, p. 238). This group is a complex of species from the Indo-Pacific (and now including California to Panama, the eastern U. S., and the west African coast, Haig, *in Litt.*) which show similar morphology, plus "a pronounced tendency toward commensalism" (Johnson, 1958, p. 97). *P. gibbesi* is considered an obligate commensal with the polychaete worm *Chaetopterus variopedatus* (Gray, 1961), the megalopa establishing the initial relationship with the worm (Gore, unpublished data). Both California species have been found commensal with *Chaetopterus* though they may not be obligate commensals.

TABLE II
Comparison of megalopa appendages in two species of Polyonyx.
Data for P. quadrangulatus from Knight (1966)
and for P. gibbesi from the present work.

Appendage	<i>P. gibbesi</i>	<i>P. quadrangulatus</i>	Appendage	<i>P. gibbesi</i>	<i>P. quadrangulatus</i>
ANTENNULE	Biramous 3 segmented peduncle, basal segment en- larged 3 segments 7 segments; 10, 10 + 3, 3 + 2, 3 aes- thetases in tiers	Biramous 3 segmented peduncle, basal segment en- larged 3 segments 7 segments; 10, 10 + 3, 3 + 2, 3 aes- thetases in tiers	MAXILLIPED 2 Propodite Endopodite	18 setae 4 segments; 9, 8, 18 20, 18 setae (in tufts on segs. 3-4)	10 setae* 4 segments; -, 8, in tufts, in tufts
Ventral ramus Dorsal ramus	3 segments 7 segments; 10, 10 + 3, 3 + 2, 3 aes- thetases in tiers	3 segments 7 segments; 10, 10 + 3, 3 + 2, 3 aes- thetases in tiers	Exopodite	2 segments; 11 termi- nal, 9 marginal setae	2 segments; 11 terminal, 7 marginal setae
ANTENNA	About 25 segments 3 segmented peduncle Small lobe on first peduncular segment	About 30 segments 3 segmented peduncle Small cylindrical branch on first pe- duncular segment	MAXILLIPED 3 Propodite Endopodite Exopodite	Serrated spine, numer- ous setae 5 segments; about 14, 17, 21, 27, 17 processes 6 terminal, 6 marginal setae	Serrated tooth, numer- ous setae 5 segments; 15, 14, 20, 12, ? processes* 6 terminal, 6 marginal* setae
MANDIBLE Distal segment	3 segmented palp About 20 setae	3 segmented palp 15-17 setae	PEREOPODS 2-4	Dactyls with 3-4 hooks; propodus may have 2 small spines distally	Dactyls with 4 fixed hooks; propodus with 2 articulated spines
MAXILLULE Endopodite Basal endite Coxal endite	2 segments; 1-2 setae 14 spines, 15 setae 10 spines, 26 setae	2 segments; 1-2 setae 10 spines, 21 setae* About 44 processes*	PEREOPOD 5	Chelate; 26 setae + 3 scythe-like hooks	Chelate; 1 spine, 2-3 scythe-like hooks + setae
MAXILLA Endopodite Basal endite Coxal endite Scaphognathite	Unsegmented, 7 setae 45 processes 49 processes About 47 setae	Unsegmented, 8 setae 41 processes* 43 processes* 48-58 setae	PLEOPODS 1-4 Endopodites Exopodites	1 subterminal seta; 4-5 terminal hooks Usually 12 setae	1 subterminal seta; 4-5 terminal hooks 12, 12-13, 13, 13 setae
MAXILLIPEID 1 Propodite Endopodite Exopodite	50 or more setae 6-7 setae 10 or 11 setae	About 50 setae 7 setae 13-20 setae	UROPODS Endopodite Exopodite TELSON	10-12 setae 18-22 setae 8 + 8 plumose setae, + spines	10-13 setae 17-22 setae 8 + 8 plumose setae, + spines

The similarity in adult and larval morphology, the geographical isolation between Pacific and Atlantic forms, plus the similarity in commensal habitat indicates that *P. gibbesi* and *P. quadriungulatus* are geminate species. Thus they are one more species pair of the many that are known to exist between Caribbean-Atlantic and Pacific coast forms (e.g., *Minyocerus kirki* and *angustus*, *Porcellana cancri-socialis* and *sayana*, etc., see Haig, 1960).

Lebour (1943) thought that Faxon's *Polyonyx macrocheles* would fit into the *Porcellana*- (and *Pisidia*) -*Polyonyx* complex, distinguished chiefly by the placement of the fifth plumose setae of the telson in the first and second zoeal stages. Knight (1966) showed that *Polyonyx quadriungulatus* adhered to Lebour's scheme and the present work confirms Lebour's suggestion for *P. gibbesi* (formerly *P. macrocheles*). The differences in antennal proportions between *Porcellana-Pisidia* and *Polyonyx quadriungulatus*, as noted by Knight (see above), apply also to *P. gibbesi*. Thus, the zoeal features such as spined antennal exopodite and its length relative to both the antennal endopodite and to the antennule allow *P. gibbesi* to be separated from *P. quadriungulatus* in both zoeal stages, as already discussed, and from the known species of *Porcellana* and *Pisidia* in the first zoeal stage. The scheme breaks down in the second zoeal stage since, in *P. gibbesi*, the antennal exopodite is $\frac{2}{3}$ as long as the endopodite while the endopodite is about as long as the antennule proper. *Polyonyx gibbesi* thus shows antennal characters (in length) similar to those shown by known species of *Porcellana* and *Pisidia* in the second zoeal stage.

The previously mentioned carapace setation may, however, allow complete separation of *Polyonyx gibbesi* in both zoeal stages from *Porcellana-Pisidia* zoeae. If it is consistent in other members of the genus *Polyonyx* then, together with the features mentioned above, it would make *Polyonyx* larvae immediately distinguishable from most other porcellanid larvae. The value of this last character must await further studies on the larvae of other genera since some western Atlantic species of *Pachycheles*, *Petrolisthes*, *Porcellana* and *Minyocerus* also have dorsal carapacial setation (Gore, unpublished data). Studies are presently being carried out on the larvae of other genera of Porcellanidae from the south Florida and Caribbean area. Each of the genera mentioned has good distinguishing features in the zoeae which, in conjunction with Lebour's characters regarding the telson, may allow them to be separated from one another (Gore, unpublished data). As larvae of these genera become better known it will be possible to construct a key for their identification and to clarify the relationships between them.

I thank Dr. Anthony J. Provenzano, Jr., Thesis Committee Chairman, for his help throughout this study and for critically reading the manuscript. I also thank the members of my thesis committee, Drs. F. M. Bayer, A. A. Myrberg, Jr., and D. M. Moore for their perceptive advice and criticism. Miss Janet Haig also read the manuscript and clarified points concerning the zoogeographical distribution of the *Polyonyx sinensis* species complex. Mrs. C. Edith Marks eased the burden of laboratory work. She and Talbot Murray also aided in recording some of the data. I must also acknowledge Dr. H. B. Moore and members of his department, for their help in collecting *Polyonyx* while engaged in his own collecting work in Biscayne Bay, Florida.

SUMMARY

1. The larval development of the porcellanid crab, *Polyonyx gibbesi*, a commensal with the polychaete worm *Chaetopterus variopedatus*, is described and illustrated. Two series of larvae were hatched and maintained in the laboratory, one fed with *Artemia* nauplii and the other starved. Members of each series were held at 10°, 15°, 20°, 25° and 30° C. At 25° C. the fed larvae hatched as pre-zoeae and molted through two additional zoeal stages to the megalopa. Duration of the pre-zoeal stage is about two hours, each of the zoeal stages usually lasts six to seven days and the megalopa lasts 12–14 days before molting to first crab. No crab stages were obtained above or below 25° C. and no megalopae were obtained below 20° C. Starved larvae died before attaining Stage II.

2. Comparison of the larvae of *Polyonyx gibbesi* with those of *Polyonyx quadriungulatus*, an eastern Pacific species, shows the zoeae and megalopae to be almost identical both in appendages and in form, numbers, and placement of setae. Similarity of morphology and habitat plus geographical isolation indicate that *P. gibbesi* and *P. quadriungulatus* are geminate species.

3. Larvae of *Polyonyx gibbesi* possess certain features which allow them to be recognized in the plankton as well as distinguished from known larvae of genera of other western Atlantic porcellanid crabs.

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COMPENSATION OF LIVER PROTEIN SYNTHESIS IN TEMPERATURE-ACCLIMATED TOADFISH, OPSANUS TAU

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In the process of temperature acclimation poikilotherms adjust their metabolic rates in order to maintain physiological activity at a more nearly constant level over a range of environmental temperatures. Thus, they attain a measure of independence of temperature (Prosser and Brown, 1961). A review of the earlier literature dealing with this phenomenon and discussion of its significance in ecology and evolution has been given by Bullock (1955). More recently, the subject has been reviewed in several symposia (Prosser, 1958; 1967; Troshin, 1967). Most studies have dealt with temperature acclimation in terms of physiological (heart rate, oxygen consumption) or behavioral parameters; it is only in the last few years that methods have become available to attempt an approach to the molecular basis of the phenomenon.

Several recent studies have presented evidence for changes in level of enzymes in response to acclimation. Freed (1965) demonstrated an increase in cytochrome oxidase activity in goldfish muscle as measured at 15° after the fish had been acclimated to 5°; decreased activity resulted from warm acclimation (25°). Similarly, increased 6-phosphogluconic dehydrogenase activity is found in gills of cold-acclimated carp (Ekberg, 1962). Hochachka and Hayes (1962) showed greater utilization of the pentose shunt in cold-acclimated trout. Changes in the lactic dehydrogenase isozyme system have been found to accompany thermal compensations in goldfish (Hochachka, 1965). Because of the probability that these activity increases are due to increased levels of enzyme protein, a study was begun on the effect of low temperature acclimation on protein synthetic capacity in fish. The toadfish was chosen as the experimental animal because of its ready adaptability to laboratory conditions, and its wide range of natural habitat (Bigelow and Schroeder, 1953; Henshall, 1891), suggesting a high degree of temperature adaptability. Studies were concentrated on the liver where (in the case of rat) a good deal has been learned about the mechanism of protein synthesis. Of equal importance to the question of the mechanism of temperature acclimation is the possibility that the phenomenon can be used to elucidate sites of regulation in the protein synthetic system *per se*. A preliminary report of this work has appeared (Haschemeyer, 1967).

MATERIALS AND METHODS

Animals

Adult toadfish, 200–300 g., were obtained from the Supply Department at the Marine Biological Laboratory, Woods Hole, and included both sexes. In the first series of experiments the fish were kept in running sea water aquaria at the Marine Biological Laboratory. With the available refrigeration system it was possible to keep groups of up to six fish at a temperature of $10^{\circ} \pm 1^{\circ}$ in running sea water during the period of low temperature acclimation. Control fish were kept at 20–22°, the laboratory temperature of the sea water supply at Woods Hole at that time. In the second series of experiments the fish were maintained in aerated static aquaria prepared with a synthetic sea salt mixture (Rila Products, Teaneck, N. J.). For low temperature acclimation the fish were kept in a 25-gal. Instant Ocean aquarium (Aquarium Systems, Inc., Wickliffe, Ohio) with temperature control at $10^{\circ} \pm 0.5^{\circ}$. Control fish were kept at room temperature, 22–24°. The toadfish were fed small *Fundulus heteroclitus* to appetite.

Measurement of in vivo protein synthesis

For determination of incorporation of C^{14} amino acids into liver protein, a rapid arterial injection route was used, as described by Cooperstein and Lazarow (1964). In cases where the incubation temperature for the measurement differed from the acclimation temperature, the fish was transferred to the new temperature $\frac{1}{2}$ –1 hr. before injection. Each fish was wrapped in cheesecloth with its tail brought around to the right of the body, and a hole was cut in the cloth over the left gill. The skin on either side of the operculum was clipped in order to expose the gill arches and 0.2 ml. of a solution containing 10 μ c. of C^{14} amino acids was injected through a 25-gauge, $\frac{5}{8}$ " needle into the branchial artery of the fourth gill arch. The isotope solution was prepared from a mixture of 15 purified C^{14} -L-amino acids (New England Nuclear Corp.) of high specific activity (1 mc./mg.); the commercial solution in 0.1 N HCl was neutralized, buffered with 0.5 M Tris, pH 7.4, and brought to 1% NaCl. After injection the gill flap was held closed for about 5 seconds to aid clotting at the needle hole, and the fish was returned to a pail of sea water for the incubation period. The entire procedure took about 2 minutes; the fish started swimming as soon as returned to water and showed no ill effects. After time intervals of 5 min. to 30 min. the fish was stunned by a blow to the head, and the liver quickly excised and transferred to cold Medium A (0.25 M sucrose, 0.05 M Tris, pH 7.4, 0.025 M KCl, 0.01 M $MgCl_2$). In some cases to determine free radioactivity in the blood as a measure of the effectiveness of the injection, approximately 0.7 ml. of blood was collected by syringe from the heart cavity and added to 0.1 ml. of 0.1 M ethylenediamine tetraacetate, K^+ , pH 8.0.

Each liver was accurately wet-weighed, minced and homogenized in 3 volumes cold Medium A in a Sorvall Omni-mixer at half of full speed. For the second series of experiments in which biochemical analyses were made, the livers were homogenized in 3 volumes of ice-cold distilled water. The homogenates were analyzed for C^{14} incorporation into protein using a filter-paper disc method, based on the method of Bollum (1959). For each homogenate 100- μ l. aliquots were

pipetted onto several discs (Whatman 3MM filter paper, 2.3 cm. diameter). After a few seconds to allow the aliquot to soak in, all discs were dropped into a washing solution of ice-cold 10% trichloroacetic acid. The amount of washing solution in this and subsequent steps was about 5–10 ml. per disc; the time for each step was about 15 min., during which the solution and discs were swirled occasionally. A second 10% trichloroacetic acid wash was done, then five washes in cold 3% perchloric acid, two in cold 95% ethanol, and two in absolute ether. The discs were air-dried, placed in glass vials and counted in a Packard Tri-Carb liquid scintillation counter using a toluene solution of 4 g./l. 2,5-diphenyloxazole and 0.5 g./l. 1,4-bis-2-(5-phenyloxazole)-benzene.

Free radioactivity in the liver homogenates was determined as follows: 1.00 ml. of the homogenate in a clinical centrifuge tube was treated with 1.00 ml. of cold 10% trichloroacetic acid and kept in ice for 15 min. The protein-nucleic acid precipitate was spun down, and 100 μ l. of the clear supernatant were pipetted into a glass vial with 10 ml. of Bray's scintillation solution (Bray, 1960) and counted in the Packard scintillation counter. Values for TCA-soluble radioactivity therefore refer to homogenate which has been diluted by a factor of two, whereas the values for protein (TCA-precipitable) radioactivity refer to the undiluted homogenate. Free radioactivity in the blood was determined as for the liver homogenate except smaller volumes were used.

Biochemical analyses

DNA, RNA, and protein analyses following Schneider (1945) were made at three concentrations of the liver homogenates prepared in distilled water. Three clinical centrifuge tubes were prepared containing 0.25, 0.50 and 1.00 ml. of the homogenate. The first two were brought to 1.00 ml. with distilled water and all were treated with 1.00 ml. of cold 10% perchloric acid to precipitate proteins and nucleic acids. After the precipitate was washed with cold 10% perchloric acid and twice with cold 95% ethanol, the nucleic acids were hydrolyzed in 3 ml. 5% perchloric acid at 90° for 15 min. The supernatant was removed and analyzed for total nucleic acids by ultraviolet absorption at 260 $m\mu$, for RNA with orcinol and for DNA with diphenylamine. The protein precipitate was collected on a pre-weighed Millipore filter, air-dried, and weighed. The diphenylamine reaction gave $OD_{600} = 0.364/\text{mg. Na DNA}$, based on standard solutions of calf thymus DNA (General Biochemicals). The DNA content of this preparation (as Na DNA, residue weight 331) was found to be 0.66 mg. per mg. of weighed material, as determined from OD_{260} of an acid hydrolysate; the conversion factor $OD_{260} = 30$ for 1 mg./ml. hydrolyzed Na DNA was calculated from extinction coefficients of isolated nucleosides (Beaven *et al.*, 1955). Recovery in the sample preparation was 75% for calf thymus DNA, and DNA values for the liver homogenates were corrected accordingly. Results are based on the slope obtained in a plot of OD_{600} vs. concentration of homogenate. A similar plot of OD_{260} for the hot perchloric acid supernatant vs. concentration yielded values for total nucleic acid concentration in the homogenate. A standard nucleoprotein consisting of purified *E. coli* K 12 ribosomes, prepared according to Nirenberg (1964), was tested in the procedure. Recovery of the ribosomal RNA was

about 75% after precipitation, washing and hydrolysis, compared with direct acid hydrolysis. Total nucleic acid as mg./ml. of liver homogenate was estimated on the basis of this figure and the conversion factor above, and the DNA content was subtracted to obtain RNA. The results in Table III are expressed in mg. per g. wet weight of liver. Measurement of RNA by the orcinol procedure was unsatisfactory due to apparent interference by a liver component which contributed as much as one-half of the total orcinol reading.

Free amino acid analyses were made by the method of Spies (1952) although modification was required because of interfering materials in the whole liver homogenate. The homogenate was first spun at 15,000 *g* for 10 min. in a Sorvall refrigerated centrifuge to remove large particles and debris and then at 105,000 *g* for 3 hr. in a Spinco L2 centrifuge (at 4°) to remove glycogen and ribosomes. The supernatant was carefully withdrawn by pipette, avoiding the lipid layer; proteins were precipitated by addition of 70% perchloric acid to a final concentration of 5%. The precipitate was spun down in a clinical centrifuge and the clear supernatant was neutralized with NaOH and analyzed at two con-

TABLE I

Effect of low temperature acclimation (10°) on protein synthetic capacity of toadfish liver measured in vivo (running sea water aquaria). Incubation time after arterial injection = 15-30 min. Data expressed as cpm in trichloroacetic acid precipitate (on filter paper disc) divided by cpm in trichloroacetic acid-soluble supernatant

Acclimation period Days at 10°	Number of animals*	Temperature of measurement	Incorporation into protein compared to available free radioactivity in liver
0	9	20-22°	0.24 ± 0.15 (S.D.)
3	4	20-22°	0.27 ± 0.15
3	4	10°	0.09 ± 0.04
7	3	20-22°	0.48 ± 0.11

* Average body weight = 240 g.

centrations for the 230 *mμ* absorption produced by Cu⁺⁺-amino acid complexes. Each solution was read against a blank consisting of all ingredients except the CuCl₂ reagent. All optical density readings were made with a Beckman DU spectrophotometer equipped with a Gilford Model 2000 photomultiplier unit and recorder. A standard solution of L-alanine (Schwarz Laboratories, Inc.) gave a value of OD₂₃₀ = 3.2 for a concentration of 1 *μm*./ml. amino acid in the final solution.

RESULTS

Table I shows the results of the first series of experiments on the effect of cold acclimation on incorporation of radioactive amino acids into liver protein *in vivo*. The nine control animals included fed and starved (3-7 day) individuals: there were not enough animals to establish a correlation between nutrition and protein synthetic rate. The fish kept at 10° did not eat during this period, although minnows were available in the tank. As shown, three days at 10°

produced no change in liver protein synthesis, when the fish were returned to 20–22° for measurement. When the measurement was made at 10°, the fish showed a low rate of synthesis, as expected for a drop of 10°, in the absence of compensation due to acclimation. The approximate Q_{10} is 2.5, based on the values in Table I and taking the higher temperature as 21°. Values of free radioactivity in the liver supernatants ranged from 500 to 3500 cpm. In two comparable groups the average for fish incubated at 10° was slightly lower (750 cpm) than that for fish incubated at 20° (1050 cpm). Measurements of free radioactivity in the blood correlated well with that in liver for each individual; either of these quantities thus provided a measure of the effectiveness of the injection. To eliminate this factor all the data have been expressed as the ratio of cpm in protein to cpm in the trichloroacetic acid-soluble phase of the liver homogenate. The ratio did not depend on time of incubation between 15 and 30 min., although small changes would have been obscured by the large variation among individuals. Shorter incubations did show time dependence and are not included in Table I. The large scatter in the final data, as indicated by the reported standard deviations, must be

TABLE II

Effect of low temperature acclimation (10°) on protein synthetic capacity of toadfish liver measured in vivo (static artificial aquaria). Incubation temperature = 23°; time = 20–30 min.

Expt. no.	Days at 10°	Nutritional state	Number of animals	Incorporation into protein compared to available free radioactivity
1	0	Starved 5 days	4	0.21 ± 0.11 (S.D.)
2	0	Fed	3	0.23 ± 0.16
3	3	Starved 4 days	3	0.13 ± 0.03
4	14	Starved 14 days	4	0.37 ± 0.12
5	14	Fed	4	0.30 ± 0.07

attributed to population variability of unknown origin. Hormonal influences on liver metabolism following the stress of handling and injection may play a role. Adrenal and pituitary hormones have been shown to influence levels of C¹⁴ leucine incorporation into protein in rat liver following a single intraperitoneal injection of the isotope (Reid *et al.*, 1956).

After seven days at 10°, as shown in Table I, an increase in protein synthesis is observed upon measurement at 20°. It must be noted that the number of animals is small and the standard deviation in all these measurements rather large; the absolute values must be viewed accordingly. However, analysis by the standard t-test indicated the difference between the 10°-acclimated and control groups to be significant at $P = 0.05$. Thus, the results strongly suggest that liver protein synthesis in toadfish exhibits at least a partial compensation in Precht's (1958) terminology.

The results for *in vivo* protein synthesis in the second series of experiments are given in Table II. In this series more attention was given to nutrition, although again no reliable correlation between feeding and protein synthesis could be established. Fish labelled "fed" had taken minnows one or two days before the experiment; remains were found in the gut at autopsy. The fish kept at 10°

showed little interest in the minnows in the tank, but even after two weeks the toadfish showed no significant weight loss or other apparent sign of starvation. For Expt. 5 the fish were hand-fed by holding minnows in front of them and tapping the mouth until they snapped angrily. The fish were also observed carefully for behavioral changes during 10° acclimation. Unfortunately the toadfish is by nature a sluggish bottom dweller, and the specimens used here conformed to this description equally well at 10° and 20°. Respiratory movements averaged about 8 per min. at 10° and did not change over the two-week acclimation period. The only difference noted was that some of the two-week acclimated fish, after having been warmed to 20°, were hyperactive and difficult to handle during the injection procedure. Effects of low temperature acclimation on oxygen consumption in this species have not been reported, although it would be particularly interesting in view of the unusually high skin respiration of toadfish which permits them to survive 1-2 days out of water (Schwartz and Robinson, 1963).

As indicated in Table II, the two groups of control (23°) fish, fed and 5-day starved, gave comparable values for liver protein synthesis. The fact that all values in the second series are less than in the first is due primarily to the lower efficiency for counting filters of the scintillation counter used for the second series. This, of course, does not affect the results on a comparative basis. Other differences in the second series were the use of static aquaria, and the slightly larger size of the fish collected later in the season. Females in this group had large well-developed eggs. In order to test for a possible effect of temperature shock as a result of transfer from 10° to over 20° immediately before the experiment, a group of fish was again assayed after 3 days at 10°. The average value of 0.13 is low compared to the controls; it is not certain what significance this may have, particularly since it was not observed in the first series. In any case, the sudden transfer of the fish from 10° to 23° does not produce by itself the increase of protein synthesis found after the longer periods at 10°. Expts. 4 and 5 in Table II show the effect of 14 days of 10° acclimation on liver protein synthesis. Comparison of the average of Expts. 4 and 5 with the average of 1, 2 and 3 yields an increased protein synthesis of about 75% as a result of low temperature acclimation (significant at $P = 0.2$ by the t-test). This is in good agreement with the results of Table I.

Biochemical analyses were also performed in the second series to determine if any major alterations of the cellular constituents involved in protein synthesis occurred as a result of 10° acclimation. Of particular importance is the free amino acid pool of liver because the amount of incorporation of the injected isotope will depend on the specific activity of the pool as well as on the rate of protein synthesis. If the pool size does not vary greatly, then the present method of measurement, based simply on cpm per unit volume of homogenate, is justified. On the other hand, if the amino acid pool in cold-acclimated fish is low, then the specific activity will be higher for the same amount of isotope, and incorporation of label into protein will be increased accordingly (this assumes that the rate is not directly dependent on pool concentrations, in which case, the effect would cancel out). As shown in Table III, although there is some decrease in free amino acid concentration in the cold-acclimated fish (Expts. 4 and 5), it is not sufficient to account for the 75% increase in incorporation into protein

shown in Table II. In fact, the observation that fish at 10° for 3 days (Expt. 3) show the same low pool value suggests that the decrease may be related to the low temperature treatment but not to the acclimation phenomenon. The present values for free amino acid concentration in a fish liver are similar to those reported for rat liver. The sum of the isolated amino acids of rat liver (one day starved or protein-fed animals) reported by Wiss (see Tarver, 1963) amounts to 24 μ moles/g. liver.

Individual liver-to-body-weight ratios generally showed the expected increase with body weight (Robinson *et al.*, 1960); average ratios in the cold acclimated fish are slightly lower than controls, probably because the fish did not eat during most of the period. This is also indicated by the slight increase of DNA per gram of tissue. Protein yields, however, did not change. Although no analysis was done for lipid and glycogen, it was noted during the centrifugation steps used in preparation for amino acid analysis that the lipid layer formed in the 15,000 *g*

TABLE III

Biochemical analyses for livers of control and cold-acclimated toadfish (incorporation data in Table II)

Expt. no.	Body wgt. average	% Liver	mg./g. tissue (wet)			μ m./g. tissue
			body	DNA	RNA	Protein
1	241	2.8	2.36	3.85	135	29
			± 0.27	± 1.40	± 15	± 3
2	314	4.0	2.28	4.15	112	26
			± 0.30	± 1.30	± 6	± 3
3	250	3.7	2.26	4.05	116	23
			± 0.25	± 2.00	± 9	± 2
4	279	3.1	2.81	5.45	110	23
			± 0.19	± 1.95	± 7	± 2
5	260	2.8	2.48	3.80	133	21
			± 0.25	± 1.95	± 17	± 2

centrifugation was almost entirely lacking in cold-acclimated fish and the glycogen pellet obtained at 105,000 *g* was reduced in size. This may account for the slight loss in liver mass in the cold-acclimated animals.

The values obtained for total RNA per g. liver showed wide variation, as indicated by the large standard deviation. Examination of the individual data revealed that values clustered around two levels; about 6 mg./g. and about 2 mg./g. Averaging of the high and low values separately yielded values of 5.70 ± 0.70 mg./g. for 11 fish and 2.15 ± 0.20 mg./g. for 7 fish. These are both low compared with values obtained by the same method for rat liver (Haschemeyer and Gross, 1967) of about 8 mg./g. in total homogenates. The finding of the higher RNA content for the seven fish which contained eggs suggested a correlation of liver RNA content with the sex of the animal. The levels of all tissue constituents measured differ appreciably from those reported for goldfish liver (Das, 1967), when compared in the same units. In that study, however, the fish were well-fed, with resultant hypertrophy of the liver in the low temperature group and concomitant increase of protein and decrease of

DNA in relation to tissue mass. In spite of the higher rate of protein synthesis, such hypertrophy was not observed in the toadfish liver, probably because the fish ate little on their own initiative during the low temperature period.

DISCUSSION

The purpose of these experiments was to determine what type of compensation might occur in liver protein synthesis of a marine fish, following varying periods at a temperature at the lower end of the animal's normal environmental range. Protein synthesis was assessed in terms of the incorporation of C^{14} amino acids into protein compared to the available free radioactivity in the liver. The procedure involving injection into the arterial blood system permitted rapid uptake of labelled amino acids by the liver; thus incorporation into protein could be measured after short intervals, before degradation or export would be expected to significantly affect the levels of labelled proteins. Rapid pulse measurements are well advised in view of the short half-lives of some liver proteins (in rat), particularly inducible enzymes; tryptophan pyrrolase (peroxidase), for example, is degraded with a half-life of about 2.3 hr. (Feigelson *et al.*, 1959). Other proteins are relatively long-lived; ribosomal protein probably has a half-life of five days, like that of liver ribosomal RNA (Loeb *et al.*, 1965).

The data presented here establish that on the average, toadfish acclimated under laboratory conditions to a temperature of 10° exhibit an increase in liver protein synthesis of about 75% over control fish acclimated to $20-23^{\circ}$, when both are measured at the higher temperature. The biochemical analyses indicated little alteration in tissue components of liver involved in protein synthesis, under the acclimation conditions used in these experiments. Most important, no significant change in free amino acid pools occurred, which would produce some uncertainty in the use of incorporation data as a measure of protein synthesis. Although Das and Prosser (1967), in a study of goldfish acclimation reported during the course of the present work, were unable to establish a significant effect in liver in short-time experiments, this may have been due to the use of intraperitoneal injection. They found, however, that accumulation of label in protein over long periods was increased in cold-acclimated fish, consistent with a greater protein synthetic rate.

The present results support the hypothesis that a control in the protein synthetic pathway may provide a common basis for increased levels of enzymes in respiratory metabolism and other essential pathways responsible for physiological adaptation to low temperatures. Such a change must of course be accompanied by associated changes in degradation rates or rates of protein export from the liver to arrive at a new equilibrium level of proteins. Although a common site for control is suggested by the magnitude of the effect, this does not mean that all proteins are necessarily equally affected; for example, an increase in rate at a step common to all proteins will not affect proteins whose synthesis is rate-limited at another step. The possibility of selective control has been suggested by the finding that different codons are used in the insertion of leucine (Weisblum *et al.*, 1965) and arginine (Weisblum *et al.*, 1967) into the α chain of rabbit hemoglobin. In the case of leucine it was shown by *in vitro* protein

synthesis with *E. coli* transfer RNA that a minor species of leucine-accepted tRNA was required to insert leucine into a particular site on the α chain. Thus, the level of a "rare" transfer RNA or its associated aminoacyl-tRNA synthetase may limit the synthesis of some proteins, even during a state of generally increased protein synthesis. Another way in which proteins may be selectively controlled results from special requirements for release from the messenger RNA-ribosome complex or their influence on polyribosome activity. It has been suggested that in hemoglobin the dependence of globin synthesis upon the availability of the prosthetic group is due to a control over release from the protein synthetic site (Gribble and Schwartz, 1965). Another study shows increased size and stability of reticulocyte polyribosomes in response to hemin addition (Grayzel *et al.*, 1966).

Thus, the increased protein synthetic rate in toadfish liver provides a simple mechanism for obtaining increased levels of many proteins, without requiring a specific derepression of genes, as in the Jacob-Monod model (1961) of bacterial enzyme induction. Although the present results refer only to liver, it is quite likely that other tissues exhibit the same phenomenon. Das and Prosser (1967) found evidence for temperature compensation of protein synthesis in gill and muscle tissue of goldfish. Mews (1957) and Jankowsky (1960) have reported increased protein synthesis during cold adaptation of frog skeletal muscle. In order to investigate the possibility of a common translational-type control over protein synthesis, subject to variation with acclimation temperature, we have turned to the study of protein synthesis and the components of the protein synthetic system *in vitro*. These studies will be reported in a separate communication.

This investigation was supported by grant #GB 5194 from the National Science Foundation. The author is indebted to Dr. Jerome Gross for helpful discussions and to Miss J. Levy for expert technical assistance. The toadfish injection procedure was kindly demonstrated by Dr. J. Moran and Mr. J. Jackson. This is publication No. 449 of the Robert W. Lovett Memorial Group for the Study of Diseases Causing Deformities.

SUMMARY

Liver protein synthesis was measured with a rapid arterial injection procedure in control (20–23°) and cold-acclimated (10°) toadfish. The results were expressed as the cpm of radioactive amino acids incorporated into protein compared to cpm of free radioactivity in the liver homogenate, to correct for variability in injections and in uptake of amino acids by the liver. The results show that, when measured at 20–23°, the 10° acclimated fish possess liver protein synthetic capacity about 75% greater than fish maintained at the higher temperature. Two series of experiments, in which fish were maintained in running sea water aquaria or in static artificial sea water aquaria, gave comparable results. The livers were analyzed for DNA, RNA, protein and free amino acids. The levels of these constituents resembled those in mammalian liver with the exception of RNA, which was lower. Under the conditions of these experiments no significant changes were observed in the constituents measured as a result of two-week cold acclimation.

The possibility of a common translational-type control over protein synthesis to account for increased enzyme levels in cold temperature acclimation is discussed.

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PIGMENT COMPOSITION OF SIPHONALES ALGAE IN THE BRAIN CORAL *FAVIA*¹

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Many types of brain corals found on the Great Barrier Reef (*Favia*, *Porites*, and *Goneastrea*) contain a subsurface layer of a green filamentous alga, which forms a curved zone, 0.5–1.0 cm. wide, some 1–3 cm. beneath the surface layer of brown zooxanthellae. This green algal layer was first observed in corals from the Great Barrier Reef by Marshall and Stephenson (1928), and was found by Odum and Odum (1955) to contribute even more plant biomass to coral communities than the well-known coral zooxanthellae. The studies of Odum and Odum emphasized for the first time the importance of both types of coral symbionts to the productivity of reef communities.

Samples of green layers taken from the brain coral *Favia* were studied in the present work, and were found to consist mainly of the alga *Ostreobium*: probably *Ostreobium Reineckii* Bornet.² Preliminary observations on the nature of the algal pigments were made by Drs. L. Muscatine and F. T. Haxo (personal communication), who separated chlorophylls *a* and *b* from the green layer by thin layer chromatography and identified the fractions by absorption spectra. These observations were of interest taxonomically, since *Ostreobium* had been placed in the phylum Chrysophycophyta by Scagel (1966), in the family Phyllosiphonaceae within the Chlorococcales by Christensen (1962) and Parke and Dixon (1964), and in the family Phyllosiphonaceae within the Siphonales by Fritsch (1948) and Taylor (1957). The presence of chlorophylls *a* and *b* in the alga *Ostreobium* definitely excluded membership within the Chrysophycophyta, but could not distinguish between the other taxonomic possibilities.

A detailed re-examination of the full complement of photosynthetic pigments in the green layer of the brain coral *Favia pallida* Dana³ was begun on the basis of these preliminary observations. These studies also formed part of a general study of the physiology and photosynthetic capacity of the deep algal layer (Halldal, 1968; Shibata and Haxo, unpublished data). The pigments were examined by two-dimensional paper chromatography, and identified by R_f values and absorption spectra. The alga in the green layer of the brain coral *Favia* was found to contain the major pigments of the order Siphonales belonging to the class Chlorophyceae.

¹ Research begun on the University of California Research Vessel R/V "Alpha Helix" during the 1966 Expedition to the Great Barrier Reef, North Queensland, Australia. The ship-board work was supported by the National Science Foundation of the U. S. A. These studies were carried out in collaboration with Drs. F. T. Haxo, P. Halldal, and K. Shibata.

² Identified by Dr. W. Randolph Taylor and Dr. M. Nizamuddin.

³ Identified by Dr. E. C. Allison.

METHODS

1. *Extraction of pigments*

(1) *Coral*: Small pieces of coral containing green layers well separated from the zooxanthellae layer were chiselled out from the coral, and freed from any surface brown zooxanthellae. The coral was extracted with methanol for 1–2 hours in the dark in the presence of MgCO_3 , to prevent possible acidification of the extract during the long extraction period. Several changes of methanol were made until no further pigment was released and the coral layer was colorless. The combined methanol extracts were clarified by centrifugation at 1–2000 *g* for 5 minutes, and the pigments were transferred to diethyl ether by adding an equal volume of ether to the methanol extract and washing once or twice with a volume of 10% NaCl solution 5–10 times that of the methanol + ether extract. All the pigments migrated to the ether layer, which was collected, concentrated to a small volume by evaporation under nitrogen, and used directly for chromatography.

(2) *Algae*: Two other representatives of the order Siphonales (*Halimeda* sp. from Princess Charlotte Bay, North Queensland, and *Codium* sp. from Port Hacking, New South Wales) were studied in order to obtain pigments for comparison with those in the green layer. *Codium* species from the same locality were previously analyzed by Strain (1965). The tissues of the algae were extracted by homogenizing in methanol with added MgCO_3 , and the extracts prepared for chromatography as above.

2. *Chromatography*

(1) *Solvents*: Solvents used in all cases were A. R. Grade, and were not further purified.

(2) *Paper*: Pigments were chromatographed on Whatman No. 3 (or No. 3 MM) paper, using the two-dimensional solvent system of Jeffrey (1961). This procedure separates the chlorophylls and major carotenoid fractions from each other, but may not fully resolve all carotenoid isomers.

(3) *Thin Layer*: Thin layers of Al_2O_3 and MgO (3:1 w/w) were used to separate α - and β -carotenes. The solvent system was 4% ethyl acetate in hexane (Chapman, 1966). Standard α -carotene for reference was obtained from the cryptomonad *Chloromonas* sp. and β -carotene was isolated from the green flagellate *Dunaliella tertiolecta*. The absorption maxima in petroleum ether (60° – 80°) were 476, 448, and 424 for α -carotene, and 482, 451, and 430, β -carotene.

3. *Identification of pigments*

Pigments were identified by R_f values, and by absorption spectra of pigment fractions eluted from paper chromatograms in different solvents. Absorption curves were taken with recording spectrophotometers (Beckman DB, and Unicam SP 700).

4. *Determination of chlorophylls a and b*

Ratios of chlorophylls *a*:*b* were determined in extracts in 90% acetone, using the equations of Humphrey and Jeffrey (in preparation):

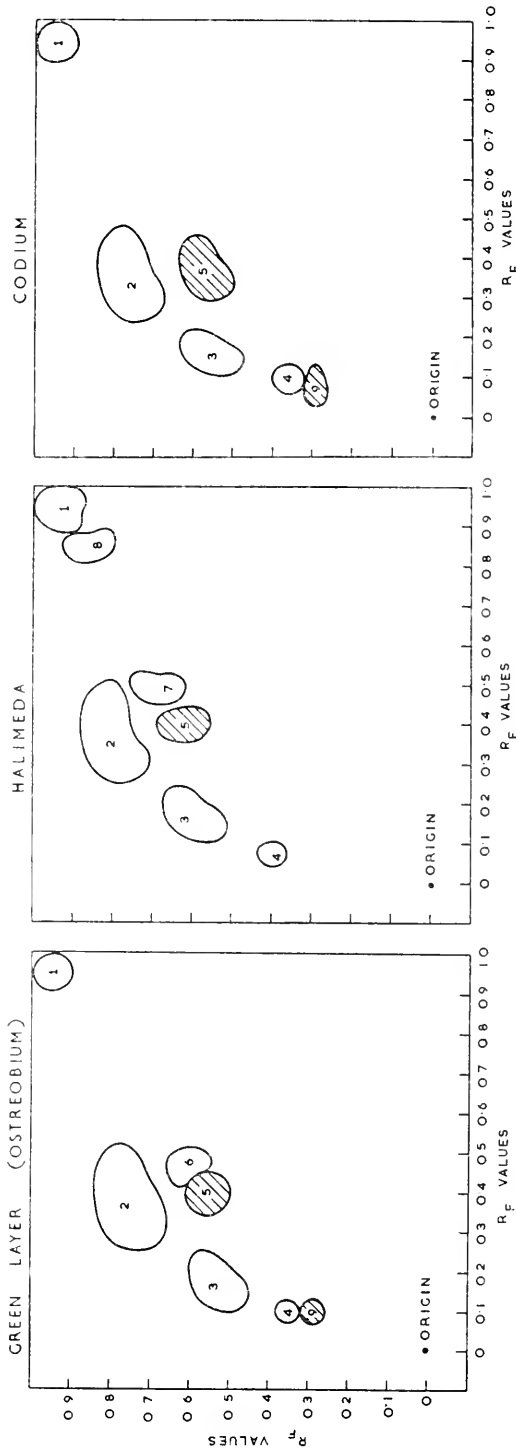


FIGURE 1. Two-dimensional paper chromatograms of pigments from *Halimeda*, *Codium*, and the green layer, *Ostreobium*. 1. Carotenes (yellow). 2. Chlorophyll *a* (blue-green). 3. Chlorophyll *b* (olive-green). 4. Neoxanthin (bright yellow). 5. Siphonaxanthin (pink-orange). 6. Unknown (yellow-orange). 7. Violaxanthin (pale yellow). 8. Lutein (yellow). 9. Siphonaxanthin (pink-orange).

$$\text{Chlorophyll } a = 13.50 E_{663} - 2.91 E_{645};$$

$$\text{Chlorophyll } b = -4.33 E_{663} + 21.20 E_{645};$$

where chlorophyll = concentration of chlorophyll in $\mu\text{g./ml.}$, E = extinction in liters/gm. cm. in a 1-cm. cell.

RESULTS

Figure 1 shows chromatograms of pigments from the green layer (*Ostreobium*) together with those of two well-known members of the Siphonales, *Halimeda* and *Codium*. Each organism contained chlorophylls a and b , a carotene zone, several yellow xanthophylls and one or more prominent pink-orange xanthophylls. Table I gives R_f values of the pigments from each alga, with tentative identification. Table II gives absorption maxima of the pigment fractions from each of the three organisms, compared with published maxima of authentic samples. The yellow

TABLE I
R_f values of pigments from Halimeda, Codium, and the green layer, Ostreobium
Paper chromatography in two solvent systems

Fraction	Pigment	Color	R _f values					
			4% n-propanol/pet. ether			30% CHCl ₃ /pet. ether		
			<i>Halimeda</i> *	<i>Codium</i>	<i>Ostreobium</i>	<i>Halimeda</i> *	<i>Codium</i>	<i>Ostreobium</i>
1	Carotenes	Yellow	0.95	0.94	0.94	0.94	0.94	0.94
2	Chlorophyll <i>a</i>	Blue-green	0.80	0.75	0.74	0.38	0.35	0.36
3	Chlorophyll <i>b</i>	Olive-green	0.59	0.55	0.52	0.17	0.15	0.16
4	Neoxanthin	Bright-yellow	0.38	0.36	0.35	0.06	0.09	0.09
5	Siphonoin	Pink-orange	0.59	0.58	0.56	0.37	0.37	0.40
6	Unknown	Yellow-orange	—	—	0.61	—	—	0.46
7	Violaxanthin	Pale-yellow	0.64	—	—	0.50	—	—
8	Lutein	Yellow	0.80	—	—	0.86	—	—
9	Siphonaxanthin	Pink-orange	—	0.30	—	—	0.05	—

* *Halimeda* pigments showed slightly higher R_f values in this solvent. It is known that other compounds (e.g., lipids) can force R_f values to higher levels during development (Sestak, 1958). R_f values are therefore not absolute, but give relative orders of separation.

xanthophylls, lutein, violaxanthin, and neoxanthin, which are normal members of the Chlorophyceae, were identified by R_f values and absorption maxima in *Halimeda*, but only neoxanthin was found in *Codium* and in the green layer. The green layer contained, however, large amounts of an unknown yellow-orange xanthophyll. The two pink xanthophylls, siphonoin and siphonaxanthin, which are characteristic of the Siphonales, were found both in *Codium* and in *Ostreobium*, but only siphonoin was present in *Halimeda*. It appears that some variations in the full complement of pigments which have been described for the group (Strain, 1958) are possible.

The carotene fraction from the organisms gave one zone on paper chromatography, and showed absorption maxima close to those of α -carotene (Table II). To determine more specifically the presence of α -carotene, extracts were chromatographed on thin layers of alumina/magnesium oxide (3:1) with 4% ethyl acetate in hexane as solvent (Chapman, 1966). In this system, α - and β -carotenes had R_f values of 0.67 and 0.41, respectively. The carotene fractions from *Halimeda*,

TABLE II

*Absorption maxima of pigments from Halimeda, Codium, and the green layer, Ostreobium
(Fractions separated by paper chromatography)*

Fraction	Color	Organism	Absorption maxima (nm)	Solvent	Identification
1	Yellow	<i>Halimeda</i> <i>Codium</i> <i>Ostreobium</i> Published maxima	423, 443, 470 417, 445, 475 423, 445, 472 422, 445, 475* 430, 450, 480*	diethyl ether	α -carotene β -carotene
2	Blue-green	<i>Halimeda</i> <i>Codium</i> <i>Ostreobium</i> Published maxima	426, 661 429, 661 429, 661.5 430, 662† 428.5, 660.5‡	diethyl ether	Chlorophyll <i>a</i>
3	Olive-green	<i>Halimeda</i> <i>Codium</i> <i>Ostreobium</i> Published maxima	453, 643 453, 643 455, 644 455, 644† 452.5, 642‡	diethyl ether	Chlorophyll <i>b</i>
4	Bright yellow	<i>Halimeda</i> <i>Codium</i> <i>Ostreobium</i> Published maxima	414, 437, 464 416, 440, 468 trace only 414, 437, 466*	diethyl ether	Neoxanthin
5	Pink-orange	<i>Halimeda</i> <i>Codium</i> <i>Ostreobium</i> Published maxima	460 462, (457, 481 pet. ether) 465.8 465**, (454, 480** pet. ether)	ethanol	Siphonein
6	Yellow-orange	<i>Halimeda</i> <i>Codium</i> <i>Ostreobium</i>	not present not present 449, 470.1	ethanol	Unknown
7	Pale yellow	<i>Halimeda</i> <i>Codium</i> <i>Ostreobium</i> Published maxima	415, 439, 468 not present not present 417, 442, 471*	ethanol	Violaxanthin
8	Yellow	<i>Halimeda</i> <i>Codium</i> <i>Ostreobium</i> Published maxima	420, 446, 476 not present not present 420, 446, 476*	ethanol	Lutein
9	Pink-orange	<i>Halimeda</i> <i>Codium</i> <i>Ostreobium</i> Published maxima	not present 449, (450, 477 pet. ether) trace only 450, (451, 480** pet. ether)	ethanol	Siphonaxanthin

* Losev 1964 (including supplementary data of Strain, 1938; and Karrer and Jucker, 1950).

† Smith and Benitez (1955).

‡ Strain, Thomas and Katz (1963).

** Strain (1958).

Codium, and the green layer of *Favia* separated into two zones on the thin layer system: one major fast-running yellow zone corresponding to α -carotene ($R_f = 0.67$), and one minor slower-running orange zone corresponding to β -carotene ($R_f = 0.41$). The presence of α -carotene as the major hydrocarbon carotenoid was therefore established in the three algae by the thin layer technique.

Chlorophyll *a* and *b* ratios were determined in *Halimeda*, *Codium*, and the green layer. The chlorophyll *b* content was relatively high in the three algae, being two-thirds that of chlorophyll *a* (chlorophyll *b*:chlorophyll *a* = 0.66, 0.67, and 0.79 in *Halimeda*, *Codium*, and the green layer, respectively). In higher plants and the green algae chlorophyll *b* is normally one-third that of chlorophyll *a* (*b*:*a* = 0.3).

DISCUSSION

The pigments of representative members of the order Siphonales have been studied by Strain (1958, 1965). He found that algae of this order within the Chlorophyceae contained the normal pigments of the green algae and higher plants—having chlorophylls *a* and *b* and the xanthophylls lutein (with or without zeaxanthin), violaxanthin and neoxanthin. In addition, these algae contained some special carotenoids, namely the pink-orange xanthophylls siphonein and siphonaxanthin, and α -carotene accompanied by small amounts of the β -isomer. The two free-living members of the Siphonales studied here showed this general pattern, although the full complement of xanthophylls expected was not present in either organism. *Halimeda* contained the yellow xanthophylls lutein, violaxanthin and neoxanthin, but only one pink-orange xanthophyll, siphonein, whereas *Codium* possessed both siphonein, siphonaxanthin and neoxanthin, but lacked lutein and violaxanthin. Both algae contained α -carotene as the major carotene, with small quantities of the β -isomer.

The green subsurface algal layer (*Ostreobium*) in the brain coral *Favia* showed a similar carotenoid pattern. α -carotene and siphonein were major components, with traces of siphonaxanthin; of the yellow xanthophylls, only neoxanthin was detected. A prominent unidentified yellow-orange xanthophyll, with absorption maxima at 470 and 449 nm in ethanol, was also present. From the evidence it appears that the filamentous alga *Ostreobium* inhabiting the brain coral *Favia* is appropriately grouped with the Siphonales, without however possessing the full complement of xanthophylls which have been described for the group. Strain (1965) examined 14 members of the Siphonales, and found only one species (*Caulerpa filiformis*) in which the full complement of pigments was not present. In this species both siphonein and siphonaxanthin were missing.

Both *Halimeda*, *Codium*, and the green layer (*Ostreobium*) contained relatively large amounts of chlorophyll *b*, approaching two-thirds to three-quarters the content of chlorophyll *a*. This is in contrast to other members of the Chlorophyceae and higher plants, where chlorophyll *b* is only one-third that of chlorophyll *a*. A wider survey would be needed to ascertain whether this high proportion of chlorophyll *b* is a characteristic of the Siphonales.

Two samples of the *Favia* green layer were extracted—one from 2–3 cm. below the surface of the coral, and the other from a depth of 6 cm. The first sample showed no trace of chlorophyll decomposition products, indicating that the algae

were in a physiologically healthy state. The second sample, taken deep within the coral, showed small zones of chlorophyll decomposition products (pheophytins, chlorophyllides, and pheophorbides). It is evident that in the very deep layers the cells eventually become moribund, with consequent decomposition of the chloroplast pigments.

The pigment evidence provides some guidance to the taxonomic affinities of *Ostreobium*. The presence of chlorophylls *a* and *b* definitely places this alga within the Chlorophyta, and excludes membership within the chlorophyll *c*-containing Chrysophycophyta. Furthermore, *Ostreobium* contains siphonein and siphonaxanthin, two xanthophylls which are found only in members of the Siphonales. On the basis of present evidence, it therefore seems appropriate to group *Ostreobium* within the order Siphonales in the class Chlorophyceae.

SUMMARY

1. The photosynthetic pigments of the green subsurface layer (*Ostreobium*) of the brain coral *Favia* were studied by two-dimensional paper chromatography. The pigments found were chlorophylls *a* and *b*, α - and β -carotene, siphonein, traces of siphonaxanthin and neoxanthin, and an unknown yellow-orange xanthophyll.

2. The pigment composition of *Ostreobium* closely resembled that of two members of the Siphonales, *Halimeda* and *Codium*. Therefore, this alga may be appropriately grouped within the Siphonales.

3. The three algae, *Halimeda*, *Codium*, and *Ostreobium* contained a high proportion of chlorophyll *b*, from two-thirds to three-quarters that of chlorophyll *a*.

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PHOTOSYNTHETIC PIGMENTS OF SYMBIOTIC DINOFLAGELLATES (ZOOXANTHELLAE) FROM CORALS AND CLAMS¹

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The brown symbiotic algae, or zooxanthellae, that live in great numbers in the tissues of some marine invertebrates, are considered on morphological evidence to be closely related to the dinoflagellates (Klebs, 1884; Kawaguti, 1944; Pringsheim, 1955). In the animal tissues the cells are in a non-motile vegetative state, without flagella or well-defined girdle, but upon isolation from the host tissue some zooxanthellae develop into typical gymnodinoid swimmers (Zahl and McLaughlin, 1957; McLaughlin and Zahl, 1959). Freudenthal (1962) has studied transformations in great detail in zooxanthellae from the jellyfish *Cassiopea* sp., and he has created a new genus, *Symbiodinium microadriaticum*, within the Dinophyceae, to accommodate this zooxanthella form. Droop (1963), however, has pointed out that "it is by no means certain that all zooxanthellae are dinoflagellates," and McLaughlin and Zahl (1966) warned that "caution must be exercised in the matter of speciation."

Knowledge of the photosynthetic pigment composition of algae can, when taken together with morphological evidence, provide a firm basis for the recognition of group affinities. This biochemical approach becomes increasingly powerful as gaps in our knowledge of the pigment composition of rare as well as known algal groups are filled. Studies of zooxanthellae pigments have, however, been relatively few. Present knowledge of the pigment composition of zooxanthellae rests largely upon studies by Strain, Manning, and Hardin (1944) of the pigment composition of the free-living dinoflagellate *Peridinium cinctum* and the endosymbiont of the sea-anemone *Bunodactis xanthogrammica* (now *Anthopleura xanthogrammica*; cf. Ricketts and Calvin, 1962). Using chromatography on columns of powdered sucrose they found that the pigment composition of the zooxanthellae exactly paralleled that of *Peridinium* in possessing chlorophylls *a* and *c*, β -carotene, and the xanthophylls peridinin and neo-peridinin, diadinoxanthin and dinoxanthin. It was concluded from the pigment evidence that the *Anthopleura* zooxanthellae and the dinoflagellate *Peridinium* "may belong to the same or related plant groups." However, identity of the former as cryptomonads from morphological evidence could not be excluded at the time. Subsequent biochemical analyses of cryptomonads by Haxo and Fork (1959) and Mallams *et al.* (1967) showed that these algae possess a unique com-

¹ Most of the experimental work described in this paper was carried out on board the University of California research vessel, R/V "Alpha Helix," in Princess Charlotte Bay, North Queensland, Australia from July to September, 1966. The Expedition was supported by the U. S. A. National Science Foundation. These studies were carried out in collaboration with Drs. P. Halldal and K. Shibata.

plement of biliproteins and carotenoids, and thus any relationship between zooxanthellae and cryptomonads seems now to be completely excluded.

Earlier, Heilbron, Jackson and Jones (1935) crystallized a red-orange pigment from the sea-anemone *Anemonia sulcata* which they called sulcatoxanthin. Strain *et al.* (1944) recognized that sulcatoxanthin was probably identical with peridinin, especially since this anemone is known to contain large numbers of zooxanthellae (Stephenson, 1935).

During the present Expedition¹ a unique opportunity was provided to examine the photosynthetic pigments of the exotic and little studied tridacnid clam zooxanthellae, and an assortment of coral endosymbionts. The pigments were screened by a simple two-dimensional paper chromatographic method (Jeffrey, 1961), which gave reliable separations of the chlorophylls and carotenoids, and which could be used, together with spectral analysis, as a simple aid to the identification of the pigments present. The pigments of the dinoflagellates *Amphidinium* and *Gymnodinium* spp. were used as standard reference material. The zooxanthellae from

TABLE I
Corals used in zooxanthellae survey

<i>CLASS</i> : Anthozoa; <i>SUB-CLASS</i> : Zooantharia; <i>ORDER</i> : Scleractinia
i) <i>Acropora</i> sp. ii) <i>Seriatopora</i> sp. iii) <i>Favia</i> sp. iv) <i>Fungia</i> (two species) v) <i>Pocillopora</i> sp.
<i>SUB-CLASS</i> : Alcyonaria; <i>ORDER</i> : Coenothecalia vi) <i>Helipora</i> sp.
<i>SUB-CLASS</i> : Alcyonaria; <i>ORDER</i> : Alcyonacea vii) <i>Xenia</i> sp. (two species)
<i>CLASS</i> : Hydrozoa; <i>ORDER</i> : Milleporina viii) <i>Millepora</i> sp.

five species of tridacnid clams, eight zooantharian and alcyonarian corals and one hydrozoan coral were found to have an identical pigment pattern to that of the dinoflagellate *Amphidinium*. The organisms contained chlorophylls *a* and *c*, β -carotene, peridinin and neo-peridinin, dinoxanthin and diadinoxanthin, and three other unidentified xanthophylls present in very small quantities.

MATERIALS AND METHODS

MATERIALS

Clams and corals were collected from the Great Barrier Reef just outside Princess Charlotte Bay, North Queensland. Species of tridacnid clams studied were *Tridacna crocea*, *Tridacna gigas*, *Tridacna squamosa*, *Tridacna deresa*, *Tridacna maxima*, and *Hippopus hippopus*. Identifications were made according to the descriptions provided by Rosewater (1965). Nine corals from both major classes and sub-classes were studied (Table I).

METHODS

A. *Preparation of zooxanthellae*

Zooxanthellae were isolated from clam mantle tissue and corals according to methods worked out by Muscatine (1967).

1. *Clams*. Mantle tissue containing zooxanthellae was excised, and freed as much as possible from supporting muscle tissue. The mantle was rinsed several times in filtered sea water, and cut into small pieces. The tissue was homogenized in sea water in a Waring Blendor for one minute, and the resulting suspension was filtered through six layers of cheesecloth, to free the cells from tissue debris. The dark brown suspension of algal cells was washed three times by centrifuging in filtered sea water at about 2500 *g* for five minutes. The cells were finally resuspended in sea water to a homogeneous suspension. Packed cells could be stored frozen without pigment deterioration for several weeks.

2. *Corals*. Zooxanthellae from hard corals were released by crushing the tissue into small pieces in aluminium foil with hammer or pliers, and rinsing continuously in sea water to wash out the cells. The suspension was filtered through cheesecloth, and the cells washed as before by centrifuging 2-3 times in filtered sea water.

Zooxanthellae from the tentacles of soft corals (*Xenia* spp.), and from the tentacles of *Fungia*, were collected by excising the tentacles and grinding them gently in sea water in a glass Potter-Elvehjem homogeniser. The zooxanthellae, which were released intact by this treatment, were filtered through cheesecloth, and washed by centrifugation as above. In every case, beautiful, clean suspensions of undamaged zooxanthellae were obtained.

B. *Preparation of pigment extracts*

Zooxanthellae were extremely difficult to extract in either acetone or methanol at room temperature. Pigments were readily released, however, if the packed cells were first suspended in a little distilled water, and frozen for 4-8 hours before methanol extraction. This procedure apparently weakened the tough cell wall, and allowed subsequent extraction to proceed readily, without alteration of the pigments. The cells were extracted about 2-3 times in methanol, until the residue was colorless. The methanol extract was clarified by centrifugation, mixed with an equal volume of diethyl ether, and shaken once or twice with a volume of 10% NaCl solution 5-10 times that of the methanol-ether extract. This saline washing caused the pigments to migrate to the ether layer, and methanol and methanol-soluble impurities were washed out in the aqueous phase. The ether layer was then concentrated for chromatography by evaporation under a stream of nitrogen. The ether extracts could be stored in the deep freeze for several days without deterioration of the pigments.

Some of the hard corals were extremely difficult to crush for the collection of zooxanthellae. Small pieces of these corals were extracted "whole" with methanol for several hours at room temperature, in the presence of a little MgCO₃, to prevent acidification. Extraction was continued until the coral skeleton was colorless. Pigments were not damaged by this treatment, as was shown by subsequent chromatography.

The dinoflagellates, *Amphidinium* and *Gymnodinium*, were used as standard sources of dinoflagellate pigments. *Gymnodinium* was used mainly as a source of standard dinoxanthin, since in *Amphidinium* the amount of this pigment was extremely small. Cultures were grown in Medium f (Guillard; see Grant, 1967) for about two weeks at 5000 lux. The cells were harvested by continuous centrifugation, and pigments were readily extracted from the packed cells with small volumes of 90% acetone. After 2-3 extractions the residue was colorless. Pigments were transferred from acetone to ether as above, and used directly for chromatography. Samples of *Amphidinium* cells were freeze-dried, for dispatch from Cronulla to Princess Charlotte Bay. The dry powder could be stored in the dark at room temperature for several weeks, without deterioration of the pigments.

C. Paper chromatography

Pigment extracts were chromatographed on 22-cm. squares of Whatman No. 3 (or No. 3 Mm paper in two dimensions, according to the method of Jeffrey (1961)). Solvents used were AR Grade, and were not further purified. Solvent systems used were 4% *n*-propanol in light petroleum (60-80° C.) for the first dimension, and 30% CHCl₃ in light petroleum for the second dimension. Absorption spectra of pigment fractions were obtained by running a number of chromatograms, and eluting the pigment spots. Absorption spectra were taken with a Beckman DB spectrophotometer (for work done on the "Alpha Helix"), a Unicam SP 700 (for work done subsequently at Cronulla), and a Carey Model 14 (for work done subsequently at La Jolla). The latter instruments were calibrated with Hg lines and were considered accurate to 1 nm.

D. Quantitative determination of pigments

1. *Spectrophotometric.* Routine quantitative determinations of chlorophylls *a* and *c* were carried out on pigment extracts, either in methanol, 90% acetone, or in diethyl ether. Extinctions were measured at the red maxima of the two chlorophylls, and the concentration calculated from the provisional equations of Humphrey and Jeffrey (in preparation).

$$\begin{array}{l}
 \text{In ether:} \\
 \text{In 90\% acetone:} \\
 \text{In methanol:}
 \end{array}
 \begin{array}{l}
 \left[\begin{array}{l}
 \text{chl } a = 10.5 E_{662} - 1.0 E_{628} \\
 \text{chl } c = -8.1 E_{662} + 64.3 E_{628}
 \end{array} \right. \\
 \left[\begin{array}{l}
 \text{chl } a = 13.31 \times E_{670} - 0.27 \times E_{630} \\
 \text{chl } c = -8.37 \times E_{668} + 51.72 \times E_{670}
 \end{array} \right. \\
 \left[\begin{array}{l}
 \text{chl } a = 13.8 E_{668} - 1.3 E_{635} \\
 \text{chl } c = -14.1 E_{668} + 67.3 E_{635}
 \end{array} \right.
 \end{array}$$

where chl = concentration of chlorophyll in ug./ml. and E = extinction in liters/gm. cm. in a 1-cm. cell.

2. *Chromatographic.* The percentage composition of each carotenoid fraction was determined by running several chromatograms simultaneously, and eluting the spots immediately in the appropriate solvent. Solutions were made up to a

measured volume, centrifuged to remove any paper fibers, and the extinctions read without delay. In this, and in all the chromatographic work, operations were carried out in dim light, or in complete darkness. Photo-decomposition of pigments was thus kept at a minimum.

Concentrations of pigments were calculated using the following extinction coefficients:

Chlorophyll <i>a</i> in acetone	90 l./gm.cm.	Vernon (1960)
Chlorophyll <i>c</i> in methanol	15.2 l./gm.cm.	Jeffrey (1963)
Carotene in ether	250.5 l./gm.cm.	Goodwin (1955)
Peridinin in ethanol	132.5 l./gm.cm.	present work

The concentrations of all other carotenoids, whose extinctions were unknown, were calculated using the extinction coefficient of carotene.

E. Thin layer chromatography

Since paper chromatography does not resolve carotene isomers, thin layers of $\text{Al}_2\text{O}_3 + \text{MgO}$ (3:1 w/w) were used to characterize the carotene fractions. Using as solvent 4% ethyl acetate in hexane (Chapman, 1966), α - and β -carotenes were separated with R_f values of 0.67 and 0.41. α -Carotene for reference was obtained from the cryptomonad *Chloromonas* sp., and β -carotene from the green flagellate, *Dunaliella tertiolecta*.

The pink-orange xanthophyll (fraction 9), which remained at the origin in the paper chromatographic system was separated from chlorophyll *c* by chromatography of the "origin material" on thin layers of polyethylene, using 90% acetone as solvent. The pink xanthophyll ran just behind the solvent front ($R_f = 0.95$) with chlorophyll *c* at R_f values of 0.3 and 0.4. The xanthophyll was eluted with ethanol, for spectral analysis.

A thin layer chromatography system, which completely duplicated the paper chromatography method, and which gave even better resolution of all the dino-flagellate pigments, was subsequently developed after returning from the Expedition. This method (to be described in detail in a separate communication) uses plates of specially prepared sucrose, with 0.6% *n*-propanol in petroleum ether (60–80° C.) and 12% chloroform in petroleum ether, as the two-dimensional solvent system. Extracts of *Amphidinium*, and extracts of zooxanthellae from the clam *Tridacna crocea* and the coral *Pocillopora* were chromatographed using this system.

F. Crystallization of peridinin

Approximately 6 mg. crystalline peridinin were isolated from *Tridacna gigas* as follows. The mantle tissue was excised from a healthy giant clam, freed substantially of colorless animal tissue and immediately deep frozen. The frozen zooxanthellae-laden mantle tissue weighing some two Kg. was chopped into small pieces and exhaustively extracted in the cold with 95% ethanol. The total pigment was transferred to benzene with the addition of saturated NaCl solution, and the washed and dried pigment solution evaporated to dryness *in vacuo* at 30° C. The pigment was dissolved in 80% aqueous methanol, and the

initial separation of the peridinin from other pigments and colorless contaminants was made by passage of the methanolic solution through powdered polyethylene. The peridinin fraction was recovered as the leading zone. Crude peridinin from several such columns was transferred to benzene and then chromatographed on columns of powdered CaCO_3 , developed with 1–2% acetone in benzene, according to the method of Pinckard *et al.* (1953). Prior to crystallization, the almost pure peridinin was rechromatographed on thin layer plates of silica gel G employing 30% acetone in hexane as developing solvent.

Crystallization of peridinin was carried out according to the procedures of Pinckard *et al.* (1953), once from benzene-hexane and once from ether-hexane mixtures. The homogeneous microcrystalline product was scarlet to warm reddish brown in color, disc-shaped and free of colorless contaminants. After drying *in vacuo* at 65.5°C . the product weighed a little over 6 mg.

The crystals were readily soluble in methanol, ethanol, acetone, ether, and benzene, and virtually insoluble in hexane. Methanolic solutions were found

TABLE II
Extinction coefficients of peridinin isolated from Tridacna gigas

Solvent	Maxima (nm)	Extinction coefficient $E_{1\text{cm.}1\%}$
Acetone (single broad maximum)	466	1340
90% Acetone (single broad maximum)	469	1330
Methanol (absolute) (single broad maximum)	469	1360
Ethanol (absolute) (single broad maximum)	472	1325
Pyridine (single broad maximum)	475	1180
Diethyl ether	454,* 475	1450
Benzene	467,* 494	1290
Hexane-ether (9:1)	454,* 483	1470
Chloroform	470,* 490	1290
Hexane	454, 484	—

* Extinctions taken at these maxima.

to be strongly hypophasic. The phase distribution in the system 65% MeOH/hexane was 96:4.

The melting point of the crystals was determined in capillaries sealed under N_2 employing a Thomas Hoover capillary melting point apparatus. Averaged values for two determinations were: softened at 130.7°C . (cor.) and complete melt at 134.9°C . (cor.). These values agree well with the total melt point of 130°C . reported by Heilbron *et al.* (1935) for sulcatoxanthin (*i.e.*, peridinin) from *Anemonia sulcata*.

For estimation of the extinction coefficient 1.040 mg. peridinin were dissolved in 200 ml. acetone and spectral absorption determined in a 1-cm. path length cuvette in the Cary 14 Spectrophotometer. From this measurement the $E_{1\text{cm.}1\%}$ at 466 $m\mu$, the wave-length of maximum absorption in acetone, was calculated to be 1340. The ash content of the sample is not known and the extinction coefficient is considered to be minimal. This value is substantially higher than the

extinction coefficient of 840 (acetone) reported by Parsons and Strickland (1963) for peridinin from *Amphidinium carterci*. Relative extinctions of peridinin in other solvents were determined by evaporating aliquot samples of the above acetone solution to dryness in a rotary evaporator *in vacuo* at room temperature and dissolving in the appropriate solvent. Extinction values for peridinin in different solvents, and absorption maxima, are given in Table II.

By way of further characterization of peridinin from *Tridacna gigas*, iodine-catalyzed trans-cis isomerizations similar to those performed by Pinckard *et al.*

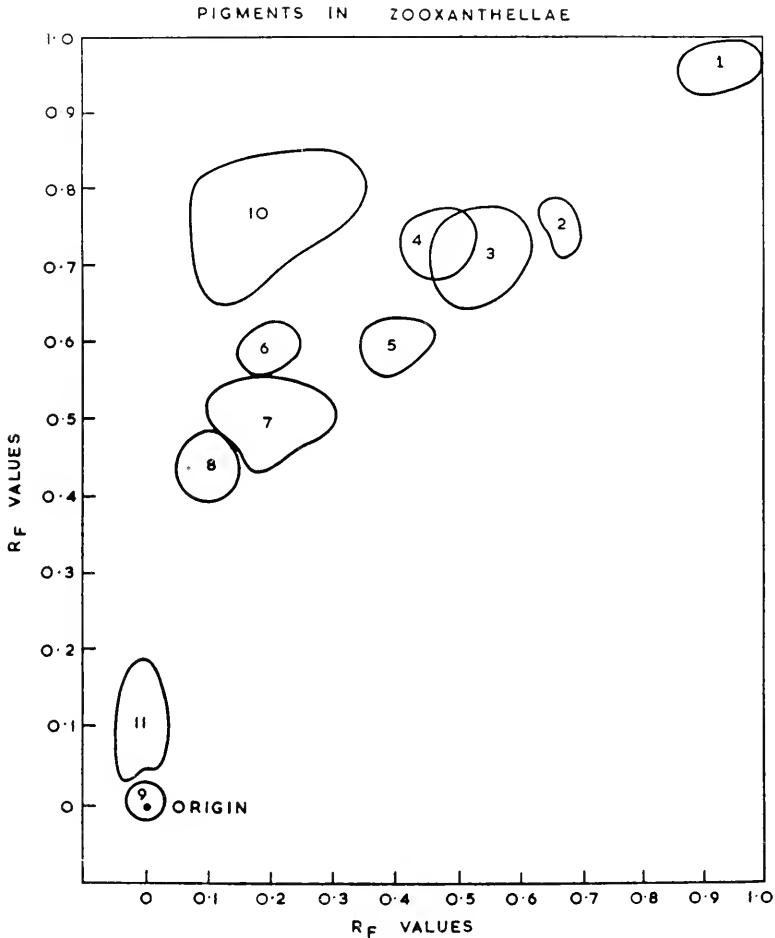


FIGURE 1. Two-dimensional paper chromatogram of pigments in zooxanthellae from *Tridacna crocea*. Chromatographic solvent systems: 1st dimension, 4% *n*-propanol in petroleum ether (60–80°); 2nd dimension, 30% chloroform in petroleum ether. 1, β -carotene (orange). 2, Unknown (pale orange). 3, Diadinoxanthin (yellow). 4, Dinoxanthin (yellow). 5, Unknown (yellow). 6, Neo-dinoxanthin (yellow). 7, Peridinin (brick red). 8, Neo-peridinin (brick red). 9, Unknown (pink orange). 10, Chlorophyll *a* (blue green). 11, Chlorophyll *c* (light green).

(1953) for crystalline peridinin isolated from *Prorocentrum micans* were carried out. In all respects examined the properties of the two preparations were identical, *e.g.*, spectra before and after iodine catalysis of benzene solutions; number, adsorption position and spectral properties of isomers separable on CaCO_3 columns; and percentage composition of an equilibrium mixture of stereo-isomers (57% all-trans and 43% combined cis-isomers).

For comparison with one of the original sources used by Strain *et al.* (1944), peridinin was isolated from zooxanthallae from the sea anemone *Anthopleura xanthogrammica*. (Live sea anemones were collected intertidally at La Jolla.) Peridinins from the sea anemone and *Tridacna gigas* were found to be chromatographically homogeneous in the thin layer system, silica gel G-30% acetone in hexane.

RESULTS

Figure 1 shows a typical two-dimensional paper chromatogram of the pigments of zooxanthellae from *Tridacna crocea*. Identical chromatograms were

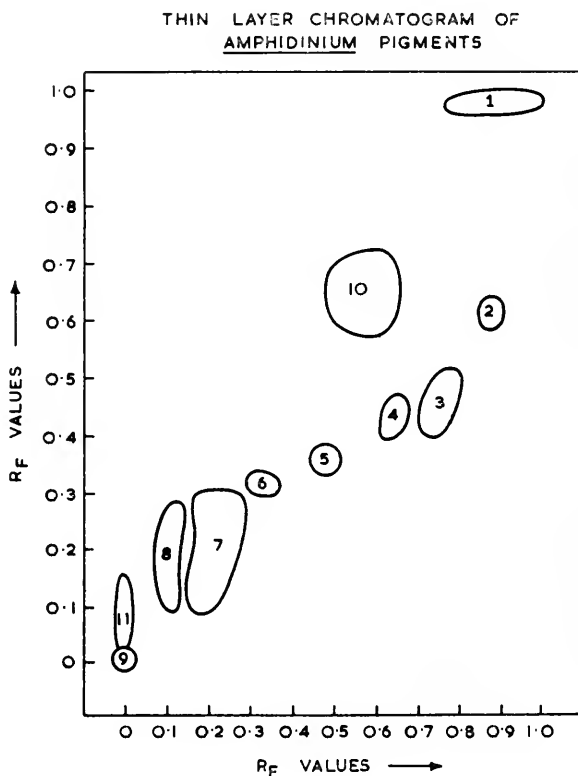


FIGURE 2. Sucrose thin layer chromatogram of pigments in the dinoflagellate *Amphidinium* sp. Chromatographic solvent systems; 1st dimension, 0.6% *n*-propanol in petroleum ether (60–80°); 2nd dimension, 12% chloroform in petroleum ether. Pigment fractions as in Figure 1.

also obtained from the other four clam zooxanthellae, the nine coral zooxanthellae, and the dinoflagellate *Amphidinium*. In every case chlorophyll *a* was accompanied by chlorophyll *c*, and the major carotenoids were always β -carotene, peridinin, neo-peridinin, dinoxanthin, and diadinoxanthin. Average R_f values for dinoflagellate pigments in the two solvent systems are given in Table III. The carotene fraction was shown to consist only of the β -isomer by chromatography of the carotene fraction from zooxanthellae and *Amphidinium* on thin layers of alumina and magnesium oxide.

Figure 2 shows a typical chromatogram of dinoflagellate pigments separated on specially prepared thin layers of powdered sucrose. Again, identical patterns of pigments were obtained with *Amphidinium* and zooxanthellae extracts from the clam *Tridacna crocea* and the coral *Pocillopora*. Since the thin layer method will be described in detail in a separate communication, only results obtained with the paper method, which was used on the Expedition, will be presented here.

TABLE III

R_f values of pigment fractions from zooxanthellae and *Amphidinium* using paper chromatography

Fraction	Pigment	Color	R_f Values	
			4% <i>n</i> -propanol in pet. ether	30% CHCl ₃ in pet. ether
1	Carotene	Orange	0.96	0.93
2	Unknown	Pale orange	0.75	0.67
3	Diadinoxanthin	Yellow	0.71	0.54
4	Dinoxanthin	Yellow	0.73	0.48
5	Unknown	Pale yellow	0.59	0.40
6	Neo-dinoxanthin	Pale yellow	0.59	0.19
7	Peridinin	Brick red	0.49	0.19
8	Neo-peridinin	Brick red	0.43	0.10
9	Unknown	Pink-orange	0	0
10	Chlorophyll <i>a</i>	Blue-green	0.76	0.21
11	Chlorophyll <i>c</i>	Light green	0.10	0

Absorption spectra of all pigment zones were analyzed in detail in zooxanthellae from the two clams *Tridacna crocea* and *Hippopus hippopus*, the coral *Pocillopora* and the dinoflagellate *Amphidinium*. The absorption maxima are listed in Table IV, and are compared with absorption maxima of pigments from the dinoflagellate *Peridinium cinctum* given by Strain, Manning and Hardin (1944).

Absorption properties of peridinin from *Tridacna gigas* and *Amphidinium* were checked on more accurate spectrophotometers after the Expedition. The values for peridinin in ethanol obtained on the Expedition with the Beckman DB spectrophotometer were from 473–475 nm (Table IV). Using the same paper chromatography method, the maximum of peridinin in ethanol from *Amphidinium* was 474 nm (Unicam SP700), and from *Tridacna gigas* was 473 nm (Carey Model 14). The absorption maximum of *Tridacna gigas* peridinin prepared as an all-trans sample and then crystallized, was 473 nm; prepared as above but rechromatographed on CaCO₃ to obtain the fresh all-trans fraction was 472 nm.

TABLE IV

Absorption spectra of pigments from zooxanthellae of corals, clams, and dinoflagellates

Fraction	Organism	Absorption maxima (nm)	Solvent	Identification
Fraction 1 (orange)	<i>Tridacna crocea</i>	426, 448, 475	diethyl ether	<i>β-Carotene</i>
	<i>Hippopus hippopus</i>	428, 449, 475	diethyl ether	
	<i>Pocillopora</i> sp.	428, 448, 475	diethyl ether	
	<i>Amphidinium</i> sp.	429, 450, 475	diethyl ether	
	<i>Peridinium cinctum</i> *	429, 450, 478 not given	hexane	
Fraction 2 (pale orange)	<i>Tridacna crocea</i>	429, 450, 476	ethanol	<i>Unknown</i>
	<i>Hippopus hippopus</i>	429, 450, 478	ethanol	
	<i>Pocillopora</i> sp.	427, 450, 477	ethanol	
	<i>Amphidinium</i> sp.	429, 452, 477	ethanol	
	<i>Peridinium cinctum</i> *	not present		
Fraction 3 (yellow)	<i>Tridacna crocea</i>	425, 447, 477	ethanol	<i>Diadinoxanthin</i>
	<i>Hippopus hippopus</i>	426, 448, 477	ethanol	
	<i>Pocillopora</i> sp.	425, 447, 476.5	ethanol	
	<i>Amphidinium</i> sp.	425, 448, 478	ethanol	
	<i>Peridinium cinctum</i> *	448, 478	ethanol	
Fraction 4 (yellow)	<i>Tridacna crocea</i>	418, 442, 470	ethanol	<i>Dinoxanthin</i>
	<i>Hippopus hippopus</i>	418, 441, 468	ethanol	
	<i>Pocillopora</i> sp.	418, 441, 469	ethanol	
	<i>Amphidinium</i> sp.	418, 443, 472	ethanol	
	<i>Peridinium cinctum</i> *	441.5, 471	ethanol	
Fraction 5 (pale yellow)	<i>Tridacna crocea</i>	408, 427, 455	ethanol	<i>Unknown</i>
	<i>Hippopus hippopus</i>	410, 428, 455	ethanol	
	<i>Pocillopora</i> sp.	408, 427, 455	ethanol	
	<i>Amphidinium</i> sp.	408, 427, 455	ethanol	
	<i>Peridinium cinctum</i> *	not present		
Fraction 6 (pale yellow)	<i>Tridacna crocea</i>	425, 441, 467	ethanol	<i>Neo-dinoxanthin</i>
	<i>Hippopus hippopus</i>	425, 440, 466	ethanol	
	<i>Pocillopora</i> sp.	420, 440, 465	ethanol	
	<i>Amphidinium</i> sp.	422, 440, 467	ethanol	
	<i>Peridinium cinctum</i> *	438, 466	ethanol	
Fraction 7 (brick red)	<i>Tridacna crocea</i>	475	ethanol	<i>Peridinin</i>
	<i>Hippopus hippopus</i>	473	ethanol	
	<i>Pocillopora</i> sp.	475	ethanol	
	<i>Amphidinium</i> sp.	475	ethanol	
	<i>Peridinium cinctum</i> *	475	ethanol	

* Data for *Peridinium cinctum* taken from Strain, Manning, and Hardin (1944).

† Present, but insufficient for spectral analysis.

TABLE IV—Continued.

Fraction	Organism	Absorption maxima (nm)	Solvent	Identification
Fraction 8 (brick red)	<i>Tridacna crocea</i>	468	ethanol	<i>Neo-peridinin</i>
	<i>Hippopus hippopus</i>	465	ethanol	
	<i>Pocillopora</i> sp.	465	ethanol	
	<i>Amphidinium</i> sp.	467	ethanol	
	<i>Peridinium cinctum</i> *	464	ethanol	
Fraction 9 (pink-orange)	<i>Tridacna crocea</i> †	460-466 not present	ethanol	<i>Unknown</i>
	<i>Hippopus hippopus</i> †			
	<i>Pocillopora</i> sp.†			
	<i>Amphidinium</i> sp.			
	<i>Peridinium cinctum</i> *			
Fraction 10 (blue-green)	<i>Tridacna crocea</i>	409, 428, 661	ether	<i>Chlorophyll a</i>
		410, 428, 662	acetone	
	<i>Hippopus hippopus</i>	409, 428, 663	ether	
	<i>Pocillopora</i> sp.	409, 428, 663	acetone	
	<i>Amphidinium</i> sp.	408, 428, 660	ether	
	<i>Peridinium cinctum</i> *	not given		
Fraction 11 (light green)	<i>Tridacna crocea</i>	448, 582, 634	methanol	<i>Chlorophyll c</i>
	<i>Hippopus hippopus</i>	448, 583, 634	methanol	
	<i>Pocillopora</i> sp.	448, 584, 634	methanol	
	<i>Amphidinium</i> sp.	451, 584, 635	methanol	
	<i>Peridinium cinctum</i> *	not given		

TABLE V

Percentage composition of carotenoids in zooxanthellae from *Tridacna crocea*, and the dinoflagellate *Amphidinium* sp.

Fraction number	Pigments	% of Total carotenoids	
		<i>Tridacna crocea</i>	<i>Amphidinium</i> sp.
1	β -carotene	3.0	2.5
2	Unknown	1.5	1.7
3 + 4	Diadinoxanthin and dinoxanthin	12.6	10.5
5	Unknown	1.3	0.3
6	Neo-dinoxanthin	3.1	0.6
7 + 8	Peridinin and neo-peridinin	77.0	84.0
9	Unknown	1.5	0.4
	Total	100.0	100.0

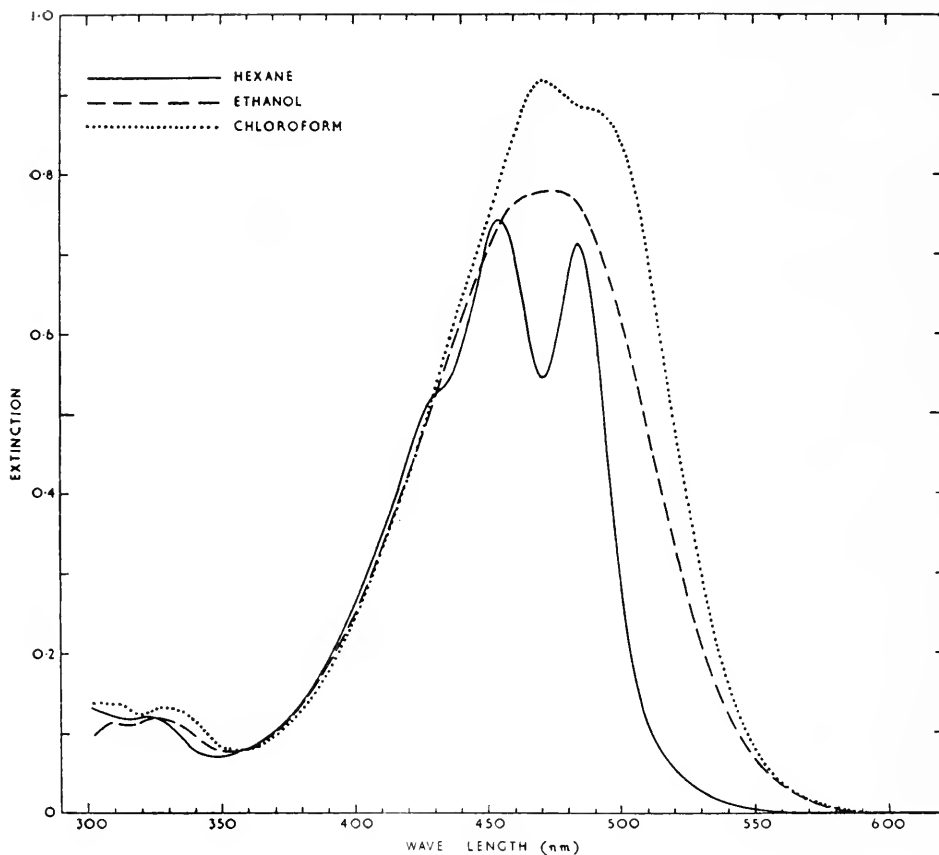


FIGURE 3. Absorption spectra of all-trans peridinin isolated from *Tridacna gigas*.

Peridinin apparently isomerizes fairly rapidly and crystallization of the all-trans fraction may not have excluded traces of the cis-isomers. Cis-trans isomerization of peridinin in benzene solution with iodine caused the maxima to shift from 496 and 468 nm (all-trans) to 493 and 466 nm (equilibrium mixture) after 10 minutes. There was no further spectral change at 120 minutes.

Full absorption curves of peridinin in ethanol, chloroform and hexane are shown in Figure 3. This peridinin was a freshly prepared all-trans fraction from *Tridacna gigas* mantle tissue. Peridinin isolated by paper chromatography from *Pocillipora zooxanthallae* gave similar absorption curves in the three solvents.

The relative proportions of the carotenoids were analyzed by paper chromatography in zooxanthellae from *Tridacna crocea* and in the dinoflagellate *Amphidinium*. Table V shows that in both cases the major xanthophyll fraction, peridinin and neo-peridinin, constituted some 77–84% of the total carotenoids. The other xanthophyll fractions were, by contrast, only a few per cent of the total.

The relative concentrations of chlorophylls *a* and *c* in the zooxanthellae are

given in Table VI. In coral zooxanthellae, the content of chlorophyll *c* was about one-tenth that of chlorophyll *a*. In zooxanthellae from clams, and in the dinoflagellates *Amphidinium* and *Gymnodinium*, the chlorophyll *c* content approached two-thirds that of chlorophyll *a*.

No animal pigments could be detected in the clam mantle tissue itself, identical chromatograms and identical absorption spectra being obtained both with extracts of whole mantle tissue and with isolated zooxanthellae. However, R_f values of the dinoflagellate pigments extracted directly from the mantle were always a little higher than those from zooxanthellae processed separately. That the increased R_f values were due to animal tissue factors extracted with the zooxanthellae was indicated by the effect of adding colorless methanol-acetone extracts of white mantle muscle (without zooxanthellae) to extracts of isolated zooxanthellae cells. R_f values of all pigment fractions became increasingly higher

TABLE VI

Ratio of chlorophyll c:chlorophyll a in zooxanthellae from corals and clams

	Organism	Chlorophyll <i>c</i> : <i>a</i>
Corals	<i>Pocillopora</i> sp.	0.1
	<i>Fungia</i> sp.	0.07
	<i>Acropora</i> sp.	0.2
	<i>Millepora</i> sp.	0.06
Clams	<i>Tridacna crocea</i>	0.6
	<i>Tridacna gigas</i>	0.3
	<i>Hippopus hippopus</i>	0.6
Dinoflagellates	<i>Amphidinium</i> sp.	0.6
	<i>Gymnodinium</i> sp.*	0.8

* Taken from Jeffrey (1963).

with increasing amounts of animal extract present. This phenomenon, of the presence of colorless factors (probably lipids) influencing the R_f values of photosynthetic pigments, is well documented for other plant pigment systems (Sestak, 1958). Consistent R_f values were, however, always obtained from zooxanthellae which were first separated from the mantle tissue before extraction.

DISCUSSION

Strain, Manning and Hardin (1944) were the first to study in detail the photosynthetic pigments of the free-living dinoflagellates. Using cultures of *Peridinium cinctum*, and separating the pigments on columns of powdered sugar, the components found were chlorophylls *a* and *c*, β -carotene, a major brick-red xanthophyll peridinin and its neo-isomer, and four minor yellow xanthophylls, diadinoxanthin, dinoxanthin, neo-diadinoxanthin, and neo-dinoxanthin. These pigments

were also found in the dinoflagellates *Amphidinium carteri* (Parsons, 1961) *Prorocentrum micans* (Pinckard *et al.*, 1953) and *Gymnodinium* (Jeffrey, 1961) and in zooxanthellae from the sea anemone *Anthopleura xanthogrammica* (Strain, Manning and Hardin, 1944).

Peridinin and dinoxanthin are specific to the dinoflagellates, and have not been found in any other classes of algae. The other pigments (chlorophylls *a* and *c*, β -carotene and diadinoxanthin) are not by themselves taxonomically definitive pigments, since they also occur in other algal groups (Strain, 1958). In the present survey of photosynthetic pigments in zooxanthellae from corals and clams, peridinin was found in every organism tested, and this compound had the same chromatographic and absorption properties as peridinin isolated from the dinoflagellate, *Amphidinium*. In addition, dinoxanthin was found as a minor pigment in all zooxanthellae and dinoflagellate preparations, together with the full complement of chlorophylls and carotenoids originally described by Strain, Manning and Hardin (1944) for *Peridinium cinctum*. The absorption maxima of the pigment fractions from different zooxanthellae corresponded closely to each other and to the published maxima (Table IV). Where small discrepancies were present low extinctions were combined with broad peaks, with consequent difficulties in obtaining accurate maxima. Identification of pigment fractions from their spectral properties was, however, strengthened by the identical R_f values and chromatographic patterns which were obtained.

Peridinin isolated by paper chromatography showed absorption maxima closely similar to freshly prepared all-trans crystalline peridinin. The absorption maxima of peridinin in ethanol from *Tridacna gigas* were at 473 nm (paper chromatography), 473 nm (crystalline "all-trans"), and 472 nm (freshly prepared all-trans), using the Carey spectrophotometer in all cases. With the Beckman DB spectrophotometer, the range of values for peridinin isolated by paper chromatography was 473–475 nm (Table IV). Cis-trans isomerization studies in benzene solution showed that the all-trans fraction had absorption maxima at higher wave-lengths than the individual cis-isomers, or the equilibrium mixture.

The extinction coefficient in acetone of crystalline peridinin from *Tridacna gigas* was substantially higher than that obtained by Parsons and Strickland (1963) for peridinin isolated from *Amphidinium carterei*. The melting point, however, agreed well with the melting point of sulcatoxanthin (peridinin) isolated from the sea anemone, *Anemonia sulcata* (Heilbron *et al.*, 1935). Further, the isomerization behavior of *Tridacna gigas* peridinin resembled similar preparations obtained from *Prorocentrum micans* (Pinckard *et al.*, 1953).

In addition to the major dinoflagellate pigments described, three new xanthophylls were also found in extracts of zooxanthellae and *Amphidinium*. These were readily distinguished both on paper chromatograms and on the sucrose thin-layer plates. Absorption maxima of these fractions taken in ethanol did not appear to correspond to any previously described xanthophyll. Due to the very small quantities present, and the limited duration of the Expedition, it was not possible to analyze these pigment fractions further. Strain, Manning and Hardin (1944) noted a "flavoxanthin-like" xanthophyll in some of their fractions in amounts too small to be analyzed. This may correspond to one or more of the minor xanthophylls described here. Neodiadinoxanthin, found by Strain, Manning,

and Hardin (1944) in *Peridinium cinctum*, could not be located in any of the present preparations studied.

Quantitative data on the carotenoid composition of zooxanthellae and *Amphidinium* showed that the major xanthophyll fraction, peridinin and neoperidinin, constituted some 77–84% of the total carotenoids. Diadinoxanthin was always more prominent than dinoxanthin, and together the concentration was about 10–12%, whereas the other xanthophylls were present in very small quantities which varied from 0.3% to 3.0% of the total. Peridinin in these zooxanthellae (Halldal, 1968; Shibata and Haxo, unpublished data), as well as peridinin from *Gonyaulax polyedra* (Haxo, 1960) appears to have an accessory pigment function in photosynthesis, but the function of the minor carotenoid components, if any, is unknown.

The ratios of chlorophylls *a* and *c* varied depending on whether the zooxanthellae came from a clam or coral host. Clam zooxanthellae resembled the free-swimming dinoflagellates in having a chlorophyll *c* content almost two-thirds that of chlorophyll *a*, whereas in coral zooxanthellae the chlorophyll *c* content was only about one-tenth that of chlorophyll *a*. Further studies would be needed to ascertain whether this was a real difference between clam and coral zooxanthellae. Burkholder and Burkholder (1960) analyzed the chlorophyll content of some alcyonarian corals by spectrophotometric methods and found that chlorophyll *c* was equal to chlorophyll *a*. Using more accurate extinction coefficients the chlorophyll *c* was probably about one-half that of chlorophyll *a*.

No chlorophyll degradation products were ever present on chromatograms of freshly prepared zooxanthellae extracts. This suggests that not only were the extraction and chromatographic procedures reliable, but that the zooxanthellae populations were in a healthy state. Zooxanthellae breakdown must therefore occur at some other locus in the animal host than that sampled, or senescent cells may be discharged from the host tissues and do not accumulate. Traces of pheophytins, chlorophyllides and pheophorbides, indicating the presence of senescent cells, would have been readily detected on the paper chromatograms, had they been present.

Zooxanthellae were further indicated to have dinoflagellate affinities by the presence of gymnodinioid swimmers in suspensions of zooxanthellae which were left overnight in sea water. Although these were not studied in detail, they clearly resembled the swimmers described for *Cassiopea* zooxanthellae, by Freudenthal (1962).

The two-dimensional paper chromatography method provided a simple but useful screening technique for determining the pigment composition of a large range of zooxanthellae. The method was previously used to study the photosynthetic pigments of a variety of different classes of marine algae both in unialgal culture (Jeffrey, 1961; Jeffrey and Allen, 1964), and in natural phytoplankton populations (Jeffrey, 1965). However, for resolution and identification of very closely related carotenoids (*e.g.*, isomers of carotene) thin layer techniques using organic or inorganic adsorbents must be used (Chapman, 1966). In the present work the nature of the carotene fraction was established using thin layers of aluminium and magnesium oxide, and the unknown pink xanthophyll (fraction 9) was separated from chlorophyll *c* on thin layers of polyethylene. The sucrose thin-layer method

had a resolution similar to the separations obtained with the paper method, except that the separation of each component (particularly the xanthophylls) was more complete, and the method may be used for sensitive quantitative analyses.

The present investigation of photosynthetic pigments in clam and coral zooxanthellae establishes that these symbiotic algae contain the full and characteristic complement of dinoflagellate pigments. On the basis of present knowledge of pigment distribution within algal groups, designation of these zooxanthellae as dinoflagellates is clearly justified.

SUMMARY

1. The photosynthetic pigments of the brown symbiotic algae (zooxanthellae) isolated from five tridacnid clams and nine corals were found to be identical with the pigments of the dinoflagellate *Amphidinium*. Identifications were carried out by two-dimensional paper chromatography and by absorption spectrophotometry. Both zooxanthellae and dinoflagellates contained chlorophylls *a* and *c*, β -carotene, peridinin, neo-peridinin, dinoxanthin, neo-dinoxanthin, diadinoxanthin, and three minor xanthophyll fractions not previously described.

2. Peridinin, crystallized from *Tridacna gigas*, or isolated by paper chromatography from *Tridacna crocea*, *Pocillopora* or the dinoflagellate *Amphidinium*, showed similar absorption characteristics in different solvents. The extinction coefficient of crystalline peridinin in acetone was $E_{1\text{ cm.}}^{1\%}$ 1340.

3. Clam zooxanthellae (and the dinoflagellates *Amphidinium* and *Gymnodinium*) contained a much higher proportion of chlorophyll *c* than coral zooxanthellae.

4. No chlorophyll decomposition products were found in any freshly prepared zooxanthellae preparations.

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PACEMAKER PROPERTIES OF TUNICATE HEART CELLS

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The primary pacemaker regions are located at the cardiovascular junctions and they alternate in periods of activity which change the direction of peristalsis (see Kriebel, 1968a). It will be demonstrated below that isolated sectors of a cardiovascular junction show about equal pacemaker capabilities and when more than one region is independently active, arrhythmia results. The region which usually functions as the pacemaker was determined and its interaction with latent pacemaker regions is discussed below.

The myocardium is composed of a single layer of musculoendothelial cells, except for a line of undifferentiated cells located opposite the cardio-pericardial raphe; these undifferentiated cells resemble those forming the ring of undifferentiated cells at the ends of the heart (see Millar, 1953). Many investigators have demonstrated that narrow rings of tunicate hearts can beat rhythmically (Bancroft and Esterly, 1903; see Krijgsman, 1956, for a review). However, rings of the heart contain both the raphe and the undifferentiated line of cells, which raises the question whether the myocardial cells themselves have pacemaker properties or whether pacemaker activity is limited to the undifferentiated cells. To investigate this question pacemaker capabilities of pieces of myocardium were examined after both the undifferentiated line and the raphe had been removed.

METHODS

Hearts of adult *Ciona intestinalis* (from Woods Hole, Massachusetts and from California) were used for most experiments. For comparative purposes, hearts of adult *Chelysoma productum*, *Ascidia callosa* and *Corella willmeriana* (from Friday Harbor, Washington) were also used.

Heart contractions in intact animals were observed with the aid of a dissecting microscope and recorded on a kymograph drum. Dissections were performed in sea water at 10° C. *Ciona* heart action potentials were recorded *in situ* by applying suction electrodes to the raphe so that portions of both pericardium and myocardium were sucked into the electrode openings. Electrical activity of the pacemaker regions was recorded by placing as many as four electrodes on the myocardium near the cardiovascular junction. In a second method of recording action potentials, hearts were opened and positioned over a Plexiglass plate containing a row of small electrode openings spaced 1-mm. apart (see Kriebel, 1967a, for details).

Small pieces of heart were spread onto a microscope slide moistened with sea water between two ridges of petroleum jelly. A coverslip holding a drop of sea

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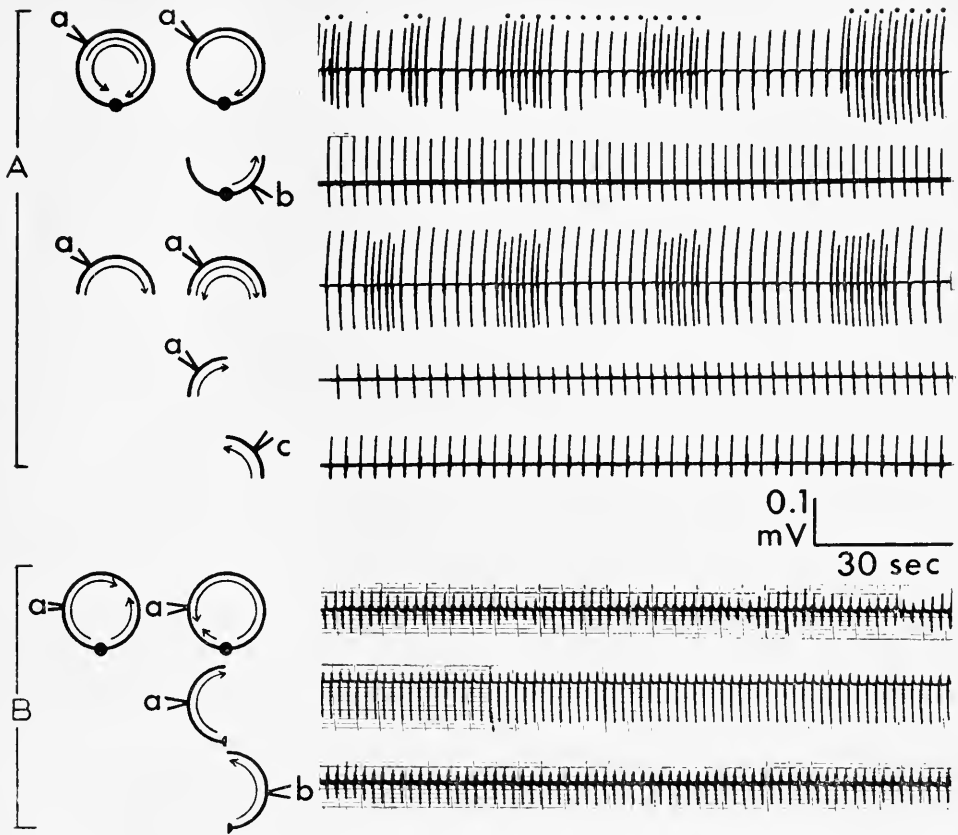


FIGURE 1. Action potentials recorded from isolated pacemaker regions. These hearts were cut in two and collapsed. To the left of each trace the pacemaker region is represented in cross-section either as a circle or, after sectioning longitudinally, as parts of a circle (the raphe is the dot). The electrode positions are alphabetized. The arrows represent directions of conduction which were determined with 3 additional electrodes (not shown). Two arrows passing each other indicate alternating directions of conduction resulting from the activity of two pacemakers (which gives rise to a doubling in beat frequency). Two arrows meeting indicate that the waves of excitation resulting from the activity of two pacemakers cancelled each other.

A. Trace 1: Note the periodic doubling in beat frequency. This results from the activity of an ectopic center (action potentials indicated by dots) alternating with the primary pacemaker. The diagrams show directions of conduction during the doubling of beat frequency (left diagram) and when only one center was active. Trace 2: The heart arm has been split in two. Note that this split heart has the same beat frequency as either center in trace 1. Trace 3: Note that the pattern of the doubling in beat frequency was the same as in trace 1. Traces 4 & 5: The myocardium as shown in trace 3 was split. Note that the frequencies of these segments were about the same as that in trace 2 and the frequencies of each center in traces 1 & 3, indicating that all segments of the ring of pacemaker cells had about the same pacemaker capabilities (amplitude was altered by a change in suction).

B. Note the periodic, gradual change in signal amplitude in trace 1. A change in signal amplitude resulted from a change in the direction of conduction and the activity of two pacemaker centers. The waves of contraction opposed each other as shown in the diagrams to the left of trace 1. The heart was split in two as shown in the diagrams to the left of traces 2 & 3 and the spread of excitation was unidirectional in each segment.

water was placed over the heart piece, which was pressed until flattened. Sea water (0° C.) was perfused onto one end of the slide and removed at the opposite end by aspiration. With this method of perfusion, pieces of heart tissue continued to contract for several hours. Visual observations of contractions were made with a phase contrast microscope.

RESULTS AND DISCUSSION

A. *Patterns of pacemaker activity*

With the exception of conditions to be discussed below, only the end regions of the heart showed pacemaker properties, *i.e.*, contractions always started at one or the other end, or at both ends of the heart; they did not originate anywhere else (see Kriebel, 1968a, for details).

Collapsed hearts *in situ* or in isolation often showed a periodic doubling in beat frequency in one end of the heart. By splitting the heart wall longitudinally, it was possible to separate strips that had the slower rhythm observed in the periods between the times when "doubling" occurred (Figs. 1A and 2A).

In some hearts, generally collapsed, a second active pacemaker could be excited by locally heating a pacemaker region; *i.e.*, in addition to the general increase in frequency, the warming often resulted in a sudden doubling in the number of beats per unit time (see Kriebel, 1968a, for details). This can be readily explained as being due to an ectopic center whose beats occurred between those of the usually active pacemaker.

A second pattern of beating was detected when recording with four electrodes placed around the ostium of an intact but isolated heart, but in contrast to the above pattern of doubling, beat frequency remained the same. In these cases the signal sequence changed, indicating that the direction of conduction had changed and that two pacemakers were active, one on each side of the raphe. The change in the direction of conduction usually changed the signal amplitude (Fig. 1B). When these preparations were split longitudinally, conduction occurred only in one direction and the signal amplitudes remained constant. The alternating dominance between two pacemaker centers in one end of the heart can be compared to pacemaker competition of both ends in the intact heart (Kriebel, 1968a). These results demonstrate that the raphe functionally isolates the pacemaker regions just as it isolates the cells of the general myocardium (Kriebel, 1967a, 1967b).

Arrhythmia could be abolished in isolated collapsed hearts by filling them through a cannula and in hearts *in situ* by mechanically stimulating the animal to contract, which raised the blood pressure.

By cutting or crushing parts of the cardiovascular junction containing the pacemaker, it was ascertained that a very small portion, not more than 8% of the circumference of the ostium (6 mm.), was necessary to maintain regular pacemaker activity. During recording from pacemaker regions of collapsed hearts it was often observed that two, sometimes three, sizes of action potentials appeared, each with its own regular frequency. It was particularly revealing that one or more action potentials of a certain series would drop out (Fig. 3C) as if an ectopic center had either failed to reach threshold for active responses or that conduction to the region of recording had failed (conduction is decremental and low at the ends of

the heart, which would decrease the safety margin for conduction; Kriebel, 1967a).

Sometimes the electrical records from isolated hearts showed even more complex rhythms. Careful study revealed that the complexities could be resolved into superimposed series of activities of two pacemaker centers, each showing periods of acceleration and deceleration (Figs. 2B and 3A).

B. *Properties of the middle pacemaker (the C center of v. Skramlik, 1938)*

Already mentioned previously, under normal circumstances, only the end pacemakers are active. However, when animals with exposed hearts were mechanically stimulated to contract, the blood pressure increased to such an extent that no blood was expelled and the end pacemakers either became irregular or stopped (Kriebel,

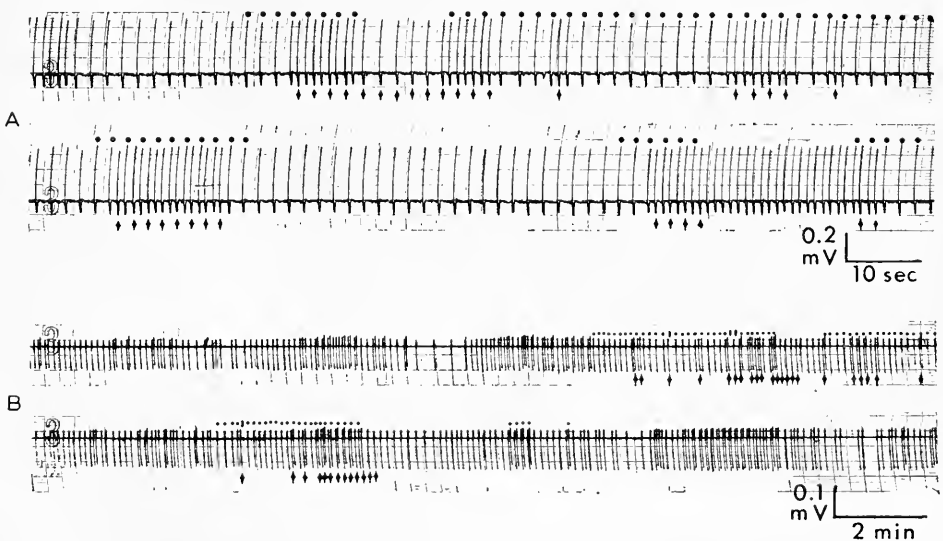


FIGURE 2. Action potentials from *Ciona* pacemaker regions. A. Traces 1 and 2 are continuous. There is little or no slowing in frequency but the frequency either doubled or decreased by half in a rhythmical pattern. Half of the action potentials recorded during the periods of doubling are in phase with those occurring during the preceding or subsequent period of low frequency beating. In trace 1, the dots indicate the action potentials that are in phase, during the doubling in beat frequency, with those preceding doubling. The diamonds indicate the phase of the remaining action potentials. Note that after the period of doubling, the action potentials were out of phase with those preceding the doubling. This activity can be explained by postulating two centers with slightly different frequencies so that a shift in phase would periodically permit both of them to drive the heart. In trace 2, one center was active all the time (dots) and the second center injected extra systoles (some action potentials are indicated by diamonds); 20° C. An opened heart arm. B. Traces 1 and 2 are continuous. In sections of this record, some action potentials in phase are indicated with diamonds or with dots and it is readily seen that the greatest frequencies result from two centers. However, the pacemakers also appear to miss beats as shown in the middle region of the first trace (cf. Fig. 3B). During the periods of low beat frequencies, a pacemaker accelerated and decelerated. This is shown just after the diamond symbols in the second trace (cf. Fig. 3A). During the periods of doubling, one center occasionally missed beats, presumably because the myocardium was refractory resulting from the prior activity of the dominant pacemaker (these misses are indicated by bars in sequence with the dots); 20° C. Heart arm containing some blood.

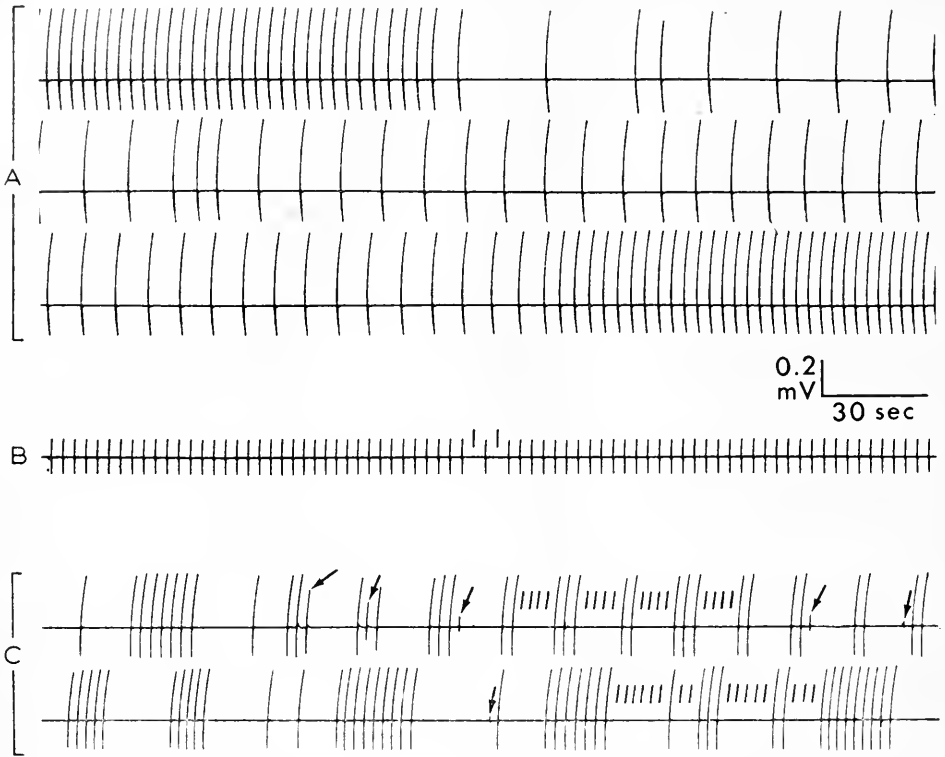


FIGURE 3. Action potentials from isolated pacemaker regions. A. These 3 records are continuous from a half-heart partially filled with blood (electrode in middle of half-heart). In the first trace, the pacemaker decreased in frequency before it stopped. In the second trace and the first half of the third trace the pacemaker gradually accelerated in a manner similar to that *in situ*. However, the beat frequency almost doubled in the middle of the third trace without a gradual increase. This probably resulted from the activity of a second pacemaker center. B. This record is from a collapsed heart arm. In the middle of the record note that two beats were missed (indicated by bars). Note that the action potentials occurring after the misses are in phase with those occurring before the misses. C. Continuous record of action potentials from a collapsed heart arm (electrode near pacemaker region). The intervals between the action potentials of the greatest amplitude are almost constant regardless of the duration of activity. The arrows point to contractions originating from ectopic centers. The bars indicate the missed beats of the pacemaker.

1968a). A few minutes after the end pacemakers had stopped, the C center became active and waves of contraction passed from the middle bend toward both ends of the heart. The frequency was always lower than that of the end pacemakers (in all four species studied).

Localized heating applied to an active C center was not as effective in increasing beat frequency as when applied to an end pacemaker. The Q_{10} of the C center in *Ciona* was found to be about 1.4 and that of the end pacemaker about 2.3 (5–25° C. range; Kriebel, 1968a). It was possible to reverse the direction of contraction by raising the temperature of the end of the heart with the inactive pacemaker or by

lowering the temperature of the end with the dominant pacemaker (Kriebel, 1968a). But when both end pacemakers were cooled to about 3° C. (so that their beat frequency dropped to only a few beats/minute) and the temperature of the middle region containing the C center was raised to about 20° C. the C center did not begin activity, although the end-to-end wave of contraction accelerated as it passed along the middle segment of the heart. Therefore a change in tension in the heart wall appears to be the only stimulus in the intact heart which will start the C center. Yet when the middle regions were isolated, the C center began to beat after a few minutes. This suggests that the change in a C center from a dormant to an active pacemaker requires that it not be driven for several minutes. The change from a dormant to an active state in the end pacemakers, after a reversal pause, requires only a few seconds (1–3 beat intervals).

The activity of the C center has been observed by many authors (Bancroft and Esterly, 1903; Hecht, 1918; Benazzi, 1935; Bacq, 1935; Sugi, *et al.*, 1965; Krijgsman, 1956).

C. *Pacemaker properties of myocardial cells*

There are many reports which demonstrate that for many species of tunicates small rings of heart tissue pulsate (see Krijgsman, 1956). However, in rings of heart it has not been determined whether the myocardial cells or the undifferentiated cells are the pacemakers. To answer this question, small pieces of tissue devoid of the raphe and the cells of the undifferentiated line were examined with a phase contrast microscope. They were observed to contract rhythmically, indicating that the cells of the general myocardium have pacemaker properties. Little or no activity was observed initially after isolating a small piece of tissue but after a few minutes 20–50% of the cells contracted at frequencies ranging from a few beats/min. to 12 beats/min. (intact *Ciona* hearts beat at about 20 beats/min.). Adjacent cells could beat independently. A few minutes later, the cells in small areas were contracting together until all the cells of the piece of myocardium contracted synchronously. The transition from contractions of small localized areas to synchronized beating of the entire piece of tissue was usually so rapid that the recruitment of additional fibers was seldom observed.

As in cultured chick heart cells (Smith and Berndt, 1964), there was no morphological difference between beating cells and quiescent cells. Synchronous beating of small groups of cells sometimes persisted for over an hour (also observed by Smith and Berndt, 1964). Occasionally single cells pulsed independently while all neighboring cells contracted together; however, after a period of time the aberrant cells usually followed their neighbors. Independent cell activity can be expected since the safety margin for conduction is very low and the excitability of the tissue is low (Kriebel, 1967a). For example, only local contractions were produced by stimulation with suction electrodes with tip openings less than 20 μ in diameter (Kriebel, 1967a). This means that several cells must be depolarizing simultaneously in order to generate a propagated wave of excitation. In the intact heart, dormant (or latent) pacemaker cells were being driven by the dominant pacemakers. This is substantiated by the fact that no intracellular pacemaker potentials were recorded from the cells of the general myocardium in the intact

heart (Kriebel, 1967a, 1967b). The transition from dormant to active states in the cells of the general myocardium requires more time than is required by the end pacemaker (after a pause during reversal) or by the C center. Since one cell cannot drive the heart, the appearance of an ectopic center depends on synchronized activity of several cells, each of which is contributing to the pacemaker current. Assuming that the pacemaker cells at the ends of the heart are functionally similar to the cells of the general myocardium, ectopic centers in the general myocardium are probably similar in size (number of cells) to those at the ends of the heart. Thus, the question arises: what factors integrate the pacemaker cells so that only one center is active in the normally functioning heart? Spread of excitation is by local current flow (Kriebel, 1967a, 1967b, 1968b). At the ends of the heart, the safety margin for conduction is very low because conduction velocity is low and decremental in nature (Kriebel, 1967c). Consequently, any factor that lowers the excitability of the tissue will lower the safety margin for conduction and in turn permit ectopic centers to develop (*cf.* Hoffman, 1965, for vertebrate hearts). Since arrhythmia was more frequent in isolated hearts, it seems reasonable to conclude that the safety margin for conduction was lower in sea water than when the hearts were in blood. However, in collapsed hearts (both *in situ* and isolated), arrhythmia was usually abolished by increasing the blood pressure. This indicates that stretch may increase the safety margin for conduction in the pacemaker region. Either one pacemaker center could drive the other potential pacemaker cells in the end pacemaker region so that they become dormant pacemakers or all of the cells in the end pacemaker region show pacemaker potentials but are driven by one center.

This work was supported in part by a NONR 1497 (OO) contract to the Marine Biological Laboratory and grants NIH T 1 GM 1194 and NIH 571 MH 6418 11. It was part of my Ph.D. thesis, 1967, Department of Zoology, University of Washington, Seattle, Washington 98105. I wish to thank Dr. R. Josephson for supplying the facilities at the Marine Biological Laboratory and Dr. E. Florey and Mrs. Laura Kriebel for help in preparing the manuscript.

SUMMARY

1. The primary pacemakers in the tunicate heart are located near the cardiovascular junctions close to the raphe that connects the V-shaped heart to the pericardium. However, small isolated pieces of the ring of myocardial cells at the ends of the heart were found to have nearly equal pacemaker capabilities. Arrhythmia in one primary pacemaker region was found to result from activity of two or more centers.

2. Following isolation of a small piece of tissue in sea water, 20–50% of the cells were observed to pulsate and usually within a few minutes all cells contracted synchronously. Reversals in the direction of conduction in strips of myocardium without any primary pacemaker region were observed.

3. The C center located in the middle of the heart was found to be dormant during normal heart activity but in opened animals it was activated by increasing the blood pressure.

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CHOLINOCEPTIVE AND ADRENOCEPTIVE PROPERTIES OF THE TUNICATE HEART PACEMAKER

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Functionally, tunicate hearts resemble those of the higher chordates since conduction is from cell to cell by local current flow (Kriebel, 1967b, 1968c, 1968d) and the pacemakers are myogenic (Millar, 1953; Kriebel, 1968b). However, cardiovascular control in tunicates has not been extensively studied.

There are no continuously active nerves that maintain the beat frequency either above or below the intrinsic frequency of the pacemaker (something like vagal tone in the higher vertebrates) since ganglion extirpation does not alter the beat frequency (Day, 1921; Bacq, 1935). Stimulation of the ganglion is without effect on the heart rate (Schultze, 1901). However, it is not excluded that there are cardiovascular nerves that are active only during specific conditions.

If regulatory nerves are present, it seems likely that the transmitters would be the same as in the higher chordate. Although acetylcholine and cholinesterase are present in tunicates (Florey, 1963). Ach at low concentrations has no effect on the beat frequency of intact isolated hearts (see Krijgsman, 1956, for a review; Krijgsman and Krijgsman, 1959). Adrenaline has been reported to accelerate the heart beat, but high concentrations were required (10^{-3} g./ml.; Scudder *et al.*, 1963).

Since there are no transverse channels (or gaps) between cells, the myocardium is an effective barrier to substances that do not penetrate cell membranes (Kriebel, 1968d). The results presented here show that Ach and adrenaline do not cross the heart wall and that receptive sites are present only on the lumen surface.

METHODS

Hearts of large adult *Ciona intestinalis* (from California) were exposed by cutting through the test and body wall into the coelomic cavity (see Kriebel, 1968a). Hearts ranged from 24 to 45 mm. in length. In some preparations the blood vessels were ligated before cutting in order to free the heart, whereas in others, the vessels were simply severed. In all preparations the pericardium was opened. Many isolated hearts were ligated in the middle region so that the activity of one primary pacemaker could be studied without the influence of the opposite pacemaker. Hearts were opened so that the lumen surface was exposed to the bath either by cutting along the raphe, or, during electrical recording, by pressing a razor blade through the myocardium. The latter method of opening hearts did not disturb the tissue that was sucked into suction electrodes (Kriebel, 1968a).

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Hearts were carefully secured with small hooks to the bottom of a 5-ml. bath (10° C. or 21° C.) to insure that constant dimensions in the heart wall were maintained during addition of the drug solutions and the washing procedure. Slight alterations in tension changed the amount of tissue sucked into the electrode openings and thereby altered the signal amplitude as well as the excitability of the pacemakers (Kriebel, 1968a). Consequently, control experiments, which consisted of adding, mixing and removing sea water, were performed before and after the drug additions to validate the drug actions. Hearts with pacemakers that initially or later became sensitive to alterations in the bath volume were not used.

The effect of drugs on the beat frequency was tested on hearts, first intact, then opened. A solution of the drug in sea water at a temperature equal to that of the bath was added with a pipette. The final concentrations were predetermined. Mixing was accomplished within 15 seconds with an eye dropper. To remove a drug solution at least 100 ml. of sea water of the desired temperature were perfused through the bath.

Acetylcholine chloride (Merck), L epinephrine bitartrate (Nutritional Biochemical Corporation), d-tubocurarine chloride pentahydrate (Burroughs Wellcome & Co.) and atropine sulphate (The Norwich Pharmacal Co.) were used.

RESULTS

Since the effects of drugs on the pacemaker frequencies of intact and opened hearts were to be compared, and since the amount of intracardiac blood affects the beat frequency (Kriebel, 1968a), intact hearts were studied in the collapsed condition (*i.e.*, they contained no blood). Thus, when the hearts were opened, the pacemakers were not subjected to a change in tension. However, arrhythmia frequently occurred in collapsed hearts (Kriebel, 1968a, 1968b). A common pattern of arrhythmia was a doubling in the beat frequency due to the activity of an ectopic center (as shown in Fig. 1A, trace 1).

Acetylcholine (10^{-8} to 10^{-4} g./ml.) and adrenaline (10^{-6} to 5×10^{-4} g./ml.) were without effect on the beat frequency of intact isolated *Ciona* hearts (and half-hearts; with and without the blood vessels tied off, full of blood or collapsed). However, when the hearts were opened, forming a flat sheet of tissue, Ach at 10^{-8} to 10^{-7} g./ml. or adrenaline at 10^{-5} g./ml. initially stopped the primary pacemaker (Figs. 1 and 2). The pacemakers did not again stop when the bath was stirred or when another threshold aliquot of drug was added. The concentration could be slowly raised to about 100 times threshold for Ach and about 10 times for adrenaline with no effect on the beat frequency of the primary pacemaker. Therefore, the response of the pacemakers to these agents exhibited tachyphylaxis, *i.e.*, they became desensitized. However, at initial concentrations greater than $100 \times$ threshold for Ach and $10 \times$ threshold for adrenaline the primary pacemakers were irregular and ectopic centers developed (Ach at 10^{-5} g./ml., Fig. 2; and adrenaline at 5×10^{-4} g./ml., Fig. 1).

Even when vessels were not ligated; intact isolated hearts were insensitive to the drugs, presumably because the blood vessels sealed when they were severed. When the vessels were opened and washed, intact hearts became sensitive to Ach and adrenaline. However, the concentrations required to arrest the hearts were

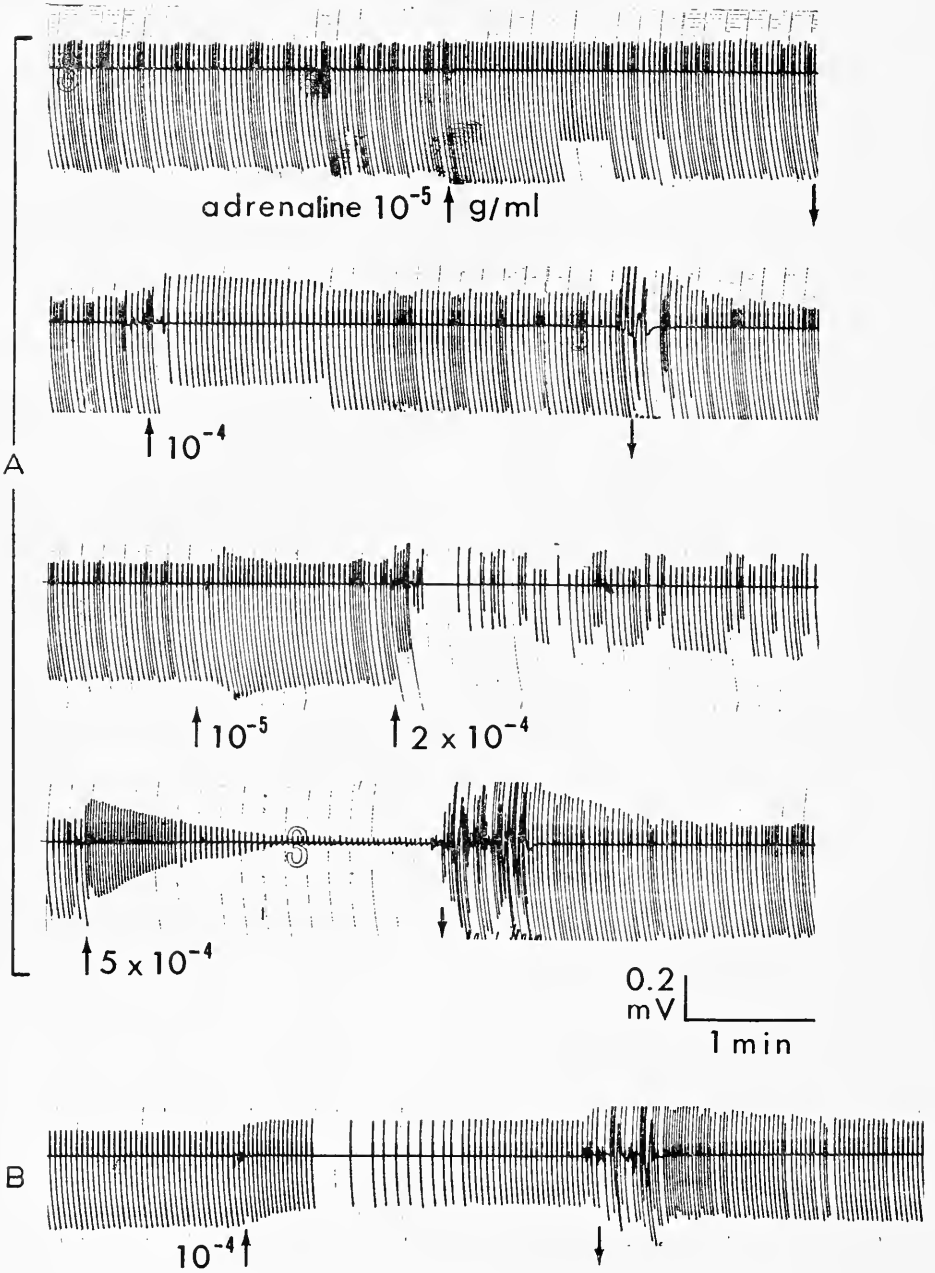


FIGURE 1.

higher than required after the hearts were opened. This indicates that diffusion through the collapsed ostia is slow and in addition it is likely that tachyphylaxis occurred.

Atropine antagonized the effect of Ach (Fig. 2). Threshold concentrations of d-tubocurarine (10^{-7} to 3×10^{-7} g./ml.) either slowed or stopped a pacemaker center. Complete stoppage of the heart occurred with slightly greater concentrations and primary pacemakers did not recover. The effect was additive to that of Ach.

Adrenaline sometimes stopped the heart for a few beats permitting an ectopic center to develop which alternated with the primary pacemaker center, doubling the beat frequency (Fig. 1A, trace 4; *cf.* Scudder *et al.*, 1963; Sugi and Matsunami, 1966). Doubled beat frequency also occurs frequently in deflated hearts and can be produced by locally heating the pacemaker region (Kriebel, 1968a, 1968b). If a primary pacemaker and an ectopic center were active in a heart arm, a threshold concentration of adrenaline (10^{-5} g./ml.) often stopped one center, resulting in halved beat frequency (Fig. 1A, trace 1).

High concentrations of adrenaline (5×10^{-4} g./ml.) decreased conduction velocity (Kriebel, 1967a). This can explain the decrease in signal amplitude (also found by Scudder *et al.*, 1963). After addition of a high concentration of adrenaline, conduction velocity continued to decrease for 1 to 3 minutes. During this time, amplitude of action potentials also gradually decreased (Fig. 1A, trace 4). Although beat frequency decreased under the influence of adrenaline, groups of beats at a relatively high frequency occurred. By the shape of the recorded wave form they could be attributed to one pacemaker and it appears that beats were missed between the groups (as seen in Fig. 1A, trace 3). Because conduction velocity is very low and because it is decremental at the ends of the heart (Kriebel, 1967a) a further decrease in the conduction velocity due to the action of adrenaline would decrease the safety margin for conduction. A reduced safety margin could give rise to missed beats and permit the development of ectopic centers.

FIGURE 1. Effect of adrenaline on the beat frequency of opened *Ciona* hearts. The electrograms were recorded with suction electrodes placed on the primary pacemaker region. The downward arrows indicate washing; 20° C. A. Trace 1: The first half of this record shows a rhythmical doubling in beat frequency attributed to the activity of an ectopic center which alternated with the activity of the primary pacemaker. After the application of adrenaline note that the doubling in beat frequency was stopped for over 1 minute. Trace 2: After the application of adrenaline at a concentration of 10^{-4} g./ml. the heart stopped, the doubling in frequency was abolished and for about 2 minutes the heart contracted at a lower frequency. However, it was visually determined that the direction of contraction had reversed. That is, an ectopic center at the cut end of the heart arm now drove the heart. The activity of the ectopic center is indicated by the change in the signal amplitude. Trace 3: With adrenaline 10^{-5} g./ml. the doubling in beat frequency was stopped as in trace 1. With adrenaline 2×10^{-4} g./ml. the heart did not recover. Signals of different amplitude indicate a change in the direction of conduction and thus the activity of different centers. Note that the signals of the same amplitude sometimes occurred in groups (see text). Trace 4: Continuous with trace 3. Adrenaline at 5×10^{-4} g./ml. doubled the beat frequency for about 40 seconds. Then the beat frequency suddenly dropped to about half, at which time it was visually observed that the wave of contraction started at the cut end of the half-heart. B. This electrogram shows the typical effect of adrenaline on an active pacemaker. The slight increase in frequency just after drug application was probably due to an increase in the bath temperature. Mixing was 30 sec. after arrow.

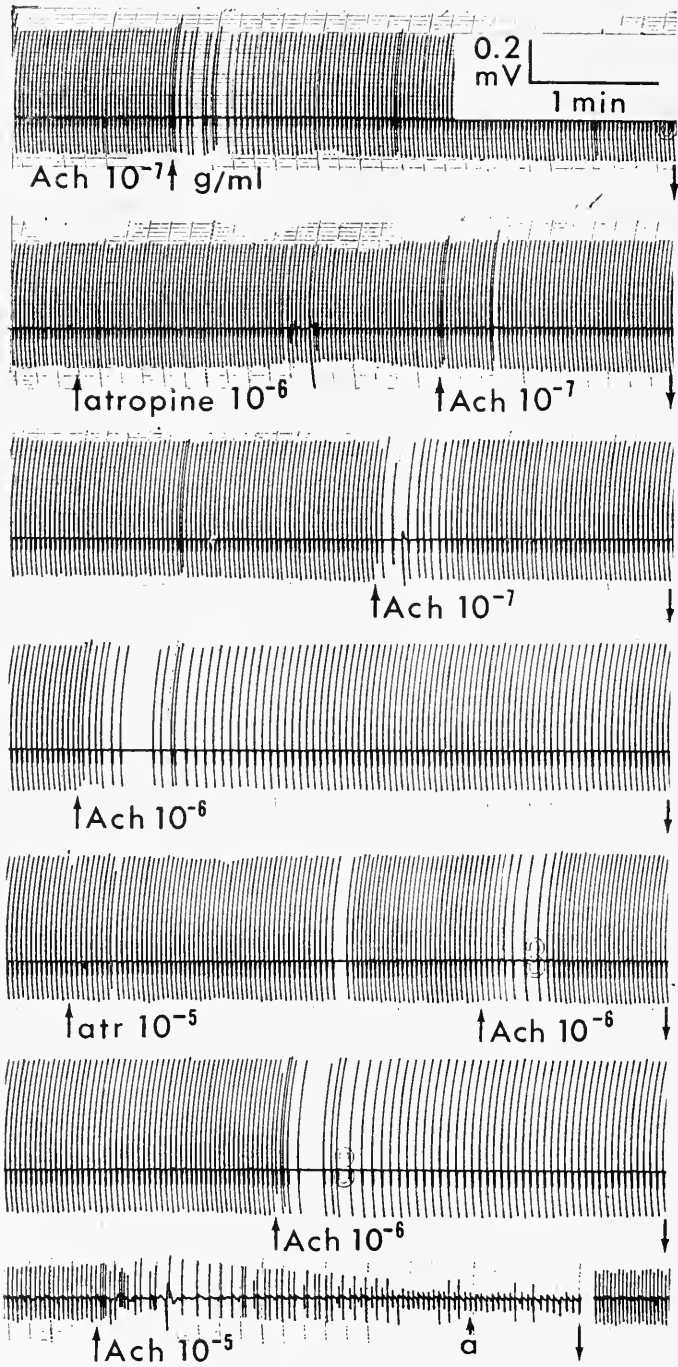


FIGURE 2.

DISCUSSION

Tight junctions join all cells together at their apical borders and exclude extracellular space so that there are no transepithelial diffusion channels between cells (Kriebel, 1968d). The ineffectiveness of drugs on intact hearts shows that the heart tube is permselective and that cholinceptive and adrenoceptive sites are on the lumen surface.

The failures of other investigators to observe the effects of low concentrations of Ach and adrenaline are readily explained by the insensitivity of the unopened heart, the sealing of the cut ends of the heart and by the presence of tachyphylaxis (*cf.* Bacq, 1934; Krijgsman and Krijgsman, 1959; Scudder *et al.*, 1963; Sugi and Matsunami, 1966).

The absence of an accelerating effect of adrenaline, as found in other chordate hearts, may be an indication of the absence of cardioacceleratory nerves.

On the other hand, the typical cholinceptive properties of the myocardium (inhibition caused by Ach and Ach-block by atropine) favors the interpretation that the tunicate heart may receive cardioinhibitory innervation (Florey, 1951) and this innervation is cholinergic. In this connection it is interesting that Bone and Whitear (1958) could trace nerves as far as the pericardium.

This work was supported by grants 5 T O 1 G M 01194, 2 R 0 1 N B 01451 and NIH 5T1 MH 6418 11. It was used as part of my Ph.D. thesis, 1967, Department of Zoology, University of Washington, Seattle, Washington 98105. I wish to thank Dr. E. Florey and Mrs. Laura Kriebel for help in preparing the manuscript.

SUMMARY

1. Many previous investigators have reported that acetylcholine has little or no effect on the beat frequency of intact but isolated hearts. However, it was found that when hearts were split open, Ach at low concentrations (10^{-8} g./ml.) stopped the heart beat for up to a minute. Atropine blocked Ach.
2. D-tubocurarine at a concentration of 3×10^{-7} g./ml. stopped the beat of opened hearts. Its effect was additive to that of Ach.
3. Adrenaline (10^{-5} g./ml.) could stop pacemakers. However, a doubling in beat frequency was frequently observed with higher concentrations. Doubling in beat frequency resulted from an ectopic center which alternated with the primary center, thus driving the heart at twice its original frequency. In hearts which showed arrhythmia resulting from activity of more than one pacemaker center,

FIGURE 2. Effect of acetylcholine on the beat frequency of an opened *Ciona* heart. The electrograms were recorded with a suction electrode placed on the myocardium near the end of the heart (*i.e.*, on the primary pacemaker region). The downward arrows indicate washing. Temperature 20° C. Traces 1-3 show that atropine blocked a threshold concentration of Ach. Traces 4-6 show that atropine partially blocked the initial stoppage caused by a high concentration of Ach (10^{-6} g./ml.) and greatly diminished the time required to regain the original beat frequency. Trace 7 is from the same heart but the electrode position had been slightly changed altering the signal amplitude. Note the different amplitude after application of Ach. It was visually observed that the smallest signals (a) corresponded to contractions starting at the cut end of the half-heart.

adrenaline at a low concentration (10^{-5} g./ml.) usually stopped one or the other of the centers, decreasing the beat frequency by half.

4. Hearts recovered in low concentrations of either Ach or adrenaline. However, at higher concentrations (Ach 10^{-6} , adrenaline 5×10^{-4} g./ml.) arrhythmia was induced and hearts did not recover. The occurrence of arrhythmia is discussed with respect to the decremental nature of conduction in the ends of the heart and to the observation that all cells near the ostia are latent or dormant pacemakers.

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INTER-PHYLOGENETIC SPECIFICITY IN THE BONDING OF AMINO ACIDS TO tRNA¹

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The first step in protein synthesis is generally considered to be the reaction of a free amino acid (AA) with adenosine triphosphate (ATP) and an appropriate "activating" enzyme to form an "activated" amino acid (probably 5'-aminoacyl adenylate: AA ~ AMP) and pyrophosphate (Hoagland, Zamecnik and Stephenson, 1957).

This activated amino acid reacts with the 2' or 3' OH of a low molecular weight ribonucleic acid (tRNA) to release AMP and produce an aminoacyl tRNA which subsequently transports the amino acid into the appropriate part of the protein being synthesized. Evidence for this unique RNA was adduced more or less simultaneously and independently by Holley (1957) and by Hoagland *et al.* (1957), as well as indirectly by Hultin (1956) and Hultin and Beskow (1956). The structures of several tRNA's have been determined and it seems quite likely that more than one tRNA accepts a particular amino acid, although there is only limited evidence that more than one aminoacyl tRNA ligase exists for each amino acid (Yu, 1966; Barnett, Epler and Brown, 1967).

It is presumed that some sort of base pairing of the aminoacylated tRNA with genetic nucleic acid (mRNA, virus RNA or even DNA) must account for the extraordinary precision with which amino acids are assembled into proteins (Loftfield, 1963) or with which phenotypes reflect different genotypes. There is some evidence that the genetic code is universal and corresponding evidence that the insertion of an amino acid into a particular locus depends only on the tRNA to which it is attached. Berg and Ofengand (1958) showed that the activating enzymes also catalyzed the transfer of amino acid to tRNA. Hecht, Stephenson and Zamecnik (1959) showed that most, if not all, tRNA's terminated in an adenylic-cytidylic-cytidylic trinucleotide and several recent studies on structure show that the secondary structures may be similar for all tRNA's. It was the purpose of this work to establish whether activating enzymes from widely different sources could aminoacylate tRNA's from other phyla. Although the results are not gratifyingly unequivocal, we find that interaction between heterologous pairs exists where others have found it absent. In some cases, the interaction of heterologous pairs of tRNA and enzyme, as measured by our methods, is better than between homologous pairs. Our techniques show that the aminoacyl tRNA is chemically active however obtained, and that there is no chemical evidence of a multiplicity of aminoacyl tRNA's.

¹ Some of this material has been presented at the Tuesday Evening Seminar Series at the Marine Biological Laboratory (Loftfield and Eigner, 1967a). This work has been supported by U.S.P.H.S. Grant CA 08000.

MATERIALS AND METHODS

(A) L-1-[^{14}C]-valine and L-1-[^{14}C]-leucine (20 c.p.m. per μmole) were prepared by the Bucherer hydantoin synthesis (Loftfield and Eigner, 1966). TRIS (tris (hydroxymethyl)-aminomethane), ATP (adenosine triphosphate), and other reagents were purchased from commercial sources.

(B) Preparation of enzymes—*E. coli*. Ten g. of *E. coli* strain B cells were suspended in 50 ml. of .02 *M* TRIS buffer pH 7.5 containing 600 mg. of reduced glutathione per liter. The cells were ruptured by high pressure passage through the orifice of a Ribi cell fractionator. The suspension was then centrifuged two hours at 27,000 *g*. The supernatant fluid possessed a high concentration of both valine and leucine tRNA ligases and was generally used as such. Occasionally, the enzymes were further purified by column chromatography on DEAE cellulose (Berg, Bergmann, Ofengand and Dieckmann, 1961).

Toadfish. One gram of toadfish liver was homogenized in a Potter-Elvehjem homogenizer with 1 ml. of Medium A (0.35 *M* sucrose, 35 mM KHCO_3 , 4 mM MgCl_2 and 25 mM KCl). The suspension was centrifuged at 24,000 *g* for 30 minutes. The supernatant phase was removed and the sediment extracted with a second 1-ml. portion of Medium A. All of the valine and leucine activating enzymes appeared in the supernatant phases which were used as such.

Starfish. The gonads, male or female, appeared to be the best source of amino acid activating enzymes. Ripe gonads were minced and stirred gently one-half hour in fresh sea water to release eggs or sperm. The water with eggs or sperm was decanted and the suspension of gonad tissue was homogenized in a Potter-Elvehjem apparatus with an equal volume of Medium A containing 600 mg./l. of reduced glutathione. The suspension was centrifuged 10 minutes at 20,000 *g* and the supernatant phase was used as the enzyme source.

Yeast. Fresh bakers yeast was cultured and harvested in log phase by Dr. J. Scaletti of The University of New Mexico. Fifty ml. of packed cells were suspended in 100 ml. of 0.02 *M* pH 7 phosphate buffer containing 600 mg./l. of reduced glutathione and forced through a Ribi press at 35,000 lbs./in.² The suspension was centrifuged 20 minutes at 20,000 *g* and $(\text{NH}_4)_2\text{SO}_4$ was added to 80% saturation. After 20 minutes at 4° C., the precipitated protein was collected by centrifugation at 20,000 *g* for 20 minutes. The precipitate was dissolved in 5 ml. of the above buffer, 1 g. of Bentonite was added and the slurry was allowed to stand at room temperature 30 minutes. The Bentonite was removed by centrifugation at 20,000 *g* and the enzyme solution was dialyzed against the phosphate glutathione buffer for six hours at 4°. The RNA contamination could be reduced by treating each ml. of the enzyme solution with 10 μmoles of MgCl_2 , 600 μg . of glutathione and 10 mg. of streptomycin for 30 minutes at 0°. The precipitate was removed by centrifugation and the solution again dialyzed against the phosphate glutathione buffer.

Each enzyme preparation was routinely examined for activity using the [^{14}C]-amino acid hydroxamate technique (Loftfield and Eigner, 1963).

(C) tRNA's.—*E. coli* tRNA and yeast tRNA were gifts from Schwartz Bio-research.

Starfish.—Eighty ml. of ripe starfish eggs were suspended in 80 ml. of 0.001 *M* pH 7.5 TRIS buffer together with 16 ml. of 0.1 *M* MgCl_2 . The suspension was

homogenized at 0° for 5 minutes in a Waring Blendor. Undissolved material was removed by centrifugation for 10 minutes at 16,000 *g*. The supernatant phase was gently mixed for one hour at room temperature with an equal volume of 90% phenol. Centrifugation separated the mixture into two phases. To the upper phase were added 0.1 volume of 20% potassium acetate and 2.5 volumes of ethanol. After 16 hours at -10°, the precipitate was collected by centrifugation, washed once with 67% ethanol and drained. The precipitate was then suspended in 1 *M* NaCl and centrifuged at 15,000 *g* for 30 minutes to sediment most of the ribosomal RNA. The supernatant was dialyzed against water, treated with 0.5 *M* TRIS at pH 8.8 for 45 minutes to hydrolyze aminoacyl tRNA, neutralized with acetic acid and dialyzed again. Although the solution showed a spectrum typical of RNA, it was very viscous, suggesting the presence of a highly polymerized contaminant. Therefore the solution was stirred two minutes at 20° with one volume of 2.5 *M*, pH 7.5 potassium phosphate buffer and one volume of methyl cellosolve. The mixture was separated into two phases by centrifugation at 15,000 *g* for 10 minutes at 4°. tRNA was precipitated from upper phase by adding potassium acetate and alcohol as above.

Toadfish.—Forty grams of toadfish liver were homogenized and worked up in much the same way as the starfish eggs. Although the product had a slight gray-brown color, it had an ultraviolet spectrum that was typical of tRNA. Like each of the other tRNA's prepared, the ratio of O.D.₂₆₀/O.D.₂₈₀ was approximately 2.0.

Using homologous enzymes, the maximum aminoacylation of each tRNA was: in μ moles per mg.: *E. coli*, leucine 1.0, valine 1.2; yeast, leucine 0.37, valine 0.35; starfish, leucine 0.45, valine 0.45; toadfish, leucine 0.59, valine 0.36.

(D) The aminoacylation of tRNA was carried out as previously described (Loftfield and Eigner, 1967a) and as summarized in the legend of Figure 1.

RESULTS

There has been an abundance of work on the cross-reaction of activating enzymes and tRNA's of different species. For the most part only bacterial, yeast and vertebrate sources have been used and examination has been confined to a few amino acids. In general, there seems to have been some cross-reaction when yeast or vertebrate sources of enzyme and RNA were used but none or little when bacteria provided one of the ingredients. Benzer and Weisblum (1961) showed that the extent of cross-reaction varied with the amino acid used. We (Loftfield and Eigner, 1963) demonstrated that if sufficient care is taken, pairs reported to have no cross-reaction can be shown to react, though the rates may be less than 1% of the homologous pairs.

Stulberg and Novelli (1962; p. 422) correctly point out some of the errors in making hasty statements that interspecies interaction does not obtain. Frequently so-called "tRNA" has not been shown to accept amino acids from the homologous enzyme. Thus many of our preparations of starfish "tRNA" had the physical and chemical properties of tRNA without being able to bind covalently significant amounts of several amino acids even using starfish enzyme. Other workers have occasionally reported no effort to determine whether the aminoacylated tRNA was biochemically or chemically active.

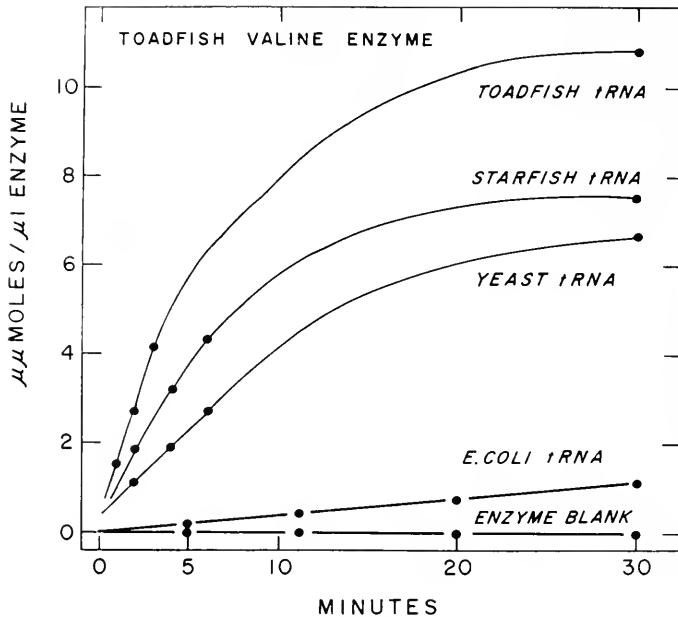


FIGURE 1. A typical experiment that shows the rate of valyl tRNA formation by the toadfish valine activating enzyme with four different species of tRNA. The reactions were carried out at 25° with 10 mM ATP, 0.1 M TRIS (pH 7.5), 30 μ M [14 C]-valine, 0.5 mg. tRNA per cc., and 6 μ l. enzyme solution per tube (final vol. per tube 225 μ l.; four aliquots of 50 μ l. each were taken at times indicated). The enzyme blank contains everything except added tRNA. The aliquots were worked up as previously described (Loftfield and Eigner, 1967a). The 30-minute observations correspond to the incorporation of 0.35 μ mole of valine to 1 mg. of yeast tRNA and to 0.45 μ mole of valine into 1 mg. of starfish tRNA, exactly the same saturating values as found with homologous enzyme. On the other hand, the *E. coli* tRNA is only about one-tenth saturated.

Hence the current work has been undertaken with considerable care to establish that aminoacylation of tRNA was actually taking place. Every figure noted in Table I represents a rate of reaction drawn from at least two different experiments in which the extent of reaction had been determined at no fewer than three time intervals. Figure 1 illustrates the kind of data used and the difficulties involved in comparing rates. For instance, many of our enzyme preparations contained an RNase which slowly destroyed the substrate RNA; this was clearly a greater problem if the reaction was slow or if larger amounts of enzyme solution were being used. Thus in some heterologous reactions similar to Figure 1, the tRNA was never saturated with amino acid. However, the addition of ATP and enzyme homologous with the tRNA at 30 minutes was ineffective. The receptor competence of the tRNA had been destroyed during the incubation. This was especially evident when enzyme and tRNA were pre-incubated without the amino acid. Depending on the species and preparation, the competence of the tRNA to accept amino acid steadily deteriorated. No two preparations of tRNA accepted, even from homologous sources, the same amount of leucine or valine and we (Loftfield

and Eigner, 1963, 1965) have previously shown that either aminoacylated tRNA or damaged (oxidized) tRNA inhibits the reaction. Even products of tRNA digestion have been shown to be inhibitory (Hayashi and Miura, 1966; Letendre, Michelson and Grunberg-Manago, 1966). We have presented in Table I relative initial rates of reaction where destruction of substrate, product inhibition and enzyme inactivation are minimized.

With due regard for all these difficulties, Table I contains material of interest. In keeping with earlier observations, most *E. coli* enzymes of established activity interact poorly if at all with non-bacterial tRNA's of established competence while there is a fair to better than natural interspecies reaction when the enzymes and tRNA's are derived from yeast or the higher organisms. (At best, this is only a weak generalization; starfish valine enzyme reacts poorly if at all with *E. coli* tRNA while the leucine enzyme of starfish is as good as the *E. coli* enzyme.)

TABLE I

Relative rates of homologous and heterologous enzymic formation of aminoacyl tRNA's. The rates are expressed as per cent of the rate obtained for homologous tRNA and enzyme under standard conditions

Enzyme		tRNA			
		<i>E. coli</i>	Toadfish	Starfish	Yeast
<i>E. coli</i>	val	100	2	1.5	0.5
	leu	100	<0.1	<0.5	2
Toadfish	val	3	100	50	27
	leu	<0.1	100	47	3
Starfish	val	<1.	100	100	83
	leu	100	40	100	40
Yeast	val	40	200	140	100
	leu	45	300	65	100

Most striking is the observation that yeast enzymes specific for both valine and leucine are more active towards starfish and toadfish tRNA than they are towards yeast tRNA. Equally notably, yeast enzymes for both amino acids are quite active in transferring leucine or valine to *E. coli* tRNA while *E. coli* enzymes are almost inert towards yeast tRNA. Partly to the contrary, starfish enzyme is very active towards *E. coli* tRNA only with leucine while *E. coli* enzyme is poor in transferring either leucine or valine to starfish tRNA.

There is a concern whether some tRNA normally specific for one amino acid is being aminoacylated by another. Although there are several reports of multiple tRNA's specific for a single amino acid there are to date, only a few suggestions of a multiplicity of the aminoacyl tRNA ligases (Yu, 1966; Barnett *et al.*, 1967). Nonetheless we deliberately used crude enzyme preparations rather than purifying the enzyme so as to lose no active component. If our enzymes were transferring amino acids indiscriminately to a variety of the heterologous tRNA's, we might

expect that some tRNA's would accept more leucine or valine than they did from the homologous enzyme. In every case, we found that the heterologous aminoacylation yielded only the same extent of reaction as the homologous. (Note the legend in Figure 1.) No evidence was found to indicate that the heterologous reaction was less specific than the homologous reaction.

The most valid criterion of whether a heterologously aminoacylated tRNA was identical with the same tRNA aminoacylated with a homologous enzyme would be to compare the behavior of the two tRNA's in the synthesis of a specific protein. Unfortunately all cell-free syntheses of protein are suspect. The use of either natural messenger RNA (*i.e.*, to synthesize hemoglobin or β -galactosidase) or so-called synthetic mRNA's like poly-uridylic acid (to synthesize polyphenylalanine) leads to data in which high backgrounds, autolysis, RNase activity, salt

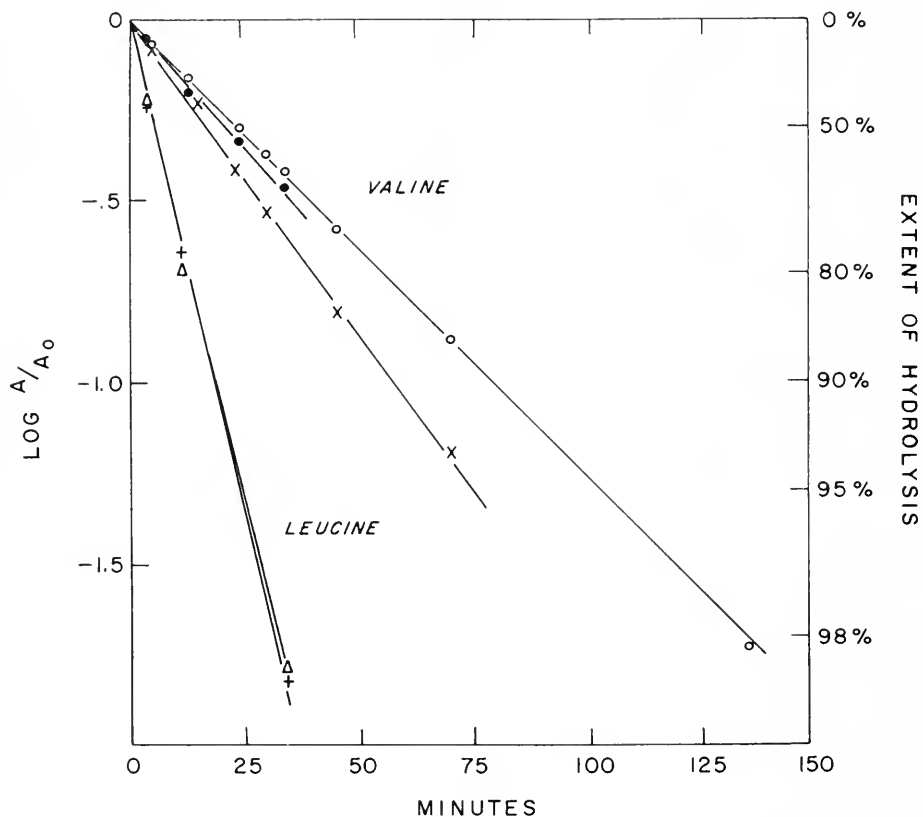


FIGURE 2. Rate of stripping of [14 C]-valyl and [14 C]-leucyl tRNA. The tRNA was labeled completely in each case at 25°, then made 0.16 *M* in Na_2CO_3 (pH 10.3) and brought to 35° to initiate the stripping reaction. Other conditions same as Figure 1, but larger amounts of enzyme were used. [14 C]-leucine was used with *E. coli* enzyme and *E. coli* tRNA (Δ), and toadfish enzyme and toadfish tRNA (+). [14 C]-Valine was reacted with *E. coli* enzyme and *E. coli* tRNA (\circ), toadfish enzyme and toadfish tRNA (\times), and toadfish enzyme and starfish tRNA (\bullet).

concentrations, etc., are prominent contributors to an ambiguous interpretation. Accordingly, we have used a much simpler criterion of the nature of the aminoacyl tRNA bond. Raacke (1958) pointed out that, although aminoacyl tRNA compounds are essentially simple esters, they are far more reactive than the aliphatic esters of amino acids. We chose to measure the rate of deaminoacylation of the aminoacyl tRNA's formed by either homologous or heterologous enzymic action. Figure 2 is illustrative of the kind of data obtained at pH 10.3 for a variety of these hydrolyses.

The first thing to be noted in Figure 2 is that the first order rate of hydrolysis is absolutely constant, *i.e.*, the semi-logarithmic plot shows no deviation from linearity. In every case, the hydrolysis was followed as far as the available [^{14}C]-aminoacyl tRNA permitted. Thus in the case of *E. coli* valyl tRNA the rate of hydrolysis remains constant until 97% complete hydrolysis. In no case is there evidence of a change in rate which certainly would have resulted from differences in chemical activities of the several leucyl tRNA's or valyl tRNA's. The graph shows clearly that the technique is sensitive to differences in rate of as little as 10%. Raacke showed that amino acids esterified to positions other than the unique 2' or 3' adenosyl terminal positions of tRNA would be a hundred times less reactive. Correspondingly, our data show that no more than 3% of the valine or leucine is bound to a relatively unreactive position such as a 5' hydroxyl or one of the amino or enol residues. Whether the tRNA has been aminoacylated by homologous or heterologous enzyme, the rates of hydrolysis are well within the range established for a particular amino acid. It is noteworthy that the leucyl derivatives are some four times more labile than the valyl derivatives. This agrees with the well recognized differences in rates of hydrolysis of leucyl and valyl peptides and is consistent with the steric inhibition generalizations of Newman (1950). In fact, the relative rates of hydrolysis of the leucyl esters are somewhat less than would have been anticipated from the application of Newman's "Rule of Six" to the straightforward attack of a hydroxyl ion or other nucleophile on the carbonyl bond.

Particularly to be noted are the clearly discernible differences in rates of hydrolysis of the several valyl tRNA's. Although these are much more stable than the leucyl tRNA's, the analysis is sensitive enough to detect differences which presumably result from differences in the structure of the tRNA.

The extraordinary activity of the yeast enzyme towards toadfish tRNA warranted a closer examination. It is quite conceivable that the high relative rates of aminoacylation are a consequence of our failure to saturate the homologous enzyme tRNA pairs. For instance if the K_m for binding toadfish tRNA to yeast enzyme is very low while the K_m for binding the yeast tRNA to yeast enzyme is high, we might expect that we should observe nearly maximal aminoacylation rates with toadfish tRNA while we were still observing only a fraction of the V_{\max} with yeast tRNA. Figure 3 shows that this interpretation is quite incorrect. An Eadie plot of rate data collected at various concentrations of tRNA shows that the K_m for yeast valine tRNA on yeast enzyme is $2.2 \times 10^{-7} M$ while the K_m for toadfish tRNA is $1.9 \times 10^{-6} M$. There is probably a ten-fold poorer binding of the toadfish tRNA which is more than compensated for by a six-fold increase in the reactivity of the toadfish tRNA as measured by V_{\max} . If the observations of

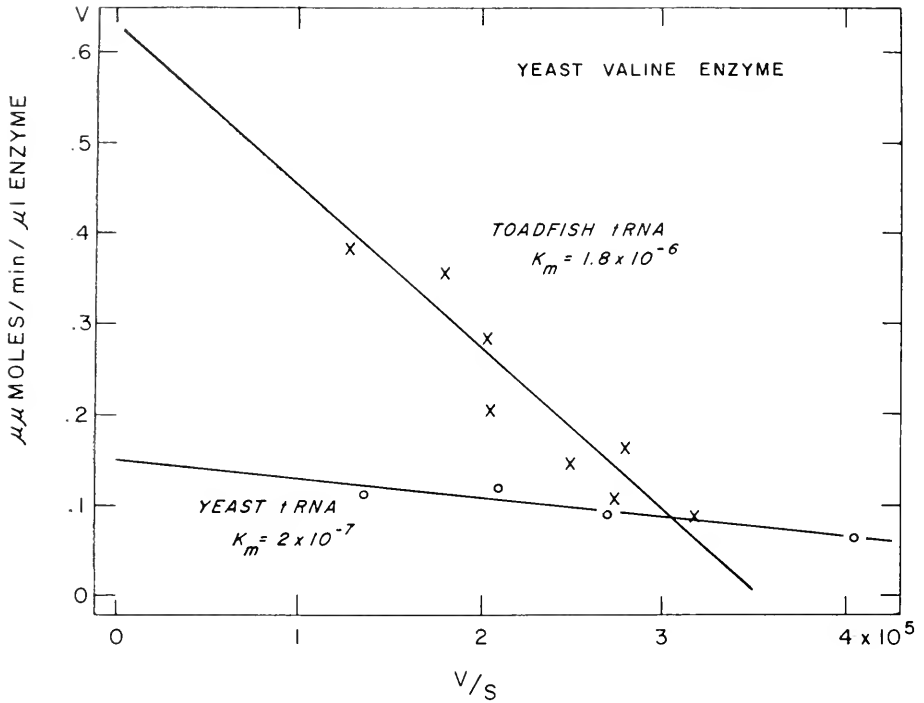


FIGURE 3. Eadie plot of formation of [14 C]-valyl tRNA by the yeast valine enzyme. The graph shows that yeast tRNA (\circ) has a K_m of 2×10^{-7} M and a V_{max} of $0.16 \mu\mu$ moles/min./ μ l. enzyme. Toadfish tRNA (\times) has a K_m of 1.8×10^{-6} but a V_{max} of $0.63 \mu\mu$ moles/min./ μ l., four times greater than the V_{max} of yeast tRNA with yeast enzyme. Conditions as in Figure 1, except yeast tRNA varied from 0.4 to 2.0 mg./cc. and toadfish tRNA varied from 0.46 to 4.9 mg./cc.

Table I had been made with tRNA concentrations of 5 mg. per ml., the toadfish tRNA would have appeared to be three or four times more active than yeast tRNA.

DISCUSSION

It has been noted by ourselves and others that there is inter-species interaction of tRNA and the corresponding ligase. It has previously been thought that most non-bacterial tRNA's would interact with other non-bacterial enzymes while bacterial tRNA would not react with non-bacterial enzymes. We (Loftfield and Eigner, 1963) have previously shown some cross reaction between *E. coli* tRNA or activating enzymes and the tRNA or enzymes derived from yeast and ascites tumors even when the rates of aminoacylation were extremely low. The present observations extend the generalization to enzymes and tRNA's from four widely differing sources. As noted previously, the failure to observe interaction as measured by an immeasurably low reaction rate does not mean that there is no interaction. Binding of the heterologous tRNA to enzyme may occur, but may be in

an unproductive mode or the rate of reaction may be so low as to escape observation in the presence of adventitious ribonucleases.

Table I shows that either valine or leucine reacts measurably for every pair of enzymes and tRNA's examined. There is no obvious pattern as to which amino acid is more likely to cross-react. For instance, *E. coli* enzyme puts more valine than leucine onto toadfish and starfish tRNA but more leucine onto yeast tRNA.

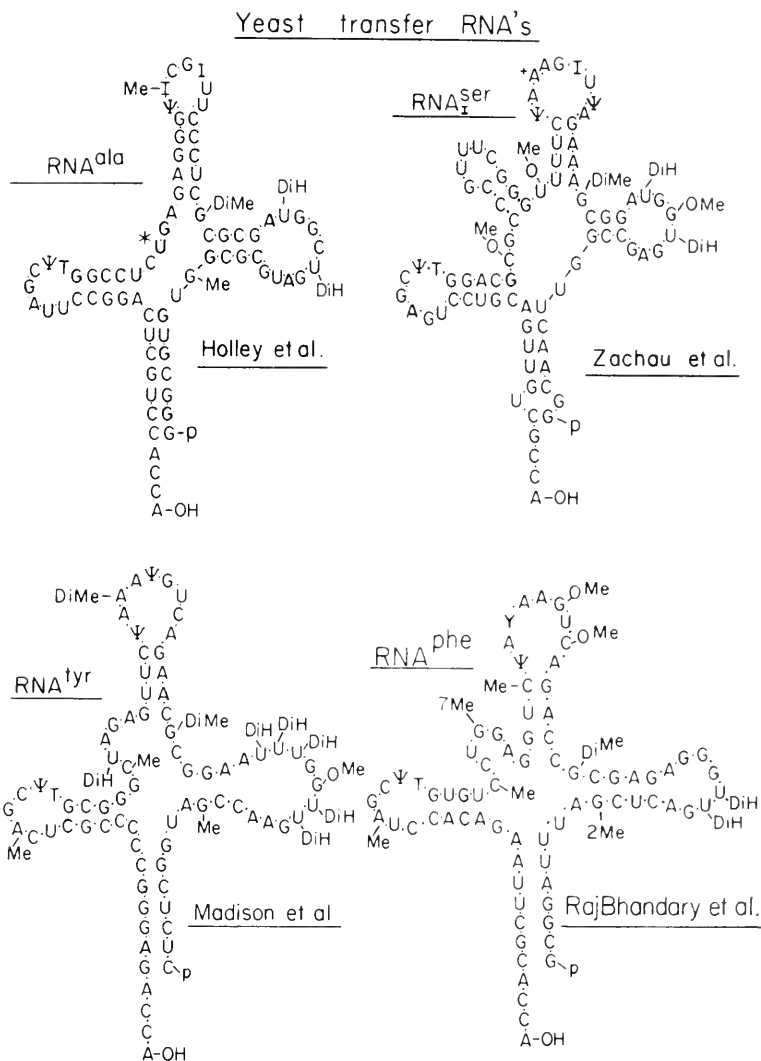


FIGURE 4. Possible conformations of four of the five tRNA's of known sequence. The sequences at the bottom (—OH end) and on the left loop and on the upper loop (which contains the presumptive anti-codon), are very unlike and may be responsible for the interphylogenetic amino acid specificity of the tRNA-ligase interaction.

Previously we have shown a similar irregularity in the actual K_m 's for the binding of yeast or *E. coli* tRNA's to valine activating enzymes. In the present study this observation too is repeated and extended. Toadfish valine tRNA is bound to yeast enzyme ten times more weakly than yeast tRNA but is apparently many times more reactive once on the enzyme.

Beyond this, we have made no observations that would suggest any heterogeneity in the formed aminoacyl tRNA. In no case did the heterologous reaction proceed further than the homologous and in every case the rate of hydrolysis of the product followed first order kinetics for a homogeneous material as far as the reaction could be followed.

All these observations are consistent with the view that all valine specific tRNA's share some features in common which are not present in tRNA's specific for other amino acids. It would be easy to imagine that this common feature is the anti-codon specific for valine. However, there is no evidence that this is the case. Each tRNA molecule possesses about 80 nucleotides so there is abundant possibility for the existence of many other highly specific combinations of nucleotides. It should be noted that five yeast tRNA's have had their structures completely elucidated (alanine, Holley *et al.*, 1965; serine I and II, Zachau *et al.*, 1966; tyrosine, Madison *et al.*, 1966; phenylalanine, RajBhandary *et al.*, 1966). In every case, the suggested anti-codon contains one of the so-called minor bases and is very remote from the adenosine residue which is to be aminoacylated. Moreover, among the four sequences shown in Figure 4, one can detect only a few patterns that are present in every case, *i.e.*, C-C-A at the receptor end and A-G-C- Ψ -T-G in the left hand loop. In all other areas, there are a sufficient number of differences to permit the possibility of specific enzyme binding. If the clover leaf structures shown are correct, one would expect that in the paired stretches, the bases would be turned inward and inaccessible to enzyme recognition leaving the non-pairing stretches accessible. There are abundant differences even there. To date no sequences have been established for non-yeast tRNA's, but it is obvious that some differences must exist, for *E. coli* tRNA contains substantial amounts of thioridine which is not found in yeast tRNA.

One must conclude that all tRNA's specific for a particular amino acid contain one or more features in common and that variations in the ability to be aminoacylated with heterologous enzymes reflect either differences in the binding of tRNA to enzyme or binding in a non-productive mode. In either case, these variations must be due to structural variations outside the enzyme recognition area of the tRNA and to differences in the corresponding parts of the enzymes.

Surprisingly, the cross reactions are not at all reciprocal. Thus, starfish leucine enzyme esterifies *E. coli* tRNA excellently while *E. coli* leucine enzyme appears inert towards starfish tRNA. Both yeast enzymes are active towards *E. coli* tRNA while the reverse pair do not react well. Toadfish tRNA is very active with yeast leucine enzyme while the reverse pair is sluggish.

Some generalization may be valid. Both of the *E. coli* ligases discriminate strongly in favor of homologous (*E. coli*) tRNA while *E. coli* tRNA frequently reacts well with ligases from other sources. Species specifically in these reactions appears to reside more in the enzymes than in the tRNA's, perhaps a consequence of great inflexibility of some of the enzymes.

SUMMARY

1. Amino acid activating enzymes (aminoacyl tRNA ligases) specific for valine and leucine tRNA have been partially purified from four widely differing phyla, namely yeast, *E. coli*, starfish and toadfish.

2. The rates of aminoacylation of the four tRNA's have been determined using both the homologous enzyme and the three heterologous enzymes. In most cases there was appreciable cross-reaction, the *E. coli* enzyme and tRNA being in general least disposed to cross-react with the others.

3. In some cases the heterologous pair react more rapidly than the homologous. In one case this has been established as being a consequence of a much higher V_{\max} that overcomes a poorer enzyme tRNA association.

4. Both homologously and heterologously esterified tRNA's appear to be homogeneous as measured by the kinetics of hydrolysis.

5. We interpret these observations as indicating that all tRNA's specific for a particular amino acid have at least one common polynucleotide sequence, probably in an area of the chain which is not base-paired. The marked differences in the rates of esterification and in the rates of hydrolysis of the aminoacyl tRNA's reflect other structural variations in tRNA's that do not include the enzyme recognition site. In particular *E. coli* enzymes are least able to adapt themselves to heterologous tRNA's.

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ISOLATION AND COMPOSITION OF A LOW DENSITY LIPOPROTEIN FROM THE EGGS OF *ARBACIA PUNCTULATA*¹

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High and low density lipoproteins have been isolated from the egg cells of vertebrate species (Wallace, 1965; Fujii, 1960). Relatively few studies of invertebrate egg lipoproteins have been made. Wallace, Walker and Hauschka (1967) have isolated high density lipoproteins from the eggs of six species of crustaceans but no low density lipoproteins have been reported from invertebrate eggs. The present work describes the isolation and analysis of a water-soluble low density lipoprotein² from sea urchin eggs which contains 74% of lipid and has a molecular weight of about 10×10^6 . This protein bears a remarkable resemblance to the low density lipoproteins of mammalian plasma.

MATERIALS AND METHODS

The purple sea urchin, *Arbacia punctulata*, was collected during the months of July and August in the waters of Woods Hole. Eggs were obtained by injection of 0.5 M KCl into the body cavity adjacent to the lantern and were centrifuged twice in filtered sea water to remove the jelly coat.

The lipoprotein was isolated by mixing the eggs gently for 30 minutes at 0° with 20 volumes of 0.05 M sodium phosphate, pH 6.8. All subsequent operations were carried out at 0–10°. The mixture was centrifuged for 40 minutes at 100,000 *g* and any material floating to the top of the tube was removed. The remaining clear supernatant fluid was adjusted to a solvent density of 1.21 by the addition of NaBr and centrifuged overnight at 100,000 *g*. The top 10% of each tube was then harvested, a simple matter because of the color imparted to the lipoprotein by echinochrome. The combined top fractions were then diluted with 4 volumes of NaBr, density 1.21, and re-centrifuged overnight. The resulting top fractions constituted the final purified product. Before analysis, the lipoprotein was dialyzed against at least 2 changes of solvent, generally 0.1 M NaCl. Generally, prolonged storage was avoided but 1:10,000 merthiolate was added if preparations were to be kept longer than a week in the refrigerator.

After extraction with chloroform-methanol by the method of Folch *et al.*

¹This work was supported by U.S.P.H.S. grant HE-05285. Analytical ultracentrifugation was carried out by Mr. L. Robinson Hyde whose assistance is gratefully acknowledged. A preliminary report of this work has been made (Marsh, 1965).

²High density lipoproteins are here arbitrarily defined as those of hydrated density greater than 1.1 and less than 1.2, while low density lipoproteins are defined as those with densities less than 1.1 and greater than 1.01.

(1957), total lipid determinations were made by the method of Marsh and Weinstein (1966), using tristearin as a standard. Protein was measured by the Lowry method (1951) with bovine plasma albumin as a standard. Phospholipids were determined by Bartlett's method (1959) and chromatography of the phospholipid fraction was carried out as described by Abramson and Blecher (1964). Total protein-bound carbohydrate was measured on the lipid-free protein after washing the precipitate with cold 5% trichloroacetic acid to remove chloride ions. The protein precipitate was then heated for 2 hours at 100° in 1 N H₂SO₄ and the carbohydrate content of the extract measured by the anthrone method (Ashwell, 1957) using galactose as a standard. Additional extraction of the precipitate or treatment with 2 N HCl for 3 hours at 100° did not significantly increase the carbohydrate yield. The neutral carbohydrate fraction contained 96% mannose and 4% arabinose as determined by gas-liquid chromatography of the alcohol hexacetates (Sawardeker, Sloneker and Jeanes, 1965). The amino acid composition of the delipidized protein was measured with an amino acid analyzer

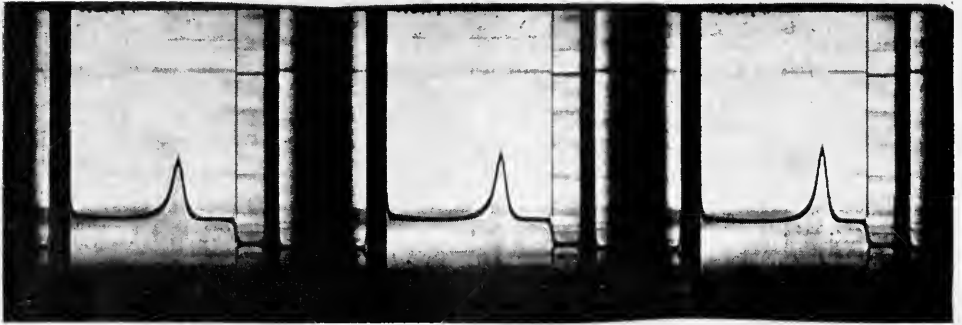


FIGURE 1. Sedimentation of *Arbacia* egg low density lipoprotein in 0.03% NaBr, pH 6, at 42,040 rpm. Sedimentation is from right to left, the interval between pictures is 4 minutes, and the bar angle 60°.

after hydrolysis in 6 N HCl for 23 hours at 110° in a sealed tube. Tryptophan was determined colorimetrically (Opienska-Blauth, Charezinski and Berbec, 1963) on a solution of the delipidized protein in 0.1 N NaOH. The fatty acid composition of the lipid moiety was determined by gas-liquid chromatography of the methyl esters after saponification in methanolic KOH (James, 1960). Identification of the acids was made by comparison of the retention times with known standards. Free and esterified cholesterol was measured by the method of Sperry and Webb (1950).

For the determination of molecular weight, the flotation rate (*S*_f) was measured in 24% NaBr (density 1.154 at 24°). The diffusion constant was measured in the same solvent at 6° in the electrophoresis cell of the Beckman-Spinco apparatus. No extrapolation to zero concentration was made in the measurement of *S*_f and *D*. However, the concentration employed was 0.5% and according to the data of Wallace, Walker and Hauschka (1967), the correction would be of the order of 2%. An approximate value for the partial specific volume (0.915) was obtained by determining the density of a solution of NaBr in which about

TABLE I
Lipid-to-protein ratio of Arbacia egg lipoprotein

Preparation number	Lipid:protein ratio
1	3.20
2	2.84
3	2.29
4	2.71
5	3.10*
6	2.65
Mean \pm S.E.M.	2.80 \pm 0.14

* Measured 24 hours after fertilization. In preparation 4, the method of determining total lipid was checked by gravimetric analysis and the resulting lipid:protein ratio was 2.78.

half the molecules sedimented and half floated at the same time at 59,780 rpm in the analytical ultracentrifuge. It was estimated that the figure for partial specific volume was only accurate within $\pm 20\%$. With this information and appropriate temperature corrections, the molecular weight was calculated by the Svedberg equation.

RESULTS

The isolated lipoprotein appeared free of ordinary proteins. When floated in the analytical ultracentrifuge at density 1.2, no sedimenting protein was seen. When dialyzed against water, the protein sedimented as a single peak with an $S_{20,W}$ of 15.2 (Fig. 1). The lipid-to-protein ratios of 6 different preparations are given in Table I. When subjected to column chromatography on Sephadex G-200, the lipoprotein emerged with the solvent front, indicating a molecular weight higher than about 500,000, and no residual protein was eluted from the column. On disc electrophoresis with 5% polyacrylamide gel, the lipoprotein failed to enter the gel, another indication of high molecular weight. No protein was found in the gel proper after staining. Electrophoresis on cellulose acetate in 0.05 M barbital buffer at pH 8.6 revealed a single fairly broad band which migrated 1 cm. per hour towards the anode, a migration rate one-half that of a control sample of bovine plasma albumin. The lipoprotein was not retained on a column of carboxy-methyl Sephadex at pH 6.0, indicating an isoelectric point below that pH.

TABLE II
*Lipid composition of Arbacia egg lipoprotein**

Lipid class	% of Total lipid
Neutral lipid	62.8
Phospholipid	28.0
Sterol	8.2
Sterol ester	1.0

* The values shown are averages of duplicate determinations on two separate preparations.

TABLE III
*Composition of Arbacia egg lipoprotein**

Component	% of total weight
Protein	28.5
Lipid	65.2
Carbohydrate	5.8
Echinochrome	0.5
Protein-bound P	Absent

* Preparation 4 of Table I. Echinochrome was measured spectrophotometrically. Protein-bound P was measured after sulfuric acid-hydrogen peroxide digestion.

Under the conditions of isolation, the lipoprotein accounted for 70–90% of the lipid in the supernatant after the initial centrifugation in 0.05 *M* buffer. In two experiments, the yield of lipoprotein was 15.7 and 17.2 mg. of lipid per ml. of packed eggs, which accounts for approximately $\frac{1}{3}$ of the total lipid of the cell. The overall lipid composition is given in Table II.

A complete analysis of one lipoprotein preparation is given in Table III. No protein-bound phosphorus was detected (Table III).

The fatty acid composition of the total lipid is shown in Table IV and the amino acid composition of the protein is shown in Table V. For purposes of comparison, the amino acid composition of human low density (B) lipoprotein as calculated from the data of Margolis and Langdon (1966) is also given.

No quantitative analysis of the lipid classes present was made, but judging by the extent of phosphate color given on thin layer plates, the major phospholipid was phosphatidyl choline, with phosphatidyl serine and ethanolamine present in lesser amounts. The lysophosphatides of these were also detected, along with

TABLE IV
Fatty acid composition of Arbacia egg lipoprotein lipid

Fatty acid	Relative retention time*	% of Total
12:0	0.18	0.2
14:0	0.31	7.1
14:1	0.41	3.1
16:0	0.55	25.4
16:1	0.68	14.7
?	0.83	0.5
18:0	1.00	6.8
18:1	1.19	9.7
18:2	1.57	4.2
20:0	1.82	2.6
18:3	2.08	10.9
?	2.42	2.2
22:1	3.90	11.2
?	5.40	1.3

* Relative to methyl stearate, on a 4 ft. column of polyethylene glycol succinate at 167° and 7.5 lbs. pressure of argon. The Pye instrument with an argon ionization detector was used.

TABLE V

*Amino acid composition of Arbacia egg lipoprotein, compared with human plasma low density lipoprotein**

Amino acid	% of Total <i>Arbacia</i> lipoprotein	Human B-lipoprotein*
Aspartic	11.1	10.9
Threonine	6.8	5.8
Serine	7.1	6.6
Glutamic	10.8	14.0
Proline	2.6	3.0
Glycine	2.6	2.6
Alanine	5.4	5.4
$\frac{1}{2}$ Cystine	6.1	0.6
Methionine	—	2.1
Isoleucine	5.8	5.8
Leucine	9.5	9.5
Tyrosine	3.1	4.1
Phenylalanine	6.8	6.6
Lysine	5.3	8.0
Histidine	4.8	3.0
Arginine	8.6	4.8
Tryptophan	1.5	0.9

* Data for human plasma B-lipoprotein calculated from the data of Margolis and Langdon (1966).

phosphatidic acid, but these were all believed to be breakdown products since no special precautions were taken. There were two unknown spots present. Although it had the same R_f as tristearin in two solvent systems on thin layer chromatography, the neutral lipid was not positively identified as triglyceride, since marine lipids are known to contain alkoxydiglycerides.

Some physical constants of the *Arbacia* egg lipoprotein are given in Table VI, and it can be seen that the molecule has a very high molecular weight, of the order of ten million.

DISCUSSION

Hen's egg yolk contains three main lipoproteins, a low density lipoprotein; lipovitellin (a high density lipoprotein as we have defined it) and a lipovitellin-phosvitin complex (Fujii, 1960). The lipoproteins of several classes of marine eggs described by Fujii (1960) and the crustacean egg lipoproteins isolated by

TABLE VI

*Some physical constants of Arbacia egg lipoproteins**

S _{20,W} = 15.2
D = 0.79×10^{-7} cm. ² sec. in 24% NaBr at 6°
Sf = 20.1 in 24% NaBr at 24°
Mol. wt. = 10×10^6

* See texts for details of measurements.

Wallace *et al.* (1967) and by Zagalsky *et al.* (1967) are high density lipoproteins. With the isolation of a low density lipoprotein in the present work, it now appears that both vertebrate and invertebrate eggs, as well as mammalian plasma, contain lipoproteins of both high and low density.

The *Arbacia* egg low density lipoprotein described here has striking similarities to that of human plasma. Its molecular weight falls in the same range of from 2 to 20 million. It is not a phosphoprotein, as are the hen's egg lipovitellins. In this respect, it resembles the crustacean egg lipoproteins. Like human plasma lipoproteins, it contains covalently bonded carbohydrate. No report of the carbohydrate content of the crustacean egg lipoproteins was made by Wallace *et al.* (1967) nor were the other egg lipoproteins isolated by Fujii (1960) analyzed for carbohydrate. Zagalsky, Cheesman and Ceccaldi (1967) have shown that the carotenoid-containing lipoprotein of the eggs of *Plesionika edwardsi*, molecular weight of about 600,000, contains 27% lipid and that the lipid-free protein contains 2.6% of carbohydrate. Mannose and glucosamine were shown to be present. The amino acid composition of the *Arbacia* egg low density lipoprotein is strikingly similar to that of human B-lipoprotein, except for the absence of methionine. Both proteins do not have large amounts of non-polar residues, and the implications of this have been discussed by Margolis and Langdon (1966).

The lipid composition again resembles that of human B-lipoprotein, since the neutral lipids made up the main component (Table II). The fatty acid composition is not remarkable, resembling marine lipids in its content of C-20 and C-22, especially the latter.

Recently, Malkin, Mangan and Gross (1965) have described a crystalline 27S protein isolated from *Arbacia* eggs, having a molecular weight of 894,000 which they believe to be derived from the breakdown of yolk granules. No information concerning lipid content was given. It is possible that their protein is a high density lipoprotein, similar to that having a molecular weight of 2.5×10^6 isolated by Fujii (1960) from cuttle-fish eggs.

Removal of lipid from the *Arbacia* lipoprotein by organic solvents at low temperature (-20°) renders the protein insoluble in water. This is quite typical behavior for human plasma B-lipoprotein. Although on one occasion it appeared that the resulting protein was soluble in 6 *M* urea (Marsh, 1965), we have not been able to repeat this observation, probably owing to denaturation problems. The minimal number of peptide chains has not been determined by end group analysis as yet.

The similarity between the storage low density lipoprotein of *Arbacia* eggs and human plasma low density lipoproteins leads to the suggestion that the latter may have evolved from storage lipoproteins. Future studies of the comparative biochemistry of egg cell lipoproteins should lead to much valuable information about water-soluble lipoproteins in general.

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THE DEVELOPMENTAL POTENTIALITY OF THE
LIVER-RNA-TREATED POSTERIOR
PRIMITIVE STREAK IN THE
CHICK EMBRYO¹

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Previous experiments have shown that RNA-treated posterior third of the primitive streak, here abbreviated to TPS, was capable of developing into highly organized kinds of tissue. The type of tissue produced varied according to the kind of RNA used (Sanyal and Niu, 1966). Brain RNA induced the formation of neural tissue. Kidney and heart RNAs seldom induced neural formation but caused the TPS to self-differentiate into tubular and vesicular structures, respectively. On the other hand, liver (L) RNA not only induced epiblasts to become neural tissue but also made the TPS self-develop into some organized but as yet unidentifiable tissue. The presence of the latter tissue indicated the need for prolonged cultivation. In this respect, the technique (New, 1955) used was limited. In the experiments now to be reported the L-RNA-treated TPS was implanted into the coelom of 2-2 ½-day chick embryo (Hara, 1961). The choice of intracoelomic grafting was made on three grounds: (1) the implant would have the time needed to develop its acquired capability, (2) the implant would be in contact with host's incompetent cells and thus would only undergo self-differentiation, and (3) the implant would be made to the host at an early stage when visceral organs, particularly endodermal derivatives, had not yet appeared. Both L-RNA and L-protein (serum albumin, abbreviated S.A.) were employed to treat the TPS. The aim of this paper is to present data showing that: (a) S.A. has no effect on the development of TPS, (b) L-RNA stimulates both growth and differentiation, and (c) RNase-treated L-RNA reduced gut formation but promotes growth of the implants and development of feather buds even better than RNA.

MATERIALS AND METHODS

Preparation of TPS

Fertilized eggs of White Leghorn chickens were obtained from Shaw Hatchery, West Chester, Pa. They were incubated for 15-18 hours at 38° C. in a forced draft incubator. The primitive streak of the blastoderm, 1.2-1.6 mm. in length and 0.2 mm. in width, was excised and stretched on agar dishes containing modified

¹ Supported in part by grants from The Population Council, New York, and National Science Foundation.

² State Department Scholar from Finland for the year 1965-1966.

Locke solution (a mixture of 100 ml. NaCl (9.43 g./l.), 3.7 ml. KCL (1.2 g./100 ml.), 2.1 ml. CaCl₂ (2.36 g./100 ml) and 1 ml. glucose (0.2 g./100 ml.). The posterior third, 0.4–0.5 mm., was cut off and immediately transferred to the Locke solution. Accumulated pieces were divided into groups and transferred to the Locke solution (2 ml.) with or without S.A. (4 mg./ml.), L-RNA (O.D.₂₆₀ m μ 80/ml.) or RNase-treated L-RNA (1 mg. RNase in 2 ml. with O.D. 160). They were kept in a cold room (2–4° C.) with an occasional stir for 15–18 hours (overnight). Bovine S.A. (fraction V) was obtained from Armour and pancreatic RNase through Worthington.

Implantation of TPS

Both untreated (control) and treated TPS were implanted into the coelomic cavity of 2–2 $\frac{1}{2}$ -day-old chick embryos according to the procedure of Hara with a minor modification, namely, when the host was slightly older, the graft was implanted into the posterior rather than anterior end of the cavity. This modification was made for the sole purpose of increasing the number of the recipient embryos. There was no difference in the rate of recovery, growth and differentiation between the anterior and posterior grafts. All implants were harvested at the 7th–9th day and fixed in Bouin's solution. Sections were 8 μ thick and stained with hematoxylin and eosin.

Preparation of RNA

RNA was isolated from calf liver. Thanks to Messrs. Collis and Owen of Cross Brothers Meat Packing Co., the liver was quickly removed from the slaughtered calf, sliced and put in ice-cold sucrose solution (0.25 *M* + CaCl₂, 0.003 *M*). The procedure of isolation has been published elsewhere (Hillman and Niu, 1963).

The glycogen of the RNA preparation was removed by Spinco centrifugation (30,000 RPM for 30 minutes) and the phenol extracted by repeated ether (or ethanol) washings, or by bubbling N₂ through. The UV absorption spectrum of L-RNA was typical of nucleic acids with maximal absorption at 260 m μ and minimal 230. The ratio of A₂₆₀/A₂₃₀ exceeded 2 and that of 280/260 varied between 0.45 and 0.5. Contamination of protein, DNA and polysaccharides was routinely estimated. The amount was less than 1%. The test with orcinol yielded a green color. When a preparation did not meet the criteria given here, it would not be used for experimentation.

For functional studies of RNA, it is of prime importance to isolate the RNA within the shortest time and to use it as soon as possible. Lyophilization and storage for longer periods in a deep freezer resulted in the loss of activity except the property pertaining to growth.

Digestion of L-RNA was carried out in 2 ml. of Locke solution containing RNA and 1 mg. of RNase. The mixture was incubated at 37° C. for 1 hour. This resulted in an increase of O.D. by 35%. To an aliquot of the sample, an equal volume of ice-cold 4% perchloric acid was added. The acid-precipitable material was the undigested portion of RNA. It amounted to 15–20% of the original.

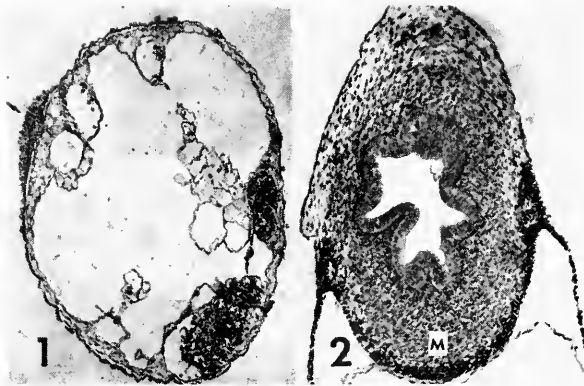


FIGURE 1. A section through the implant of control. Note the presence of endodermal epithelial lining on the outer surface (at 10:00 o'clock) and two islands of blood cells on the lower left. $\times 120$.

FIGURE 2. A section through the L-RNA-treated implant showing intestinal vesicle. M, circular muscle. $\times 120$.

RESULTS

Development of the Untreated TPS (controls)

A total of 60 fragments was excised. After being kept in the modified Locke solution at 2–4° C. for 15–18 hours, they were implanted into the coelomic cavity. The explants were loose and fell readily into tiny pieces, thus causing some difficulty during the process of implantation. Histological examination revealed, however, that cells of the central mass were normal and those in the outermost layer became swollen or even broken. Of the 60 controls, 23 (38%) were recovered.

The implants are either free or attached to the host tissue. Histological examination of the controls showed that the structures produced were limited to tissue with or without epidermal and/or endodermal epithelial lining. There was no appreciable difference between the untreated (control) and the S.A.-treated series, thus showing that the slight damage to the outermost layer of cells, caused by keeping the explants overnight in the cold room, had no adverse effect on their developmental potentiality. To further demonstrate this point,

TABLE I
Developmental potentiality of the control posterior primitive streaks

Conditions of explant	Growth		Differentiation				
	Attachment to host	Size increased 3 or more times	Total	Epidermal	Epidermal-teather buds	Endodermal	Gut
Fresh controls (57)	14	3	7	5	1	5	0
Aged controls (60)	14	6	9	7	3	7	1

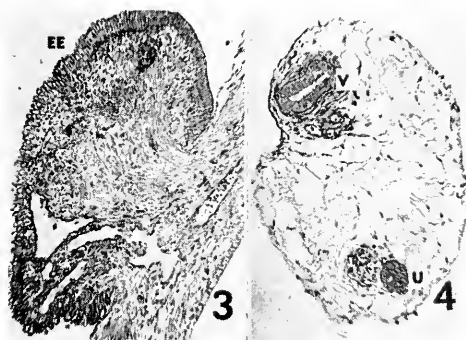


FIGURE 3. A section through a tiny portion of a big skin vesicle, developed from a L-RNA-treated TPS. The attached mass of connective tissue (see also Fig. 7) is covered by a distinctive layer of endodermal epithelium, EE. $\times 100$.

FIGURE 4. A section through a poorly developed vesicle of the L-RNA-treated implants. V, intestinal vesicle and U, unidentified tissue. $\times 100$.

10 anterior primitive streaks including Hensen's node were kept in the cold room for 15–18 hours. The explants showed similar loose consistency as noted in the TPS. All developed into large or very large grafts containing skin, brain, intestine, cartilage, etc.

Usually the implants were small but compact. When somewhat larger, they contained some connective tissue, blood (Fig. 1) and occasionally smooth muscle fibers. In cases when they were attached to the host, they were covered with poorly organized, vacuolated, spherical, or more or less elongated cells. These cells formed a definite layer in some cases and had the appearance of intestinal epithelium (arrow in Fig. 1).

TABLE II

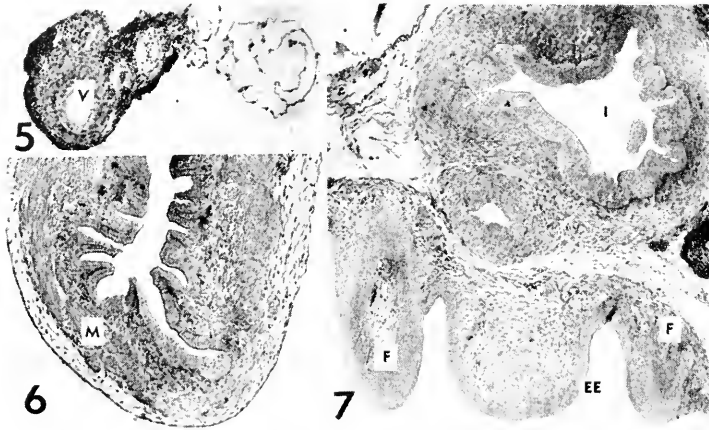
Growth and differentiation of the posterior third primitive streak treated with and without liver protein (S.A.) RNA and its hydrolytic products (A = total number, B = percentage of the total & C = percentage of the recovered)

Experimental series	Untreated (control)			Treated with S.A.			Treated with RNA			Treated with RNase-digested RNA		
	A	B	C	A	B	C	A	B	C	A	B	C
1. Implants	60	—	—	33	—	—	46	—	—	29	—	—
2. Recovered	23	38	—	13	39	—	27	59	—	23	79	—
3. Attached to host tissue	14	23	61	8	24	62	21	46	76	12	41	52
4. Size increased 3 or more times	6	10	26	2	6	15	16	35	59	14	48	61
5. Differentiation	9	15	39	6	18	46	20	43	74	14	48	56
6. Epidermal	7	12	30	3	9	23	14	30	52	9	31	39
7. Epidermal with feather buds	3	5	13	2	6	15	9	20	33	9	31	39
8. Endodermal	7	12	30	5	15	29	17	37	63	7	24	30
9. Gut	1	2	4	2	6	15	9	20	33	2	7	9

TPS were also implanted immediately into the coelomic cavity after extirpation. Of the 57 fresh controls, 37 (65%) were recovered which is superior to the 38% of the aged controls mentioned above. However, when the recovered implants were examined for growth and differentiation, the aged controls were slightly better than the fresh controls (see Table I). Although the reason for this improvement is not known, the observation undoubtedly adds support to the contention that slight damage to the explants had no adverse effect on their developmental potentiality.

Development of the S.A.-treated TPS

Thirty-three explants were kept in Locke solution with S.A. added. At the time of implantation, they were compact and looked healthy. However, the growth



FIGURES 5 and 6. Sections of two vesicles from the L-RNA-treated implants showing variations of intestinal structures. V, poorly developed intestine; M, circular muscle. $\times 100$.

FIGURE 7. A section through a big skin vesicle obtained from RNase-treated RNA series. The lumen of the vesicle is toward the bottom. The attached mass has developed into distinctive intestinal tubes (I). EE, epidermal epithelium. I, intestinal tube, and F, feather follicle. $\times 120$.

and differentiation of this series were remarkably similar to those of the control (Table II). It seems, therefore, that the developmental potentiality of the control and S.A.-treated TPS is similar and thus considered to be intrinsic.

Development of the L-RNA-treated TPS

At the time of implantation, the explants were similar to those of the S.A. series. A total of 46 implants were made and 27 (59%) were recovered. Most of them were attached to the host. More than half showed good growth. The overall differentiation was 43% of the total. About half (14) of the available grafts (27 cases) contained epidermal epithelium. Nine of the 14 appeared

in the form of large vesicles with feather buds, *i.e.* 64% of epidermal vesicles developed feather buds. Of the 27 recovered implants, 17 developed into tissue with endodermal epithelial lining (Figs. 3-5) and 9 into intestinal vesicles (Fig. 2) or well formed intestinal tubes (Fig. 6). It should be noted that differentiation of both epidermal and endodermal epithelial cells was usually found in the same graft. These grafts were well differentiated and rather large.

Development of the TPS treated by RNase-digested L-RNA

Incubation of L-RNA with RNase resulted in a loss of 85% of RNA. The RNase-treated L-RNA (RNase, RNA and its hydrolytic products) was used to treat 29 explants. Before implantation, they were compact and healthy. Twenty-three implants were recovered, thus providing the highest rate of recovery (79%) in our experimental series. The rate of growth was also the best (Fig. 7). The overall rate of differentiation and the formation of epidermal epithelium were similar to the RNA series. However, the development of endodermal epithelium and gut particularly decreased from 37% to 24% (a loss of 35%) and from 20% to 7% (a loss of 65%), respectively.

The effect of plain RNase on TPS was also tested. Seven explants were kept in Locke solution containing RNase. None of the implants was recovered after 10 days in the peritoneal cavity of the developing chick embryos.

DISCUSSION

The TPS used for implantation was taken from the region where epiblasts and hypoblasts merge into a mass of cells. They are destined to give rise to connective tissue, and epidermal and endodermal epithelium (Willier and Rawles, 1931a), but are capable of being induced to develop into tissues other than epithelial cells. However, when they were grafted into the developing blastoderm, earlier experiments showed that 72% of the implants could not be located at the site of implantation and the remaining 28% appeared as mesenchyme, seldom developing into a neural tube. In the foregoing experiments, the explants were grafted into the coelomic cavity. The rate of recovery was improved (38%, Table II). Among the chemically treated series, S.A. did not alter the rate of recovery (39%), nor development. L-RNA and its hydrolytic products increased the recovery rate up to 59% and 79%, respectively. The improvement was accompanied by better growth and differentiation.

The implant of control groups developed into connective tissue with or without epidermal epithelium and/or endodermal epithelial lining. The formation of ectodermal (feather buds) or endodermal derivative (gut) was rare. Therefore, the developmental potentiality of TPS was better realized in the coelomic cavity of 2 $\frac{1}{2}$ -day chick embryo than with the *in vitro* technique (New, 1955). This finding is in accord with the recent observation that the type of medium used is extremely important in the study of differentiation (Spratt and Hass, 1967).

The developmental potentiality of the control and chemically treated TPS was compared categorically in terms of total implants. It can be seen in Table II that the data of the control resemble the S.A.-treated series. Both differ significantly from RNA and RNA-digest series. The latter two series stimulate

both growth and differentiation. Of particular interest is the finding that gut formation is increased by RNA, and feather bud formation can better be facilitated by the hydrolytic products than RNA itself. Feather bud is one of the ectodermal derivatives. Its frequency of development in the epidermal vesicles is 43% of the control, 66% S.A., 64% RNA and 100% RNA digest series. Furthermore the RNA digests cut down the developing rate of endodermal epithelium (from 37% to 24%) and gut (from 20% to 7%), but increase the rate of recovery (from 59% to 79%) and the number of sizable implants (from 35% to 48%). Therefore, the reduction in development of endodermal structure is related to the loss of RNA (about 85%) and the digests are responsible for the gain in feather bud formation and growth of the implants as well. The action of the latter is not specific and belongs to the same category as reported by Sengel (1964). They perhaps contribute to the enrichment of the nutrients. It may be relevant to mention here that much of the exogenous RNA remains intact in the recipient cells (Niu, Niu and Guha, 1968) and some are being hydrolyzed which, in turn, may be responsible for the non-specific action of RNA.

Gut was found once in 60 implants of the controls and twice in 33 S.A. and 29 hydrolyzed RNA series, respectively. This low frequency could hardly be used to argue for the action of S.A. or RNA digests on gut differentiation. Instead we consider this gut-forming ability as one of the intrinsic properties of TPS. L-RNA was capable of increasing the rate of gut formation three-fold (from 6% to 20%). This amounts to 45% of the cases with differentiation.

A double check on the developmental potentiality was made by using recovered implants (see Column C of Table II). The rate of attachment (row 3) is compared with that of the growth (row 4) and differentiation (row 5) of the same series. It can be seen that the rate of growth and differentiation is notably lowered in the control and S.A. series. On the other hand, the RNA and RNA-digest series have maintained at about the same level as the attachment. The types of differentiation are derivatives of the ectoderm and endoderm. The ectodermal derivatives of the control are similar to the S.A. series (rows 6 and 7), but strikingly different from the RNA and RNA-digest series. As to the endodermal derivatives (rows 8 and 9) the rate of development is approximately the same in the control, S.A. and RNA-digest series *but* less than half of the RNA series. In other words, the RNA used in the present experiment stimulates the formation of endodermal derivative, and this function requires the intact macromolecule of RNA. The promotion of ectodermal formation by RNA is non-specific because hydrolytic products of RNA can do almost equally well.

Liver is an organ derived from the ventral diverticulum of the gut. In the chick, its development is rather complicated and requires an inducing action from the heart rudiment (Willier and Rawles, 1931b). The fact that L-RNA can indeed stimulate gut formation is a step forward in our study on the possibility of the directed formation of specific tissue (Hillman and Niu, 1963; Niu and Leikola, 1966; Sanyal and Niu, 1966). Should L-RNA carry the message of the liver (Niu, Cordova and Niu, 1961; Zimmerman, Zoller and Turba, 1963), one immediate question is the condition under which the development of gut and liver can separately be achieved. Experiments dealing with this and other related problems are in progress.

SUMMARY

The posterior third of the primitive streak was implanted into the 2 $\frac{1}{2}$ -day chick embryo. Both growth and differentiation of the untreated implants were similar to those of the serum albumin series. Treatment with liver RNA, however, resulted in a significant increase in the rate of growth and differentiation. When liver RNA was incubated with RNase, the products furthered the growth and feather bud formation, but reduced the differentiation of the endodermal derivatives, especially gut.

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THE BIOLOGY OF FERTILIZATION AND BROOD PROTECTION IN SPIRORBIS (LAEOSPIRA) MORCHI¹

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The described species of the serpulid genus *Spirorbis* are hermaphroditic and exhibit brood protection. Alexander Agassiz (1866) appears to be the first investigator to offer information on the biology of fertilization in the genus. Agassiz, who examined *Spirorbis spirillum*, claimed that the embryos reached quite an advanced stage within the body of the parent before the brood sac was deposited within the cavity of the tube. Fewkes (1885) examined what he thought to be *Spirorbis borecalis* and observed that the eggs were laid in strings within the parental tube; however, he was not able to confirm Agassiz' claim that development initially occurs within the body of the parent. According to Schively (1897), who worked with what she believed to be *Spirorbis borecalis*, the eggs pass into the body cavity and then into the operculum where fertilization takes place and a capsule is secreted. Schively further claimed that once the egg capsule was secreted, the capsule passed through an opening in the operculum and was placed in the "mid-dorsal furrow." It seems quite likely that both Fewkes and Schively were working with *Spirorbis spirillum* and not *Spirorbis borecalis*. Zur Loye (1908) believed that fertilization in *Spirorbis borecalis* occurred externally within the parental tube. Abe (1943), working with a species which he identified as *Spirorbis argutus*, reported that fertilization probably took place internally. Recently, Gee and Williams (1965) have reported that fertilization in *Spirorbis borecalis* and *Spirorbis pagenstecheri* occurs externally and have presented evidence that self-fertilization is possible in both species.

Brood protection in the genus *Spirorbis* takes place either within the parental tube or within a modified operculum. As far as is known, the mode of brood protection is species-specific. Thorson (1946) has claimed that *Spirorbis granulatus*, which is characterized as an operculum brooder (Canllery and Mesnil, 1897; Bergan, 1953b), broods in the tube and suggested that the mode of brood protection varies within the species; however, Thorson (personal communication) believes that he may have been mistaken as to the identity of the species. Of the two types of brood protection, opercular brood protection is the most specialized and has been considered a recent development in the evolution of the genus (Elsler, 1907; Borg, 1917; Gravier, 1923). Bergan (1953a) observed the presence of a pore in the operculum of *Spirorbis granulatus*, but was of the opinion that it was too small to allow the passage of eggs and did not pursue its significance. To date, there has been no attempt to explain how spawned oocytes are transferred to the opercular brood pouch.

¹ Supported, in part, by predoctoral fellowship 1-F1-GM-20, 593-01 from U.S.P.H.S.

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The present paper deals with the biology of fertilization and opercular brood protection in *Spirorbis (Lacospira) morchi* Levinsen. The observations on fertilization are an elaboration of those presented earlier in abstract (Potswald, 1964).

MATERIALS AND METHODS

Spirorbis morchi adults were collected periodically throughout the year, from 1960 to 1963, in Argyle Creek on San Juan Island, Washington. The animals were found on rock and shell and often in association with *Spirorbis (Paradexiospira) vitreus*. In Argyle Creek, *Sp. morchi* breeds the year around; consequently, developmental stages were always available.

In order to determine whether animals raised in strict isolation are capable of self-reproduction, individual larvae were isolated and cultured. This was accomplished by placing larvae in polyethylene ice-cube trays, one larva per cube, and allowing them to settle. Each cube was given a number, thereby allowing complete records to be kept for each isolate. The sea water used in these cultures was filtered to prevent larvae, that might be in the sea water system, from settling in the containers used to store the water, and was then allowed to stand for a minimum of three weeks before using. Such water was considered to be sperm-free and will be subsequently referred to as sperm-free sea water. The sea water was removed periodically, by means of a pipette attached to an aspirator, and replaced. A new pipette was used for each cube, thus eliminating the possibility of cross-contamination by introducing sea water from one cube into that of another. Initially, food was supplied in the form of *Nitzschia* sp.; however, the cultures fouled very rapidly even upon the addition of small numbers of diatoms. It was found that the animals maintained themselves on protozoa and bacteria present in the water so the addition of diatoms was discontinued.

Larvae used in the isolation experiments were artificially released from brood pouches. Opercular brood pouches, which contained actively moving larvae, were selected and removed with #5 watchmaker's forceps. Once removed, the opercular ampullae were torn open with forceps or size 0 insect pins. The larvae thus released showed the same behavioral responses and settled in about the same length of time as naturally released larvae.

The histological techniques used in the present study were reported previously (Potswald, 1967). In addition, the chloranilic acid method of Carr *et al.* (1961) was used for the demonstration of calcium.

OBSERVATIONS ON FERTILIZATION

Attempts at controlled fertilization

In the summer of 1961, while in residence at the Friday Harbor Laboratories, several experiments were carried out in an attempt to bring about controlled artificial fertilization. Adults with gravid female segments were removed from their tubes, individually isolated in syracuse dishes, and gently washed with two or three changes of pasteurized sea water to remove any extraneous sperm that may be adhering to the body. Male and female segments were then jabbed a few times with a size 0 insect pin which caused the longitudinal muscles of the abdomen to contract strongly and spasmodically, resulting in the release of gametes through

ruptures in the body wall. After a minute or two, the oocytes were removed by means of a pipette and transferred through several changes of pasteurized sea water to prevent possible polyspermy. Periodic examination with the compound microscope revealed that in over half the animals tested, activation had been initiated, as manifested by the lifting of a fertilization envelope; however, in none of the cases observed was there germinal vesicle breakdown accompanied by polar body formation. The same results were obtained when sperm from another individual were added. In all cases, the sperm were active and the oocytes, as judged by morphological criteria, appeared to be "ripe." It is possible that physiological maturation of the oocytes precedes natural spawning by only a matter of hours as has been shown by Howie (1961) for *Arenicola marina*.

Isolation of larvae

A second approach involved the use of artificially released larvae which were individually isolated and raised in sperm-free sea water (see Materials and Methods). Seventy larvae were isolated in the spring and summer of 1961. The mor-

TABLE I
Brooding record for Spirorbis morchi adults raised from larvae in isolation

Animal	Date isolated as larva	Date found brooding	Age at first brooding	No. of embryos	Remarks
I-12	6/18 61	9 06 62	15 mos.	3	Non-viable brood
I- 6	4 07 61	9/20 62	17 mos.	4	Non-viable brood
I-15	7/21 61	12 8 62	17 mos.	5	Non-viable brood
III- 4	7 03 61	2 22 63	14 mos.	3	Larval release & settlement
		9 06 62		4	Larval release & settlement
		10 22 62		5	Larval release & settlement
		12 13 62		?	Operculum & brood shed before completion of development
		3 09 63		?	Non-viable brood

tality rate was quite high during the first year and as of August, 1963, there were only 16 survivors. Of these, only four produced broods and as can be seen from the data summarized in Table I, two of the four isolates produced more than one brood.

Among the four animals which were able successfully to mature and spawn gametes, only a small number of eggs per brood were produced. This small number, and the fact that only $\frac{1}{4}$ of the survivors were able to spawn at all, can probably be attributed to nutritional factors. As mentioned earlier, attempts to supply food, in the form of diatoms, were not successful, and therefore discontinued. Obviously, settled animals can be maintained on bacteria and protozoa present in the sea water; however, only a few will be able to mature a small number of gametes under these conditions. At the time the cultures were discontinued (March, 1964), 12 of the original isolates still survived and ranged from 31 to 33 months in age. One of the isolates was sectioned and although it had the adult number of segments, all abdominal segments were completely agametic.

The cause of non-viable brood formation is not known. In the above cases, it is quite possible that mature sperm were not formed and, consequently, the oocytes not fertilized. It should be pointed out, however, that non-viable broods are often found in animals taken from the field. The fact that viable larvae, capable of normal settlement and metamorphosis, were produced from animals isolated as larvae is conclusive proof that in *Sp. morchi*, reproduction in isolation is possible. Had more effort and time been spent in finding a proper food source, it might be assumed that all of the isolates would have produced viable broods, probably over quite an expanse of time.

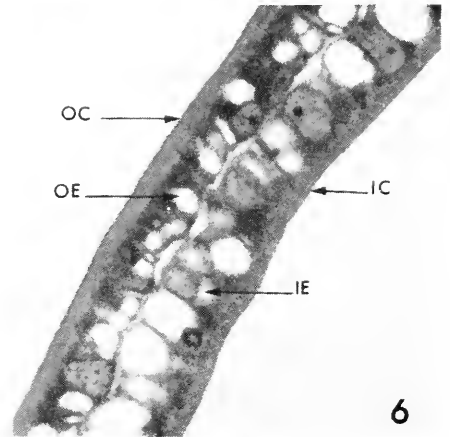
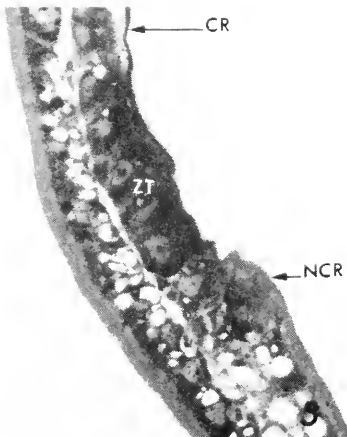
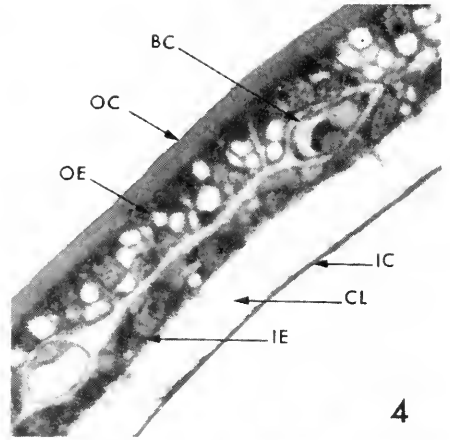
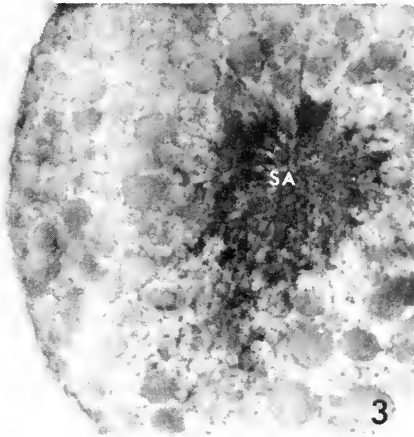
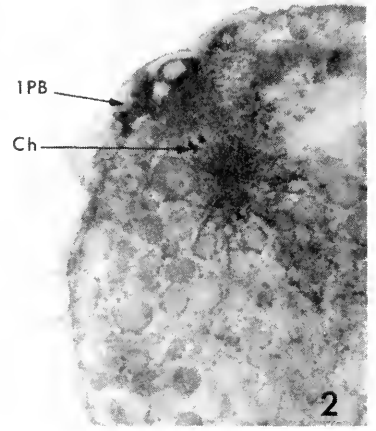
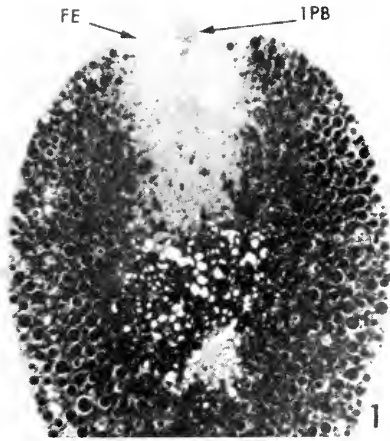
Isolation of adults

To test sperm viability, contents of male segments were artificially released, placed in filtered sea water, kept at about 12° C., and periodically examined with the compound microscope. Mature sperm remain active during the first eight

TABLE II
Isolation of Spirorbis morchi adults

Animal	Date of isolation	Date of spawning	Time elapsed since isolation in days	Remarks
1	6/18/62	7/05/62	17	Arrested development
2	6/18/62	6/23/62	5	Larvae released 7/24/62
*3	6/18/62	6/22/62	4	Larvae released 7/24/62
4	6/19/62	6/22/62	3	Arrested development
5	6/19/62	7/09/62	20	Arrested development
6	6/22/62	6/25/62	3	Arrested development
7	6/26/62	7/01/62	5	Larvae released 7/26/62
8	6/29/62	7/01/62	2	Larvae released 8/01/62
9	6/29/62	7/04/62	5	Arrested development
10	7/05/62	7/13/62	8	Larvae released 7/12/62
*11	7/12/62	7/26/62	14	Arrested development
12	2/07/63	2/14/63	7	Arrested development
13	2/07/63	2/17/63	10	Arrested development
*14	2/07/63	2/17/63	10	Larvae released 3/25/63
15	2/07/63	3/13/63	34	Arrested development
		4/06/63	58	Fixed 4/06/63
16	2/07/63	2/19/63	12	Larvae released 3/23/63
		3/27/63	48	Larvae released 4/25/63
17	2/07/63	3/12/63	33	Arrested development
18	2/07/63	2/13/63	6	Arrested development
		3/20/63	41	Larvae released 4/19/63
19	2/07/63	2/17/63	10	Arrested development
20	2/07/63	2/12/63	5	Larvae released 3/20/63
		3/27/63	48	Arrested development
21	2/07/63	3/20/63	13	Arrested development
22	2/07/63	3/09/63	30	Larvae released 4/13/63
23	2/07/63	3/06/63	27	Arrested development
*24	8/12/63	8/14/63	2	Larvae released 9/24/63
*25	8/12/63	9/05/63	25	Larvae released 10/17/63
		11/7/63	84	Non-viable
26	8/19/63	8/22/63	3	Arrested development

* Animal left in tube.



FIGURES 1-6.

hours after release, but become completely non-motile within 24 hours. "Plates" of spermatids remain active for three days but start to disintegrate on the fourth day without completing metamorphosis. With these facts in mind, adult *Sp. morchi*, taken from the field and in most cases removed from their tubes, were isolated in either 10-ml. beakers or polyethylene ice-cube trays containing pasteurized sea water. Daily observations were made to determine the length of time elapsed after isolation for spawning to occur. Of 63 isolates, 26 spawned, and of these, 13 produced broods of normal larvae. The results are summarized in Table II.

It will be subsequently explained that *Sp. morchi* adults removed from their tubes are not always capable of transferring all of their spawned oocytes into the opercular ampulla, and therefore, a number, and often all, of the oocytes are spawned freely into the sea water. Development outside of the brood pouch is almost always suspended, presumably because of the growth of bacteria and protozoa which adhere to the sticky fertilization envelope. In most instances, the term "arrested" where it appears in the above table refers to such cases.

In the adult isolation experiments, newly spawned oocytes were removed intermittently from culture and examined with the compound microscope. Sperm penetration was never observed and only a partial sequence of events can be given. In the earliest stage observed, the vitelline envelope was just elevating at the animal pole as the fertilization envelope. The germinal vesicle at this time appears to be still intact and lies close to the surface under the plasma membrane of the animal pole. About $2\frac{1}{2}$ hours after elevation of the fertilization envelope, extrusion of the first polar body begins and is completed in about 30 minutes. The second polar body is given off about two hours after the first and with its extrusion, the first polar body divides. In live material, fine filaments can be seen extending from the surface of the polar bodies to the fertilization envelope where they apparently attach; the filaments have not been seen in fixed material. Because of the large amount of heavily pigmented yolk, male and female pronuclei have not been observed following polar body extrusion. The fertilization envelope remains closely

FIGURE 1. Section through a fertilized oocyte showing the lifting of the fertilization envelope (FE), and first polar body (1 PB). Epon; Richardson's stain. (300 ×)

FIGURE 2. Section through an oocyte spawned after 58 days in isolation, showing the first polar body (1 PB) and two chromosomes (Ch) arranged in second meiotic metaphase. Paraffin; haematoxylin-eosin. (750 ×)

FIGURE 3. An adjacent section to that in Figure 2 showing a sperm aster (SA). Paraffin; haematoxylin-eosin. (750 ×)

FIGURE 4. Cross-section through the calcified region of the operculum showing the outer cuticle (OC), outer epithelium (OE), inner epithelium (IE), calcified layer (CL), inner cuticle (IC), and blood capillary (BC). Epon; Richardson's stain. (1125 ×)

FIGURE 5. Cross-section through the operculum in the zone of transition (ZT) between the calcified region (CR), and the non-calcified region (NCR). Epon; Richardson's stain. (750 ×)

FIGURE 6. Cross-section through the non-calcified region of the operculum showing the outer cuticle (OC), outer epithelium (OE), inner epithelium (IE), and inner cuticle (IC). Epon; Richardson's stain. (1125 ×)

applied to the egg surface except at the animal pole where the perivitelline space is about 14 microns (Fig. 1). The above observations were made at a temperature of 12–13° C.

Five oocytes spawned after the parent had been in isolation for 58 days (animal #15, second spawning) were fixed and sectioned. Cytological examination revealed that the first polar body had been given off and the chromosomes were in second meiotic metaphase; a sperm aster was present in the cytoplasm (Figs. 2, 3). Although actual fusion of pronuclei has not been observed, it is, nevertheless, concluded that *Sp. morchi* is not only capable of reproduction in isolation but of self-fertilization. The fact that both polar bodies are given off in oocytes spawned in isolation supports this conclusion and makes it seem unlikely that parthenogenesis is occurring.

OBSERVATIONS ON OPERCULAR BROOD PROTECTION

Structure of the operculum in Sp. morchi

Second of the branchial tentacles in position, in relation to the dorsal midline, the operculum arises from the left branchial lobe (see zur Loye, 1908, Fig. 12). Typically, the operculum contains a spacious cavity, the ampulla, and is capped by a characteristically bilobed calcareous plate. Careful observation of a living operculum with a dissecting microscope reveals the presence of a distinct though small pore in the wall of the ampulla. It can be determined with direct lighting that, except for the ampullar side, the operculum is almost completely calcified. At the base of the operculum is a short stalk, the peduncle, which often has a groove running around its circumference. Below the groove, when present, the peduncle is swollen. This swelling is the replacement operculum and will become a functional operculum after the extant operculum is shed. The size of the operculum in mature adults can be quite variable but averages about 700×330 microns at its longest and widest dimensions. No attempt has been made to correlate opercular size with, for example, age classes, but it is possible that such a correlation may exist.

From the outside in, the operculum consists of an outer cuticle and epithelial layer followed by an inner epithelial layer and cuticle. There is a distinct space separating the two epithelial layers; however, a basement lamina has not been demonstrated. The histology of the calcified region of the operculum is somewhat different from that of the non-calcified portion.

In the calcified region (Fig. 4), the outer cuticle is about 2–2.2 microns thick and has a striated appearance. The outer surface of the cuticle is fuzzy, suggesting the presence of minute projections. The cuticle is strongly PAS-positive, indicating the presence of a polysaccharide, and stains blue with Heidenhain's Azan. The outer epithelial layer varies in height from 3 to 6 microns and is cuboidal. Protoplasmic processes extend from the apices of the cuboidal cells into the outer cuticle and give the latter its striated appearance. Nuclei are basal in position, and contain a single nucleolus and irregular pieces of chromatin. Several vacuoles are located within each cell of the outer layer but their significance is not known. Separating the outer epithelial layer from the inner epithelial layer is a space of about 3 microns which widens to about 6–7 microns at the sites of blood capillaries which penetrate between the two tissue layers. The inner epithelial layer is about

half as thick as the outer and contains few vacuoles. Nuclei of the inner layer are more elliptical in shape than those of the outer layer. The inner cuticle is about 1 micron thick and very dense; separating it from the inner epithelium is a space which varies from 6 microns along the sides of the operculum to about 28 microns at its apex. Coarse protoplasmic processes extend from the inner epithelium across the space and attach to the cuticle. Sections of opercular brood pouches fixed in

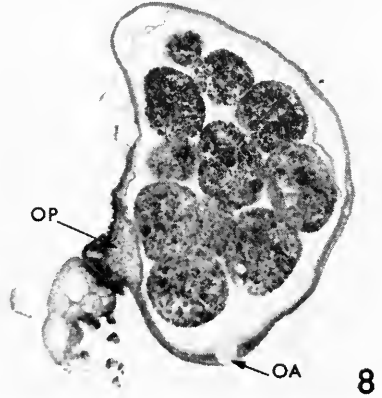


FIGURE 7. Cross-section through the opercular aperture (OA). Note that the epithelium surrounding the aperture is columnar. Paraffin; iron haematoxylin. (400 \times)

FIGURE 8. Sagittal section through an operculum containing a brood. Note the opercular aperture (OA), and opercular peduncle (OP). Paraffin; iron haematoxylin. (75 \times)

FIGURE 9. Sagittal section through an operculum containing a moribund brood. Note the developing replacement operculum (RO). Paraffin; haematoxylin-eosin. (75 \times)

FIGURE 10. Cross-section through the branchial crown showing that the opercular aperture opens to the center of the crown. Paraffin; haematoxylin-eosin. (75 \times)

neutral formalin and treated with the chloranilic acid method reveal that the space between the inner epithelium and cuticle is the site of calcification. The surface of the inner epithelial layer, facing the cuticle, gives a very strong reaction and presumably it is this layer that is responsible for calcium secretion.

There is a sharp zone of transition between the calcified and non-calcified regions of the operculum (Fig. 5). In the non-calcified portion, the outer cuticle and epithelium are similar to those of the calcified region (Fig. 6). The space separating the outer epithelium from the inner epithelium averages 1-3 microns and there are fewer blood capillaries than are found in the calcified region. The inner epithelium varies from about 7 to 9 microns in thickness and is also cuboidal in appearance. Nuclei are tall, having a height almost equivalent to that of the entire cell. One or more large intracellular vacuoles are present, but often disappear when the operculum contains a brood. In both epithelial layers of the calcified and uncalcified regions alike, the cells contain within their cytoplasm several dark-staining spherical bodies, each of which encloses a less dense internum. The dark-staining bodies are removed in the solvents used in routine paraffin embedding which suggests that they are lipid in composition. The inner cuticle of the uncalcified region is like the outer except the surface is less fuzzy and there is a thin membrane-like layer applied to it.

Around the opercular aperture the epithelial layers become considerably thickened (Fig. 7). The cuboidal nature of both outer and inner epithelium changes to columnar. The height of the outer cells averages about 13 microns and that of the inner about 22 microns. Nuclei are elongated in shape and the cytoplasm is almost completely devoid of vacuoles. The interspace between outer and inner epithelial layers is obliterated. Muscle and nerve fibers are not demonstrable.

The outer cuticle and epithelium of the operculum are continuous with those of the peduncle (Fig. 8). The inner cuticle and epithelium forms the floor of the operculum and is not continuous with any portion of the peduncle; consequently, the only opening into the opercular brood pouch is by way of the opercular pore or aperture. There is a basement lamina below the columnar epithelium of the peduncle and also below the inner epithelium of the operculum where the latter forms the roof of the peduncle. Composing the core of the peduncle are longitudinal muscle fibers and blood capillaries embedded in an ill-defined connective tissue containing fibers which stain blue with Azan. According to Hanson (1949) in the peduncle and branchial tentacles of *Sp. corrugatus* and *Sp. militaris* three nerves can be demonstrated using the Azan method; however, in the present material, using the same technique, only one nerve (the internal branchial nerve, using Hanson's terminology) could be demonstrated. At the top of the peduncle, or base of the operculum, muscle fibers can be observed to penetrate between the inner and outer epithelial layers of the operculum, but they can not be followed for any distance. As mentioned earlier, there is often a groove running around the circumference of the peduncle. This may properly be referred to as the zone of detachment because it is here that the operculum becomes disconnected from the peduncle when it is periodically shed. A new or replacement operculum makes its appearance below the old operculum and is generally well-formed before the latter is shed (Fig. 9). Experimental extirpation of the operculum results in its renewal in the same position.

Brooding behavior in Sp. morchi

The primary or first operculum formed at metamorphosis of the larva does not serve as a brood pouch but is replaced by a secondary operculum which does serve in that capacity. This conclusion is based on observations made on animals raised from isolated larvae. It cannot be stated exactly how many consecutive broods an operculum can accommodate before it is replaced, but it is certainly at least three. As pointed out by Gravier (1923), opercular replacement is not peculiar to operculum brooders but also occurs periodically in species which brood in the tube. Based on counts made on 115 broods, the average number of embryos per brood in *Sp. morchi* is 15; however, within a population, a range of from 3 to 45 embryos per brood is not unusual. The embryos are not surrounded by a common membrane within the operculum as described in *Sp. granulatus* by Bergan (1953a).

On three different occasions larval release was observed in the laboratory. Larvae emerge *via* the opercular pore which opens wide enough to allow the simultaneous release of two larvae at a time. After release of the larvae, the pore appeared to close somewhat but it could not be ascertained whether or not this closure was complete. In all three cases, within 12 hours of larval release a new brood had been deposited in the operculum and the opercular pore was tightly closed. From these observations, it is not possible to determine whether the pore opens actively during larval release or merely passively as the result of force applied by

TABLE III

Number of eggs spawned free and/or successfully transferred to the brood pouch in Spirorbis morchi adults removed from their tubes

Animal	Numbers of eggs spawned	Numbers of eggs placed in brood pouch	Numbers of eggs spawned free into the sea water
1	8	0	8
2	12	3	9
3	37	18	19
4	13	2	11
5	21	0	21
6	5	5	0
7	16	7	9
8	15	0	15
9	10	0	10
	11	0	11
10	5	0	5
	8	0	8
11	4	0	4
12	7	0	7
	8	6	2
13	13	0	13
14	13	0	13
	3	0	3
15	8	0	8
16	6	6	0
17	7	4	3
18	9	1	8

the moving larvae; however, it is certain that the pore closes after receiving a brood and this would seem to suggest an active process. It should be pointed out that in a virgin operculum the pore is tightly closed prior to deposition of the first brood and this would suggest that opening, like closing, of the pore is under active control.

Earlier it was mentioned that adults removed from their tubes are inefficient in transferring spawned eggs into the opercular brood pouch. In most instances, counts of the number of eggs spawned free and the number placed in the opercular brood pouch were made. These data are tabulated in Table III.

In all cases, the oocytes spawned free into the sea water were activated and commenced development. Failure to transfer spawned oocytes into the brood pouch has never been observed in control animals left in their tubes.

Although egg transfer has never been observed, a reasonable hypothesis can be constructed based upon the aforementioned observations. First, it has been established that the only entrance into the brood chamber is *via* the opercular pore; consequently, the oocytes must be spawned to the outside and then transferred into the operculum. The opercular pore, in some unexplained way, is capable of opening and closing. Activation of the oocyte apparently occurs outside of the operculum. An animal removed from its tube is inefficient in transferring its eggs into the operculum; consequently, the confines of the tube are necessary. It seems reasonable to suppose that gametes are released *via* rupture of the body wall (Potswald, 1967) into the space formed by the ventral fecal pellet groove and wall of the tube. The oocytes are here fertilized and are swept, by means of ciliary action, across the achaetous zone, forward along the dorsal surface of the thorax, and finally into the center of the branchial crown which is completely withdrawn into the tube at spawning. Having reached the branchial crown, the fertilized eggs pass into the operculum by means of the opercular pore which opens to the center of the crown (Fig. 10).

Inefficiency in egg transfer in the case of adults removed from their tubes is not total. As can be seen from an examination of the data, a number of animals were successful in transferring a portion of their brood and, in fact, two were 100% effective in transferring relatively small broods. *Spirorbis morchi* has a well developed thoracic cloak which extends posteriorly and attaches ventrolaterally on about the sixth abdominal segment. The under surface of the cloak is ciliated and could conceivably simulate, to some degree, the confines of the missing tube. The latter explanation would hold true especially for animals which assume an abnormal position, *i.e.*, ventral surface of the thorax facing the substratum, in which case the cloak would serve as a natural trough into which the spawned oocytes could fall. It is common for animals removed from their tubes to assume such a position.

A major problem yet to be solved involves the controlling factors operating in opening and closing of the opercular pore. Perhaps control is endocrine in nature and correlated with spawning. At the level of light microscopy, histological investigation failed to reveal muscle fibers in the area of the pore. In an attempt to determine the possible cytological basis of contraction, a study of the opercular epithelium, at the ultrastructural level, is presently being undertaken.

DISCUSSION

Functional hermaphroditism is found in nearly all phyla of invertebrates (Coe, 1943). Generally, when eggs and sperm develop concurrently in the same animal there are mechanisms involved which insure that self-fertilization does not take place. Nevertheless, there are documented cases of self-fertilization in various invertebrate groups. Smith (1950) has demonstrated, quite conclusively, self-fertilization in *Neanthes lighti*, an hermaphroditic, viviparous nereid which occurs along the California coast, from estuaries and sloughs of higher than normal salinity to rivers where salinity may be less than 2% that of sea water. Within the phylum Mollusca there are many instances where self-fertilization has been demonstrated, especially among the pulmonate gastropods where individuals may reproduce by self-fertilization generation after generation when experimentally isolated (see Coe, 1943, 1944 for review). Nyholm (1951) has shown that in *Labidoplax buskii* (Echinodermata, Holothuroidea) artificial self-fertilization can occur on a limited scale during the height of the breeding season. Finally, in the Arthropoda there is evidence that self-fertilization is possible in some barnacles (Barnes and Crisp, 1956; Barnes and Barnes, 1958).

In the examples cited it is obvious that eggs and sperm from the same individual are physiologically compatible; however, in most of the cases mentioned, cross-fertilization is probably the normal mode of reproduction, or at least it occurs much of the time. Whenever selfing is occasional or periodic, it need not cause any essential change in the evolutionary pattern of the species; the gene pool continues to be a reality. If, however, self-fertilization becomes obligatory rather than facultative, pure lines or clones are produced which do not exchange genes; the genotype of each line becomes a closed system capable of changing only through mutation or through reversal to biparental sexuality (Dobzhansky, 1951). In the latter case, the species, by virtue of selfing, robs itself of much evolutionary plasticity and actually gives up the benefits of sexual reproduction.

The observations presented in this report on the biology of fertilization in *Sp. morchi* are, to some extent, comparable to the observations of Gee and Williams (1965) on *Sp. borealis* and *Sp. pagenstecheri*. Although Gee and Williams did not rear animals from settlement to maturity in isolation, they did obtain data from isolated adults indicating that *Sp. borealis* and *Sp. pagenstecheri* are capable of self-fertilization. Gee and Williams make the assumption that cross-fertilization occurs when a number of worms are placed together and point out that in both species the viability of the progeny produced in association is noticeably greater in comparison with the progeny produced in isolation. This decrease in viability may be due to semi-lethal recessive genes. An apparent decrease in viability is also observed when *Sp. morchi* is allowed to reproduce in isolation.

Dasgupta and Austin (1960) report that they have evidence which suggests that the hermaphrodite serpulids *Spirorbis* and *Filograna* have been derived from triploids with a loss of a single chromosome. According to their study, *Serpula crater* has a diploid number of 14, and considering the "ancestral serpulid" as being $2n = 14$ then *Spirorbis* and *Filograna* are $2n + n - 1 = 20$. Dales (1963), presumably on the basis of the latter work, has made the interesting suggestion

that self-fertilization may have arisen because of the difficulty of chromosome pairing. Studies such as Dasgupta and Austin's which suggest the evolution of a group through polyploidy are often based, as is theirs, on chromosome numbers alone without study of chromosome morphology, and, as White (1954) points out, hermaphroditic forms capable of self-fertilization do not seem to show any more polyploidy than those in which cross-fertilization is obligatory. Although, at present, there is no concrete evidence that *Spirorbis* normally cross-fertilizes, there are cogent arguments in favor of this view. One of the best arguments, aside from the genetic implications discussed above, is that offered by Gee and Williams (1965) who argue that the gregarious settling behavior exhibited by *Spirorbis* larvae (Knight-Jones, 1951) would seem to allow cross-fertilization in a population. Conceivably, an animal, just prior to spawning, could draw into its tube, by means of the respiratory current, sperm from an adjacent animal which has already spawned. It is concluded that *Spirorbis* is capable of self-fertilization and this may be of some advantage to a sedentary hermaphroditic species which has a non-planktotrophic dispersal phase; however, cross-fertilization is probably the common mode of reproduction in a natural population.

Elsler (1907) described opercular brood protection in *Sp. corrugatus* Montagu and *Sp. pusillus* de Saint Joseph as taking place between the calcareous plate and the opercular epithelium. Borg (1917) found that brood protection in *Sp. pagenstecheri* occurs in the same fashion as described by Elsler. Apparently, the structure of the operculum in the last-mentioned species must be quite different from that in *Sp. morchi* and is deserving of further study. Bergan (1953a) describes brood protection in *Sp. granulatus* as occurring in the opercular ampulla as it does in *Sp. morchi*. Bergan also observed the presence of the opercular pore; however, he was of the opinion that it was too small to allow the passage of eggs and did not attempt to determine its function. In the present study on *Sp. morchi*, it has been observed that the opercular pore is capable of opening and closing, and, in fact, is the only entrance into the opercular ampulla. The mechanisms by which the pore operates have yet to be elucidated.

Elsler (1907), Borg (1917), and Gravier (1923) all speculated that opercular brood protection is a recent development as compared with brood protection in the tube. According to Borg, brood protection in the operculum has been found only in species having three thoracic segments and never in species having $3\frac{1}{2}$ or four thoracic segments. This would fall in line with the idea proposed by Caullery and Mesnil (1897) that the genus *Spirorbis* has been evolved from other Serpulidae by a gradual incorporation of thoracic segments into the achaetous region. Although a very interesting hypothesis, and one worthy of consideration, it would be of interest to know what advantage, if any, opercular brood protection has over brood protection in the tube. Perhaps a comparative physiological study of embryos developing under the two different modes of brood protection might shed some light on this problem.

I wish to thank Dr. Robert L. Fernald, Director of the Friday Harbor Laboratories, for his persistent encouragement and help throughout the course of this study. Dr. W. Siang Hsu and Dr. Paul L. Ilg are thanked for their many helpful suggestions.

SUMMARY

1. Data obtained from *Spirorbis morchi* larvae reared from settlement to maturity in isolation, together with observations on isolated adults, provide evidence that *Sp. morchi* is capable of self-fertilization. Self-fertilization in *Spirorbis* is believed to be facultative and not obligatory.

2. The histology of the opercular brood pouch in *Sp. morchi* is described, and it is concluded that the only opening into the brood pouch is by means of an opercular pore.

3. Observations on the brooding behavior of *Sp. morchi* have led to the conclusion that spawning takes place when the animal is completely withdrawn into its tube.

4. An explanation as to how oocytes are deposited into the opercular brood pouch is advanced.

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TEMPERATURE RELATIONSHIPS OF POIKILOTHERMS AND THE MELTING TEMPERATURE OF MOLECULAR COLLAGEN

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Biochemical and biophysical data are beginning to accumulate for various proteins of a wide range of animals. These data include the amino acid analyses, molecular weights and dimensions of the molecular units, as well as their thermal and chemical properties. Of these, the protein of connective tissue, collagen, is probably the most widely studied from a comparative point of view (Harkness, 1961; Gross, 1963). It is the main fibrous component of the skins and tendons of the vertebrates and occurs in animals of most of the other phyla. It can be prepared in a fairly pure form from the tissues of most animals and is identified by its characteristic high angle x-ray diagram and its distinctive amino acid composition. In the native state (*e.g.*, tendon, skin, cuticle) or in its molecular state¹ in dilute solution, it exhibits fairly sharp, reproducible melting points. The melting of collagen is predominantly a first order phase transition (Garrett and Flory, 1956), and as such is virtually independent of time and takes place over a small temperature range.

These melting points, *i.e.*, the bulk melting temperature T_s and the molecular melting temperature T_D , are known to correlate with the upper limit of the environmental temperature (where these are available) of the animal concerned (Gustavson, 1956; Leach, 1957; Rigby, 1967a). Thus, all mammals with body temperatures similar to man have the same molecular melting temperature in water, $\sim 36^\circ$ C. (body temperature 37° C.) while the collagen of cod (a cold-water fish) melts at 15° and that of tuna fish (warm water) melts at 27° C.

For many collagens, T_s and T_D also correlate with the sum of two of the amino acid residues present in the molecule—proline and hydroxyproline (see Harrington and von Hippel, 1961). However it has recently been found (Rigby, 1967a; Fujimoto and Adams, 1964) that some earthworms and the parasitic worm *Ascaris* each contain two distinct collagens with respect to amino acid composition, particularly the sum of proline and hydroxyproline, yet T_s and T_D for each pair is identical (Rigby, 1967a).

It is perhaps advisable at this stage to describe one of these collagens as "collagenous" since it differs in some respects from the generally accepted definition of a collagen. Thus the cuticle of these worms (the collagen in question) while it shows the typical high angle x-ray diagram, and has an amino acid

¹ The molecular unit in vertebrate collagen is composed of three polypeptide chains, each in the form of a left-handed helix wound into a super helix about a common axis. The dimensions of this unit are *ca.* $2800 \text{ \AA} \times 14 \text{ \AA}$, and the molecular weight is *ca.* 300,000 (see Harrington and von Hippel, 1961). The dimensions and molecular weights of invertebrate collagens appear to be different and not yet fully characterized (Maser and Rice, 1962; Josse and Harrington, 1964).

composition similar in pattern to vertebrate collagen, is secreted by epidermal cells rather than by fibroblasts, and does not exhibit the 600–700 Å axial repeat in the electron microscope. However from the point of view of this paper it is not crucial whether the cuticular protein is a true collagen or not, since, as we have already mentioned, it has the same values for T_s and T_D as the body collagen of the same worm (which collagen has all the accepted characteristics of collagen).

In this paper we should like to point out correlations between the melting properties of the collagens, and the behavior and physiological properties of a number of animals, particularly earthworms and parasitic worms. It is not meant to suggest that there is any causal relation between the melting of collagen and sudden changes in physiological behavior (although this is possible), but to emphasize the fact that the thermal properties of at least two proteins allow one to predict some of the temperature relations of the animal.

MATERIALS AND RESULTS

Bulk collagen was prepared from the following worms: *Ascaris lumbricoides* and *Macracanthorhynchus hirudinaceus* which are parasitic worms from the small intestine of the hog, and three earthworms, *Allolobophora caliginosa*, *Digaster longmani* and *Pheretima megascolidioides*. *Digaster* is a giant earthworm found in the Kyogle district of New South Wales and attains a length of 5 feet or more. *Pheretima* is a common Japanese worm and was kindly supplied by Dr. K. Inukai, of the Department of Chemistry, Kyoto University, Japan.

All these worms with the exception of *Macracanthorhynchus* have an outer covering, the cuticle, which is almost pure collagen (with the reservation as to definition noted above). This was used as one sample. After removal of the cuticle, collagen can be prepared from the body wall of the worms. This was the second source of collagen from each worm. In the case of *Macracanthorhynchus* the body wall could easily be separated into two layers after soaking strips of wall in water overnight. The inner layer is collagen (high angle x-ray, 600–700 Å repeat and amino acid composition) (Rigby, 1967a). The method of purification of each tissue consisted in successive extraction for 24 hours in 0.1% trypsin, 10% NaCl and saturated NaH_2PO_4 solution, then washing in 0.9% saline. Full details of the preparative procedures, purification, identification and chemical composition of these samples have been given elsewhere (Rigby, 1967a).

The thermal stability of each collagen was determined by measuring the melting point of the bulk material in physiological saline, and the melting point of the bulk material in hydrochloric acid at pH 1. The melting point in saline, denoted by T_s , is commonly referred to as the shrinkage temperature. The melting point in HCl at pH 1 has been shown (Rigby, 1961, 1967b) to agree with the temperature of the helix coil transition of molecular collagen in dilute aqueous solution at neutral pH. This temperature is denoted by T_D . The principle of our method is the detection of the sudden increase in force when it melts, in a sample which is prevented from changing its dimensions. In saline it is crystalline aggregates of molecular units which melt, while in acid solution, swelling of the structure and subsequent dispersion of molecular units allows single molecules or small groups to melt.

The results relevant to this paper are given in Table I. The temperatures are the means of at least four determinations; the error is $\pm 1^\circ$ C.

At this point it is worth noting that for a number of soft collagenous tissues T_s and T_D are independent of the degree of purification. Thus, in the present case, strips of fresh earthworm body wall (*i.e.*, with cuticle removed) give the same values for T_s and T_D as does the collagen extracted from same by the methods described above. While it does not follow that only the collagen is melting it indicates that in soft tissue, at least, some of the physical properties of collagen are unaffected by its association with other components. This point is of particular physiological interest as the following discussion attempts to show.

The question as to the correlation of T_s and T_D with amino acid composition—in particular, with the total pyrrolidine content—is not considered to be relevant to this discussion, but it has been dealt with fully elsewhere (Rigby, 1967a).

TABLE I

Melting temperatures in physiological saline ($^\circ$ C.) of the collagen of various worms. T_s is the melting point of bulk, native sample, T_D the melting point of molecular unit. B, body. C, cuticle. These transitions were measured on the bulk native material (Rigby, 1961, 1967b)

	<i>P. megascolidioides</i>	<i>Digaster longmani</i>		<i>A. caliginosa</i>		<i>As. lumbricoides</i>		<i>Muraecanthorynchus</i>
	C	C	B	C	B	C	B	B
T_s	37	34	40	34	40	56	59	62
T_D	22	22	22	22	22	40	38	42

DISCUSSION

The first thing to be noticed from Table I is that all the earthworms have the same value of T_D for both cuticle and body collagen. Also, the hog parasitic worms and hog intestinal wall collagens all have similar values of T_D . The values of T_s tend to be the same for each group as well, although a number of factors affect T_s without affecting T_D . In any case it is of more fundamental interest to consider T_D . It will be seen that T_D for the parasitic worms is close to that of their environmental temperature, namely that of the hog. This does not appear to be a coincidence, for it is well known that the T_D values of the collagens of the mammals, man, rat, sheep, and ox, are all virtually the same as that of their deep body temperature, as was mentioned in the introduction. This information is summarized in Figure 1 for a wide range of animals and environments. The simplest explanation of this correlation is of course, natural selection, although why the two temperatures should almost coincide is a question which requires further study. It might be thought, for example, that T_D would be well above the maximum environmental temperature.

The second point of interest in Table I is the value of 22° C. for T_D of all the earthworm collagens. According to Figure 1 this would imply an upper limit of about 22° C. for the environmental temperature of these worms in their natural habitat. There are two kinds of evidence in the literature which are

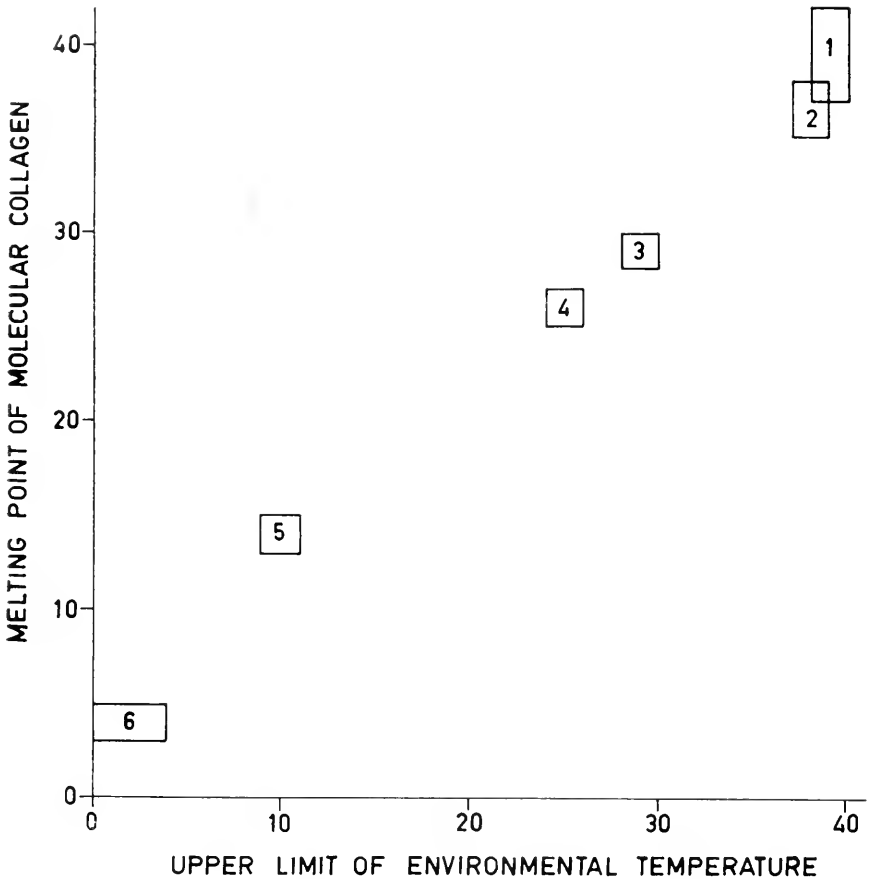


FIGURE 1. The relation between the melting point (T_D), of the collagen molecule from various animals, and the approximate temperature of the upper limit of their environment. 1. *Macracanthorhynchus*, *Ascaris* and hog; 2. rat, human, cow; 3. snail (Newell, 1966) (*Helix aspersa*); 4. tuna skin (Dakin, 1953); 5. cod skin (Tait, 1952); 6. Antarctic "ice-fish." The reference numbers are for environmental temperatures. The Antarctic "ice-fish" is the subject of a detailed paper in preparation.

consistent with this view. These are (i) the thermal preference range of the worm and (ii) the relation between the environmental temperature and the body temperature of the worm. We have only been able to examine collagens of two of the worms studied by others: viz: *Pheretima megascalidioides* and *Allolobophora caliginosa*. In both cases we must assume that temperature was the only factor under observation and secondly, point out that our *Allolobophora* were of Australian origin while the published work to be discussed was performed with North American and Egyptian animals.

We turn first to studies of temperature preference ranges and consider *A. caliginosa* particularly, since we have data for the collagen of this animal. Grant (1955) and El-Duweini and Ghabbour (1965) have studied the thermal

preference ranges of this worm, as well as *E. foetida* and *P. hupciensis* and *P. californica* and *Alma* sp., respectively. Although the lower limit of this range varies, Grant (1955) quotes 23° C. as the upper limit of each animal. The results of El-Duweini and Ghabbour (1965) are more difficult to interpret since they used *A. caliginosa* from two regions of Egypt. Both groups of worms showed a wide temperature preference range but in both cases there was a sharp drop in the number of worms preferring temperatures above 28° C., this number was about 10% of the total. For *Alma* sp. 88% of the population preferred the range 23–28° C. with the peak at 25° C. *Ph. californica* were almost equally distributed between 26° C. and 35° C. Grant (1955) allowed 24 hours before deciding on a worm's temperature preference, whereas El-Duweini and Ghabbour (1965) allowed 0.5 hour and it could be questioned whether theirs are equilibrium results. However for *A. caliginosa* a T_D value of 22° C. appears to be correlated with the temperature of the upper limit.

Considering next body temperature, we draw attention to the work of Kim (1930) who studied *Pheretima megascolidioides*. His main conclusion was that between 6° and 23° C. the body temperature of *Pheretima* was the same as that of the surrounding medium, but beyond 24° C., for unknown reasons, its body temperature varied erratically. His results imply an upper limit of temperature preference of 23–24° C., and this again (as with *A. caliginosa*) correlates with T_D .

Further, on the subject of sudden physiological changes due to temperature increase such as the one just described, it is relevant to mention the results of Hatai (1922) concerning muscular contraction in several Japanese earthworms. He found rhythmic contraction to be normal and unaffected by temperature up to 23° C., beyond which the form of contraction altered suddenly.

The preceding results and discussion show that earthworms from a number of species and habitats have collagens with identical thermal stability, and, in turn, similar upper limits for their temperature preference range. This value, 22° C., may be characteristic of the family.

It is now of interest to examine the so-called upper lethal temperature. The definition of this temperature is arbitrary and values quoted for a given animal are thus widely spread. Assuming as before that the upper lethal temperature depends only upon temperature and that other factors, such as water content, are kept constant it would seem that the upper lethal temperature could be defined by extrapolation of a time-temperature relation from high temperatures until time and temperature become independent. In other words, above the true lethal temperature, time and temperature would be approximately inversely proportional; below, no relation would be evident.

In the literature on lethal points, El-Duweini and Ghabbour (1965) quote 39° C. for *A. caliginosa* with similar figures for *Ph. californica* and *Alma* sp. Their definition was the temperature at which more than half the sample dies after an exposure of 0.5 hour. Wolf (1938) and Hogben and Kirk (1944), using *L. terrestris* and times of 400 minutes and 12 hours, respectively, gave lethal temperatures of about 29° C. Grant (1955) using an interpolation method and exposure times up to 48 hours gives values of 26° C., 25° C. and 23° C. for *A. caliginosa*, *P. hupciensis* and *E. foetida*, respectively.

It would appear that the high values quoted by El-Duweini and Ghabbour

(1965) are due to the short exposure time and the inverse relation between time and temperature discussed above, although a factor could be that the worms have some measure of acclimatization to the hot Egyptian climate as mentioned by these authors. Of greater significance, however, is the fact shown by Table I that the thermal shrinkage of earthworm bulk collagen takes place between 35 and 40° C. The values of Grant (1955) are quite close to the upper limit of the preferred temperature range and this is consistent with the data of Kim (1930) where it was seen that above this upper limit definite physiological irregularities become apparent. In other words irreversible changes have begun to occur in the animals' tissues, but long times may still be required before death occurs. One further point which should be mentioned here is that whereas any molecular melting which takes place in bulk collagen is reversible as long as there is no mechanical stress present and T_D is not exceeded by 5–10° C., the melting which occurs at T_S is irreversible except, possibly, in the case of cross-linked collagens such as elastoidin. Thus, so far as collagen is concerned the temperature could exceed T_D for short periods of time without necessarily affecting the animal adversely.

Summarizing, we can say that for the earthworms discussed, the melting point of the collagen molecule (the basic unit of their connective tissue) is very close to the value of the upper limit of their preferred temperature range and in turn to the upper lethal temperature. They are similar to mammals in this respect and to parasitic worms of mammals which at some stage of their life cycle exist at much lower temperatures. Whether or not collagen is uniquely involved in determining these limits cannot be said at this stage, although the fact that it is a protein with a very low rate of turnover (with certain exceptions, Harkness, 1961) and sharp melting point is perhaps significant. It should be pointed out, however, that it is likely that all the animal proteins begin to denature at temperatures above T_D . Finally it has recently been shown by Newell (1966) that the oxygen uptake of various poikilotherms increases abruptly at temperatures which correspond with the upper limit of the environmental temperature, and in one case for which collagen has been isolated (*Hclir aspersa*), to the T_D for this collagen, *viz.* 27° C. (Rigby 1967b).

SUMMARY

The molecular unit of collagen, the main fibrous protein of connective tissue, undergoes a first order phase transition at a characteristic temperature, when heated in physiological saline. For the collagen of mammals this temperature is known to almost coincide with the deep body temperature of the animal, and for a number of poikilotherms, with the upper limit of their environmental temperature. We have measured the transition temperature of the collagens of a number of worms and now report that:

1. The molecular unit of the collagen of two hog intestinal worms undergoes this transition at temperatures close to that of the body of their host.
2. For a number of earthworms for which there are precise data on thermal preference limits, the molecular unit of the collagen of these animals has a transition temperature which is very close to the upper limit of the thermal preference zone.

3. There is no evidence of any causal relation between the transition temperature of molecular collagen, and the abrupt physiological and behavioral changes which become apparent at the same temperature. However, the results are direct evidence of a specific protein having easily measurable thermal properties which reflect thermal properties of the whole animal.

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MIGRATORY ORIENTATION OF AMBYSTOMA MACULATUM:
MOVEMENTS NEAR BREEDING PONDS AND
DISPLACEMENTS OF MIGRATING
INDIVIDUALS

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Migrating spotted salamanders, *Ambystoma maculatum*, tend to enter and leave breeding ponds near the same point (Shoop, 1965). This paper confirms the original observation and presents results of observations and experiments designed to further elucidate the problems of migratory orientation of spotted salamanders.

METHODS

Animals from two breeding ponds (A and B) in Wellesley, Massachusetts were studied from 1964 to 1967. Nylon-coated, screen fences 40 cm. high, buried 15 cm. in the soil, surrounded the ponds; deep can traps were buried at intervals along both sides of the fences. Other fences from 10 to 200 m. long with traps along both sides were located at varying distances (15 to 175 m.) from the fence surrounding Pond A. Details concerning Pond A and methods are described by Shoop (1965); Pond B is in mixed hardwood-pine forest adjacent to a golf course.

Traps were checked regularly throughout the day and night during the breeding season and at dawn during the rest of the year except when snow covered the traps. The trapping method exposed the animals to increased predation by raccoons; several salamanders were killed before a wire with electrical charger was added around the outer face of the fencing about 15 cm. above the ground. Some animals were released on the Wellesley College campus in a 40-m.-diameter arena with traps along the inner face of a 0.5-m.-high surrounding fence. All animals were individually marked by toe-clipping. Statistical procedures follow Batschelet (1965).

CONFIRMATION OF EARLIER OBSERVATIONS

The relation between point of entry and exit of Pond A by animals during 1964, 1965, and 1966 and mean vector values, 99% confidence intervals for mean vectors, and angular deviations are presented in Figure 1. In all cases corrections for trap intervals are included in the vector determination. The r values vary from 0.42 to 0.55, indicating some dispersion but non-random distribution. In all three cases the mean exit vector falls within the consistent direction (Shoop, 1965) in relation to the point of entry into the pond, and the 99% confidence intervals for the mean vectors overlap. The data for the three years are not

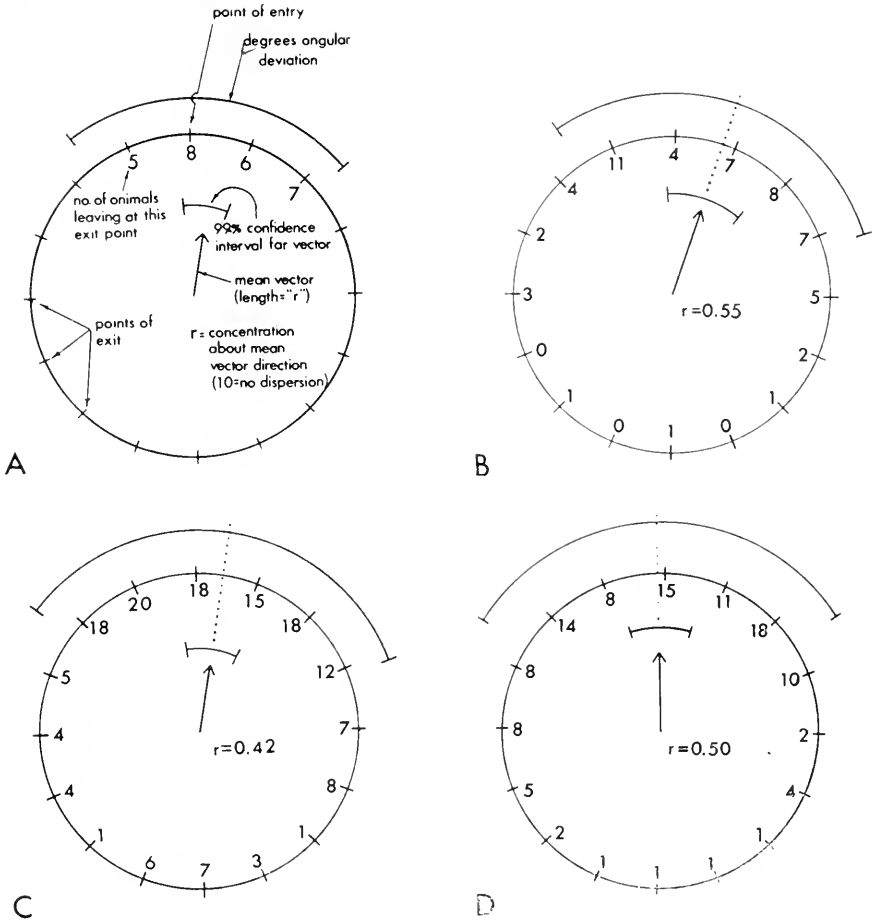


FIGURE 1. Relation of exit point at breeding pond border to point of entrance (top of circle is point of entry, not necessarily north): A, explanation; B, 1964; C, 1965; D, 1966. The 99% confidence intervals of the mean vectors overlap indicating highly significant correlation between years.

significantly different. Animals collected only at entry or exit, or animals used for displacement studies in 1966 were not included in the vector determination.

When interpreting the data of movements into and out of the breeding pond, certain aspects of the experimental design should be considered. The pond was circular with a steep bank, a rather uncommon type of vernal pond. This allowed for close placement of the surrounding fence with traps at the sixteen compass points, and since all recorded directions are in relation to the center of the pond, the data may be more refined than is biologically significant. In reality, a migrating salamander may not move toward the center of the pond but merely attempt to intercept the border. An animal moving from a point east (90°) of the pond center may intercept the border anywhere from 0° to 180° and therefore have a

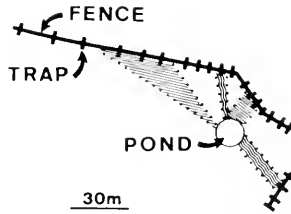


FIGURE 2. Corridors of travel of 4 animals (1967) selected to avoid overlap and show range of variability ($N=36$). One-half distance to next trap included in corridors. The fences and traps border forest areas.

“successful” migration. Conversely, the animals may leave the pond anywhere between 0° and 180° and be on the east side of the pond. Considering the amount of dispersion, this type of orientation may actually be more typical of the biologic situation. That a statistically significant number of animals did choose consistent headings in relation to the point they entered the pond indicates that the animals tend to use the same track even though their terrestrial retreats may not be closest to the entry point. Additionally, animals determined as taking “poor” headings may correct for their “error” after leaving the pond.

Studies utilizing fences at varying distances from breeding ponds in 1966 and 1967 seem to confirm the above suppositions. When migrating in a meadow to and from the pond many animals tend to limit their movements when within 100 m. of the pond to a narrow corridor 10 to 30 m. wide. Some corridors used by the salamanders in 1967 are represented in Figure 2. In 1967 36 animals were caught at the forest border and pond border at entry and exit (4 captures each). The corridor used in crossing the meadow to and from the pond may have little relation to the movement of an animal in the forest. Figure 3 shows capture points of an animal moving in an easterly-westerly corridor at a great angle to a line drawn between capture points at the forest border and in the forest. Time between the last two captures was 6 days.

Of the animals collected in successive years, most seem to travel in the same corridor to and from the pond (Fig. 4), although some apparently use different corridors. A pattern of different corridors for entry and exit (seen in 3 animals) may be repeated in successive years (Fig. 5A). A few animals seem to move in no definite pattern (Fig. 5B).

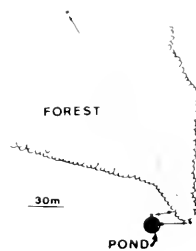


FIGURE 3. Movements of a single individual, 1967. ● = capture point; → = direction of movement; . . . = shortest distance between capture points.

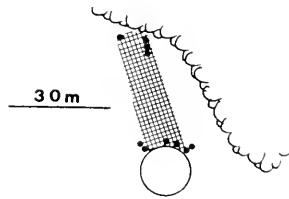


FIGURE 4. Capture points of a single individual, 1965-67. Dots around pond represent captures on entry and exit, 1965-67. Dots at forest border, entry and exit points 1966-67. Shaded area is probable corridor of travel.

EXPERIMENTS INVOLVING DISPLACEMENT OF MIGRATING SALAMANDERS

Displacements in the field

Since migrating animals tend to use the same track into and out of a breeding pond, they may possess an ability to utilize a directional tendency (Griffin, 1952, "Type II orientation," or Schmidt-Koenig, 1965, "compass orientation"); if familiar landmarks or local sensory cues are unavailable or not utilized, they may move in a prescribed direction or its reciprocal (along a directional axis). Assuming that only a compass sense is involved, the animals should fail to reach the pond in the following situation: animals are captured at the pond border, carried 90° around the pond, and then taken in a straight line away from the pond and released at a location where familiar landmarks are obscured. For example, if an animal that had migrated from a terrestrial retreat east of the pond to the east border of the pond, was moved to a point north of the pond, it would travel east or west and miss the pond. The following experiments tested this possibility.

Twelve animals captured at night in the ENE trap at Pond A border were placed in opaque containers and transported at dawn to four locations. Three animals were released 20 m. from ENE; three at 135 m. from ENE; three at 45 m. from NNW; and three at 175 m. from NNW. Of six animals collected in trap NNW, three were released 20 m. from ENE and three 45 m. from NNW. Results of this experiment are presented in Table I. One animal released 135 m. from ENE was killed by raccoons on return to the pond fence.

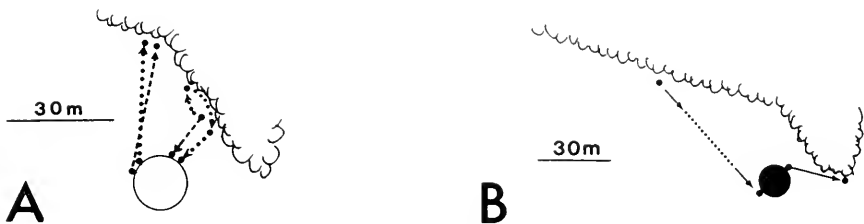


FIGURE 5. A, Movements of a single individual, 1966 (dashed line), 1967 (dotted line). Looping movements (1967) due to dry weather with animal returning to forest. Note different exit and entry corridors. B, Movements of an individual showing no regular pattern, 1967.

One of three males caught at the pond border in trap NNW and transported to a point in a meadow 45 m. NNW of this trap behaved unexpectedly when released under cloudy skies and a NW wind (*ca.* 10 knots). Instead of crawling under cover as animals released in daylight usually do, the salamander, out of sight of the pond or its bordering trees, extended its limbs and slowly moved its head from side to side. Within a few seconds it began to move uphill at a heading of 153°. After moving approximately 3 m. in a straight line, it stopped, repeated the limb extension and lateral head movements, changed its heading to 170° and continued this heading with occasional pauses, moving over the ridge and down to trees bordering the pond where it burrowed under leaves west of the pond center. Slight detours around obstructions were made, but the resulting path of travel on the 34 m., 170° route varied less than 1 m. from a straight line. This animal paused nine times during the 27-minute excursion with a rate of travel during the active portions of about 2 m. per minute.

Except for animals released 20 m. from ENE, all the returning animals had to move some distance uphill before reaching the pond. Animals released 175 m. from NNW were in a different drainage system and had to cross a 9-m.-high

TABLE I
Results of releases of animals originally captured at pond border

Point of capture at pond	Point and direction of release	Returns to pond no. released
ENE	20 m. ENE	3 3
ENE	45 m. NNW	3 3
ENE	135 m. ENE	2 3
ENE	175 m. NNW	3 3
NNW	20 m. ENE	2 3
NNW	45 m. NNW	3 3

ridge with 35° slopes. The time required to return to the pond varied from two to ten days but extremely dry weather also slowed normal migration of other animals at this time. One of the two animals not reaching the pond returned the following year.

In a related experiment late in the 1966 breeding season, 57 animals moving from many directions and collected on arrival at another pond (B) were released at points 200 and 250 m. from Pond B. A salamander returning from the 200 m. release point could encounter any of three other ponds never known to harbor breeding salamanders; return from the 250 m. release point involved crossing a lawn, a large, steep (35° slope) 25-m.-high ridge, and possibly encountering a permanent pond. Two days after the releases the area between the pond and the release points burned, adding strong odors to the area.

Of 28 animals released 200 m. from the pond, 13 were eventually recaptured at the pond border, 7 in the same season, 6 more one year later. Nine of 29 released at 250 m. were eventually recaptured at the pond border, one the same season, one in the autumn, and 7 the following spring. Returns of the same season tended to enter the pond at points nearest the release point. In

both cases returns after two years approximated the percentages of returns by non-displaced animals.

Since the ability of some animals to return from distances up to 250 m. was established, four animals (two males, two females) were collected at dawn at Pond B border early in the 1967 breeding season, placed in covered containers and carried in a truck for one day. At night, phosphorescent vinyl bands were placed around the bodies of the animals, and they were released on a golf course fairway during a heavy rain, 500 m. from Pond B. Return to the pond involved crossing over 100 m. of lawn, encountering up to four temporary ponds, one road, and a small stream.

At release all four animals took headings within 15° of the pond direction. They were turned around by hand but they immediately headed in the pond direction (uphill) and moved approximately 25 m. before being lost from view. After 11 days one of the animals (♀) arrived at the pond border trap nearest the release site. The other three animals were not seen again.

If orientation to a breeding pond is a simple geotaxis, rheotaxis or random search, some migrating animals collected while moving to a breeding pond and displaced close to another breeding pond should then move into the "new" pond. Negative results would suggest that geotaxis, rheotaxis, or random search must be coupled with unknown qualitative factors of the pond itself or that these taxes are not the major factors used in finding the pond.

To test for the above taxes, nine salamanders captured as they moved to Pond B were carried at night in covered containers to sites near Pond A. Three animals were released in a short-grass meadow at each of three compass points (0° , 90° , and 270°) 10 m. from the Pond A border. In two cases, the pond was downhill from the release site; in the other, slightly uphill. None of the animals (6 males, 3 females) moved to the new pond (A). Eight of nine control animals (captured at the Pond A border) moved to Pond A. The following year three of the displaced animals (all males) were recaptured at Pond A, and two of these animals were seen the succeeding year.

Six males transported from the Pond B border and released in the center of Pond A stayed within Pond A from four to 22 days. Their exit directions bore no relationship to the directions from which they attempted to enter their original pond (B) or the direction to Pond B. One of these animals returned to Pond A the following year and was captured while migrating in a nearby wooded area the succeeding year but not at the Pond A border.

Displacements to an arena

In an effort to remove all possible local cues surrounding breeding Pond B, 106 migrating animals were transported to the laboratory and maintained in running lake water aquaria for periods varying from three to eight days. These animals were released one or more times in a 40-m.-diameter arena on the Wellesley College campus under cloudy and clear skies at night during the normal migration period. No significant differences in performances between males and females or between cloudy and clear conditions were noted. A total of 547 releases was made, and the capture points around the inner face of the fence were determined.

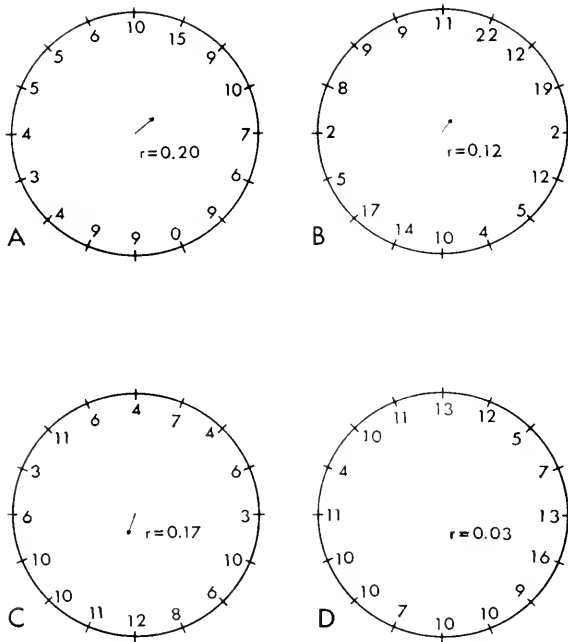


FIGURE 6. Representation of capture points of animals released in arena about one mile from home pond. A, Relation of headings of animals released facing 90° to the left of their heading when captured in the field; top of circle represents original heading in field, not north. B, Headings of animals released by removal of covering container, in relation to original heading in field; top of circle represents original heading. C, Compass headings of animals released facing ENE; top of circle is N, SSE is direction to home. D, Compass headings of animals released by removal of covering container; top of circle is N, SSE is direction to home.

The data obtained by releasing animals facing 90° to the left of their original heading in the field, or released at chance headings (by removing a covering container), in relation to direction of original field heading are random ($r \leq 0.20$; Fig. 6 A, B). Pooled results of releases of animals faced ENE or released by removal of a covering container (Fig. 6C, D, respectively) are random ($r < 0.20$) and show no relation to the true homeward direction, SSE. These results demonstrate that under the conditions of these experiments the animals have no preferred direction in relation to the way they were traveling when collected or in relation to the direction of the home pond approximately one mile away.

DISCUSSION

Whitford and Vinegar (1966) displaced *Ambystoma maculatum* adults up to 420 ft. (ca. 128 m.) from a breeding pond and found they were able to return to the pond, apparently aided by a rheotactic, or perhaps olfactory, response, thereby adding support to a finding of rheotactic orientation of *A. maculatum* by Finneran (1951). Under the conditions of my experiments rheotaxis apparently plays little or no role in locating the home breeding pond.

If the four animals displaced 500 m. from their home pond had no sensory contact with that pond or known routes of travel to it, they may have used a true navigation ability (Type III orientation of Griffin, 1952; reverse displacement or bicoordinate navigation of Schmidt-Koenig, 1965) in their initial oriented headings. The life-time movements of spotted salamanders are unknown; therefore displacements to unfamiliar territory are contingent upon thorough studies before true navigation can be established.

The random headings of animals released in the arena about 1 mile away from the home pond may parallel homing studies with *Taricha rivularis* in California (Twitty, 1959, 1961). Oldham (1967) reported similar results in displacements of green frogs, *Rana clamitans*. David L. Grant (personal communication), who is continuing the studies of *Taricha* orientation, utilizes a much larger "arena" to determine headings of animals displaced large distances. Apparently *Taricha* requires some time and space to determine and maintain correct headings to the home area, but in a relatively small arena *Taricha granulosa* can utilize a sun compass to move on land in a direction that would take the animals to a home shore (Landreth and Ferguson, 1967).

Laboratory conditions may affect the performances of *A. maculatum* released in an outdoor arena. Since handling and laboratory storage of females may prevent ovulation in some unknown way (Shoop, 1967), a physiologic barrier resulting from these conditions has been demonstrated. Perhaps a similar "upset" was responsible for the results of the arena release experiments.

Since spotted salamanders leaving a breeding pond often maintain a very similar or identical track away from the pond as they took to it, the animals could conceivably run through an imprinted cue system obtained as they left the ponds as juveniles and reverse it on returns to the pond, although day-length, temperature, sun altitude, and star patterns are different. Whether they always migrate every year only in familiar territory (home range) remains unknown. Uzzell (personal communication) and I have observed other ambystomatids migrating to areas where temporary ponds were not yet formed, as is often the case with the terrestrial egg-layer of the group, *A. opacum*. I observed a population of *Ambystoma talpoideum* moving into an area where a breeding pond was previously located but road construction between breeding seasons had obliterated the previous habitat. Heusser (1960) made a similar observation of migrating toads (*Bufo bufo*). While local visual or olfactory cues may play important roles in orienting to a breeding pond, in these instances, other cues were probably operative.

Many salamanders show a decided preference for a breeding pond and return year after year. Others may adopt a new breeding pond if displaced. Until lifelong records of movements or home ranges and perceptive abilities of amphibians are known, definitive statements regarding navigation and sensory mechanisms utilized in orientation will be open to question.

I thank Patricia Moehlman, Robert Rice, and Phyllis Phelps for aid during several portions of the study. E. Willow Reed, Martha Garrett, and Ann Scott assisted in the field. Dr. Edwin Gould criticized an early manuscript. I am pleased to acknowledge the financial support of the U. S. Atomic Energy Commission (Contract AT(30-1)3554) and Wellesley College.

SUMMARY

1. Migrating spotted salamanders entered and departed from breeding ponds at or near the same point. This pattern was repeated for three consecutive years.
2. Animals migrating in a meadow to and from the breeding pond often traveled in a corridor 10 to 30 m. wide. Several variations from this plan were noted.
3. Spotted salamanders displaced up to 500 m. may return to the home breeding pond.
4. Migrating adults transferred from the border of one breeding pond to within 10 m. of another breeding pond did not move to the new pond as did controls.
5. Migrating adults transported from the border of one pond to the center of another failed to orient to the old pond or in the direction they were moving when captured.
6. Some adults adopted a new breeding pond when displaced.
7. Animals moved to a laboratory and then released outdoors in an unfamiliar area failed to orient to their pond or in the direction they were moving when originally collected. Handling and laboratory conditions may have altered normal behavior.
8. Since lifelong records of salamander movements are lacking, certainty of navigation ability may be questioned.
9. Sensory bases for orientation remain poorly known.

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POLAROTAXIS IN COPEPODS. I. AN ENDOGENOUS RHYTHM
IN POLAROTAXIS IN *CYCLOPS VERNALIS* AND ITS
RELATION TO VERTICAL MIGRATION

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In many terrestrial arthropods, the ability to detect polarized light is used to find geographical directions by means of sky polarization patterns which change with the sun's position in the sky (von Frisch, 1948; Pardi and Papi, 1953a, 1953b; Birukow, 1954; Vowles, 1955; Wellington, 1955; Papi, 1955). Such direction-finding is actually a special extension of the sun compass and requires that the organism possess a physiological clock to correct for the sun's movement through the sky and the concomitant changes in sky polarization.

The functional significance of polarized light sensitivity for planktonic animals has yet to be demonstrated. However, many such organisms can perceive polarized light (Baylor and Smith, 1953; Jander and Waterman, 1960) and polarization patterns are indeed present in the aquatic habitat (Waterman, 1954; Waterman and Westell, 1956; Ivanoff and Waterman, 1958; Ivanoff, Jerlov and Waterman, 1961). It is reasonable to speculate that these aquatic organisms use the polarization pattern of their environment for orientation just as do their terrestrial counterparts. This paper presents evidence that the natural pattern of polarization in the aquatic habitat is used as a frame of reference for swimming orientation in the copepod *Cyclops vernalis* and that the orientation of this animal to the plane of polarization changes rhythmically throughout the day in an ecologically meaningful way.

GENERAL BEHAVIOR

Material and Methods

As experimental animals, the fresh-water copepod *Cyclops vernalis* Fischer, collected in Clark's Pond near New Haven, Connecticut, was cultured at 20° C. in an aerated five-gallon aquarium in a darkroom having a 12-hour light, 12-hour dark cycle of illumination. Animals had lived in this culture for more than a month before experiments were begun.

To measure phototactic responses, 20 copepods were placed in a rectangular, clear Plexiglas vessel. One-half of this vessel was lined and covered with black Plexiglas to shield this half of the container from the overhead tungsten filament light source. An intensity of light set at 1100 lux was chosen to be the value incident at the water surface since this is a daylight intensity level commonly found in natural waters (Harris and Wolfe, 1955). The phototactic sign of the copepods was determined by measuring their preference for the light or dark half of the vessel at various times of day. The number of copepods in the light half of the vessel was recorded every 20 seconds over a period of 10 minutes, permitting an

average to be calculated. These observations were made at 2-hour intervals over the course of the 12-hour light period.

To ascertain geotactic responses, 20 copepods were placed in a 500-ml. graduated cylinder in complete darkness; there was no light-dark cycle. The copepods were observed with a red light at all times of the day to determine their distribution in the vessel. The number of copepods in the top third, middle third and bottom third of the vessel was recorded at 20-second intervals for 10 minutes. These observations were made at 2-hour intervals during the course of a 24-hour day.

Results

The copepods were photonegative at all times during the 12-hour light period, spending 90% of the time in the dark. When an occasional animal would venture into the lighted half of the vessel, it remained at the bottom and seldom swam upwards toward the light. There was no evidence for changes, rhythmic or otherwise, in phototactic sign.

The copepods were also geopositive at all times of day, spending 85% of the time on the bottom of the graduated cylinder. Again, there was no evidence for changes in geotactic sign at any time of the day.

It was also observed that when swimming horizontally in the water, *Cyclops vernalis* directed its ventral surface upward toward the overhead light. If light was shined from below, the ventral surface then faced downward, still toward the light. This behavior indicated that a ventral light reflex was present. When swimming vertically upward, the anterior end always led with the long axis of the body vertical. When the copepod moved downward, it either actively swam with its anterior end downward and long axis of the body vertical or it passively sank with its body axis horizontal.

Cyclops vernalis tended to cling to the sidewalls and bottom of any experimental vessel and often adhered to the water-air interface when one was present. It sporadically swam freely in the water. This behavior in the laboratory agrees with the field behavior as reported by Pennak (1966). Thus *Cyclops vernalis* is essentially restricted to vegetation in the littoral zone of lakes and ponds, being seldom found in the open water. It is a browsing species that often leaves the substrate to cruise about in spaces among the vegetation. This suggests that the copepods in the laboratory culture were in normal physiological condition.

POLAROTACTIC BEHAVIOR

Methods

The apparatus used to measure polarotaxis was similar to that employed by Bainbridge and Waterman (1957). A shallow cylindrical Plexiglas cell, 26 mm. deep and 108 mm. in diameter, containing 20 copepods, was illuminated from above by a vertical beam of linearly polarized light whose plane of polarization could be altered at will. The light originated from a tungsten filament lamp and passed successively through a focusing lens system, a heat filter, a depolarizer, a rotatable Polaroid linear polarizer, the experimental cell and finally through two lenses which produced an image of the copepods on a sheet of paper on the bench beneath.

The intensity of light was 1100 lux at the upper surface of the experimental cell.

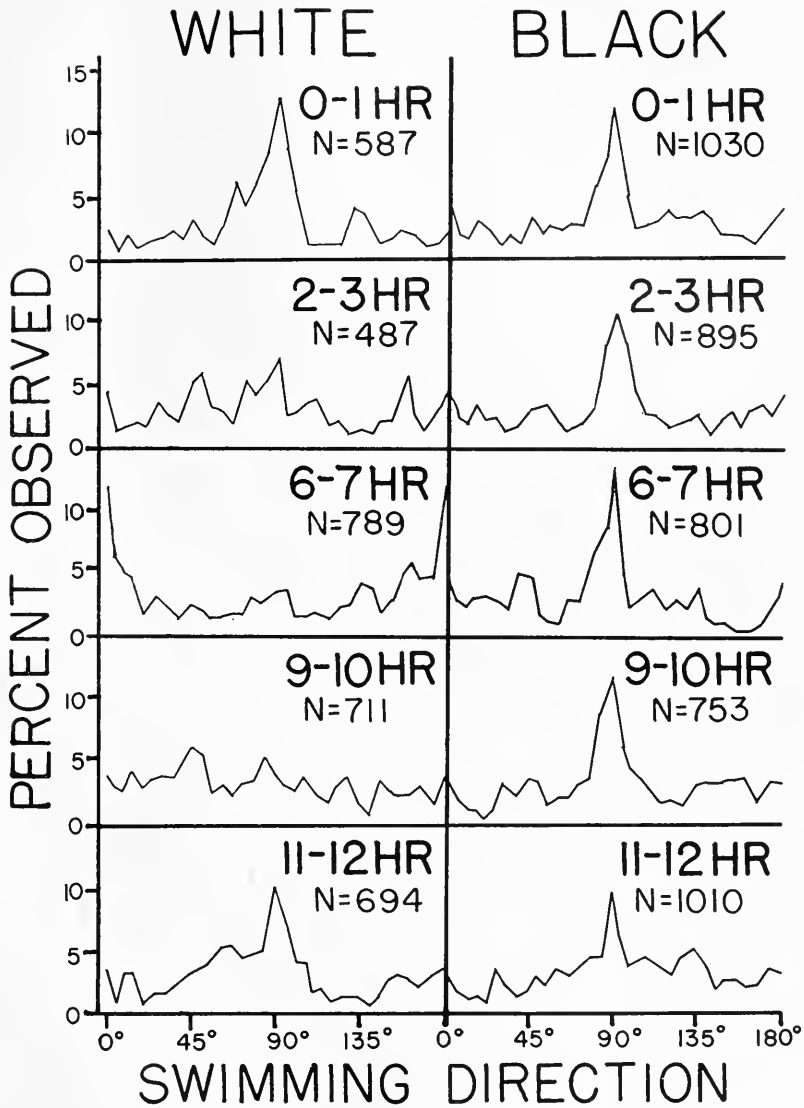


FIGURE 1. Polarotaxis at different times during a 12-hour light cycle. Abscissa gives the angle between swimming path and plane of polarization. Ordinate gives the percentage of total swimming attempts. N gives the total number of swimming attempts. WHITE and BLACK denote the color of the sidewalls and diaphragm of experimental cell.

The vertical outer walls of the cell were covered with either black or white construction paper to minimize direct reflections from the walls. In addition, the walls were shaded from direct light with a circular black or white diaphragm having an aperture 20 mm. less than the inner diameter of the cell. The meniscus at the margin of the water surface was eliminated by pressing a flat glass plate against

the upper edge of the cell walls, thus sealing off the completely full vessel without an air-water interface. The water was filtered twice to reduce the differential scattering of polarized light by a turbid medium (Bainbridge and Waterman, 1958; Waterman, 1960).

Using this apparatus, the paths of the swimming copepods could be traced by following their projected images with a pencil on a piece of paper on the bench beneath the experimental cell. The plane of polarization was first set to one of four positions and noted on the paper along with the tracings. After a number of observations had been made, the plane of polarization was then changed at random to a new position and more tracings were made. A total of four different e -vector positions was used for each trial to eliminate the effect of any intensity or spatial cues in the cell.

The angle between the path of the copepod and the plane of polarization was then measured with a protractor at 1-cm. intervals along the length of each path tracing. The frequency distribution of the resulting angles indicates any relationship between the course of the copepods and the plane of polarization. Thus, if a copepod tended to swim perpendicular to the plane of polarization, a peak in the orientation readings would appear at 90° on the plot. The frequency distributions can be statistically evaluated as either random or with significant peaks by a chi-square test (Bainbridge and Waterman, 1957).

As indicated previously, *Cyclops vernalis* tended to cling to the bottom and sides of the experimental cell and only sporadically swam freely in the water. In order to get a sufficient number of tracings in a 30-minute period, 20 copepods were placed in the cell to insure that one or two individuals would always be swimming. The paths of the copepods were recorded at 2-hour intervals throughout the 12-hour light period; no readings were attempted during the 12-hour dark period of the illumination cycle. In the following discussion, therefore, a time of 0-1 hours indicates the first hour after the light went on (experimental "dawn") and a time of 11-12 hours indicates the last hour before the light went off (experimental "dusk"). During each 1-hour period of observation, both black and white sidewalls and diaphragm were used to discover if any light contrast reactions were present as reported in several animals by Jander and Waterman (1960).

Results

When black sidewalls and diaphragm were used, *Cyclops vernalis* oriented at right angles to the plane of polarization at all times during the 12-hour light period. However, when white sidewalls and diaphragm were used, the same copepods in the same vessel at the same times of day now showed a regular 12-hour rhythm in polarized light orientation. Under these conditions, there was a 90° orientation to the plane of polarization at the beginning and end of the light period and a 0° orientation midway through the light period. These orientation peaks were statistically significant at the 0.5% level using a chi-square test. At intermediate times, however, orientation was poor and not significant statistically, appearing to be transitional between 90° and 0° orientation. The results for one such day are shown in Figure 1. The same results, using other groups of 20 copepods, were obtained on two other days when the experiment was repeated.

POLAROTACTIC BEHAVIOR UNDER SIMULATED NATURAL CONDITIONS

Introduction

In natural waters, the angle between the plane of polarization and the horizontal depends upon the line of sight of the observer, the position of the sun in the sky and the depth of the observer in the water (Waterman, 1954; Waterman and Westell, 1956). The plane of underwater polarization is always horizontal in horizontal lines of sight toward and away from the sun's bearing, but is tilted toward the sun in all other azimuths. The amount of tilting between the plane of polarization and the horizontal is maximal at right angles to the sun's bearing and is approximately equal to the sunlight's angle of refraction at the surface of the water. However, there is a decrease in the tilting of the polarization plane as depth increases.

At sunrise and sunset, when the sun is near the horizon, the angle of the plane of polarization may be as great as 45° along horizontal lines of sight perpendicular to the sun's bearing. At midday, when the sun is nearest the zenith, the plane of polarization along a line of sight perpendicular to the sun's bearing reaches its minimal tilt from the horizontal. In fact, when the sun is in the zenith, the *e*-vector is horizontal in all lines of sight. This then is the simplest pattern of natural underwater polarization and the most easily imitated in the laboratory.

Verheijen (1958) and Waterman (1961) have emphasized that much of the experimental work on light reactions of aquatic animals has been conducted under conditions atypical of those in open water. In the previously described experiment, the plane of polarization was directed vertically from above the experimental cell. This experiment might simulate polarotaxis in shallow waters where overhead sky patterns are directly visible over about 90° . However, such conditions of polarization are limited to shallow waters and could not model the effect of lateral illumination from underwater polarization in deeper waters.

Methods

In order to approximate more natural conditions in the study of polarotaxis, the following setup was employed: Fifty copepods were placed in a rectangular battery jar, 156 mm. deep by 100 mm. long by 80 mm. wide. Sheets of Polaroid were placed on each side of the jar so that the plane of polarization was horizontal along all lines of sight at all times of day. This simulates the simplest natural pattern described above. The jar was illuminated from above by unpolarized light. Both the vertical and horizontal illumination was provided by tungsten filament lamps, but their intensities were set in two different patterns.

In surface waters the light intensity from above is much greater than the light intensity from the sides. To approximate this in the laboratory, the light sources were adjusted such that the intensity from above was 1100 lux inside the battery jar and the intensity from the sides was 100 lux inside the battery jar. As depth increases in natural waters, the ratio of the vertical to horizontal illumination decreases (Jerlov, 1951). To approximate the pattern of light intensity found in deeper water, the light intensities from above and from the sides were made equal inside the battery jar and set at either 1100 lux or 100 lux.

The intensity of light was controlled with a rheostat and measured with a luxmeter. Since changing intensity with a rheostat can produce changes in the wave-length of the emitted light, any behavioral changes induced by different intensity patterns might be due to different reactions of the organism to different wave-lengths of light. To establish that this was not the case, the intensities were also varied by changing the distance of the light source from the battery jar; this procedure does not change the wave-length. It was found that similar results were obtained in this set of experiments and in all others requiring differential intensities regardless of the manner in which intensity was varied.

The swimming copepods were observed visually and their paths were tallied in the following way: On a clear piece of plastic a circle was drawn and divided into 16 equal parts by 8 lines drawn as diameters through the center. One of these diameters was chosen to represent an up-down axis; the diameter perpendicular to it represented a left-right axis. The other diameters represented intermediate directions. When a copepod began swimming freely in the water, the center of this circle was visually aligned to the copepod's starting point so that one of the axes of the circle closely approximated the path of the swimming animal. The paths were then recorded over a period of a half hour and a frequency diagram plotted. The angle between the path of the copepod and the plane of polarization could be estimated only roughly with this method. Any given excursion of a copepod from rest into the open water had to be approximately aligned to one of the reference axes on the plastic sheet; the error involved could be as great as 11° , but was generally much less. The copepods were observed at approximately 6-hour intervals throughout the course of the 12-hour light period.

Results

Using horizontal polarization to simulate natural conditions, the effect of the angular distribution of light intensity on polarotaxis was studied (Fig. 2). With a contrasting ratio of vertical to horizontal intensity (1100 lux from above, 100 lux from the side), polarotaxis was always primarily 90° to the horizontal plane of polarization with no rhythmic responses present. However, with a uniform angular distribution of light intensity (equal intensity from both the sides and above) of either 1100 lux or 100 lux, a rhythm was present at both intensities. Compared with a control group where no polarization was present, these copepods swam more vertically (at right angles to the plane of polarization) at the beginning and end of the 12-hour light period and more horizontally (in the plane of polarization) at the middle of the light period. This indicates that a uniform angular distribution of light intensity over a range of absolute intensities induces the diurnal rhythm in the polarotactic response.

Swimming in unpolarized light is basically horizontal at all times of day, but orientation is not precise. This is true in uniform intensity patterns of either 1100 lux or 100 lux and in contrast intensity patterns as well. The addition of horizontally polarized light both increases the precision of horizontal swimming and induces a periodic vertical swimming component. The control group further demonstrates that there are no rhythms in orientation in unpolarized light. Polar-

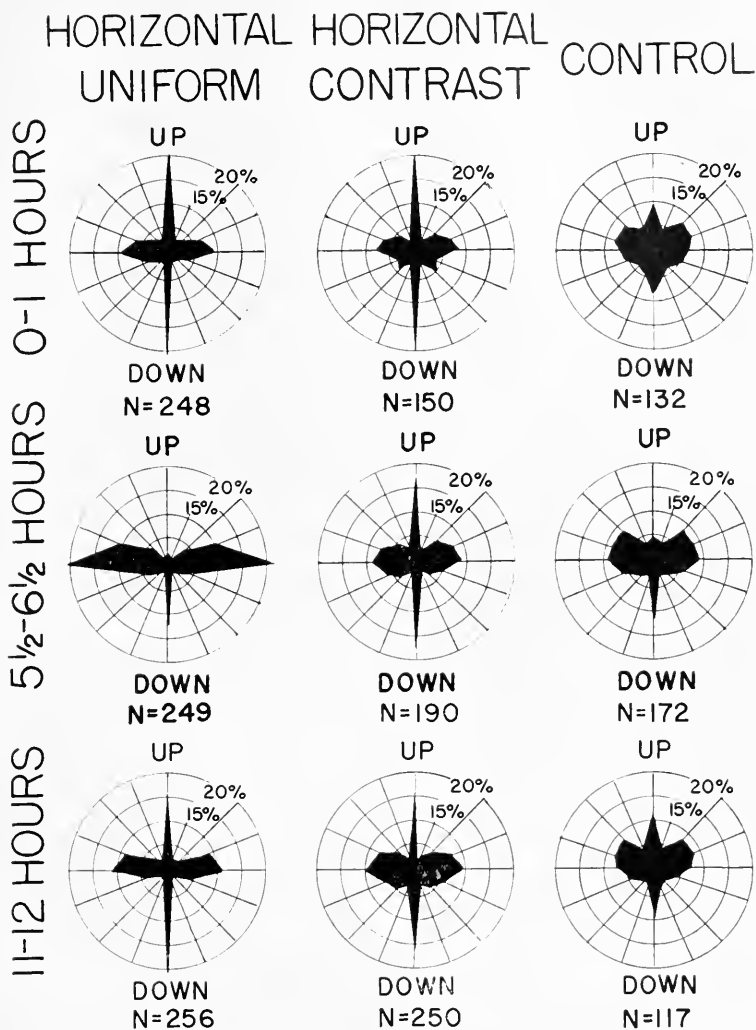


FIGURE 2. Orientation to horizontally polarized light at different times during a 12-hour light cycle. Radial axes represent direction of swimming with respect to gravity. Concentric circles represent the percentage of total swimming movements. N gives the total number of swimming movements. HORIZONTAL means horizontally polarized light was used; the plane of polarization is perpendicular to the UP-DOWN axis. UNIFORM means the light intensities from above and from the sides were equal. CONTRAST means the light intensity from above was greater than from the sides. CONTROL group had no polarization; angular distribution of intensity had no effect on this group.

ization must be present (along with a uniform angular distribution of intensity) to elicit a diurnal rhythm in the orientation behavior of these copepods. Furthermore, contrast and uniform angular distributions of light intensity elicit the same response (predominantly horizontal swimming) in unpolarized light, but each elicits a different response in polarized light.

In order to demonstrate conclusively that the observed rhythm in swimming behavior was not influenced by gravity responses, the lateral patterns of polarization were varied. The intensity in all cases was 1100 lux inside the battery jar from both above and from the sides. If the light from the sides was polarized in a vertical plane, instead of horizontally as above, the diurnal rhythm in polarotaxis still persisted, but now the copepods swam horizontally with respect to gravity (but still 90° to the plane of polarization) at the beginning and end of the light period and vertically with respect to gravity (but still 0° to the plane of polarization) midway through the light period.

If the plane of polarization was set at an angle of 45° to the horizontal on all sides of the jar, the copepods swam at an angle of 45° upwards and downwards with respect to gravity at all times of the day. The rhythm did not appear under these conditions since swimming at either 0° or at 90° to the plane of polarization both result in swimming 45° to the horizontal with respect to gravity. The frequency diagrams for these experiments are not figured in the paper since they can be adequately described verbally.

POLAROTACTIC BEHAVIOR IN CHANGING INTENSITIES OF LIGHT

Introduction

All the experiments described so far indicate that polarized light merely orients swimming direction more precisely, either in a vertical, horizontal or 45° to the horizontal direction, but causes no net displacement of the population, *i.e.*, migration. Inspection of Figure 2 shows that given a population of *Cyclops vernalis* swimming primarily horizontally in unpolarized light, the addition of horizontally polarized light will alter the swimming behavior by inducing a vertically directed swimming component. However, this vertical component is equally divided into upward and downward fractions, causing no migration in the population.

Experiments on other planktonic organisms (Ewald, 1912; Clarke, 1930, 1932; Johnson, 1938; Johnson and Raymond, 1939) indicate that changing light intensities rather than constant illumination cause net displacements of the populations: increasing intensity causes a downward migration and decreasing intensity causes an upward migration when the light source is overhead. Similar changes have been observed in the deep scattering layer in nature (Hersey and Backus, 1962). This sound-scattering layer, composed of marine organisms, migrates upward at night and downward during the day, being greatly influenced by light intensity.

Methods

Since polarized light of constant intensity causes no migrations of the swimming *Cyclops vernalis*, the effects of intensity changes with and without horizontal polarization were investigated to see if net displacements could be elicited. Observations were made with either decreasing or increasing light intensities from above and/or from the sides to approximate the conditions at sunset and sunrise, respectively.

In cases of decreasing intensity, the initial intensity of light was 1100 lux and decreased by steps of 200 lux every ten minutes until a final intensity of 100 lux was reached. In cases of increasing intensity, the initial intensity was 100 lux and increased by steps of 200 lux every ten minutes until a final intensity of 1100 lux was reached. In addition, either the overhead and lateral lights changed intensity at the same rate (uniform angular distribution of intensity) or the lateral light remained constant at 100 lux while the overhead light changed in intensity (contrast angular distribution of intensity). The intensity of light was controlled manually with a rheostat and measured with a luxmeter as described previously.

Results

When polarization is not present, intensity changes caused no net displacements in the population; swimming remained primarily horizontal in all cases.

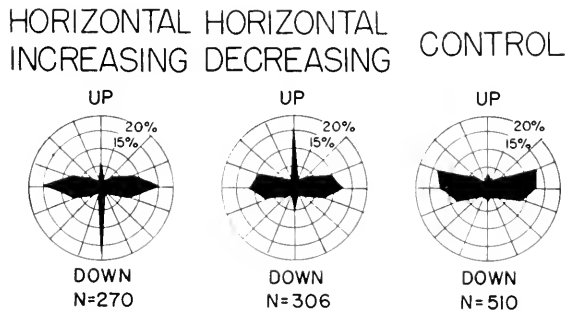


FIGURE 3. Orientation in changing light intensities. Method of diagramming is same as in Figure 2. HORIZONTAL means horizontally polarized light was used. INCREASING and DECREASING describe the change in light intensity. CONTROL group had no polarization; changing intensity of either type had no effect on this group. Time of day and pattern of intensity distribution had no effect on intensity responses.

Without polarization, then, orientation behavior in changing intensities was similar to that found with a constant intensity.

However, when the lateral light was polarized horizontally (Fig. 3), increasing intensity caused a net downward movement in vertically swimming individuals whereas decreasing intensity caused a net upward movement in vertically swimming individuals. Horizontally swimming individuals were not affected.

These results in horizontally polarized light held regardless of whether overhead and lateral lights changed intensity at the same or at different rates. The time of day also had no effect on the results; changing intensities always caused a vertical displacement of the population as long as polarized light was present. This indicates that the cue of changing intensity is stronger than the rhythm in polarotaxis that occurs when intensity is held constant: The 0° orientation to the plane of polarization that occurs at midday in a uniform intensity distribution of unchanging intensity is changed to a 90° orientation when changing intensities

are employed. However, since intensities are normally constant at midday in nature, the rhythm probably persists under natural conditions.

DISCUSSION

Ecological significance

From the experiments with *Cyclops vernalis* it is clear that experimental conditions including unpolarized light at constant intensity in all directions at all times of day are highly artificial. Extremely interesting patterns of behavior that would never occur under such highly artificial light regimes are elicited when certain aspects of natural conditions of illumination are simulated, even

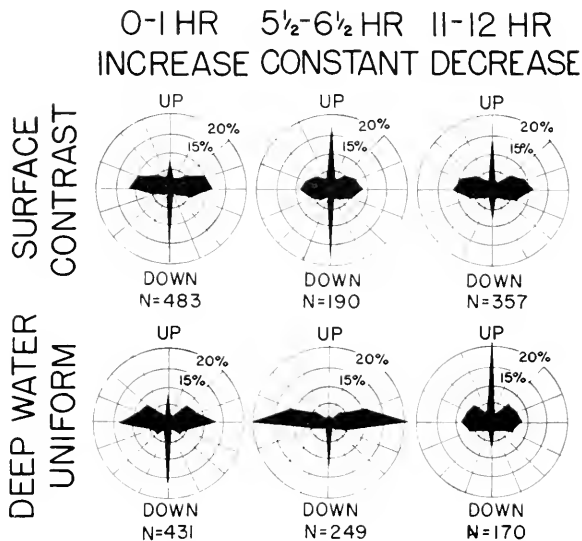


FIGURE 4. Summary of experiments using horizontally polarized light at different times of the 12-hour light cycle. Method of diagramming is same as in Figure 2. INCREASE, CONSTANT and DECREASE describe change in light intensity. CONTRAST describes intensity distribution found at the SURFACE of water. UNIFORM describes intensity distribution in DEEP WATER.

crudely. In order to understand the ecological significance of the various behavioral reactions of *Cyclops vernalis* to light, the various parameters of the experimental illumination must be given a natural interpretation.

In this study, a horizontal polarization of light along all lines of sight was used to approximate the simplest of natural patterns of polarization in the aquatic habitat. It must be emphasized that this is a pattern that occurs only in low latitudes around noon. Increasing intensities of light were used to simulate conditions occurring during the early morning hours whereas decreasing intensities were used to approximate the conditions of illumination at dusk; constant intensities were used to represent conditions at midday. The effects of depth on light

intensity patterns were crudely mimicked by changing the intensity ratio of the overhead and lateral lights and by the use of black or white sidewalls and diaphragm in the experiments using polarized light from above.

A summary of the experimental results is presented in Figure 4. Only those studies performed under conditions of illumination approximating those found in nature are used; those studies that elucidated the behavioral reactions but which have no natural counterpart (such as increasing intensities at the end of the light period) are omitted.

When all the experimental results are arranged in a natural sequence, the following trends appear: In the morning, swimming has a net downward component both at the surface and in deep water. A population behaving in this manner would tend to sink in the water. At midday, those copepods that sank into deep water in the morning would now move primarily horizontally in the water, maintaining their position in deep water. However, those copepods that remained at the surface during the morning would still tend to move vertically up and down at midday. Copepods moving vertically downward from the surface might eventually reach a more uniform angular distribution of light intensity, causing them to fan out horizontally.

In this way, the vertically swimming copepods found at the surface at midday would gradually be "trapped" by the deep-water angular distribution of light intensity, eventually causing this surface population to redistribute in deeper water. In the evening, swimming has a net upward component both at the surface and in deep water. This would cause the population to rise toward the surface in the evening. These trends in swimming behavior are consistent with the patterns of movement shown by many planktonic Crustacea in their vertical migrations. This implies that the parameters of illumination examined in this study may be very important for the orientation and release of migratory behavior in the aquatic habitat.

Physiological significance

If the foregoing interpretation of the experimental results with *Cyclops vernalis* is correct, then the physiological significance of the various parameters of illumination can be summarized as follows: The angular distribution of light intensity is a visual cue giving information as to the copepod's relative depth in the water. The plane of polarization is used as a frame of reference for orienting locomotor movements in the water where other orienting cues such as landmarks are sparse. By swimming at right angles to the plane of polarization, the copepod will move predominantly vertically at most times of day; by swimming in the plane of polarization, the copepod will move horizontally much of the time. Thus the polarization pattern of the aquatic habitat provides the copepod with vertical-horizontal reference coordinates with which to orient. Changing intensity of illumination provides migratory cues that lead to a net displacement of the population only when it is correctly oriented. Increasing intensity is a cue to swim downward, whereas decreasing intensity is a cue to move upward. The diurnal rhythm in polarotaxis that occurs only in deep-water uniform intensity distributions functions as a behavioral "trap" to keep the copepods in deep water during midday.

This investigation was supported by a National Aeronautics and Space Administration traineeship awarded through the Department of Biology at Yale University. Appreciation is expressed to Professor Talbot H. Waterman for the stimulation, guidance and comments that made this study possible.

SUMMARY

1. A series of laboratory experiments on the fresh-water copepod, *Cyclops vernalis*, revealed an endogenous rhythm in its orientation to polarized light. The copepod swam perpendicular to the plane of polarization at the beginning and end of a 12-hour diurnal light period, but swam parallel to the plane of polarization midway through this light period.

2. The rhythm was not present under all experimental conditions, being released only when the intensity of light from above was equal to that from the side. Under other experimental conditions, the copepods swam perpendicular to the plane of polarization at all times during the light period.

3. A pattern of horizontal polarization, crudely simulating the simplest natural underwater pattern, was found to provide *Cyclops vernalis* with vertical-horizontal reference coordinates for spatial orientation.

4. Changing intensities of polarized light induced net displacements of the population, thus effecting vertical migration in the laboratory.

5. The ratio of overhead to lateral light intensity provided a visual cue indicating water depth. The diurnal rhythm in polarotaxis operated only with deep-water intensity ratios of horizontal and vertical light and served to orient deep-water individuals in a horizontal plane at midday, thus inhibiting vertical swimming patterns (which occur at dawn and dusk) that might carry them to the surface at that time.

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POLAROTAXIS IN COPEPODS. II. THE ULTRASTRUCTURAL BASIS AND ECOLOGICAL SIGNIFICANCE OF POLARIZED LIGHT SENSITIVITY IN COPEPODS

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The ability to perceive linearly polarized light and to determine its plane of polarization has been demonstrated both behaviorally and electrophysiologically in a great number of species, mostly in arthropods but also in cephalopod mollusks (Waterman, 1966a). The mechanism of polarized light perception is localized at the retinal level and, in the cases adequately known, apparently depends on a two-channel system comprising a pair of dichroic analyzers oriented at right angles (Waterman, 1966b; Waterman and Horsch, 1966; Eguchi and Waterman, 1966, 1967).

In a previous paper (Umminger, 1968), polarotaxis was behaviorally demonstrated in the copepod *Cyclops vernalis*. This was the first reported case of polarized light sensitivity in an organism possessing a naupliar eye exclusively. Earlier investigators (Stockhammer, 1959; Jander, 1966) had failed to find polarotaxis in copepods.

The purposes of this paper, then, are two-fold: First, an ultrastructural examination of the naupliar eye of *Cyclops vernalis* was undertaken to determine if there was a two-channel system of microvilli present as in polarotactic arthropods with compound eyes. Secondly, a great number of copepod species were examined for polarotactic behavior to see if the ability to perceive polarized light is universal or limited to only a few species of copepods. If polarotaxis is not universally present in copepods, this would explain the failure of Stockhammer and Jander to find it in the species they studied.

FINE STRUCTURE OF THE NAUPLIAR EYE

Methods

To determine whether the polarotactic behavior of *Cyclops vernalis* Fischer could be explained by some structural component of the naupliar eye, the fine structure of the eye was examined. The entire copepod was first fixed for two hours with 5% glutaraldehyde buffered in 0.1 M Sorenson's phosphate solution at pH 7.4 (Sabatini, Bensch and Barnett, 1963). After washing with buffer, the copepods were cut in half and the metasomes post-fixed for another two hours in 2.5% osmium tetroxide, also buffered at pH 7.4 with Sorenson's solution. Fixation was followed by acetone dehydration, transfer to propylene oxide and embedding in epoxy resin (Luft, 1961). Sections were cut with a Porter-Blum MT-2 microtome and stained with uranyl acetate and lead citrate (Reynolds, 1963). Observations were then made in a Philips 200 electron microscope.

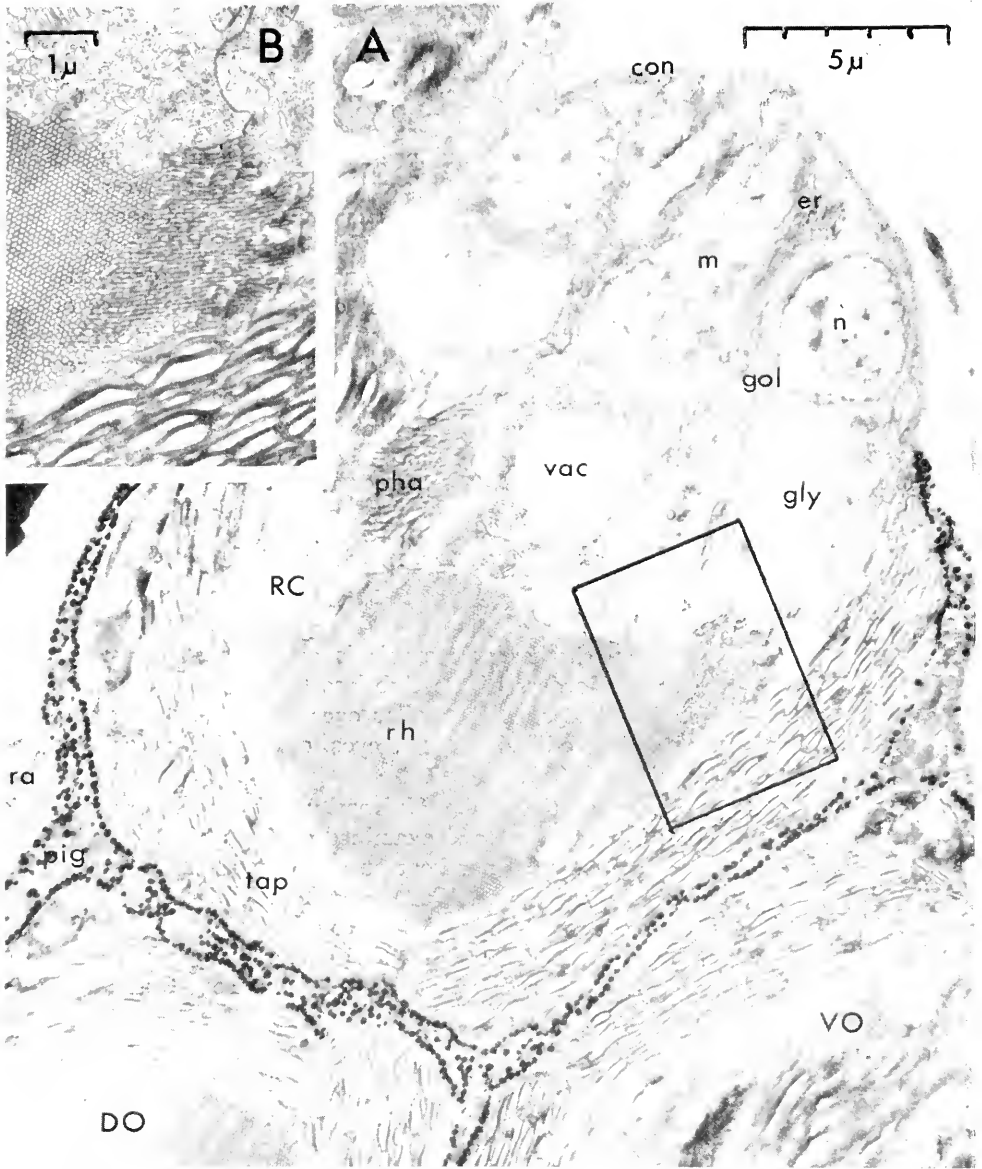


FIGURE 1. Longitudinal section through a dorsal ocellus of *Cyclops vernalis*. A. Portions of the other dorsal ocellus, DO, and the single ventral ocellus, VO, can be seen at the bottom. The two dorsal ocelli and the ventral ocellus make up the tripartite naupliar eye. con = conjunctiva, er = endoplasmic reticulum, gly = glycogen granules, gol = Golgi complex, m = mitochondrion, n = nucleus, pha = phaosome, pig = pigment cell, ra = retinular cell axons, RC = retinular cell, rh = rhabdom, tap = tapetum, vac = vacuole. Box represents area enlarged in B. B. Higher-power detail of the rhabdom.

Results

The "eye" of *Cyclops vernalis*, as is typical of most copepods, is composed of three ocelli on the anterodorsal region of the protocerebrum. The two dorsal ocelli each measure about $25\ \mu$ in diameter and point in a lateral direction. The single ventral ocellus points ventrally towards the rostrum.

The dorsal ocellus (Fig. 1) is composed primarily of several reticular cells whose inner ends have projections of microvilli forming the light-sensitive rhabdom. Of particular interest is the arrangement of these microvilli: those of one reticular cell meet those of another reticular cell at right angles (Fig. 1B). The reticular cells contain phasomes, much endoplasmic reticulum and many glycogen particles. A reflecting tapetum forms a hemispherical cup around the bases of the light-sensitive reticular cells whereas a thin conjunctiva covers their distal surfaces. Surrounding the tapetum, which is tightly packed with crystals, is a screening pigment cell containing granules of melanin.

Light enters the ocellus through the conjunctiva and passes through the distal parts of the reticular cells. It then strikes the rhabdom which has its microvilli perpendicular to the incoming light.

POLAROTACTIC BEHAVIOR IN COPEPODS

Material and Methods

In an attempt to discover whether polarotaxis is widespread in copepods, a number of species from several orders of the Subclass Copepoda were examined. Several of the species studied had simple naupliar eyes as does *Cyclops vernalis*, whereas others had elaborately modified naupliar eyes, often with cuticular lenses and movable parts (Vaissière, 1961).

In experiments performed at the Duke University Marine Laboratory, the copepods were freshly captured each day and examined at times of natural sunrise, midday and sunset; none were cultured nor kept on a 12-hour light, 12-hour dark cycle as was *Cyclops vernalis* (Umminger, 1968). At Yale University, specimens of *Tisbe furcata* (Baird) were cultured and kept on a 12-hour light, 12-hour dark cycle of illumination. In addition, the fresh-water *Diaptomus shoshone* Forbes was studied after it had been flown to New Haven from Colorado.

All species were tested with linearly polarized light in experiments identical to those performed on *Cyclops vernalis* in a previous paper (Umminger, 1968).

Results

Polarotactic behavior like that of *Cyclops vernalis* could not be elicited in all the species studied. Several species oriented to polarized light whereas other species showed no polarotactic responses whatsoever. Using linearly polarized light from above, *Caligus rapax* Milne Edwards, *Centropages hamatus* (Lilljeborg) and *Diaptomus shoshone* Forbes all swam at right angles to the plane of polarization. *Pontella mcadiei* Wheeler and *Tisbe furcata* (Baird) both oriented 0° , $\pm 45^\circ$ and 90° (4 peaks) to the plane of polarization. *Labidocera aestiva* Wheeler oriented 0° to the plane of polarization. In all these species, however, the angle

of orientation was the same at all times of day and under all conditions of illumination. There were no rhythms in polarotaxis.

The species showing no polarotactic responses were *Pseudodiaptomus coronatus* Williams, *Acartia tonsa* Dana, *Eucalanus monachus* Giesbrecht, *Corycaeus speciosus* Dana, *Oncaca venusta* Philippi, *Oithona similis* Claus, *Oithona nana* Giesbrecht, *Oithona spinirostris* Claus and *Euterpina acutifrons* (Dana).

DISCUSSION

Polarized light perception and eye structure

In many arthropods, particularly Crustacea, and cephalopod mollusks, where polarized light perception has been best demonstrated, there is always a regular arrangement of two sets of retinal microvilli which comprise the rhabdom, one set being perpendicular to the other (Eguchi and Waterman, 1966; Waterman, 1966c). If the dichroic molecules of visual pigment are oriented so that their major axis of absorbance is parallel to the long axis of the microvilli, then the orthogonal arrangement of microvilli functions as a two-channel retinal analyzer of polarized light (Waterman, 1966b; Waterman and Horch, 1966; Eguchi and Waterman, 1967). Thus the set of reticular cells with the long axes of their microvilli parallel to the plane of polarization will be maximally stimulated by the incoming polarized light, whereas the remaining reticular cells with microvilli having their long axes perpendicular to the plane of polarization will be minimally stimulated. If both sets of microvilli are at a 45° angle to the plane of polarization, then both will be stimulated to the same degree. If the organism compares the stimulation in one set of microvilli to the stimulation in the other set, it has instantaneous information as to its orientation with respect to the plane of polarization.

However, if an organism has only one set of parallel microvilli, it might also orient to polarized light, but it could not get an instantaneous bearing. In this case, the organism would have to compare the stimulation received by this one set of parallel microvilli at one point in time with the stimulation received at another point in time. Thus, if the organism were oriented so that the long axes of the microvilli were parallel to the plane of polarization, the retinal stimulation would be maximal. Then, if the organism tilted or turned a bit, the microvilli would no longer be parallel to the plane of polarization, thus reducing retinal stimulation. The comparison of successive stimulations in time would allow the organism to orient so that retinal stimulation was maximal; in this way the microvilli would always be parallel to the plane of polarization. However, a very real disadvantage to this system is that changing light intensities might not be discriminated from the *e*-vector responses. A decrease in light intensity might confuse the organism if it interpreted such a change as a deviation from its position of orientation to the plane of polarization. One would expect, then, that such a one-channel system is not ideally suited for polarotactic behavior.

The fine structure of the eye of *Cyclops vernalis* clearly shows regions of mutually perpendicular microvilli, suggesting that this geometric arrangement is responsible for the copepod's perception of polarized light. Furthermore, both sets of microvilli are perpendicular to the incoming light as is typically the case

of compound or camera eyes sensitive to polarized light. When the copepod is swimming in either of its usual positions (horizontally or vertically), the laterally pointing dorsal ocelli (Fig. 1) will be oriented so that one set of microvilli is parallel to the naturally horizontal plane of polarization whereas the other set is perpendicular to this plane. Therefore, the vertical and horizontal directions of the microvilli correspond to the vertical and horizontal axes of the copepod's normal spatial orientation, a situation closely similar to that in decapod crustacean compound eyes (Waterman and Horsch, 1966; Eguchi and Waterman, 1967) and in cephalopod camera eyes (Tasaki and Karita, 1966).

Unfortunately, no electron micrographs are available for the other species of copepods studied, but light microscopy by Vaissière (1961) suggests that *Centropages typicus* has parallel rows of microvilli in its naupliar eye whereas *Corycaeus clausi* has no regular arrangement of microvilli, the entire rhabdium being curved so that the microvilli are not parallel. In the experimental studies presented here, *Centropages hamatus* is polarotactic whereas *Corycaeus speciosus* is not. This additional information suggests that polarotactic behavior is correlated with a

TABLE I
Relation of polarotaxis to copepod ecology

Species	Polarotactic	Medium	Habitat	Vertical migration	Food habits	Eye structure
Calanoida						
<i>Labidocera aestiva</i>	Yes	SW ¹	Pelagic*	Yes ^{††}	Omnivore ^{@ @}	Complex
<i>Pontella meadii</i>	Yes	SW	Pelagic*	—	Omnivore ⁺	Complex
<i>Centropages hamatus</i>	Yes	SW	Pelagic*	Yes ^φ	Omnivore ^{@ @}	Complex
<i>Diaptomus shoshone</i>	Yes	FW ¹¹	Pelagic**	Yes ^{φφ}	Predator**	Simple
<i>Pseudodiaptomus coronatus</i>	No	SW	Pelagic*	Yes [#]	Probably herbivore	Simple
<i>Acartia tonsa</i>	No	SW	Pelagic*	Yes ^{# #}	Omnivore ^{@ @}	Simple
<i>Eucalanus monachus</i>	No	SW	Pelagic*	—	Probably herbivore	Simple
Cyclopoida						
<i>Cyclops vernalis</i>	Yes	FW	Littoral†	—	Predator ⁺⁺	Simple
<i>Corycaeus speciosus</i>	No	SW	Pelagic*	Yes ^ε	Predator ⁺	Complex
<i>Oncaea venusta</i>	No	SW	Pelagic*	—	Probably herbivore	Simple
<i>Oithona similis</i>	No	SW	Littoral*	Yes ^{εε}	Predator ^{\$}	Simple
<i>Oithona nana</i>	No	SW	Pelagic*	Yes ^φ	Herbivore ^{SS}	Simple
<i>Oithona spinirostris</i>	No	SW	Pelagic*	Yes [@]	Predator*	Simple
Harpacticoida						
<i>Tisbe furcata</i>	Yes	SW	Littoral*	—	Herbivore ⁻	Simple
<i>Euterpina acutifrons</i>	No	SW	Pelagic*	—	Scavenger [∴]	Simple
Caligoida						
<i>Caligus rapax</i>	Yes	SW	Pelagic*	—	Exoparasite*	Complex

¹ Sea water; ¹¹ fresh water.

*Wilson, 1932; **Brooks, J. L., personal communication; †Pennak, 1966; ††Parker, 1902; ^φHansen, 1951; ^{φφ}Pennak, 1944; ^εGrice, 1957; [#]Schallek, 1942; ^εRoehr and Moore, 1965; ^{εε}Bogorov, 1946; [@]Moore and O'Berry, 1957; ^{@@}Anraku and Omori, 1963; ⁺Personal observation; ⁺⁺Dziuban, 1964; ^{\$}Marshall and Orr, 1966; ^{SS}Murphy, 1923; ⁻Battaglia and Bryan, 1964; [∴]Lebour, 1923.

geometrically regular arrangement of microvilli within the eye, a condition essential to the dichroic rhabdom analyzer mechanism.

Of the copepods studied, five had very elaborate naupliar eyes with cuticular lenses and often with movable parts (Table I). Of these five species with such complex eyes, four gave evidence of perceiving polarized light. Of the eleven species with simple naupliar eyes, only three responded specifically to polarized light. Since polarized light can be perceived by both simple and complex naupliar eyes and since its perception clearly does not depend on lenses or other gross structural modifications, then this again points to the ultrastructure of the eye as the site of polarized light perception. Actually, the tendency for polarized light perception to be more prevalent in forms with complex eyes may be explained by the fact that all these species are predators (Table I).

Polarized light perception by copepods

Although polarized light sensitivity has been reported for cephalopod mollusks with camera eyes (Moody and Parriss, 1960, 1961; Jander, Daumer and Waterman, 1963; Tasaki and Karita, 1966), for arthropods with compound eyes (Baylor and Smith, 1953; Jander and Waterman, 1960; Waterman, 1966a) and for insect larvae (Wellington, 1953, 1955; Waterman, 1966a), mites (Baylor and Smith, 1953) and spiders (Papi, 1955; Waterman, 1966a) with simple ocelli, it has only once been reported for an organism with a naupliar eye exclusively (Umminger, 1968). However, since Stockhammer (1959) was unable to find polarotaxis in copepods of the *Cyclops* group and since Jander (1966) could not demonstrate polarized light sensitivity in the copepods *Macrocylops albidus*, *Cyclops strenuus* and *Labidocera* spp., it appeared that at least some copepods did not exhibit polarotaxis. The present paper confirms definitely that the naupliar eye can perceive polarized light, but that this ability is not to be found universally in copepods.

The failure of Stockhammer and Jander to find polarotaxis in copepods could be due either to organismal or to experimental factors. In the former case, the species investigated actually may not have been able to perceive polarized light. Since the structure of the naupliar eye varies so greatly with the Crustacea as a whole (Elofsson, 1963) and even within the Copepoda (Vaissière, 1961), it is entirely possible that the species examined by Stockhammer and Jander did not possess eyes with the regular two-channel arrangement of microvilli necessary for polarized light perception. This appears to be the reason why *Corycaeus speciosus* showed no polarotaxis in the present paper.

For example, Fahrenbach (1964) has made an ultrastructural study of the eye of *Macrocylops albidus*. The eye contains great masses of microvilli running parallel, but not perpendicular, to one another in a tightly packed array. As previously discussed, such a one-channel system is not ideally suited for polarized light detection and, indeed, Jander (1966) observed no polarotaxis in this species.

Moreover, even if a species did possess the appropriate eye structure and did perceive polarized light, it is entirely possible that the copepods did not behaviorally react to it because they do not use polarized light as a sensory cue in their orientation behavior; *i.e.*, they perceive, but ignore, polarized light.

In addition to such organismal factors, several experimental factors might also explain the failure of Stockhammer and Jander to observe polarotaxis in copepods. Thus the time of observation may have been wrong. That is, there may have been a rhythm in polarotaxis such that observations were made only during the unoriented phase of behavior. This explanation seems likely for those copepods of the *Cyclops* group. If other members of the genus *Cyclops* show a rhythm as does *Cyclops vernalis* (Umminger, 1968), then orientation peaks would appear at dusk, midday and dawn, the times when most investigators are least likely to be observing copepods. At other times, the copepods would be in a transitional state and appear unoriented.

Another experimental factor which may explain the failure to obtain polarotactic responses is as follows: The copepods may have sensed the polarized light, but the correct combination of illumination parameters (absolute intensity, wave-length, angular distribution of intensity, changing intensity, etc.) was not employed to release the corresponding polarotactic behavior.

Ecological significance of polarotaxis

In a previous paper, a polarotactic rhythm in *Cyclops vernalis* occurred in an ecologically meaningful way, implicating it as a behavioral mechanism in vertical migration (Umminger, 1968). In the present paper, however, all the copepods with polarotactic behavior showed no rhythm in their responses. This may be due either to the absolute lack of such rhythms or to the failure of the experiments to provide the correct combination of illumination (or other) parameters needed to release the rhythm. In the experiments with *Cyclops vernalis*, for example, the rhythm was released only in the presence of a uniform angular distribution of light intensity. If such an esoteric cue is needed as a prerequisite for the rhythm in this case, then equally unsuspected conditions may be necessary to release the rhythms in the other copepods with polarotactic behavior.

Moreover, as discussed above, the lack of demonstrated polarotaxis in other species of copepods examined in this paper does not necessarily mean that the copepods in question cannot perceive linearly polarized light, but may indicate the failure of the experimental conditions to elicit a response to it. On present evidence, however, one must assume that orientation rhythms are not present in some copepods with polarotaxis and that other copepods do not have polarotactic behavior.

If one accepts the observed differences in polarotaxis as real, then some clue as to their ecological significance might come from comparing the ecologies of polarotactic and non-polarotactic species (Table I).

Obviously polarotaxis has no relation to habitat since it is found in both salt-water and fresh-water forms and is common in both pelagic and littoral copepods. Furthermore, natural vertical migrations have been reported in forms without polarotaxis as well as in forms with polarotaxis. Therefore, although polarotaxis may be useful for orientation in vertical migration as with *Cyclops vernalis*, polarized light sensitivity may not be necessary for such migrators.

One rather striking similarity among polarotactic copepods is their tendency to be either predatory or omnivorous. Of the seven copepods showing polarized light sensitivity, six are either predators or omnivores. This suggests that the

ability to perceive polarized light might be of greater advantage to a copepod with predatory habits than to one that is strictly herbivorous. This may be related to the findings of Lythgoe and Hemmings (1967) who reported that polarized light sensitivity in clear waters enables distant objects to be more clearly seen and causes contrasts in intensity to be much sharper. Such an improvement in visual ability would seem to be more advantageous to a predator than to an herbivore. One might expect, then, the findings that polarized light sensitivity is most common in predators.

The improved visual ability imparted to predators by polarized light sensitivity does not preclude, however, the additional use of this sensitivity for spatial orientation and migration. The very fact that the predators actively orient to the plane of polarization demonstrates that they do more than passively perceive it; they use it to modify their spatial orientation as well.

This investigation was supported by a National Aeronautics and Space Administration traineeship awarded through the Department of Biology at Yale University. I wish to express my deepest appreciation to Professor Talbot H. Waterman for the stimulation, guidance and critical review of the manuscript that made this study possible. Thanks is also given to Dr. Eisuke Eguchi for his skill and patience in teaching me the techniques of electron microscopy used in this paper. The author is grateful to Dr. John L. Brooks for supplying one of the experimental species. Acknowledgement should also be given to the Director and staff of the Duke University Marine Laboratory for making available to me their facilities where most of the experiments with marine animals were performed.

SUMMARY

1. The ability of *Cyclops vernalis* to perceive polarized light has an ultrastructural basis in the presence of mutually perpendicular microvilli in its naupliar eye.
2. Laboratory investigations with several species of copepods were conducted to determine the extent of polarized light sensitivity in this group. Polarotaxis was not found to be universally present in copepods, but apparently depended on the presence of mutually perpendicular microvilli in the rhabdom of the naupliar eye.
3. Polarotaxis showed no correlation with the presence of a complex, rather than simple, naupliar eye.
4. The ability to perceive polarized light was more prominent in predatory than in herbivorous species, suggesting that its function might be to enable the copepods to discern distant objects more clearly, in addition to its role in orientation during vertical migration.

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THE BIOLOGICAL BULLETIN

PUBLISHED BY THE MARINE BIOLOGICAL LABORATORY

THE LARVAE OF A DISCINID (INARTICULATA, BRACHIOPODA)

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The first account of a discinid larva was given by Mueller (1860). This larva from the sea off Santa Catarina, Brazil, had a pair of thin, approximately circular valves with no hinge, a lophophore with a median tentacle and 4 pairs of cirri, a pair of eyespots, a pair of statocysts, a pedicle, 5 pairs of stout mantle setae and a large number of fine mantle setae. Further specimens (Mueller, 1861), also with 4 pairs of cirri, remained free-swimming for about 5 or 6 days before they became attached to the bottom and sides of glass vessels in the laboratory.

Simroth (1897) reported two inarticulate larvae, one with 3 pairs of cirri and two bunches of extremely fine setae and the other with 4 pairs of cirri and 3 widely separated stout mantle setae. The former, he believed (p. 4), would not develop any more cirri during its pelagic larval life. This belief, the morphological differences between the two larvae and the fact that one was obtained near the surface and the other from a depth of 1000 to 2000 meters, led him to the conclusion that one was adapted to planktonic life at the surface, and the other to life at great depth (p. 6). He further concluded (p. 5) that the larvae belonged to two different species of a single discinid genus.

Blochmann (1898) obtained 10 larvae from the Strait of Rhio in the Pacific identical with Mueller's larvae both in structure and in the stage of development. He considered them as larvae of *Pelagodiscus atlanticus* (King) since the adults of this worldwide species were reported from both the Atlantic and the Pacific. However, in his alum-carmin-stained specimens, he only found nephridia on either side of the stomach where Mueller (1860) had previously seen only statocysts.

Yatsu (1902) described a discinid larva off Japan with a pair of statocysts, 5 pairs of stout setae and a large number of fine mantle setae. He mentioned the possibility (p. 107) that the free-swimming discinid larvae at the stage of 4 pairs of cirri made their way towards the coast shortly before settlement.

Eichler (1911) obtained two large larvae from a depth of 3000 meters at 66 degrees S. Lat. and 90 degrees E. Long. These also had 4 pairs of very long cirri, a pair of statocysts, 4 pairs of stout setae and a great number of fine setae. He also

attributed these to *Pelagodiscus atlanticus*, adding (p. 384) that they had given up pelagic life and were changing over to a sessile mode of life.

Ashworth (1915) found 6 larvae of *Pelagodiscus* from a depth of 40 fathoms in the Indian Ocean. These also had 4 pairs of cirri and led him to believe (p. 66) that the younger stages might live on the sea bottom and hence escaped capture.

In the present study a complete series of larvae was obtained. The smallest was even smaller than the smallest ever reported (Simroth, 1897) and the largest exceeded in size all the largest larvae ever reported, except for the two of Eichler (1911). Moreover, the larvae of medium size in the present series showed the transitional stages between the two larvae described by Simroth (1897), thus serving to provide evidence for the probable identity in species of the two larvae of Simroth. The nature of the organs lying on either side of the stomach was also studied, since these were differently interpreted by Mueller (1860) and Blochmann (1898). The morphological changes during the larval stage were described. The larvae were compared with those previously found and their similarities and differences discussed.

MATERIALS AND METHODS

Thirty-three larvae of a discinid were obtained by surface plankton hauls carried out in the daytime in the shallow water of the Johore Strait off the north coast of Singapore Island. Two of them were caught on August 28, 1952 (Table I, nos. 15 and 17), one each on September 21, 1952 (not in Table), September 28, 1952 (Table I, no. 2) and June 12, 1953 (Table I, no. 1). The others, which were obtained on November 16, 1952, formed a continuous series of transitional developmental stages and are presumed to belong to the same species. Before the adults are discovered in the vicinity of Singapore, these larvae are provisionally identified as those of a discinid on the grounds of the resemblance of some of the stages in the present series to similar larvae reported by Mueller (1860), Simroth (1897) and others.

The shell and setae of each larva were measured with a calibrated ocular micrometer under a monocular microscope. Shell dimensions were taken with the larva lying on its dorsal or ventral valve. The shell length was the shortest distance between the anterior and posterior edges of the shell along the sagittal plane, when both edges were in the same plane of focus. Similarly, the width of the shell was the distance between the right edge and the left edge at the widest part of the shell when both edges were in focus.

The number and type of setae and the number of paired cirri in the larvae were recorded. The absence or presence of statocysts was noted.

All observations were carried out on live specimens in the Department of Zoology, University of Singapore.

RESULTS

A moderately young discinid larva consisted of a dorsoventrally flattened body enclosed between two shell valves. These were slightly convex externally, not hinged but merely held together by muscles and other soft tissues. The visceral cavity occupied a small central part in the posterior half of the space between the

TABLE I
Shell dimension and the type, number and length of setae in discidinid larvae with 4 pairs of cirri from the Johore Strait

Specimen no.	Shell valve				Length in μ of embryonic setae						Type and number of setae*									
	Dorsal		Ventral [†]		No. 1		No. 2		No. 3		No. 4		No. 5		Embryonic		Early larval		Late larval	
	Length μ	Width μ	Length μ	Width μ	Left	Right	Left	Right	Left	Right	Left	Right	Left	Right	Left	Right	Left	Right	Left	Right
1	462	500	349	455											0		1, 1		24, 24	
2	401	479	337	434											0		1, 1		0	
3	382	437	326	407											0		2, 2		0	
4			347	429																
5			346	424																
6	311	364	291	364											0		4, 5		0	
7	311	364	288	356											0		2, 2		0	
8	308	358																		
9	282	330	265	323		1213		1289							0, 2		3, 3		0	
10	271	326	259	318	1198		1213, 1243								2, 1		3, 3		0	
11	253	302	245	297	1107		1228								2, 0		2, 2		0	
12	242	284	239	276	1152, 1198		1167, 1213				917				2, 3		0		0	
13	241	288	243	288	121, 1137		1152, 1122								0		3, 2		0	
14	233	259	219	251											1, 2		2, 2		0	
15	217	242	212	238	773		1228			819					3, 3		(1), (1)		0	
16	216	251	208	245	1092, 1092		1084, 1084			819, 834					4, 4		3, 3		0	
17	216	236	197	232																
18	213	242	207	236	1152, 1152		1289								2, 3		3, 3		0	
19	207	239	199	228	1152, 1122		1228, 1198								3, 3		0		0	
20	184	212	179	204						796, 781					2, 2		3, 3		0	
21	179	199	173	193	1167		1213								0, 2		0		0	
22	179	196			1092, 1107		1182, 1137			788					4, 2		0		0	
23	168	183	156	173	1175, 1152		1175, 1167			872					2, 3		0		0	
24	161	183	160	184	1198, 1213		1243, 1228			773, 773					4, 5		1, 2		0	
25			157	169		1031				803					1, 3		0		0	
26	158	173	147	161	1213, 1182		1243, 1243			788					2, 4		0		0	
27	143	151	130	138	1122, 1137		1228, 1198			879, 879					4, 3		0		0	
28	140	145	130	136						788					2, 1		0, 1		0	
29	124	125	104	111	1137		1228, 1228			819, 849					3, 4		0		0	
30	115	124	112	118	1084		1122, 1152			894, 894					2, 3		0		0	
31	108	115	98	99	1078, 1078		1038			728					3, 2		0		0	
32															4, 1		0		0	

shell valves and was almost filled with the stomach and the intestine (Fig. 1). A cone of tissue projected laterally from the lateral body wall at the level of the stomach. This setal cone contained the bases of the long straight, brittle setae, which will be referred to as the embryonic setae in the present study. This cone also housed the muscles to move these setae.

The rest of the space between the shell valves not occupied by the visceral cavity constituted the mantle cavity. In the front part of this was a contractile columnar projection of the anterior body wall. This column contained the oesophagus,

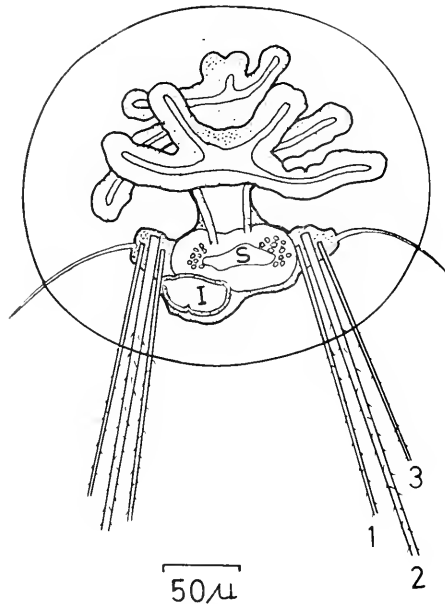


FIGURE 1. Ventral view of a young discinid larva (no. 15, Table) with embryonic setae (1-3) and a curved seta issuing from each setal cone. The stomach (S) and intestine (I) almost fill up the visceral cavity.

pharynx and muscles and continued anteriorly into the lophophore. Muscular movements of the column and lophophore brought about extension and expansion or retraction and contraction of both column and lophophore.

The larval lophophore consisted of a central disk roughly horizontal in position with the more or less centrally placed mouth on the ventral side. At the periphery of the disk 9 lophophoral processes radiated out. One of these, the tentacle, pointed anteriorly in the sagittal plane, and the others, the cirri, were more lateral in position. All the larvae in the present series had 4 pairs of cirri. Each cirrus on one side of the mouth usually corresponded in position and size with the one directly opposite on the other side, thus forming a pair.

A thin layer of mantle tissue lined the inner surface of the mantle cavity. In widely separated regions along the lateral border of the ventral mantle typically 4 stout curved setae appeared on each side of the medium-sized larvae. From each

lateroposterior border of the dorsal mantle appeared another. These 5 curved setae will be referred to as the early larval setae.

In the oldest larvae yet another kind of setae appeared in the lateral border of the mantle. They were shorter and more flexible than the other two types and will be referred to as the late larval setae.

The lengths and widths of both the dorsal and the ventral shell valves were tabulated in Table I, in which the larvae were arranged in descending order of magnitude of the length of the dorsal shell valve wherever possible. Data on the different kinds of setae on the left and the right side of the body were also included.

Both shell valves were transversely oval in shape in the youngest larvae, although occasionally one valve or the other was circular, *e.g.*, the dorsal valve of no. 29 and the ventral valve of no. 31 in Table I. However, while the dorsal shell valve maintained approximately the same oval shape during the entire larval stage, the ventral one changed shape. The convex posterior edge of this in the youngest larvae gradually became less and less curved until it became almost a straight line when the larva attained about $219\ \mu$ in ventral valve length (no. 14). The posterior edge remained straight even at $243\ \mu$ ventral shell-valve length (no. 13), but with further growth it gradually developed a slight but noticeable bay near the median line, when the ventral-valve length reached $265\ \mu$ or so (no. 9). This bay became more prominent with further larval growth to $326\ \mu$ (no. 3) and deeper still in larger larvae.

This change in shape of the ventral shell valve was presumably caused by a slowing down of deposition of shell material at the posterior end. The deceleration was greatest near the sagittal plane, becoming less laterally.

With the exception of larvae nos. 13 and 24, in both of which the two shell valves were equal in both length and width, and of no. 6 in which the two shell valves were equal in width, all the larvae had unequal shell valves. In each of them the dorsal shell valve was slightly longer and wider than the opposing ventral one. This difference was slight throughout this series with the exception of the largest larvae (nos. 1 to 3), in which the difference was considerable.

The external surface of both shell valves was smooth. There was no observable growth line with the exception of larva no. 2. In this there was an oval area, $168\ \mu$ long and $183\ \mu$ wide, which appeared as a prominent growth line on the dorsal shell valve.

In the smallest larva (no. 31 in Table I) the alimentary canal consisted of the mouth, pharynx or oesophagus, stomach, intestine and the anus. In an older larva such as no. 15 the intestine was constricted near the distal end to form another chamber. The highly extensible oesophagus was long and slender when the lophophore was everted during swimming. When this was retracted the inner end of the oesophagus projected into the stomach in the form of a valve-like fold, almost completely closing the opening of the oesophagus into the stomach. The stomach was a large sac with its lateral diameter greater than its anteroposterior one. The high epithelium laterally indicated distinct regions of phagocytosis. The intestine was a thin-walled tube with a distinct constriction separating it from the stomach. It was directed to the right lateroposteriorly in young larvae but either laterally or anterolaterally in older specimens. The anus, not easily distinguishable except during defecation or after, was a slit about 2 to $2.5\ \mu$ in diameter.

Defecation was observed in larva no. 19. It was preceded by a flattening of

the stomach in the anteroposterior direction. The stomach and the intestine then shrank visibly and progressively as particles from the hind part of the intestine escaped through the anus. Meanwhile the constriction between the stomach and the intestine also diminished in diameter. Presumably during defecation muscular contraction of the wall of both the stomach and the intestine was responsible for the expulsion of the faecal material.

Statocysts, absent in young larvae, occurred in the oldest larvae nos. 1 to 3. Each of these had a statocyst situated dorsally in the visceral cavity at each posterolateral corner of the stomach. Each statocyst had a central cavity, in which danced 15 to 25 statoliths. The cilia on the inner epithelium of the statocysts were also observed to stir vigorously.

The pedicle occurred in larva no. 6 and in the larger larvae as an oval vesicle posterior to the stomach. It was absent in younger larvae.

The embryonic setae were the first to appear among the three different kinds of setae found during the larval stage in a discinid. They were observed in young larvae of dorsal shell length of 179μ and under (nos. 21-32 in Table I) before another kind of setae were formed, with the exception of larvae nos. 24 and 28. They were equal in number on the two sides of the body, unless lost by accident or some other cause. The usual number appeared to be 4 per setal cone, although 5 had been observed in one larva (no. 24). In every setal cone the 4 or 5 setae were not of the same length and diameter, but were very regularly arranged in a definite order. The embryonic seta nearest the midline (no. 1 seta in Fig. 1 and Table I) was the second longest and second thickest among the four on each half of the body. The next one, occupying a more lateral position, was invariably the longest and the stoutest (no. 2 in Fig. 1 and Table I). The next one was the shortest and the thinnest (no. 3 in Fig. 1 and Table I). The outermost one (not shown in Fig. 1) was slightly longer and slightly thicker than the one medial to it. In larva no. 24 of Table I a fifth embryonic seta occurred outside the other four. This was the shortest and the finest in the larva.

Short spines were found on all the embryonic setae in approximately four irregular, longitudinal rows starting from a region about halfway between the base of the setae and the posterior edge of the shell (Fig. 1). Those on the coarser setae appeared to be stouter than those on the finer ones. They seemed to occur throughout the length of the setae.

The lengths of the different embryonic setae on both sides of the body were summarized in Table I. Due to the widely differing diameters of the first medial two, they could be distinguished from each other. The third and the fourth setae were, however, not easily distinguishable when removed from the larva. The diameter of each decreased very gradually from base to tip. The color was darker than that of other setae that appeared later. The embryonic setae were stiff and brittle. They broke off when they collided with hard objects on a slide. In older larvae they decreased in number and were absent in the oldest larvae.

The early larval setae appeared in an older larva during its early planktonic life. They occurred singly in widely separated regions along the mantle border. They were curved, stout setae, tapering rapidly in the last quarter or so to a sharp point. The stoutest among them was situated at each posterolateral region of the ventral shell valve and was equipped with stout spines. The maximum number of early larval setae seemed to be five on each side, although this was attained only

on one side of larva no. 6. Of these, four issued from the ventral mantle, and one from the dorsal mantle. Many larvae had only 3 or fewer setae on each side. The stoutest seta was 300 to 350 μ long, the others, only half as long.

Larva no. 15 (Fig. 1) had a curved seta, morphologically indistinguishable from an early larval seta except in length, in diameter and in its point of origin. It was more slender and shorter. It issued not from the mantle border but from the setal cone. Whether it was produced during the embryonic or the larval stage is not known.

The late larval setae were the last to appear in the larva and were found only in the oldest larva (no. 1). In this there were about 24 of them evenly spaced in the lateral region of the mantle between the most anterior and the most posterior early larval setae. Finer and more flexible than these, they tapered gradually to a fine point.

Rotational movements of the dorsal shell valve were observed in a submerged larva lying with its ventral valve motionless at the bottom of a microscope slide. Protrusion and retraction of the lophophore, bending, extension and shortening of the individual cirri occurred now and again.

In swimming the anterior edges of the shell gaped open, the lophophore emerged from the shell and the tentacle and cirri spread out radially. The cilia on both the tentacle and cirri beat vigorously to drag the larva forward with the two bundles of primary setae trailing behind. On collision with another object the cilia immediately stopped beating and the larva sank. During forward progression the flat body regularly rotated about the anteroposterior axis and the two bundles of embryonic setae were sometimes parallel to each other and sometimes pointing outward from each other at an angle. On several occasions each bundle was simultaneously moved medially past each other to form a cross as the larva swam forward. The various positions of these setal bundles indicated a fine control by the muscles in the setal cone. While the animal was at rest, movements of individual embryonic setae in a bundle were also observed.

Movements of the early larval setae were also observed. These were capable of independent movements. The stoutest one on each posterolateral region was especially active and was seen to make intermittent lashing movements in the gape between the two shell valves. Whether these movements were an aid in swimming is not known. Movements of early larval setae increased in intensity and frequency when they collided with objects on the slide under observation in the microscope.

DISCUSSION

The oldest larva in the present study resembled the discinid larvae described by Mueller (1860, 1861), Simroth (1897, Blochmann (1898) and Yatsu (1902). The youngest, however, though resembling the youngest discinid larva of Simroth in many ways, differed from it in the following respects. Firstly, Simroth's larva had 3 pairs of cirri instead of the 4 in the larvae of the present series. Simroth (1897, p. 4) believed that the larva with 3 pairs of cirri was in a stable state and that it would not develop additional cirri during its larval stage. Secondly, the innermost embryonic setae in Simroth's larva were bent. Thirdly, the longest embryonic seta in Simroth's larva no. 1 was the one next to the outermost, whereas in all the larvae examined in this study it was the one next to the innermost.

Presumably not realizing that the two discinid larvae he described could have been two different developmental stages in the ontogeny of a single species, Simroth (1897, p. 5) assigned them to two different species of discinids. However, the larvae in the present study consisted of a continuous series of growth stages, including those that are transitional stages between the two discinid larvae of Simroth. These stages included larvae that possessed both the embryonic and the early larval setae. The loss of the embryonic setae occurred in older larvae, thus transforming them into the more familiar so-called "Mueller's larvae." Obviously the larvae in the present study should be attributed to one single species of discinid. Their adults still remain to be discovered in the vicinity of Singapore. However, Blochmann (1898) relegated the 10 larvae from the Rho Strait and those of Mueller (1860, 1861) to *Pelagodiscus atlanticus* (King).

The succession of three different types of setae in the brief span of discinid planktonic larval stage is a unique phenomenon unknown in brachiopod ontogeny. The present study showed the presence of two bunches of long embryonic setae on the lateral cones of the body wall in the youngest larvae in which the cirri were just mere short lobes of the lophophore. These lobes or short cirri of the lophophore were similar in appearance to those of the youngest planktonic larvae of *Lingula anatina* (Lamarck) the present author had observed and also to those of embryos of *Lingula* at the end of the embryonic period (Yatsu, 1902). Since the two bunches of embryonic setae were already present in the youngest larvae obtained, and also similar in length, width and appearance to those of older larvae, they were presumably embryonic structures elaborated from the substance stored in the fertilized ovum. The present work also showed the first appearance of the early larval setae in larvae of about 140 μ dorsal-valve length. These were obviously not normally present in younger larvae. These setae are called early larval setae, since they developed in early larval life. Another kind of setae, called late larval setae, appeared only in the oldest larva in the present study and in the larvae described by Mueller, Simroth, Blochmann, Yatsu, Eichler and Ashworth. These late larval setae, formed in late planktonic life of the larvae, prepared them for settlement to the sessile life of the postlarva. They reminded one of the setae that appeared near the end of the planktonic larval period of *Lingula* (Yatsu, 1902, Chuang, 1959). Simroth (1897, p. 15) regarded both the embryonic setae in his smaller larva and the early larval setae in his other larva as flotation aids. Presumably these two types of setae could also serve a protective function by making the larvae into less easily swallowed morsels. Moreover, they could be used in steering while afloat or drifting in a current. The rapid succession of three different types of setae is presumably an indication of their great necessity to these larvae. Mueller (1861) reported the resorption of the stoutest early larval setae and the loss of others in his discinid larvae that settled down to a sessile life in the laboratory. The present study (Table I) showed a gradual loss of not only the early larval setae in older larvae, thus confirming Mueller's observations, but also of the embryonic setae.

Simroth's larva no. 1 of 220 μ greatest diameter (1897, p. 3 and Tafel I, Fig. 1) was equivalent in size to no. 20 (Table I) of the present study, but in the development of setae it was in the same stage as larvae nos. 21 to 32 with the exception of nos. 24 and 28, which apparently had an earlier development of the early larval setae. His larva no. 2 of 420 μ diameter (p. 5 and Tafel I, Fig. 2) was equivalent in shell size to no. 5 but in setal development to no. 1 of the present study. Thus

Simiroth's no. 1 was a discinid larva before the appearance of the early larval setae, while his no. 2 could well be an older larval stage of the same discinid species after the loss of the embryonic setae and the appearance of the next two types of larval setae, contrary to his (1897, p. 5) conclusion that the two larvae belonged to two different species of the same genus.

During the larval stage the shell valves changed in shape and proportion. In both shell valves there was a change from an almost circular plate of the youngest larvae to a pronounced oval with a greater transverse diameter. Next there was an ever-widening difference between the transverse and the longitudinal diameters of each shell valve during the last part of the larval life. Meanwhile the dorsal shell valve also increased in diameter much faster than the ventral. Finally there was a striking but gradual change in the posterior edge of the ventral shell valve from a convex to a concave one. This was presumably brought about by a finely adjusted modification of shell growth along the medial third of the posterior mantle edge. The formation of a bay at the posterior margin of the ventral shell valve, just as the formation of the pedicle, is a preparation for settlement to a sessile life. It is through this bay presumably that the pedicle is extruded for attachment to the substratum at the time of settlement.

Mueller (1860) reported the presence of a pair of statocysts in his discinid larvae. Although Blochmann (1898) denied their presence, they were observed by Yatsu (1902), Eichler (1911), Ashworth (1915) and the present writer.

A pair of eyespots, first reported by Mueller (1860), occurred only in the larvae found by Ashworth (1915). They were not observed by other writers.

If the sequence of appearance of organs during ontogeny can serve as an indication of their usefulness, then the pedicle, the statocysts and the late larval setae, which appeared towards the end of the larval stage, are presumably of little functional value to the larvae during the planktonic stage. They are presumably of use during settlement and the sessile life that follows.

In the ontogeny of *Glottidia* (Brooks, 1879) and *Lingula* (Yatsu, 1902; Chuang, 1959) shell size, as well as the number of pairs of cirri, increases during the planktonic larval life and either of them can be a fair index of the stage of larval development. In the ontogeny of a discinid, Table I showed that only shell size increased while the number of cirri remained constant at 4 pairs during the entire larval stage. Hence only shell size of a discinid larva could indicate the stage of development. Discinid larvae with 4 pairs of cirri need not necessarily be at the same stage of development, contrary to the belief of previous writers. In addition to shell size, the occurrence of organs or structures that develop at a definite stage during ontogeny may be a useful index of the age of the larvae. Such organs or structures include the different types of setae, the statocysts, the pedicle, the bay or notch on the ventral valve and the nephridia.

Mueller (1860, p. 79) rightly believed that his larvae were near the end of their larval life and on the way to adopting a sessile life. The absence of younger stages, he thought, was an indication that these had up till then remained in the shell of the adult. Ashworth (1915, p. 66), on the other hand, believed that the younger stages might live on the sea bottom and thus escaped capture. The occurrence of a series of developmental stages including the youngest larvae in the surface plankton of shallow water seems to indicate that the young stages can occur near the surface. The capture of these young and old stages near the shore also indicates

the random movements of both young and old to or away from shore, contrary to the belief of Yatsu (1902, p. 107) that the larvae shortly before settlement made their way towards the coast.

Discinid larvae were obtained from February to April (Mueller, 1860, 1861) in the South Atlantic Ocean, from July to August or September (Yatsu, 1902), off Japan in the Northern Pacific and in October (Ashworth, 1915) in the northern part of the Indian Ocean. They were captured in the tropical West Pacific in July (Blochmann, 1898) and in June, August, September and November by the present author. This indicates the restriction of breeding to the summer and autumn. More intensive collection may possibly extend the breeding season in tropical waters.

SUMMARY

1. Thirty-three discinid larvae were obtained one meter below the water surface in the daytime in the shallow part of the Johore Strait, north of Singapore.

2. The oldest among them, 462μ in longitudinal diameter, resembled the "Mueller's larva" discovered by Mueller and referred to *Pelagodiscus atlanticus* (King) by many subsequent authors.

3. The youngest among them, 108μ long and 115μ wide in dorsal shell, resembled Simroth's smaller discinid larva of 220μ diameter in many essential features.

4. The larvae in this study showed a gradual transition in many morphological characters, indicating that they were a series of developmental stages in the ontogeny of a single discinid species.

5. The two bunches of long embryonic setae were gradually replaced by the curved early larval setae in the medium-sized specimens. Towards the close of the larval stage the third kind of setae appeared from the mantle border.

6. The dorsal shell valve maintained its oval shape during the entire larval stage. The posterior edge of the ventral one, however, gradually changed from convex to concave, passing through a stage when it was almost a straight line.

7. The statocysts and the pedicle appeared in older larvae.

8. The number of cirri remained constant throughout the larval stage.

9. The larvae were compared with the discinid larvae previously recorded. Their developmental changes and the breeding season of discinids were discussed.

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REPRODUCTIVE BEHAVIOR IN THE SPIDER CRAB, *LIBINIA EMARGINATA* (L.)¹

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Crustaceans have complex behavior patterns. Among these are patterns concerned with reproduction and more specifically mating (reviewed by Schöne, 1961). Mating behavior patterns vary among different species and in a number depend upon the physiological condition of the female for their initiation. For example, in some species, mating is correlated with the pre-adult molt of the female and the transferred sperm are stored in the seminal receptacles of the female. Among the portunids, the male protects the pre-molt female and mating occurs immediately after she molts. This behavior is initiated by a pheromone released into the water by the pre-molt female (Ryan, 1966). In some brachyurans (*e.g.*, *Callinectes*, *Carcinus*), mating is apparently dependent on a condition of ripeness of the ovary (Chidester, 1911; Broekhuysen, 1937). Among Majidae, Schöne (1968) reported that *Maja verrucosa* males protect the females beneath them prior to mating. In some brachyurans submissiveness of the female when seized is necessary for mating to occur (Chidester, 1911). Finally, the mating posture varies among species.

The initiation and probably other aspects of the mating behavior in these crustaceans evidently require rather specific communications between potential mates. In land forms this is often visual and auditory (Salmon and Atsrides, 1968) while in aquatic forms it may also be chemotactic. Chemotactic communication would simply involve one or more pheromones as in the one clear-cut example described by Ryan (1966). In view of the interest in mechanisms of communication in Crustacea as well as other organisms it seemed of interest to report observations on mating behavior of the spider crab, *Libinia emarginata* (L.), and to describe for the first time a heretofore unrecognized male-female association, here called "obstetrical behavior."

MATERIAL AND METHODS

From June to early September, 1967, male and female *Libinia emarginata* obtained from the Marine Biological Laboratory Supply Department, were kept in an aquarium (approximately 3' × 5' × 9") through which sea water constantly flowed. This arrangement was initially selected to provide storage for the crabs for another purpose. However, as specific behavior patterns began to be recognized, immature animals were removed and only mature animals (5-6 males and 8-14 females) were

¹ Contribution No. 090 from the Institute of Molecular Evolution. This investigation was supported in part by NIH grant 5T1 HD26-06, Fertilization and Gamete Physiology Training Program and by A5429 from the Institute of Arthritis and Metabolic Diseases. The author wishes to thank Mrs. Patricia Golmont for the drawings which were made from specimens and selected movie frames.

retained. Routine feeding (*Mytilus* and *Spisula*) was initiated, females were marked with insoluble dyes for individual identification, and their brood chambers were examined periodically. The crabs were observed for reproductive behavior at intervals during most of the day and evening.

OBSERVATIONS

Ovigerous females were observed from early June until early September. The females are apparently able to produce at least 3-4 consecutive broods during the period studied. In these crabs spermatozoa enclosed in spermatophores are transferred to the seminal receptacle of the female at mating. At the time of oviposition,

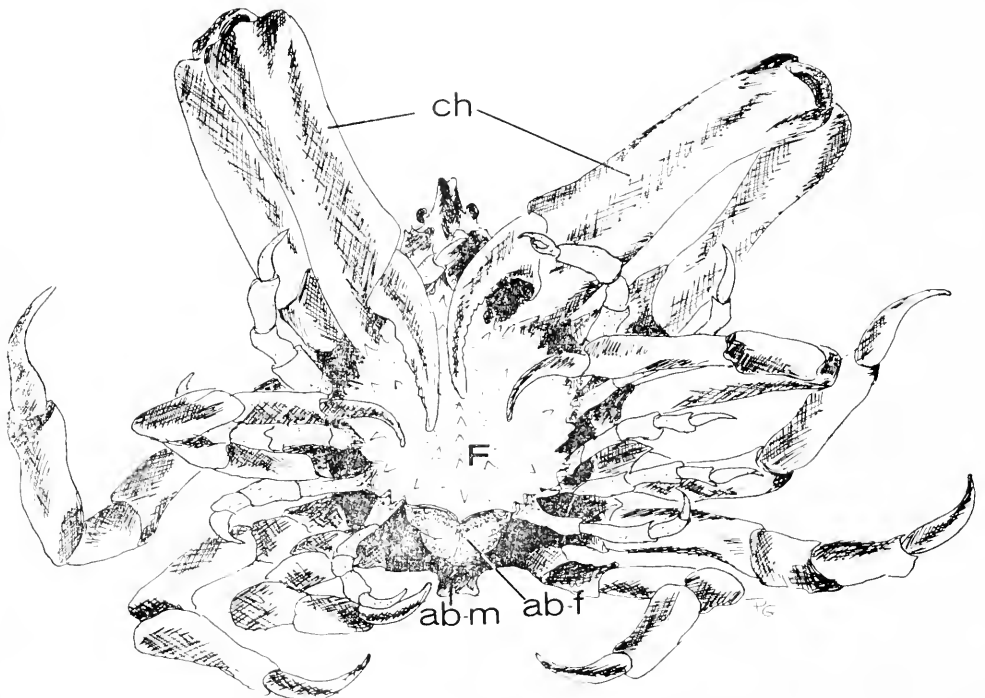


FIGURE 1. Mating position. Chelae (ch) of the male are bent as he holds the female (F) in the rostral region and rotates onto his carapace. Abdomens of male (ab-m) and female (ab-f) are extended.

the eggs pass the seminal receptacle where they can interact with spermatozoa, and are then discharged into the "brood chamber" where the egg mass becomes attached to the pleopods. The last known date of oviposition in this study was September 1. As has been reported by Costello *et al.* (1957), the eggs at the time of oviposition are a bright orange-red in color. The color changes from orange red to brown as development proceeds in the egg mass. Zoeae were apparent within the egg chorion a few days prior to hatching. In the case of at least six marked females development to the swimming zoea was completed in 25 days. In nine of thirteen

recorded cases, the females had completed oviposition of a new brood in less than 12 hours after releasing zoeae. In four other cases, oviposition occurred after more than 12 hours following zoea release.

Two definite and distinct types of behavior were associated with the release of the zoeae, *i.e.*, mating and obstetrical behavior.

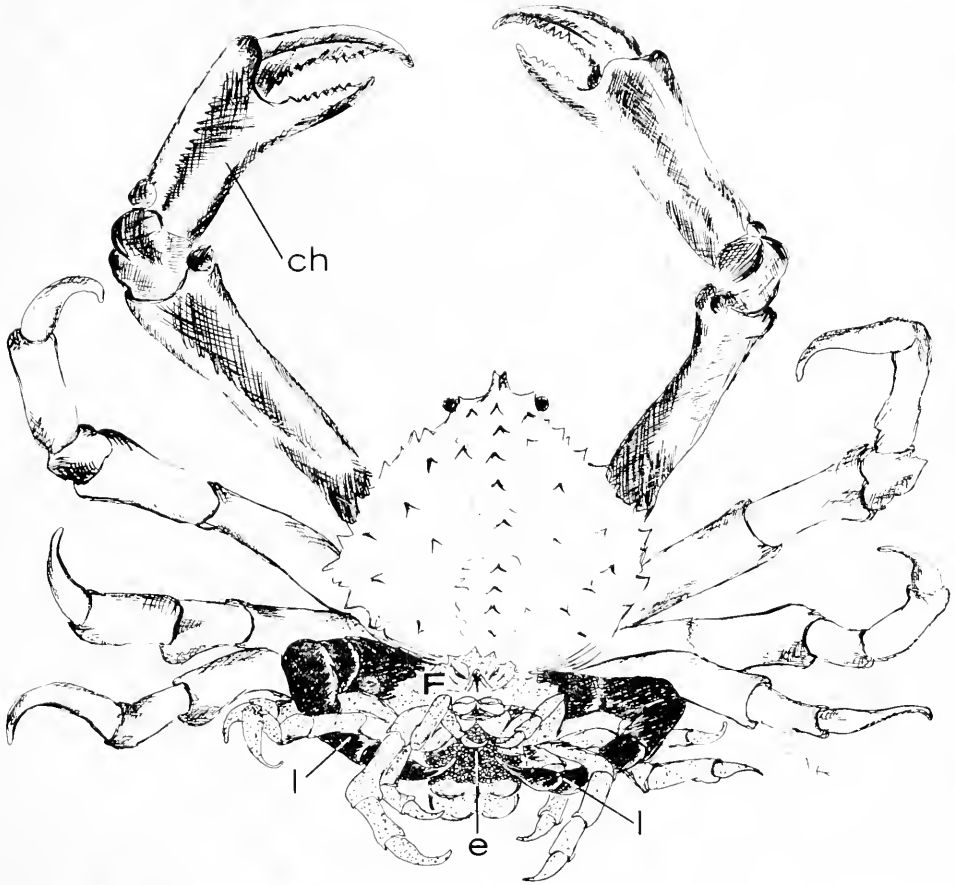


FIGURE 2. Obstetrical position. Female (F) held at right angles to body of the male. The fifth pair of walking legs of the male (l) are inserted into the brood pouch filled with eggs (e) containing zoeae about to hatch. Chelae (ch) of the male are extended at this time.

Mating

In the aquarium containing females about to release zoeae, the males can be observed walking about on the tips of their dactyls. They appear to be actively searching. Encounters at this time with non-gravid females or females which have either just released their larvae or are about to release larvae generally result in mating. Females with eggs in the early stages of development were never observed to mate.

In six recorded cases, the males first grasped the legs of the female with his chelae and positioned her beneath him. His chelae were bent in the manner shown in Figure 1 as he grasped the female in the rostral region. The abdomen of the female was pulled down and bent back at the same time the male's abdomen was lowered. Copulation then began and the male frequently rotated over onto his carapace holding the female above him. Once copulation had been completed, the male resumed a walking position, released his hold on the head of the female, once again grasped her by the legs and released her.

Obstetrical behavior

Immediately preceding the release of zoeae by a female, males were seen actively searching. Males were very aggressive at this time and frequently fought with each other. In sixteen recorded cases, when a female about to release zoeae was encountered, the male captured her and placed her beneath him. Two males might fight for possession of a female and sometimes lose her to a third male. Eventually, a male grasped the female by her legs with his chelae, moved her posteriorly, and positioned her body at right angles to his own. The tips of his fifth walking legs were inserted beneath the abdominal flap to hold her in this position (Fig. 2). The male then frequently backed into a protected area and stood with chelae outstretched. In this position he would fight with other mature males or drive away immature males or females. While being held in this position, the female fanned her abdomen and zoeae were released. Following zoea release she cleaned the remaining egg cases from her pleopods with her chelae. When she had finished the cleaning operation the male released her. Juvenile males and females as well as females with broods at early stages of development were never held in such a position by the male.

DISCUSSION

Libinia emarginata is abundant in coastal waters of the north Atlantic. Bumpus (1898a, b) reported the collection of breeding animals from May through early August while Thompson (1899) collected larvae as late as September 4. Costello *et al.* (1957) stated that development from oviposition to release of larvae takes about one month. In the present study, development was completed in 25 days and the last observed date of oviposition was September 1.

In *Libinia emarginata* there seems to be little sexual display other than the searching walk of the male. Mating position seems similar to that observed in other brachyurans (see review, Schöne, 1961). Although copulation occurs only in newly molted females in many crustaceans, mating in adult *Libinia* occurs between broods and is not associated with molting. Whether the female first mates at the time of the final molt to adulthood is not known. Mating between broods is apparently not needed to produce viable embryos. In *Menippe mercenaria* many broods are produced between intermolts and the sperm are carried over at molting (Cheng, 1968). In *Libinia emarginata* females have been observed to mate with more than one male, suggesting that storage of sperm from several males is possible.

Chemical or mechanical stimuli have been implicated in reproductive behavior

in many arthropods. The presence of a ^{mobile}pheromone in the portunids has been established (Ryan, 1966). In *Libinia*, the predictability of the onset of mating and obstetrical behavior, the active searching by the males, the increased fighting among males and the selectivity of the animals to be placed in the obstetrical position suggests that the behavior patterns are initiated by some chemotactic substance, i.e., pheromone(s). Direct experimental evidence for the presence of a pheromone has not yet been obtained. Submissiveness on the part of the female in *Libinia emarginata* may be necessary for mating and for the obstetrical behavior.

Assuming the action of a pheromone(s), it must either come from the female or be released by or with the zoeae at hatching. Production of such a substance and initiation of the behavior patterns seem to be independent of time of day since they occurred at all times of day and night although with greater frequency during the evening hours. In the laboratory situation this could be influenced by artificial conditions such as the fluorescent lighting or crowding. However, predictability, selectivity of participants and the fact that behavioral patterns are not interrupted by feeding or grooming activities suggest that this behavior may occur in nature.

What evolutionary advantages are there in the development of such behavioral patterns? The obstetrical behavior may have evolved to protect the female as the zoeae are being released although females are able to release zoeae without the presence of a male. In *Maja*, the male protects the female beneath him while the zoeae are released, (Schöne, 1968). *Libinia emarginata* males have been observed standing over and protecting females which had only two or three legs. They seemed unable to place such females in the obstetrical position. The distinctiveness of the behavior in *Libinia* may in part be accounted for by the great differences in size between male and female.

Among Majidae, both *Libinia emarginata* and *Maja* have a new egg mass in the brood pouch a short time after zoea release. In *Maja*, the crabs mate prior to zoea release. In *Libinia* it seems possible for a female to produce a new egg mass without mating again although mating frequently occurred either before or after zoea release. Production of a pheromone at this time may insure mating before the next brood. Among Majidae where no further molts occur after the adult stage has been attained, mating at the time of zoea release may insure carryover of sperm from one season to the next.

SUMMARY

1. Data concerning reproductive behavior and development in the spider crab, *Libinia emarginata* (L.) are recorded. Ovigerous females were collected from early June to early September. Females are apparently able to produce at least 3-4 consecutive broods of 25 days development each during a breeding season.

2. Two distinct reproductive behavior patterns are associated with the release of zoea on the 25th day of development, i.e., mating and obstetrical behavior. These male-female associations are stereotyped behaviors. The position of the bodies of the male and female differ in the two behavior patterns.

3. The predictability of onset of these behaviors and the specificity of the patterns suggest their possible initiation by pheromone(s).

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THE OCCURRENCE OF DINOFLAGELLATE LUMINESCENCE AT WOODS HOLE¹

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The use of sensitive photomultiplier radiometers has brought about recognition of the very widespread occurrence of bioluminescence in the oceans. Flashing has been recorded in nearly all parts of the oceans and in almost all locations studied (Clarke and Wertheim, 1956; Clarke and Kelly, 1965). At night using proper equipment luminescence may be recorded continuously in the upper layers of the ocean, usually within the euphotic zone and above the thermocline (Clarke and Kelly, 1965; Yentsch, Backus and Wing, 1964). However, the organisms responsible for the near-surface luminescence have been definitely identified only in exceptional circumstances such as during unialgal blooms (Harvey, 1952) or in very eutrophic embayments such as Phosphorescent Bay in Puerto Rico (Clarke and Breslau, 1960; Seliger *et al.*, 1962).

Many authors have suggested but not shown that the organisms responsible for this common near-surface luminescence are the dinoflagellates, very common phytoplankters with carbohydrate cell walls and two flagella, one encircling the cell at the midline and the other directed posteriorly (Clarke and Kelly, 1965; Harvey, 1952; Kelly and Katona, 1966; Schiller, 1937). The work reported here is one of three studies conducted under diverse conditions and at diverse locations to determine the relationship between the occurrence of luminescent dinoflagellates and the common oceanic bioluminescence.

In the work described here the dinoflagellates and bioluminescence present in Woods Hole Harbor between November, 1965, and November, 1966, were studied. The species capable of luminescence were identified, their concentrations in the water were measured throughout the year, and the amounts of luminescence in the water were determined at least once a week by testing samples brought into the laboratory darkroom. The populations were also tested for susceptibility to light inhibition of luminescence and for the presence of an endogenous diurnal control of the amount of luminescence produced.

In a preliminary study made during August and September, 1965 (Kelly and Katona, 1966), water was brought into the laboratory at Woods Hole Oceanographic Institution and the luminescence in the water was measured. The species of armored dinoflagellates capable of luminescence were identified and although cell concentrations were not measured, these organisms appeared to be present in sufficient numbers to explain the luminescence in the water. In addition, the popula-

¹ Presented as part of a doctoral thesis at Harvard University. Contribution no. 1900 from Woods Hole Oceanographic Institution. Research supported in part by National Science Foundation grant 2435.

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tions displayed an endogenous rhythm in luminescence and in sensitivity to light inhibition of luminescence. It became apparent that a study over an extended period of time at one location was required to show if luminescent dinoflagellates are usually present in sufficient numbers to explain the amounts of luminescence recorded in the water, and whether endogenous diurnal rhythms of luminescence are present throughout the year.

LUMINESCENCE OF ISOLATED CELLS

Individual dinoflagellates were tested to determine which species were capable of contributing to the luminescence recorded in whole water samples. Cells were isolated by micropipette into 0.5 ml. of water and stimulated in front of a photo-

TABLE I

Dinoflagellates examined for flashing. Numbers refer to months in which tested, from 8, 65 to 8, 66. One to ten specimens were tested on each date given

Species	Luminescent		Fluorescent		Photo-synthetic pigments
	yes	no	yes	no	
<i>Ceratium fusus</i>	8, 65; 9; 10				always
<i>C. lineatum</i>		9; 10; 11; 12; 1		12	always
<i>C. tripos</i>		9; 10; 11			always
<i>Glenodinium lenticula</i>		9; 10; 11; 3			never
<i>Gonyaulax digitale</i>	9; 8, 66				always
<i>G. spinifera</i>	8, 65				always
<i>Gymnodinium splendens</i>		10			always
<i>Peridinium</i> spp.	varies				varies
<i>P. claudicans</i>	8; 10	12; 8, 66		12; 8, 66	never
<i>P. conicum</i>	6; 10		9, 66		never
<i>P. depressum</i>	4				never
<i>P. divergens</i>	11		12		never
<i>P. globulus</i>	10; 11				never
<i>P. granii</i>					?
<i>P. leonis</i>	9; 10; 11; 8, 66				never
<i>P. mite</i>		8, 65			?
<i>P. oceanicum</i>	9; 10				never
<i>P. oblongum</i>		8, 66		8, 66	never
<i>P. pallidum</i>	9; 12		12; 8, 66		never
<i>P. pellucidum</i>		4			never
<i>P. pentagonum</i>	8, 66		8, 66		never
<i>P. punctulatum</i>	9; 8, 66		8, 66		never
<i>P. subinermis</i>	11; 12; 1; 2; 3		2		never

multiplier radiometer. Methods were described in detail by Kelly and Katona (1966). The results are given in Table I. Kelly and Katona (1966) and Sweeney (1963) listed other dinoflagellate species known to be luminescent. All species showed similar intensity (approximately 10^{-4} μ V per cm.² at 10 cm. from the sensing surface) and duration of flashing. Occasionally a prolonged flash was recorded, lasting as long as 3-4 seconds, and the specimen could never be recovered afterward, probably because it splashed against and adhered to the side of the

sample tube. Usually, however the flash duration did not exceed 10 milliseconds. Specimens of *Peridinium subinermis* and *Peridinium pallidum* remained alive and capable of flashing for as long as 7 to 21 days while isolated in the dark in test tubes without any food other than what little might be in the 0.5-ml. Millipore-filtered sea water that contained them. None of the cells divided, although culture medium and microalgal suspensions were added to several of the samples.

Some of the species were tested for UV fluorescence under an American Optical Co. fluorescence microscope (see Table I). Long-wavelength UV produced only the typical ruby-colored fluorescence of chlorophyll-*a* (in diatoms and *Ceratium lineatum*, the only photosynthetic dinoflagellate examined). Short-wavelength UV produced a continuous blue-green fluorescence in all the luminescent dinoflagellates examined. Fluorescence originated in cytoplasmic particles ranging in size from less than 0.001 μ m. to 0.003 μ m. Nine species (3 or more specimens of each) were examined in this way (see Table I). All specimens of the 6 luminescent species showed the blue-green fluorescence while the 3 non-luminescent species did not. The long duration of the fluorescence indicated that the organisms were not merely stimulated to luminesce by the UV irradiation.

The fluorescence passed through a blue-green 480 nm. wavelength interference filter without appreciable visible attenuation but was attenuated by 470 and 490 nm. filters. The fluorescence therefore peaks quite sharply at or near 480 nm., which is the luminescence peak for several species. Eckert (1966) reported the "microsources" of luminescence in *Noctiluca scintillans* fluoresced in this range.

It seems likely that the blue-green fluorescence is an indication of luminescent capability. The peak at 480 nm. suggests that the fluorescence is due to the chemical that is the source of luminescence. This is not surprising since the luminescent compound must be easily excited to an active state and it should be expected that light output from this excited molecule would have similar spectral composition whether the excited state is produced by an enzymatic reaction or by UV irradiation. It also seems that luminescence probably originates in the fluorescent cytoplasmic particles, as it does in *Noctiluca scintillans*. Several other organisms such as copepods (Clarke *et al.*, 1962) and *Cypridina* (Harvey, 1952) have fluorescent luminescent organs.

The bioluminescence of several dinoflagellate species merits further discussion: *Ceratium fusus* is notable for being the only species of *Ceratium* that has been definitely shown to be luminescent (Sweeney, 1963; Kelly and Katona, 1966). Nordli (1957) maintained luminescent cultures of this species, but they were lost and he was unable to culture luminescent forms again. In the present work few of the cells tested were luminescent. This suggests that only certain cells are luminescent and that luminescence is an inherited characteristic. An inheritable ability for luminescence is also suggested by the existence of both luminescent and non-luminescent clonal strains of *Noctiluca scintillans*. *Ceratium tripos* has been reported by several authors to be luminescent (Harvey, 1952) but it was not so in Woods Hole and Nordli's (1957) cultures of this species were not luminescent. This places the earlier reports in doubt, although there is the possibility that both luminescent and non-luminescent forms exist as in *Ceratium fusus*. The luminescence of *Peridinium claudicans* was also variable; it was luminescent during certain months and not others in Woods Hole. The taxonomy of *P. claudicans*, however, is confused and more than one species may have been included under this name.

TABLE II
Summary of flashing rates (flashes per 30 seconds) and luminescent cell concentrations (cells per liter) during 1965-1966 at Woods Hole

	Dec.			Jan.			Feb.			Mar.			Apr.			
	7	15	21	27	3	12	25	31	8	15	23	2	16	23	29	6
1. Mean	160	76	100	76	84	111	190	223	117	189	110	176	144	149	146	152
2. Maximum	179	94	128	94	109	186	208	241	159	238	181	195	198	162	165	185
3. Minimum	141	56	66	65	61	163	200	81	138	74	160	58	137	130	121	144
4. Mean, 1100-1400 hrs.	169	74	117	77	97	76	177	221	99	164	96	189	—	143	146	134
5. Mean, 2000-2300 hrs.	158	89	79	79	74	158	199	215	148	197	138	177	167	156	136	173
6. Luminescent cells	71	68	42	21	54	141	243	892	602	993	446	640	237	118	157	52

	Apr.			May			June			July			Aug.			Sept.			Oct.			
	12	20	27	4	10	17	31	7	16	21	28	5	13	16	27	3	13	30	7	13	28	11
1. 179	94	150	208	107	112	152	136	66	8	8	31	19	50	2	10	41	25	34	83	217	199	137
2. 196	117	208	208	107	152	205	164	84	18	18	40	26	78	5	17	87	41	45	104	260	206	177
3. 78	78	115	115	84	93	119	81	49	44	1	10	15	32	0	5	12	3	8	62	201	189	90
4. 175	98	180	180	98	135	173	—	70	66	—	34	18	38	3	16	67	33	—	85	196	199	—
5. 194	93	124	124	98	97	—	157	64	66	—	40	22	29	—	9	18	7	42	89	250	—	158
6. 70	23	35	35	26	51	11	45	13	12	1	2	8	40	0	40	172	160	292	20	33	29	54

VARIATION OF LUMINESCENCE IN WATER SAMPLES

Fifteen-liter surface water samples were brought into the laboratory darkroom from near the harbor dock of the Woods Hole Oceanographic Institution. Dates of sampling are shown in Figure 1 and are given in Kelly (1968); luminescence stimulated by bubbling air through the water was measured about every week throughout the year. The method and equipment were described by Kelly and Katona (1966). All measurements were performed in a darkroom at sea-surface temperature. For consistency the equipment and procedures were carefully kept the same throughout the entire program. Samples were collected after dark the day before measurements commenced around midday. Measurements were taken every hour to detect any endogenous diurnal rhythms that might have been present (see Kelly and Katona, 1966). Observations were also made on the effects of light inhibition of luminescence. They are described below.

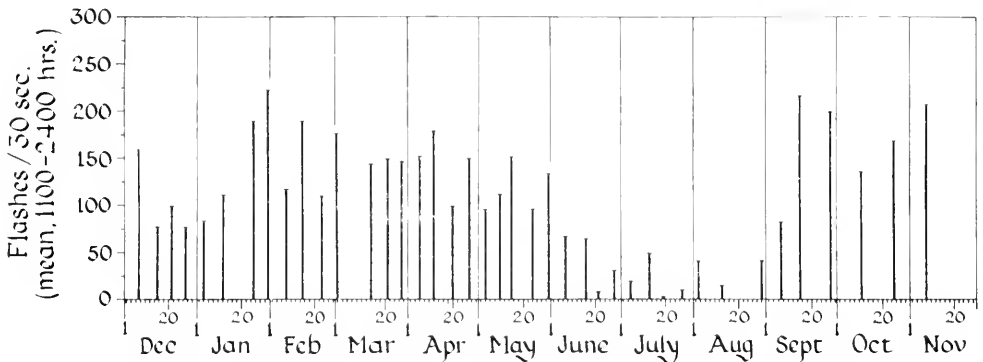


FIGURE 1. Bioluminescence recorded in water samples from Woods Hole Harbor during 1965 and 1966.

The amounts of luminescence found throughout the year are shown in Table II and Figure 1. These are compared below to the numbers of luminescent cells found in the water. The detailed measurements were too numerous to report here but are given by Kelly (1968). Surprisingly no significant endogenous rhythms were found. As shown in Table II, the averages of the flashing rates during midday and after dark were similar: 139.6 and 136.8 flashes per 30 seconds, respectively, through the end of May. These averages suggest a lack of any significant endogenous day-night rhythms in flashing. There was usually some variation within any day's observations as shown in the examples in Figure 2. However, the variation was usually random, and it has been impossible to determine any cause for it. There were increases in flashing rates after dark on the 12th and 25th of January and the 8th and 23rd of February, but these were probably fortuitous since they were not repeatable. Figure 2 shows typical daily measurements from November through May plotted against time of day. The curve marked by \blacktriangle (from Kelly and Katona, 1966) is typical of the endogenous rhythm found in August and September, 1965. The one marked by \triangle shows the measurements of January 12, 1966, which were the most suggestive of an endogenous rhythm during 1966. The one shown by \circ , taken January 25, 1966, is more typical and

shows no rhythm. The last curve, ●, is a plot of the means and its lack of variation indicates that hourly variations on any given date were random.

The endogenous rhythms found by Kelly and Katona (1966) in August, 1965, were thus not present during the rest of the year. The genus *Gonyaulax* was dominant during the August, 1965, studies, but luminescent forms of the genus were not often found during the rest of the year. Also, *Gonyaulax* is the only genus reported to have an endogenous luminescence rhythm in culture. It would

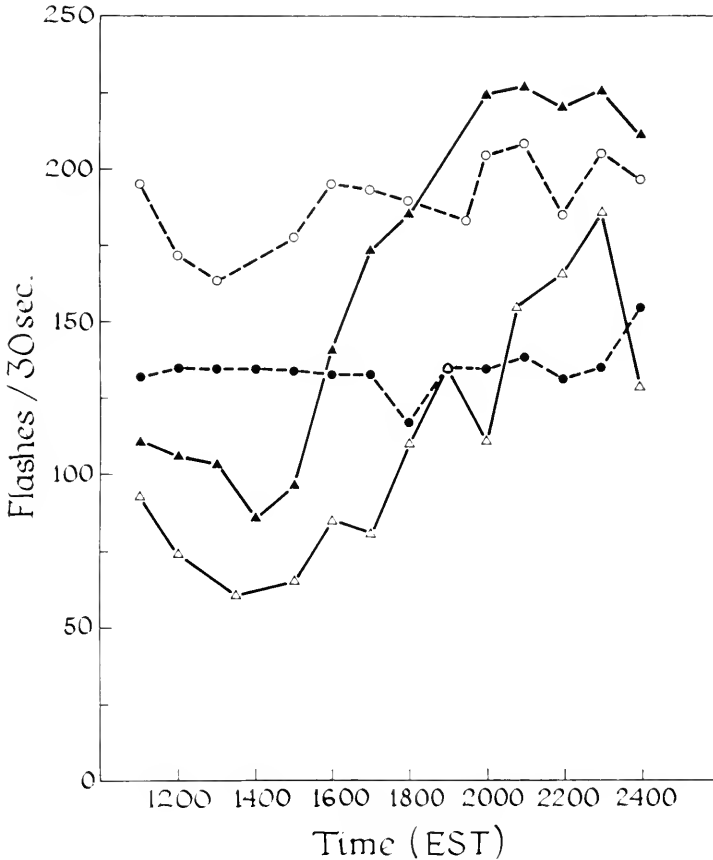


FIGURE 2. Luminescent flashing in samples of water from Woods Hole Harbor measured from noon to midnight on various dates. See text for explanation.

appear therefore that endogenous diurnal rhythms of luminescence are uncommon insofar as the situation in Woods Hole is typical.

Numerous observations were made that indicated dinoflagellate flashing is nearly always inhibited by light. This would explain the daytime decreases in luminescence usually recorded in the natural environment (Clarke and Kelly, 1965; Kelly and Katona, 1966). Kelly and Katona (1966) found light inhibition of flashing as well as an endogenous diurnal rhythm in sensitivity to that inhibition

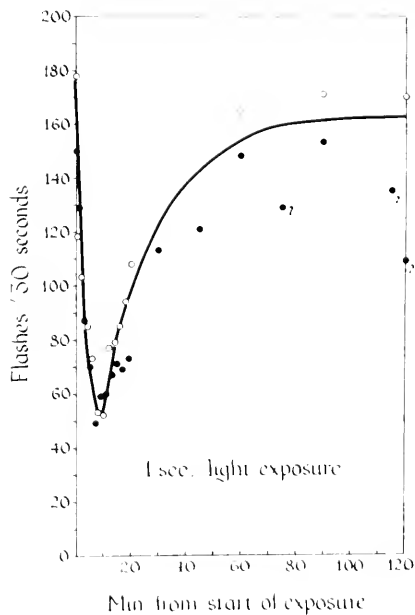
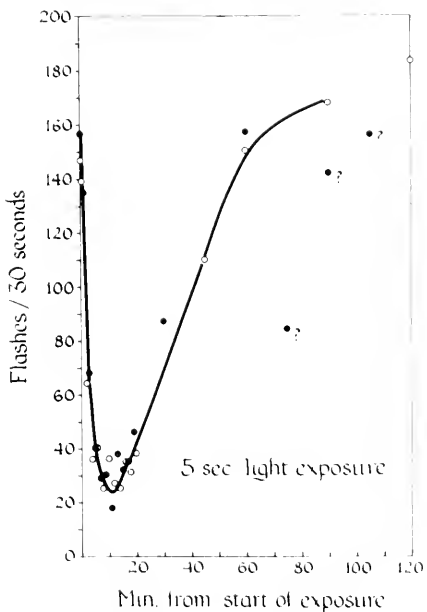
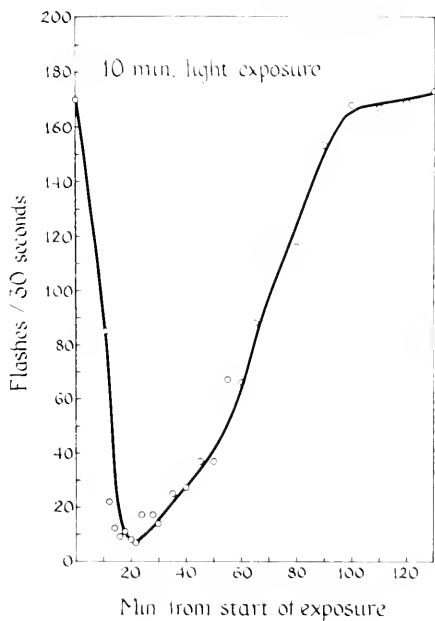


FIGURE 3. Light inhibition of luminescence in water collected from Woods Hole Harbor. See text for details.

during August, 1965. In the subsequent work casual observations of flashing inhibition were made at least once a month by turning on the lights in the dark-room (350 lux at the water surface, G.E. "cool white" bulbs, approximately 4100° K color temperature) at the end of a day's observations and measuring the effect on flashing rate. Inhibition was always found, although the amount of inhibition varied greatly.

Light inhibition of luminescence was quantitatively measured on April 13, 1966, in water collected on April 11 and kept continuously in the dark. Results are shown in Figure 3. The water was exposed to the room lights (350 lux as above) for periods of 10 minutes, 5 seconds, and 1 second. All of these exposures gave considerable inhibition. In the 5-second and 1-second observations two barrels of water were used simultaneously (the white and black circles in Figure 3). Recovery was always complete within 100 minutes of the start of exposure. Interestingly, minimal flashing rates were found 10 minutes after the end of exposure in all cases, including during the 10-minute observation, *i.e.*, inhibition appears to commence when the lights go on. An explanation of this must await better understanding of the physiology of these organisms.

VARIATION OF DINOFLAGELLATE POPULATIONS

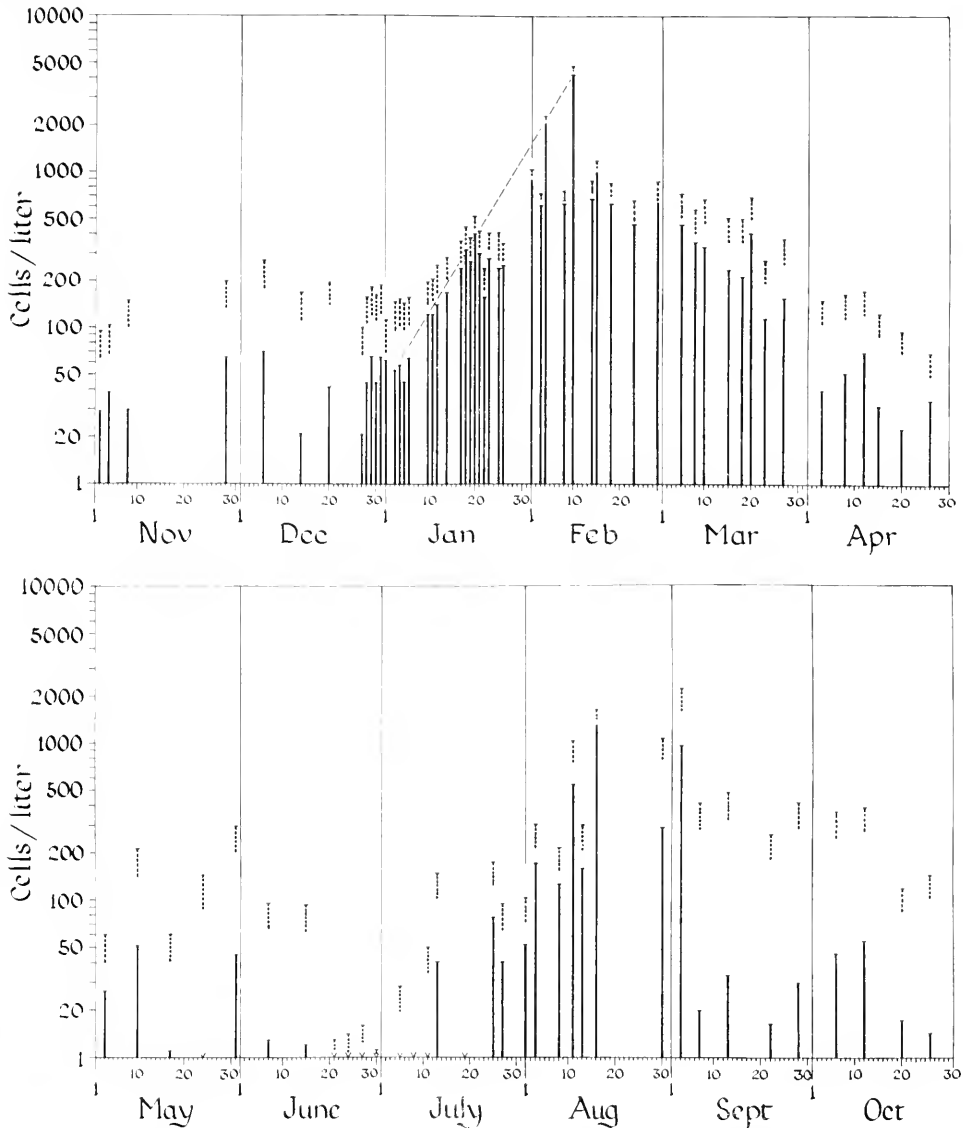
It was necessary to know the concentrations of the luminescent cells so as to determine if sufficient numbers were present to account for the recorded luminescence. Since very few cells per liter may produce measurable quantities of luminescence, and since usual methods of determining phytoplankton populations depend on small (usually 200-ml.) water samples (Braarud, 1958), it was necessary to use a new counting technique.

A quantitative 10-liter water sample was taken from the Woods Hole Oceanographic Institution dock and a surface tow was taken for taxonomic use with a 0.035-mm. mesh aperture, 20 cm. diameter net. Dates of sampling are shown in Figures 4 and 5. The 10-liter quantitative samples were passed through a 400-mesh (0.035-mm. mesh aperture) stainless steel screen. The retained material was very carefully washed from the screen with 15 to 25 ml. of Millipore-filtered sea water from a squeeze bottle and the concentrated samples were preserved with Rhode's iodine or with formalin. The samples were allowed to settle for at least 24 hours, the supernatant was removed, the residue was transferred with washing into a 10-ml. graduated conical centrifuge tube, and the samples were again allowed to settle. The supernatant was again removed, the sample was transferred to a watchglass, and the centrifuge tube was rinsed with sufficient water to bring the sample volume to 1 ml. The sample was strongly agitated in the watch glass and a 0.1-ml. aliquot was transferred to a microscope slide using a hypodermic syringe. The syringe was rotated continuously to prevent the organisms from settling in the barrel during transfer. The dinoflagellates on the slide were tabulated and counted by species under the 10-power objective of a compound microscope. A second or third 0.1-ml. aliquot was examined if necessary to count enough organisms. Since the organisms in 10 liters were concentrated into 1 ml., 0.1 ml. of the concentrate contained the organisms in 1 liter of the original sea water.

Specimens from the net tows were drawn and identified. The taxonomy is discussed by Kelly (1968) and will be published elsewhere. Although immediate

sight identification of dinoflagellates is difficult, it was possible to recognize the species in a sample during counting if detailed studies of the two samples were made first.

There were possible sources of error in the sampling technique but with care all proved negligible. Sample losses during transfer and during removal of the super-



FIGURES 4 and 5. Dinoflagellate concentrations found in samples of water from Woods Hole Harbor during 1965 and 1966. Dashed lines: total dinoflagellates. Solid lines: luminescent species. The straight dashed line points out an apparent logarithmic increase in populations that were nearly all *Peridinium subinermis*.

nantant were not significant since examination of the supernatants, glassware, and screen very rarely showed specimens left behind. Inconsistencies due to transfer to the slide and measurement of the sample aliquots were negligible since results were repeatable (see Table III). Similarly, these repeatable results argue against inconsistencies in the original sampling and screening. Some cells may have passed through the screen; indeed, Wheldon (1939) found considerable loss of specimens when collecting on 0.065-mm. aperture mesh bolting silk. However, many species are smaller than 0.065 mm., but few are smaller than the 0.035-mm. aperture mesh used here. Since no armored dinoflagellates could be found on

TABLE III
*Dinoflagellate concentrations in cells per liter measured at various times of day
in Woods Hole Harbor on November 29, 1965*

Species	Time			
	1530	1730	2100	2300
<i>Ceratium fusus</i>	10	8	9	8
<i>C. lineatum</i>	77	78	74	84
<i>C. macroceros</i>	4	5	5	5
<i>C. tripos</i>	38	36	39	28
<i>Glenodinium lenticula</i>	13	17	17	23
<i>Gonyaulax</i> spp.	1	1	1	1
<i>Peridinium claudicans</i>	3	6	6	4
<i>P. divergens</i>	3	3	2	2
<i>P. leonis</i>	3	3	2	2
<i>P. oceanicum</i>	2	2	1	1
<i>P. ovatum</i>	6	5	4	6
<i>P. pallidum</i>	37	36	33	37
<i>P. subinermis</i>	2	2	3	3
<i>Phalacroma</i> spp.	1	1	1	1
Total of all species	199	203	196	204
Total of luminescent species	64	66	61	63
Liters counted	2	2	2	2

cleared Millipore filters through which screened water was passed, and since water passed through the screen rarely showed bioluminescence, loss through the screen must have been usually insignificant.

Possible sources of error are misidentification of species of small size or confusing shape and normal random sampling errors. Misidentification was rare, and was insignificant in the overall results. Extra aliquots were counted so as to increase the sample size when the specimens were few enough to give significant random counting errors.

The concentrations of total armored dinoflagellates and of luminescent dinoflagellates are shown in Table II and in Figures 4 and 5. Thirty-two species were present of which 16 were luminescent. The species are listed in Table I; their concentrations are given in Kelly (1968). Maximal numbers of dinoflagellates occurred during January, February, and March, and during August and September, with decreases in between. The concentrations are compared below with the

amounts of bioluminescence recorded. Variations in cell concentrations during a single day (November 29, 1965) are given in Table III. The uniformity of these results shows a lack of tidal and diurnal fluctuation in populations in Woods Hole Harbor and verifies the reproducibility in the counting method.

CORRELATION OF LUMINESCENCE WITH DINOFLAGELLATE CONCENTRATIONS

The regression lines of luminescent dinoflagellate concentrations and flashes recorded per 30 seconds were calculated and are plotted in Figure 6.

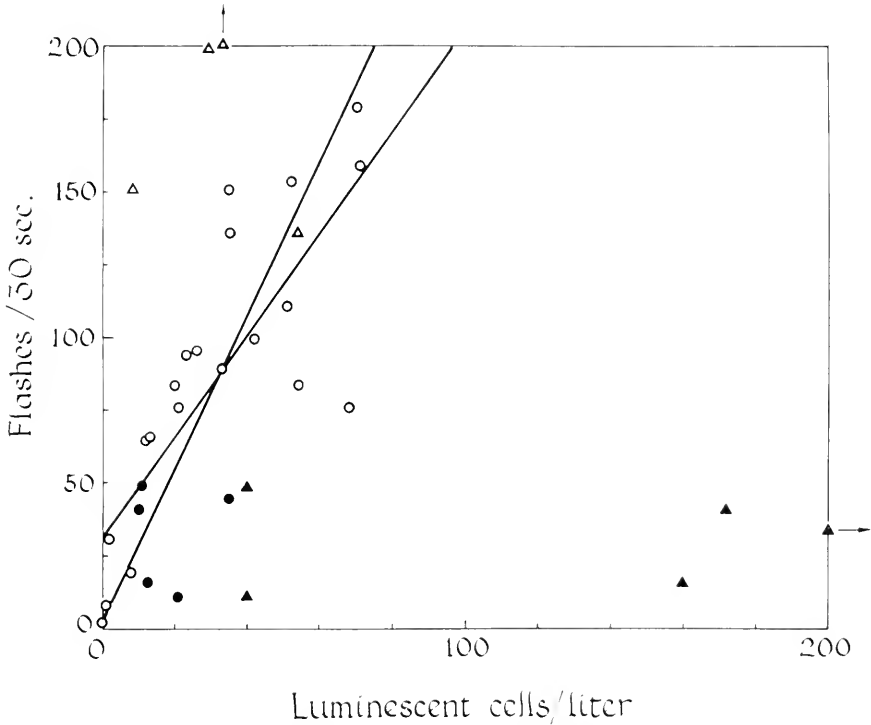


FIGURE 6. Correlations of dinoflagellate concentrations and luminescence in samples from Woods Hole Harbor during 1965 and 1966. See text for details.

On a few occasions described below the data were obviously invalid and not included in the calculations. The high flashing rates on May 17 and between September 13 and October 11 were due to luminescent cells too small to be collected and counted and hence were deleted from the correlations. They are designated by Δ in Figure 6. The luminescence of the small cells was realized because luminescence was recorded in water that passed through the collecting screen. Flashing in the screened water varied from 53 to 103 flashes per 30 seconds compared to 136 to 217 flashes per 30 seconds in the unscreened water. However, the flashing rates in the screened water were probably not closely related to the number of luminescent small cells originally present since some may have been retained on

the screen and many that passed through were probably damaged. A liter of water collected on September 15 was passed through the screen and centrifuged. Many diverse microflagellates were collected, but it was impossible to determine which were responsible for the luminescence.

The flashing rates from July 13 and from July 27 through August 30 (designated by ▲) were low for the number of apparently luminescent cells counted. It was at first feared this discrepancy might be due to some failure in technique such as contamination of the samples with toxic materials, but visual observation while swimming off the dock at night showed qualitatively less luminescence than would have been expected from the populations of *Gonyaulax* spp. counted in the samples. If flashing rates for those dates are plotted against luminescent cell counts not including *Gonyaulax* spp. they fall within the expected range (● in Figure 6). The data from those days were not used in the regression calculations and it seems likely that the *Gonyaulax* spp. populations were not luminescent. This was realized after the populations had declined, too late to directly test the luminescence of the cells.

As the number of flashes increased they tended to overlap on the records and thus at higher flashing rates many of the flashes could not be counted. Those luminescence records could not be expected to have a direct relationship with the number of luminescent cells present. Because of this, samples with more than 100 cells per liter were not included in the regression calculations.

The equations of the regression lines shown in Figure 6 are as follows:

$$Y = (1.75 \pm 0.88)X + (31.5 \pm 21.2) \quad (1)$$

$$X = (0.38 \pm 0.68)Y + (-0.5 \pm 35.5) \quad (2)$$

X and Y represent luminescent cells per liter and mean flashes per 30 seconds, respectively. The errors are given for a 90% confidence interval. The calculations show a correlation coefficient of 0.81 between flashing rates and cell concentration, which with this sample size assures that there is less than a 0.001 probability that there is no correlation.

It is impossible to give a simple ratio relationship between luminescent dinoflagellate population density and flashing rates because of the scatter in the regression curves. This scatter indicates that factors other than simple population density influenced the amount of flashing recorded, which is not surprising considering the diverse environmental conditions and species studied. Species may differ in the number of cells capable of luminescence. For example, not all specimens of *Ceratium fusus* were capable of flashing, while nearly all of *Peridinium subinermis* were. Similarly a particular species may differ in its ability to luminesce under different environmental conditions, *i.e.*, the ability may be influenced by available energy, by nutrient concentrations, or by temperature. Also, different species or a particular species under different environmental conditions might differ in threshold to stimulus, and hence the constant stimulus used in the experiments might more readily produce flashing in some populations than in others.

It is nonetheless apparent that there was a strong relationship between the amount of flashing and the density of the luminescent populations present in the water. Except in the few instances when small protozoa were mainly responsible

for the luminescence, armored dinoflagellates were present in sufficient numbers to explain the flashing recorded.

DISCUSSION

From this work it may be concluded that for the period studied dinoflagellates were the only major cause of bioluminescence in Woods Hole Harbor, except on rare occasions when microflagellates were present that were too small to be taken on the collecting screen. The question arises whether the armored dinoflagellates are able to account for the luminescence that is found almost everywhere in the surface regions of the oceans. Extensive studies at other locations (Kelly, 1968; to be reported separately) indicate that dinoflagellates are responsible for luminescence in many diverse oceanic environments. Dinoflagellates are common organisms (Lebour, 1925; Schiller, 1937) and are evidently commonly luminescent.

The question also arises whether dinoflagellate luminescence has evolved separately in several diverse dinoflagellate genera, or whether the luminescent species are derived from a common ancestor. Because of the diverse biochemical systems of luminescence in other marine organisms such as euphausiids, ostracods, shrimp, and bacteria (Harvey, 1952; Johnson and Haneda, 1966) it seems likely that bioluminescence evolved several times in the oceanic environment. This argues a strong selective pressure for bioluminescence in the sea. Several features of dinoflagellate luminescence, however, argue that the systems are biochemically and hence genetically related.

The four species whose spectral outputs have been studied have similar emission curves (Fig. 7). *Pyrodinium bahamense*, *Gonyaulax polyedra*, and *Pyrocystis lunula*, very diverse species, all have a spectral peak between 470 and 480 nM., rising sharply from between 450 and 435 nM. and dropping off by 550 nM. (Taylor *et al.*, 1966; Hastings and Sweeney, 1958; Swift and Taylor, 1967). *Noctiluca scintillans* has a similar light output but its spectrum was not as accurately measured (Nicol, 1958). The spectral differences between these species are very slight and can probably be attributed to differences in technique. Although the emission curves of other organisms peak near this wavelength, they usually have a different shape (Johnson and Haneda, 1966).

The very similar emission spectra of widely diverse dinoflagellate species argue for the same light-producing enzyme-substrate reaction throughout the group since the bonding energies and structure of the light-producing molecules determine their output spectra. In addition, Hastings and Bode (1961) found that cross-reactions of *Gonyaulax monilata* and *G. polyedra* luciferin and luciferase produced light, as did cross-reactions between *G. polyedra* and *N. scintillans* (Hastings, personal communication).

Dinoflagellate luminescence spectra are similar to sea-water light transmission curves and this suggests an adaptation of luminescent dinoflagellates to the environment: an evolutionary adaptation on the molecular level. Light transmission of most offshore sea water is at a maximum between 470 and 480 nM., *i.e.*, daylight at depths below about 10 or 20 M. is blue (Jerlov and Koczy, 1951). The luminescence and maximal visual sensitivity of other marine organisms are usually in a similar color range (Nicol, 1958) although the biochemical mechanisms of their luminescence certainly differ. Although McElroy and Seliger (1962) suggest a

purely biochemical function, these spectral similarities seem to suggest an adaptation of the luminescence of marine organisms to the blue maximum transparency of sea water or to the blue sensitivity of marine animals' eyes.

The question is thus raised as to whether the identical emission spectra of dinoflagellates are due to inheritance without change from a highly adapted parent species, or whether the spectra arose independently through parallel adaptation to

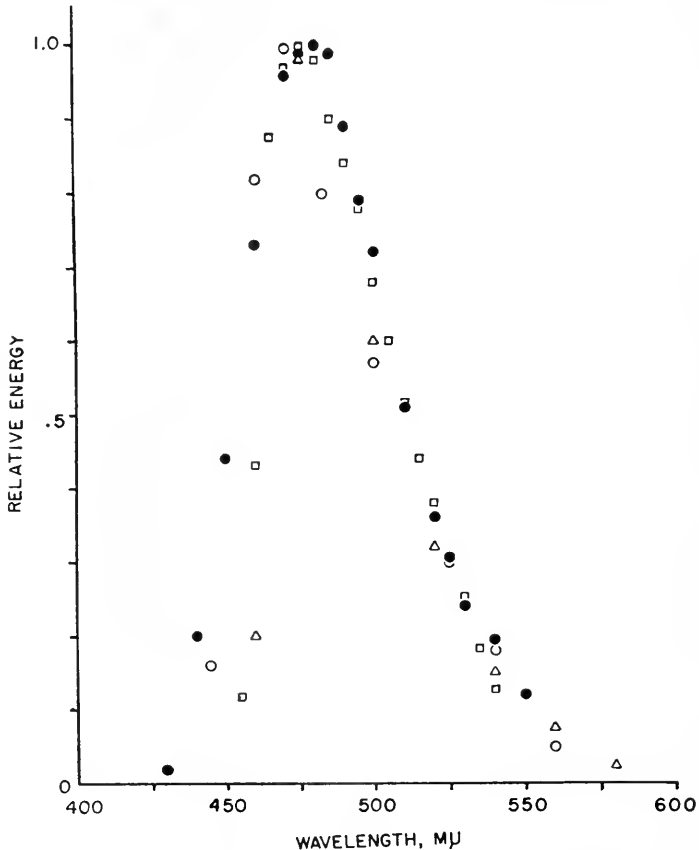


FIGURE 7. Emission spectra of luminescent dinoflagellates. ○ *Noctiluca scintillans* (Nicol, 1958); □ *Pyrodinium bahamense* (Taylor *et al.*, 1966); ● *Gonyaulax polyedra* acetone extract (Hastings and Sweeney, 1957); △ *Pyrocystis lunula* "cysts" (Swift and Taylor, 1967).

identical environmental characteristics. The cross-reactions between the species and the fact that the spectra of these species show more similarity to each other than to unrelated marine forms suggest very similar or identical chemical structures and derivation from a single parent stock. This may well have been a primitive gymnodinioid form similar to the swarm cells of *Noctiluca scintillans*. The retention by these forms of the genetic information allowing them to luminesce at the expense of energy utilization indicates that luminescence probably continues to serve some adaptive function. Further study of luminescence may have con-

siderable bearing on our understanding of both the phylogeny of dinoflagellates and the adaptive values of luminescence.

If the ecology of luminescence in the ocean is to be understood, we must be able to predict the spatial and temporal distribution of dinoflagellates and must understand their luminescence characteristics. This means that much more work is needed on the trophic dynamics of phytoplankton in the ocean, on the biogeography of dinoflagellates, on their taxonomy, and particularly on their characteristics in culture. To date only a very few luminescent species have been cultured, and the only common colorless species included in these is *Noctiluca scintillans*. It thus appears that an understanding of the characteristics and distribution of luminescence must await better understanding of other fields of marine biology.

Because of the diverse species of luminescent dinoflagellates and because of their diverse characteristics and adaptations, I greatly doubt that any single model will serve to predict or describe the distribution of luminescence. Rather, the problem will require considerable descriptive understanding of the variation of species distributions, of photoinhibition of luminescence, of endogenous rhythms, of the luminescence abilities of different species, of the energy sources for those species and of the relationships between their distribution and the environment. Such an understanding will be slow in coming and will depend on a number of diverse fields of study. I believe, however, that the present work shows dinoflagellate luminescence to be very common and to deserve attention from workers in many fields.

I would like to thank Dr. George L. Clarke for encouragement and constructive criticism and for providing necessary facilities. I would also like to thank Drs. Johan Hellebust and David Wall and Mr. Barry Dale for frequent help and suggestions. Mr. Paul Willis made numerous measurements while I was away from Woods Hole.

SUMMARY

1. Bioluminescence in the water in Woods Hole was measured once a week from November, 1965, to November, 1966, and was related to the occurrence of armored luminescent dinoflagellates.

2. Thirty-two species of dinoflagellates were identified, 23 of these were tested for luminescence as isolated cells. Sixteen species were found to be luminescent.

3. Population densities of dinoflagellate species were measured by counting individuals screened from water samples. The concentrations of luminescent species varied from nearly 1000 cells per liter in February, 1966, to less than 5 cells per liter during June and July, while total dinoflagellates ranged from more than 4000 cells per liter to 5 cells per liter.

4. Amounts of luminescence radiometrically recorded in air-bubble-stimulated water samples correlated with the numbers of luminescent armored dinoflagellates present. Exceptions to this correlation were found when population densities were too high for the radiometric recording techniques to be accurate and during May and August, 1966, when the luminescent organisms were too small to be collected and counted.

5. Endogenous diurnal rhythms of flashing were found only during August, 1965 (Kelly and Katona, 1966).
6. Light inhibition of flashing was found whenever it was tested (at least once each month), although sensitivity to light inhibition varied greatly—as little as 1 second of exposure to 350 lux illumination reduced flashing by $\frac{2}{3}$.
7. The ecology and evolution of luminescent dinoflagellates are discussed. It is suggested that dinoflagellates are responsible for most near-surface luminescence, that luminescent forms are derived from a single common ancestor, that the ability serves an adaptive function, as yet undetermined, and that much more work on a number of areas will be required before a detailed ecological understanding of dinoflagellate luminescence is obtained.

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THE GENERAL ECOLOGY AND GROWTH OF A SOLITARY ASCIDIAN, *CORELLA WILLMERIANA*¹

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The diverse assemblage of marine organisms growing on boat hulls, floating docks and other man-made installations is commonly termed the fouling community. Many practical investigations into the development and control of these communities have been undertaken (see Anon., 1952, for review). Ecological study of the fouling communities has been hampered, however, by the lack of information on the life cycles, growth and survival rates, and other properties of the individual species forming this complex (Grave, 1933; Berrill, 1950; Abbott, 1957; Moore, 1958).

One group which may form an important, even dominant, part of the fouling complex, but which has received very little attention, is the Ascidiacea. Scattered ecological data on this group can be found in a few papers, for example those by Huntsman (1921) on *Chelyosoma*, and Just (1934), Hüus (1937), and Millar (1953) on *Ciona*. Although certain aspects of ascidian population ecology have been studied by Millar (1952, 1954) and Goodbody (1961a, 1961b, 1962, 1963, 1965), a detailed study of the growth of many individuals of a single population is lacking.

Of the thirteen species of solitary ascidians that occur in the Puget Sound region of the Pacific Northwest coast of the United States, one of the more common occurring on marina floats is *Corella willmeriana* Herdman (which will hereafter be referred to as *Corella*). This small (2–3 cm. in height), fairly transparent animal breeds throughout the year. The eggs are shed into the atrial chamber, where they are fertilized and develop into the free-swimming tadpole stage before being released. The aims of this study were to determine the growth rate of the young post-larval stages of *Corella* in the field, the approximate size at sexual maturity and the average life span. Observations on predation and on interspecific competition for space have been included, since it is the relative interplay between all these factors that determines the establishment, persistence, and general success of any species population characteristic of fouling communities.

MATERIAL AND METHODS

Study area and apparatus

Field observations were carried out at the Bremerton Yacht Club on Phinney Bay, adjacent to the city of Bremerton, Washington, from April 24, 1966, to April 15, 1967. These floats, although composed of such diverse materials as creosoted

¹ From a thesis submitted in partial fulfillment of the requirements for the M.S. degree, University of Washington, 1967.

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and fiberglass-covered styrofoam, wood, and rubber (old airplane fuel tanks), harbor a luxuriant growth of marine animals, including usually a large population of *Corella*. In order to study the growth rate and settling preferences of the young post-larval stages of *Corella*, two frames of polyvinyl chloride were constructed (Fig. 1). Each frame held 32 8.3×10.2 cm. glass plates 1 mm. in thickness, arranged on four sides of the frame, four pairs of plates per side. The plates were suspended vertically in pairs so that organisms could settle on only one surface of

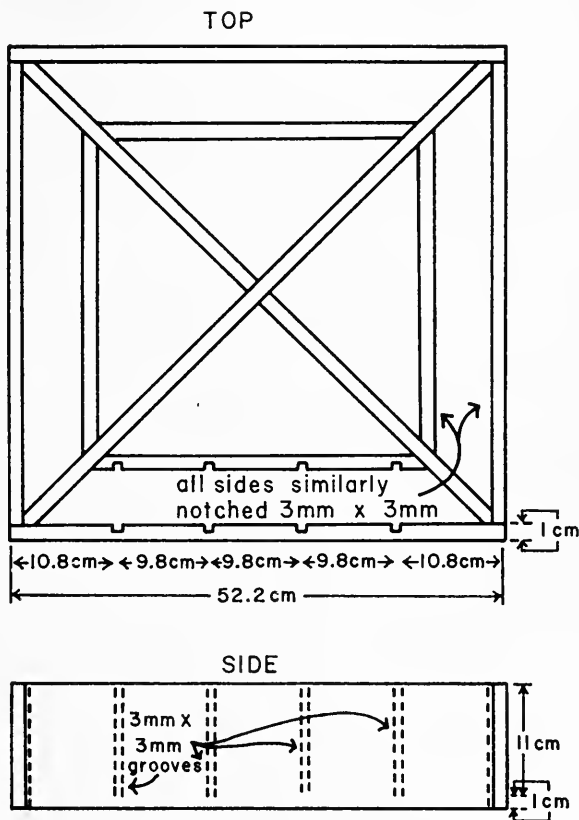


FIGURE 1. Diagram of settling frame.

each plate. The frames (designated A and B) were suspended about 24 cm. below the water surface by nylon rope and plastic-coated wire from the walkway on the floats.

On April 23-24, 1966, a large number of new creosoted styrofoam floats were placed in the water at the Bremerton Yacht Club by the management. Since this was the date on which the two frames were initially submerged, a good opportunity was provided for comparing each month the extent of fouling of the floats and frames. It was hoped that this would give some indication of the effect (or lack of effect) of different substrates on settlement intensity and pattern.

Replacement schedule for the glass plates

A record of the first month's growth of the newly settled *Corella* was obtained for each month of the year by replacing monthly four plates from each frame. For submergence times of 2–11 months, two plates were removed from each frame each month. The remaining eight plates in each frame were left in the water for 12 months. The schedule of the particular plates to be replaced was decided before the frames were placed in the environment by using a random numbers table.

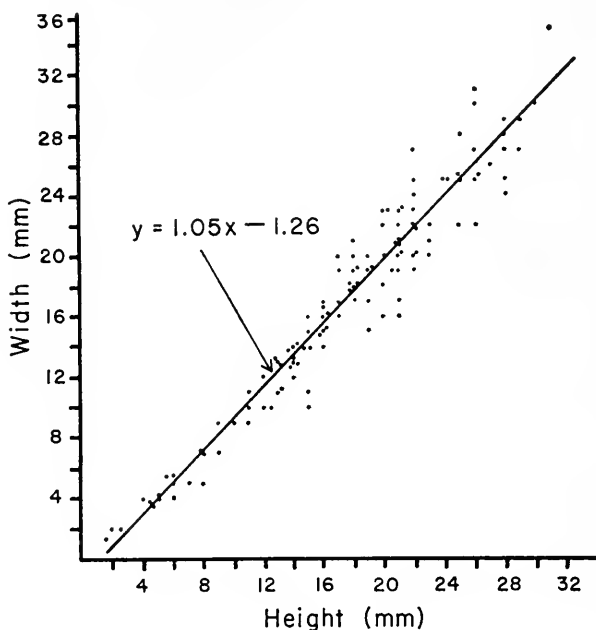


FIGURE 2. The relationship of height to width of 124 *Corella* from natural populations. The line was fitted by linear regression.

Pomerat and Weiss (1946) indicated that the surface texture of the substrate is an important determinant of the species and numbers of animals that will settle. In order to determine to some extent whether the tadpoles preferred to settle on smooth or roughened glass, four of the eight plates replaced each month had a smooth surface exposed for settlement, as did eight of the sixteen plates left in the frames for the entire year. The remaining plates were roughened on one side with coarse carborundum paper.

On 8/20/66 it was discovered that both the nylon rope and plastic coated wire supporting frame A had been cut. The frame was retrieved from the bottom a month later and after cleaning, a new set of glass plates was installed and the frame submerged on 9/25/66. The schedule of replacement of the glass plates was begun again from the beginning, but time did not permit an additional year of study of frame A from the date of its resubmergence. Thus only from April through July was settlement on the two frames comparable.

As each plate was removed from its frame it was immediately isolated in a clear water-filled plastic sandwich box. These were transported back to the laboratory in an ice chest and examined with a binocular dissecting microscope without ever being removed from the boxes. In this way the organisms which had settled on the plates were disturbed as little as possible, and if anything was dislodged, it was kept with that plate. During the replacement of plates and examination of the frames each month, the frames were raised to the surface of the water but were never removed from the water, thus minimizing any undue disturbance and possible tearing off of organisms. The water in which the frames were suspended was fairly sheltered, and tidal currents, though present, were not swift. Therefore, many *Corella* and other organisms were able to remain tenuously attached to the frame.

Method of size measurement of Corella

The measure of individual size used in the growth studies was determined as follows. The height of *Corella* shows a linear relationship (Fig. 2) to its width, as measured across the two siphons. Therefore, although both height and width are a valid measure of size, the latter was used because it was easier to make and entailed a minimum of disturbance to the animals. Newly metamorphosed *Corella* possesses paired atrial siphons (which later fuse to form one atrial siphon) and a single branchial siphon. For newly metamorphosed individuals, the width was measured from the branchial siphon across one of the atrial siphons.

RESULTS AND DISCUSSION

General description of changes occurring on settling frames

The two settling frames were placed in the water on April 24, 1966. During the first two months of submergence, large clumps of filamentous diatoms, hydroids, and barnacles developed. By June 19 a few specimens of *Corella* were large enough to be seen easily on the frames, the largest measuring 4.5 mm. in width. After three months' submergence the number of *Corella* had increased greatly; the largest were 12 mm. in size and were sexually mature, as evidenced by the full sperm duct in all the largest individuals and the presence of a few eggs floating in the atrial chamber of some. Large masses of hydroids (mostly *Obelia* spp.) and the colonial ascidian *Distaplia occidentalis* were also present, though the number of barnacles and filamentous diatoms had declined. In August, frame A was missing, but *Corella* was very abundant on frame B, and the largest individuals, now four months old, measured 30 mm., a size near maximum for float populations of this species. *Distaplia* colonies were numerous, and the colonies of another colonial tunicate, *Diplosoma macdonaldi* (formerly *D. pisoni*) had begun to appear.

At the September sampling, most of the hydroids on frame B had disappeared, as had most of the largest *Corella*. Some empty entire and torn tests were found still attached to the frame and to some of the glass plates submerged since April. The cause of death of most of the *Corella* is not known. It was most likely not environmentally caused, because hundreds of 1-2-month-old *Corella* on the frame and plates appeared perfectly normal and healthy. A small percentage of the

mortality was caused by flatworms of the genus *Eurylepta*; this will be discussed later.

Very few sexually mature specimens of *Corella* were observed on frame B in October; most of the population measured about 1 cm. in width or less. The largest of these had become sexually mature by the end of November, and throughout December, January, and February large clumps of mature *Corella* dominated the surface of frame B and those plates which had been in the water for 4-6 months. Very little growth of *Distaplia* occurred during the winter months; most of the individuals had disappeared or had died back to an apparently quiescent base. On the other hand, colonies of *Diplosoma* grew rapidly between December and March, covering most of the bare space (including a large part of the plates replaced each month) and overgrowing the *Corella* to some extent. Between November and February, when large numbers of *Corella* tadpoles were settling on the clean plates each month, many specimens 300 μ or less in size were found beneath the spreading *Diplosoma* colonies. The overgrown individuals near the edge of the colonies were often still alive, but those near the center of the colonies had died. Figure 3 shows three tracings made of a glass plate removed from frame B and suspended in running sea water in the laboratory for two months. The *Diplosoma* has eventually grown around the *Corella* without growing over them; in all cases there is a space surrounding the *Corella*. These events are similar to those which occurred on the floats and frame B at Bremerton; *Diplosoma* usually grew around rather than over organisms larger than 1 cm. in height.

By March a few of the largest *Corella* had disappeared from frame B, especially those nearest the top edge of the frame and plates. This may have been due to low surface salinity, caused by the heavy rainfall during the winter months which resulted in a surface layer of fresh water that could be seen floating over and mixing to some extent with the salt water underneath. At the completion of these observations in April, 1967, nearly all the adult *Corella* had disappeared from frame B. This second wave of mortality is believed to be due to the large masses of filamentous diatoms that had accumulated on frame B starting in March and increasing tremendously in April. Thick mats of the diatoms were found attached to the siphonal end of the remaining adult *Corella*, and in some the branchial basket was clogged to a variable extent. Large sheets of *Diplosoma* were still present, and *Distaplia* had reappeared. Hydroid growth was negligible.

The changes occurring on the creosoted floats were quite similar to those occurring on frame B, with a few exceptions. When initially submerged, in April of 1966, barnacles settled thickly on the floats (nearly all of these fell off a few months later), but relatively few settled on the frames and virtually none on the glass plates. Most of those that did settle on the frames had fallen off after three months. Large populations of *Corella* colonized both the frames and the new floats during the summer of 1966, but after the large adult mortality in September, younger individuals did not replace them so rapidly or completely on the floats as on frame B. During the winter months the anemone *Metridium senile* became very dense on the floats and grew tall enough to extend beyond the *Corella* present there. On frame B, in contrast, specimens of *Corella* were always larger and more numerous than *Metridium*. It appears that the smooth surface

of the polyvinyl chloride frames provided a substrate somewhat more suitable for *Corella* than for some of the other species present in the area.

After frame A was resubmerged in September, filamentous diatoms and hydroids were the first to appear, though not so thickly or in so short a time as in the spring. The diatoms were gone by November, although the hydroids (mainly *Obelia* spp.) continued to increase in numbers and length steadily until

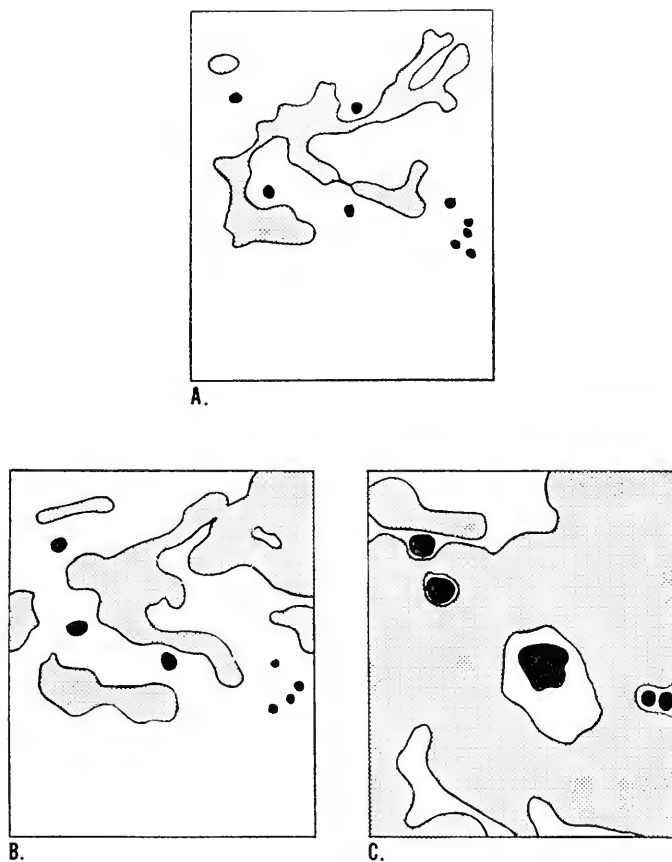


FIGURE 3. Diagram showing growth of *Corella* (solid black circles) and the colonial ascidian *Diplosoma macdonaldi* (stippled) on a glass plate in the laboratory. Unutilized space is blank. A, 8/23/66; B, 9/6/66; C, 10/29/66.

April, when the project was terminated. Throughout the winter months large parts of the frame and plates were covered by broad flat colonies of *Diplosoma*. *Corella* was not observed on frame A or its plates until January. In February about three or four mature individuals 2–2.5 cm. in size were observed, suggesting that there were a few young unnoticed *Corella* present before January. Even in March only about a dozen were observed, and most of them had not yet reached reproductive size. By April these were mature and healthy-looking. Almost

no filamentous diatoms had accumulated, possibly because frame A was in a shadier location and subject to faster water currents than frame B. Large numbers of laminarians and *Agarum* sp. were found growing on the upper edges of frame A in March, and although most of them were removed so as not to cover the frame completely, they had grown back again in April; several had reached a length of one meter. *Distaplia* colonies appeared in November but grew very little until the beginning of March. By the end of April the colonies had increased in size, and at the termination of observations the most abundant organisms on frame A were *Diplosoma*, *Distaplia*, *Corella* and *Obelia*.

It is obvious that an important difference between the two frames developed after the resubmergence of frame A. Before that there had been some quantitative differences, but in general they were comparable. During the winter months, how-

TABLE I

Surface water temperature and size after one month of growth of the largest 10% of Corella at the Bremerton Yacht Club (April 1966–April 1967)

Date	Total no. settled	Mean size of largest 10%	H ₂ O temp. (° C.)
April 24	—	—	11
May 21	0	—	12
June 18	1	1.4 mm.	15
July 16	0	—	16
Aug. 20	71	1.7 mm.	17
Sept. 25	1164	1.9 mm.	14.5
Oct. 22	10	1.5 mm.	11.5
Nov. 20	604	.356 mm.	11
Dec. 18	439	.500 mm.	9.5
Jan. 22	525	.507 mm.	8
Feb. 19	284	.402 mm.	8
March 19	89	.394 mm.	8
April 15	6	.430 mm.	10

ever, while frame B contained a large breeding population of *Corella*, frame A remained nearly completely free of *Corella*. Floats nearby frame A possessed at least a few *Corella*, so theoretically there should have been tadpoles in the plankton. Table I presents the total number of *Corella* settling on the plates removed monthly from frame B, and it will be seen that not until a breeding population had become established (July–August) on the frames did there appear large numbers of tadpoles settling on the plates. After the first adult mortality in September, only a few tadpoles settled in October. But beginning in November, after a second breeding generation had developed, there were again large numbers of tadpoles settling. When the second wave of adult mortality occurred in the spring of 1967, the number of tadpoles settling decreased sharply, in spite of the fact that the *Corella* on frame A and some of the marina floats were still alive and healthy. Thus it seems to be extremely local recruitment of tadpoles that accounts for the presence of nearly all of the individuals on frame B. Nair (1962) has made a similar observation for compound ascidians in his extensive paper on marine fouling.

Seasonal fluctuations in settlement and growth rate of young Corella

Because of the small size of the plates and the fact that frame A had to be restarted, the variation on plates from frame B submerged for longer than one month was too great to permit any realistic evaluation of the growth rate of *Corella* older than one month. Fair numbers of tadpoles settled on the plates which were replaced monthly, although this number fluctuated widely from month to month (Table I).

In agreement with Pomerat and Weiss (1946), there was a significant difference between the numbers of individuals settling on smooth and roughened plates each month. For the eight months considered appropriate for making these paired observations, 662 settled on smooth and 1405 on roughened plates. The hypothesis of random settlement was rejected by χ^2 ($p \ll 0.01$), indicating that *Corella* either preferentially settled on or adhered to the roughened surfaces. Between plates of different surface texture, no significant difference in individual size of the

TABLE II
Number of newly settled Corella 200 μ in size or less on plates submerged for varying lengths of time

Month of analysis	Duration of submergence	Number of <i>Corella</i>
Nov.	1 mo.	549
	7 mo.	0
Dec.	1 mo.	200
	2 mo.	48
	4 mo.	4
	8 mo.	0
Jan.	1 mo.	169
	3 ¹ / ₂ mo.	14
	9 ¹ / ₂ mo.	5

largest 10% settling each month could be demonstrated, by an analysis of variance. Therefore, size measurements from all four one-month plates from frame B were pooled each month. Only the largest 10% were used, based on a statement by Moore (1958) that such a measure of the few largest should give a reasonable indication of growth for the entire period.

One might question the validity of assuming that the monthly size measurements actually represent in most cases an entire month's growth. Many projects have been carried out to determine the normal sequence in fouling of denuded or clean surfaces submerged in the sea (Scheer, 1945; Anon., 1952). Although this problem is far from being solved, it does appear that certain types of organisms settle first (usually bacteria, diatoms and suctorians) and other organisms settle later, even if ready-to-settle larval stages have been present in the water for the entire time. *Corella* tadpoles, however, apparently prefer to settle on bare surfaces, as evidenced by the number of newly settled tadpoles observed on plates submerged for varying lengths of time (Table II). (Similar data have been collected on a larger scale by Goodbody (1965) for *Ascidia nigra*, which also shows a marked

preference for unfouled surfaces.) The indications are, then, that *Corella* will settle immediately on clean surfaces such as glass, supporting the assumption that the largest specimens represent a full month's growth. The close relationship between size of the breeding population on the frames and number of young *Corella* on the one-month plates is further support for this assumption. It might have been better to submerge panels on which tadpoles had been allowed to settle in the laboratory and for which the exact age was thus known, but this was unfeasible for the present study and would have introduced other problems of analysis.

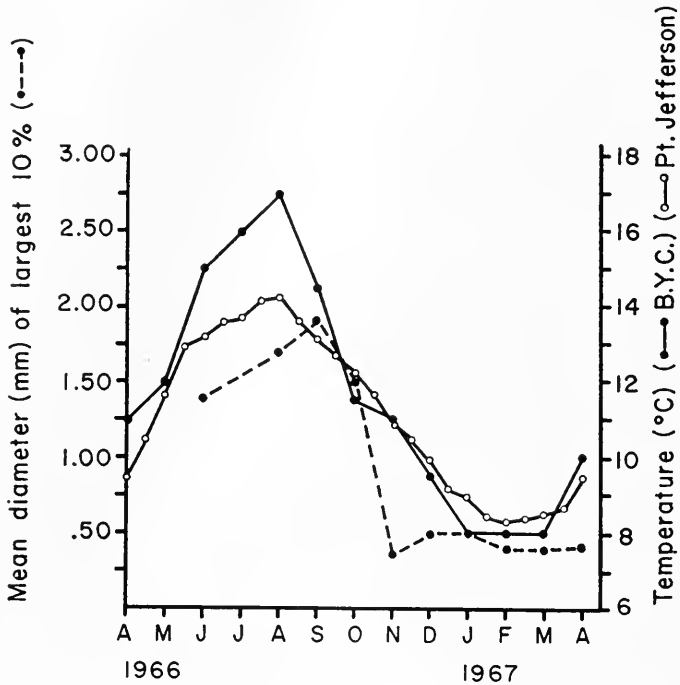


FIGURE 4. A comparison of surface water temperatures at Pt. Jefferson and the Bremerton Yacht Club with the size attained by *Corella* after one month of growth.

In order to determine whether there were seasonal fluctuations in the growth rate of the young stages of *Corella*, the average size attained in one month by the largest 10% of the newly settled individuals for each month throughout the year was measured (Table I). It should be noted that some of the figures are more reliable than others, being based on larger sample sizes. For instance, in October very few mature individuals were present on frame B, and a total of only ten *Corella* was counted on all four of the one-month plates removed that month. In order to obtain a reasonable size value for such a small sample, the size given (1.5 mm.) is an average for the three largest individuals. In April of 1967 only six *Corella* settled on the one-month plates, and the size given is the largest of those six. The mean size for November is abnormally small, probably

because it represents less than a whole month's growth. The breeding stock of *Corella* on frame B was very small until those individuals which were still immature in October grew to reproductive size, sometime before November 20. Thus recruitment of the tadpoles to the plates throughout the month of November was unlikely; the 604 individuals on the one-month November plates probably represent settlement for only the prior two weeks or so.

In Figure 4 the size values for the first month's growth of *Corella* for each month of the year have been plotted against temperature. Since surface temperature readings were taken only once a month at Bremerton, it was thought that these readings might show enough variability to mask average seasonal trends. Therefore, along with the Bremerton temperature readings are plotted average bimonthly surface temperature readings for Point Jefferson (Megia, 1956). The Point Jefferson and Bremerton curves agree fairly well, although as might be expected, higher summer temperatures occur in the inshore waters at Bremerton. Figure 4 shows that there is a strong correlation between the initial month's post-larval growth and temperature. Seasonal food supply is very likely also an important factor here, but it was not measured in the present study. The correlation between morphological changes occurring after metamorphosis and the rate of growth of *Corella* is currently being investigated.

Predation on Corella

Eurylepta leoparda Freeman, a cotylean polyclad flatworm, was found inside a few adult *Corella* on the settling frames and floats at Bremerton during the September, November, December, March, and April visits. The worms were 1–3 cm. in length; laboratory observations (about 20–25) showed that they always entered the *Corella* by rolling into a tube as they passed through the branchial (incurrent) siphon (never the atrial siphon). Once inside the *Corella*, the flatworms unrolled themselves and usually ingested the branchial basket first; then they might either leave the individual through a siphon or continue devouring the remainder of the internal organs within 3–7 days, finally leaving only an empty test. Ingestion occurred by extrusion of the pharynx and sucking in of the food material. Generally only one flatworm was found inside a *Corella* at Bremerton, but occasionally there were two, and in the laboratory three or four often moved into the same *Corella*.

There are very few references concerning predation on ascidians. The polyclad flatworm *Cycloporus papillosus* feeds on the colonial tunicates *Botryllus* and *Botrylloides* (Jennings, 1957), and several species of molluscs have been observed to feed on simple and colonial ascidians (Thompson and Slim, 1959; Miller, 1961; Barrett and Yonge, 1964; Ghiselin, 1964). Goodbody (1963) states that young *Ascidia nigra* are "undoubtedly" eaten by flatworms and by young polychaetes. To my knowledge, there is only one recorded observation of a flatworm feeding on adult simple ascidians. Crozier (1917) reported that *Pseudoceros crozieri* was found in the branchial sac of *Ascidia curvata* and *A. nigra* as well as on colonies of *Ectinascidia turbinata*. According to Crozier, each of the three sets of flatworms taken from the three species of ascidians would feed only on the particular species on which it had originally been found, that is, there were three "physiological varieties" of *Pseudoceros crozieri*. A few feeding experi-

ments carried out with *Eurylepta leoparda* indicate that it will feed only on adult *Corella* and not on *Ascidia callosa*, *Chelyosoma productum*, or *Diplosoma macdonaldi*, three other species of ascidians presented as food. Conversely, other species of polyclads that have been collected have not been observed to feed on *Corella*. Interestingly, *E. leoparda* has been found only with the Bremerton population of *Corella*. In other local areas, such as Edmonds and Friday Harbor, this flatworm is apparently absent from the *Corella* populations, which show much more marked fluctuations in numbers than does the Bremerton population.

CONCLUSIONS

Corella's position in the successional sequence in fouling communities appears to be the following. It is apparently a primary colonizer, since the tadpoles have been shown to settle preferentially on clean, unfouled surfaces. It is a "fugitive" species; the tadpoles settle on new or denuded surfaces and grow quickly to sexual maturity. As they die off, however, colonial ascidians may take over the space, overgrowing small *Corella*, particularly during the winter, and thus preventing the maintenance of stable populations of *Corella*. Tadpoles tend to settle in the immediate vicinity of the adult *Corella*, however, thus prolonging the length of time that *Corella* can exist before being replaced.

The dominant organisms on the older floats (submerged for five years or more) are *Metridium senile*, *Eudistylia vancouveri*, *Mytilus edulis* and *Pyura haustor*. All of these species, with the exception of *M. senile*, settled sparsely on the frames; only at the termination of observations were they beginning to become more evident. Judging from observations of the older floats, had the frames been left submerged these species would probably have continued to increase in abundance. On the older floats there is, however, a seasonal sloughing off of some of these organisms. These spots are usually then colonized by *Corella*. The populations of *Corella* in shady locations are able to survive the spring period of diatom overgrowth better than other populations and may be partially responsible for the summer recolonization by *Corella* of the denuded areas.

SUMMARY

1. A one-year field study of the ecology of the solitary ascidian *Corella willmeriana* Herdman was conducted between April, 1966, and April, 1967, at the Bremerton Yacht Club, Bremerton, Washington, where two polyvinyl chloride frames containing glass plates were examined at monthly intervals.

2. The results indicate that *Corella* is a primary colonizer, preferring to settle on clean surfaces. Growth is rapid during the summer, when sexual maturity, corresponding to a size of 12 mm., may be attained in three months and life span is approximately five months. Individuals grow at a slower rate and live longer during the winter; the life span then is seven or eight months.

3. Very young specimens of *Corella* are frequently overgrown during the winter by the colonial ascidian *Diplosoma macdonaldi*. The causes of death of adult *Corella* are not completely known, although a small percentage of them are eaten by the polyclad flatworm *Eurylepta leoparda*. A luxuriant spring growth of filamentous diatoms may cause death of adult *Corella* by smothering them.

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FEEDING AND OXYGEN UPTAKE OF THE TROPICAL SEA URCHIN *EUCIDARIS TRIBULOIDES* (LAMARCK)^{1,2}

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Feeding and oxygen uptake of echinoids have been studied by a number of investigators (for a recent review see Boolootian, 1966), but little work has been done on cidaroid urchins. These urchins are particularly interesting since they belong to an ancestral group from which all echinoids evolved (Durham and Melville, 1957).

There are over 150 living species and varieties in the family Cidaridae (Mortensen, 1928). Most of these live in the bathy-benthic zone, but some live in the littoral zone and a few live at great depths. In the West Indian region there are 12 species of cidaroids, one of which is littoral, the rest being found between 30 and 2000 meters (Mortensen, 1928). The littoral species, *Eucidaris tribuloides* (Lamarck), is relatively abundant in some habitats off southeastern Florida, particularly on coral reefs and rocky areas. It is easily maintained in aquaria where it grows at rates at least comparable with, and sometimes greater than those in the field (McPherson, in press). It is thus well-suited for laboratory studies.

This study was made to determine some of the factors which are important in the metabolism of *Eucidaris tribuloides*. Such parameters as temperature, season, size, and nutrition and their effect on oxygen uptake have been considered. Food preference and rate of feeding have been measured under laboratory conditions. This work will form a basis for studies on metabolism of deep-water cidaroids.

METHODS AND MATERIALS

Urchins used in respiration and feeding experiments were collected on Margot Fish Shoal, a "patch" coral reef several miles east of Elliot Key, Florida. After they were brought in from the field they were allowed to acclimate to aquarium conditions for at least several days before being used in experiments.

The food preference of *Eucidaris tribuloides* was tested for the boring sponge *Cliona lampha*, turtle grass (*Thalassia testudinum*), several algae (including *Dictyota* sp., *Liagonea* sp.), and fish strips (mullet, *Mugil cephalus*). Eight rocks were arranged in a circle within a clean holding tank. The rocks were placed so that each was separated from the next by about 10 to 15 cm. The diameter of the circle of rocks was approximately one meter. Four of these rocks were infested with the boring sponge *Cliona lampha* or were placed on the other foods to be tested. Alternating with these were four rocks of similar size, but without any evident algal or

¹ Contribution No. 966 from the Institute of Marine Sciences, University of Miami.

² This investigation was supported by the Department of Health, Education, and Welfare under Grant No. WP-00573.

animal growth. Sea water flowed into the tank, circulated around the tank and flowed out at a drain in the center of the circle of rocks. Urchins were placed in the center of the circle and the following day their positions were recorded in relation to the food. Those touching a non-food rock (control) were recorded as R. Those not touching any rock or food were recorded as 0. This was repeated on several days for each food with fresh urchins. The position of each rock was moved one place each day in the circle.

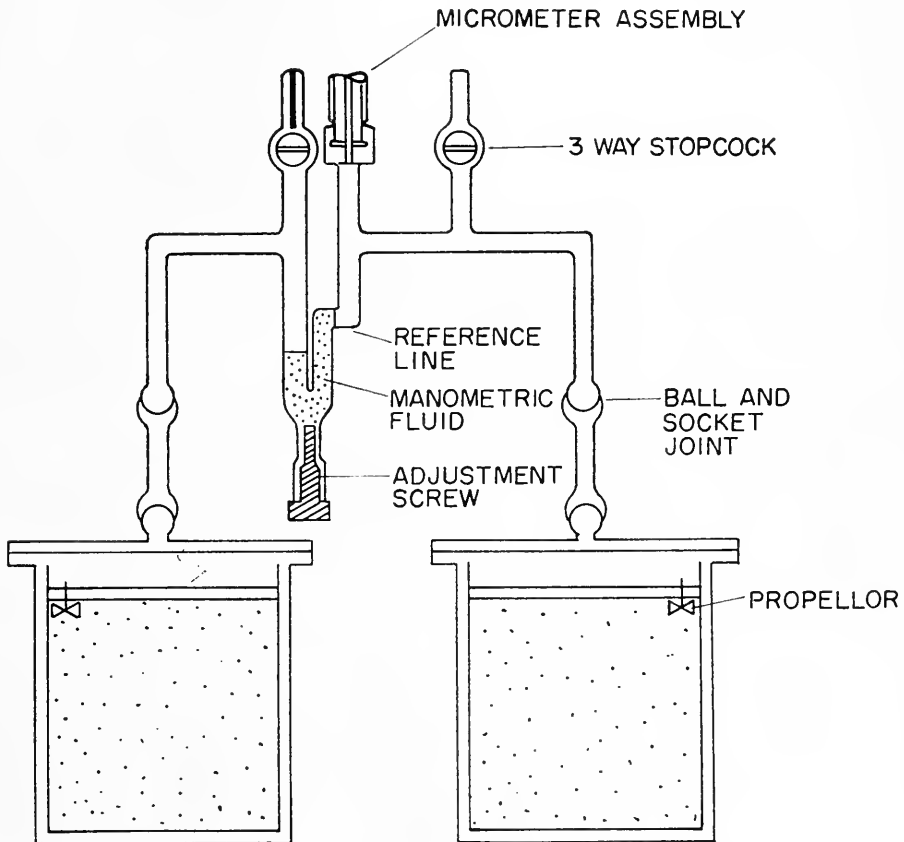


FIGURE 1. Schematic drawing of modified Scholander volumetric respirometer.

Before determining feeding rates urchins were maintained in aquaria two to three months and were offered an excess of rock infested with *Cliona lampa*. Feeding rate was determined in the laboratory on these individuals held in 950-cc. polyethylene cages. Weighed portions of rock infested with *Cliona lampa* were put into the cages and at the end of eight days the rock was reweighed (wet) and the amount eaten determined by differences between first and second weighings. Controls indicated that weight loss of rock during the experimental periods was not significant. Flow of sea water into aquaria was maintained in all feeding experiments at about two liters per minute.

Oxygen consumption was measured with a modified Scholander volumetric respirometer (Fig. 1). This consisted of a differential syringe manometer modified for attachment to two glass vessels (manometer obtained from RGI Inc., Vineland, New Jersey). The micrometer syringe had a total capacity of 200 microliters in 0.2 microliter divisions. The manometer was attached to the vessels by ground glass ball and socket joints. Vaseline was used as a lubricant for the joints and for the ground glass top and lip of the vessels. Hexanol with rodamine B dye was used as the manometer fluid.

The vessels contained a plastic ring with a magnetic stirrer. During respiration experiments small motors outside the vessels were used to stir and aerate the water. Water movement appeared to be an important factor for the well-being of urchins (Moore and McPherson, 1965).

The total volume of a vessel and its connecting tube was about 955 cc. The plastic ring had a volume of 120 cc. The air volume was maintained at about 65 cc., so that there was about 770 cc. in the vessel for sea water and for the urchin. The volumes of the urchins used ranged from about 5 to 80 cc., leaving 765 to 690 cc. of sea water, respectively. One vessel contained the experimental animal while the other (control vessel) contained only sea water. Carbon dioxide released by the urchin would be absorbed in the sea water, and would not increase significantly in the gas phase.

The general procedure for a respiration experiment was as follows:

- (1) Sea water was put in both vessels and well aerated at a constant temperature (in a controlled-temperature bath which held the temperature to $\pm 0.01^\circ$ C.) for several hours.

- (2) An urchin was placed in one vessel.

- (3) The magnetic stirring apparatus was placed in both vessels and the water levels adjusted.

- (4) The vessels were closed and connected to the manometer and then immersed into the controlled-temperature bath, so that the major air spaces were submerged.

- (5) The stop cocks were left open so that both vessels were connected to the atmosphere.

- (6) The sea water in the vessels was stirred with the magnetic stirrers.

- (7) After about 60 minutes the vessels were closed to the atmosphere and readings were made every five or ten minutes for 30 minutes. In a few cases readings were made over periods longer than 30 minutes. The standard error for the mean oxygen uptake in cc. per hour per gm. dry weight, determined from 27 readings on six urchins at 20° C., was ± 0.03 .

- (8) Determinations of oxygen uptake were made during the day, between the hours of 0900 and 1400.

After respiration and feeding rate experiments the test diameter of each urchin was measured. A few of these urchins were saved for later tests, but most were examined to determine sexual condition, gut content, and tissue weight. Sex and the presence or absence of ripe sex cells were determined by microscopic examinations of gonadal smears. Gonads of each individual were removed, measured volumetrically in a graduated cylinder and dried in an oven. Gonad index was

calculated from the gonadal volume and test volume following the method of McPherson (in press). In order to determine the total dry tissue weight, the remaining parts of the urchin, with the exception of the coelomic fluid, were decalcified with 20% formic acid. An ion exchange resin (WIN 3000) was added to facilitate the removal of calcium from the tissue, following a method used for decalcification of teeth at the University of Miami Medical School. A test indicated that the dry weight of the coelomic fluid was negligible (less than 1% of the total weight of the somatic tissue). Tissue of the urchin was dried for 24 hours at 105° C.

The ash-free dry weight of both the urchin and its food, rock infested with *Cliona lampha*, was measured in several cases. The rock was weighed and decalcified by the same method described above for *Eucidaris tribuloides*. The decalcified and dried residue of either of these was burned in a muffle furnace at 500° C. for 24 hours, and then reweighed. The difference between the dry weight before and after burning was used as an estimate of the ash-free dry tissue.

RESULTS

1. Food preference

In a starved state *Eucidaris tribuloides* would consume practically anything including wooden and even fiber-glassed sides of holding tanks. Under normal feeding conditions, however, these urchins showed a preference for the sponge

TABLE I

Food preference of Eucidaris tribuloides in the laboratory with the foods: Cliona lampha rock (C), Thalassia testudinum (T), algal rock (A), and fish strips (F).

R = clean rock; O = urchins not in contact with any of the above items.

Chi square (χ^2) used to test significance of differences between the number of urchins associated with a food (rock) and those associated with a clean rock (control)

Total number tested	Percentages			χ^2
	C	R	O	
169	66	15	19	32.4
	T	R	0	
145	26	28	47	0.1
	A	R	0	
57	26	18	56	1.4
	F	R	0	
65	32	43	25	1.1

Cliona lampha (rock) over *Thalassia testudinum*, algal-covered rocks, and strips of fish (mullet) (Table I). Sixty-six per cent of those urchins tested were associated with the *C. lampha* as compared with 24% not associated with this food. In other tests 26% were associated with *T. testudinum* compared with 74% not associated with food; 26% were associated with algal covered rocks compared with 74% not associated with food; and 32% were associated with fish strips compared with 68% not associated with food. Because of this preference in the laboratory and because

hard material was an important constituent of the gut content of urchins from the field, rock infested with *Cliona lampha* was selected as the food to be used in feeding rate and oxygen uptake experiments.

2. Feeding rate

Figure 2 gives the weight in gm. of wet rock (*Cliona lampha*) ingested per day for well-fed urchins during the period 1 to 9 February 1966. Urchins feed primarily on the surface of rock infested with *Cliona lampha*, ingesting both rock and

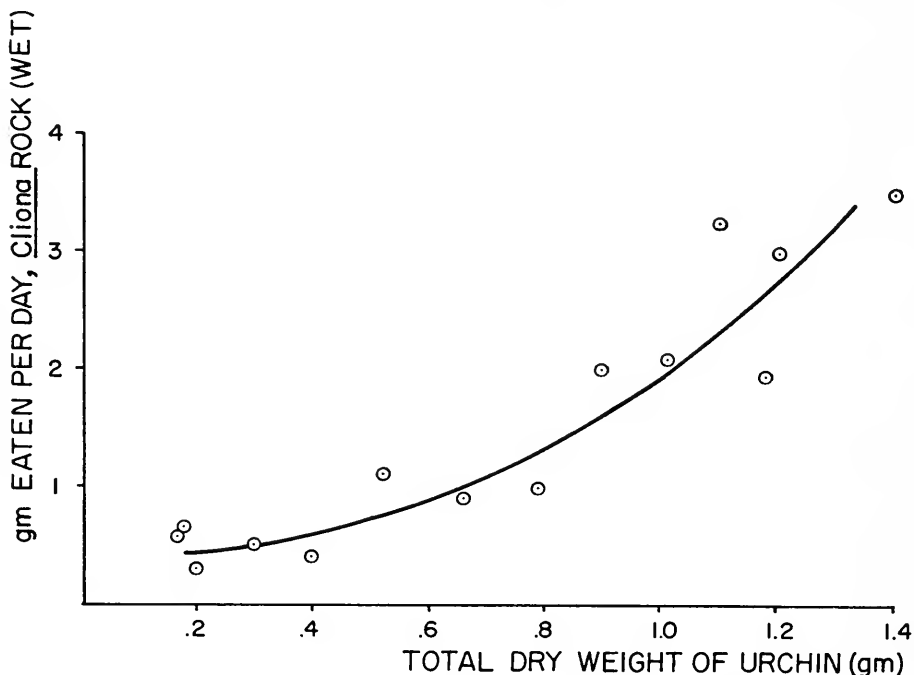


FIGURE 2. Feeding rate of well-fed individuals of *Eucidaris tribuloides* on rock infested with the boring sponge *Cliona lampha*. Experiment was made in aquaria during winter. Mean temperature 17° C. Line fitted by inspection.

sponge. Since the sponge tissue is concentrated at the surface and decreases with depth into the rock (Neumann, 1966), the urchins obtain relatively large amounts of tissue for the rock consumed. When the ash-free dry weight of the urchin and the rock consumed were determined, the intake corresponded roughly to 5% of the ash-free dry weight of the urchin per day.

3. Preliminary experiments on oxygen uptake

Tests were run to determine if changes in oxygen partial pressure or metabolic products would significantly change the rate of oxygen uptake of *Eucidaris tribuloides*. This was done by measuring oxygen consumption over periods of time

considerably longer than normal respiration experiments. If changes in either of these factors significantly affected respiration during the normal tests then their effects should have been even more obvious in longer tests. The actual decrease in oxygen dissolved in the sea water that occurred during a test was measured in two cases. The micro-Winkler method described by Slack (1965) was used to measure the dissolved oxygen.

Urchins held in respirometers for varying periods of time over two hours showed some slight variation in oxygen consumption, but there appeared to be no significant trends in these tests. This was taken as an indication that any changes in the concentration of oxygen, carbon dioxide, metabolites or waste products in the respirometer water during these periods did not result in a measurable effect on the respiration of the urchin.

There was a 6% decrease in the dissolved oxygen in the respirometer water during an hour as indicated by micro-Winkler measurements. This small decrease in the dissolved oxygen in the respirometer water in spite of the oxygen consumed by the urchin is explained by the fact that there is a large source of oxygen in the respirometer vessel (compared with that used by the urchin). As the oxygen in

TABLE II

*O₂ consumption of Eucidaris tribuloides (well-fed and starved individuals).
Urchins fed Cliona lampra rock. Temperature 30° C.*

T.D. (mm.)	Well-fed	cc. O ₂ per hour				
		Days of starvation				
		1	2	3	4	5
42	1.23	1.00	0.96		0.98	
26	0.46	0.34	0.34			
20	0.23	0.20				
28	0.65	0.65		0.28		
36	0.86	0.61		0.59		0.60
30	0.74	0.57	0.58			0.55

the water is removed by the urchin it is replaced rapidly by oxygen in the air over the water due to the action of the magnetic stirrer. As stated, the volume of air over the water in the respirometer was about 65 cc. Since air contains about 200 cc. of oxygen per liter, each vessel before a run would contain about 13 cc. of oxygen in the air and about 5 cc. of oxygen dissolved in the water, depending on the temperature. If an average urchin removed 0.5 cc. of oxygen per hour, the loss at the end of two hours would be 1.0 cc. or only about 6% decrease in the total oxygen. This value is very close to the observed decreases.

4. Oxygen uptake and nutrition

The effect of nutrition on oxygen consumption was studied by comparing the respiration of well-fed and starved urchins. Well-fed urchins were defined as those which were exposed to the food *Cliona lampra* and had this material in their

gut. Starved urchins were defined as those which had been offered *Cliona lampa* in the laboratory, and which were then removed from this food for varying periods of time and held in clean polyethylene cages.

Respiration was measured individually on six well-fed urchins. These urchins were starved, and on one or more successive days their oxygen consumption was

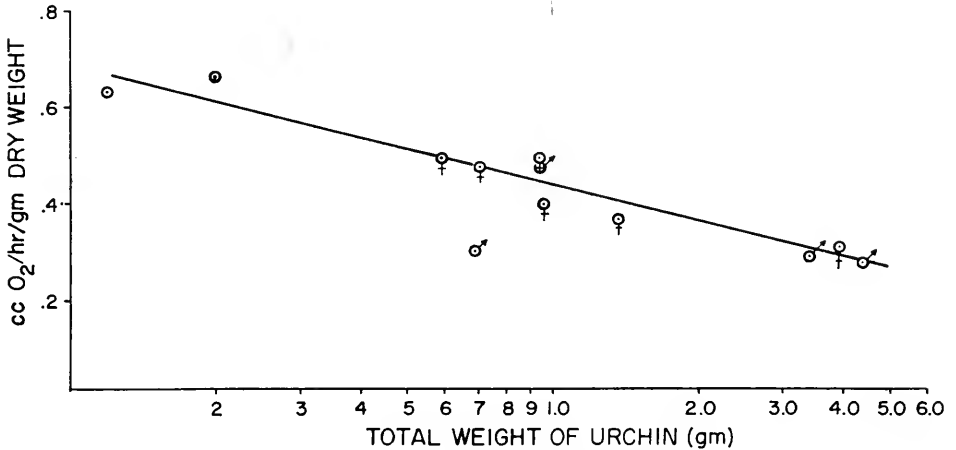


FIGURE 3. Relative oxygen uptake of *Eucidaris tribuloides*. Experiment was made in winter at 20° C.

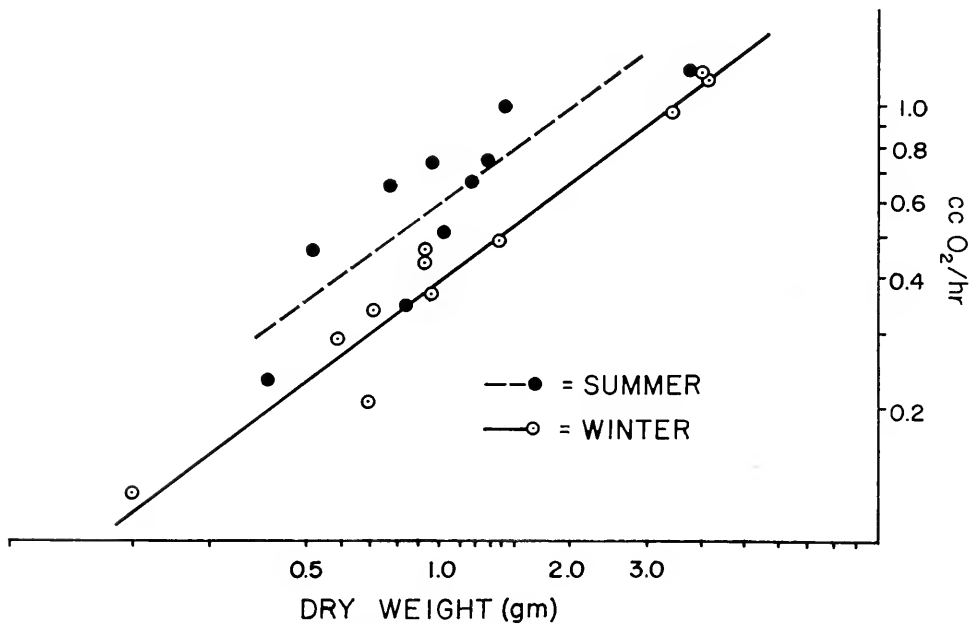


FIGURE 4. Oxygen uptake of *Eucidaris tribuloides* plotted against body size during summer at 30° C. ($b = 0.65$) and during winter at 20° C. ($b = 0.70$). Urchins well-fed.

remeasured. This test indicated that oxygen uptake decreased markedly during the first day of starvation, but showed little change over the next few days (Table II). The mean oxygen uptake decreased 22% from the original rate during the first day. Since oxygen uptake was relatively stable during at least the first one to five days of starvation, this interval of time was selected as a standard starvation period for further work.

5. Oxygen uptake and reproduction

Gonadal condition and sex were determined for most of the urchins in which oxygen consumption was measured. There was no indication that these factors significantly affected respiration. In addition there was no significant correlation between relative oxygen uptake and gonad index.

6. Oxygen uptake and size

Relative oxygen consumption plotted against body weight showed the expected trend of decreasing oxygen uptake with increasing body size (Fig. 3). The smallest urchins tested respired about twice as fast as did the largest urchins, the latter being about 30 times as large as the former.

The relationship between metabolic rate and body size may be expressed in the formula (Bertalanffy, 1957):

$M = a W^b$, where $M =$ cc. O_2 per hour consumed by the urchin

$W =$ weight of urchin

$b =$ log-log slope of the line which relates O_2 uptake to weight

$a =$ the point at which the above line intercepts the ordinate M .

Figure 4 gives the oxygen consumption of *Euclidaris tribuloides* plotted against size at 20° and 30° C. (winter and summer). The slopes of the resulting lines are 0.70 (winter, 20° C.) and 0.65 (summer, 30° C.). These values indicate an oxygen uptake which is almost proportional to surface area (or $\frac{2}{3}$ power of weight). In some echinoderms "b" has been shown to decrease with increasing temperature, indicating that larger animals consume proportionally less oxygen than smaller animals as the temperature increases (Farmanfarmaian, 1966). This may be related to decreased ambient oxygen or increased oxygen demand but insufficient supply.

7. O_2 uptake and temperature

Two aspects of the effect of temperature on respiration were considered. First, the short-term effect of temperature change on oxygen consumption was measured for some individuals in order to determine the time required for respiration to reach a relatively stable rate. Urchins were taken from water with an ambient temperature of 28–29° C. and placed in respirometers at temperatures of 10°, 15°, 20°, and 30° C. The oxygen uptake of urchins placed in water which was cooler than ambient fluctuated during the first 40 minutes, and after this remained relatively stable (Fig. 5). Oxygen uptake was low during the first 10 minutes, then rose to a maximum in 20 to 30 minutes, and then decreased to an intermediate value after about 40 minutes. The oxygen uptake of urchins tested at temperature near

ambient showed little change in respiratory rate over the same interval (Fig. 5). Second, the effect of temperature on the stabilized rate of oxygen consumption was measured at the following temperatures: 10°, 11°, 15°, 20°, 24°, 26°, and 30° C. Urchins were placed in the respirometers at these controlled temperatures and allowed to acclimate for one hour, after which oxygen determinations were made. The urchins used in these tests were within the size range 28 to 35 mm. After each test the dry weight of the urchin was determined and the respiration was expressed as cc. oxygen per hour per gm. dry weight (Figs. 6 and 7). The mean

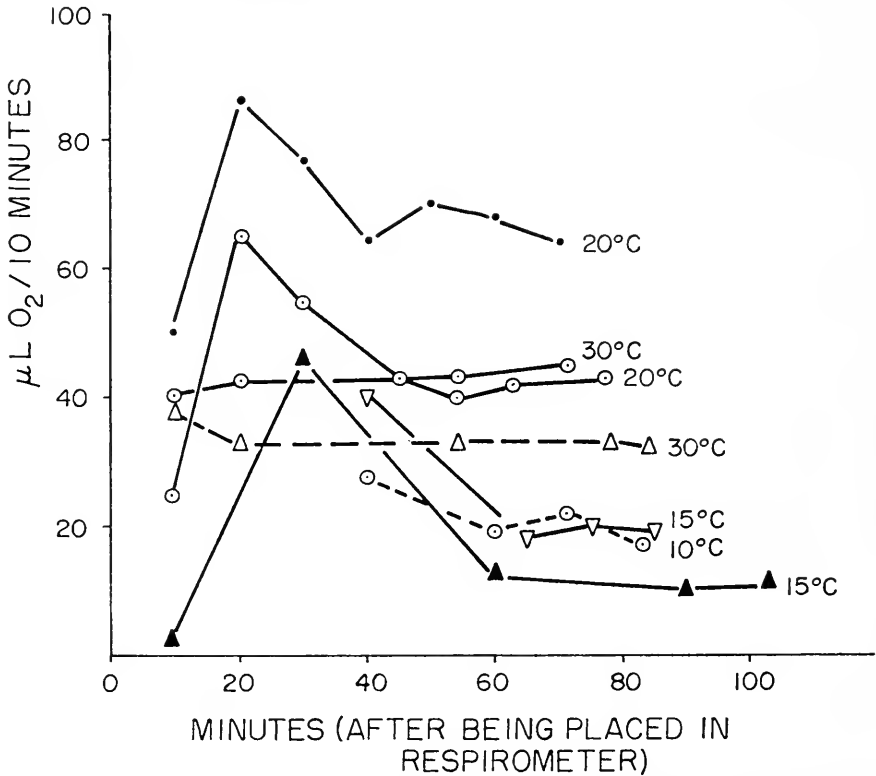


FIGURE 5. The immediate effect of temperature change on the oxygen uptake of *Eucidaris tribuloides*. Urchins were moved from ambient water with a temperature of 28–29° C. to water in the respirometer at different temperatures.

oxygen consumption for each temperature and the Q_{10} for each 5° C. interval between 10° and 30° C. were then calculated. Both starved and well-fed urchins were used (but these were considered separately).

The stabilized rate of oxygen consumption of *Eucidaris tribuloides* at different temperatures was measured in both summer and winter to determine if there were seasonal acclimation. The log of the mean oxygen consumption for each of the temperatures given above was plotted following the method given by Prosser and Brown (1961), and the degree of translation and rotation of the resulting curves for winter and summer was described (Figs. 6 and 7).

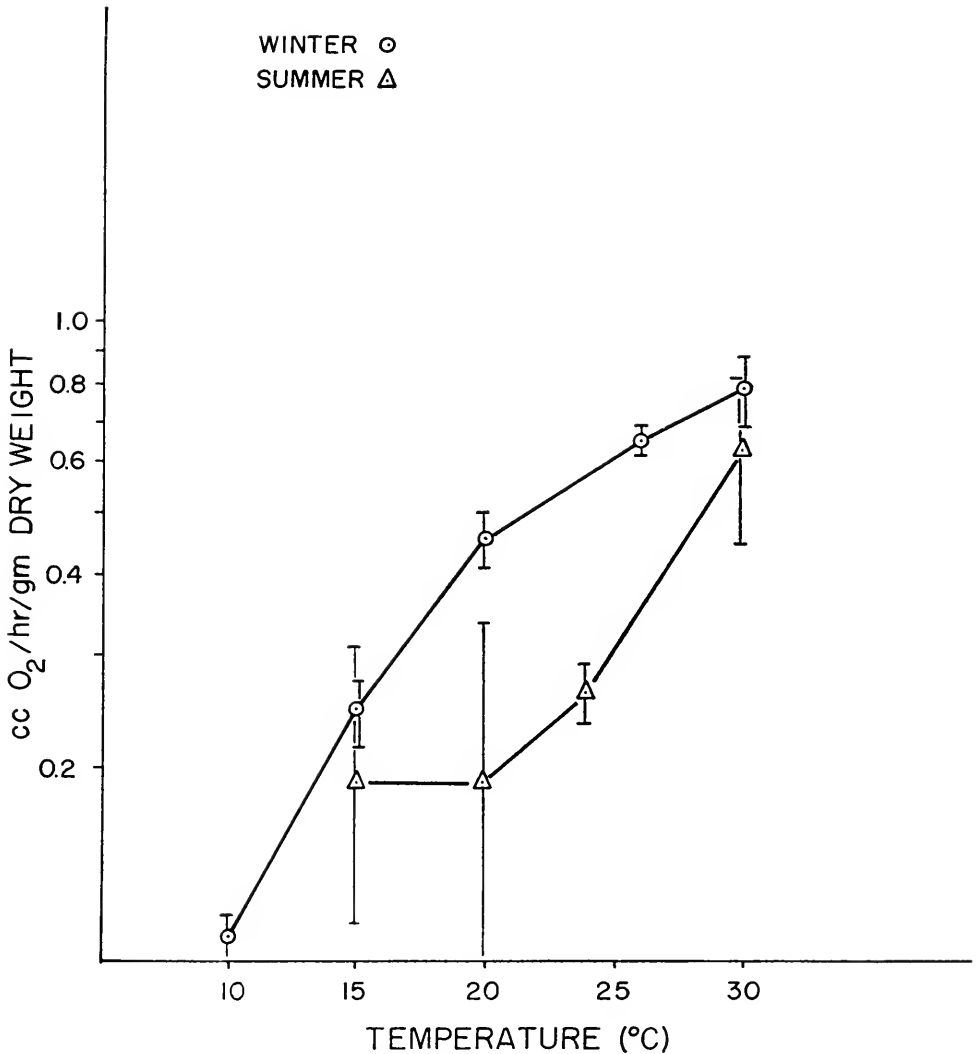


FIGURE 6. Mean oxygen uptake of individuals of *Eucidaris tribuloides* at stabilized rates for different temperatures. Urchins well-fed. The vertical lines give the 95% confidence intervals.

The Q_{10} of well-fed urchins increased rapidly in summer and decreased in winter as the temperature was increased from 15° to 30° C. The Q_{10} of starved urchins over a similar temperature range was more variable and manifested less overall change than that of the well fed individuals. In summer it increased 1.1 units in the temperature range 15° to 30° C. as compared with 3.4 units for the well-fed urchins. In winter it was essentially the same at high and low temperatures (Table III). These facts indicate that in both summer and winter well-fed

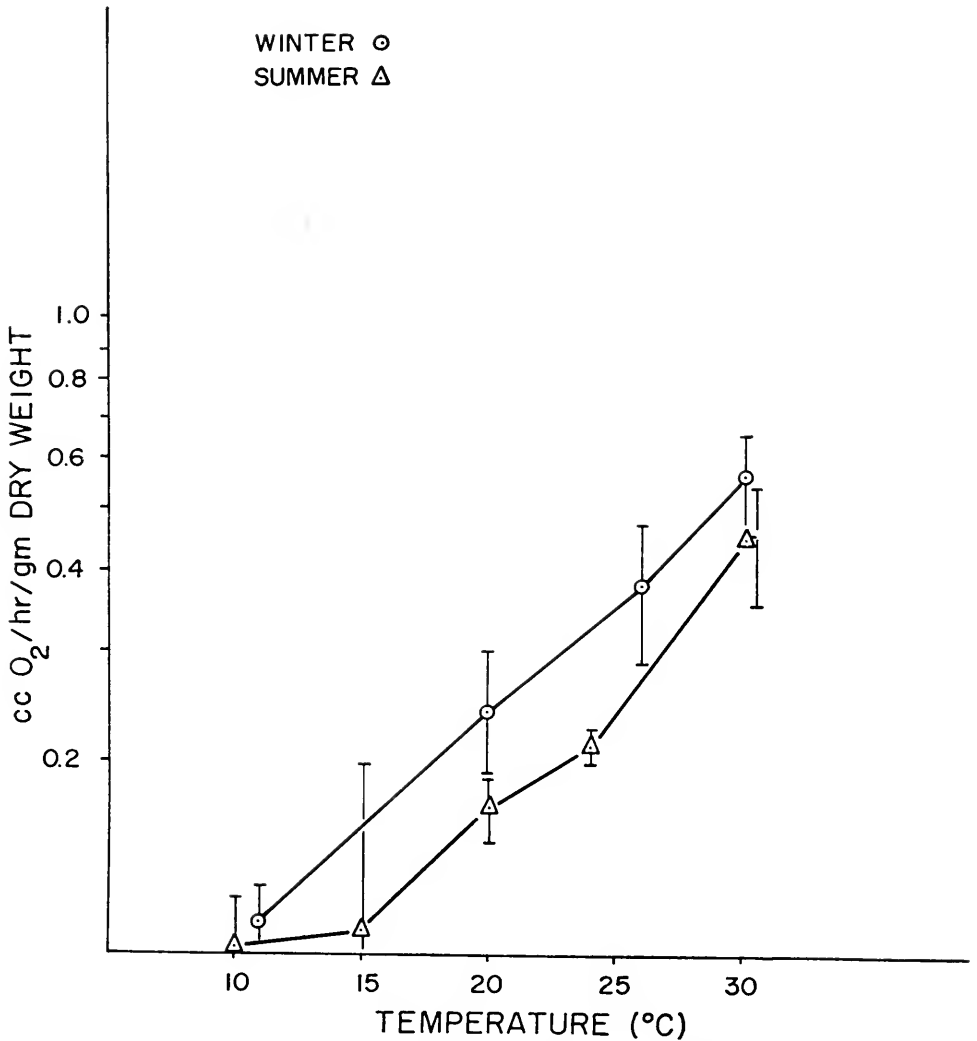


FIGURE 7. Mean oxygen uptake of individuals of *Eucidaris tribuloides* at stabilized rates for different temperatures. Urchins starved. The vertical lines give the 95% confidence intervals.

urchins were probably more sensitive to temperature changes in their stabilized respiratory rates than were starved urchins.

Both starved and well-fed urchins exhibited partial seasonal acclimation in which oxygen consumption curves were well above summer curves at the corresponding temperatures. The acclimation was not complete since the mean summer oxygen consumption was higher at the summer ambient than the mean winter oxygen consumption at the winter ambient. The difference between the mean summer and winter oxygen consumption curves of the starved urchins was less

than that between the mean summer and winter curves of the well-fed urchins. This suggests that the process of feeding or digestion may play an important part in acclimation over normal environmental temperature ranges.

At 30° C., which is close to the summer ambient, the oxygen consumption curves tended to converge while at the winter ambient of 20° C. they were widely separated. This pattern resembled what Prosser and Brown (1961) have called translation and clockwise rotation (convergence at higher temperatures). At temperatures below the winter ambient there was a second convergence of oxygen consumption curves, so that around 10° C. there was little difference in the O₂ consumption of winter or summer urchins (no acclimation).

TABLE III

*Q*₁₀ of well-fed and starved urchins (*Eucidaris tribuloides*) in winter and summer. Determined from changes in the mean oxygen uptake given in Figures 6 and 7

Well-fed		Starved	
Temperature, ° C.	<i>Q</i> ₁₀	Temperature, ° C.	<i>Q</i> ₁₀
Summer		Summer	
15-20	1.0	10-15	1.2
20-35	2.2	15-20	2.4
25-30	4.4	20-25	1.7
		25-30	3.5
Winter		Winter	
10-15	5.2	11-15	2.5
15-20	3.4	15-20	2.1
20-25	1.8	20-25	2.3
25-30	1.6	25-30	2.7

DISCUSSION

Eucidaris tribuloides feeds on hard particles of limestone rock deriving nourishment from associated organisms (McPherson, in press). Since boring and encrusting sponges are prevalent on rocks in tropical seas these might well be important in the nutrition of this urchin. The preference of *E. tribuloides* in aquaria for the boring sponge *Cliona lampha* might be an indication of the importance of sponge in the diet of this species.

The relatively high rate of feeding observed in the laboratory indicates that *Eucidaris tribuloides* is a potential erosive agent. Urchins apparently fed throughout the year as indicated by the presence of food in individuals collected monthly in the field, and by observations of feeding in aquaria throughout the year (McPherson, in press). Urchins, of the mean size of those found on coral reefs, consumed about one to two gm. of wet carbonate rock per day in laboratory experiments. The feeding rate in the field, however, is probably less than that measured in the laboratory, as indicated by a slower growth rate in the former area than in the latter (McPherson, in press). Lewis (1964) stated that carbonate ingestion by *Diadema antillarum* may be of some importance in the process of erosion of certain zones of the coral reef. In this habitat he determined that *D. antillarum* ingested a mean weight of at least two gm. of carbonate per day. This intake in

an area of high density, such as that reported by Randall *et al.* (1964) of 13.4 urchins per square meter off the Virgin Islands, would result in the removal of a considerable amount of coral rock. Erosion caused by *E. tribuloides*, however, would probably be much less than that caused by *D. antillarum* since the density and feeding rate of the former are probably much lower than that of the latter. McPherson (in press) reported a maximum mean density for *E. tribuloides* of two per square meter on French Reef, Florida. In other places off southeastern Florida densities were much less than that observed on this reef.

Under the experimental conditions used in this study food, size of organism, and temperature were important factors influencing the oxygen uptake of *Eucidaris tribuloides*. This is not surprising since these factors are all generally important in the metabolism of invertebrates (Zeuthen, 1947). It is interesting, however, that *E. tribuloides* does manifest some control over its oxygen uptake when temperature changes. Sudden changes in temperature resulted in a rather unstable rate of oxygen consumption, but this only lasted some minutes, and in an hour the rate of respiration became relatively stable. Slow changes in temperature, such as those which occur seasonally, had only a slight effect on respiration since the urchins partially acclimated during this time. Partial control of respiration would allow these animals to maintain some independence from environmental temperature.

Farmanfarmaian and Giese (1963) found that *Strongylocentrotus purpuratus* from ambient water of 14–19° acclimated to water at 5° C. The oxygen consumption curves were similar to those described for seasonal acclimation of *Eucidaris tribuloides* with the cold-acclimated urchins having rates above the warmer living individuals (translation). The two oxygen consumption curves, for acclimated and non-acclimated urchins, also tended to converge at higher temperatures (clockwise rotation) as did those of *E. tribuloides*. Boolootian and Cantor (1965) reported that *Arbacia punctulata* acclimated to low temperatures did not have a higher O₂ consumption than urchins maintained at higher temperatures. They stated that this suggested that *Arbacia* and *Strongylocentrotus* have different physiological mechanisms for thermal acclimation.

There was little variation in oxygen uptake between individuals of *Eucidaris tribuloides* at temperatures well below their winter ambient (10° and 15° C.). Even urchins tested in different seasons and in different nutritional conditions had roughly the same oxygen consumption for that particular temperature used in the experiment. Processes involved in nutrition and acclimation presumably had ceased or decreased to such low levels that their effect on respiration was negligible.

SUMMARY

1. In the laboratory *Eucidaris tribuloides* showed a food preference for rock infested with the boring sponge *Cliona lampha* over turtle grass (*Thalassia testudinum*), several algae, and fish strips.

2. In the laboratory, urchins consumed wet rock infested with *Cliona lampha* at rates ranging from less than one to over three gm. per day depending on the size of the urchin.

3. Oxygen consumption of urchins was measured in the laboratory with a modified Scholander volumetric respirometer.

4. Oxygen uptake decreased significantly during the first day of starvation, but then remained relatively stable during several successive days of starvation.

5. There was not a significant correlation between gonad index or sex of the urchin and relative oxygen uptake.

6. Relative oxygen consumption decreased with increasing size of the urchin. Values for "b" indicated an oxygen uptake which was almost proportional to surface area at 20° and 30° C.

7. Oxygen uptake fluctuated during the first hour when urchins were placed in water several degrees colder than their ambient water. At the end of this hour, however, the rate of oxygen uptake had become relatively stable. Well-fed urchins manifested greater changes in their stabilized rates of oxygen uptake (Q_{10}) at different temperatures than did starved urchins.

8. Urchins manifested partial seasonal acclimation of oxygen uptake as indicated by the fact that winter oxygen consumption curves were well above summer curves at the corresponding temperatures. The acclimation was probably not complete since mean summer oxygen uptake was higher at the summer ambient than the mean winter oxygen uptake at the winter ambient.

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DYNAMICS OF ECDYSONE SECRETION AND ACTION IN THE FLESHFLY *SARCOPHAGA PEREGRINA*¹

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Ecdysone, the growth and molting hormone of insects, provokes a developmental response when it accumulates to "threshold titer" at certain "critical periods." Meanwhile, the titer of juvenile hormone dictates whether the developmental response will be molting or metamorphosis. These two generalizations epitomize the prevailing theory of ecdysone's action.

In support of the theory there is a wealth of evidence that the secretion of ecdysone is necessary for molting or metamorphosis. But whether the developmental response is triggered by a certain threshold titer of ecdysone is by no means assured. There remains the possibility that ecdysone is promptly and progressively utilized and inactivated after its release into the blood. Under that circumstance, the effects of the hormone might be cumulative within the target-organs without any marked change in its concentration in the blood or the insect as a whole.

Precisely this state-of-affairs appears to be the case when ecdysone provokes puparium formation of mature larvae of the fleshfly, *Sarcophaga peregrina*. In the present report we document the swift inactivation of ecdysone after its secretion into the blood. Moreover, we show that the overt developmental response, puparium formation, is triggered, not by the accumulation of ecdysone itself, but, rather, by a summation of the latent, covert effects of the hormone.

MATERIALS AND METHODS

1. *Experimental animals*

All experiments were carried out on larvae of *Sarcophaga peregrina*. This species was chosen for study because of its large size and the ease with which one can obtain unlimited numbers of carefully timed larvae. Thus, as documented by Ohtaki (1966) and diagrammed in Figure 1, the activation of the endocrine system is opposed or prevented as long as mature larvae are stored in contact with water. This inhibition is relieved when "wet larvae" are transferred to dry conditions. Therefore, the secretion of ecdysone can be timed from the "0-hour of dryness" until the initiation of puparium formation which, at 25° C, routinely takes place 16–17 hours later. Meanwhile, at any time during this period one

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can obtain homogeneous groups of larvae by timing their exposure to dry conditions.

All the larvae used in the present study were reared in gallon glass jars containing pig liver. After three days of feeding at $24.5 \pm 1.5^\circ \text{C}$, the mature larvae emptied their crops and tried to crawl away from the food. For two further days they were stored in the same jars, to which water was added to maintain wet conditions. Then they were collected, washed in tap water, and placed in clean gallon jars containing sufficient water to cover the bottom. After two further days of storage at 24.5°C , the wet larvae were blotted with filter paper and either used immediately as "0-hour larvae" or aged in dry jars for two or more hours.

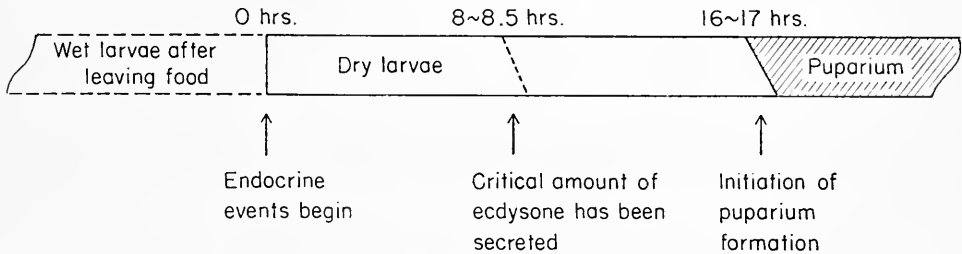


FIGURE 1. The timing of developmental events when mature larvae of *Sarcophaga peregrina* are transferred from wet to dry conditions at 25°C .

2. The *Sarcophaga* assay

After exposure to dry conditions for 0, 2, 4, or 6 hours, homogeneous groups of larvae were ligated just behind the sixth segment. The majority of assays were performed on "standard test abdomens" which were prepared as follows:

Larvae were ligated after exposure to dry conditions for six hours. Twenty-four hours later, individuals showing puparium formation in front of, but not behind, the ligature were collected and aged an additional 16 hours under dry conditions. Larvae showing any trace of puparium formation behind the ligature were discarded; the number that did so ranged from 1 to 10%. The others were immediately used as "standard test abdomens."

Each assay was performed on a homogeneous group of 10 to 20 abdomens. The body anterior to the ligature was cut away and $10 \mu\text{l}$ of the test solution were injected by means of a 30-gauge glass or stainless steel needle sealed to a $100\text{-}\mu\text{l}$ microsyringe; the latter was controlled by an Agla micrometer. The needle was inserted through a loose ligature which was tightened immediately after the injection. When tests were performed on entire larvae, the injection was made through a ligature placed across the anterior tip just behind the mouthparts.

3. The "puparium index"

All assays were controlled 24 hours after injection. As recommended by Karlson (1956), each abdomen was scored as having undergone complete, marked, slight, or no puparium formation. These reactions were equated to 100, 75, 50, and 0%, respectively, for the purpose of calculating the "puparium index."

4. *Ecdysone solutions*

Synthetic α -ecdysone was obtained from Drs. John Fried and John Siddall of the Syntex Research Laboratory. The crystalline hormone was weighed, dissolved in 1 part warm isopropanol, and diluted with 9 parts water. Twofold serial dilutions were prepared in 10% isopropanol. One series of experiments made use of the phytoecdysones, β -ecdysone and ponasterone A, obtained through the courtesy of Profs. K. Nakanishi and T. Takemoto, of Tohoku University.

5. *Collection and bioassay of larval blood*

By means of iridectomy scissors a small sliver of integument was cut from the dorsal tips of the abdomens of a homogeneous group of ten washed larvae, care being taken not to damage the gut or other viscera. The larvae were immediately placed in a tilted Petri dish into which the blood was expressed by the spontaneous contractions of the cut larvae. The blood was immediately drawn into a 100- μ l syringe and 10 μ l injected into each of a series of test abdomens.

6. *Extraction and concentration of ecdysone from the blood*

The method was a modification of that described by Kaplanis *et al* (1966). Blood was collected from ten larvae, as just described. The volume was recorded and the blood immediately ejected into a conical centrifuge tube containing 4 ml of a 1:1 mixture of acetone and absolute ethanol. In this same manner additional blood was collected until a total of 1.5 ml had been obtained from a homogeneous group of approximately 100 individuals. The mixture was stirred at room temperature and then centrifuged at 1000 *g* for five minutes. The supernatant was decanted and the precipitate extracted twice more with 2-ml volumes of the solvent mixture. The supernatants were pooled and filtered with suction through a sintered medium-pore filter into a 50-ml round-bottom flask. The filter was rinsed with the solvent mixture and the extract reduced to dryness *in vacuo* on a rotary evaporator with temperatures rising to 50° C.

Crude extracts of this sort are extremely toxic when injected into larvae. Therefore, further purification was necessary. To this end, the contents of the flask were rinsed with 70% methanol into a small separatory funnel and washed with an equal volume of petroleum ether (*b.p.* 37–60° C). The methanolic hypophase was drained into a small round-bottom flask and reduced to dryness at 50° C as just described. By means of *n*-butanol the contents were rinsed into a separatory funnel and an equal volume of aqueous 2% sodium carbonate was added. The mixture was stirred, centrifuged, and the butanolic hypophase collected. The aqueous epiphase was twice re-extracted with 0.5 volume of butanol. The alcoholic phases were combined and washed twice with 0.5 volume of water. The aqueous phases were collected and re-extracted with 0.25 volume butanol and the latter washed with 0.5 volume of water. The butanolic phases were once again combined, placed in a round-bottom flask, and reduced to dryness on a rotary evaporator with temperatures rising to 65° C. With small volumes of methanol the contents of the flask were rinsed into a vial and the solvent evaporated in a stream of nitrogen. The extracted material was redissolved in 0.15 ml water, corresponding to one-tenth the original blood volume. This tenfold

concentrate was assayed by the injection of 10 μ l into each of a series of isolated abdomens.

By the addition of known amounts of α -ecdysone the combined efficiencies of the extraction and purification procedures were found to be 90%. Therefore, in the present report, all extractions (except where otherwise noted) have been corrected for 10% loss.

7. *Extraction and concentration of ecdysone from the entire animal*

One to four larvae were homogenized in Ringer's solution and extracted with 4 ml of a 1:1 mixture of acetone and absolute ethanol. The mixture was centrifuged and the precipitate washed twice with a total of 4 ml of the solvent mixture. The supernatants were combined and the ecdysone extracted as described above for blood.

To harvest any ecdysone that might be complexed to proteins, the precipitate was reduced to dryness and treated with "Pronase." The reaction mixture consisted of 1 ml of 0.04 *M* Tris buffer (pH 7.5), 0.0066 *M* calcium chloride, and 1 ml of a 0.15% aqueous solution of the enzyme. After incubation at 40° C for 24 hours, the mixture was centrifuged and the supernatant decanted. The precipitate was twice washed with acetone-ethanol and the supernatants combined. The latter was purified and assayed as described above for blood. Additional extractions were also carried out on anterior and posterior ends of ligated larvae.

RESULTS

1. *The effects of massive injections of ecdysone*

As illustrated in Figure 1, 50% of larval abdomens undergo puparium formation when isolated after 8.5 hours of dryness. However, they require an additional 8 hours to do so. The significance of this latent period was examined in groups of 0-hour larvae injected with α -ecdysone, β -ecdysone, or ponasterone A. As summarized in Table I, even the largest doses failed to reduce the latent period to less than 8.5 hours.

2. *Ecdysone secretion during exposure to dry conditions*

Graded doses of α -ecdysone were injected into abdomens ligated after 0, 2, 4, and 6 hours of exposure to dry conditions. Each dose was tested in a homogeneous group of 20 abdomens. As shown in Figure 2A, the longer the preliminary exposure to dryness, the more sensitive were the abdomens to the injected ecdysone.

In Figure 2B the critical doses provoking a puparium index of 50% have been plotted as a function of the hours of exposure to dry conditions prior to ligation. These values define a *Sarcophaga* unit for abdomens isolated after 0, 2, 4, and 6 hours, respectively. Sensitivity to injected ecdysone increases at a rate that may be equated to about 0.007 μ g per hour. It will be recalled that when ligation is postponed to the 8–8.5 hour, 50% of abdomens undergo puparium formation without any injection. Taken as a whole, these findings suggest that, at the outset of exposure to dry conditions, the endocrine system is activated and the ring-glands initiate the secretion of ecdysone at a fairly steady rate of about 0.01 μ g per hour.

On the basis of Figures 2A and 2B one might suppose that abdomens isolated after 6 hours of dryness would be optimal for the biological assay. Unfortunately, this is not the case because up to 10% of the "6-hour abdomens" give false positive assays—*i.e.*, they undergo puparium formation without any injection.

TABLE I
Acceleration of puparium formation by injection of ecdysones into 0-hour larvae

Substance injected	Amount (μg)	Number injected	Time for 50% to initiate puparium formation (hrs)
Controls (solvent only)			
H ₂ O		20	17
10% Isopropanol		20	21
α -Ecdysone*	0.5	10	8.5
	1	10	8.5
	2	10	8.5
β -Ecdysone*	0.25	10	8.8
	0.5	10	9
	1	10	9
	2	10	9
Ponasterone A**	0.5	10	8.5
	0.7	10	8.5

* In 5 μl H₂O.

** In 5 μl 10% isopropanol.

Figure 2A also records the sensitivity of "standard test abdomens" which, as described above, are prepared by the further ageing of abdomens ligated after 6 hours of dryness. It will be observed that 95% of these abdomens have become less sensitive than at the time of their isolation. However, about 5% retain great sensitivity without showing any false positive assays. A puparium index of 50% is provoked by the injection of 0.035 μg α -ecdysone, which we define as a "*Sarcophaga* unit" for standard test abdomens (Ohtaki *et al.*, 1967). This is the same average value that characterizes abdomens isolated and immediately used after 2 hours of dryness.

3. Ecdysone assays of the blood

On the basis of the foregoing results as well as Fraenkel's (1935) classic study of *Calliphora*, we anticipated that detectable amounts of ecdysone would be demonstrable in the blood. This prospect was tested by injecting standard test abdomens with blood obtained from donors at four successive stages—namely, larvae after 0, 6, and 12 hours of dryness; and 16-hour larvae showing the initiation of puparium formation. Each standard test abdomen received 10 μl of donor blood.

All assays were uniformly negative. So, by reference to Figure 2A, it appears that at no stage in puparium formation is as much as 0.0013 μg α -ecdysone present in 10 μl of blood. And since a mature larva contains 35 μl blood, the

total ecdysone in the entire blood could scarcely exceed about 0.005 μg , *i.e.*, 15% of a *Sarcophaga* unit for standard test abdomens.

4. Extraction of blood ecdysone

In order to bring this low titer within the range of the biological assay, we undertook the extraction and partial purification of blood ecdysone as described under Methods. The extract was redissolved in distilled water as a tenfold concentrate of the original blood, and 10 μl were injected into each of a series of standard test abdomens.

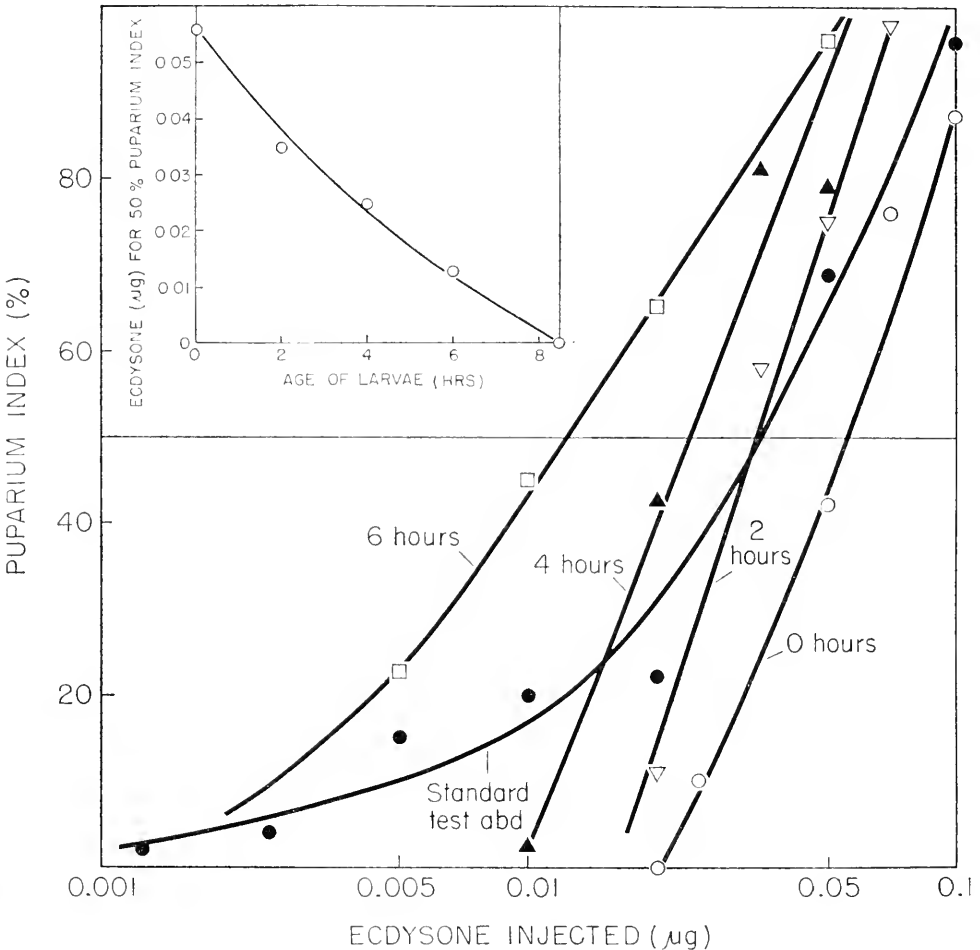


FIGURE 2A. The developmental responses (computed in terms of the puparium index) are plotted as a function of the amount of injected α -ecdysone. The curves correspond to abdomens ligatured after the indicated hours of exposure to dry conditions. For description of "standard test abdomens," see text.

FIGURE 2B. The inset in the upper left corner records the amount of injected ecdysone required for a 50% response of each type of abdomen.

Three concentrates of the blood of 0-hour larvae were assayed on a total of 41 standard test abdomens with the following result: 2 abdomens showed a trace of puparium formation; the other 39 were negative. This result corresponds to a puparium index of about 2%—a value which, as indicated in Figure 2A, is at the extreme limit of the calibration curve for standard test abdomens. Consequently, it appears that about 0.001 μg ecdysone is present in 10 μl of blood concentrate. On the assumption of 90% efficiency of the extraction procedure, we estimate that a total of 0.0004 μg ecdysone is present in the 35 μl of blood of a 0-hour larva, corresponding to about 1% of a *Sarcophaga* unit for standard test abdomens (see Table II).

The experiment was repeated on concentrates prepared from the blood of larvae at the 8th hour of exposure to dry conditions; 27 of 37 test abdomens showed no detectable reaction; 10 showed traces of puparium formation. This corresponds to a puparium index of 13% which, by reference to Figure 2A and on the assumption of 90% recovery, indicates the presence of about 0.0016 μg ecdysone in the entire blood of an 8-hour larva; *i.e.*, about 5% of a *Sarcophaga* unit (Table II).

TABLE II
Extraction of ecdysone from Sarcophaga peregrina

Stage	Ecdysone in:	
	Entire blood (μg)	Entire animal (μg)
0-hour larva	0.0004 (1%)*	0.0013 (4%)
8-hour larva	0.0016 (5%)	0.0025 (7%)
17-hour (white puparium)	0.005 (15%)	0.02 (57%)

* The bracketed figures record the corresponding percentage of a *Sarcophaga* unit as ascertained in assays performed on standard test abdomens.

In like manner the blood was collected from 100 individuals which had formed white puparia after 17 hours of exposure to dry conditions. Assays of the concentrated extract gave a puparium index of 34%, corresponding to about 0.005 μg α -ecdysone (15% of a *Sarcophaga* unit). This value is in accord with the upper limit of blood ecdysone calculated above in Section 3.

5. Ecdysone in the entire larva

The presence of so little ecdysone in the blood suggested that the hormone might be bound to the tissues. To test this possibility entire larvae were homogenized and extracted. The results summarized in Table II show that only about 4% of a *Sarcophaga* unit is present in an entire 0-hour larva, including the ring-gland itself. Even more surprising is the finding that only about 7% of a unit is present in an 8-hour larva, *i.e.*, at a stage where 50% of individuals have received all the ecdysone they require for puparium formation. The highest titer is found in white puparia where each individual contains about 57% of a *Sarcophaga* unit.

6. Recovery of injected ecdysone from the blood

Known amounts of α -ecdysone were injected into 0-hour larvae. Thereafter, at regular intervals, groups of injected larvae were sacrificed and their blood collected and assayed by injecting 5 μ l into standard test abdomens. The puparium index was calculated in each case and equated to absolute units of α -ecdysone by reference to Figure 2A. The percentage of the injected ecdysone remaining in the blood was then calculated for each group.

The results summarized in Table III demonstrate that more than half of the injected hormone disappears from the blood within the first hour and that only traces remain after 8 hours.

TABLE III
Biological assays of blood obtained from larvae injected with α -ecdysone*

Ecdysone injected into blood donors (μ g)	Time between injection and blood collection (hrs)	Number of test abdomens	Puparium index (%)	Equivalent ecdysone (μ g)	% remaining in blood
2	1	10	100	>0.1	>35
	3	10	100	>0.1	>35
	5	20	62	0.047	16
	8	20	10	0.006	2
1	1	20	69	0.05	35
	3	20	23	0.015	11
	8	20	0	<0.001	<1
0.5	1	20	16	0.01	14
	3	20	8	0.004	6
	8	20	2	0.001	1

* Each standard test abdomen received 5 μ l blood from donor larvae.

7. Recovery of injected ecdysone from entire larvae

One μ g of α -ecdysone was injected into a series of 0-hour larvae. After exposure to room temperature for 0.1, 1, 3, or 8 hours, the entire larvae were homogenized and their ecdysone extracted and assayed in the usual way. Ninety per cent of the injected ecdysone was recovered from larvae extracted immediately after injection thereby testifying to the efficiency of the extraction and purification procedure. After correcting for this 10% loss, we found that one hour after the injection, only 50% was recovered; after three hours, 23%; and after eight hours, 2%.

The experiment was repeated except in this case the injected larvae were separated into blood and carcass. The blood was assayed by injecting 5 μ l into standard test abdomens. The carcasses were cut open, rinsed in several changes of Ringer's solution, and then extracted and assayed in the usual manner.

The results summarized in Table IV once again document the rapid disappearance of injected ecdysone. And, here again, as in Table III, we note that the lower the dose, the more rapid is the relative rate of disappearance.

We entertained the possibility that ecdysone might be progressively bound to cellular proteins and in this manner rendered insoluble in the acetone-ethanol extraction mixture. To examine this possibility, 1 μg α -ecdysone was injected into each of ten 0-hour larvae. After three hours at room temperature, pairs of larvae were homogenized and five extracts prepared in acetone-ethanol. The precipitates were suspended in buffer and treated with "Pronase" as described under Methods. After incubation at 40° C for 24 hours, the reaction mixtures were once again extracted with acetone-ethanol to harvest any ecdysone that had been bound to proteins. These latter extracts were purified and assayed in the usual manner in a total of 54 standard test abdomens. All tests were uniformly negative.

TABLE IV
Recovery of injected α -ecdysone from blood and carcass of 1- to 8-hour larvae

Ecdysone injected (μg)	Time from injection (hr)	Number of animals	% of injected ecdysone recovered in		% not recovered
			Blood	Carcass*	
1.0	1	1	35	16	49
	3	2	11	13	76
	8	4	<1	<1	ca 99
0.5	1	1	14	9	77
	3	3	<3	<3	ca 95
	8	4	<1	<1	ca 99

* Corrected for 10% loss in the extraction procedure. Each experiment was performed in triplicate on the indicated number of animals. The average results are cited.

In control experiments, Pronase was found to have no deleterious effects on ecdysone. For example, two homogenates were prepared from uninjected larvae and 1 μg of α -ecdysone was added to each homogenate just prior to Pronase incubation. After 24 hours at 40° C, the reaction mixture was extracted with acetone-ethanol and purified in the usual way. The biological assays indicated the recovery of 100% of the added ecdysone.

For these several reasons the swift disappearance of injected ecdysone is accountable only in terms of its equally swift inactivation within the living insect.

8. *Stability of injected ecdysone at low temperature*

Three 0-hour larvae were placed at 2° C for 24 hours and then injected with 1 μg α -ecdysone. After an additional 24 hours at the low temperature, the larvae were homogenized and extracted. Biological assays of the extract showed the recovery of 100% of the injected hormone. Clearly, the inactivation of ecdysone is blocked by low temperature.

9. *Stability of injected ecdysone in the absence of oxygen*

Nine 0-hour larvae were placed in a covered Buchner funnel and exposed to a stream of nitrogen for 1 hour. Each individual was then injected with 1 μg

α -ecdysone. After three additional hours in nitrogen, the entire larvae were homogenized and extracted. Assays showed the recovery of 90% of the injected ecdysone. Control experiments performed on entire larvae exposed to air showed the recovery of only 20% of the injected ecdysone. In additional control experiments, 0-hour larvae underwent full and complete recovery after 4 hours of exposure to nitrogen.

10. *Inactivation of ecdysone by larval fragments*

The gut was removed from four 0-hour larvae. These larvae were then cut into small fragments and placed in 2 ml Ringer's solution containing 1 mg streptomycin sulfate. The mixture was subdivided between two small flasks, one of which contained 3 mg of the potent anti-tyrosinase, phenylthiourea (PTU). One μ g of α -ecdysone was added to each flask and the latter incubated at 25° C on a gyratory table. After 24 hours the ecdysone was extracted and assayed. The recovery was 40% in the mixture containing PTU and 22% in the mixture not containing PTU.

The experiment was repeated using a shorter incubation time and cut-up larvae from which the gut was not removed. After 8 hours, 94% of the added ecdysone was recovered from the mixture containing PTU and 70% from the mixture not containing PTU.

11. *Experiments on homogenates and subcellular fractions*

When the just-mentioned experiment was repeated using crude homogenates prepared from 0-hour larvae, 100% of the added ecdysone was recovered from all the reaction mixtures after 24 hours of incubation. The same was true when ecdysone was incubated with blood collected from 0-hour larvae.

On the assumption that ecdysone inactivation might be blocked by the presence of one or more endogenous inhibitors, the crude homogenate was fractionated prior to testing. To this end, two 0-hour larvae were cut into fragments and rinsed for 2 minutes in 2 ml 0.01 M HEK buffer (pH 7.4) made up in 20% sucrose (Kafatos, 1968, page 1258). The fragments were collected and homogenized in 2 ml of sucrose-free HEK buffer. The homogenate was centrifuged at 1000 *g* for 3 minutes. The supernatant was collected and centrifuged for 50 minutes at 200,000 *g*, and the particulate fraction was combined with the low-speed sediment. The insoluble fraction was diluted with 2 ml Ringer's solution. One mg streptomycin and 1 μ g α -ecdysone were added to each reaction mixture and the latter incubated at 25° C on a gyratory table. Eight hours later, the ecdysone was extracted and assayed. Recovery of added hormone was 100% for both reaction mixtures. The experiment was twice repeated with the same result. Thus, as in the case of crude homogenates, ecdysone inactivation was not brought about by any of the subcellular fractions. This same negative result was observed in additional experiments in which 1 mg/ml of NAD or NADH was added to the reaction mixture prior to incubation.

DISCUSSION

1. *Ecdysone titer in relation to puparium formation*

At the outset of exposure to dry conditions a 0-hour larva contains 1.3 nanograms of ecdysone, most of which may be sequestered in its ring-gland (Table II). Fifty per cent of abdomens isolated at this time undergo puparium formation when injected with 55 nanograms of ecdysone (Fig. 2). Since the abdomen makes up 55% of larval mass, we calculate that an entire larva deprived of its ring-gland would require the injection of approximately 100 nanograms. When the ligation is postponed until the 8th hour of dryness, the ring-gland has already secreted sufficient ecdysone to cause 50% of abdomens to form puparia without any injection. Despite this fact, the amount of ecdysone in an 8-hour larva is only 2.5 nanograms (Table II). No less than 97 nanograms of ecdysone seem to have disappeared.

This same paradox is seen in the extractions and assays which Shaaya and Karlson (1965) and Shaaya (1967) have carried out on entire *Calliphora* larvae. From their data we calculate that at no stage prior to puparium formation does the titer of ecdysone exceed 26% of a *Calliphora* unit.

2. *The half-life of injected ecdysone*

All our experiments direct attention to the lability of α -ecdysone after its injection or its secretion by the ring-gland. Thus, as we have seen, within 1 hour after the injection of 1 μ g, 50% had been inactivated; and after 8 hours, about 98% had been inactivated. The rate of inactivation was even higher in larvae injected with 0.5 μ g. Therefore, for mature larvae of *Sarcophaga peregrina* at 25° C, we can state that the time for half-inactivation of α -ecdysone is less than 1 hour.

3. *The inactivating mechanism*

As described in Section 11 of Results, no trace of inactivation could be detected when α -ecdysone was incubated for 24 hours with larval blood or with crude homogenates prepared from 0-hour larvae. The same negative results were observed in tests performed on the soluble and the particulate fractions. By contrast, clear-cut inactivation occurred when α -ecdysone was incubated with larval fragments; *i.e.*, under conditions where the integrity of the cells was largely preserved. This demonstrates that the inactivating mechanism is present in at least some and perhaps most cells. As mentioned in Section 10 of Results, inactivation was brought about by larval fragments, irrespective of whether the gut was present or absent. Moreover, in additional experiments not mentioned above, inactivation of injected ecdysone took place in both anterior and posterior parts of ligated larvae. Inactivation by the anterior end was substantially faster than by the posterior end.

Our inability to demonstrate the inactivating mechanism in crude or purified homogenates implies the lability of the system or the requirement of co-factors and incubation conditions that were not satisfied. In Sections 8 and 9 of Results we observed that inactivation is blocked in larvae exposed to low temperature or to anaerobic conditions. Therefore, chemical and, more particularly, oxidative processes appear to be principally concerned.

4. *The dynamics of ecdysone action*

The rapid inactivation of ecdysone, demonstrated for the first time in the present study, directs attention to the accumulation, not of the hormone itself, but of the covert effects of the hormone. This new interpretation is summarized in the flow-sheet diagrammed in Figure 3. Ecdysone is secreted into the blood, taken up by the tissues, and reacts with the hormonal receptors to exert its primary action, whatever that may be. As a result of this primary reaction, a concatenation of biochemical and biophysical events is set in motion which comprise the covert effects of the hormone. The latter undergo, as it were, spatial and temporal summation within the target organs and finally trigger the overt effects, *i.e.*, the initiation of molting or metamorphosis. Meanwhile it may be noted that the covert

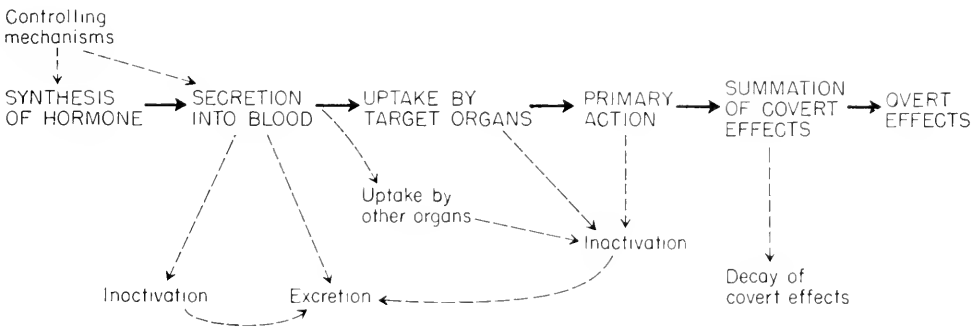


FIGURE 3. A revised theory of the hormonal action emphasizing the accumulation of the covert effects of ecdysone.

effects are subject to decay if their temporal accumulation is interrupted short of the discharge level. This fact is documented in Figure 2A in terms of the lessened sensitivity to ecdysone injection of standard-test abdomens as compared to 6-hour abdomens from which they were derived. Moreover, the accumulation and implementation of the covert effects require a certain finite time. Thus, as shown in Table I, even the largest doses of α -ecdysone, β -ecdysone, or ponasterone A required at least 8.5 hours to provoke the initiation of puparium formation. This minimal "latent period" apparently corresponds to the time required for the covert effects of ecdysone to accumulate to the discharge level.

It has not escaped our attention that this conception of the dynamics of ecdysone action may be applicable to hormones in general.

5. *Transfusion versus parabiosis*

The literature pertaining to insect endocrinology is replete with carefully controlled experiments in which hormonal effects have been conveyed between donors and recipients by joining them in parabiosis. By contrast, there are few claims to accomplishing this same goal by the injection or transfusion of blood. One of these rare claims ranks among the most famous experiments in the history of insect physiology; namely, Fraenkel's (1935) induction of puparium formation by the injection of "pupation blood."

During the past 33 years the literature is silent in confirming or refuting this historic experiment. Now, for reasons set forth in the present study, it appears that Fraenkel's results were "false positives" such as were described above in Section 2 of Results. This seems all the more likely since Fraenkel's experiments were not controlled by the injection of any materials other than "pupation blood." Needless to say, our very ability to resolve this paradox owes a great deal to Fraenkel's insightful pioneering work.

We gratefully acknowledge the critical reading of the manuscript by Professor Lynn M. Riddiford.

SUMMARY

1. Ecdysone is in a highly dynamic state after its injection or its secretion by the ring-gland of *Sarcophaga peregrina*. Hormonal activity is rapidly destroyed by an inactivating mechanism which is present in the tissues but not in the blood.

2. Inactivation is blocked by low temperatures or anaerobic conditions—a finding that implicates chemical and, more particularly, oxidative reactions. The mechanism in question could be demonstrated in larval fragments but not in crude or fractionated homogenates.

3. When injected into mature larvae, 1 μ g of α -ecdysone loses 50% of its activity in 1 hour and 98% in 8 hours. Lower doses show even briefer "half-lives."

4. The rapid inactivation of ecdysone can account for its low titer in both the blood and tissues. Thus at the "critical period" for puparium formation, the entire larva contains only 2.5 nanograms, corresponding to only 7% of a *Sarcophaga* unit.

5. The evidence points to the accumulation, not of the hormone itself, but the covert biochemical and biophysical effects of the hormone. The covert effects undergo spatial and temporal summation within the target organs and finally discharge the overt developmental response.

6. The role of the blood is to serve, not as a reservoir, but as a pipeline through which ecdysone flows from the ring-gland to its sites of action and swift inactivation.

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LIGHT ORGAN FINE STRUCTURE IN CERTAIN ASIATIC FIREFLIES

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Control of effector activity by nerve has been studied exhaustively in muscle and to a lesser extent in glands and in electric organs. Coordinated light production by congregations of specialized photocytes represents an entirely different kind of effector activity, in which neural control, though long suspected, has only recently been established directly. Thus far it is only in the light organ or lantern of certain fireflies of the beetle family Lampyridae that nerve endings have been demonstrated (Smith, 1963), excitatory volleys in peripheral lantern nerves recorded (Case and Buck, 1963; Buonamici and Magni, 1967; Magni, 1967) and pharmacological evidence of neurosecretory excitation adduced (Smalley, 1965). However, the firefly lantern is a particularly favorable material in which to study excitation-effector coupling because it excels all other photophores in versatility and precision of control and because there exists a large body of detailed information about the biochemistry, physiology and morphology of the light-emitting system.

Recently recordings have been made of the flashing of fireflies of the genus *Pteroptyx* in Bornea and Thailand (Buck and Buck, 1968). These insects can produce flashes lasting only about 30 msec. and can flash repetitively at frequencies of more than 30 per second. Because this is at least three times as sharp a control as in any previously studied firefly, and because electron microscopic studies on species from the tropical orient have not previously been reported, the fine structure of the *Pteroptyx* lantern was studied. For comparison we present also some data for the firefly genera *Luciola* and *Pyrophanes* from the same regions.

MATERIAL AND METHODS

The species studied were *Pteroptyx valida* and *P. malaccac* from Thailand and *Pyrophanes appendiculata* and *Luciola* sp. from Sarawak.² The comparative histology of these and other Asiatic species is being reported separately by Mrs. Miriam McLean of our Laboratory, but she has kindly allowed us to introduce the account of ultrastructure with semidiagrammatic sketches made from her slides (Figs. 1 and 2).

¹The authors acknowledge with gratitude the extensive technical assistance of Mr. Charles Hanna. (One of us [J. B.] is grateful to The American Philosophical Society [Penrose Fund grant no. 3862] and to the National Geographical Society for travel grants.)

²Identifications were kindly made for us by Mrs. L. P. Ballantyne at the University of Queensland, Brisbane, Australia.

Most of our material for electron microscopy was fixed in the field, and of necessity the fixation schedule was modified from the usual, since microscopes, hoods and temperature controls were not available. The abdomens of the specimens were slit open and the whole insects were dropped into 6% glutaraldehyde in Millonig's buffer at ambient (tropical) temperatures. Some specimens were left in glutaraldehyde and others were transferred after an hour and a half to buffer con-

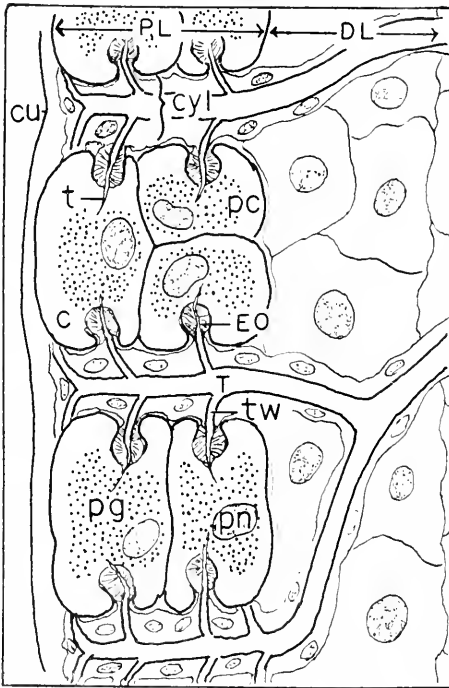


FIGURE 1.

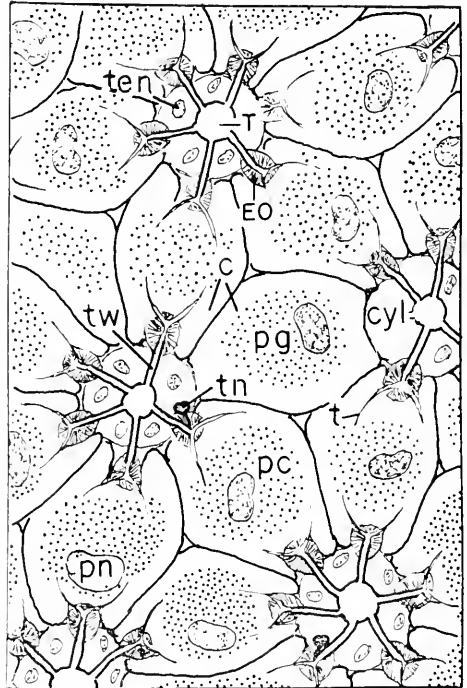


FIGURE 2.

FIGURE 1. Semidiagrammatic drawing of a transverse section through the lantern of *Pteroptyx*, showing portions of the photogenic (PL) and dorsal (DL) layers and three cylinders (cyl) containing tracheal trunks (T). c, cortex of photocyte, pc; EO, end organ; pg, region of photocyte granules; pn, nucleus of photocyte; t, tracheole; tw, tracheal twig; cu, cuticle.

FIGURE 2. Semidiagrammatic drawing of a horizontal section through part of the photogenic layer of a *Pteroptyx* lantern. cyl, cylinder; T, tracheal trunk; tw, tracheal twig; t, tracheole; ten, tracheal epithelial nucleus; tn, tracheolar cell nucleus; EO, end organ; pc, photocyte; c, cortex of photocyte; pg, region of photocyte granules; pn, photocyte nucleus. $\times 650$.

taining a small amount of thymol. They were in transit for from ten days to six weeks before reaching this laboratory, where they were post-fixed in Millonig's OsO_4 , dehydrated in alcohol, and embedded in Maraglas. Of these specimens, only *Luciola* showed satisfactory fixation. The other tissues were fixed poorly, but well enough to permit comparisons.

Later, we were very fortunate in obtaining some live specimens of *Pteroptyx*

malaccae.³ Lanterns of some of these specimens were fixed in cold 6% glutaraldehyde for one and a half hours, and then treated as above.

GENERAL STRUCTURE

A surprising finding was the similarity in gross lantern morphology not only among the three Asiatic genera but between them and the common North American genera *Photinus* and *Photuris*. As seen by light microscopy (Figs. 1 and 2) the organs consist of a photogenic layer usually two cells thick, lying immediately

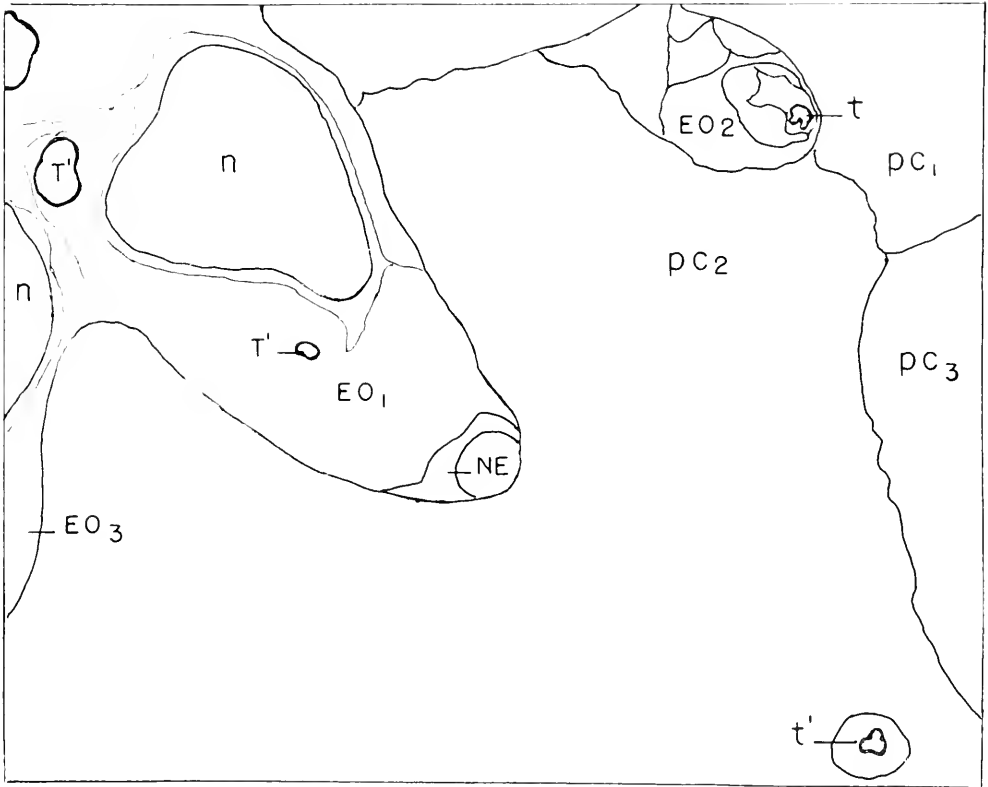


FIGURE 3A. Outline of cell boundaries in Figure 3B as an aid to orientation. EO₁, EO₂, and EO₃, end organs; pc₁, pc₂, and pc₃, photocytes; T', tracheal twig; t and t', tracheoles; n, twig cell nuclei. NE, terminal portion of nerve.

internal to the transparent ventral abdominal cuticle and backed by a dorsal ("reflector") layer several cells thick. Running vertically through both layers are tracheal trunks. In the photogenic layer each trunk runs within a wide tunnel or cylinder into which it gives off a multitude of short tracheal twigs. In horizontal

³ Live *Pteroptyx malaccae* were obtained by Dr. Douglas Gould of the Walter Reed Army Medical Center unit of the Seato Medical Research Project, Thailand, and brought to Bethesda by Dr. Robert Gordon of our Institute.

section (Fig. 2), it is seen that these cylinders are arranged in triangular symmetry, each surrounded by a radial rosette of polyhedral photocytes. At the periphery of the cylinders each twig terminates in a minute "tracheal end organ" which protrudes into the body of a photocyte. From the end organs issue delicate tracheoles which can be followed but a short distance as they penetrate an external "differentiated zone" or cortex, staining differently from the interior cytoplasm. In the Asiatic species the number of tracheoles originating from a single tracheal twig is

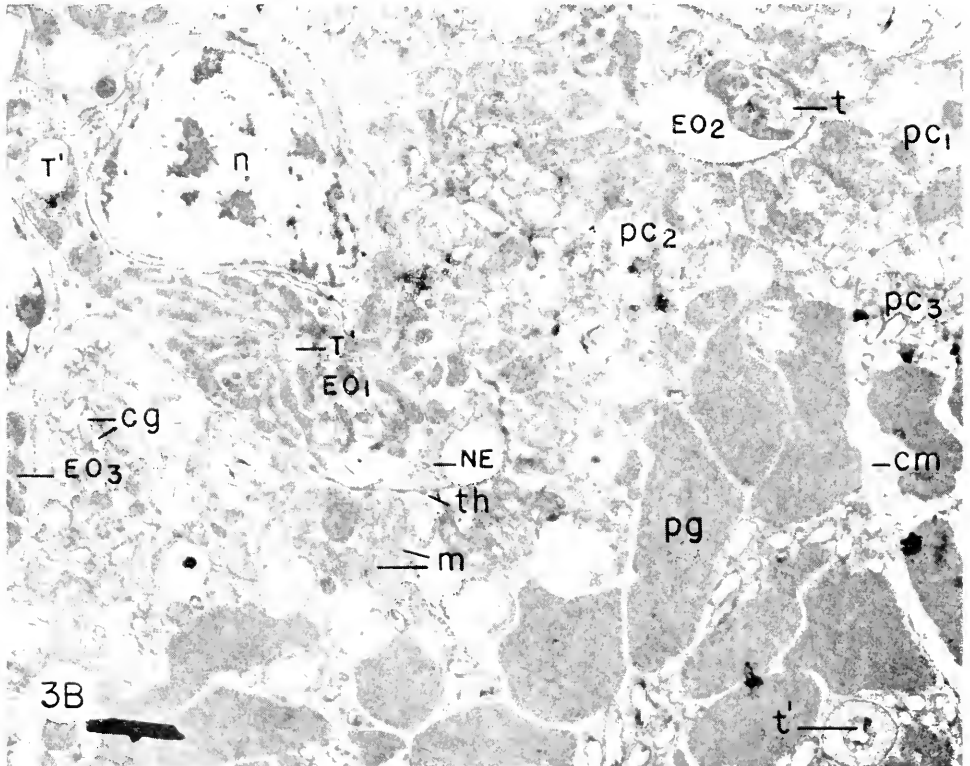


FIGURE 3B. Low power electron micrograph of region of contact between a cylinder and the photogenic layer, showing parts of three end organs (EO₁, EO₂, EO₃) and of three photocytes (pc₁, pc₂, pc₃). T', tracheal twig; t, tracheole in tracheolar cell; t', section of tracheole within photocyte cytoplasm enclosed by sheath of tracheolar cell cytoplasm and tracheolar cell membrane; n, nucleus of tracheal twig cell; NE, terminal portion of nerve; th, electron-dense thickenings along contact of end organ and photocyte membranes; m, photocyte mitochondria grouped around end organ; cg, clear-centered granules in cortical region of photocyte; pg, photocyte granule; cm, cell membrane of photocyte. *Luciola* sp. - 6600.

apparently always two, in contrast to the three or four of *Photinus* and the two or three (Smith, 1963) or three to seven (Kluss, 1958) reported for *Photuris*.

Previous electron microscopic studies of *Photinus* (Beams and Anderson, 1955) and *Photuris* (Kluss, 1958; Smith, 1963) have shown that the tracheal "end cell" is actually a highly complex structure consisting of an inner tracheolar cell en-

sheathing the tracheoles and a concentric outer cell enclosing the tracheal twig and its branching point and enveloping the tracheolar cell. Since this outer cell is very different from the usual tracheal end cell of insect tissues, we shall call it the "twig cell." Also in the interest of clarity we shall adopt Kluss' "tracheal end organ" to describe the entire complex, and we shall use "cortex" instead of "differentiated

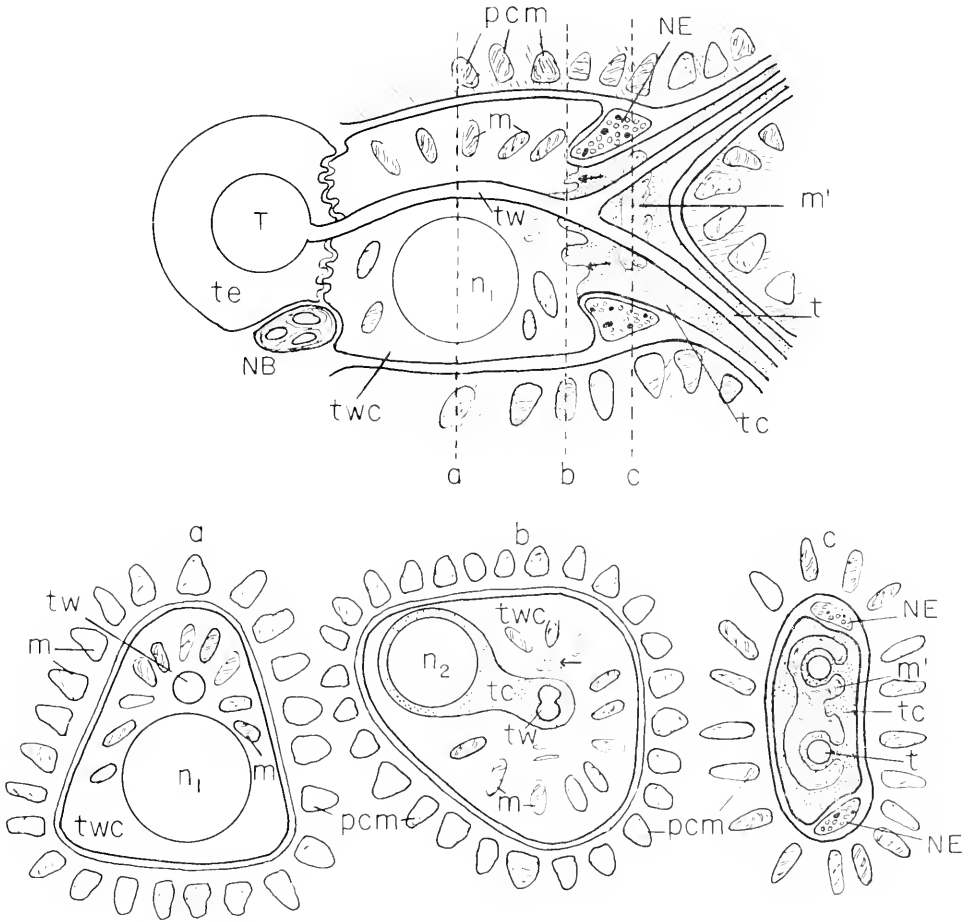
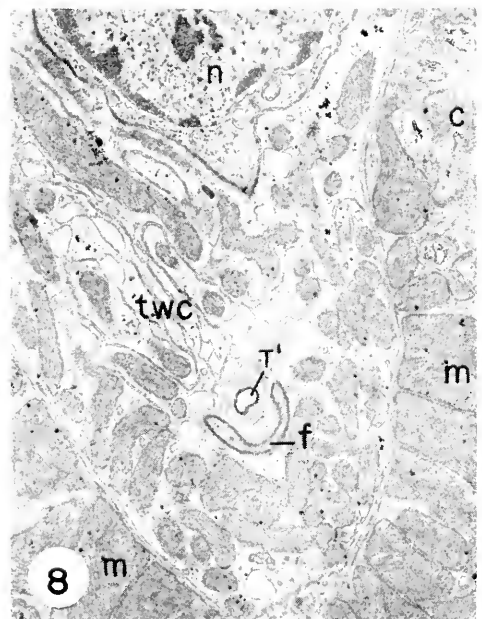
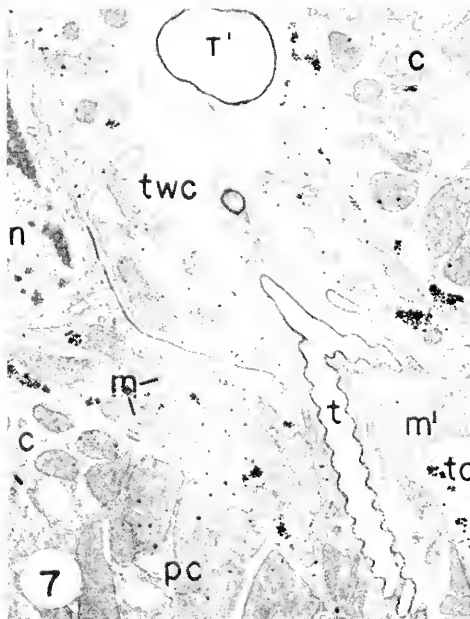
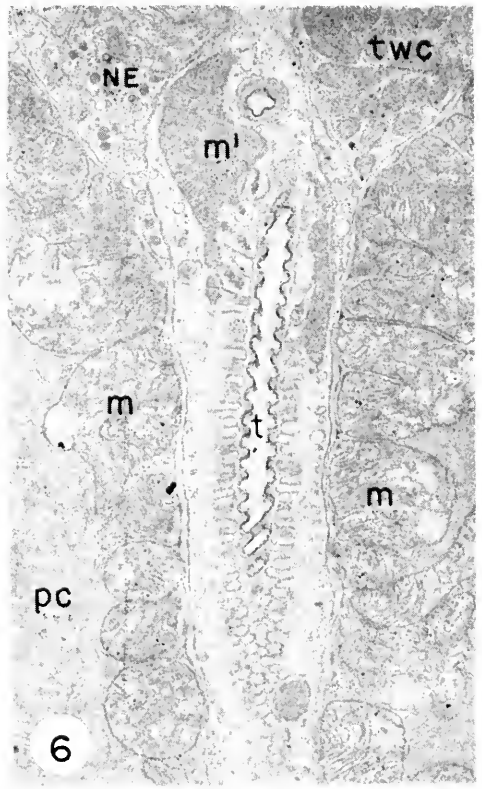
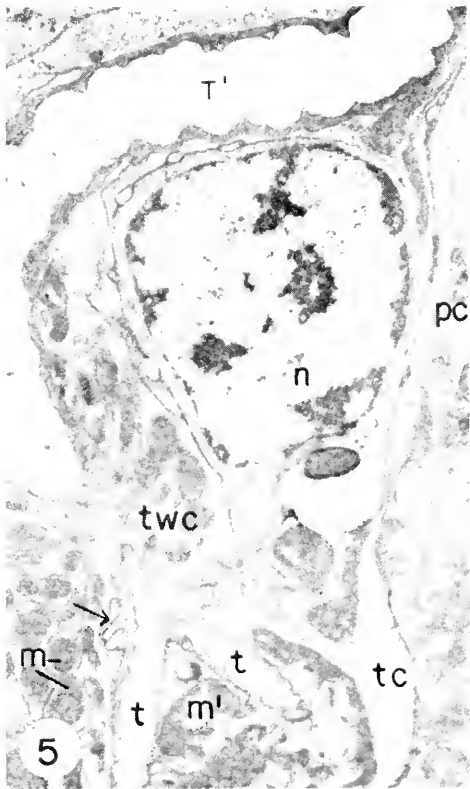


FIGURE 4. Diagram of an end organ to show the relationship of its components (twig cell, twc; tracheolar cell, tc; nerve endings, NE) to each other and to the photocyte. Transverse sections taken at the levels indicated by broken lines are represented in a, b, and c. T, trachea; tw, tracheal twig; t, tracheole; te, tracheal epithelial cell; m, mitochondria of twig cell; pcm, mitochondria of photocyte; m', large mitochondrion enveloping origins of tracheoles; NB, nerve branch approaching end organ; n₁, nucleus of twig cell; n₂, nucleus of tracheolar cell.

layer" or zone for the specialized peripheral zone of cytoplasm which is seen at all external or free faces of the photocytes—*i.e.*, where a photocyte abuts on the dorsal layer, on a cylinder, or on the ventral epithelium, but not where two photocytes touch.



FIGURES 5-8.

At the electron microscope level Asiatic lantern topography is also quite similar to that in *Photinus* and *Photuris*. We shall accordingly not describe the fine structure in general but concentrate on the several important structural differences between the two groups of species. Figure 3B gives a low power view of the region of contact between cylinder and photogenic layer. Parts of two end organs and of three photocytes are shown. The protrusions of the photocytes, which interdigitate tightly with the end organs, consist exclusively of cortical cytoplasm containing closely packed mitochondria and clear-centered granules. A nerve terminal shows near the distal tip of the end organ close to the twig cell-photocyte apposition plane. (The clear space under the NE label is an artifact.) The line of apposition between end organ and photocyte is marked by many electron-dense thickenings which are lacking on stretches of membrane between photocytes. A section of a tracheole is seen within the photocyte cytoplasm still accompanied by a sheath of tracheolar cell cytoplasm and the infolded photocyte membrane (*t'*).

TRACHEAL END ORGAN

Because of its presumed involvement in excitation of luminescence (see below), the tracheal end organ cannot be discussed in isolation from the structures with which it makes direct contact, particularly the photocyte.

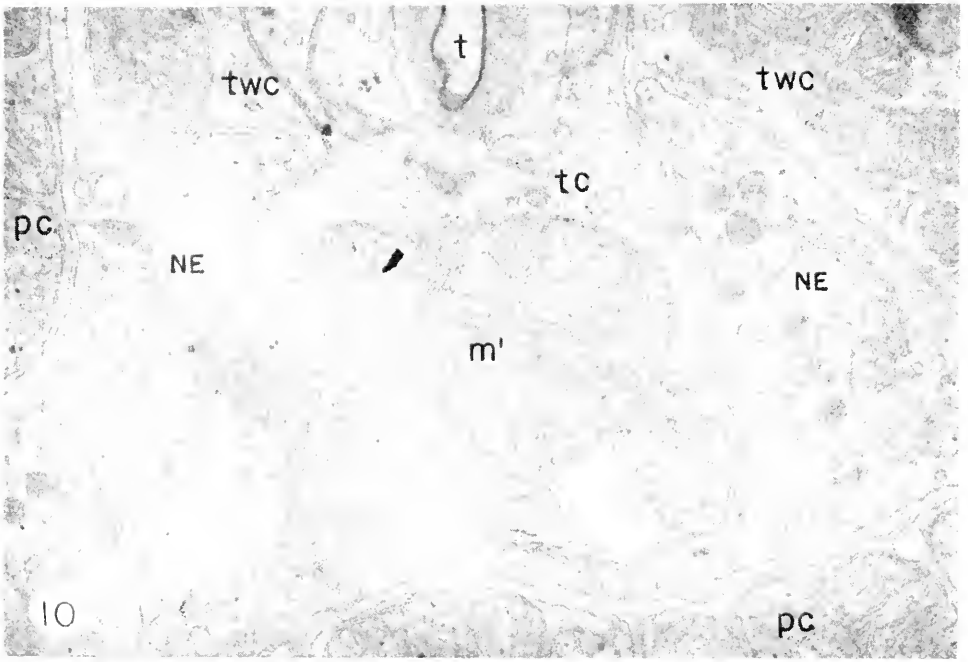
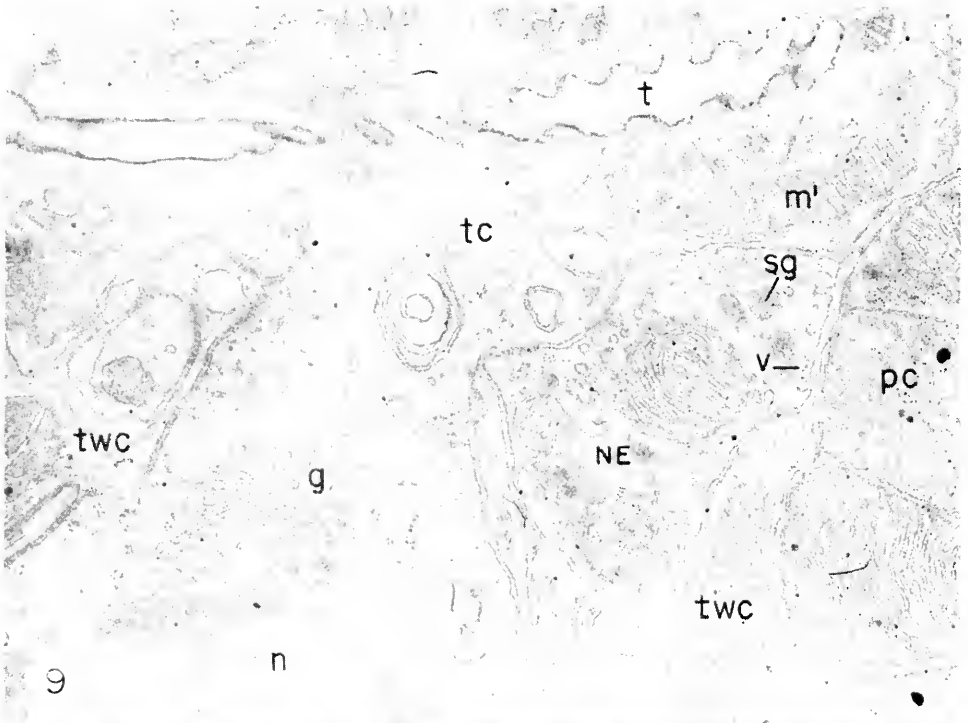
The end organs of *Pteroptyx*, *Pyrophancs* and *Luciola* conform in general to the pattern described in detail by Smith (1963). The relationship of its parts can be followed by reference to the diagram (Fig. 4). The twig cell membrane is thrown into profuse finger-like radial folds in which lie elongated mitochondria (Figs. 5, 7, m). The cell membrane also interdigitates distally with the tracheolar cell membrane (arrows, Figs. 4 and 5). However, the Asiatic species differ from *Photuris* in a number of important respects, chief among which is the point of termination of the nerves. In *Photuris*, the ends of the finest peripheral nerves lie between the tracheolar and twig cells and the neuronal membrane is apposed equally to the membranes of these two cellular components of the end organ. It is thus separated from the photocyte by the twig cell cytoplasm and membranes, a distance of usually not less than a micron. In the oriental species the bell-shaped twig cell overlaps only the proximal portion of the tracheo-

FIGURE 5. Longitudinal section through an end organ of *Luciola* sp. The tracheal twig (*T'*) curves around the nucleus (n) of the twig cell (twc) before branching into two tracheoles (*t*). A special large lobed mitochondrion (*m'*) in the tracheolar cell (*tc*) envelops the tracheoles just beyond their origin. Twig and tracheolar cell membranes interlock (arrow). pc, photocyte; m, mitochondria of twig cell. $\times 11,400$.

FIGURE 6. Section through an end organ of *P. malaccac* in the plane of one tracheole (*t*). Note especially the lobes of the large tracheolar cell mitochondrion (*m'*) protruding into cuticular folds of the tracheolar wall, and the array of photocyte mitochondria (*m*) along the tracheole. twc, twig cell; NE, nerve ending; pc, photocyte. $\times 11,400$.

FIGURE 7. Longitudinal section through an end organ of *P. malaccac* showing the origin of the two tracheoles (*t*). *T'*, tracheal twig; twc, twig cell; tc, tracheolar cell; *m'*, large mitochondrion of tc; n, nucleus of tc; c, cortex of photocyte (pc); m, mitochondria of twig cell. $\times 11,400$.

FIGURE 8. Transverse section through an end organ of *P. malaccac*, proximal to branching point of tracheal twig (*T'*). n, twig cell nucleus; m, mitochondria, and c, cortex, of photocyte; f, fold of tracheolar cell cytoplasm. $\times 11,400$.



FIGURES 9-10.

lar cell, rather than covering it completely. Consequently the nerve terminals can and do make contact not only with the tracheolar cell and twig cell membranes but with the photocyte membrane as well. A very careful search of several hundred micrographs failed to disclose any structure analogous to the tight junction or subsynaptic web of the neuromuscular junction of vertebrates. The neuronal contact with the membrane of the photocyte appears to be somewhat less close than that with the cells of the end organ (Figs. 9-12).

The end organs of Asiatic fireflies differ markedly from their American counterparts in having a single, very large, branched mitochondrion wrapped around the two tracheoles just distal to their origin from the tracheal twig (Figs. 11, 15, m'). There seems to be a tendency for the nerve terminal to be very intimately associated with this mitochondrion, both laterally and distally (Figs. 9-12, 15).

As in *Photinus* and *Photuris* the end organs are closely associated with the photocytes. Fingers or lobes of cortical photocytoplasm interdigitate with end organs, and the tracheoles with their sheaths of tracheolar cell cytoplasm push in between each pair of contiguous photocytes. Smith (1963) showed that the mitochondria of the cortex tend to congregate at the photocyte membrane. This effect is much more pronounced in the Asiatic fireflies, in which the mitochondria are larger, tend to a conical shape with base against membrane, and are often crowded tightly together in lateral contact (Figs. 6 and 15). This orientation is particularly striking along the proximal portions of the tracheoles. As the latter penetrate beyond the cortex of the photocytes, they are often surrounded by a looser group of more nearly spherical mitochondria, which may sometimes be replaced by photocyte granules.

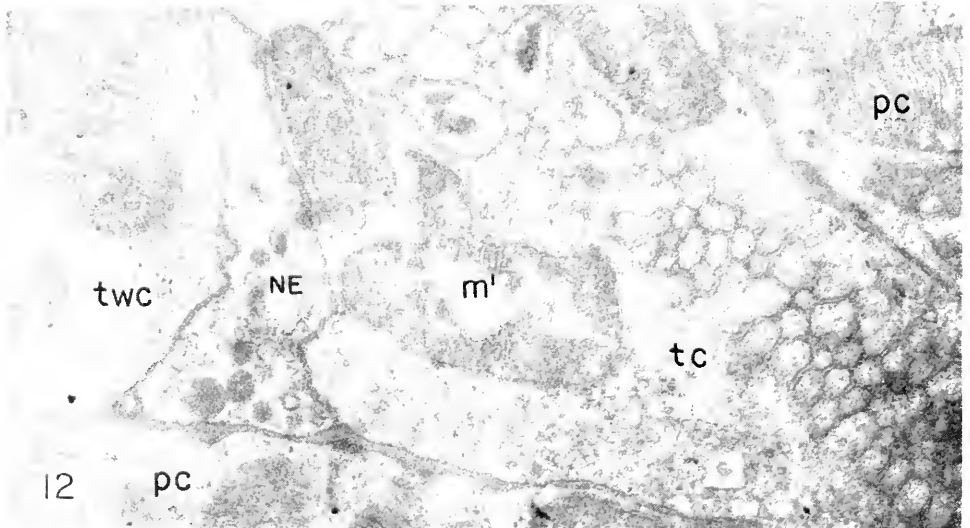
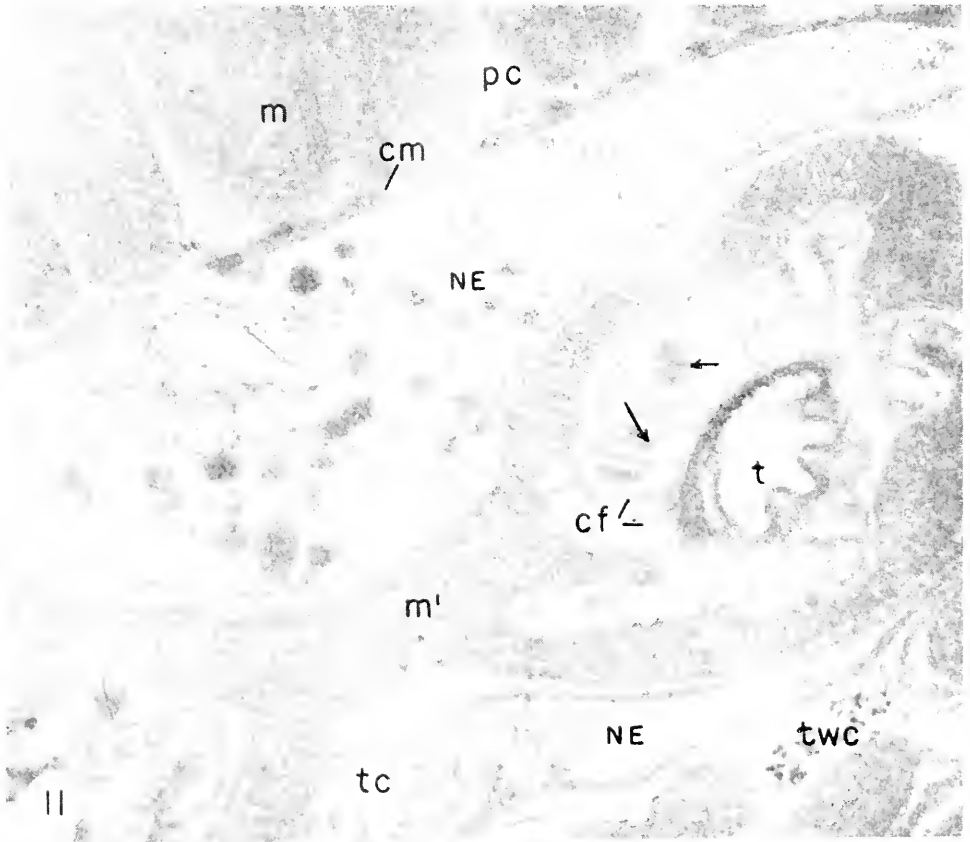
TRACHEOLES

Whatever they may be in other insects and other tissues, firefly lantern tracheoles appear to be more than merely taenidialized tubes for gas conduction. We shall therefore understand the term to include not only the cuticular lining but also the cytoplasmic sheath from the tracheolar cell which covers the intima throughout its course.

The tracheoles in the Asiatic fireflies show the same minutely villous honeycombed outpocketings of the sclerotized lining (Fig. 14) as they do in *Photuris* (Smith, 1963), and the same sheath of cytoplasm, in which mitochondria are sometimes intercalated between the luminal protrusions (Fig. 11, arrows). Whereas *Photuris* tracheoles follow an exclusively extracellular course between the apposed cell membranes of contiguous photocytes, the *Pteroptyx* tracheoles penetrate deeply into the cytoplasm, sometimes even approaching the photocyte nucleus

FIGURE 9. Section through an end organ of *P. malaccac*, parallel to one tracheole (t), showing terminal portion of nerve (NE) in contact with three types of cells (twc, tc, pc). m', special tracheolar cell mitochondrion; n, nucleus of tracheolar cell; g, Golgi body; v, vesicle; sg, granule. $\times 34,000$.

FIGURE 10. Section through end organ of *P. malaccac*, showing three contacts of massive terminal portion of nerve (NE). t, tracheole; twc, twig cell; tc, tracheolar cell; pc, photocyte; m', special tracheolar cell mitochondrion. $\times 34,000$.



FIGURES 11-12.

(Fig. 13). The penetration is not truly intracellular, since a sheath of the photocyte membrane accompanies the tracheolar intrusion (Fig. 13), but contact with the photocyte interior is otherwise intimate.

The lumen of the tracheoles seems everywhere to have a diameter of about 0.3μ , and the cytoplasmic sheath, once free of the tracheolar cell body, does not taper appreciably. The question arises as to whether the tracheoles end blindly or anastomose with tracheoles from an adjacent cylinder as claimed by Buck (1948) on the basis of light microscopic evidence from *Photinus*. Electron microscope preparations leave the matter still moot. Beams and Anderson (1955) saw no evidence of anastomosis in *Photinus pyralis*; Kluss (1958) states that *Photuris* tracheoles do run continuously from one cylinder to the next; Smith (1963) does not mention tracheole termination. Our own preparations have not disclosed anything identifiable as an ending, but neither have they provided anything like the continuous several micron-long stretch of tracheole midway between cylinders that would be needed to support tracheolar anastomosis with any confidence.

PHOTOCYTE GRANULES

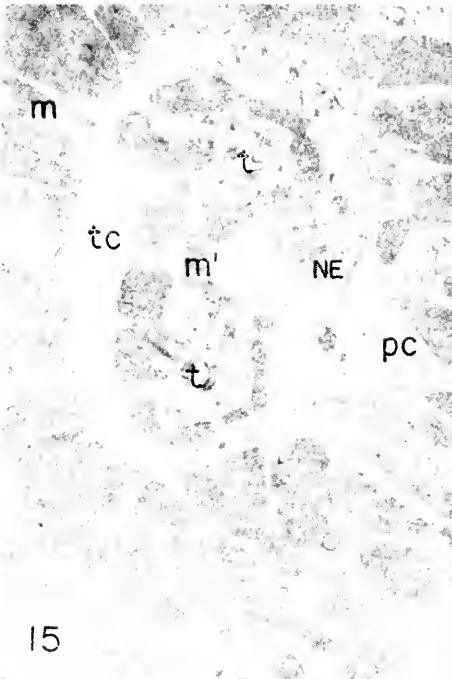
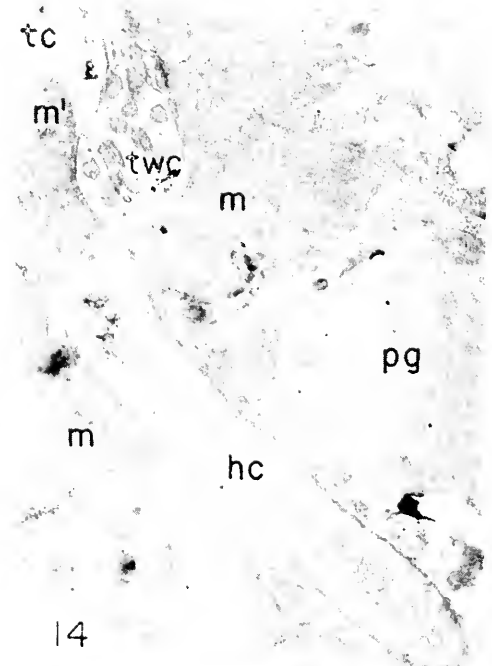
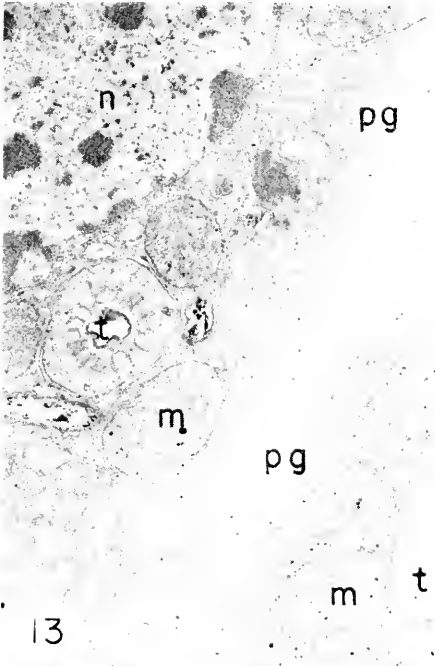
The large granules that are the most prominent inclusion of the interior cytoplasm of photocytes are also the most enigmatic. For about a century they have been referred to as "photogenic granules" or the equivalent without a scrap of direct evidence that they are involved in light emission. In our oriental species they tend to be elongated, rather than spherical, and to be larger ($4.5 \times 2.5 \mu$) than in *Photinus* and *Photuris*. We have only rarely seen single interior vacuoles or vesicles, sometimes connected to what looks like a slender duct, that were occasionally seen in *Photuris* photocyte granules by Smith (1963). Instead we find heavy concentrations of typical microtubules, 230 \AA o.d., 150 \AA i.d. (Fig. 16). There is some evidence of the tubules curving in conformation with the contour of the granule, but no real hint as to what their function may be.

INTERCELLULAR CONTACT

Smith described quite striking "adhesion plates" connecting the surface membranes of contiguous photocytes at intervals. We have not observed such structures in our material. However, we find rather regularly spaced triangular nodes of enhanced electron density along the line of contact between end organ and photocyte. In the domain of tracheolar cell-photocyte membrane contact the nodes appear to be about twice as numerous as in the region of twig cell-photocyte contact (Figs. 11, 12).

FIGURE 11. Transverse section through end organ of *Luciola* distal to branching of tracheal twig (cf. Figure 4c). Nerve endings (NE) make contact with twig cell (twc), tracheolar cell (tc) and photocyte (pc), and also follow the contours of the large tracheolar cell mitochondrion (m'). t, tracheole; m, mitochondria of photocyte clustered around end organ. Note protrusions of tracheolar cell mitochondrion into cuticular folds of tracheolar wall (arrows). $\times 54,000$.

FIGURE 12. Longitudinal section through end organ of *Luciola* showing contact of nerve endings (NE) with twig cell (twc), tracheolar cell (tc), and photocyte (pc). m' = special tracheolar cell mitochondrion. $\times 34,000$.



FIGURES 13-16.

DISCUSSION

Comparative histological data strongly indicate that tracheal end organs are present in all firefly species able to produce sharp flashes and absent in species able only to emit a long-lasting glow (Buck, 1948). This information, when combined with the facts that nerves terminate in the end organs (Smith, 1963) and that the light emitted by the organ seems to come from the region of cylinder-photocyte contact (review in Buck, 1966), strongly indicates that the end organ-cortex region is in some sense a neuroeffector junction. It is by no means clear, however, that the junction conforms to the pattern seen in neurally excitable muscle or gland or in intern neuronal synapses.

The evidence for neurosecretory excitation of firefly luminescence has rested primarily on the occurrence of neuroeffector facilitation during repetitive electrical stimulation (Buck and Case, 1961) and on the close morphological similarity, pointed out by Smith, between the two types of inclusions seen in the peripheral nerve terminals of the lantern and the "neurosecretory droplets" and "synaptic vesicles" of the vertebrate neurohypophyseal junction (Palay, 1956). The case has recently been strengthened by direct evidence of involvement of noradrenalin in excitation (Smalley, 1965). Nevertheless, neither in Smith's preparations of the *Photuris* lantern nor in ours from the Asiatic species does the membrane of the nerve terminal show the tight membranal junctions with another cell membrane and sub-synaptic web which are characteristic, for example, of many neuroeffector or neuronal synapses (De Robertis, 1967).

The discovery that the nerve terminals in *Pteroptyx* and its Asiatic neighbors make direct contact with the presumed effector cell of the lantern, the photocyte, only partly relieves the difficulty that, as in *Photuris*, they seem also to be closely associated with both the tracheolar and twig cell membranes. A conceivable role of tracheoles in excitation as well as in gas conduction is suggested by their profuse distribution, deep inter- and intra-cellular penetration, and the special structure of their enveloping cytoplasm. On the whole, the present ultrastructural evidence leaves the whole question of the triggering of bioluminescence in the firefly lantern in an unsatisfactory conceptual state, but certainly indicates that the junction is of a quite new type.

FIGURE 13. Portion of a photocyte of *P. malaccae* showing two sections of tracheole (t) within the cytoplasm, one very close to the nucleus (n) of the photocyte. Note that the membranes of tracheolar wall and of photocyte can both be seen. Mitochondria (m) cluster around tracheole. Photocyte granules (pg) contain numerous microtubules in cross section. $\times 11,400$.

FIGURE 14. Part of an end organ and tracheole of *P. valida*, poorly fixed, but demonstrating the honey-comb structure (hc) of the tracheolar wall. tc, tracheolar cell; twc, twig cell; m', tracheolar cell mitochondrion; m, photocyte mitochondria; pg, photocyte granule. $\times 11,400$.

FIGURE 15. Section through a tracheolar cell of *Luciola*, beyond the overlap of the twig cell (cf. Fig. 4c). t, tracheole; tc, tracheolar cell; m', tracheolar cell mitochondrion; NE, nerve ending; pc, photocyte with many mitochondria (m) radiating around tracheolar cell. $\times 11,400$.

FIGURE 16. High magnification electron micrograph of a portion of two photocyte granules of *Pyrophanes*, showing cross-sections and longitudinal sections of numerous microtubules. $\times 80,000$.

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REPRODUCTIVE CYCLE OF THE SURF CLAM, *SPISULA SOLIDISSIMA*, IN OFFSHORE NEW JERSEY

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Atlantic surf clams, *Spisula solidissima* (Dillwyn), were collected from offshore New Jersey during a three and one-half-year period to obtain their gonads for histological treatment in a study of the annual reproductive cycle. Documentation of the annual reproductive cycle on the basis of gonad distention (Westman and Bidwell)¹ and excision of gametes (Allen, 1951, 1953; Schechter, 1941) is incomplete and fragmentary. Neither gonad distention nor excision of gametes assesses the events of gamete development within the gonad. Also, parasites are a possible source of error if the technique of observing gonad distention is used. I have seen infected surf clams with distended gonads when those of uninfected individuals were shrunken. Gonad color has been used as an indicator of sex and gonad development of the sea scallop, *Placopecten magellanicus* (Posgay and Norman, 1958; Merrill and Burch, 1960). The color of surf clam gonads containing mature sex products, however, is not consistent. Ovaries of females are sometimes pink, but may also be white, as are the testes (Schechter, 1941).

Periodic histological examinations of gonad tissue during several successive years has been a valuable method of determining the annual frequency and duration of the reproductive cycle in many marine invertebrates (Giese, 1959). The reproductive cycles of the soft-shelled clam, *Mya arenaria* (Pfitzenmeyer, 1962, 1965; Shaw, 1962, 1965; Ropes and Stickney, 1965) have been recently established by describing the cytological events within the gonad. Such events within the gonads of surf clams from an area of intensive fishing off the New Jersey coast have been related to water temperature. The present study of seasonal gonad development was undertaken to determine the annual frequency and duration of spawning—information basic to understanding larval occurrence and time of setting of juvenile surf clams.

MATERIALS AND METHODS

Surf clam gonads were collected from port landings (1962 and 1963) and commercial and research vessels (1964 and 1965). In all years, samples were taken as soon as practicable after the clams were removed from the water, because the surf clam readily reacts to artificial spawning stimuli (Stickney, 1963) and some released sex products soon after capture. All samples were from

¹Westman, J. R., and M. H. Bidwell, 1946. The surf clam. Economics and biology of a New York marine resource. State of New York Conservation Dept. Unpublished manuscript on file at Bureau of Commercial Fisheries Biological Laboratory, Oxford, Maryland 21654.

the vicinity of Barnegat Lightship and were caught by hydraulic dredges operating at depths of 18 to 32 meters. A sample was taken monthly during the winter and biweekly during the spring, summer, and fall. Each sample comprised 25 live clams with shells 120 mm. long or longer. The portion of the gonad ventrad to the heart was removed after fixation of the entire visceral mass in Bonin's solution, thereby preventing the loss of eggs and sperm, and minimizing the possible variability in development within a single gonad. Standard histological techniques were employed to prepare sections 7 μ thick for staining with Delafield's hematoxylin and eosin.

After microscopic examination of the section, each specimen was assigned to a category of gonad development. The proportions of clams in each category, regardless of sex, were recorded for individual samples and the samples were combined each month to simplify presentation. In graphic presentations of the results, the combined data are plotted at a date midway between the sample dates.

The possible influence of temperature on gonad development and spawning was considered. Surface and bottom sea-water temperatures at Barnegat Lightship were obtained from Chase (1965, 1966) for 1962 and 1963 and from unpublished records for 1964 and 1965.

ANATOMY AND SPAWNING

Male and female surf clams are distinguishable by microscopic examination of the gonads because they are unisexual or dioecious (Coe, 1943). Hermaphroditism is rare and then is apparently a developmental accident (Ropes, 1966).

The gross features of a surf clam's reproductive anatomy include anastomosing gonadal alveoli which lie ventrad to the pericardial chamber and overlay the dark brown digestive gland or liver mass, and surround the coiled intestine. In the distended gonads before spawning, the root-like alveoli are crowded close to each other and are filled with "late ripening" and "ripe" sex cells; their connective tissue walls are stretched thin; and they occupy much of the visceral mass. After spawning the alveoli contract, their walls thicken, and mesenchyme is interspersed between them; the dark brown digestive gland is then the most prominent organ and the visceral mass is flaccid.

The alveoli lead into a pair of gonoducts, each of which has an enlarged portion that serves as a reservoir for ripe sex cells before spawning (Stickney, 1963). The gonoducts terminate at the genital apertures on minute papillae that protrude from the surface of the posterior foot retractor muscle on each side of the visceral mass.

The passage of water for respiration and digestion conducts sex cells from the clam into the surrounding water during spawning. Water drawn through the incurrent siphon flows into the infrabranchial chamber, around the gills, past the genital papillae, into the epibranchial chamber, and then out through the excurrent siphon. Spawning is often accompanied by rhythmic contractions of the adductor muscles which pull the two valves together. Gametes are either emitted into the water by the muscular activities or flow out. Oocyte maturation follows spawning and fertilization (Costello *et al.*, 1957; Stickney, 1963) but meiosis occurs in the gonads of male surf clams and functional sperm are developed before spawning.

CATEGORIES OF GONAD CONDITION

In this study the more or less continuous reproductive process was divided into five phases as described below. The demarcation between phases is not sharp—the divisions are convenient rather than natural.

*Female gonads*1. *Early active phase*

In the early active phase ovogonia appear at the periphery and are embedded in the alveolar walls, often before the gonads are completely empty of ripe oocytes. The round and oval nuclei of ovogonia contain distinctly basophilic nucleoli and the nucleoplasm has thin, irregularly shaped, dark-staining threads (Fig. 1). Cytoplasm surrounds the nuclei but is indistinct and of irregular shape. The ovogonia are possibly undergoing the pre-meiotic stages similar to those described for the desert snail, *Eremina desertorum* (Fahmy, 1949) and the soft-shelled clam, *Mya arenaria* (Coe and Turner, 1938). The alveolar walls are not completely contracted and lumina are evident in most gonads.

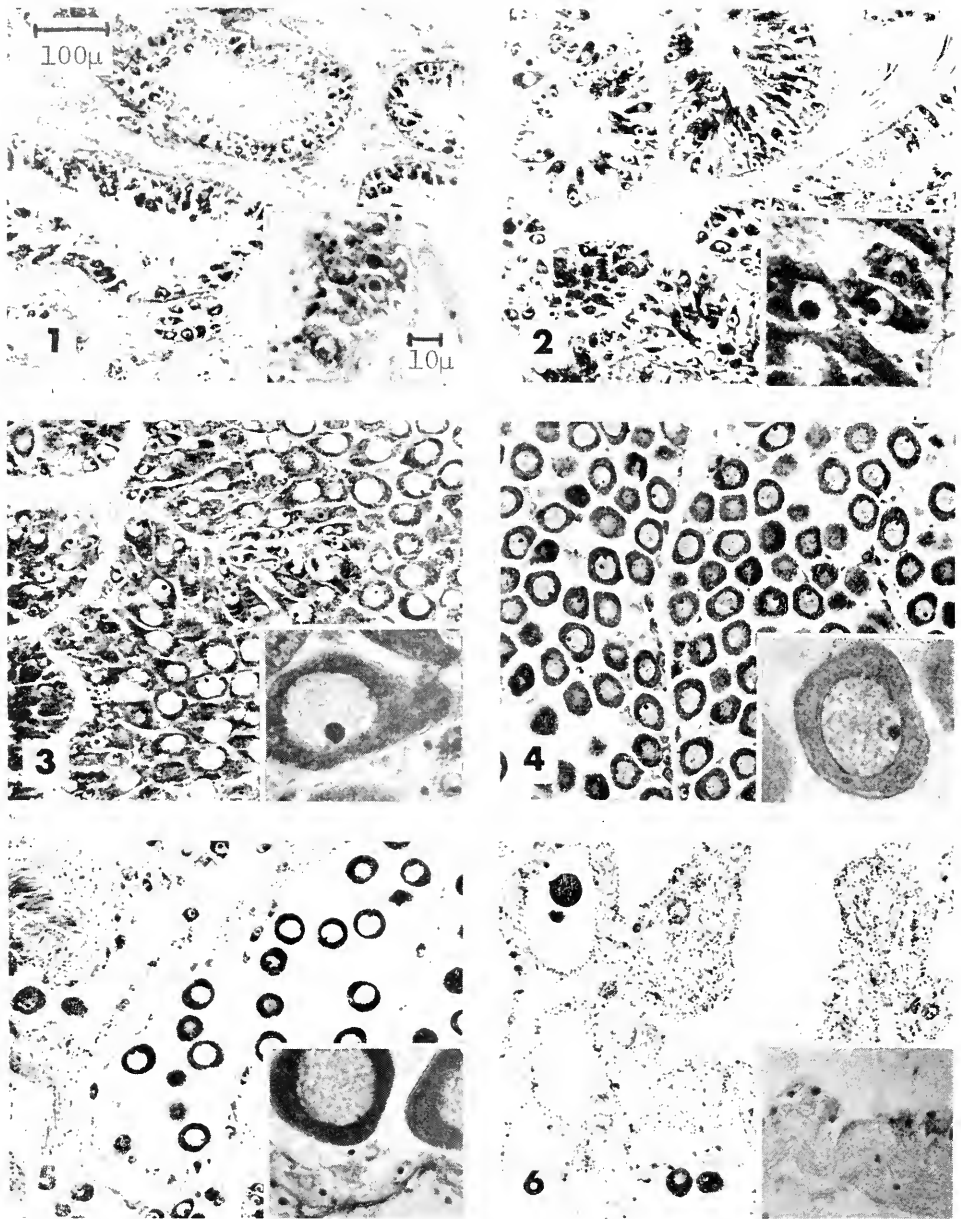
The early oocytes also have mostly round and oval nuclei containing basophilic nucleoli, but the nucleoplasm is homogeneous and almost clear (Fig. 2). The oocytes lie at the periphery of the alveolar walls and the cytoplasm extends into the lumen of the alveolus, where a rounded cap is formed. The shape of the oocytes may be square, triangular, cylindrical, or hemispherical. All of these oocytes are attached to the basement membrane by a stalk. The diameters of the nuclei range from 5 to 20 μ and average 11.1 μ .

2. *Late active phase*

Oocytes in the late active phase are larger than early oocytes and many are round (Fig. 3). The nuclei contain basophilic nucleoli and the nucleoplasm is homogeneously granular. The nuclei, surrounded by cytoplasm, are the most conspicuous characteristic of the cells. Some oocytes are attached to the basement membrane of the alveoli by a thin stalk, but most are free in the lumina. The basement membrane is thin—apparently due to the increased volume of cells within the alveoli. The diameters of the nuclei range from 14 to 32 μ and average 24.1 μ .

3. *Ripe phase*

In ripe clams, oocytes are free in the lumina of the alveoli. The vitelline membrane contains the cytoplasm and a large nucleus (Fig. 4). These oocytes are very similar to those in the late active phase except for the presence of prominent amphinucleoli (Allen, 1953). Each amphinucleolus consists of an almost transparent, granular nucleolus and a small opaque, basophilic nucleolus. Ripe gonads typically have a dense appearance because the alveoli are crowded together and are filled with large oocytes. The diameters of the nuclei range from 22 to 33 μ and average 28.5 μ .



FIGURES 1-6.

FIGURE 1. Section of gonad tissue from a female surf clam, early active phase of the first annual reproductive cycle. All of the large photomicrographs are reproduced at the same magnification and a scale ($100\ \mu$) is provided. The small photomicrographs inserted in the lower right-hand corner of each figure are enlargements of portions of the gonad all at the same magnification and a scale ($10\ \mu$) is provided. Collection date: 21 January 1964.

4. *Partially spawned*

A few large, ripe ovocytes are free in the lumina of some alveoli of partially spawned clams; other alveoli are completely void of ripe ovocytes (Fig. 5). Ovogonia and early ovocytes appear in the contracted and thickened alveolar walls. The spaces between alveoli contain mesenchyme and the whole gonad has a loose, flaccid appearance. In some clams, ripe ovocytes are being cytolized.

5. *Spent*

In typical spent clams, the open lumina of alveoli are usually void of ripe ovocytes. The thickened alveolar walls contain ovogonia and early ovocytes (Fig. 6). Mesenchyme occurs in the spaces between the loosely arranged alveoli.

Male gonads

The criteria for the five categories of gonad condition in the male are presented below:

1. *Early active phase*

Clams in the early active phase contain cells undergoing the initial meiotic stages of development. Darkly stained spermatogonia are in the thickened alveolar walls (Fig. 7). Primary spermatocytes about 3 to 4 μ in diameter proliferate toward the lumina from a close contact with the alveolar walls and form single cell columns (Fig. 8). Their large, round and oval nuclei are surrounded by thin irregularly shaped cytoplasm. Two basophilic nucleoli within some cells indicate that cell division is taking place. The nucleoli are less basophilic and smaller than those in the earliest ovocytes.

2. *Late active phase*

The remaining spermatogenic stages are seen in clams in the late active phase. Secondary spermatocytes of about 2 to 3 μ in diameter with dark basophilic chromatin threads that nearly fill the nuclei are numerous and the surrounding cytoplasm is irregular in shape and very indistinct (Fig. 9). Small and strongly basophilic spermatids measuring about 1.5 μ in diameter form dense masses near the centers of the alveoli. A transformation of the spermatids results in the appearance of sperm in the lumina. The sperm form weak columns with the tails oriented toward the centers of alveoli. The sperm heads are about 1 μ in diameter.

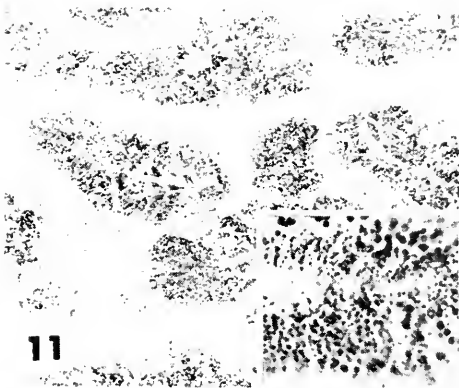
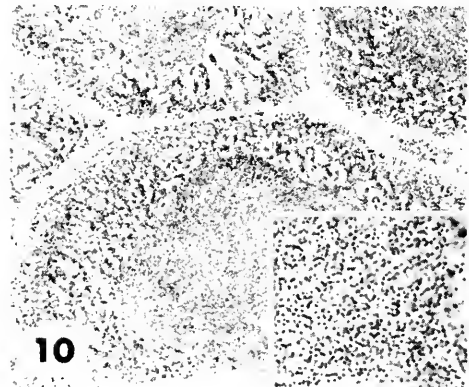
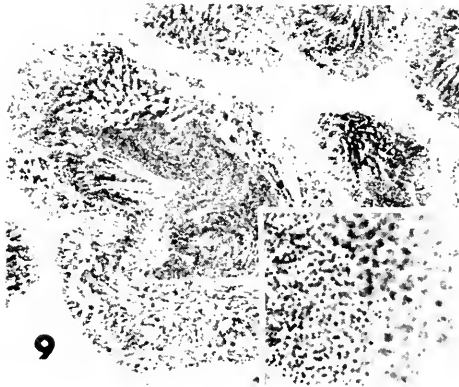
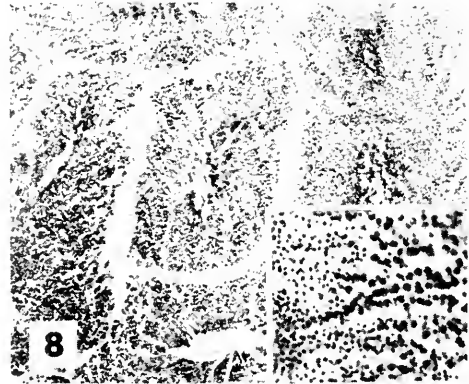
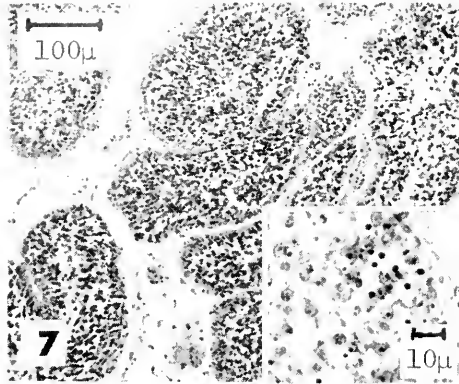
FIGURE 2. Same from a female surf clam, early active phase, showing enlarging ovocytes. Collection date: 18 February 1964.

FIGURE 3. Same, from a female surf clam, late active phase, showing large oval ovocytes. Collection date: 18 February 1964.

FIGURE 4. Same, from a female surf clam, ripe phase. Collection date: 26 June 1964.

FIGURE 5. Same, from a female surf clam, partially spent phase. Collection date: 28 August 1964.

FIGURE 6. Same, from a female surf clam, spent phase. Collection date: 13 August 1964.



FIGURES 7-12.

FIGURE 7. Section of gonad tissue from a male surf clam, early active phase of the first annual reproductive cycle. Collection date: 21 January 1964.

FIGURE 8. Same, from a male surf clam, early active phase, spermatozoa form columns. Collection date: 18 February 1964.

FIGURE 9. Same, from a male surf clam, late active phase. Collection date: 3 June 1964.

3. *Ripe phase*

Mature sperm form dense masses in the alveoli of clams in the ripe phase (Fig. 10). The sperm appear to be swirling in the centers of some alveoli, in others their tails are oriented toward the centers of alveoli, and in still others they form a homogeneous mass. Cells in early stages of spermatogenesis at the periphery of alveoli are much less numerous than the sperm.

4. *Partially spawned phase*

Partially spawned male clams still contain spermatozoa within the centers of alveoli, but they are substantially less numerous than in ripe clams (Fig. 11). The alveolar walls are compressed and thickened. Like the ovogonia and early oocytes at this stage, spermatogonia and primary spermatocytes are developing along the basal membrane before most gonads are completely void of spermatozoa.

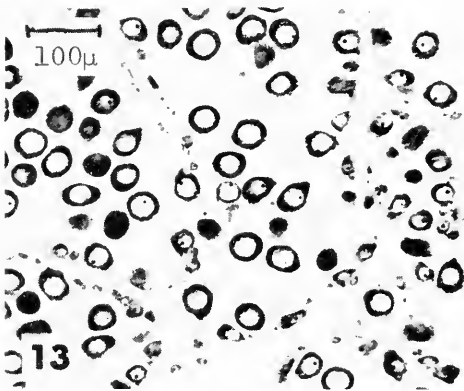


FIGURE 13. Section of gonad tissue from a female surf clam, ripe phase of the second annual reproductive cycle. Collection date: 24 September 1964.

FIGURE 14. Section of gonad tissue from a male surf clam, ripe phase of the second annual reproductive cycle. Collection date: 24 September 1964.

5. *Spent phase*

The alveoli of spent male clams contain no spermatozoa or very few and the lumina are characteristically open (Fig. 12). Primary spermatogonia are proliferating from the alveolar walls.

SECOND ANNUAL CYCLE

A second annual maturation of New Jersey surf clams was observed during three of the four years and was essentially a repetition of the sequence in gonad conditions already presented. The second maturation seemed to be of minor

FIGURE 10. Same, from a male surf clam, ripe phase, a dense mass of spermatozoa is formed. Collection date: 15 July 1964.

FIGURE 11. Same, from a male surf clam, partially spawned phase. Collection date: 29 July 1964.

FIGURE 12. Same, from a male surf clam, spent phase. Collection date: 29 July 1964.

importance, however, since fewer oocytes appeared in the gonadal alveoli (Fig. 13). Counts of the number of oocytes in 1.25 mm.² of gonadal tissue sections of ripe clams from the two reproductive cycles in 1964 showed an average density of 44.2 oocytes for 32 ripe clams from the first cycle and 26.4 oocytes for 20 ripe clams from the second cycle. The sperm in male gonadal tissues also appeared to be less numerous during the second cycle (Fig. 14).

ANNUAL REPRODUCTIVE CYCLES IN RELATION TO WATER TEMPERATURE

An account of the gonad conditions in 1514 surf clams and bottom water temperatures taken in the area sampled for clams is presented in chronological order. Gonad collections were initiated in June, 1962, and continued through 1965; water temperature data were available for the period 1962-65. Unless otherwise specified, all references to water temperature are for the bottom.

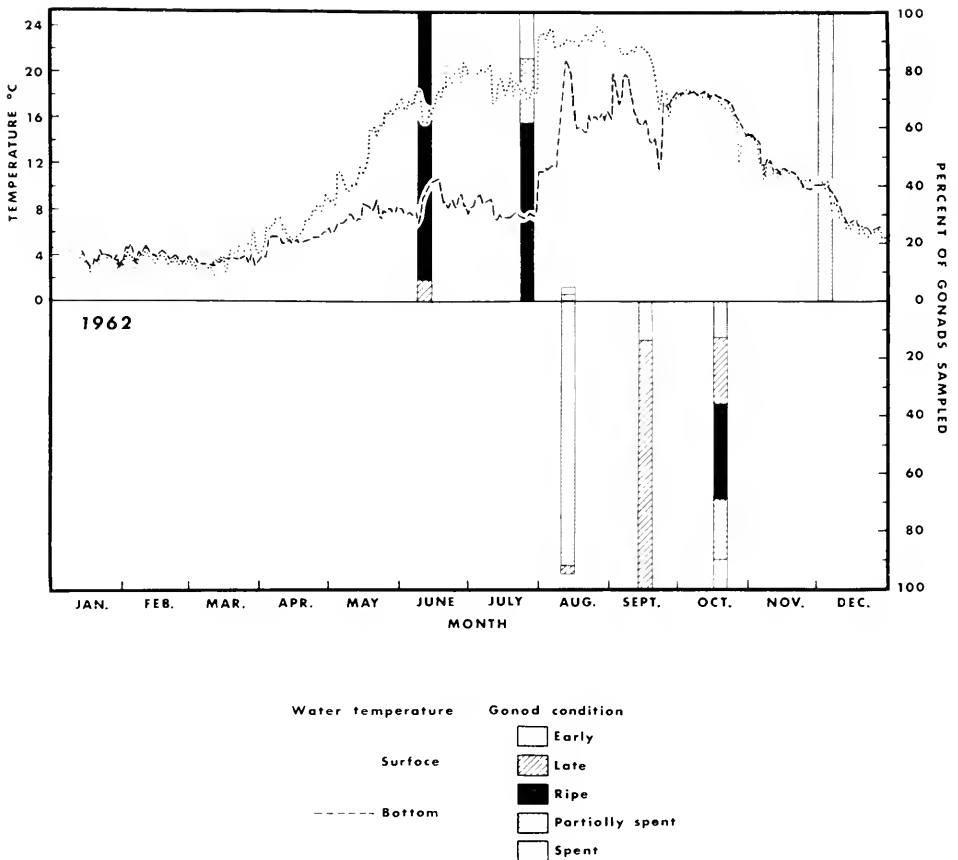


FIGURE 15. Gonad conditions of *Spisula solidissima* from New Jersey and sea-water temperatures at Barnegat Lightship in 1962. The bars above the centerline represent a single reproductive cycle and those below the centerline represent a subsequent cycle.

Reproductive cycle and water temperature in 1962

The June and July samples contained mostly ripe clams (Fig. 15). Spawning began by about mid-July when 37% of the clams were in the partially spawned and spent condition, and was completed by mid-August when 94% of the clams were in the early active phase. All clams were in the early and late active phases during September. All phases of gonad condition were represented in the October sample. A second spawning was apparent in mid-October when 32% of the clams were partially spawned and spent. All clams collected in early December were in the early active phase.

The temperatures were lowest (2.2° C.–5.2° C.) in late January, February, and early March. A warming trend followed, and 8° C. was reached by mid-May. The temperature reached 10.7° C. in mid-June, but most temperatures were between 7° C. and 9° C. during the 1½ months preceding the mid-July spawning. Spawning began 1 to 2 weeks before a rapid temperature rise, from 8° C. to 21° C., during late July to mid-August. Clams in the early active phase of a second reproductive cycle were most numerous when the temperature had reached its seasonal high (21° C.). The second spawning was in October, during a period when temperatures dropped from a high of 18° C. to about 10° C.

Reproductive cycle and water temperature in 1963

All of the clams were in the early active phase in January (Fig. 16). Gonads were in the late active phase in 4% of the clams by mid-February and in 56% of the clams by mid-March. Ripe clams constituted 92% of the mid-May sample and 98% of the mid-June sample. Spawning began in early July when 60% of the clams were partially spawned or spent. During August, 75% of the clams were still partially spawned or spent. The first annual spawning must have been completed by the end of August, because by mid-September 96% of the clams were in the late active and ripe phase. In mid-October collections, 28% of the clams were partially spawned and spent—evidence of a second spawning. Most (92%) of the clams were in the early active phase by early December.

Winter temperatures fell to a low of 2° C. during February. A warming trend after March (bottom temperatures are not available for March) was gradual and without marked fluctuations. The temperature on the bottom reached 10° C. by early June and 11° C. by late June. Temperatures reached 12° C. in mid-July. Spawning in July and August preceded a rapid rise in temperature from 9.6° C. to 18.9° C. during the early part of September. The beginning of a second reproductive cycle coincided with the rapid rise in temperature. The second spawning in October was during a period when temperatures dropped from 17° C. to about 15° C.

Reproductive cycle and water temperature in 1964

All clams were in the early active phase in January (Fig. 17); 63% were in the late active phase in February and this condition predominated in March and April. Gonads were ripe in 79% of the clams in May and in all of the clams in June. Spawning occurred in early July (43% of the clams were in the partially spawned and spent condition in mid-July). Fifty-three per cent of the clams were

partially spawned and spent in mid-August and the rest were in the early active phase. A progressive development of the gonads during September and October was followed by a second spawning late in the year; 68% of the clams were partially spawned and spent in early November. In early December, 80% of the clams were in the early active phase.

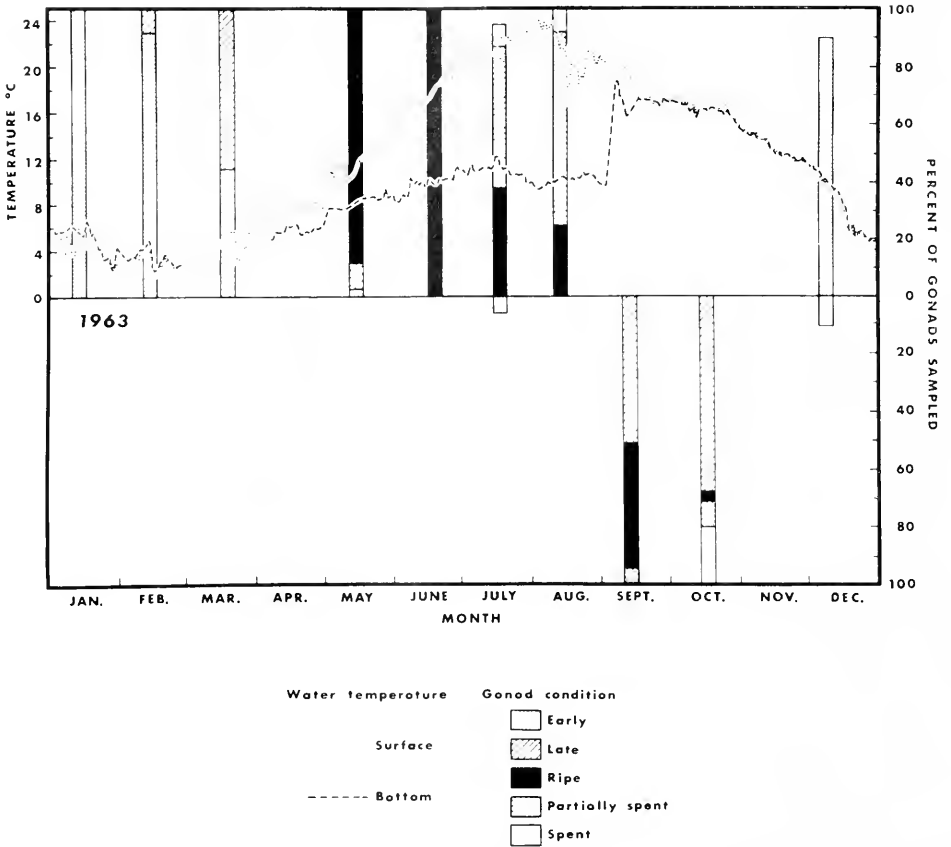


FIGURE 16. Gonad conditions of *Spisula solidissima* from New Jersey and sea-water temperatures at Barnegat Lightship in 1963.

The period of low winter temperature from January through March included a drop to 2° C. in February and was followed by a warming trend in April. In May and June the bottom temperatures fluctuated between 7° C. and 9° C. A temperature of 11° C. was reached in early July and was a possible inducement to spawning. All of the clams were completely spawned out in August after the rapid rise in temperature (7.4° C. 18.6° C.) during late July. The second reproductive cycle was underway in 47% of the clams after the rapid rise in temperature. The second spawning was during a period when temperature dropped from 16° C. to about 12° C.

Reproductive cycle and water temperature in 1965

All clams were in the early active phase during January and February (Fig. 18). The late active phase appeared in 76% and 62% of the clams during March and April, respectively. Ripe clams made up 94% of the late June sample and continued to predominate in the July, August, and September samples. In July

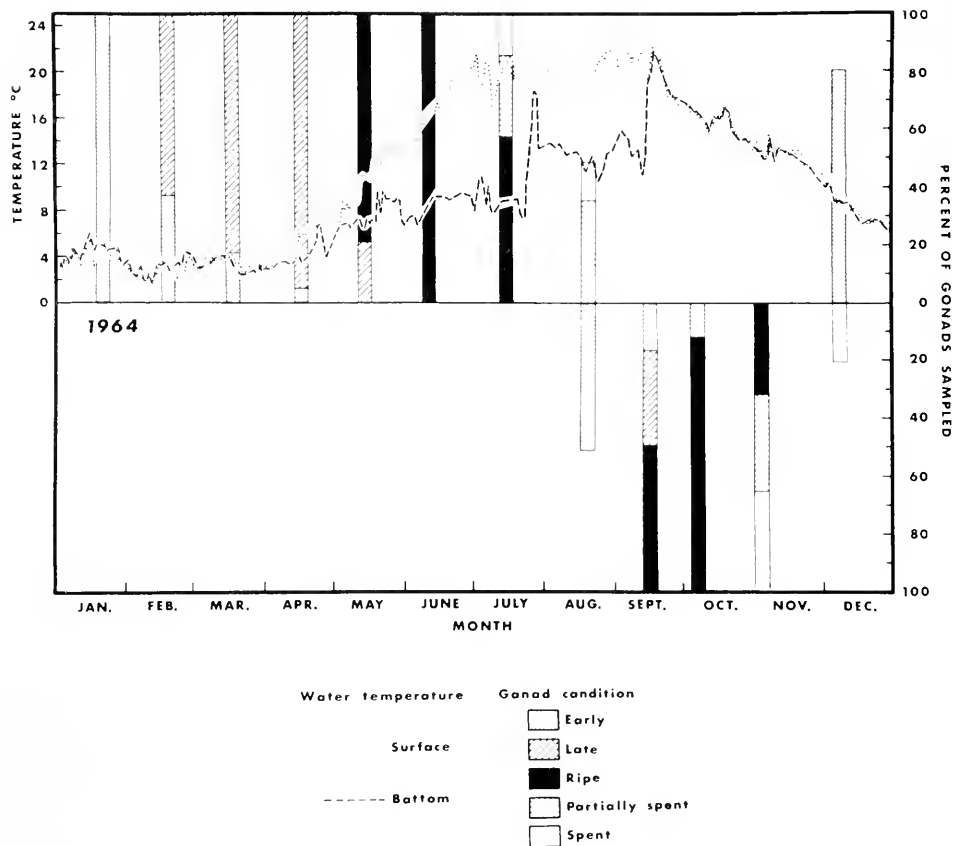


FIGURE 17. Gonad conditions of *Spisula solidissima* from New Jersey and sea-water temperatures at Barnegat Lightship in 1964.

and August only 7% to 11% of the clams were partially spawned and from mid-September to mid-October 29% to 52% of the clams were partially spawned or spent. Thus a mid-year spawning observed during the 3 previous years of the study was delayed 2 months in 1965. A few clams (7% to 13%) were in the spawning condition in late November and mid-December. From mid-October through December, however, many clams (48% to 93%) were in the early active phase. No second reproductive cycle was observed.

Low winter temperatures were approximated from the surface observations because bottom temperatures were not available until early May. Warming to

about 6° C. in May was followed by temperatures of about 8° C. in June and July; fluctuations of 2° C. occurred but the temperature did not exceed 10° C. Temperatures of 11° C. were recorded on three occasions in August. Temperature rose sharply from 10° C. to 18° C. in mid-September. Unlike the sequence in the 3 earlier years, spawning apparently occurred after the sharp rise in temperature.

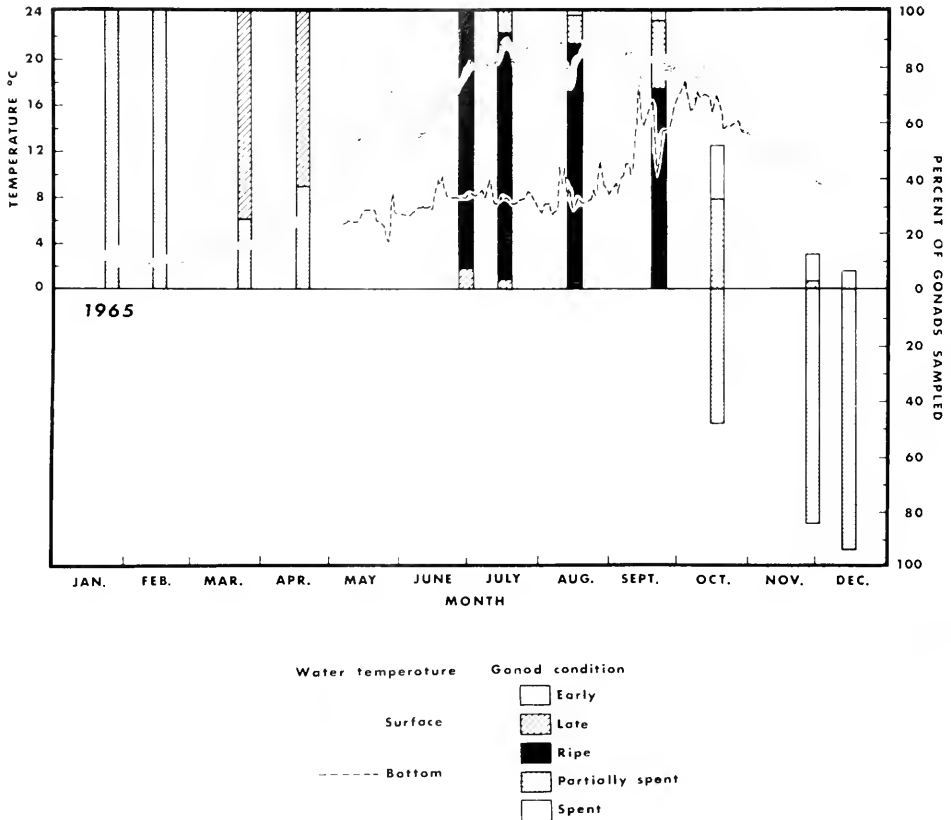


FIGURE 18. Gonad conditions of *Spisula solidissima* from New Jersey and sea-water temperatures at Barnegat Lightship in 1965.

DISCUSSION

In this study the term "reproductive cycle" includes the earliest detection of sex cells during periodic histological examinations of gonad tissue, the growth of female sex cells and multiplication of male sex cells which eventually fill the gonadal tubules, and the disappearance of sex cells from the tubules during the spawning period. The reproductive cycle can be separated into the gametogenic period (*i.e.*, the early, late, and ripe phases) and the spawning period (*i.e.*, partially spawned and spent phases). No period of inactivity was apparent in the sections of surf clam gonads, since sex cells for a succeeding reproductive cycle were detected in

spent individuals (Figs. 6 and 12). In fact, separating the reproductive cycles of some marine invertebrates into an "inactive phase," "rest period," "quiescent period," or "vegetative phase" may only be a convenience (Ropes and Stickney, 1965). Biological processes maintaining life and early gametogenesis could very well be taking place during the inactive phase and be undetected by the histological technique used.

Examinations of the slide preparations of gonads provide an accurate basis for describing the reproductive cycle of surf clams from offshore New Jersey. Sex cell development began during December and, concurrent with the enlargement of the gonads, was followed by the occurrence of numerous partially spawned and spent clams during July and August each year from 1962 through 1964. A second maturation of sex cells followed the mid-year spawning, but gonad distention was minimal. Partially spawned and spent clams in the mid-October to early November samples in 1962 through 1964 was evidence of a second annual spawning. Although a biannual reproductive cycle may be typical for surf clams from offshore New Jersey during most years, a single reproductive cycle and spawning in 1965 was an important exception. The time and frequency of spawning each year, then, can vary.

Detection of a biannual reproductive cycle in surf clams from offshore New Jersey would be difficult without microscopic examinations of gonads throughout several calendar years. Westman and Bidwell (1946, see footnote 1) reported a mid-year spawning of surf clams from offshore Long Island, New York, by observing gonad distention and shrinking after spawning, but their method of visually determining gonad condition might not be precise enough to identify a second annual spawning, if one occurred. Allen (1951, 1953) observed a scarcity of eggs in surf clams from Woods Hole, Massachusetts, during 1 to 2 weeks in mid-summer coincident with a period of excessive heat. A possible explanation is that the clams had spawned or, since he continued to collect eggs for a 2- to 3-month period after mid-summer, these eggs may have been produced during a second annual reproductive cycle.

Temperature frequently influences the reproductive cycle and spawning of marine invertebrates (Giese, 1959; Loosanoff and Davis, 1963). During the gametogenic period, the surf clams in the beds off New Jersey were surrounded by gradually increasing, but fairly stable temperatures for the first six months in the three early years of the study. A progressive development of the gonads to a turgid and ripe state during the period of warming to temperatures of 12° C. or a little less preceded the first annual spawning. In 1965, however, the water temperatures during the gametogenic period were generally colder (8° C. or less) and more erratic (highs to 11° C.) than in the previous years. Gametogenesis was slightly delayed by the lower temperatures—94% of the clams were in the ripe phase by late June, 1965, whereas 100% were in the ripe phase by mid-June of 1962, 1963, and 1964.

Temperature was not clearly a stimulus to spawning of surf clams in their natural beds off New Jersey. Spawning by some clams did not accompany abrupt temperature changes in all 4 years, although more clams spawned before the abrupt temperature change in 1962-64 than in 1965. Spawning by some bivalves can be effectively delayed by holding them at low temperatures. Conversely, gonad development to a ripe condition has been hastened by holding bivalves in warm sea water.

The ripe bivalves have been spawned "out of season" and have produced normal larvae (Loosanoff and Davis, 1963).

I have held surf clams in 20° C. water from late February to mid April, 1967 and compared their gonadal development with that of clams from the parent population in Chincoteague Inlet, Virginia. Gonads of all clams in the inlet were in the late active phase in late February and remained in this condition until after April. Water temperatures during the experimental period gradually increased from 5° C. to 11° C. In contrast, 10 gonads from clams held in the warmed water were all in the ripe phase by mid-March. On 11 April, a female spawned spontaneously in one of the containers and 2 partially spawned individuals were found when gonad sections of 10 clams were examined. These three specimens, then, spawned without the stimulus of a thermal shock and were perhaps like the ripe individuals from the beds off New Jersey that spawned before an abrupt temperature change.

The surf clam apparently develops its gonads to a ripe condition within a short period of time, contains ripe sex cells for several months of the year, and responds readily to relatively simple spawning stimuli. A rapid gonad development may account for the report by Carson (1945) that the surf clams of the mid-Atlantic coast spawn in the spring and throughout the summer. Allen (1951, 1953) obtained eggs of New Jersey coast clams from early spring to late autumn. Schechter (1941) collected ripe surf clams from areas near Woods Hole, Massachusetts, throughout the summer. Loosanoff and Davis (1963) found morphologically ripe eggs and spermatozoa in surf clam gonads from mid-January to the end of June. Conditioning the clams in the early months of the year by placing them in warmed water produced ripe specimens. Later in the season, when the natural waters had reached 12° C., no conditioning was needed, and thermal stimulation alone was sufficient to induce spawning. In contrast, other species of bivalves in the laboratory require various physical and chemical stimuli such as electrical shocks, rhythmic changes in water temperature, the addition of ammonium hydroxide to the water or injected into the gonad, or an introduction of egg or sperm suspensions to the water to induce spawning—and even then some species do not respond (Galtsoff, 1961, 1964; Loosanoff and Davis, 1963; Stickney, 1964).

The importance of nervous and hormonal interrelationships in bivalves has been reviewed by Galtsoff (1961, 1964) and Giese (1959). Products of neurosecretory cells inhibit spawning, and bivalves become sensitive to environmental spawning stimuli after the cells disappear. Degeneration of the neurosecretory cells begins a few days before spawning. Nagabhushanam (1963a) recognized two types of neurosecretory cells in the cerebral and visceral ganglia of *Spisula solidissima*. The cells in surf clams appear morphologically similar to those in the oyster, *Crassostrea virginica*, and the mussel, *Modiolus demissus* (Nagabhushanam 1963b, 1963c, 1964). Experiments with the latter two species link the cells with the hormonal control of the gametogenic cycle and spawning. Stimuli such as low salinities, electrical shocks, and high temperature influenced the secretions.

Both Galtsoff (1961) and Stickney (1964) believed that ripeness, rather than specific temperature, influences the response of some bivalves to spawning stimuli. Ripeness and sensitivity to spawning stimuli seem to precede the natural spawning period of surf clams by many months. I classified surf clams from New Jersey as ripe for at least two months each year before the first annual spawning. Some

were induced to spawn in late May and early June, 1965, by simply placing them in running sea water aboard a research vessel. At four locations where the clams were caught, the water temperature increased as little as 9.5° C. and as much as 11.5° C. from bottom to surface. Changes in hydrostatic pressure or shocks from being caught by the jet dredge might stimulate spawning, but none of the clams from a fifth location spawned when the temperature change was only 0.8° C. A response to the stimuli of an abrupt change in temperature, well in advance of the seasonal time of spawning, suggests that the hormonal control of spawning can be overridden in surf clams. Nevertheless, a minimum temperature threshold for a spawning response is probably necessary.

Sections of surf clam gonads from geographic locations other than the commercial fishing areas off the New Jersey coast were examined, since a biannual reproductive cycle and spawning may not be typical throughout their range. In monthly samples from a more northern habitat—Sagadahoc Bay, Maine—during May through September, 1962, only a single reproductive cycle was apparent. Partially spawned and spent clams in the August and September samples were evidence of a late summer and early fall spawning.

In other bivalves, the reproductive cycle changes with latitude. Pfitzenmeyer (1962, 1965) and Shaw (1962, 1965) observed a biannual reproductive cycle in soft-shell clams, *Mya arenaria*, from Chesapeake Bay, Maryland, the southern part of their range. Ropes and Stickney (1965) reported on a single annual cycle in soft-shell clams from areas north of Cape Cod, Massachusetts, and discussed the available evidence of a biannual cycle and spawning south of Cape Cod. Porter (1964), who described the seasonal gonadal changes in hard clams, *Mercenaria mercenaria*, from North Carolina, found two annual spawning periods. Both of these species apparently have only one reproductive cycle each year, but two in southern waters. The change in the breeding season with respect to latitude and an extension of the season as one goes south has been observed in many temperate marine invertebrates (Giese, 1959).

I wish to gratefully acknowledge the cooperation of the members of the surf clam industry and many of the commercial fishermen who welcomed me aboard their vessels to sample clams from their catches at sea. The technical assistance of Virginia A. Liddell, Sally V. Otto, Brenda Spriggs, and Geraldine J. Fitzgerald aided me in the preparation of the many slides for the study. Dr. Rudolf S. Scheltema, Woods Hole Oceanographic Institution, Woods Hole, Mass., reviewed the manuscript and made many helpful suggestions. Dean F. Bumpus, Woods Hole Oceanographic Institution, Woods Hole, Mass., furnished the unpublished temperature records at Barnegat Lightship for 1964 and 1965.

SUMMARY

1. A biannual reproductive cycle was observed in *Spisula solidissima* gonads during 3 of the 4 years in which samples were taken from offshore New Jersey.

2. The biannual cycle was characterized by a major mid-year spawning and minor late-year spawning. A second annual reproductive cycle may be neither typical for surf clams throughout their geographical range nor always an annual event in clams from New Jersey.

3. A delayed spawning and single annual cycle in 1965 was related to colder environmental temperatures than were observed in the three earlier years of the study.

4. Abrupt temperature changes were not clearly a cause of spawning in the natural populations of surf clams. Partially spawned clams were found in the samples in advance of abrupt temperature changes. Temperature, however, clearly delays or hastens the gametogenic cycle and stimulates spawning of ripe surf clams in the laboratory.

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THE GROWTH AND SIZE DISTRIBUTION OF CURRENT YEAR CLASS *LOLIGO PEALEI*¹

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Loligo pealei (Lesueur, 1821) is the common squid between Cape Cod and Cape Hatteras (Verrill, 1882). It is abundant inshore during the warmer part of the year and is concentrated near the Continental Shelf break in the winter time (Summers, 1967). In spite of its ubiquity, its apparent importance in pelagic food chains and its significance as a source of "squid giant axons," the life history of *L. pealei* has not been fully described.

Verrill (1882) and Haefner (1964) both estimated the growth rate of this species by equating different year classes with size-frequency modes. By this method, Verrill suggested that the largest individuals were three or four years old and Haefner demonstrated two distinct year classes. Their analyses were subjective when the size classes overlapped each other and contained the assumption that *L. pealei* grows at a continuous, if not uniform, rate through its life span.

Verrill's report was based upon a synthesis of preserved specimens and was presented as a description of the growth rate of *L. pealei* off the "southern coast of New England." His tabulated data included measurements of approximately 3378 individuals, more than 2600 of which were collected in Vineyard Sound, south of the Cape Cod peninsula. Haefner cited morphometric data on 1165 individuals taken in Delaware Bay during the summer of 1958. His size-frequency information was pooled over the collecting period and consequently cannot be used to demonstrate short-term growth. Neither study fully accounted for the sampling bias of the collecting gear.

The abundance of *L. pealei* in Vineyard Sound during the warmer part of the year is known from Verrill's collections and from the work of Sumner, Osburn and Cole (1913). The latter dredged specimens throughout the waters surrounding Woods Hole, Massachusetts, and indicated a concentration in the southwestern part of Vineyard Sound. During the past five summers, the fishing vessel CAP'N BILL IV, under charter to the Marine Biological Laboratory, has trawled for this squid on 181 different occasions in the southwestern part of Vineyard Sound.

The present paper attempts to more precisely describe the early growth and size distribution of *L. pealei* in Vineyard Sound. The study was facilitated by dealing primarily with the numerous, rapidly growing current year class (individuals less than one year old) and by the fact that Vineyard Sound is near the northern range limit of this species where the breeding season is relatively

¹ Contribution number 156 from the Systematics-Ecology Program, Marine Biological Laboratory. This research has been supported by NSF Grant GB-4509 to the Systematics-Ecology Program and by the Marine Biological Laboratory.

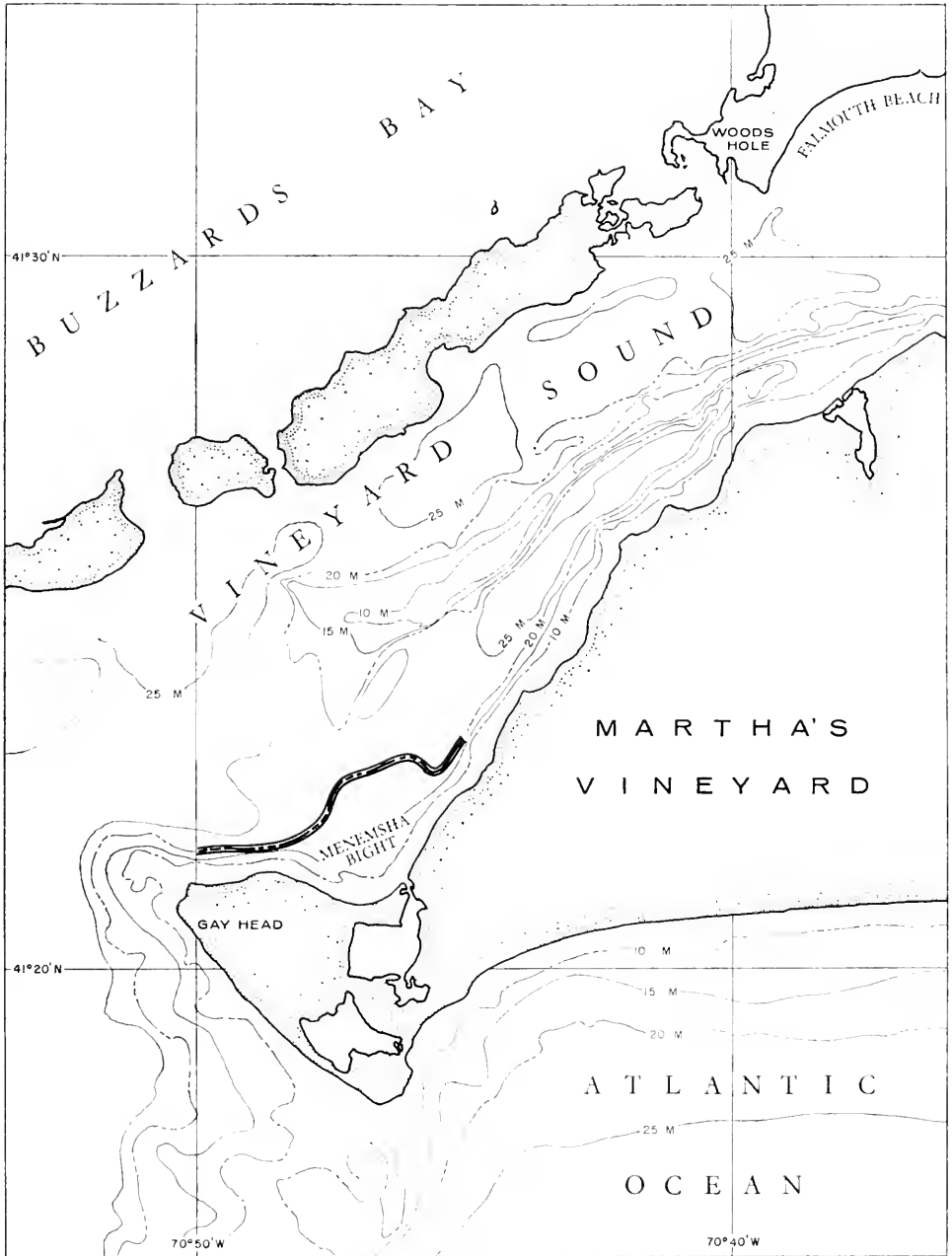


FIGURE 1. A contour map of Vineyard Sound. Menemsha Bight is located at $41^{\circ} 22' N$, $70^{\circ} 48' W$. The mean water depths are given in meters; the sampling region is shown by a heavy 20-m. contour line in Menemsha Bight.

short. Sampling was restricted to one location and procedures were consistent throughout the study in order to determine seasonal changes. This work was undertaken as a part of a continuing study of the ecology of *L. pealei*.

METHODS

The growth rate and longevity of *L. pealei* could not be determined in the laboratory because this animal can not be reared in captivity. A program of tagging and recapture of squid in the field did not seem practical short of a large scale operation. The remaining option was to repeatedly determine the size-frequency of a natural population and infer growth from the changes between sampling dates. This method was employed knowing that it introduced the least experimental bias in growth estimates and required the greatest care in sampling from a single population.

The site chosen for this study was Menemsha Bight (41° 22' N, 70° 48' W), a shallow open embayment at the southern extreme of Vineyard Sound, west of Martha's Vineyard Island. As illustrated in Figure 1, Menemsha Bight is partially protected from the open waters of the Atlantic Ocean by the Gay Head promontory. The Bight has an unobstructed, gently sloping sandy bottom and normally does not have tidal currents in excess of 3 km./hr. There is little temperature stratification in Vineyard Sound due to significant water movement and the annual surface temperature cycle ranges from 0 to 22° C. in February and August, respectively. Sea-water salinities remain consistently at 31 to 32‰ through the year. The sampling region is shown by a heavy 20-m. contour line in Figure 1. This region was accessible for otter trawling for a linear distance of approximately 5 km.

Squid were taken with a #35 otter trawl towed by the stern rigged, 65-foot (20-m.) research vessel A. E. VERRILL (named for the author cited above). Except for the addition of a 1.25-inch mesh (3.2-cm.) stretched measure nylon liner in the cod end, this was a typical New England commercial groundfish net. The trawl doors weighed 500 lb. (225 kg.) apiece and each was separated from the net by 12 fathoms (22 m.) of groundlines and legs. The headrope and footrope measured 52 and 72 feet (16 and 22 m.), respectively and the forward part of the net was made of 42-thread nylon twine with 5-inch mesh (13-cm.) stretched measure in the wing and square sections and 4½-inch mesh (11-cm.) stretched measure in the belly and cod end. The headrope had 12 8-inch (20-cm.) aluminum floats spaced along its length and the foot rope was protected by 4- and 6-inch (10- and 15-cm.) rubber rollers.

All of the samples were taken near midday (earliest 1031 hours, latest 1422 hours) and the tows averaged 50 minutes in duration. The towing speed was 6.5 km./hr. except on the first two sampling dates when a slightly slower speed was employed. The contents of the net were dumped into large tubs on the deck and sorted immediately. The entire squid catch, or a representative sample of several hundred individuals, was measured and sexed as soon as possible after capture. No preserved specimens were included in the results.

The dorsal mantle length was recorded for every individual in the sample; squid less than 5 cm. were measured to the nearest whole millimeter, larger squid were measured to the nearest whole centimeter. The ventral mantle wall was slit open and the sex was recorded for all but the smallest, immature individuals. Males

were considered mature when spermatophores were present; this was always accompanied by a thickening and white coloration of the vas deferens. Females were considered mature when the ovary was expanded and loose eggs were found in the oviduct.

RESULTS

A total of 1619 individuals were measured from samples collected in Menemsha Bight on seven consecutive dates ranging from August 18 to November 16, 1967. The numbers of squid caught on November 2, 16 and 22 were approximately 1000, 5 and 0, respectively, and none were taken through March of the following year. *L. pealei* was the only species of cephalopod taken. Many species of fish were caught, some in abundances of several hundred pounds per tow.

It is apparent from the size-frequency data for squid taken on the first sampling dates that current year class *L. pealei* were represented by a few small individuals mixed in with the larger size classes. This class grew and became more dispersed in later samples and eventually merged with the larger classes in late September. The current year class was initially characterized by a narrow size range and illustrated a marked skewness toward the larger sizes. Examination of the various size-frequencies showed that the current year class and the older classes were adequately approximated by lognormal size distributions.² Thus, it was possible to more rigorously separate the size classes on lognormal probability paper using the method described by Cassie (1954, 1962). Appropriate age categories could be assigned to these size classes by contrasting the collections on different dates.

The best fit size distributions for six sampling dates are shown in a lognormal probability presentation in Figure 2. Age classes are indicated by brackets to the right of the best fit lines in this figure. The amount of class overlap can be seen in Figure 2 by the horizontal coincidence of different age classes on a single sampling date. It should be noted that Cassie's method of size class separation is based upon size distributions and is little affected by exceptional individuals or the location of the modal size. The latter generally occur at a point below the cumulative 50% in Figure 2.

The current year class made up an increasing proportion of the samples during the time covered by this study. The percentage of current year class individuals in the measured samples is shown for various dates in Figure 3. The dashed line refers to a sizeable catch of *L. pealei* (308 individuals) taken with a 16-foot (5-m.) shrimp trawl under the same conditions give above. There were no current year class squid in this sample and the one plus year class had 5th, 50th and 95th percentile lognormal fit mantle lengths (equivalent to the ends and middle of the lines shown in Figure 2) of 6.8, 9.3 and 12.7 cm., respectively. The dotted line on November 16 indicates the last date of squid capture for the 1967 season.

Figure 4 shows a growth scheme for the current year class *L. pealei* based upon the best fit lognormal size distributions illustrated in Figure 2. The bars

² A lognormal size distribution is one in which the logarithm of the variate (in this case, mantle length) is normally distributed. Size distributions in this report are all positively skewed, implying that the modal size lies closer to the lower range size than the upper range size. A positively skewed lognormal distribution restricts the variate to values greater than zero.

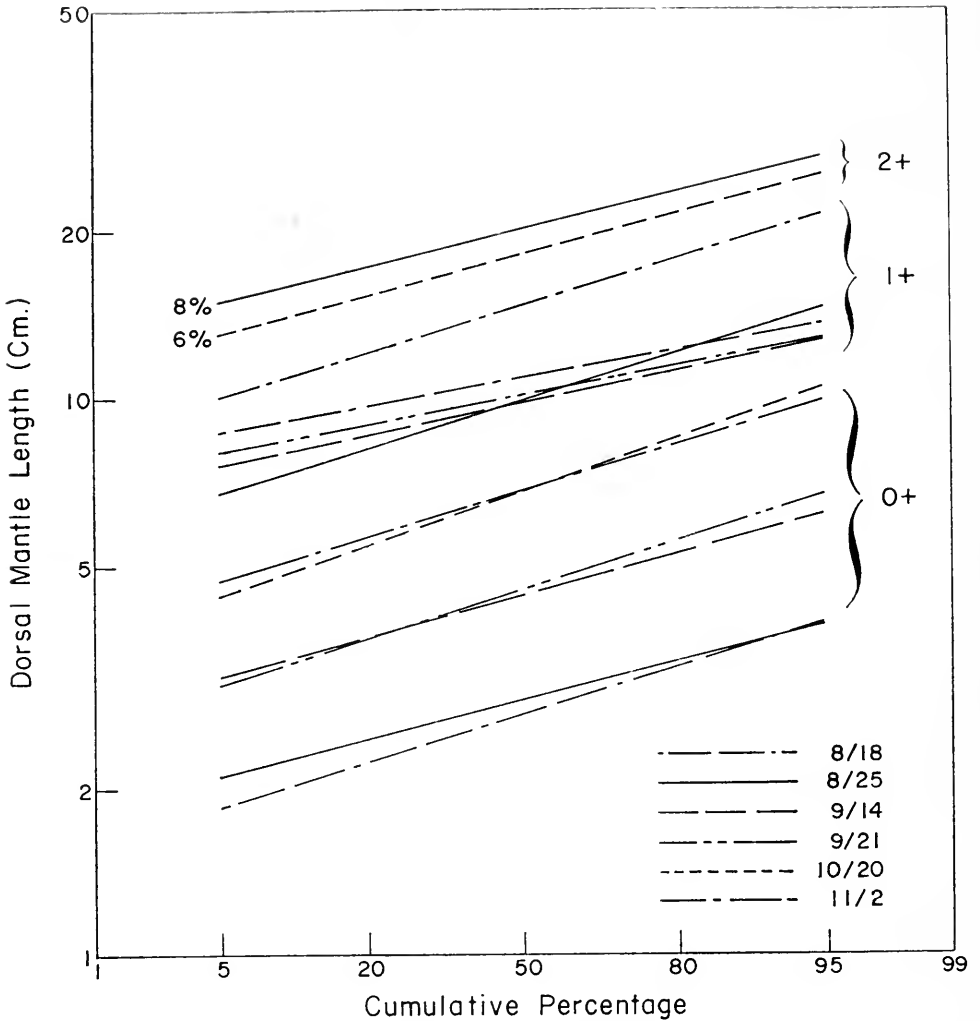


FIGURE 2. Best fit lognormal size-frequencies for different age classes of *L. pealei* resulting from six collections. This is a lognormal probability presentation of the data. Age classes in years are shown by brackets to the right of the fit lines. Sampling dates are indicated by the line codes given in the figure legend. The percentage occurrence of two plus year class individuals in the samples is shown by numerals to the left of the fit lines.

extend from the 5th to the 95th percentile sizes and the dots represent the median, or 50th percentile size. Linear regression lines are given each of these percentages and drawn out from the hatching size (2 mm. mantle length) to the date of last squid capture, November 16. The stippled area in this figure represents Verrill's conclusion on the growth rate of *L. pealei* off the "southern coast of New England." These reports do not agree in date, growth rate nor size distribution.

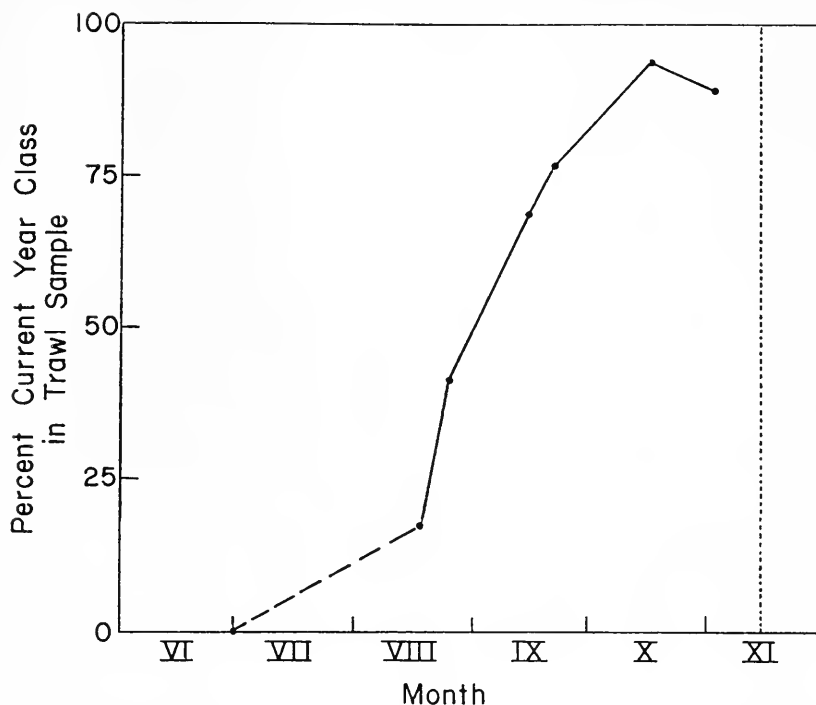


FIGURE 3. Percentage of current year class individuals in trawl samples for various dates. The dotted line on November 16 indicates the last date that *L. pealei* was captured in Menemsha Bight before its fall migration. A dashed line connects with data from a collection made with a shrimp trawl.

DISCUSSION

The accuracy of the foregoing estimates of growth rate and size distribution is dependent upon how well the measured samples represent the natural squid population. Owing to the mobility and seasonal migration of *L. pealei*, one can readily question the adequacy of daytime, otter trawl samples from a single depth zone and location. Furthermore, the size bias introduced through the collecting gear needs to be established in interpreting the results.

The consistency of the squid population in Menemsha Bight can only be implied by the regularity of sample size-frequencies and by comparison with other sources. The latter produced no significant differences when applied to records of miscellaneous collections on comparable dates in 1967 from Buzzards Bay, Vineyard Sound and Nantucket Sound, resulting from otter trawling, fishtrap hauls, squid jigging and fish stomach contents. Winter offshore collections of *L. pealei* show an increase in the numbers of individuals trawled in daylight hours and a positive correlation between depth and mean mantle length. This last factor may explain the size constancy of the one plus year class shown in Figure 2, by suggesting that this class may have been migrating through the sampling region during the study.

L. pealei does migrate vertically. For instance, newly hatched fry congregate near the surface in the best lit part of an aquarium and, during the warmer part of the year, adult squid are frequently seen swimming near the surface at night. The #35 otter trawl fishes in the first one to two meters above the bottom and it may well miss squid in upper strata. I suspect that the largest squid either swim high enough above the bottom to be missed by this net or are capable of avoiding

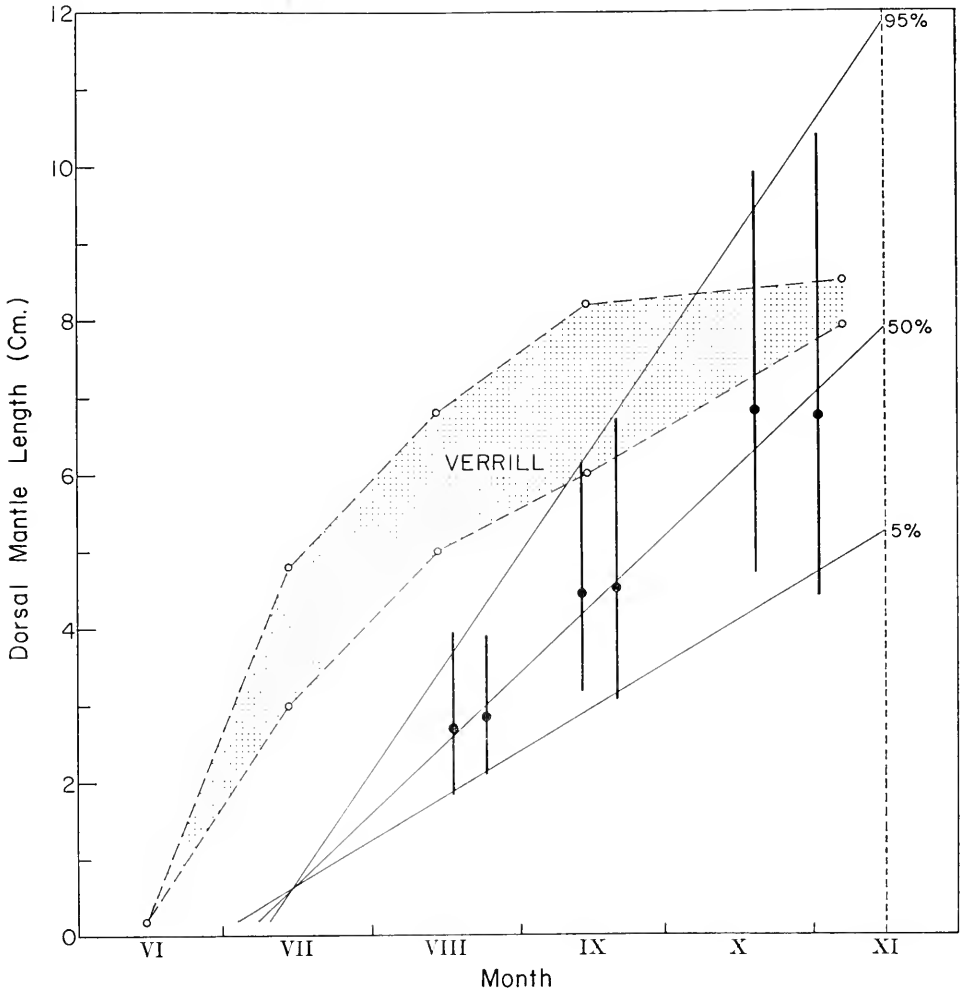


FIGURE 4. Two growth schemes for current year class *L. pealei*. The dorsal mantle length in centimeters is shown as a function of time. Verrill's data (1882) are indicated by a stippled envelope including the typical size range of his collections. The present study is represented by bars and dots showing the 5th, 50th and 95th percentile lognormal fit mantle lengths on six sampling dates. Linear regression lines are given for each of these percentile sizes. The dotted line on November 16 indicates the last date that *L. pealei* was captured in Menemsha Bight before the fall migration.

it, as the biggest individuals I have measured (including one male with a dorsal mantle length of 46.5 cm.) were taken in a fishtrap.

Selectivity

The selectivity of an otter trawl refers to the fact that it may retain individuals of different sizes with different efficiencies. Obviously, small individuals can slip through the net meshes and larger ones will be retained. Thus, selectivity is specified by the size *vs.* retention characteristics of a particular mesh size and species. This matter is discussed for several species of fish by Clark (1963).

Otter trawl selectivity can be estimated by comparing the catches of netting with different mesh sizes either on the same trawl or on similar nets fishing under comparable conditions. Such a comparison was made between the current year class *L. pealei* taken with the #35 net in Menemsha Bight on August 18 and two collections made by the Bureau of Commercial Fisheries R/V ALBATROSS IV which trawled in the same area at midday on August 17 and 18. The ALBATROSS IV used a #36 otter trawl with a $\frac{1}{2}$ -inch mesh (1.5-cm.) stretched measure nylon liner in both the cod end and upper belly; it was 4 feet (1.2 m.) longer in each wing than the #35 net and more heavily protected for fishing on rough ground, but otherwise the nets were identical.

There were a total of 319 current year class individuals in these three collections, only 21 of which were caught in the #35 net. A comparison did not show a size bias in the #35 net sample and suggested that the entire size class straddled the transitional retention zones of the two trawl liners which had caused the #35 net to be uniformly less efficient in retaining this collection of squid. The 5th, 50th and 95th percentile lognormal fit mantle lengths for the ALBATROSS IV collections of August 17 and 18 and the A. E. VERRILL collection of August 18 were: 1.7, 2.2, 2.9; 1.9, 2.6, 3.4 and 1.9, 2.7, 4.0 cm., respectively.

The selectivity of the #35 net is probably the single most important factor in the explanation of the sample size class composition illustrated in Figure 3. As the current year class *L. pealei* grew, it was more efficiently retained by the net, and made up a larger proportion of the total catch. By assuming that the current year class made up 100% of the natural squid population and replotting Figure 3 on the basis of mantle length *vs.* per cent current year class in the samples, the 50% retention point for the #35 net was estimated to occur at a mantle length of 3.7 cm. This is consistent with the previous comparison and with Clark's data for fish.

Size distribution

Lognormal size-frequency distributions are not uncommon in juvenile animals and tend to occur in older animals where several age classes overlap in size and where mortality has reduced the abundance of the older classes. The mathematical philosophy of this distribution is contained in a monograph by Aitchison and Brown (1957); I will suggest some of the factors which may relate to the present case.

Net selectivity can cause the truncation of a size-frequency distribution if the smaller individuals are retained less efficiently than the larger ones. The A. E.

VERRILL collection of August 18 lay almost entirely below the 3.7 cm., 50% retention point, of the #35 net, and could be lognormally distributed as a result of net selectivity. Theoretically, the #36 net collections of the same and the previous day should have been influenced by a 50% retention point smaller than 3.7 cm. by the ratio of the liner mesh sizes (approximately 1.0 cm. mantle length when the twine diameters are taken into consideration). For practical purposes, the #36 net should not have shown a selectivity bias in the current year class collections of those dates. The size-frequencies of the ALBATROSS IV collections were strongly lognormal, indicating the condition of the natural population. This is borne out by the maintenance of lognormality in later collections and in older year classes.

A sexual dimorphism in the size-frequency relationship accompanies the onset of maturity in *L. pealei*. Haefner placed the inception of this dimorphism at a mantle length of 9.5 cm. and Verrill concluded that it took place at an age of eight months or a size of 6.2 to 18.8 cm. Thus, each of the older year classes shown in Figure 2 is a sum of two overlapping sex classes; the mature males generally larger than the females. The introduction of additional classes may help to explain the lognormality of the summed older year classes taken in contrast to the current year class. It was this contrast which made Cassie's method useful in the present study.

The relative abundance of the sexes did influence the size-frequencies of the samples. There were almost three times as many females as males in the one plus year class taken with a small trawl on June 30. This ratio was reduced to 2:1 on August 18 and remained approximately 1:1 thereafter. The two plus year class was composed mostly of mature males on August 25. Non-current year classes on October 20 and November 2 were poorly represented and scattered in size; as a result, their placement as two plus and one plus year classes, respectively, is tentative and based upon maturities. The current year class remained sexually immature throughout the sampling period, as did approximately one-half of the one plus year class. Spent females were rare in the collections.

Growth rates

The upper size limit in Verrill's growth scheme, as illustrated in Figure 4, was apparently constructed from the interface between mature and immature individuals listed in his tabulated results. It was his assumption that *L. pealei* matures in one year, an assumption that requires an individual to be a mature participant in 3 to 4 breeding migrations if it lives to that age. The size class separation shown in Figure 2 suggests that about half of the one plus year class does not mature short of the second fall migration. This, undoubtedly, explains part of the difference in growth schemes shown in Figure 4. It is instructive to note that the west coast species, *Loligo opalescens*, was reported to die shortly after breeding (McGowen, 1954; Fields, 1965; Hobson, 1965). Such mortality has not been demonstrated for *L. pealei*, but it is inconsistent with Verrill's growth scheme if it does occur.

As shown in Figure 4, the linear regression lines fit to the 5th, 50th and 95th percentile sizes intersect on July 16 close to the hatching size, 2 mm. mantle length (see Arnold, 1965). Without taking into account the changes in body

proportions which occur soon after a squid hatches, one can use this extrapolation to suggest that the probable hatching date for the current year class squid used in this study was approximately July 9, 1967. If allowance is made for a relative increase in mantle length to total length soon after hatching, the probable hatching date would be a few days later. Surface water temperatures in Vineyard Sound ranged from 15 to 19° C. between mid-June and mid-July, 1967. A collection of freshly laid squid egg capsules taken in 14 m. of water northwest of Gay Head on September 19, 1967, hatched in 20 days in the laboratory at water temperatures of 16 to 18° C. Thus I estimate that the egg deposition which produced the current year class squid in this study took place about June 19, 1967.

TABLE I
Early growth rate of three loliginid squid

Species	Location	Month hatched	Growth rate, mm./month to given age	Reference
<i>L. vulgaris</i>	North Sea	VII	20 mm., 6 months*	Tinbergen and Verwey, 1945
<i>L. vulgaris</i>	Mediterranean Sea	VI or VII	16.3 mm., 3 or 4 months 13.6 mm., 5 or 6 months	Mangold-Wirz, 1963
<i>L. opalescens</i>	Monterey Bay, California	V	6 mm., 12 months	Fields, 1965
<i>L. pealei</i>	Vineyard Sound, Massachusetts	VI	30-48 mm., 1 month 25-34 mm., 2 months 20-27 mm., 3 months 16-17 mm., 4.75 months	Verrill, 1882
<i>L. pealei</i>	Menemsha Bight, Vineyard Sound, Massachusetts	VII	17.8 mm., 4 months†	Summers, this paper

* Tinbergen and Verwey reported ventral mantle lengths which are slightly shorter than the dorsal mantle lengths given by the other authors.

† This is the growth rate of the median, or 50th percentile size. The 5th and 95th percentile sizes have growth rates of 11.2 mm. and 27.5 mm. per month for the first four months, respectively. These three growth rates are illustrated by the linear regression lines in Figure 4.

Verrill gave the date of first significant hatching as "the second week in June." The difference between his date and the one given above may reflect a difference in years (1875 *vs.* 1967), different collecting sites, his acquaintance with Long Island Sound or a combination of these factors. Adult squid were abundant in Vineyard Sound by June 1, 1967, when several fishermen began catching them regularly. Sizable accumulations of squid egg capsules were taken in the Bureau of Commercial Fisheries fishtrap in Buzzards Bay on June 3 (Mr. C. L. Wheeler, personal communication) and from Falmouth Beach on June 17, 1967. Verrill's mid-June hatch seems distinctly early for the 1967 season.

I take these isolated collection of egg capsules and the narrowness of the current year class size distribution to be an indication that *L. pealei* breeding activity is highly localized and that the breeding season is made up of a number

of these events. This is a logical consequence of Arnold's observation on the mating behavior of this species (1962), which indicated the importance of a visual stimulus in egg deposition.

The early growth rate of three loliginid squid is given in Table I. The first four entries are based upon modal analyses. With the exception of Fields' data, all of the entries are roughly comparable over the first few months of growth.

SUMMARY

1. This paper reports estimates of the growth rate and size distribution of current year class *L. pealei* based upon the population statistics of daytime, otter trawl collections in Menemsha Bight during the second half of 1967.

2. The selectivity of a #35 otter trawl with a 1.25-inch mesh (3.2-cm.) stretched measure cod end liner was estimated by a 50% retention of *L. pealei* with a dorsal mantle length of approximately 3.7 cm.

3. The current year class, summed older year classes and individual older year classes of *L. pealei* were found to have lognormal size-frequency distributions.

4. The current year class could readily be separated from the older year classes by the use of Cassie's method on lognormal probability paper.

5. The growth rate of a median individual in the current year class of 1967 was found to be 1.8 cm. dorsal mantle length per month for the first four months past hatching.

6. Approximately one-half of the one-year-old *L. pealei* did not mature sexually before the second fall migration. None of the current year class *L. pealei* matured before the first fall migration.

7. The results indicate that *L. pealei* egg deposition is isolated in time and location, and repeated throughout the Vineyard Sound area from at least June through September.

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FORM AND HABIT IN SPECIES OF MALLEUS (INCLUDING THE
"HAMMER OYSTERS") WITH COMPARATIVE OBSERVA-
TIONS ON ISOGNOMON ISOGNOMON

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Probably no tropical bivalve shells have been better known than those of the well-named "hammer oysters." The black species appears as *Ostrea malleus* in the 10th edition of the *Systema naturae* of Linnaeus. The genus *Malleus* was erected by Lamarck in 1799 for the inclusion of this species and later for the white *M. albus*. These shells were highly prized by early collectors; Dance (1966) records the sale of shells of *M. malleus* for 32 guilders in Holland and 240 francs in Paris during the eighteenth century. The distinctive, indeed exaggerated, hammer shape of these two species has not proved characteristic of the genus (as defined by Thiele (1935)) unless those smaller species without the anterior and posterior extensions of the hinge line be placed in separate genera, such as *Parimalleus* (Iredale, 1939).

Despite long knowledge of and interest in the shells, effectively nothing is known about the mode of life of any species of *Malleus*. Coming from coral reef areas they have been regarded as being attached to them. Thus Iredale (1939) in his account of the bivalves collected during the course of the Great Barrier Reef Expedition refers to *M. malleus* as an inhabitant of coral reefs but states that *M. albus* is not a reef shell but "the mainland species." Although some account of the structure of species of *Malleus* has been given by Pelseneer (1911) and Kühnelt (1938), no one appears ever to have observed these animals in life.

Personal opportunity of so doing was afforded during the sixth cruise of the Stanford University Research vessel "Te Vega" which the author accompanied from Singapore to the Solomon Islands from January to March, 1965. Some additional, and very valuable, information was obtained during some four weeks spent on the Belgian "De Moor" Expedition to the Great Barrier Reef of Australia during September to October, 1967. Acknowledgments for these facilities are given later.

Detailed examination of the smaller, *M. regula* (Forskål), without extended hinge line, provided the basis for conclusions on the adaptations and habitats of *M. malleus* (Linn.) and *M. albus* Lam., specimens of which were obtained from various sources and which undoubtedly represent successively further stages in adaptive evolution, essentially from epifaunal to infaunal life. The striking fact that *M. regula* may occur in vast numbers mingled with similar populations of *Isognomon isognomon* (Linn.), a species of a not very closely related genus which has also taken to vertical posture after byssal attachment, involved an illuminating parallel study. Both species exhibit most interesting consequences of the mono-

¹ From the Te Vega Expedition, Cruise 6.

EXPLANATION OF LETTERING

A	anus	MI	mantle isthmus
ABR	anterior byssal retractor	ML	multivincular ligament
AD	adductor	N	limit of nacreous region
AE	anterior extension of shell	OBG	opening of byssal gland
AF	accessory foot	OCL	outer calcareous layer
AOL	anterior outer ligament layer	OF	outer fold of mantle margin
AOM	epithelium secreting anterior outer ligament	P	periostracum
AU	auricle	PAR	pallial retractor
BG	byssal groove	PBR	posterior byssal retractor
BN	byssal notch	PC	pericardium
By	byssus	PE	posterior extension of shell
CT	ctenidium	PMC	promyal chamber
E	exhalant current	POL	posterior outer ligament
F	foot	POM	epithelium secreting posterior outer ligament
FL	fusion layer	PR	prismatic layer
I	inhalant current	R	rectum
ICL	inner calcareous layer	RI	ridge associated with pallial retractor
IF	inner fold of mantle margin	U	umbo
IL	inner ligament layer	V	ventricle
LP	labial palp		
MF	middle fold of mantle margin		

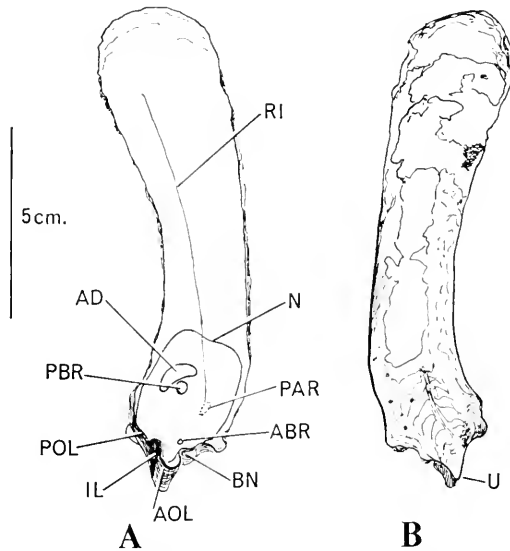


FIGURE 1. *Malleus regula*, right valve, **A**, inner, **B**, outer view. For explanation of lettering in this and all other figures see above.

myarian condition, description and discussion of which supplement a series of prior studies on other monomyarians.

The above statement explains the general plan of this paper in which descriptions of structure, illuminated where possible by discussion of function, precede an account of mode of life and general biology.

THE GENUS *MALLEUS* LAMARCK 1799*Malleus regula* (Forskål)

This species, widely distributed and very common in the tropical Indo-Pacific, was first personally observed at Palau Gaya, one of a half-circle of three precipitous islands forming the remains of the rim of a large extinct, now flooded, volcanic crater in Darvel Bay on the east of Sabah (North Borneo) which was visited on the *Te Vega* at the end of January, 1965.

Shell

This is straight (Fig. 1), about four times as long as it is broad and attains lengths of some 12 cm. Old shells tend to be very irregular with the outer surface (Fig. 1B) usually covered with calcareous algae and other encrusting growths. It stands more or less erect, firmly attached by byssus threads to a rocky, often partially mud-covered, surface. The prominent posteriorly directed umbones (U) are situated immediately on the posterior side of the byssal notch (BN). These mid-dorsal structures thus lie in the middle of the topographically under surface. This is true of all the species considered (including *I. isognomon*), which for this reason are figured with the umbonal (*i.e.*, "dorsal"), as well as the byssal (*i.e.*, "ventral"), surface undermost as they invariably occur in life.

The internal surface of the shell valve is shown in Figure 1A, and that of *M. malleus* in Figure 3. The nacreous or proximal area (delimited by N) occupies some quarter of the surface, the distal remainder (apart from some thin superficial periostracum) consisting of prismatic (outer calcareous) layer formed by the mantle margins. Conditions are precisely as in *Pinna* (Yonge, 1953b). In both, the mantle is capable of great extension, and corresponding withdrawal, and is able to make rapid repair following the frequent damage species of both genera experience. Within the space occupied by the visceropedal mass (where the nacreous, *i.e.*, inner calcareous, shell layer is secreted by the mantle) lie the scars of the somewhat crescentic adductor (AD) and of the larger posterior and the smaller anterior byssal retractors (PBR, ABR). There is also the fainter scar of the pallial retractor (PAR). From this a low ridge (RI) runs along the prismatic (topographically upper) extension of the shell dividing this lengthways into anterior and posterior regions, corresponding, as shown in Figure 2A, to inhalant and exhalant regions, the former twice as wide as the latter.

Ligament

As shown in more detail in Figure 2A (also in figures of *M. malleus* and *M. albus*) this consists of a centrally placed convex inner layer (IL) secreted by the mantle isthmus (MI) with a shorter anterior, and a longer posterior, outer layer (AOL, POL) secreted in the embayments by the outer surface of the outer fold of the mantle margins. Progressive stages in the growth of the ligament are shown in Figure 2 indicating "ventralward" displacement of the hinge line which is characteristic of cemented bivalves [*e.g.*, *Hinnites*, *Spondylus*, *Etheria*, *etc.* (Yonge, 1951, 1962)] and would seem to be due in the case of this byssally attached species to a similar close adherence of the hinge region to a rock surface. In other byssally attached bivalves, *e.g.*, the Mytilidae, the hinge is not pressed against the substrate.

The ligament is bounded anteriorly on the right valve by the deep byssal notch (Fig. 1, BN). Further reference to the ligament is made in the accounts of *M. malleus* and *M. albus*.

Mantle cavity

The appearance of the animal when fully expanded after removal of the left shell valve and the mantle lobe is shown in Figure 2A. The extent of the nacreous layer within which lies the relatively small visceropodal mass is indicated by the broken line (N). The mantle lobes have extended above this carrying the ctenidia (CT) with them to fill the prismatic extension. Any stimulus causes contraction of the pallial retractor (PAR), the presence of which was first noted by Pelseneer (1911), and consequent withdrawal of the mantle lobes, if complete then within the confines of the nacreous region. There is no pallial line in these monomyarians. Of the three marginal folds, the outer one (Fig. 2B, OF), which secretes an extremely thin but very elastic periostracum (P) as well as the prismatic layer (PR), is obscured by the larger middle fold (MF) which bears a single row of small tentacles as does the better developed inner fold (IF) where they are alternately larger and smaller. By apposition or separation these inner folds, or pallial curtains, control entrance into or exit from the mantle cavity.

The lobes are nowhere fused although closely applied to one another at the tips of the ctenidia somewhat to the posterior side of the end of the distal extension (Fig. 2A). This represents the point of division between inhalant and exhalant chambers. The functional advantage of a wide inhalant region, well raised above the bottom and extending along both anterior and distal surfaces, is apparent.

Promyal chamber

In addition to the extensive exhalant chamber distal to the adductor, water also passes posteriorly by way of a promyal chamber (PMC) corresponding to that initially described in *Crassostrea* by Nelson (1938) and now realized as one of the distinctive characters of that genus compared with *Ostrea*. It represents a secondary connection, on the dorsal side of the adductor, between the anterior and posterior ends of the mantle cavity and, as in *Crassostrea*, runs along the right side. After passage through the ctenidia, the flow of water, along the course indicated by the broken arrows in Figure 2, is assisted by cilia lining the chamber and facilitated by displacement of the rectum to the left side as described and figured by Pelseneer (1911) although without knowledge of its significance. Extrusion is then direct (Fig. 2A, E₁) or, if the mantle margins are applied in this region, in more distal regions of the exhalant chamber.

Nelson pointed out that the promyal chamber in *Crassostrea* increased the pumping rate and also permitted life in greater concentrations of sediment. He correlated its presence with greater elongation in *Crassostrea* and consequent greater distance between the hinge line and the adductor compared with the more rounded *Ostrea*. All of these arguments apply equally to *Malleus*, particularly the last although, as noted later, there is no promyal chamber in the effectively equally elongated *Isognomon isognomon*. The chamber in *M. regula* and *M. malleus* is relatively smaller than in *Crassostrea* owing to the presence of the large

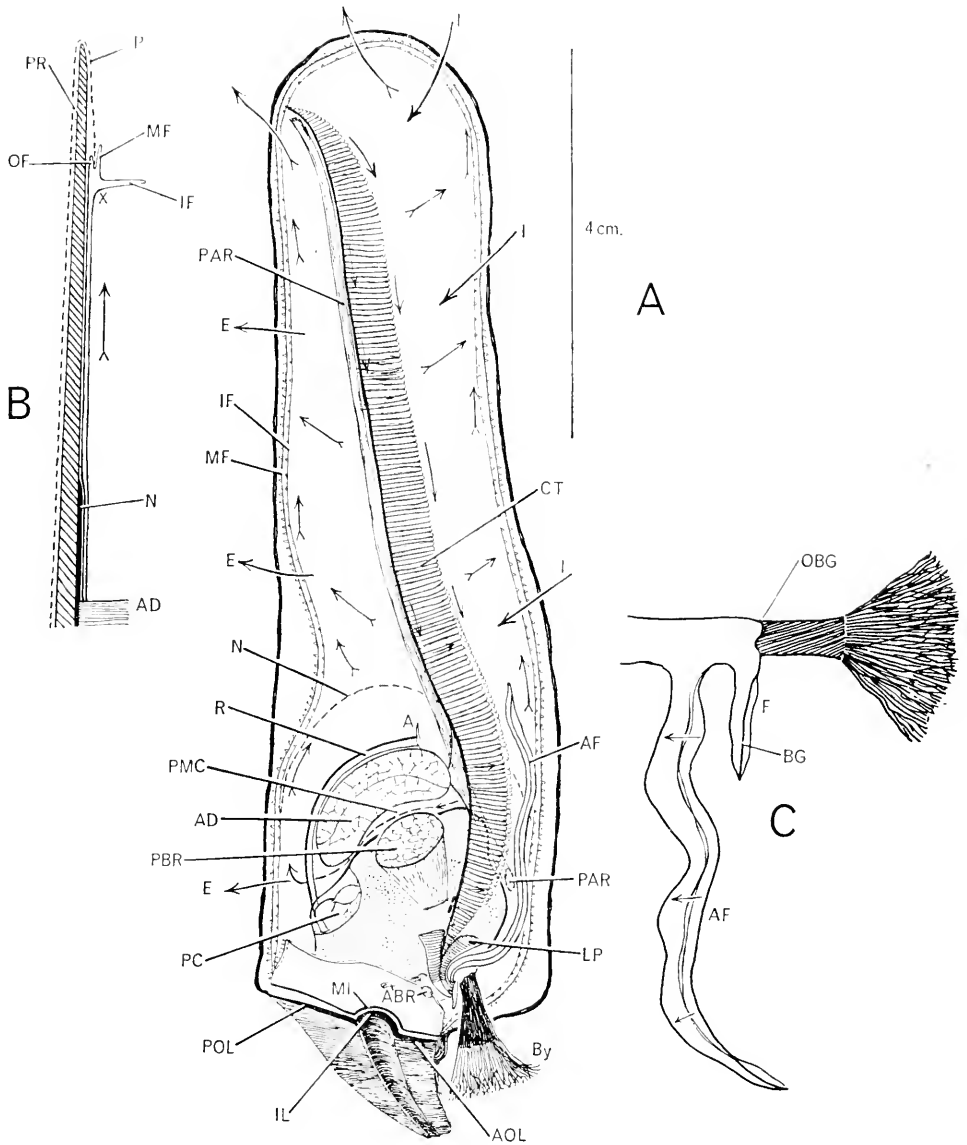


FIGURE 2. *M. regula*. **A**, fully expanded animal within right valve, left mantle lobe removed. Plain arrows indicate direction of collecting currents on the ctenidia, also of inhalant currents; feathered arrows direction of cleansing currents. **B**, Section through valve and mantle distal to adductor showing three marginal mantle folds with periostracum (broken line), outer prismatic layer (oblique lines) and inner nacreous layer (black) of the valve. Waste material carried marginally and then, where marked with x, conveyed distally. **C**, foot, with byssal groove and byssus, and accessory foot viewed laterally. Arrows indicate direction of ciliary beat on extremely active accessory foot.

posterior byssal retractor (PBR) absent in the Ostreidae (and also in adult *M. albus*).

Ctenidia and palps

The greatly extended filibranch ctenidia are attached proximally by way of a suspensory membrane to the anterior surface of the adductor (Fig. 2A) and at the distal extremities to the inner fold of the mantle margins. Attachments between the inner demibranchs and between the outer demibranchs are all by way of interlocking cilia as in other Pteriacea including *Pinna* (Yonge, 1953b). The structure of the gill in *M. malleus* and *M. albus* is described by Ridewood (1903) and Atkins (1936, 1937, 1943) although, despite statements to the contrary, the ctenidium is heterorhabdic with principal filaments. There is fusion between filaments in the region of the "ventral" margins but not terminally. Cilia on the frontal surfaces beat "ventralward" except for those on the inner surface of the outer demibranch which beat towards the axis. There are five oralward currents, three "dorsal" and two "ventral," although there is only a groove along the margin of the inner demibranch. No evidence of antagonistic frontal currents, such as Atkins (1936) found in the related *Pteria*, was noted. There are both longitudinal muscles in the axis and lateral longitudinal muscles on either side as described and figured by Atkins (1943). They must function in co-ordination with the pallial retractor.

The rather small, rather elongate palps (LP) call for no comment. They are symmetrical on the two sides and without fusion of the lips such as occurs in certain Pectinacea. The mouth is relatively large.

Foot and retractors

As described and figured by Pelseneer (1911), the foot in *Malleus* is unique in being divided into two portions. Both project anteriorly and are ventrally grooved; the first is elongated, the more distal bears the byssal opening. The former, designated "bandförmigen Tentakel," was found by Kühnelt (1938) in *M. albus* and considered a possible taste organ. It is here termed the accessory foot (AF) and observations in life have extended knowledge and provided better understanding of possible function.

As shown in Figures 2A and C, the foot (F) is concerned exclusively with the planting of byssus threads which issue from the opening of the byssal gland (OBG) and are directed along the ventral groove. The result is the formation of an elaborate byssus (By), the threads consolidated basally to form a massive stalk. This is the sole function of this region in adult life. It could be concerned with locomotion in early post-larval life as it is in the Mytilidae; it retains that function throughout life in the closely allied *Pinctada* (Herdman, 1904). The bivalve foot may also be a cleansing organ as in certain cemented genera, e.g., *Spondylus* and the Chamidae (Yonge, 1967a) and most strikingly in the Anomiacea where the massive, calcified byssus is planted direct from byssal opening while the elongate foot is concerned exclusively with cleansing the mantle cavity (becoming locomotory again in the limpet-like *Enigmonia*; Yonge, 1957). In the Pinnidae the foot is solely concerned with byssal attachment but a unique pallial organ in the *exhalant*

chamber probably clears that region, especially following the frequent damage these vertically disposed bivalves experience (Yonge, 1953b).

In *Malleus* the function of cleansing has been taken over by the extremely long and active accessory foot which moves with complete freedom throughout the lower regions of the inhalant cavity continually turning and twisting, extending and withdrawing. It is grooved "ventrally" and everywhere ciliated but owing to the constant movement it was impossible to follow the direction of beat except on the outer surface where it is always away from the groove. The function of the cilia appears to be to keep the foot clean, not to direct particles in any particular direction. Transverse sections reveal the presence of a thin layer of circular muscle and within this much greater amounts of longitudinal muscle divided into groups by inward-running strands of fine connective tissue. Blood sinuses are numerous, extension is clearly by blood pressure and contraction of circular muscle, retraction by contraction of the longitudinal muscle.

This is a unique structure which cannot be homologized with any part of the divided foot in the Gastropoda. Its presence is to be correlated with the vertical posture in *Malleus* and the danger of sediment accumulations in the restricted pocket at the base of the inhalant chamber with consequent blockage of palps and mouth (see Fig. 2A). Similar dangers to other similarly disposed anisomyarians, e.g., the Pinnidae and *Pedum spondyloideum* which live, respectively, in sand and in cavities within reef-building corals, are met by way of the unique waste canal in the former (Yonge, 1953b) and of a powerful anterior cleansing current through the byssal notch in the latter (Yonge, 1967b). The small anterior and much larger posterior retractor muscles (ABR, PBR) are symmetrically disposed and effectively entirely concerned with the byssus. Movement of the accessory foot is by intrinsic agencies.

Cleansing

The general course of the cleansing currents in the mantle cavity is indicated by the feathered arrows in Figure 2A. All material in both chambers is carried distally by way of currents running along the inner side of the inner mantle fold (Fig. 2B) and ejected in the distal extremities; there is no accumulation of pseudo-faeces in the inhalant chamber. Cleansing in the base of the inhalant chamber is effected by the accessory foot. Longish faecal pellets discharged from the anus (A) are also conveyed distally for disposal.

Visceropodal mass

Much of this has been described; other anatomical features (shown in Fig. 2A) include the posteriorly disposed pericardium (PC) with the rectum passing through the ventricle and both displaced to the left as described by Pelseneer (1911) in consequence of the development of a promyal chamber. The course of the alimentary canal is simple. The gonad covers most of the visceral mass although all specimens opened—in early February—appeared empty (unlike the associated *Isognomon*), implying recent spawning.

Observations on small individuals

Observations made on animals between 1 and 2 cm. long and up to 6 mm. wide were illuminating. Apart from the byssal notch on the right valve, they are equi-valve, almost completely straight-sided and with the prismatic distal region completely plastic and readily bent in any plane. An instance was noted where the distal region had turned at an angle of 90° in the plane of the valves (it could equally be at right angles to this). The adaptive advantage to individuals which may settle under rocks and can only survive by growing around their margins is obvious. The end result is *always* that the distal regions point upward, opening to receive food and oxygen. This region is initially incapable of closure; contraction of the adductor pulls together the rigid nacreous, but not the distal prismatic, region. As in *Pinna*, this shell layer has a high proportion of conchiolin. However, accompanying withdrawal of the mantle lobes will effectively seal the nacreous region. On later relaxation the mantle lobes could be seen, through the translucent shell, slowly "creeping" back into the distal extension. It is probable that the periostracal sheet, which must be stretched during withdrawal, assists this process. Increased growth brings added danger of damage. However the distal shell is capable of indefinite repair (as in the similarly constructed Pinnidae and most strikingly in *M. malleus*; see Fig. 5A) so that larger shells become increasingly irregular while their full exposure leads to overgrowth by encrusting organisms.

Malleus malleus (Linn.)

This is the black hammer oyster, the longest known and best known species. During the cruise of the *Te Vega*, living specimens were obtained from shallow water near Zamboanga in the Philippines and through the kindness of divers from deeper water in the harbor at Rabaul, New Britain. Subsequently others were examined which had been collected by diving off Lizard Island and off Low Isles by Dr. A. Bastin of the *De Moor* Expedition. In all cases the animals came from relatively clean sandy bottoms in which they were largely buried, only the most distal regions projecting. Specimens measured in Rabaul were 18 to 21 cm. long with a "hinge line" up to 21 cm.; in Australia up to 22 cm. long by as much as 24 cm.

The interior of an adult shell is shown in Figure 3E. The general appearance is similar to that in *M. regula* with a relatively smaller nacreous region bearing the same four muscle scars. There is a well developed byssus but here *both* valves are notched, although to a greater depth on the right. The significant difference, apart from the greater extent of the prismatic region distally, is the formation of similar areas in line with the hinge to produce a very long posterior, and a somewhat shorter anterior, extension (PE, AE). It will be noted (Fig. 3) that the nacreous region is *not* concerned in this. The mantle lobes pass into and are withdrawn away from these extensions exactly as they do in the distal region. While the posterior extension does resemble (on an hypertrophied scale) the posterior auricle in a pectinid, this is not true of the anterior extension which is separated from the hinge and ligament by the byssal notch. There is thus no anterior extension of the ligament but this is not true posteriorly as revealed by the transverse sections—actually of *M. regula* but more suitably discussed in

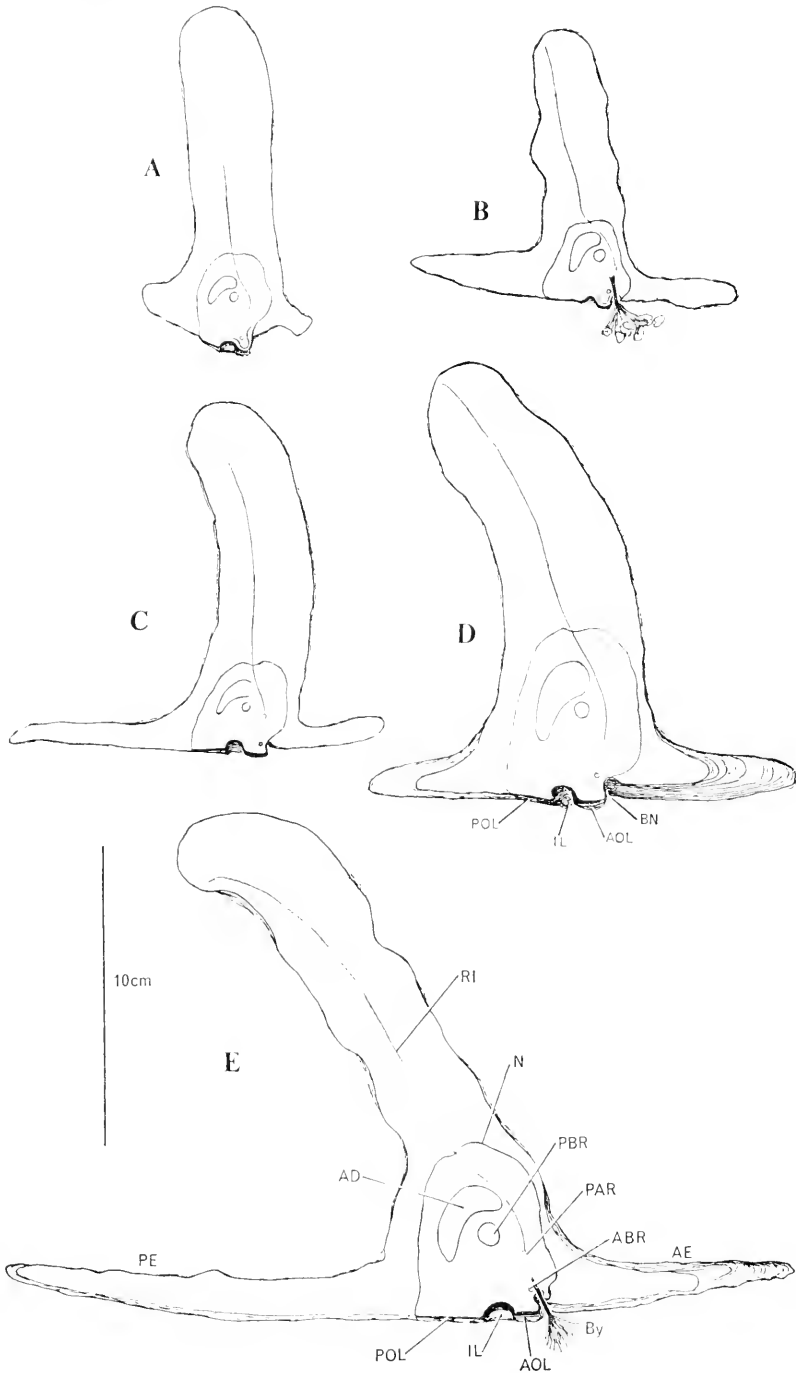


FIGURE 3. *M. malleus*, A-E, interior views of right valves showing various stages in growth with full development of anterior and posterior extensions of the "hinge line" shown in adult (E).

relation to *M. malleus*—shown in Figure 4. These indicate the structure successively from anterior to posterior (*a-g*) of the anterior outer, the inner and the posterior outer ligament layers (AOL, IL, POL) with their respective secreting epithelia (AOM, MI, POM). There is a posterior movement in growth indicated by the intrusion of the anterior outer layer into the inner layer (*b, c*) and a corresponding extension of the latter backward on either side of the posterior outer layer (*f*). Finally, posterior to the primary ligament formed by these three layers (with initially a covering periostracum) there is a secondary extension consisting of fusion layer (*h, FL*) formed by the union of the outer surfaces of the outer fold of the mantle margins, a secreting surface which elsewhere forms the outer calcareous (*i.e.*, prismatic) layer of the valves (OCL). This secondary ligament, short in *M. regula*, is of impressive length in *M. malleus* and *M. albus* (Fig. 6) owing to the great posterior extension of the hinge line in those species. It can have little or no functional significance. It corresponds to the long posterior secondary ligament in the Pinnidae (Yonge, 1953b) which is certainly functionally significant. Both represent the consequence of pallial fusion associated with posterior extension of hinge line.

Apart from the lateral extension of the mantle lobes, internal structure is similar to that of *M. regula*. Interest concentrates on the formation of the lateral extensions of the shell and on habits and life history. As shown in Figure 3, these extensions appear at varying periods in the life history. Thus specimen **A**, although longer than **B**, still has only short extensions. In **B** and **C** the shell has the hammer form and is being vigorously extended in all three directions. In **D**, however, while length increases the lateral pallial extensions have been withdrawn although without effect on the extent of shell earlier secreted. But the mantle could again extend and increase the length of the "hinge line."

The transition from the condition in *M. regula* to that in *M. malleus* clearly occurs during the life history. As discussed later, it permits a changed mode of life. Apart from its vertical disposition, *M. regula* is a typical byssally attached epifaunal bivalve. *M. malleus* is not; it lives in clean but usually coarse sand with the byssus attached to fragments on this (much as does *Pinna*). The anterior and posterior extensions to the hinge line assist in the maintenance of the shell which is always buried with only the distal quarter or less exposed. The extent of exposure is usually revealed by the presence of encrusting organisms absent from buried regions of the shell. The shell extensions provide an anchor within the sand in which this species lives. But the byssus persists throughout life.

The relatively thin shell, the three projecting extremities composed exclusively of prismatic layer with a high proportion of organic conchiolin, inevitably suffers extensive damage. It is rare to find a large shell which does not exhibit extensive areas of repair. An example (from Rabaul) is shown in Figure 5A. The animal has withdrawn from a broken extent of distal shell and formed a new one at a somewhat different angle to the basal extensions, the form of which has also been somewhat altered. Such changes may be in any plane and many adult shells are of grotesquely distorted appearance. The area occupied by the visceropedal mass is seldom, if indeed ever, affected. Conditions are precisely as in the Pinnidae, also byssally attached within soft substrates, similarly exposed to damage and with the same means of extensive and rapid repair of these exclusively pris-

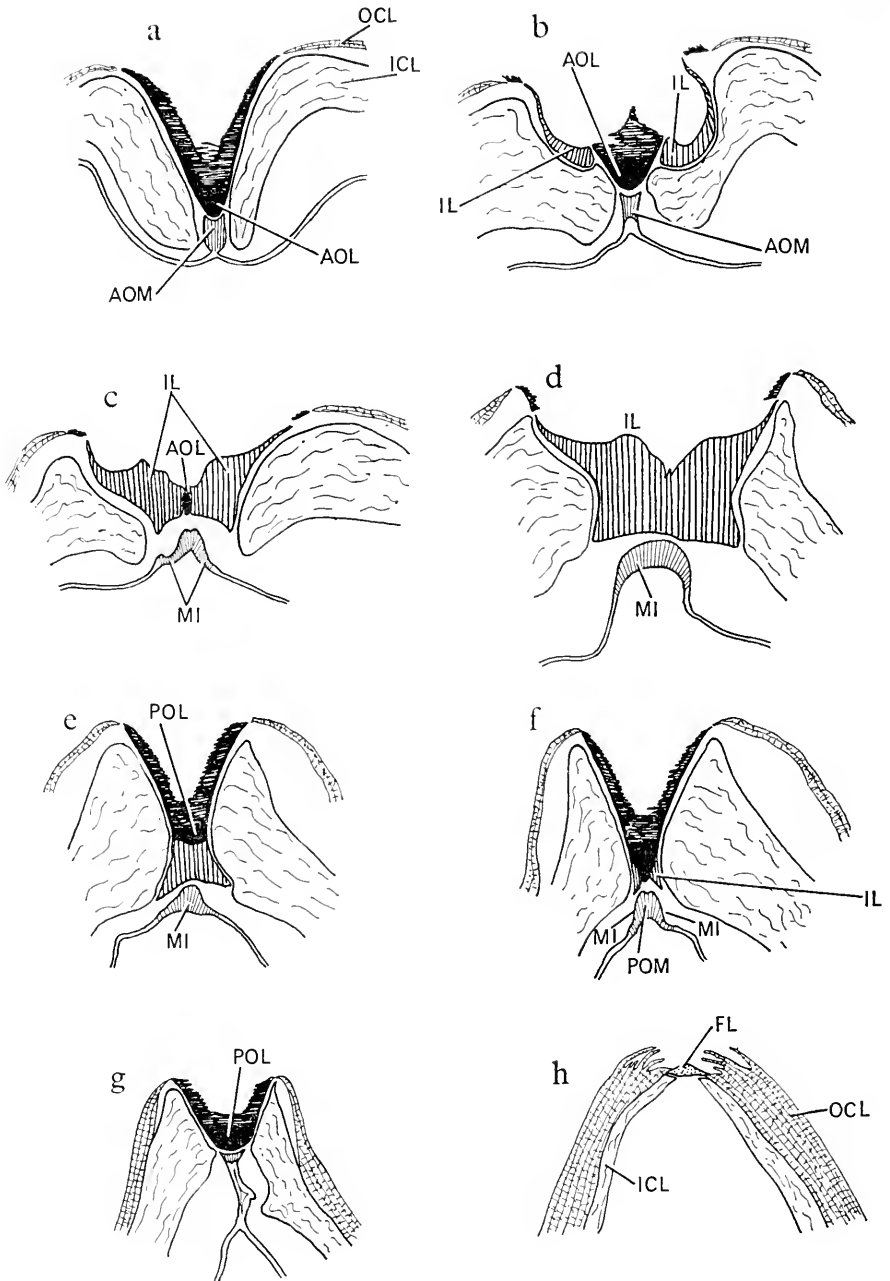


FIGURE 4. *M. regula*. Transverse sections through ligamental region from anterior end back, *a-c* in region of anterior outer ligament; *b-f* in region of inner ligament; *e-g* in region of posterior outer ligament; *h* in region of fusion layer.

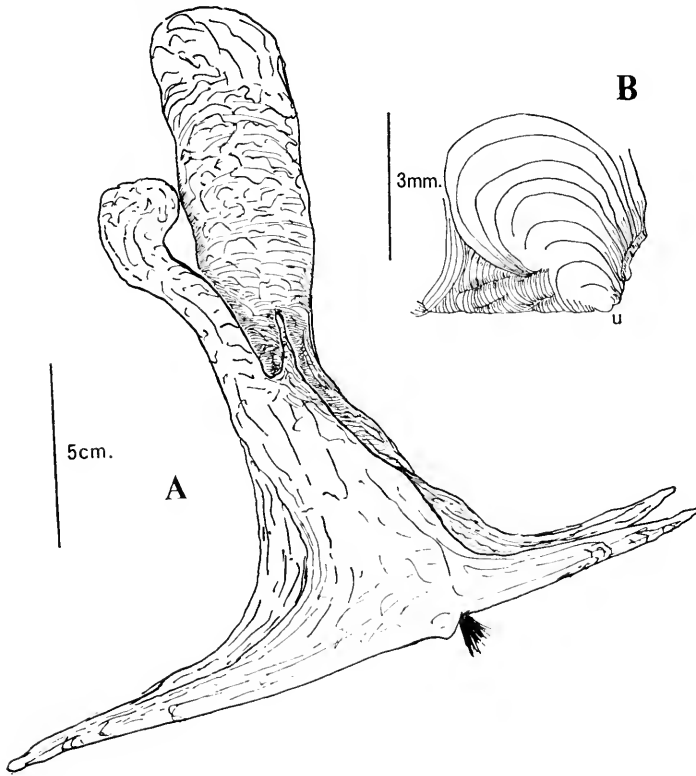


FIGURE 5. *M. malleus*, **A**, very irregular shell (from Rabaul); the mantle lobes have withdrawn from the first formed distal (prismatic) regions of the valves to form new ones at a somewhat different angle to the "hinge line." The nacreous region of the shell is not affected; **B**, umbonal region of left valve showing initial form (resembling *Pinctada*) immediately after settlement (After Jackson, 1890).

matic regions with very high conchiolin content (Bevelander and Benzer, 1948; Bevelander, 1952; Yonge, 1953b).

Malleus albus Lam.

The shell of this second hammer oyster (Fig. 6A) differs in being yellowish white and much stouter; the prismatic layer is more heavily calcified. There is less repair and the shells are therefore less irregular. They attain much the same size; the largest measured, from Bougainville in the Solomons, was 24.5 cm. long with a hinge line of 25 cm. Unlike *M. malleus*, the shell margins are deeply folded, the two valves interlocking very precisely. As originally noted by Lamarck, the adult shell lacks a byssal notch and Kühnelt (1938), in a description of a preserved specimen, found neither byssus nor posterior byssal retractor. As shown in Figure 6A, the corresponding muscle scar has disappeared, covered over by subsequent secretion of nacreous material.

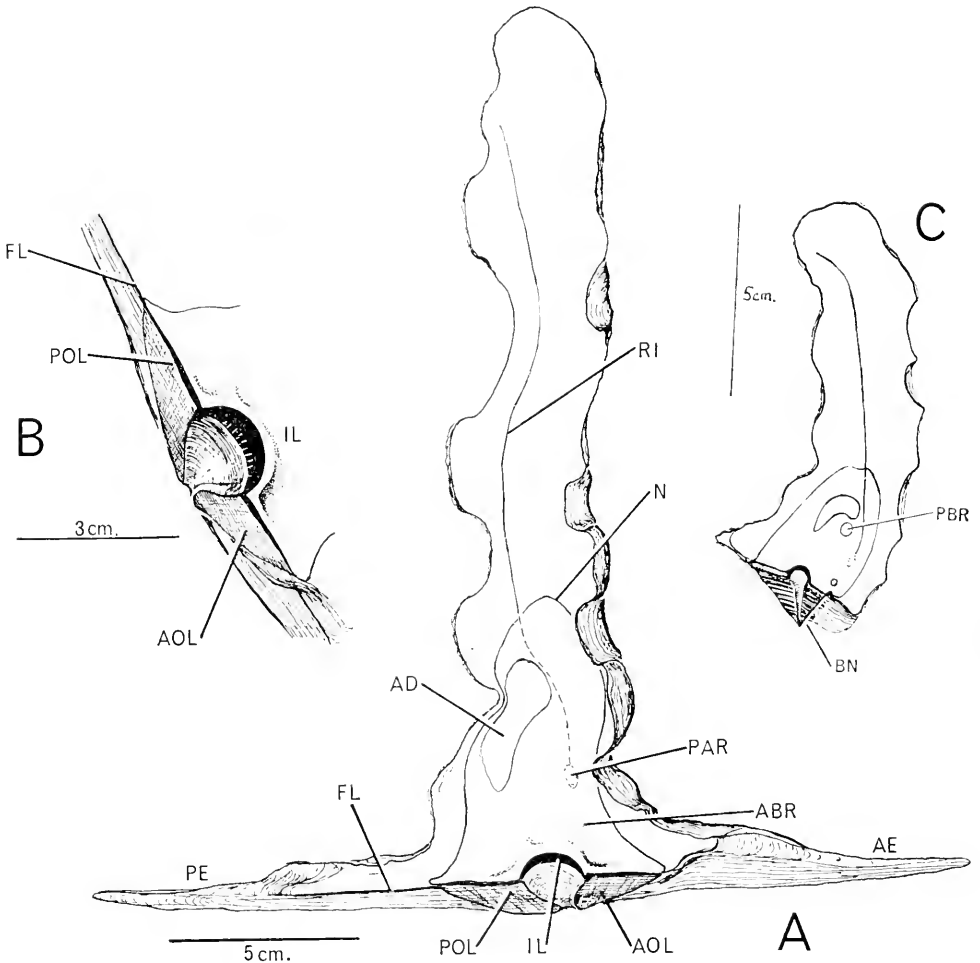


FIGURE 6. *M. albus*. **A**, interior of right valve of adult shell; **B**, ligamental region further enlarged; **C**, interior of right valve of half-grown specimen with byssal notch still present.

In a younger shell, at a stage before the hinge line has lengthened (Fig. 6C), there is both byssal notch (BN) and muscle scar (PBR). Unfortunately this shell, which was personally collected at Zamboanga in the Philippines, was empty. A living adult specimen was later collected by Professor T. F. Goreau by diving under the *De Moor* in the lee of Lizard Island within the Great Barrier Reef. The shell was 16.5 cm. long by some 12 cm. along the hinge line. There was no byssal notch, no byssus and no posterior byssal retractor although the small anterior muscle was retained. The true foot was reduced to a small protuberance with a minute opening of the former byssal gland and was situated at the base of the very large and extremely mobile accessory foot.

In *M. albus*, therefore, the animal comes to be anchored exclusively by way of the anterior and posterior extensions of the shell. Moreover it differs from *M. malleus* in living always in a substrate of muddy sand. Initial information to this effect was obtained from shell collectors at Rabaul; this was later confirmed by Mr. S. Bernik of Cairns, N.Q., who had collected specimens from muddy sand at depths of about 30 feet off Bougainville Island. The specimen from Lizard Island came from a similar substrate; it was lying somewhat obliquely with the distal third covered with sponge and other encrusting growths. Specimens of *M. malleus* from that area all came from clean sand.

LIFE HISTORY IN THE GENUS MALLEUS

In the absence of precise knowledge any discussion about life histories must be on a basis of reasonable supposition. The problem is least in *M. regula*; there the larvae must settle on a rocky surface where mud is present. Unless impeded by overgrowths of rock the shell will then grow vertically. In the two other species metamorphosis of the settling veligers must be stimulated by contact respectively with a clean sandy or sandy gravel substrate or with one of muddy sand. Despite the opinion of Kühnelt, who considers the shell extensions as concerned with stability after attachment to rock, both *M. malleus* and *M. albus* are infaunal. Settlement of the larvae of *M. regula* on mud-covered rock could well represent the ancestral habit.

Initial byssal attachment is presumably to small fragments within the substrate. Gradual descent into this as growth proceeds must initially—as in the Pinnidae—be due to progressively deeper extensions of the foot with planting of new byssus threads on to fragments further down. The progressive extension of the hinge line will certainly anchor the animals more securely but at the same time must present an obstacle (absent in the anteriorly pointed Pinnidae) to the further penetration which certainly occurs. The eventual closure of the byssal notch in *M. albus* will even prevent any downward ejection of water which could assist further penetration in *M. malleus*. However a large shell of *M. albus* could well settle into the softer substrate it inhabits by its own weight. According to Mr. Bernik shells of this species are usually obliquely disposed with two regions of the shell protruding, presumably the posterior as well as the distal extremity. Professor R. D. Purchon (personal communication) states that, near Singapore, he collected living specimens lying on the surface of muddy sand. They could have been thrown up by storms. This does not appear to happen in the attached *M. malleus*. In both species not more than a third (often less) of the distal region projects. Differences in the two infaunal species are certainly associated with their distinct habitats. Water movements will be greater in regions where clean sand or shell sand accumulates. The combined effect of byssal attachment and shell anchorage would seem to be needed; the distorted form of larger shells indicates major disturbances and the means of coping with these. Water movements are obviously less where muddy sediments accumulate—so the byssus can be discarded while a thicker shell will oppose greater resistance to undue settlement of the massive adult shell into this softer substrate.

THE GENUS *ISOGNOMON* SOLANDER 1786

This genus, with *Malleus* included by Thiele (1935) in the Family Vulsellidae of the Pteriacea, differs from that genus in the possession of a multivincular type of ligament (Figs. 7, 8, ML). This has been described by Trueman (1954) for *Pedalion* (*Perna*) *alata*. (These generic names are synonyms for *Isognomon*.) Instead of the usual single inner ligament with bounding and superficial anterior and posterior outer ligament layers, the ligament is here subdivided into a series of inner ligament layers (or resilifers) with intervening outer ligament layers as shown in Figures 7-9. This type of ligament is shared with only the allied genus *Crenatula* and might be considered, as Trueman points out, to indicate wider than merely generic separation from *Malleus*. However the anatomical studies of Pelseneer (1911), which have been confirmed, do indicate close affinity between *Isognomon* and *Malleus* while Bernard (1898) describes the presence of a minute resilifer in association with the umbones and anterior to the large inner ligament in *Malleus*. By removing the major dissimilarity, this could indicate considerable affinity between *Malleus* and *Isognomon*.

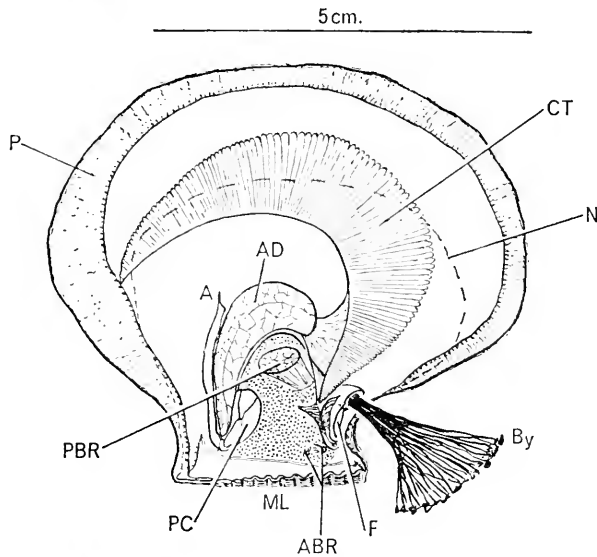


FIGURE 7. *Isognomon ehippium*, general view of animal lying within right valve, left valve and mantle lobe removed. Withdrawal of periostracal sheet formed by outer fold of mantle margin indicated (P).

Isognomon ehippium (Linn.)

A brief account is given of this species (Fig. 7), specimens of which were collected at Kieta on Bougainville Island. The rounded "oyster-like" form is characteristic of the majority of species of this genus which is widely distributed in the tropical Indo-Pacific and Atlantic (where it extends to Bermuda). This description forms a necessary preliminary to the more detailed account of *I. isognomon* where form and habit so closely parallel those in *M. regula*.

This and other rounded species are characteristically attached to rocky, often inter-tidal, surfaces by a massive byssus. This projects through a notch on the right valve on which side the animal lies precisely as do species of the pearl oysters, *Pinctada* and *Pteria* (Pteriidae). The animal has the typical monomyarian form with the viscero-pedal mass reorganized in relation to the centrally placed, although somewhat dorso-ventrally extended, adductor (AD) around which curl the ctenidia (CT). Although the nacreous region (delimited by the broken line N) is relatively larger than in *Malleus*, it is surrounded by a broad marginal prismatic region. The mantle margins retract across this retaining attachment to the edge of the shell by a delicate periostracal sheet (P). The small foot (F) is concerned, probably exclusively, with planting of the massive byssus (By). There are large posterior and small anterior pedal retractors (PBR, ABR) the latter with divided insertions. As shown by Pelseneer (1911) there is no asymmetry in the heart region, *i.e.*, no promyal chamber. As in all such monomyarians there is an extensive inhalant, and a correspondingly restricted exhalant, region. Ciliary currents were not particularly observed in this species; they doubtless correspond to those figured for *I. costellatus* by Stasek (1963) showing rejection of waste particles from both inhalant and exhalant chambers at a common point just on the exhalant side of the tip of the ctenidia.

Isognomon isognomon (Linn.)

This common Indo-Pacific species differs from *I. ephippium* and similar species in being elongated, attached in an erect position and, as noted already by Kühnelt (1938), assuming both the external appearance and mode of life of *M. regula* with which it is found. It is somewhat the larger, attaining lengths of 15 cm. and being somewhat broader. Appearance in life after removal of the left valve and mantle lobe is shown in Figure 8 which is directly comparable with that of *M. regula* (Fig. 2A.) There is obvious difference in the structure of the ligament which in *Isognomon* extends along the entire dorsal surface of the shell and in the position of the byssal notch, here anterior instead of dorsal. Consequently the massive byssus, the threads of which are not basally fused, emerges in that direction although applied to an under surface. The nacreous area is relatively much larger and (*cf.* Figs. 3 and 9) its margins less definite, the mantle lobes with the ctenidia not retracting so far. There is no pallial retractor and no corresponding ridge within the distal region of the shell and no promyal chamber. Without the accessory appendage, the foot is primarily concerned with byssal attachment but still large enough to have possible cleansing functions within the anterior extremity of the mantle cavity.

Distal to the nacreous region the mantle lobes and ctenidia are increasingly pigmented and finally dense black. The mantle margins are as in *Malleus* but thicker and relatively larger with short tentacles on the inner and middle folds. The non-plicate ctenidia are less firmly united and the filaments separate more readily (*i.e.*, are more typically filibranch) than in *M. regula*. As in that species there is no "ventral" groove along the outer demibranch and ciliation is similar. The tips of the ctenidia are attached to the inner mantle folds in the same position as in *Malleus*, the mantle cavity being similarly divided; the exhalant chamber is correspondingly greater than in the rounded *I. ephippium* (*cf.* Figs. 7 and 8).

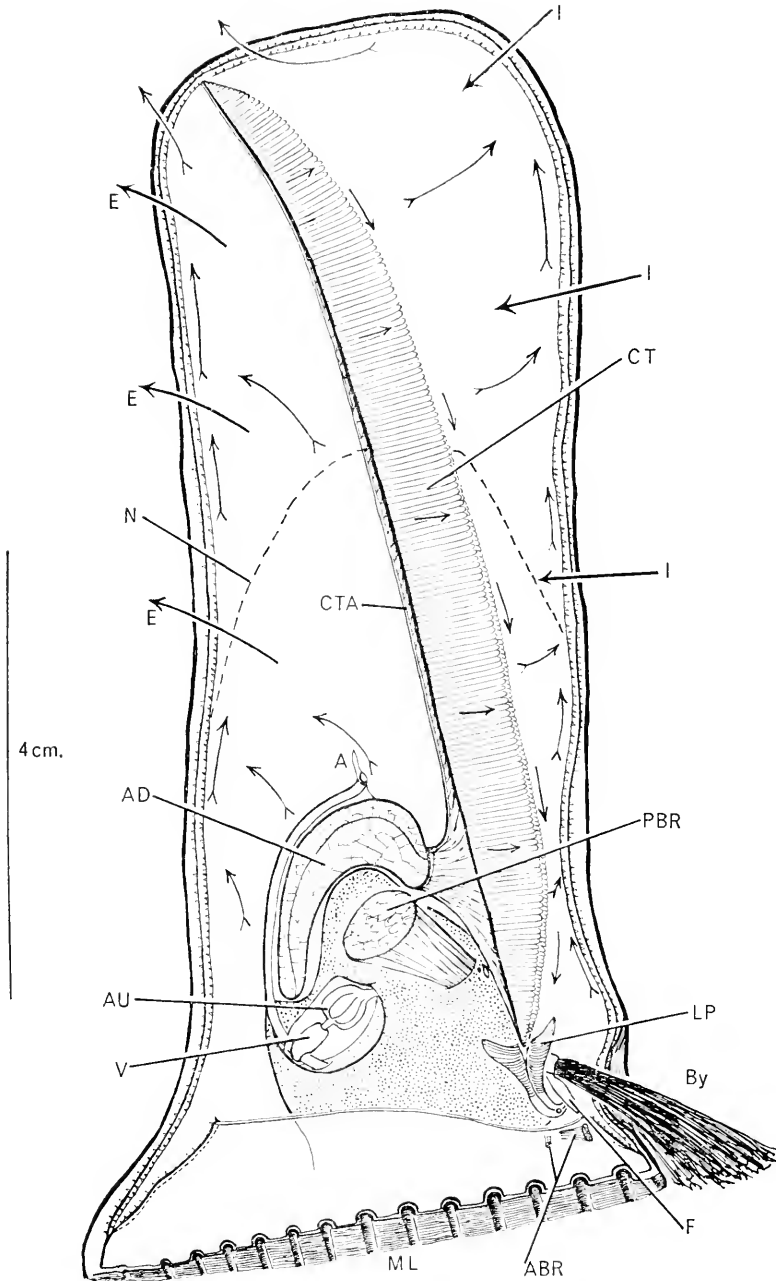


FIGURE 8. *I. isognomon*, fully expanded animal within right valve, left mantle lobe removed. Plain arrows indicate direction of collecting currents on the ctenidia, also of inhalant and exhalant currents; feathered arrows direction of cleansing currents.

The palps are relatively longer than in *M. regula*; the lips are similarly developed, without frilling or fusion. There is a long anal flap which assists in directing the faeces, long scroll-like pellets up to 5 mm. long. As in *Malleus*, all waste material in both inhalant and exhalant chambers is carried to the inner edge of the inner mantle fold and so to the posterior distal regions where it is ejected (see feathered arrows in Figure 8).

Unlike *M. regula*, the gonad, which covers the visceral mass, was obviously maturing although sperm was not ripe.

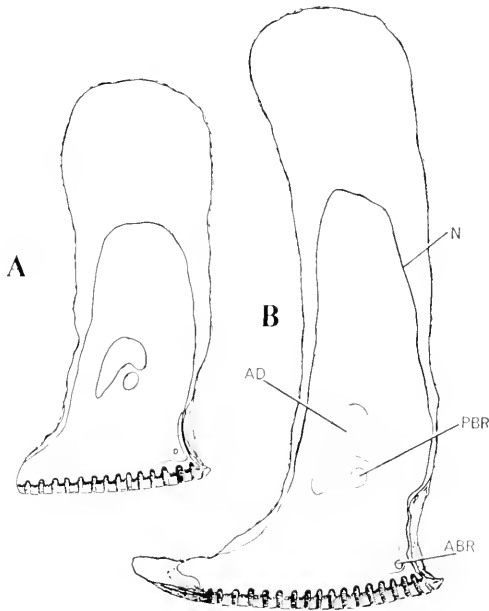


FIGURE 9. *I. isognomon*. A, B, interior views of right valves of two specimens showing stages in extension of the ligament and hinge line and of the distal region. The shells are 12 and 17.5 cm. long.

In the smallest animals collected the shell was 13 mm. long by 8 mm. wide and was oval rather than elongated like similar-sized *M. regula*. The shell is usually bluish with light-colored radiating curved rays. Young shells in dark areas tended to remain white. In some the foot was observed extended and seeking a surface for byssal attachment; when found initial threads were quickly secured. Until some 30 mm. long the shells remained oval and then displayed increasing irregularities due to the confined space in which they grew and to the effects of repair. But there is increasing elongation as form changes from the initially more rounded condition. They come to assume a striking external resemblance to *M. regula* although the hinge line is extended posteriorly to a greater extent than in that species (*cf.* Figs. 1 and 9). This process of extension continues until, as shown in Figure 9B, it may attain a length double the width of the shell. At this size the mantle tissues tend to withdraw, the ligament no longer elongates and the shell only increases in thickness.

Multivincular ligament

Two stages in the growth of the ligament are shown in Figure 11A, B. It begins as a simple opisthodontic ligament (**A**), the inner ligament layer inclined posteriorly from the umbones as in *Malleus*. But instead of this increasing steadily in size by added secretion from the mantle isthmus as in that genus (see Fig. 2A), new areas of inner ligament, or resilifers, appear at intervals posteriorly. Two of these are shown in Figure 11B; the maximum number observed in *I. isognomon* was 18. While obviously the number is related to the length of the hinge line, it is not necessarily related to the length of the shell.

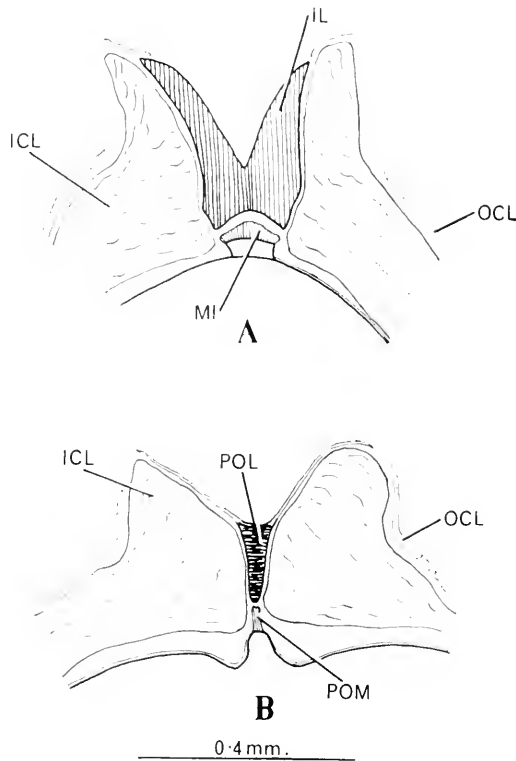


FIGURE 10. *I. isognomon*, transverse sections through ligament, **A**, through a region of inner ligament (a resilifer); **B**, region of outer ligament between resilifers.

In older individuals the original resilifer may be partially worn away or lost. Trueman (1954) describes a fusion layer but no sign of this appears in sections nor, in view of the absence of any extension of the mantle behind the primary ligament, as occurs in *Malleus* and *Pinna*, would its presence seem possible. The unsolved problem in this type of ligament is the origin of the successive areas of inner ligament, the one first formed being secreted by the mantle. Although there is as yet no certain supporting evidence, it appears most probable, as indicated diagrammatically in Figure 11C, that successive regions of the inner mantle surface

(which normally secretes the inner calcareous, *i.e.*, nacreous, layer of the valves) fuse in the mid-line to invade the epithelia secreting the outer ligament layer and produce successive mantle isthmuses which secrete resilifer 2, then 3 and so on posteriorly. The inner mantle epithelium is here assumed to have the potential of secreting inner ligament should it, like the original mantle isthmus, become situated in the mid-dorsal line, *i.e.*, when subjected to the opening and closing movements of the valves. As in *Malleus* there is a marked ventralward movement of the hinge line as shown in the transverse sections (Fig. 10).

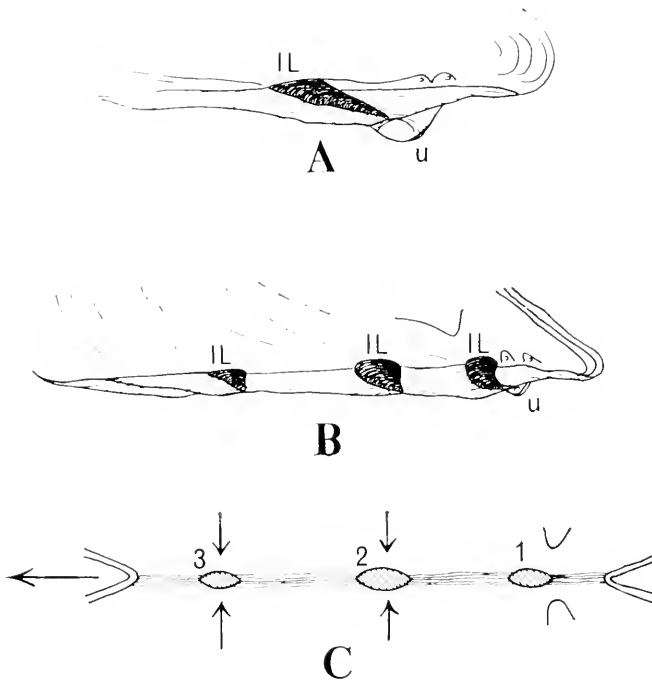


FIGURE 11. *I. chippium*, views of ligamental regions of young shells (right valves), **A**, with only the original region of inner ligament; **B**, with two others formed subsequently (both from Jackson, 1890); **C**, diagram indicating possible mode of formation of multivincular ligament, 1-3 successively formed resilifers (1 in association with umbones), arrows indicate regions of possible cross fusion of inner mantle epithelium, horizontal arrow direction of ligamental growth (original).

MIXED NATURAL BED OF *M. REGULA* AND *I. ISOGNOMON*

A remarkable mixed population of *M. regula* and *I. isognomon* was encountered in the narrow and very shallow channel between Palau Gaya and the much smaller Palau Tatagan. The general appearance of this "clam bed" at low spring tide is shown in Figure 12A with a closer view in Figure 12B.

The shallow rocky channel was everywhere covered with the densest possible growth of mixed *M. regula* and *I. isognomon*. The upright shells formed a compact "pavement" over an area of some 65 by 40 meters kept free by water move-



FIGURE 12. **A**, passage between Palau Tatagan and Palau Gaya photographed from south at low spring tide and showing exposed bed of mixed *Malleus regula* and *I. isognomon*; **B**, closer view of an area of the bed largely exposed and showing density of the mixed population.

ments from mud or sand. The maximum population counted within a foot square patch was 97, the largest *M. regula* being 12 cm. and *I. isognomon* 15 cm. long. The shells were usually irregular, due to dense crowding and frequent repair, but all pointed upward. Although the rock surface was irregular, all was smoothed out by the zonal growth of these bivalves above which occasional rocky patches projected.

As the tide exposed them sudden contractions of the "quick" muscle of the adductors produced innumerable jets of water (Fig. 12B). These ejections could be cleansing or be related to stages in mantle withdrawal. A final tightening up due to contraction of the "catch" muscle and the shells remained firmly closed until again covered with water.

Examination of samples indicated a greater proportion of *Malleus* in higher, and of *Isognomon* in lower, levels. In areas first uncovered proportions were almost 4/1, in mid and lower levels they were almost 1/1 but in the deepest level sampled, they were 1/3. Three factors could account for these slight differences in horizontal distribution; (1) tolerance to exposure (*i.e.*, effects of temperature and desiccation); (2) tolerance to presence of mud; (3) level at which the larvae settle. However the striking fact is not these minor differences but the complete mingling of two distinct species in the one habitat.

The bed constituted an interesting ecosystem. Within the shelter provided by the upright shells was a varied community of attached and free-living animals such as mussels, small cowries, crabs and weed with numerous ophiuroids moving actively between the bases of the shells. Occasional *Pinna* sp. were found in this unusual environment. Attached to the occasional patches of exposed rock was a distinct population of oysters, patelloid limpets, *Chama* sp., etc.

EVOLUTION WITHIN THE PTERIACEA

This group is more open to criticism than are the other four superfamilies—Mytilacea, Pectinacea, Anomiacea and Ostreacea—into which Thiele (1935) divided the Order Anisomyaria. Of the three constituent families, two genera of the Vulsellidae possess a multivincular ligament (but see Bernard, 1898), and so might be included in a separate family, while the heteromyarian Pinnidae have many unique features (Yonge, 1953b) and could be separated as a distinct Superfamily. However, accepting the present position, in Table I are listed certain crucial characters of the constituent genera with particular reference to habitat.

The Pteriacea occupy three distinct habitats: (1) a certainly primitive one byssally attached to rock or other hard surfaces, and secondary ones (2) without byssus within sponges (and also algae) and (3) with byssus in soft substrates although with eventual loss of this in *M. albus*. There is thus a tendency, absent in all other Anisomyaria, for return to infaunal life (the exception would be rock-boring Mytilacea, *e.g.*, *Botula* and *Lithophaga*). It could be noted in possible defense of the present classification, that both the alivincular *Vulsella* and the multivincular *Crenatula* live in sponges, and both *M. malleus* with *M. albus* and the Pinnidae occupy soft substrates.

These points are illustrated in Figure 13. From the primitive habit of byssal attachment with the animal lying on the right side found in the Pteriidae (*e.g.*, *Pinctada*) and also in most species of *Isognomon* (A) there is change to the

vertical posture found in *I. isognomon* (B) and *M. regula* (C). From the latter condition comes invasion of soft substrates with accompanying elongation of the hinge line in *M. malleus* (D) and *M. albus* (E), the second losing the byssus in adult life. From both (A) and (B) there is independent change to life within sponges (F, G) with loss of the byssus. Finally, and independently, there is the infaunal habit of the Pinnidae (H), heteromyarians with an exceptionally large byssus.

TABLE I
Pteriacea

Family	Genus	Ligament		Byssus	Habitat		
		Alivincular	Multivincular		Epifaunal to rock, etc.	Infaunal sponges	Infaunal soft substrate
Pteriidae (Mono-myarian)	<i>Pteria</i>	x	—	x	x	—	—
	<i>Pinctada</i>	x	—	x	x	—	—
Vulsellidae (Mono-myarian)	<i>Isognomon</i>	—	x	x	x	—	—
	<i>Crenatula</i>	—	x	—	—	x	—
	<i>Foramelinia</i>	x	—	x	x	—	—
	<i>Vulsella</i>	x	—	—	—	x	—
	<i>Malleus</i>	x	—	x	x	—	—
	<i>regula</i>	x	—	x	—	—	x
	<i>M. malleus</i>	x	—	x	—	—	(coarse sand)
(Pinnidae) (Hetero-myarian)	<i>M. albus</i> (adult)	x	—	—	—	—	x (muddy sand)
	<i>Pinna</i>	x	—	x	—	—	x
	<i>Atrina</i>	x	—	x	—	—	x

Elongation in Malleus and I. isognomon

The typical monomyarian (Yonge, 1953a) is rounded with the exception of occasional species or varieties in the cemented Ostreacea and Etheriidae (Unionacea), e.g., respectively, *Crassostrea gigas* and *Etheria elliptica* var. *cailliaudi* (Yonge, 1962), although in both, and especially in the latter, it is the ligamental region particularly which elongates dorso-ventrally. Among those byssally attached, elongation is confined to *Malleus* spp. and to *I. isognomon*.

The primitive condition in the Pteriacea (omitting the Pinnidae) is probably best represented by *Pinctada* and by rounded species of *Isognomon*. Unlike the Pectinacea, the visceral mass is proximal to the adductor into a dorsal embayment in which its fits (Figs. 2A, 8). Distal extension is thus readily attained by change

in growth gradients involving straightening and extension of the ctenidia. The difference between *I. ephippium* and *I. isognomon* (cf. Figs. 7 and 8) is brought about in this way and is associated with change to a vertical posture and a modified mode of life.

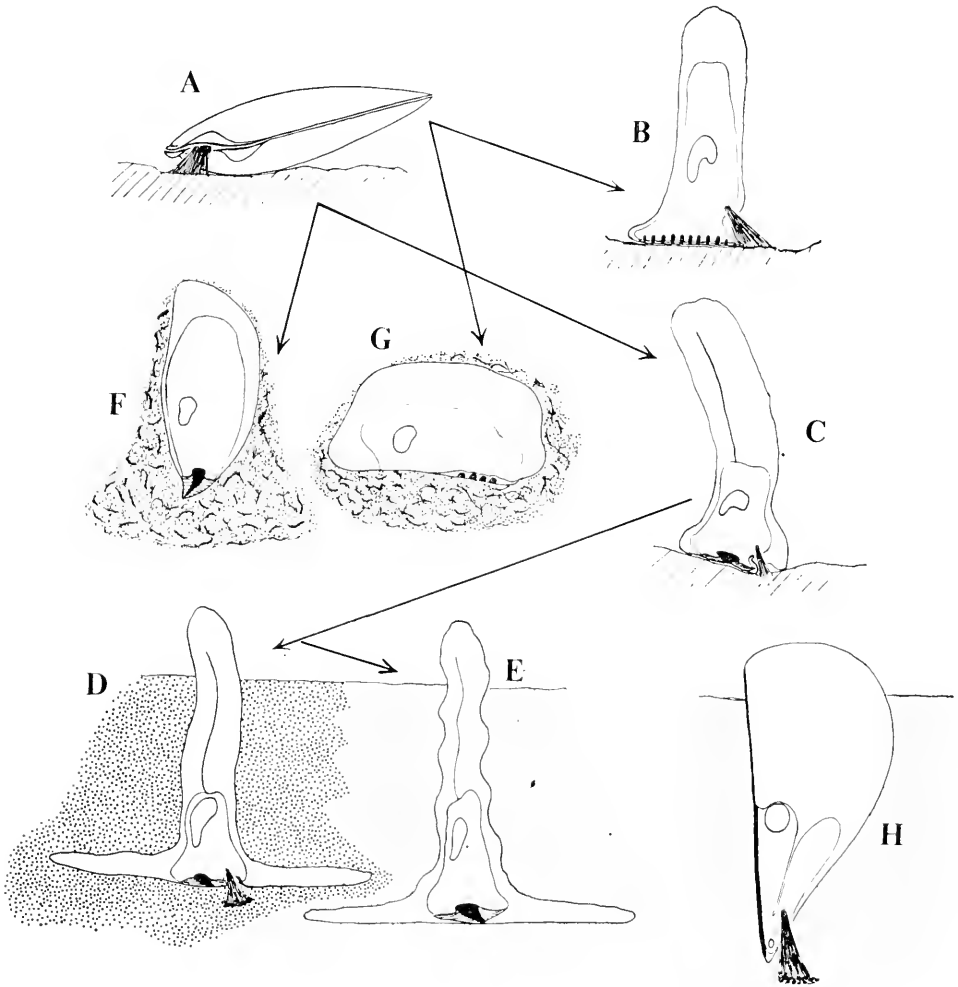


FIGURE 13. Pterinea: drawings illustrating major adaptations and habitats of representative genera and species. **A**, *Pinctada* or *Isognomon ephippium* (alivincular and multivincular ligaments, respectively), rounded lying on right valve, byssally attached to hard substrate; **B**, *I. isognomon*, elongated, vertically disposed, byssally attached; **C**, *M. regula*, as in **B**; **D**, *M. malleus*, elongated with long hinge line, byssally attached within coarse sand; **E**, *M. albus*, as in **D** but adult not attached and in muddy sand; **F**, **G**, *Vulsella* (alivincular ligament) and *Crenatula* (multivincular ligament) both within sponges; **H**, *Pinna*, elongated, byssally attached in soft substrates.

Elongation is characteristic of the genus *Malleus*. Although, as shown in Figure 5B, the first formed shell is rounded, this is quickly altered by subsequent extensions of the mantle. The major changes involved consist of:

1. Great distal extension of the marginal prismatic zone of the shell.
2. Appearance of a pallial retractor closely associated with the elongated ctenidium.
3. Movement of byssal notch to the hinge line so that byssus emerges "downward."
4. Appearance of a promyal chamber proximal to the adductor (as in *Crassostrea*) made possible by elongation and permitting more efficient water circulation in the proximal region of the mantle cavity.
5. Development of a unique accessory foot concerned with cleansing of this region of the elongated mantle cavity.
6. Loss of any previous asymmetry associated with the recumbent posture, *i.e.*, of valves, byssal notch (gradually) and pedal musculature.

Other changes, notably elongation of the hinge line and eventual loss of the byssus, are associated with change from epifaunal to infaunal life.

Comparison between *M. regula* and *I. isognomon* has already been made (*cf.* Figs. 2 and 8). There is remarkable convergence both in form and consequent habit leading to the complete mingling of populations observed in the channel between Palau Gaya and Palau Tatagan. They are sympatric species. While the certainly more highly specialized *M. regula* might be expected to be the more successful this is certainly not the case at lower levels of the bed, over all levels of which *I. isognomon* occurs. The vertical distances are, of course, very small.

This mingling of populations of two species apparently contradicts the "competitive exclusion principle" of Gause (1934) which postulates that closely related (in this case similarly adapted) species with the same ecology cannot live together. Mayr (1963) has pointed out the exceptions, namely where two species occupy different niches in the same general habitat and the case of generalized herbivores where numbers are controlled by predators or disease and *not* by limitation of food. This could certainly apply to *M. regula* and *I. isognomon*.

The two species feed in identical ways and in shallow water where phytoplankton should be rich. Although *M. regula* is certainly the more highly adapted for dealing with collection of sediment within the vertically disposed mantle cavity this does not appear to give the species any obvious advantage over *Isognomon*. (It may well be a factor in permitting related species to exploit infaunal life.) As already noted, possible differences in settlement behavior and in resistance to limited degrees of exposure (but to *very* high temperatures) could account for the greater abundance of *M. regula* at higher, and of *I. isognomon* at lower, levels. But the major factor permitting co-existence of the two species would seem to be their different spawning seasons for which adequate evidence was obtained. In February *M. regula* had recently spawned but *I. isognomon* was only preparing to do so. There could be a gap of a month, or months, between the two events. There would therefore be no competition during the most crucial stages in the life history, when the larvae are planktonic and, above all, when they are settling with the accompanying, usually very rapid, metamorphosis into the adult form with assumption of the adult habit.

It is a pleasure to express gratitude to Stanford University for the invitation to participate in Cruise 6 of the *Te Vega*, itself made possible by N.S.F. Grant G17465. Particular thanks are due to Dr. Lawrence Blinks, then Director of the Hopkins Marine Station, Pacific Grove, who was in scientific charge of that cruise, and to Dr. Dan Cohen and to all others on board. Observations were extended during September to October, 1967, during the Belgian *De Moor* Expedition to the Great Barrier Reef of Australia thanks to the invitation of Professor M. Du-buisson, Rector of the University of Liège, and the kind co-operation of the Leader of the Expedition, Professor A. Distèche and his colleagues and also of Professor T. F. Goreau, a co-worker and fellow visitor. All microscopic sections have been prepared and all figures produced by the author's research assistant, Miss J. I. Campbell, whose help has been indispensable. Dr. Robert Robertson kindly read the proofs.

SUMMARY

1. Study of the structure and mode of life of *Malleus regula* provides the basis for consideration of the "hammer" species, *M. malleus* and *M. albus*.
2. *M. regula* occurs byssally attached, vertically disposed on rocky substrates associated with mud and is widely distributed in the tropical Indo-Pacific.
3. The distal two-thirds of the elongated shell is exclusively prismatic. By means of special pallial retractors the mantle lobes can be withdrawn within the nacreous region.
4. The massive opisthodetic ligament has a short secondary extension of fusion layer.
5. A promyal chamber on the right side proximal to the adductor increases water flow into the exhalant chamber.
6. The long filibranch ctenidia provide a vertically extended food-collecting surface.
7. The foot is concerned with planting of the massive byssus which emerges through a notch in the right valve but on the under (*i.e.*, dorsal) surface. There is also a unique and very long accessory foot, ventrally grooved, everywhere ciliated and in constant writhing activity due to blood pressure and intrinsic muscle. Moving freely throughout the lower mantle cavity it can only be concerned with cleansing.
8. Pseudofaeces are ejected from the distal tip of the mantle cavity.
9. *M. malleus*, the black hammer shell, occurs vertically embedded in coarse sand or sandy gravel. Byssus threads are attached to fragments within the substrate. The great anterior and posterior elongations of the hinge line (also exclusively prismatic) are separated by the byssal notch (now affecting both valves).
10. Shells are usually excessively irregular due to the great exposure to damage and the almost unlimited powers of rapid repair by the three pallial extensions.
11. *M. albus*, the white hammer shell, is stouter and inhabits muddy sand. During growth it loses the byssus with reduction of the foot and byssal retractors and closure of the byssal notch. The accessory foot is not affected. The animal becomes anchored in the substrate exclusively by the anterior and posterior extensions of the shell.

12. Description of the typically rounded *Isognomon ephippium* leads to that of the elongated *I. isognomon* which occupies precisely the same habitat as *M. regula*.

13. It lacks the pallial retractors, promyal chamber and accessory foot of *Malleus*, the nacreous region is more extended distally and the ligament is multi-vincular (the formation of which is discussed), producing some posterior, but never any anterior, extension of the hinge line.

14. A remarkably dense mixed bed of *M. regula* and *I. isognomon* in Darvel Bay, Sabah (Borneo), is described. Different spawning periods, by preventing competition during settlement, may account for this complete intermixing of sympatric species, which, however, are generalized herbivores where numbers are not limited by food supply.

15. From a basic epifaunal habit (*Pteria*, *Isognomon*, *M. regula*), members of the Pteriacea have become adapted for infaunal life within sponges (*Vulsella*, *Crenatula*) or within soft substrates (*M. malleus*, *M. alba* and the Pinnidae).

16. There is final discussion about elongation in monomyarians, *i.e.*, in the genus *Malleus* and in *I. isognomon*.

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ABSTRACTS OF PAPERS PRESENTED AT THE
MARINE BIOLOGICAL LABORATORY
1968

ABSTRACTS OF SEMINAR PAPERS

JULY 16, 1968

Sulfonation of rat lens proteins. SEYMOUR ZIGMAN.

When rat lens proteins were sulfonated using Bailey's reagent (Na_2SO_3 , $\text{Na}_2\text{S}_2\text{O}_8$, 8 M urea, pH 9.9, HCO_3^- buffer), and then were subjected to preparative high voltage electrophoresis on Pevikon (polyvinyl chloride) plates (1.2 mm. thick) at pH 9.9, two components were obtained for both gamma crystallin and albuminoid. The mobilities of the faster-moving components (I) of both were 16 cm./hr. at 3000 v. as compared to 11 cm./hr. for the slower moving components (II). By eluting the proteins from the Pevikon with water, component I was present in both sulfo-protein fractions at twice the level as that of component II when 20-mg. samples were used. The recovery was approximately 80%.

Ultracentrifugation indicated that all four components had a 2.55 sedimentation rate. From amino acid analysis a molecular weight of 21,500 was found for each. Alanine was found to be the N-terminal amino acid of all four polypeptides. Immunochemical comparisons by precipitin test indicated cross reactivity among all four components.

Ultraviolet light extinction coefficients ($E_{1\%}^{1\text{cm}}$) were 17-18 for components I, and 20-21 for components II. These differences were in agreement with the amino acid compositions of the components which showed tyrosine levels of components II to be 20% greater than those of components I. Lysine levels of both I's were nearly 2 \times those of the II's; alanine levels of the I's were 1.5 \times those of the II's.

By vertical acrylamide gel electrophoresis, sulfo- γ I, sulfo- γ II, and sulfo-albuminoid II gave very similar patterns of 2 close bands each. However, sulfo-albuminoid I exhibited many additional bands above these two (similar to α -crystallin).

The data, including a mathematical analysis of amino acid compositions, indicate that the albuminoid of the young rat lens is composed of approximately 66% γ -crystallin and 33% α - (perhaps in the subunit form). Interactions between γ - and α -crystallins leading to an insoluble product could involve all types of protein-protein binding, and may explain species and aging differences in the makeup of albuminoid.

Supported by the Rochester Eye Bank and PHS grant #NB 03081.

Possible role of protein synthetic enzymes in the control of protein synthesis in higher organisms. AUDREY E. V. HASCHEMEYER.

Comparative studies on aminoacyl-tRNA synthetases and aminoacyl transferases have been made in rat, frog (*Rana catesbeiana*) and toadfish (*Opsanus tau*). Enzyme activity levels in toadfish liver have been measured in relation to the temperature of acclimation of the animals. Although aminoacyl-tRNA synthetase activity is unaffected by cold (10°) acclimation, transferase activity shows a 60% increase compared with levels in fish maintained at 21° . This change is similar to the apparent increase in liver protein synthesis in 10° -acclimated fish as measured by the incorporation of radioactive amino acids into protein 30 min. after arterial injection. The findings thus suggest a possible mechanism for control of protein synthesis in cold acclimation; however, it depends on whether the step catalyzed by the transferase is rate-limiting *in vivo*. Increased activity during cold acclimation may result from conversion from an inactive or dissociated form. Analogy to studies in rat liver suggests that the enzyme involved is aminoacyl transferase I.

Studies on aminoacyl-tRNA synthetases have been made in rat and frog liver in relation to the possibility of control of codon recognition during changes in protein synthesis induced by thyroid hormone. Three enzyme fractions capable of activating and transferring phenylalanine to rat liver transfer RNA were separated by ammonium sulfate fractionation and DEAE-cellulose chromatography of rat liver extracts. Activities comparable to two of these fractions were found in tadpole and frog liver extracts. Preliminary studies indicate that two different types of phenylalanine-acceptor tRNA may be charged by these enzymes.

Supported by National Science Foundation grant GB 5194.

Studies with reaction centers from Rhodospseudomonas spheroides. J. R. BOLTON
AND R. K. CLAYTON.

A reaction-center fraction isolated from *Rhodospseudomonas spheroides* chromatophores exhibits light-induced changes in its optical and electron spin-resonance (ESR) spectra. In particular, a bleaching at 870 nm (P870) has been found to be closely correlated with the appearance of an ESR signal with a g factor of 2.0023 and a peak-to-peak line width of 10 G. The ESR signal is indistinguishable from light-induced signals found in chromatophores or whole cells.

A careful measurement of the spin concentration showed that the ratio of the light-induced spins to bleached P870 molecules is 1.1 ± 0.1 . In addition the formation and decay kinetics are identical within experimental error under a variety of experimental conditions.

Previous work has shown that P870 is a bacteriochlorophyll molecule in a specialized environment and that the bleaching signifies oxidation. The present work provides strong evidence that the product of the bleaching of P870 is the radical cation of bacteriochlorophyll, P870⁺, and that this radical is the source of the ESR signal in whole cells.

The quantum yield for the bleaching of P870 in reaction centers has been measured, using actinic light of wavelengths 880, 800, 760, and 680 nm. For light absorbed at 880 or 800 nm the efficiency is close to 100%. In a coupled reaction, the oxidation of mammalian cytochrome *c* by P870⁺ proceeds with nearly the same efficiency.

The above results place definite limits on the possibilities for the identity of the primary acceptor. These possibilities are discussed.

JULY 23, 1968

The relationship between carbon atom chain length and effectiveness of n-alkylamines in inducing rhythmicity of gastropod smooth muscle. ROBERT B. HILL.

The isolated radula protractor preparation from *Busycon canaliculatum* has seemed of particular interest because it can be provoked to rhythmicity by the simultaneous presence of two naturally occurring neurohumors, ACh and 5HT. ACh alone will not provoke rhythmicity and 5HT or tryptamine alone do not provoke such regular rhythmicity as they do in the course of relaxing an ACh contracture. A disturbing feature of the induced rhythmicity as a model of natural rhythmicity is the high concentration of indolealkylamines required. However, other biogenic amines are quite ineffective in inducing rhythmicity. N-alkylamines of differing carbon chain length were tested as analogs of the alkylamine side-chain of 5HT and tryptamine. In the series from methylamine to undecylamine, pentylamine and hexylamine were found to be the only substances which consistently and significantly potentiated twitches, and they did so at the 10^{-6} M level, which may be compared to potentiation by 5HT and tryptamine at the 10^{-7} M level. Pentylamine and hexylamine relaxed acetylcholine contractures, and in so doing provoked rhythmicity, at the level of 3×10^{-8} M, whereas 5HT and tryptamine relax with rhythmicity at the level of 10^{-8} M. Acting alone, pentylamine and hexylamine induced irregular rhythmicity at the 10^{-6} M level while 5HT and tryptamine induced more regular rhythmicity at the 10^{-8} M level. Thus the 5 carbon and 6 carbon atom analogs of the alkylamine side chains of 5HT and tryptamine function at concentrations 1000 times lower than the natural neurohumors in inducing rhythmicity of the radula protractor.

Local variations of pigmentary patterns and chromatophoric responses in the fiddler crab, Uca pugilator. K. RANGA RAO AND MILTON FINGERMAN.

Dimorphic pigmentary variants can be distinguished among specimens of *Uca pugilator*. One variant has a dark patch in the metagastric area of the carapace while the other has a light patch. The variant with a dark patch occurs at Levy Bay, King Bay and Porter Island Peninsula, Florida, but occurs neither at Alligator Point, Florida, nor Chapoquoit in Falmouth, Massachusetts. The variant with a light patch occurs at the five localities mentioned above. The chromatophoric responses of both variants obtained from Florida were studied. Both variants exhibited different degrees of adaptation on black and white backgrounds. These responses varied with the size of the crabs and the area from which they were collected. Among eyestalkless individuals the melanin was maximally concentrated in the chromatophores of both variants. However, the degree of white pigment dispersion in the eyestalkless crabs varied with the size of crabs and the area from which they were collected. The red pigment was nearly maximally concentrated in the chromatophores of eyestalkless individuals of the dark patch variant, whereas it was in an intermediate state of dispersion in the light patch variant. The melanin in the chromatophores of eyestalkless individuals of the light patch variant dispersed in response to bright light, whereas the melanin in the chromatophores of eyestalkless dark patch crabs did not.

Supported by Grants GB-5236 and GB-7595X from the NSF.

Reproductive performance of insects from U. S. Biosatellite II. D. S. GROSCH.

No sweeping general cytological conclusion can be made about the effect on cells of radiation delivered during orbital flight. Both cell type studied and criterion of effect are important considerations. Enhanced, null, and antagonistic effects have been demonstrated in *Tribolium*, *Drosophila* and *Habrobracon* and all three kinds of effects can be demonstrated for cells of the ovary.

JULY 30, 1968

Formation of the first cleavage furrow in a telolecithal egg (Loligo pealii). JOHN M. ARNOLD.

First cleavage in the squid egg is meroblastic with the furrow appearing in the center of the blastodisc and proceeding toward the edges. Therefore, it is possible to obtain a temporal sequence of furrow formation by progressively taking sections from the blastodisc edge toward the center.

Electron micrographs show the earliest indications of the furrow as a distal flattened region, with longitudinal surface folds running toward the furrow. Immediately below this is a dense layer of fibrils approximately 70 Å in diameter which run parallel toward and below the base of the furrow. These fibrils appear to be derived from membrane-bound dense bodies which originate in the Golgi apparatus. The folds contain many vesicles and tubules which form an irregular tubular network. The older furrow extends downward into the cytoplasm but still has the dense fibrillar region at its base and an average of 17 longitudinal folds in an expanded region at the furrow base. Frequently associated with the furrow base are large multivesiculate bodies which occasionally open and empty vesicles toward the furrow. Below and around the furrow base there is an extensive area of smooth endoplasmic reticulum-like vesicles which are oriented toward the furrow. When cleavage is completed the base of the furrow is in contact with the yolk mass and only a few longitudinal folds remain, but the dense fibrillar layer is still evident.

A cleavage mechanism can be postulated in which the fibrils are associated into a contractile band which is linked to the plasma membrane. This band forms at its ends, is anchored at the edge of the blastodisc, and upon contraction cuts through the cytoplasm. As contraction occurs the fibrils are laterally compressed together by the resistance of the cytoplasm. This causes the longitudinal folds to form. Then the band is bent back upon itself. As the furrow divides the cytoplasm, most of the longitudinal folds unfold and contribute their plasma membrane to

the "new" surface of the blastomeres. In addition, new membranes must be added, possibly from either the smooth endoplasmic reticulum-like vesicles or from the tubular network in the folds.

This work was supported by NIH Grant HD 02688.

On the distribution of genetic variation in nature. ROGER MILKMAN.

The data on allelic variation in nature, and particularly the many recent studies of electrophoretic mobility variants of numerous enzymes, suggest a generalization comprising four parts: (1) At most gene loci in any species there are between two and five alleles that attain significant frequencies (over 1%) anywhere. (2) The achievement and maintenance of these frequencies is due to (single-locus) heterosis. (3) There are relatively few locally-adapted alleles; the common demands of membership in the species outweigh the special demands of a particular external environment within its range. (4) Certain alleles are fixed in local populations from time to time due to the limitations on the number of polymorphisms that can be maintained in finite populations. Thus distant populations of a species differ mainly in their samples of genes from a common species library.

The reunion of alleles with a history of heterosis can result in the production of some superior progeny by parents of distant origins. This phenomenon is frequent; it would not be expected if local adaptations were the preponderant determinants of genotypes.

Numerous counter-examples suggest that this generalization has many exceptions at every level from locus to species. Yet it may be a useful starting point.

The accurate estimation of the number of loci in a given species is now of fundamental importance; the vast difference in estimates generated by different methods must be resolved.

Rapidly contracting muscles used in sound production by a katydid. R. K. JOSEPHSON AND H. Y. ELDER.

The song of the katydid, *Neoconocephalus robustus*, consists of a series of chirps, each produced by a stroke of the forewings against one another, but the frequency of the wing movements and chirps is so high (150-200 per second) that to a human listener the song is a loud, continuous buzz. Wing movements at frequencies exceeding 100 per second are known for a number of insects, but such frequencies have hitherto always been associated with muscle which is termed asynchronous because there is not a 1:1 relation between muscle action potentials and contractions. Recordings from extra-cellular electrodes implanted in the singing muscles of *Neoconocephalus* reveal that there is a muscle action potential associated with each contraction; these muscles are, therefore, of the synchronous type. The contraction-relaxation time of these muscles during singing is approximately 5 msec., making them among the fastest synchronous muscles in the animal kingdom. The myofibrils of the singing muscles are thin sheets which radiate from the center of the muscle fiber. The myofibrils are densely invested with sarcoplasmic reticulum, a characteristic of very fast synchronous muscles. Mitochondria comprise approximately 44% of the volume of these muscles.

Supported by PHS Grant NB-06054 and by a grant GB-3447 from the National Science Foundation to the Department of Invertebrate Zoology, Marine Biological Laboratory.

AUGUST 6, 1968

Shark and dogfish studies: Pulsatile pressure extracorporeal blood oxygenation by means of silicone membrane "heart-lung" devices and artificial kidney. C. LLOYD CLAFF, ARMAND A. CRESCENZI AND ARNOLD J. LANDÉ.

After reading the title: Three questions may naturally come to mind: Question number one: Why use sharks and dogfish instead of dogs? Question number two: Why a membrane oxygenator? Question number three: Why pulsatile pressure? Shark blood retains 2-2½% urea, while other vertebrate blood retains only 0.01-0.03% urea. The shark therefore is especially suitable for evaluation of artificial kidney devices and oxygenating devices.

Why a membrane oxygenator? There is not a case where, in a biological system, blood is in direct contact with a gas, without a membrane at interface.

Why pulsatile pressure? Pulsatile pressure insures a new interface at every pulse—hence a better oxygenation per square meter of surface. Clinical evidence now shows steady pressure causes areas of anoxia, with resulting brain damage, and kidney complications. A nurse shark 4½ feet long was put on “bypass” at Florida Oceanographic Society Marine Station, Stuart, Florida, using Dr. Landé’s artificial gill and the Claff-Crescenzi pulsatile pressure method of oxygenation, except fluoro-carbon was used instead of oxygen, and the carbon dioxide was washed out with filtered sea water. The extracorporeal circuit was a cannulation from the caudal vein, through a DeBaKey pump, thence to a membrane artificial kidney, to a membrane heart-lung device. The oxygenated blood was returned to the body by cannula in the conus arteriosus. Venous return was assured by implanting a cannula in the heart cavity, sewing up the cavity, and connecting the cannula to a source of pulsating negative pressure.

Anesthesia was accomplished with tricaine methane sulphonate; curare 3 mg./kg. every 30 minutes; heparin 3 mg./kg. intravenous. Results: Hematocrit arterial, 30; venous, 33; pH arterial, 7.37; venous, 7.31; pCO₂ arterial, 17; venous, 22; pO₂ arterial, 350; venous, 180. Blood urea 428–420 net change of 8; no blood sugar present.

The assistance of Louis Sonstegard, Katherine McCusker and Marianne Broome is gratefully acknowledged.

*Asexual and sexual reproduction of *Pygospio elegans* (Polychaeta) in Barnstable Harbor, Massachusetts.* KATHERINE D. HOBSON AND ROGER H. GREEN.

We observed the cycle of asexual and sexual reproduction of *Pygospio elegans* Claparède during a study (1967 to 1968) of the intertidal fauna near Huckins Island. Every month, eight samples of the fauna retained by a screen of 1 mm. mesh were taken from each of two stations (60 cm. and 90 cm. above mean low water).

Pygospio elegans reproduces asexually by breaking into fragments, usually of two or three segments, each of which then regenerates a head and tail. Fragmentation was most intense (increasing the population by a factor of 10) during April and May of both years at the lower station. By June or July, the population had returned to the original level. At the upper station, intense fragmentation occurred only in the spring of 1968.

Females with egg capsules were observed from March through May, but comprised less than 7% of the pre-fragmentation population. Because at least 30% of the water in Barnstable Harbor is exchanged on each tidal cycle, a female releasing 500 larvae, which have a pelagic life of two weeks, would be replaced by only 0.02 larvae. Thus this population of *P. elegans* must be maintained by sexual reproduction of adults outside the harbor or by asexual fragmentation. The fragments would not be subject to tidal flushing because they remain in the tube of the adult during the regeneration process.

During spring in Barnstable Harbor, a large portion of the population reproduces asexually and a small portion reproduces sexually. We do not know the role of asexual reproduction in the life history of *P. elegans*, nor the extent of its importance in maintaining the population in Barnstable Harbor.

This research was supported by a grant from the Whitehall Foundation to the MBL Systematics-Ecology Program; SEP Contribution No. 163.

Limb regeneration in adult amphibians. MARC H. DRESDEN.

Investigations during the past 30 years have shown that regeneration of amputated limbs in adult amphibians requires an adequate nerve supply. In an effort to gain a more basic insight into the role of nerve in regeneration, a study of the macromolecular syntheses of the regenerating blastema was initiated.

Both forelimbs of adult newts (*Triturus viridescens*) were amputated proximal to the elbow and regeneration was allowed to proceed to the palette stage (3–4 weeks post-amputation). These animals were then denervated unilaterally (the other arm serving as a control) by resection of spinal nerves 3, 4, and 5. After various periods of time the animals were

injected intraperitoneally with a mixture of ^3H -thymidine and either ^{14}C -leucine or ^{14}C -uridine. Four to five hours later the animals were sacrificed and the radioactivity incorporated by the blastema was assayed. Within seven hours the denervated blastema begins to show a decreased capacity for uridine incorporation, followed within 25 hours by a decreased capacity for thymidine and leucine incorporation. Sham-operated animals show no change in thymidine and leucine incorporation. Thymidine incorporation decreases to about 25% of the control regenerating capacity, which is equivalent to that found in normal arms and in the proximal regions of regenerating arms. No significant effect of denervation on the incorporation of thymidine and leucine is seen in non-regenerating limb tissues.

In addition, it was found that the changes in thymidine incorporation following denervation can be mimicked *in vitro*. When the blastema derived from animals denervated 48 hours previously are incubated in a tissue culture medium containing ^3H -thymidine they incorporate about 35% as much of the isotope as do the blastema derived from undenervated animals or from animals denervated 17 hours previously (at which time thymidine incorporation has not yet begun the decrease *in vivo*).

The changes in incorporation of precursors of RNA, DNA, and protein following denervation precede any morphological changes. The magnitude of the denervation effect and its specificity for regenerating limb tissues indicates that there may be a dual control over macromolecular syntheses in amphibian tissues.

Supported by a grant (#GB 5194) from the National Science Foundation and a fellowship from the Atomic Energy Commission.

AUGUST 13, 1968

Segregation of filamentous protein structures from the axoplasm of the squid giant nerve fiber by uranyl ions. J. METUZALS.

Certain heavy metal ions show specificity for reactive groups of protein molecules. The proper choice of metals can be very useful in fractionation, crystallization of desired proteins and in labeling of active sites of proteins by virtue of such specificity. UO_2^{++} ions of a 1% uranyl acetate solution at pH 4.4 display a specific affinity to filamentous protein structures of the squid giant nerve fiber axoplasm. At pH 4.4 the uranyl ions segregate the filamentous elements of the axoplasm in a coiled, threadlike preparation surrounded by a flocculent precipitate. The uranyl ions combine probably with the carboxyl groups of the main amino acids of the protein—the glutamic and the aspartic acids. Segregation cannot be observed at pH 7.8. Affinity to the filamentous axoplasmic protein structures is displayed also by Ag ions, but not by ions of La, Co, Pb and Zn salts.

Because of the fixing effect of the uranyl ions at the macro-molecular level and the simultaneous increase of contrast, the preparations obtained by the uranyl procedure are well suited for an integrated approach using both high resolution electron microscopy and physicochemical methods. Three hierarchic levels of coiled configurations can be observed as a characteristic structural pattern of the thread preparation: coiling at (1) the light microscopic level visualized by interference optics, (2) the fine structural, and (3) the macromolecular levels disclosed by electron microscopy.

Analysis data of the axoplasmic thread, segregated by uranyl ions, show the following composition: N 10.8%; U 18.8%; P 4.6%; Ca 0.6%; Mg 0.08%; 1-2% ribonucleotides. Amino acid composition expressed as moles per cent: Ala 9.0; Arg 3.5; Asp 11.0; Cys 0.8; Glu 14.4; Gly 10.8; His 1.5; Ileu 2.5; Leu 6.8; Lys 6.4; Met 2.5; Phe 3.5; Pro 5.7; Ser 9.6; Thr 5.7; Tyr 2.5; Val 3.9.

Supported by the Medical Research Council of Canada.

Observations of living giant nerve fibers of the squid by differential interference-contrast microscopy. J. METUZALS, C. S. IZZARD AND M. GOSPODNETIĆ.

Using the differential interference-contrast microscope (Carl Zeiss), one can obtain thin "optical sections" of the living giant nerve fiber of the squid without interference by structures above or below and approach the limit of resolution. For observations with transmitted light,

the stellate ganglion with a 5-cm. length of giant nerve fiber, ligated at the distal end, was mounted in sea water. *In situ* giant nerve fibers were observed using the reflected interference-contrast microscope. One cm. of fiber was cleared of surrounding tissue and a fragment of aluminized coverslip, as a reflecting surface, was inserted between the fiber and the underlying mantle wall.

Parallel threadlike elements of 1–2 μ diameter and irregular dense bodies of varying size represent the main structural features of the axoplasm at the light microscope level. A peripheral ectoplasm and a central endoplasm can be distinguished by differences in packing and arrangement of these elements. The ectoplasm exhibits a dense parallel packing of straight threadlike elements, whereas in the endoplasm the threads often show a looser reticular array. In the ectoplasm the threadlike elements are oriented at an angle of 5–15° to the longitudinal axis of the fiber. In optical sections of the upper and the lower ectoplasmic regions, the direction of this angle is reversed with respect to the axis of the fiber, indicating that the threads are wound helically along the fiber. In certain locations of the endoplasm, sinuous threads are in register across the width of the fiber resulting in the overall appearance of large scale cross banding. The structural configurations observed in the living fibers correspond well to those observed by light microscopy of stained, thick plastic sections of material prepared for electron microscopy.

Supported by MRC of Canada and NIH grants.

Quantitative radioautography of vitellogenesis in Artemia salina. REV. J. D. CASSIDY, O. P., C. F. STARMER AND L. J. BEAUREGARD.

Genetic transcription sites were localized by grain frequencies counted on series of 0.5- μ cellular sections in differentiating ova and accessory cells of adult brine shrimp. Isotope distribution was scored after *in vivo* exposure to tritiated thymidine, uridine and actinomycin-D. Nine incorporation sites, dosage and rate of isotopic uptake, as functions of time, were tested using a multivariate general linear model in the IBM 1130 computer. The primary model related multiple sites in the vitellarium to time effect and precursor effect as a 3 \times 3 multivariate factorial design. Hypotheses tested specific incorporation of precursors at a particular time and similarities between precursors over time and time averaged over precursors. Results were evaluated using the multivariate likelihood ratio to determine treatment effects (time and precursor) on the entire multivariable observation (incorporation sites) and individual "F" ratios (adjusted according to the Bonferonni inequality) to determine treatment effects on individual incorporation sites.

Thymidine, uridine and actinomycin-D were incorporated in similar amounts, pooled over time, by the oocyte yolk platelets and nurse cell nuclei. Comparison of thymidine and uridine indicated comparable isotopic uptake by nuclei of the follicular layer and lower ratios within the nurse cell cytoplasm and the oocyte-nurse cell interface. Evaluating time effect on precursor incorporation yielded no difference in uptake from 5 to 10 and 5 to 20 min. in follicular cytoplasm and the region adjacent to yolk platelets. Tests for dissimilar incorporation of each precursor/time determined no interaction at the oocyte-nurse cell interface, follicle cytoplasm and yolk platelets; significant interaction adjacent to yolk platelets and follicular nuclei; and highly significant interaction involving the germinal vesicle, cortical ooplasm, nurse cell nuclei and cytoplasm. This cytological-cytochemical-computerized approach identified yolk platelets as a cytoplasmic center of informational template synthesis in the developing egg chamber.

Supported by USPHS Grants ES-00044, GM-13625 and NSF Grant UG-1209.

GENERAL SCIENTIFIC MEETINGS

AUGUST 19–22, 1968

Abstracts in this section are arranged alphabetically by authors. Author and subject references will be found also in the regular volume index, appearing in the December issue.

An analysis of cleavage furrow formation in the egg of Loligo pealii. JOHN M. ARNOLD.

Cleavage furrows in the meroblastic squid egg form from the center outward. The base of these furrows is associated with many 70 Å fibrils which arise from Golgi-derived, membrane-bound dense bodies. In an earlier report a theory of fibrillar contraction in furrow formation was put forth. Simply stated this theory involves a progressively distal formation of a fibrillar band below and associated with the plasma membrane. A cleavage furrow is formed when this band contracts. The experiments described below test this hypothesis.

If fibrillar contraction cuts into the cytoplasm to form the furrow, then curvature of the egg surface is necessary. Prior to first cleavage the blastodisc was flattened with a coverslip. This not only prevented furrow formation, but also stopped it once it had begun. Release of the pressure resulted in furrow formation a few minutes later. Heavy compression before cleavage prevented furrowing and resulted in a delayed, irregular cleavage pattern. Flattening one side of the blastodisc parallel to the first furrow caused unilateral delay or complete inhibition. Lateral flattening changed the shape of the blastodisc so the future ends of the first furrow were well below the upper margin of the yolk mass. When the furrow formed it indented the yolk. In normal eggs third cleavage is asynchronous with the anterior furrows forming slightly before the posterior furrows. These furrows appeared attached to the second furrow because it is distorted just at the junction with the forming third furrow. This strongly suggests the forming furrow is contracting. By using an ultraviolet microbeam it was possible to irradiate the forming fibrillar band and cause the second furrow to disappear unilaterally. A very faint birefringence, possibly the fibrillar band, is apparent in the future furrow region just prior to actual cleavage. All of these results and observations are consistent with a mechanism of furrowing by a contractile fibrillar band.

This work was supported by NIH Grant HD 02688.

A ciliary activity inhibitor extracted from the nidamental gland of Loligo pealei.

BURR G. ATKINSON AND N. A. GRANHOLM.

An extract prepared from the nidamental gland of the squid causes adjacent cilia to adhere to one another, but the effect is *not* a general clumping: groups of cilia appear attached at their tips. The overall effect of the factor is to bring the swimming movements of ciliated organisms to a halt by reducing the *effectiveness* of ciliary activity; the actual beating continues.

Histochemical examination of the nidamental gland discloses that it is primarily composed of columnar cells containing a substance which stains metachromatically with toluidine blue and which is PAS and alcian blue (pH 2.5) positive. While the squid jelly case and dry smears of the crude viscous extract exhibited similar histochemical properties, the accessory nidamental gland did not. In keeping with this histological evidence, biological activity could not be extracted from the accessory nidamental gland, but was obtained from the nidamental gland and the egg jelly case.

The viscous preparation initially extracted from the gland has a milky-white color. Low-speed centrifugation produces a clear supernatant fraction and a viscous milky-white pellet. Although the clear supernatant was used as the source of biological activity, more activity could be extracted from the pellet by homogenization and centrifugation. The pooled supernatants were centrifuged at 105,000 × *g* and the activity was recovered in the supernatant fractions. The ratio of proteins to neutral hexoses in these final supernatant fractions was 5:1. The biological activity of this fraction was not labile to dialysis against Versene or 0.1 N NaCl, nor to 10 minutes in a boiling water bath. The high-speed extract could be prepared in Millipore-filtered sea water, distilled water or CMF sea water; it could then be lyophilized and resuspended in the desired medium. Further purification of the lyophilized fraction involved a separation of the biological activity from most of the proteins on a G-25 Sephadex column.

Supported by Grant 5-TO1-HD0014-09 from the Public Health Service.

Cellular clots in Pagurus longicarpus and the possible role of microtubules. F. B. BANG.

Clotting of amebocytes was studied following the cutting of the antenna and placing the cut end in sea water. N-ethyl maleimide in concentrations as low as 0.005 mM in sea water was absorbed over a period of hours by *Pagurus* and acted as an anti-cellular anticoagulant. The effect persisted for several days. Most of the animals exposed to this concentration survived. Electron microscopy of clotted cells, following glutaraldehyde fixation and later treatment with osmium, showed numerous microtubules which apparently strengthened the second stage of clotting. These were not found one and two minutes after cutting the antenna but were found from three minutes to 11 days afterward. Several anaesthetics and drugs known to affect the microtubules of spindle figures during mitosis altered the clot of *Pagurus* when the cells were allowed to spill into a drop of an appropriate solution. These presumably acted by keeping the cells from entering into the second stage of the clot. Urethane (6%) and hexylene glycol were the most effective. The interrelationship of these findings is as yet unexplored.

Electron microscopic examination of subcellular components containing melanophorotropic activity, separated by density gradient centrifugation, from neurosecretory cells in the eyestalks of the fiddler crab, Uca pugilator. CLELMER K. BARTELL, KENNETH MAY AND MILTON FINGERMAN.

The melanin-dispersing hormone from the eyestalks of *Uca pugilator* is thought to be stored within neurosecretory granules of the sinus gland. In an effort to isolate these granules, the microsomal fractions of eyestalk extracts were prepared by differential centrifugation of tissue which had been sonicated in 0.82 M sucrose. Much of the melanin-dispersing activity of the extract was contained in the microsomal fraction. The microsomal pellet, resuspended in 1.00 M sucrose, was layered over a discontinuous sucrose gradient, from 1.00 to 1.75 M concentration, and centrifuged for two hours at $200,000 \times g$. Bio-assay of the fractionated gradient indicated the presence of two distinct fractions having different densities. The fractions were adjusted to a concentration of 0.82 M sucrose, aliquots were assayed for melanin-dispersing activity in eyestalkless *Uca*, and the remainder of each fraction was reduced to a pellet by centrifugation. The pellets were then prepared for electron microscopic examination. The less dense fraction contained an assortment of electron-lucent, vesicular membranes and an occasional neurosecretory granule. The more dense fraction was a non-homogeneous mixture of cell fragments including numerous electron-dense particles having a size and electron-density similar to the neurosecretory granules observed in sections of the intact sinus gland. In contrast, extracts prepared in physiological saline, instead of sucrose, contained only the less dense active fraction: the granules appeared to be disrupted by extraction in saline.

These results indicated that much of the hormonal activity can be isolated in association with specific particulate fractions from the disrupted cells of the eyestalks from *Uca*. The activity is probably contained within neurosecretory granules and fragments of the granules. Further refinement of the technique should produce fairly pure preparations of granules that will be useful for chemical analysis.

Supported by Grant GB-7595X from the NSF.

Morphology of the eye of the flame fringe clam, Lima scabra. ALLEN L. BELL AND GEORGE J. MPITSOS.

The electrophysiology of the vision of *Lima* has been studied by one of us (Mpitsos). The eyes give both an on-response and an off-response to stimulation with light. This report gives the preliminary results of fine structure studies which were undertaken with the purpose of locating morphological correlates to the electrophysiological data.

The eyes of *Lima* are cup-shaped organs each about 600 microns long, 200 microns wide and 150 microns deep, lying along the edge of the mantle. Longitudinal sections through the cup show that the retina consists of a pseudostratified epithelium comprised of two cell types: visual cells and pigment cells.

The visual cells have a large inner segment or cell body attached by a short stalk to an outer segment or rhabdome. These cells are 40 to 50 microns long, 10 to 15 microns wide and stand vertically on a thick basement membrane. In addition to a nucleus, mitochondria and Golgi material, the base of the visual cell is filled with a large amount of membranous material consisting of tubules and vesicles. Visual cell outer segments measure 8 to 10 microns in diameter and have numerous microvilli extending outward from their surface interdigitating with those of adjacent outer segments. Pigment cells lie between the visual cells. Their nuclei are located apically and the cells narrow as they approach the basement membrane. The apical surface of the pigment cells is at the level of the stalk of the visual cells. Thus the outer photoreceptor portion of the visual cells extends beyond the most apical part of the pigment cells.

Nerve fibers are found between the basal surface of the retinal cells and the basement membrane. Occasionally one of these fibers appears to end in a swollen bulb which contains clear vesicles 600 to 800 Å in diameter and dense cored granules 1200 to 1500 Å in diameter.

Preliminary examination of the "lens" of the eye reveals large bundles of cilia and numerous processes which appear to be neural. Nerve terminations are numerous in this region. The role, if any, played by the lens structure in the visual response is being examined.

Part of this research was supported by a Grass Foundation Fellowship, 1967, to George J. Mpitsois.

Electrical coupling of embryonic cells by way of extracellular space and specialized junctions. M. V. L. BENNETT AND J. P. TRINKAUS.

In cleavage stages of the egg of the teleost, *Fundulus heteroclitus*, current applied in one cell produces very nearly the same potential in all cells. In later stages of blastula formation, current in one cell produces a larger potential in that cell than in distant cells. However, as distance from the polarized cell is increased, the potential rapidly approaches that in the distant cells. If current is applied in the segmentation cavity of the gastrula, the same potential is recorded throughout the cavity and in all cells as well. The resistance between the cavity and the exterior is very high indicating that junctions connecting outer margins of the superficial cells (Trinkaus and Lentz, *J. Cell. Biol.*, 32: 139, 1967) seal off the cavity from the exterior and are functionally "tight." Also, the resistance of the outer membrane of superficial cells is high (ca. 100 kΩcm²), much higher than that of underlying membrane abutting on the segmentation cavity. Thus, current can pass from cell to segmentation cavity to cell with little leak to the exterior, and cells can be coupled by way of intra-embryonic extracellular space. Superficial cells also appear to be coupled by way of junctions between them because cells near a polarized cell can be at a higher potential than the underlying segmentation cavity. Coupling of embryonic cells by way of junctions is also indicated since pairs of blastomeres isolated mechanically are usually coupled and single cells reassociated *in vitro* can develop coupling. In this embryo, junctions can transmit small molecules from cell to cell over short distances, but transmission over large distances must be primarily by way of extracellular space.

Dogfish white blood cells: protein synthesis in vivo and in vitro. GRACIELA C. CANDELAS, SANFORD TEMES, JOSÉ R. ORTIZ AND SEYMOUR GELFANT.

The technique for culturing dogfish blood cells has been developed in our laboratory. Studies have indicated that the metabolism of cells in culture may be different than *in vivo*. We have studied protein synthesis of dogfish white blood cells *in vivo* and *in vitro* under a number of conditions by following the incorporation of tritiated amino acids in TCA-precipitable protein. Preliminary studies of the protein synthesizing units of these cells were also undertaken.

A comparison of the rates of protein synthesis *in vivo* and in culture medium (20% plasma, 80% elasmobranch Ringer's) revealed a sharp drop when the cells were placed in culture. The rate of protein synthesis was reduced by about 65% *in vitro*. It was found that this drop occurs within the first few minutes of culture and that the rate remains reduced throughout the first hour. However, by 24 hours of culture, the rate of protein synthesis returns to almost 100% of the *in vivo* rate.

If the cells were cultured in 100% plasma, the drop in protein synthesis was minimized. After 72 hours of culture, cells in 100% plasma displayed higher rates of protein synthesis than cells in the usual medium (20% plasma). Thus plasma was shown to be beneficial either in promoting proliferation, activating protein synthesis, or both.

Density gradient analysis of the $7000 \times g$ supernatant revealed some protein synthesizing activity in a heavy peak, but most activity was found in the light monosome peak, indicating the need for an RNase inhibitor in these studies. Analysis of the resuspended $105,000 \times g$ pellet yielded a heavy peak containing all the activity. The activity in this heavy region was insensitive to standard RNase and deoxycholate treatments.

Supported by research grant GB-2803 from the National Science Foundation.

The use of luminescent systems and image intensification as cytochemical probes.

I. The distribution of ATP in the muscle cell. G. B. CALLEJA, GEO. T. REYNOLDS AND R. J. PODOLSKY.

Because of its sensitivity to photons, the image intensifier may be used to detect the spatial distribution of photon sources not discernible by the unaided eye. The localization of microsources of light by image intensification has already been achieved in *Noctiluca*, a bioluminescent dinoflagellate. In addition, cells which are not luminescent may be made to emit light if a luminescent system is introduced into them. If the system introduced is complete except for one necessary component, not only the amount, but also the spatial distribution of this one component can be determined. Luminescent systems which possess potential cytochemical uses include firefly luciferin-luciferase for the detection of ATP, aequorin from *Aequorea* for Ca^{++} , bacterial luciferase for elements of the electron flow system, etc.

We report in this paper the preliminary results of our studies on the distribution of ATP in the muscle cell. Single fibers were isolated from the semitendinosus muscle of *Rana pipiens*. Firefly luciferin-luciferase was introduced as a microdrop onto the surface of fibers with or without sarcolemma. The preparations, viewed through a microscope attached to an image intensifier, were then photographed in the dark. We found that most of the ATP that resulted in light emission was not preferentially associated with any structure identifiable by means of the light microscope. To the limits of resolution so far achieved (the order of 1 micron), it appears that ATP is distributed uniformly along the longitudinal axis of the cell. In the intact fiber, luminescence was restricted to the cell surface. Thus, whereas the sarcolemma effectively excluded luciferase from the sarcoplasm, it did not prevent contact between ATP and the enzyme. The kinetics of emission suggests that light from the intact fiber was due to ATP either leaking out or being regenerated on the cell surface.

Supported in part by AEC contract AT(30-1)-3406.

Effect of cycloheximide on ciliogenesis, ciliary regeneration, and protein synthesis in Arbacia embryos. FRANK M. CHILD AND MATTHEW N. APTER.

Regeneration of amputated cilia in protozoa generally requires some protein synthesis, although assembly of regenerating cilia can occur in the absence of protein synthesis if the necessary proteins have been allowed to accumulate. In *Paracentrotus*, Auclair and Siegel, 1966 (*Science*, 154) conclude that protein synthesis is not required for ciliary regeneration, since puromycin will not prevent regeneration. They suggest that all of the required proteins exist in a pre-formed pool drawn upon during cilia formation. Do sea urchins differ from protozoa in the control mechanism governing the formation of cilia?

We have now shown in *Arbacia* that CH (cycloheximide, 5 mg./ml.) will significantly delay the formation of motile cilia in regenerating blastulae and early gastrulae. CH will also delay the normal formation of cilia in blastulae when CH is applied after fertilization as late as the elapse of 84% of the scheduled time of ciliogenesis.

Cycloheximide inhibits the incorporation of C^{14} -valine into TCA-insoluble protein of late blastulae and early gastrulae. Five mg./ml. of CH permits incorporation at 10% of the control rate; lower concentrations of CH inhibit less.

Since we assay the formation of cilia by sensitively detecting the first appearance of motile shafts, the delays produced by CH suggest that ciliogenesis and ciliary regeneration in

Arbacia are processes directly dependent on the synthesis of at least one protein in the half-hour or so preceding the formation of motile shafts. Inhibition of protein synthesis by CH could act by delaying the accumulation of this protein required for shaft assembly.

This work was supported by NIH grant GM-13993.

Immotility and motility of acoel turbellarian spermatozoa, with special reference to Polychoerus carmelensis. DONALD P. COSTELLO AND HELEN M. COSTELLO.

Examination of living adult *Polychoerus carmelensis* over an extended period of time (10 weeks), from beginning to height of their breeding season, has yielded information on spermatogenesis, spermiogenesis, oogenesis, and on activities in such organs as the bursa seminalis. Since *Polychoerus* can be effectively flattened, it was possible to study living specimens under phase oil immersion objectives, and to take photomicrographs of salient features.

The bi-axial filamented spermatozoan is almost invariably immotile, as seen in the distributed testes, sperm ducts, ejaculatory ducts, or when emitted into sea water. These observations are in perfect accord with the known fact that although fertilization of the eggs by the sperm of the same animal would be anatomically possible in the hermaphroditic Turbellaria, self-fertilization is unknown or very rare. Mating occurs by reciprocal copulation, and the sperm of another individual are introduced by way of the female antrum into the bursa seminalis (seminal receptacle). In this bursa, and in the paths and cavities leading from it to the mouthpieces (nozzles), the spermatozoa become intensely active, wriggling with snake-like undulations. The motion is caused by two lateral undulating membranes on opposite sides of the long flattened body of the spermatozoan. The wave motion in these begins at the posterior end and progresses forward.

We have observed active spermatozoa proximal to (nearest the bursa) the mouthpieces, where some appear to be attempting to enter the tiny tri-radiate cuticular canals. Inert, dark thread-like material is frequently seen at the distal ends of the mouthpieces. No evidence was obtained indicating that these threads are active, living sperm. A few active sperm were observed, however, passing between or around the mouthpieces. This suggests that the function of the mouthpieces is not necessarily to direct the sperm toward the eggs. They may have the function of hindering and digesting excess sperm.

Aided by a grant from the National Institutes of Health, GM-15311.

Oxygen-linked dissociation of hemocyanin. I. Experimental evidence. HENRY A. DEPHILLIPS, JR. AND K. E. VAN HOLDE.

The investigation of the oxygen binding properties of hemocyanin obtained from the squid, *Loligo pealei*, has shown a novel dependence of per cent oxygenation on molecular size. Confining our attention to a concentration of 5 mg. protein/ml., an ionic strength of 0.125 (0.1 ionic strength buffer plus 0.01 M Mg⁺⁺) and a temperature of 20° C. we find that over the pH range of 6.0 to 8.3 in the absence of oxygen (under argon) the 59S component predominates (about 95% 59S species and 5% of non-oxygen binding protein). In the same pH range for samples equilibrated with oxygen in air the 59S component predominates below pH 7.1 and above pH 8.0. Between these values, however, the amount of 59S decreases as the pH increases, to a minimum at pH 7.5, where the 19S component is the only molecular species present, and then increases to 95% at pH 8.0 once again. A similar phenomenon is observed when samples are equilibrated under pure oxygen. In this case, the 59S component is stable below pH 6.8 and above pH 7.4. The decrease in the 59S species reaches a maximum at pH 7.1 where the 19S component is the only molecular species present.

That these are true equilibrium phenomena was demonstrated in an experiment in which the relative amounts of 59S and 19S components were varied by varying only the amount of oxygen in equilibrium with the solution.

Correlation of all these results is achieved if the amount of 59S component is displayed as a function of percent oxygenation. This graph shows that this component predominates below 50% saturation and above 95% saturation. Above 50% saturation, dissociation to the 19S component occurs and is complete in the range of 75 to 90 per cent saturation.

This research was supported by Public Health Service Grants HE-11671 and HE-12326.

Excitation and inhibition of Purkyně cells in the cerebellum of Mustelus canis.

JOHN C. ECCLES, HELENA TÁBOŘÍKOVÁ AND NAKAAKIRA TSUKAHARA.

The selachian cerebellum resembles histologically cerebella of higher vertebrates. However, basket cells are absent, though in the molecular layer there are numerous stellate cells that may be regarded as undeveloped basket cells. In investigating the mode of action of the various cells, we utilized the same techniques as for the mammalian cerebellum. After immobilization by intramuscular injection of Tubocurarine, the spiracles are perfused by aerated sea water, the cerebellum is exposed and the skull is rigidly clamped. Concentric needle electrodes are employed for stimulating with currents driven by 0.2-msec. pulses of 10 to 90 volts. One stimulating electrode in the stratum fibrosum excites nerve fibers passing to cerebellar cortex, the mossy and climbing fibers, and also Purkyně axons. A surface electrode stimulates parallel fibers passing laterally from the granule cells. Glass micropipettes (3 M NaCl, 2 to 5 M Ω) and (3 M KCl, 10 M Ω) record, respectively, field potentials and intracellular potentials from Purkyně cells.

Stimulation of Purkyně axons gives field and intracellular potentials. Axonal spike potentials precede slower spike potentials of cell invasion. The depth profile suggests that antidromic invasion fails in superficial dendrites. With double stimulation there is recovery, often to low supernormality around 50 msec. Superimposed on the antidromic response there is often a rhythmic wave that by depth profile and intracellular recording is identified as compounded of all-or-nothing climbing fiber responses, initial repetitive spikes at about 250/sec. being superimposed on a 20-msec. depolarization. Climbing fibers are spontaneously active, often in bursts, phases up to 20/sec., then silences of seconds.

Parallel fiber stimulation depresses antidromic invasion of Purkyně cells, often in two phases. The initial depression of about 50 msec. is largely a sequel of the initial activation of Purkyně cells, but the later depression—from 70 to 500 msec.—gives evidence of prolonged inhibition, presumably produced by the stellate cells, which thus resemble basket cells.

This work was supported by the Grass Foundation.

The response of the red chromatophores of the prawn Palaemonetes vulgaris to cyclic 3',5'-adenosine monophosphate. MILTON FINGERMAN, ROBERT D. HAMMOND AND RENATE SCHLENZ TRUE.

Cyclic 3',5'-adenosine monophosphate has been proposed as the intracellular mediator of the action of several hormones including intermedin. The following experiments were devised to determine a possible role of cyclic 3',5'-AMP in mediating the actions of the pigment-dispersing and pigment-concentrating hormones on the red chromatophores of *Palaemonetes vulgaris*.

Solutions of cyclic 3',5'-AMP (0.15–15.00 mM) prepared in isosmotic sea water were tested *in vitro* on the red chromatophores of the epidermis adhering to the exoskeleton dorsal to the heart. The section of exoskeleton with adhering epidermis was cut into two portions, one serving as the control for the other. In proportion to the concentration in the solution, cyclic 3',5'-AMP dispersed the red pigment in chromatophores whose pigment was maximally concentrated at the outset, but had no red pigment-concentrating action on chromatophores with maximally dispersed pigment. The pigment-dispersing response to 3.75 mM cyclic 3',5'-AMP was independent of the calcium concentration (0.0–4.6 grams per liter) in isosmotic solutions. In contrast, it was previously shown that the response of these chromatophores to a constant amount of red pigment-dispersing hormone (RPDH) is directly related to the calcium ion concentration in the medium and that of all the substances tested, calcium ion permitted a response to RPDH at least twice as large as that permitted by any other substance.

These results suggest the following hypothesis of the action of RPDH. RPDH causes a rise in the calcium concentration within the chromatophore. An increase in the internal calcium concentration is the signal that activates adenylyl cyclase which catalyzes the synthesis of cyclic 3',5'-AMP. The amount synthesized is proportional to the amount of RPDH and to the amount of calcium that entered the chromatophore. Cyclic 3',5'-AMP then triggers pigment dispersion in proportion to the quantity that was synthesized.

Supported by Grant GB-7595X from the NSF.

A new and improved source of trehalase. FRANK M. FISHER, JR. AND ROBERT O. MCALISTER.

The enzyme trehalase (α - α -1-glucoside 1-glucohydrolase, EC 3.2.1.28) has been isolated from a bacterial symbiont from the midgut of *Blaberus craniifer*, a tropical cockroach. The enzyme has been purified 80-fold and some of its physical characteristics determined.

The bacteria, a species of *Aerobacter*, were grown in minimal salt media fortified with 5% trehalose (α -D-glucopyranosyl 1-glucopyranoside) and the enzyme liberated by alternate freezing and thawing with intermittent sonication. The supernatant was treated with $MnCl_2$ followed by acetone (30% V/V) at $-7^\circ C$. The acetone precipitate was resuspended in citrate buffer (pH 4.6) and subjected to further purification on a DEAE-cellulose column. Elution of the enzyme was effected by a stepwise gradient of increasing pH followed by increasing concentration of NaCl.

The purified enzyme is specific for trehalose and does not have any activity toward 16 other oligosaccharides, glycogen, starch, or any phosphate esters of glucose. The pH optimum is 4.6 and there is little inhibitory effect of heavy metals. The K_m is $2.24 \times 10^{-3} M$ and the V_{max} is about $290 \mu M$ hydrolyzed Mg^{-1} protein hour $^{-1}$. Due to the absence of other carbohydrases in the crude supernatant and the relative ease of purification, this trehalase affords a rapid and easy method of determination of this unique disaccharide.

Supported by Grants (C-239) from the Robert A. Welch Foundation and the USPHS, GM 12263.

The cysteinyl-dopa-iron pigmentation of red species. PETER FLESCH.

Red hair of man, dog, rabbit and red chicken feathers contains amphoteric yellow-orange pigments. These may be extracted at room temperature with dilute acids or alkalies as orange "protopigments" which, upon heating with acids, are converted to red, indicator-like substances. This color change is due to decarboxylation, as shown by Italian chemists. The pigment consists of protein, ferric ion, and a "chromophore" group. The latter is responsible for the indicator behavior and characteristic absorption spectrum of the pigment. Italian chemists have established that the chromophore group results from the polymerization of cysteinyl-dopa with dopa. In brilliant experiments they synthesized this grouping *in vitro*. The iron apparently links the chromophore group to the protein. In contrast to the black melanoproteins, this protein contains no cystine or dopa. The proteins obtained from the pigments of different species have remarkably similar compositions.

The cysteinyl-dopa-iron pigments are apparently widespread in nature. It is not known whether they occur in amphibians or fishes. The presence of these pigments can be established by extracting red epidermal structures with boiling acids, precipitating the pigments at their I.P. (in the neutral range for the red "pheomelanoproteins," around pH 3-4 for the chromophore) and obtaining the pure chromophore by repeated filtration through Sephadex G-25 and cation exchange resin columns. The chromophore may be recognized by its purple color in acids, reversible change to orange on alkalization and characteristic absorption bands.

Because of their unique properties and easy availability of their protein moieties, these pigments are of considerable importance to geneticists, anthropologists and biochemists. Three groups independently studied their chemical nature; hence the terminology ("trichosiderin," "pyrrotrichol," "pheomelanin," etc.) is confusing and will have to be reconciled at international meetings. Until then the term "red pheomelanoprotein" is suggested for the natural pigment and "cysteinyl-dopa-polymer" for its chromophore group.

Distribution and dispersion patterns of the dwarf tellin clam, Tellina agilis. WILLIAM H. GILBERT.

Field work for this study was done at Barnstable Harbor, Mass., during July, 1968. Random samples were taken at several stations along a transect from the mainland to Horseshoe Shoals. Samples were sieved through a 0.5-mm. mesh; specimens of *T. agilis* were counted, and shell-lengths were measured. The size-class frequency distribution indicated that 2.5 mm. separated juveniles (young-of-the-year) from adults. The upper boundary of the population distribution is around 0.5 feet above mean low water (MLW), where adult

density is approximately 50/m.². Adult densities of around 150/m.² were found down to 2.5 feet below MLW, the deepest area sampled. Juveniles were found concentrated around MLW, between the eel grass and the channel, at densities of up to 830/m.².

Juveniles exhibited an aggregated dispersion pattern; adult dispersion was not shown to differ significantly from random.

The following correlation was found between clam size and substrate composition: adults and juveniles of mean lengths 4.6 mm. and 1.8 mm., respectively, were found where the median sand particle diameter was 0.5 mm. with a large variance (degree of sorting, D.S., equal to 1.5). Significantly smaller adults and juveniles (3.5 mm. and 1.5 mm., respectively) were found in the area of juvenile abundance, where median particle size was 0.2 mm. with small variance (D.S. = 0.6). The settling velocity of juvenile tellin clams was found to approximate that of sand 0.2 mm. in diameter.

The above data suggest that hydrodynamic forces may be responsible for the intraspecific size-class zonation observed.

This work was supported by NSF funds and a Woods Hole Scholarship through the University of Massachusetts, and aided considerably by the staff and facilities of the Marine Ecology Course at the Marine Biological Laboratory.

Factors influencing the production of byssus threads in Mytilus edulis. KAREN JANE GLAUS.

Measurements were made of the pull required to remove *Mytilus edulis* from the rock substrate at three sites on Sippewissett Beach and from rock crevices in Sandwich jetty, Cape Cod, Massachusetts. At the beach, an average pull of 1.7 kg. removed animals from a site just above the mean low water mark. One kg. removed animals from a site half way up the shore and 0.5 kg. removed animals from the high shore. The pull required at the jetty was 1.0 kg. There is no significant difference between the holding power of the animals in the middle of the bed and at the jetty. All other areas differed significantly from each other.

The mean breaking modulus for byssus threads is 10×10^9 dynes/cm.² It seems logical to assume that the holding power of a mussel is related to the number of threads produced.

In the laboratory, animals were placed in slack (0.002 liters/sec.), medium (0.02 liters/sec.) and fast (0.2 liters/sec.) running sea water. The average numbers of threads produced in 12 hours were 10.6, 15.0 and 16.6, respectively.

Large (4-5 cm. in length) and small (2-3 cm.) mussels were subjected to combinations of 3 salinities and 3 temperatures for 24 hours. The salinities used were 15.9, 31.3 and 46.3‰. Temperatures were 18, 23 and 28° C. Small animals produced more threads than large animals. Large animals produced the greatest number of threads at a salinity of 31.3 under all temperature conditions. At 31.3‰, the number of threads increased as temperature increased.

Small animals produced more threads at 31.3‰ than at the lower and higher salinities. At 31.3 and 15.9 the number increased with increased temperature. At 28° C. the number of threads produced at the high salinity was significantly less than the number at 31.3. At 18° C., the number produced at the low salinity was significantly less than that at 31.3‰.

Electron microscopy of early cleavage stages in Arbacia punctulata. DANIEL A. GOODENOUGH, SUSUMU ITO AND JEAN-PAUL REVEL.

Various stages during the early development of *Arbacia punctulata* eggs were studied in the electron microscope after fixation in cacodylate-buffered Karnovsky's fixative, modified by the addition of an equal volume of sea water and trinitroresorcinol or trinitrocresol to a final concentration of 0.01%. Examination of mitotic figures shows that the chromosomes at metaphase have no particular relationship to the numerous cisternae and tubules of the endoplasmic reticulum found in the mitotic apparatus. During anaphase, however, membranous elements become closely applied to and eventually completely invest the chromosomes. At early telophase, when the chromosomes lose their condensed appearance, they resemble tiny nuclei (chromosome vesicles), each surrounded by an envelope with numerous nuclear pores and containing a nucleolus-like body.

Examination of the periphery of dividing cells at the onset of cytokinesis reveals the presence of numerous parallel filaments, less than 100 Å in diameter, which are intimately associated with the inner face of the plasma membrane in the region of the early cleavage furrow. During the first cleavage these filaments form a belt about 0.1 μ thick and 10–12 μ wide around the equator of the zygote. This belt remains associated with the leading edge and the lateral walls of the furrow as it invaginates, but appears to maintain its original width and thickness during cleavage. In later divisions a narrower filamentous belt is associated with the plasma membrane in the cleavage furrow region.

As cytokinesis proceeds, numerous folds project from the wall of the furrow into the newly formed intercellular cleft. Folds from opposite sides may make contact with each other. At the end of cytokinesis the intercellular space between the newly formed blastomeres becomes largely reduced to a narrow zone between elaborate interdigitations of the cells. In various regions the intercellular space becomes altogether obliterated, suggesting the formation of actual intercellular junctions between the blastomeres.

Supported by grants from the U. S. Public Health Service.

Pharmacological studies on electromechanical coupling in a phasic molluscan muscle.

MICHAEL J. GREENBERG, ROBERT B. HILL, HIROSHI IRISAWA AND HIROMICHI NOMURA.

The radula protractor of *Busycon canaliculatum* is a twitch muscle; each stimulus to the nerve results in a single phasic contraction. Tetanus tension is regulated both by the frequency of muscle action potentials, and by the level of membrane depolarization. Membrane potentials of the isolated muscles were measured in the sucrose gap apparatus simultaneously with tension. While low potassium solutions hyperpolarized with relaxation, high potassium solutions depolarized with contracture. Resting potentials were 50–55 mV. Acetylcholine (ACh) solutions also depolarized the muscle and produced contracture (threshold, $3 \times 10^{-7} M$). Both electrical and mechanical effects increased with ACh concentration to $10^{-8} M$.

Tryptamine ($10^{-9} M$), alone, produced only about a 4 mV. depolarization together with a just measureable tension increase. However, in the presence of a 75 mM KCl contracture, tryptamine induced further tension development but without affecting the depolarization. Tryptamine could not enhance the large contracture resulting from 150 mM KCl. Excess calcium, added during a 75 mM KCl contracture, did not affect depolarization but induced a small transient tension increase. Caffeine (10^{-3} g./ml.) had a similar though larger effect. Prior soaking in 18 mM calcium for 10 minutes greatly augmented an ensuing KCl contracture. Therefore, tryptamine may increase the tension accompanying KCl depolarization by accelerating the availability of calcium as an activator.

Added during a contracture induced by $7 \times 10^{-7} M$ ACh, $10^{-8} M$ tryptamine reduced the depolarization and relaxed tension, although ACh remained in the test solution. Tryptamine had the same qualitative effect at higher ACh levels ($10^{-5} M$), but the relaxation was less rapid. However, the characteristic oscillations of membrane potential and tension were more marked and prolonged during these slower relaxations. This phenomenon suggests a model for the origin of rhythmicity through the oscillation of a membrane made unstable by nicely balanced concentrations of opposing neurohumors.

*The morphology of a mechanoreceptor organ of the bivalve, *Spisula solidissima*.*

MARY ANN HART.

The receptor ganglion with four sensory regions is suspended between the cardinal teeth of the right valve. In addition, a small anterior sensory region and a small posterior sensory region are associated with the receptor organ. A small wedge-shaped tooth with a fragile laminar process from the left valve interdigitates with the right valve cardinal teeth and stimulates the four sensory regions of the ganglion complex and the small posterior region. A lateral tooth from the right valve stimulates the small anterior sensory region.

Application of mechanical stimuli to the sensory regions of the intact animal results in foot withdrawal. The receptor regions are phasic in their responses to single pulses and to the onset and cessation of continuously maintained pressure from the stimulating probe.

Supported by Grant NB 04989-05 from USPHS to Dr. Deforest Mellon.

Determination of rate of polypeptide chain assembly in vivo in Opsanus tau.

AUDREY E. V. HASCHEMEYER AND JANICE C. LEVY.

Previous studies have indicated that cold 10°-acclimated toadfish have a greater liver protein synthetic capacity than control (21–22°) fish, when measured at the same temperature. It was also found that aminoacyl transferase activity is elevated in livers of cold acclimated fish. The transferases catalyze steps involved in the binding of aminoacyl-tRNA to ribosomes and its addition to the growing polypeptide chain, and thus can control the rate of chain assembly. In order to determine if this rate differs in cold-acclimated and control fish, an experimental procedure was designed to measure the rate of addition of radioactive amino acids on the ribosomes and the appearance of radioactivity in completed soluble protein. Toadfish were anesthetized and strapped to a fish rack with circulating oxygenated sea water passing over the gills. Five μ c. of a mixture of C¹⁴ amino acids were injected into the hepatic portal vein. After the desired incubation time, from 1 to 8 minutes, the needle was removed, and the liver was excised, homogenized, and fractionated by differential centrifugation. Incorporation of radioactivity into protein was determined in the total homogenate and in various fractions, including the ribosomal pellets and the ribosome-free supernatant. Ratios of these quantities when plotted *vs.* time gave values of t_c , the time of assembly of an average polypeptide chain in liver, independent of the usual individual animal variations. For control fish, t_c was about 6.7 minutes at 22°; with the assumption that an average protein chain in liver is about 50,000 molecular weight, the result compares favorably with data of Dintzis on hemoglobin chain assembly (1.5 minutes per 16,000 MW) in reticulocytes and that of Loftfield and Eigner on ferritin production (6.5 min. per molecule) in rat liver. A faster rate of chain assembly at 22° was found in livers of 10°-acclimated fish.

Calcium activated bioluminescent proteins from ctenophores (Mnemiopsis) and colonial hydroids (Obelia). J. WOODLAND HASTINGS AND JAMES G. MORIN.

The biochemical system responsible for the bioluminescent flashing in *Mnemiopsis leidyi* and three species of *Obelia* (*bicuspidata*, *geniculata* and *longissima*) has been isolated and partially characterized. In each case the system involves a calcium activated protein similar to that isolated by Shimomura *et al.* from certain hydromedusae. In our experiments the active proteins were isolated by extraction in 0.04M EDTA in 0.2M Tris at pH 10. After dialysis these extracts are stable for at least several weeks. Bioluminescence can be evoked simply by the addition of excess calcium. The light emission occurs as a flash having a duration of about one second. The identity of the energy-yielding chemical step is obscure. The reaction does not utilize or depend upon the presence of molecular oxygen. Only a single component in the extracts is involved. Its molecular weight was estimated to be about 18,300 by gel filtration, this being identical with that of a sample of the active protein from the hydromedusan *Aequorea aequorea*. The material isolated from *Mnemiopsis* is similar in its response to calcium and in its molecular weight, but it has the unusual property of being rapidly inactivated by exposure to light. This feature could account for the fact that the yield of activity in extracts is greatly decreased after exposure of the living organism to light. Such is not the case in *Obelia* even though light may inhibit the *in vivo* bioluminescent flashing.

Microtubules in the axial filament complexes of acoel turbellarian spermatozoa, as revealed by negative staining. CATHERINE HENLEY, DONALD P. COSTELLO AND CHARLES R. AULT.

The mature spermatozoa of both *Childia groenlandica* and *Polychoerus carmelensis* have an anterior nucleus followed by 2 parallel rows of refractile bodies bounded on either side by an undulating membrane. The apparent differences between the two are in their differing lengths and widths: 52 × 4–5 microns for *Childia*, 400 × 1–2 microns for *Polychoerus*. Under certain specific conditions, at least, both spermatozoa are highly motile, progressing by waves passing along each of the two undulating membranes from the tip of the tail towards the head.

As seen in cross-section by electron microscopy, both have two incorporated axial filament

complexes, each about 0.18 micron in diameter and with 9 doublet microtubules arranged in a circle; these complexes lack the central pair of the 9+2 pattern. This offers evidence against an invariable correlation between motility and the presence of the central pair of microtubules. The cilia on the surfaces of both animals *do* have the 9+2 arrangement.

Late spermatids or spermatozoa of *Polychoerus* were negatively stained with 1% phosphotungstic acid and examined with the Zeiss electron microscope. Large bundles of microtubules were found, containing both doublets and singlets which were bound together at the posterior end by a cap-like structure with definite periodicity. A total of 18 doublets and varying numbers of singlets could be counted. The diameters of the doublets were ± 400 Å, those of the singlets ± 200 Å, with periodicity in the walls of both of ± 75 Å.

In spermatozoa of *Childia* similarly negatively stained and examined by EM, a large number of intact individual axial filament complexes was observed, each with 9 doublet microtubules, all 9 being attached to what is apparently the basal plate of the blepharoplast. The doublets had a repeating substructure in their walls of ca. 150 Å period, and were about 400 Å in diameter. No intact axial filament complexes were observed with more than 9 attached doublets.

Thus far, the posterior cap-like structure binding the microtubules of the axial filament complexes together in *Polychoerus* sperm has not been found in spermatozoa of *Childia*, but further work is continuing, with investigations of negatively-stained microtubules in spermatozoa of other forms, including *Chaetopterus*.

Aided by a grant from the National Institutes of Health, GM-15311. We are indebted to Mr. Heinz Rische, of Carl Zeiss, Inc., for assistance with some of the electron micrographs. The work of C.R.A. is supported by a NASA predoctoral fellowship.

The effect of electrical stimulation on the action of sulfhydryl reagents in axonal preparations. GILBERT R. HILLMAN AND HENRY G. MAUTNER.

The effects of N-ethylmaleimide (NEM) and parachloromercuribenzoate (PCMB), reagents capable of reacting with sulfhydryl groups, were studied in giant axons of squid, *Loligo pealei*, and in nerve bundles from the walking leg of lobsters, *Homarus americanus*. In order to avoid damaging the nerve membranes, the reagents were applied externally, and the nerve responses were followed using external electrodes.

PCMB and NEM both blocked the nerve response. The action of NEM was irreversible; the effect of PCMB could be reversed completely with 2-mercaptoethanol. NEM was used at 5 and 10 mM, PCMB at 0.05 mM, and 2-mercaptoethanol at 10 mM.

The effects of both reagents were potentiated strongly by electrical stimulation. In the squid axon, treatment with 10 mM NEM for ten minutes gave 30% reduction of the size of the action potential; the application of one threshold stimulus per second for thirty seconds resulted in 100% block. Similar effects were seen in lobster nerves at 5 mM NEM.

The effects of PCMB also were potentiated by stimulation, both in the lobster and in the squid. Using a small stimulus voltage with the lobster nerve, it was possible to stimulate just a few fibers in the nerve bundle. After 15 minutes of treatment with PCMB all fibers were 50%–70% inhibited. Selective stimulation of these relatively sensitive fibers 6 times a second for 2 minutes gave complete block only of these fibers; the remaining fibers stayed at a constant level of activity until the stimulating voltage was raised above their threshold for another 2 minutes.

These findings are interpreted to indicate that there is a membrane biopolymer containing sulfhydryl groups which are made available for chemical attack only during membrane excitation. It is possible that the sulfhydryl reagents act at more than one site, with some sites being insensitive to stimulation effects.

This work was supported by a National Science Foundation grant GB 4114.

Interstitial marine gastrotrichs from Woods Hole, Massachusetts. Part II. WILLIAM D. HUMMON.

A preliminary study of the gastrotrichs from Woods Hole was reported a year ago in this journal. Since then, a one-year ecological study has been completed on Crane's Beach,

located at the proximal end of Penzance Point, facing Buzzards Bay. Now included in the list of known gastrotrichs from this beach are all seven species reported last year, as well as three additional species: *Xenotrichula pygmaea* Remane, 1934, *Diplodasys ankei* Wilke, 1954, and *Macrodasys caudatus* Remane, 1927. The latter three species are here reported for the first time from North America, each having previously been listed among the fauna of both the Atlantic and Mediterranean Coasts of Europe.

Pseudostomella roscovita, noted last year from Nobska and MBL Beaches, has subsequently been found to inhabit a comparable position on Crane's Beach, where it co-occurs regularly with *Tetranchyroderma papū*. *Turbanella cornuta* has also been observed on Crane's Beach since last August, but occupying a restricted position at 5-10 cm. depth, low-tide elevation. Of the newly reported species, *Xenotrichula pygmaea* regularly inhabits the 5-40-cm. depths at all tide levels, while both *Diplodasys ankei* and *Macrodasys caudatus* occur only rarely, at 25-40-cm. depths, mid-tide elevations. Tidal elevations in both reports refer to mean high- and low-tide levels, and occurrences refer only to summer observations.

In addition to the ten species indicated above, another five, new or as yet unidentified, have been observed in samples taken from Crane's Beach. A total of 15 species thus have been found in a beach measuring only 15 m. in width from high- to low-tide levels. Data on seasonal patterns and changes in density, dispersion and co-occurrence have formed a major part of the Crane's Beach study. They will be correlated with patterns and changes in physical niche parameters in a future publication.

Contribution No. 165 from the Systematics-Ecology Program, Marine Biological Laboratory, Woods Hole, Massachusetts.

Supported by an NDEA Fellowship, University of Massachusetts, Amherst, Massachusetts.

Migration of germ cells through successive generations of pallial buds in Botryllus schlosseri. C. S. IZZARD.

Berrill (1941) states that in *Botryllus* germ cells originate in the atrial epithelium of the vesicle-stage bud and segregate from the epithelium into the adjacent mesenchyme space underlying the epidermis. In previous light and electron microscopical studies I could find no cytologically differentiated cells in the epithelium nor evidence of cellular segregation from the epithelium. The study has been extended on living sexually-mature colonies and sectioned material.

Germ cells first appear in the vesicle bud, but they are relatively large oocytes (20-40 μ diameter) in a previtellogenic phase of growth. Oocytes in meiotic prophase do not appear until 2 days later following the formation of atrial folds, and gut and pericardial rudiments.

The associated parental bud contains many previtellogenic growth-phase oocytes which, in living buds, prior to the appearance of oocytes in the vesicle bud, are washed from the gonad area into the blood. For approximately 12 hours, large numbers of these oocytes circulate throughout the adults, buds, test vessels and ampullae.

Shortly before the release of oocytes into the blood, a circulatory loop from the parental blood-circulation develops through the vesicle bud. Over a period of 24 hours, previtellogenic growth-phase oocytes passing through the loop settle on either side of the vesicle in the mesenchyme space immediately above the loop. These oocytes mature in this bud generation. The precocious appearance of germ cells and the absence of early oocyte stages in the vesicle bud are thus explained.

From their distribution in sectioned material, one can infer that a population of small (8 μ diameter), mitotic (hence not oocytic) cells, cytologically similar to early spermatogonia, also migrate from the parental gonad area to the vesicle bud. Their appearance prior to that of the testes and meiotic prophase oocytes supports the notion that they represent a primordial germ cell stock.

Supported by NIH grant GM 14891.

Aspects of carbohydrate metabolism in Crustacea. MICHAEL A. JOHNSTON AND FRANK M. FISHER, JR.

Hormonal regulation of carbohydrate metabolism by the decapod eyestalk has been investigated in two crustaceans, the fresh-water crayfish *Cambarus* sp. and the marine spider crab

Libinia emarginata. Basal concentrations of hemolymph and hepatopancreatic sugars were determined for normal animals and for those subjected to eyestalk ablation. Reduction in basal levels is noted for several sugars. In the hemolymph the decline from intact to ablated animals is 36% for glucose, 52% for reducing compounds, and 68% for total carbohydrates. Hepatopancreatic levels show similar declines for ablatants: for glucose, 44%; for reducing compounds, 24%; and for polysaccharides, 64.5%. The data suggest an influence on basal sugar concentration by an elaboration from the eyestalk.

Water balance is also affected. Normal *Libinia* show blood volumes of 0.379 ± 0.04 ml. of fluid per gram body weight; the protein content is 0.53 g./ml. of blood. Ablated animals show correspondingly larger blood volumes (0.53 ± 0.33 ml./g.) and a smaller protein content (0.33 g./ml.). Dilution resulting from ablation is clear, but such does not completely account for decreases in sugar levels.

Remarkable differences are noted in a comparison of hemolymph and hepatopancreas polysaccharide content. In the hemolymph, freely-circulating polysaccharides constitute a high 78.5% of total sugar concentrations. In the hepatopancreas, the polysaccharides are a small 6.1% of the total. Acid hydrolysis of hemolymph polysaccharides recovers 60% as glucose, suggesting that the blood rather than the hepatopancreas may be the important sugar storage site in this species.

Cambarus and *Libinia* were injected with exogenous (100-1000 μ g.) glucose. Blood samples were removed hourly for five hours and assayed for glucose. Expected glucose-tolerance curves are noted with no significant differences apparent between intact and ablated individuals. The identity of curves indicates that eyestalk hormonal elaborations have no remarkable influence over dispersion of sugar within the blood nor over transport of carbohydrates out of the circulation.

Supported by a grant GB-3447 from the National Science Foundation to the Department of Invertebrate Zoology, Marine Biological Laboratory.

The biological role of Bugula-type avicularia. KARL W. KAUFMANN, JR.

Very little work has been done to determine the biological role of the avicularia of ectopods. The most widely accepted theory is that they are used for defense. However, no one has presented sufficient evidence to demonstrate what the avicularia might specifically defend against. *Bugula simplex* and *Bugula* sp. were intensively studied from a morphological and ecological viewpoint to determine (a) what the avicularia of the two species are best suited to do, and (b) what biological role they actually perform. Structural studies of the musculature, size, strength, and orientation on the colony showed that the avicularia were best suited for grabbing animals with many small appendages and which were in the size range of 1 to 3 mm. Ecological studies of the two species showed that the gammarid amphipods *Corophium insidiosum* and *Jassa falcata* did extensive harm to the colonies by building tubes in them, were of the proper size and shape to be easily caught by the avicularia, and were hindered in their activities by the avicularia. Thus, the primary biological role of the avicularia of *Bugula simplex* and *Bugula* sp. was found to be that of defense against these two amphipods. The widespread distribution of both *Bugula*-type avicularia and tube-building organisms similar to these amphipods makes it likely that this is not a unique association.

The work was carried out under NSF grant GB-7325 to T. J. M. Schopf.

Crustacean pigmentary-effector isohormones. L. H. KLEINHOLZ.

Chromatography of acetic acid extracts of lyophilized eyestalks on G-25 Sephadex columns resolves two zones of pigmentary hormone activity; region 1B contains distal retinal pigment (DRPH) and brachyuran melanophore (MDH) activators, while Peak 4 contains the erythrochrome hormone. When Sephadex Peak 1B material is eluted from DEAE cellulose columns with collidine-acetate buffers, followed by 0.1 M HCl, 6 peaks are revealed after alkaline hydrolysis and ninhydrin reaction. The first two of these peaks, conveniently called "alpha-DRPH" and "beta-DRPH," are active. Assay injections of alpha-DRPH from 8 different preparations, in doses ranging from 0.28 μ g. to 1.5 μ g., result in average DRP indices between 0.150 and 0.233. Tests with beta-DRPH from the same preparations in 0.45 μ g. to 2.3 μ g. doses give DRP indices between 0.081 and 0.190. An index of about 0.200 characterizes the light-

adapted retina, while that for the dark-adapted retina is about 0.050. Alpha-DRPH is resolvable into two peaks (alpha-1-DRPH and alpha-2-DRPH) of activity by gradient elution with acetate buffer from CM cellulose columns. DRP indices of 0.142 and 0.174 result from doses of 3 μg . (alpha-1-DRPH) and 0.39 μg . (alpha-2-DRPH), respectively. Beta-DRPH residues, similarly chromatographed, are separable into several peaks, only the first of which is active. MDH, which disperses the pigment of brachyuran melanophores, consistently accompanies DRPH activity, and appears in the same peaks. Electrophoresis and chromatography of alpha-1-DRPH and alpha-2-DRPH on thin layer silica gel show only one spot reacting to chlorine-tolidine reagent; the melanophore and retinal pigment activities may be due to the same substance.

Supported by grant NB 02606 from the National Institutes of Health.

Neurophysiology of oculomotor neurons in teleosts. M. E. KRIEBEL, S. G. WAXMAN, G. D. PAPPAS AND M. V. L. BENNETT.

Oculomotor neurons in several teleosts were studied by intracellular recording. Motoneurons were identified by antidromic stimulation of nerves to extraocular muscles. Electrotonic spread could often be measured between pairs of neighboring cells. Short latency (<0.3 msec.) graded depolarizations were evoked in many cells by graded antidromic stimulation subthreshold for the penetrated cell. These depolarizations presumably resulted from electrotonic spread of antidromic spikes in nearby cells. Depolarizations were not evoked by stimulating heteronymous muscle nerves even where nuclei were close together. Thus, motoneurons innervating different muscles are not coupled. Interneuronal fibers were identified by orthodromic firing without PSP's and absence of spikes in response to antidromic stimulation. These fibers often showed short latency graded depolarizations upon antidromic stimulation of oculomotor nerves indicating electrotonic coupling to motoneurons. Electron microscopy demonstrated "tight junctions" at some synapses on motoneurons. Motoneurons are probably electrotonically coupled by way of presynaptic fibers which synapse on a number of cells. Most motoneuron spikes evoked by eighth nerve stimulation arose sharply from a nearly level baseline and there was little subthreshold PSP, suggesting that the spikes arose from highly synchronous and rapidly rising PSP's or propagated in from dendrites. Spikes evoked by ophthalmic nerve stimulation arose from large slowly rising PSP's, although the apparent firing level was not always constant. Brief, subthreshold spike-like potentials were observed in many cells. Some were due to electrotonic spread of spikes from neighboring cells, since they were associated with an impulse in the efferent nerve and were blocked by antidromic stimulation. Others were not blocked by antidromic stimulation and were probably due to spikes in dendrites or to PSP's. Electrotonic coupling of motoneurons is probably important in mediating the synchronous efferent volleys that can be evoked by orthodromic stimulation. Different modes of impulse initiation may operate in fast and slow eye movements.

Electron microscopy of the excretory vesicle in the cercaria of the trematode, Cryptocotyle lingua. PAUL L. KRUPA, GILLES H. COUSINEAU AND ARYA K. BAL.

In the cercaria of the parasite, *Cryptocotyle lingua*, the excretory vesicle or bladder is a large, saccate organ, located at the caudal end of the body near the junction of the tail. In thin-sectioned specimens fixed in glutaraldehyde and osmium the epithelium of the vesicle appears as a multinucleated syncytium containing numerous secretory granules or droplets, which are reminiscent of zymogen granules. The membrane-bounded granules are circular or ellipsoid in profile and contain a dense amorphous material. Some of the granules, in groups of 4 or 5, tend to be aligned perpendicular to the luminal surface of the vesicle. The number of granules increases as the parasites develop; few granules are found in intrarectal stages of cercariae, but in free-swimming cercariae the epithelium of the vesicle is charged with vast clusters of secretory granules, with their enveloping membranes in close apposition to each other. In early embryological stages of cercariae the rough endoplasmic reticulum contains irregularly-shaped saccules, but in free-swimming cercariae the cisternae of the endoplasmic reticulum are arranged in numerous tightly-packed arrays. Other structures in the cytoplasm

of the vesicle are Golgi, mitochondria, and myelin-like whorls. Cytoplasmic flaps or rugae project into the lumen of the excretory vesicle and that of the collecting and accessory ducts. Membranous elements are found occasionally in the lumen of the vesicle, but secretion of the granules into the lumen does not seem to occur in intramolluscan or free-swimming cercariae.

The trematode protonephridial system, usually consisting of flame cells, ducts, a vesicle or bladder, and a pore opening externally, has long been assumed to be excretory and osmoregulatory in function on the basis of little or no direct evidence. The presumed role of the excretory vesicle has been the temporary storage of excretory wastes carried to it by ciliated cells and ducts of the excretory tree. On morphological grounds presented in this study, however, the secretory nature of the excretory vesicle epithelium may have some importance in the successful completion of this parasite's life cycle.

This work was supported in part by The National Research Council of Canada (A-3624), the Damon Runyon Memorial Fund for Cancer Research (DRG, 918-AT) and The City College of New York Research Foundation.

Functionally excitable area of shrimp medullated giant fiber. KIYOSHI KUSANO AND MATTHEW M. LAVAIL, III.

External potential field during impulse conduction along the functionally isolated medullated giant fiber of shrimp (*Penaeus setiferus*), showing a conduction velocity greater than 90 m./sec. at 24° C. without nodes of Ranvier, was recorded with a 1 M NaCl glass capillary electrode. Recordings from most parts of the myelin-covered giant fiber surface showed no action potentials or only small monophasic positive potentials. However, at the points of giant fiber-motor fiber synapses and where giant fiber axons branch into abdominal ganglia, negativity of up to 1 mV. was recorded. By surrounding a single area of synapses or ganglia with a Vaseline partition, a positive-negative or positive-negative-positive potential could be recorded. When physiological saline in the partition was replaced with Na-free, isotonic KCl, or 10⁻⁷ g./ml. tetrodotoxin-containing saline, the negativity was abolished, and impulse conduction was blocked. Application of these solutions to a myelin-covered area did not block impulse propagation. Fine structure studies of the giant fiber at the area of synapses revealed that the myelin sheath thins and is absent in some small areas. The axon sends a branch to this region. The terminal knob of this branch shows characteristic infoldings and interdigitations with a similarly infolded motor fiber axon. Similar morphology is seen at the sites of contact between the giant fiber axon and the ganglion. Thus functionally excitable areas are present which appear to act similarly to nodes of Ranvier. They make possible impulse conduction at high velocity in shrimp medullated giant fiber. The myelin sheath is very resistive to depressants of excitation and also has a high electrical resistance. Impulse propagation therefore appears to be saltatory.

Supported by PHS Research Grant NB-06968.

*Maltase activity in developing embryos and adult tissues of the surf clam, *Spisula solidissima*.* LEONARD LASTER AND PHILIPP STRITTMATTER.

As part of a general examination of the association of enzyme activities with the appearance of organ systems during embryonic development of the surf clam, *Spisula solidissima*, in past years we assayed various oxidative and hydrolytic enzymes in embryos and adult tissues. Many of these activities did not increase significantly or changed only slightly. Because the surf clam utilizes carbohydrates as nutriment, in the present study we examined properties of the maltase activity in tissues of the adult clam and determined the appearance of this enzyme activity during the first three days of embryogenesis.

Maltase activity of adult clams is concentrated in the digestive tissues, particularly in the digestive gland. Thus, maltase specific activity in homogenates of digestive gland is approximately 100-fold greater than in homogenates of foot muscle. Unfertilized eggs and early embryos (during the first 24 hours after fertilization) are comparatively low in maltase activity. During the second day of embryonic development there is a rapid, 10-fold, increase in maltase specific activity. This increase coincides temporally with the reported appearance of the esophagus, stomach, intestine, and digestive gland during embryogenesis in this animal. Maltase activity may, therefore, serve as one biochemical marker for gastrointestinal organogenesis. In addition, our observation that digestive gland maltase activity is predominantly associated with

intracellular particles suggests that its increase during development of embryos may reflect functional differentiation of a particular organelle in cells of the digestive gland.

We are now isolating the enzyme activity for further characterization of its properties with a view toward utilizing it to study regulatory mechanisms underlying its rapid rise during the second day of development.

Composition and formation of insoluble protein in the dogfish (Mustelus canis) lens. SIDNEY LERMAN, JOAN TUTTLE AND RICHARD KOSER.

The concentration of insoluble protein (albuminoid) in the ocular lens serves as an excellent aging parameter in all species, including the dogfish. Initially, the lens is mainly composed of the three soluble protein fractions (alpha, beta and gamma crystallin) and water (60-70%), but there is a progressive increase in the insoluble protein fraction with aging. The amino acid composition of dogfish lens albuminoid is very similar to that of the gamma crystallin fraction. The tyrosine content of these two fractions is particularly significant in that it is almost twice that of alpha and beta crystallin, resulting in E_{1}^{1} values of 22-25 for gamma crystallin and albuminoid in contrast with significantly lower values in the other two crystallins. The SH and SS level of gamma crystallin derived from the mature dogfish lens is identical with the corresponding levels in albuminoid, and electrophoresis of solubilized albuminoid (by means of sulfonation) and gamma crystallin on thin layer Pevikon plates show similar R_f values. Precipitin reactions utilizing antibodies to all four lens protein fractions and the individual proteins provide further evidence of a close relationship between the gamma crystallin fraction and the insoluble protein of the dogfish lens. The foregoing experiments indicate that as the lens ages, the albuminoid (insoluble protein) fraction derives mainly from gamma crystallin by means of conformational changes, disulfide bridges between the gamma monomers, and certain other changes in gamma crystallin which enhance the process of polymerization and insolubilization. The latter is probably related to the presence of a fluorescent compound which becomes apparent as the lens ages and is localized to gamma crystallin and albuminoid. Fluorescence and UV spectra and chromatographic analyses indicate that this compound is a derivative of tyrosine and may be a form of bityrosine in which the phenolic groups are substituted rather than free. The fact that this fluorescent compound is present in a specific peptide derived from 6-hour tryptic digests of gamma crystallin would tend to rule out the possibility that it is a breakdown product derived from tyrosine or tryptophan.

Supported by M.R.C. grant MA 3208 and in part by a Fight for Sight Student Fellowship (Joan Tuttle).

The number of molecules involved in inhibiting either an enzyme or cell division.

ROBERT B. LOFTFIELD AND ELIZABETH ANN EIGNER.

We (*J. Biol. Chem.*, 1968) have shown that the inhibition of an enzyme can be represented by the equation; $\log(1/v - 1/v_0) = C + n \log I$, where v is the uninhibited rate of reaction; v_0 , the inhibited rate; C , a collection of constants; I , the concentration of inhibitor; and n is a small integral number representing the number of inhibitor molecules required to deactivate the enzyme. The value of n is easily determined as the slope of a straight line resulting from a series of experiments in which all factors are constant except for change in I . We showed that the value of n was exactly 1 or exactly 2 for a variety of enzymic reactions, evidence that precisely one or precisely two molecules of inhibitor are required to inactivate alcohol dehydrogenase or amino acid activating enzymes.

We wondered whether much more complex biological reactions can be examined by the same technique. When the accumulated work of Clowes (*Ann. N. Y. Acad. Sci.*, 1951) on the phenol inhibition of cell division of *Arbacia punctulata* is analyzed in this fashion, we find n has the exact value of 1.0 for *p*-aminophenol and dinitro-carvacrol and exactly 2.0 for 2,4,6-tribromophenol, 2,4,6-triiodophenol and 2,4-dibromophenol. There is correspondingly a strong suggestion that there is a vital center involved in cell division that is inactivated by exactly 1.0 or 2.0 molecules of the above phenols. On the other hand, some metabolic poisons such as iodoacetic acid react with a number of amino acids of widely varying function in enzyme systems of widely varying importance. Such poisons would not be expected to yield linear graphs because

there is no single target. When the iodoacetate inhibition of *Arbacia* division (Krahl and Clowes, *J. Gen. Phys.*, 1941) is plotted as above, the slope changes from about 1.0 to 4.0 in the range 0.2 mM to 33 mM.

Supported by USPHS Grant CA 08000.

Stabilization of axopodia by D₂O; a pressure-temperature study. DOUGLAS MARSLAND AND MICHAEL HIRSHFIELD.

Previously it has been shown by Tilney, Hiromoto and Marsland that the axopodia of *Actinosphaerium nucleofilum* become unstable, with concomitant beading and retraction, upon exposure to high pressure (4000–8000 psi). Moreover, the stability of the axopodium as a whole appears to be related to the stability of the complex system of microtubules found in the axial core.

The present work involves the use of two agencies, namely, deuteration and rising temperature. These are known to have a stabilizing influence on the structural integrity of the mitotic apparatus, presumably because of their effects upon the microtubular components.

With increasing concentrations of D₂O, a pronounced increase in resistance to axopodial beading and retraction has been observed. Thus at 20° C., the minimum pressure required to induce complete retraction rose from 5000 psi with no D₂O; to 6000 psi for 50% D₂O; 8000 psi for 60%; 10,000 psi for 70%; and 12,000 psi for 80%.

Experiments still in progress indicate that a further increase in axopodial stability also occurs when the experimental temperature is raised from 20° C. to 25° C. Thus the pressure requirement at the higher temperature has proved to be about 6000 psi in the H₂O medium; 8000 psi for 50% D₂O; and 9000 psi for 60%.

Work supported by grant ROI GM-15893-20, from the National Institutes of Health.

Differential effects of LAS detergent on reproductive performance of male and female brine shrimp Artemia. D. F. MOFFETT AND D. S. GROSCHE.

Linear alkylate sulfonate, the straight-chain biodegradable detergent which is the principal active ingredient in household washing products, was found to reduce reproductive performance of mating pairs of adult *Artemia* at concentration levels reported in polluted waters (A. G. Spencer, 1967, M. S. Thesis, N. C. State Univ.). The present experiment is analogous to the reciprocal cross method of revealing extra-chromosomal inheritance. Young adults were exposed in brine solution containing 5 parts per million of LAS for 8 hours. The time for 50% lethality was 22 hours. Treated and untreated shrimp were paired to give three groups: females treated, males treated, and neither sex treated. Reproductive performance was recorded for 30 days from the appearance of the first brood. Damage *via* the female appeared as significant depression in number of broods, number of zygotes per brood, total number of zygotes, and survival of larvae to maturity. The sex ratio of survivors was normal. Although fewer encysted zygotes were produced, their hatchability surpassed even the control values, suggested pre-deposit selection. The number of females with blocked ovisacs in which resorption of cyst-like material can be observed is consistent with this hypothesis. While the female was productive, the frequency of brood deposit was unchanged. However 6 of the 10 treated females averaged only 3.7 broods while the other 4 females averaged 9.3 broods. Preliminary studies with a continuous recording respirometer indicate that LAS can inhibit oxidative metabolism at such concentrations. This suggests that the agent causes somatic effects as well as cytosomal changes, perhaps dauer modification, in the female gonads.

Supported by NSF Undergraduate Research Participation and USPHS research grant ES-00044 Division of Environmental Engineering and Food Protection.

Excitatory physiology and localization of bioluminescence in Obelia. JAMES G. MORIN, GEORGE T. REYNOLDS AND J. WOODLAND HASTINGS.

A partial survey of the Woods Hole hydroids has revealed that both the hydroid and medusoid generations of three species of *Obelia* (*bicuspidata*, *geniculata* and *longissima*) are bioluminescent.

A study of the localization of light emission within the *Obelia geniculata* and *O. longissima* hydroid stage was achieved by photographs taken through an image intensifier. Luminescence was initiated by stimulating across the colony perisarc with fine silver-silver chloride electrodes. These studies show that the stolons, stems and pedicels are luminescent, while the hydranths and gonangia are not. The light emanates from discrete, irregularly spaced spots. Successive pictures show that the spots are in fixed positions along the coenosarc.

Recordings of luminescence from specific areas were made using micromanipulator-held fiber optics with a tip diameter of 0.5 mm. coupled to a photomultiplier recording system. The colonies were stimulated as before. Suction electrodes with tip diameters of 10–50 μ were used to record electrical activity from the hydranths during some experiments. In all three species suprathreshold stimulation at a rate of one or less per second produces a burst of flashes lasting from a few hundred milliseconds to several seconds. These bursts do not appear until after the second or third stimulus. The flashing frequency within this burst ranges between 4 and 16 per second and is highest at the beginning. Successive bursts show marked fatigue in flash number and intensity. The initial flash has a latency of 10 to 20 msec. and all flashes show rise times of 20 msec. and total durations of about 100 msec. Flashes within the burst facilitate markedly and then slowly fatigue. Conduction velocities computed from records of two fiber optics observed from different points within the colony give values between 10 and 20 cm./sec.

Simultaneous records of the electrical activity and luminescence show an accompanying non-facilitating potential of 200–500 μ V. and 75 msec. duration with each task. The constancy of spots in successive image intensifier pictures and the phenomenon of luminescent bursts from a small region of coenosarc give indirect evidence for repetitive flashing of individual spots.

Supported in part by AEC contract AT(30-1)-3406.

A continuous rate assay for transpeptidating enzymes. R. V. MYHRMAN AND L. LORAND.

Transpeptidases catalyze the nucleophilic attack by amines on activated carbonyl groups such as the γ -amide of glutamine residues. Enzymes of this type are already known to be important in the crosslinking of fibrin structures with γ -glutamyl- ϵ -lysine bonds in the coagulation of vertebrate blood. Liver transglutaminase (of unassigned biological function so far) and a clotting enzyme ("tissue coagulin"), found in lobster tail muscle, catalyze similar reactions.

To facilitate the search for transpeptidases, in general, we have developed a continuous fluorescence rate assay based on the incorporation of N-(5-aminopentyl)-5-dimethylamino-1-naphthalenesulfonamide into α -casein. With the transpeptidase catalyzed incorporation of this fluorescent lysine analog into the protein, there is a marked enhancement of the fluorescence of the naphthalene residue ($\lambda_A \sim 360$ m μ ; $\lambda_F \sim 540$ m μ), as if this chromophore were placed in a more hydrophobic environment. Since α -casein probably contains several incorporation sites, the fluorescence change may represent an overall average. It will be interesting to examine proteins of known tertiary structure and with a single glutamine residue, in place of α -casein, as acceptors.

In the case of the lobster muscle enzyme, a $K_{M, app}$ of about 20 μ M was obtained on the basis of the initial rate of enhancement of fluorescence at different concentrations of the amine; (24° C; pH 7.5; 0.04 M Tris; 1.5 mM calcium chloride; 0.53 mg./ml. α -casein, Mann).

Aided by grant HE-02212 from the National Institutes of Health and by a grant from the American Heart Association.

Receptor and generator potentials of ampullae of Lorenzini in the skate, Raja.
S. OBARA AND M. V. L. BENNETT.

Recording and stimulating electrodes were placed in ampullae while monitoring neural output. In some ampullae negative-going all-or-none spikes were evoked by cathodal stimuli. These receptor spikes usually had thresholds of ca. 5 mV., but could appear spontaneously. They reached 20–30 mV. in amplitude and lasted ca. 50 msec. In other ampullae cathodal stimuli evoked graded oscillations without a sharp threshold, and stronger stimuli might also

evoke spikes. Both types of responses were larger in ampullae than in their canals and probably were generated by receptor cells. Cathodal stimuli increased nerve discharge frequency but impulses had no fixed phase relations to receptor oscillations. Small brief positive-negative potentials could be recorded superimposed on receptor oscillations and were due to nerve impulses, because antidromic stimulation evoked similar potentials. Receptor spikes caused a burst of impulses in an innervating fiber followed by a silent period. Often receptor spikes produced a burst followed by greatly decreased activity in the whole afferent nerve indicating that receptor spikes generated by one ampulla could affect other ampullae. Generator potentials recorded intra-axonally close to ampullae had minimum latencies of 6-9 msec. and could outlast ampullary potential changes. Generator potentials due to brief stimuli recovered following superimposed antidromic spikes which were reduced in amplitude over a time course following the amplitude of the underlying generator potential. These results indicate prolonged transmitter action producing increased conductance. A tentative model accounting for the data is that conventional depolarizing activity of lumenal faces of receptor cells is evoked by cathodal stimuli and this activity depolarizes inner faces, causing transmitter release. The transmitter then depolarizes innervating fibers. Receptor oscillations represent asynchronous and perhaps graded activity of varying numbers of receptor cells arranged in parallel, whereas receptor spikes are due to synchronous and regenerative activity of the receptor cells.

Dogfish erythrocytes: activation of nucleic acid synthesis in a repressed cell.

THORU PEDERSON AND SEYMOUR GELFANT.

The presence of a nucleus in the mature erythrocytes of non-mammalian vertebrates raises questions about the nucleic acid metabolism of such cells. We have studied DNA and RNA synthesis in the mature, nucleated erythrocytes of the smooth dogfish, *Mustelus canis*, using cytological and biochemical methods. Dogfish erythrocytes are completely inactive with respect to nucleic acid and protein synthesis, and electron microscopic studies reveal no cytoplasmic organelle systems whatsoever in these cells. However, these fully repressed cells can be activated under certain physiological and experimental conditions.

Cytophotometric measurements of Feulgen-stained nuclei revealed that all mature erythrocytes contain the 2C amount of DNA; nuclei with increased DNA contents were not observed. This lack of DNA synthesis was confirmed by radioautography of cells labeled with DNA precursors. Further, no RNA or protein synthesis was detected in mature erythrocytes by radioautography. These findings indicate that, under normal conditions, the dogfish mature erythrocyte is a completely repressed cell.

Dilution of whole blood samples with either autologous plasma or elasmobranch Ringer's resulted in an activation of DNA and RNA synthesis in mature erythrocytes, as measured both by total isotope incorporation and by radioautography. Dilution experiments were also performed *in vivo*, in which large quantities of blood were withdrawn and replaced immediately with equal volumes of sterile Ringer's. In these experiments an activation of erythrocyte nucleic acid synthesis was also observed.

Following the removal of about 25% of the total blood volume from dogfish, there occurred a rapid activation of DNA and RNA synthesis in circulating mature erythrocytes. This activated state persisted for at least 24 hours after blood removal.

These studies show that the repressed nucleus of the dogfish mature erythrocyte can be activated, and that such an activation may be an important physiological adjustment during blood regeneration.

Supported by research grant GB-2803 from the National Science Foundation.

Effects of an ecdysone on growth and cuticle formation of Drosophila imaginal discs cultured in vivo. JOHN H. POSTLETHWAIT AND H. A. SCHNEIDERMAN.

When the imaginal discs of *Drosophila melanogaster* are implanted into larvae, they metamorphose when the larvae metamorphose. When they are implanted into adult abdomens, the discs may grow, but they do not metamorphose. The present experiments were designed to see whether injection of an ecdysone into an adult fly will cause implanted imaginal discs to metamorphose.

Whole leg discs from mature third instar larvae were injected into fertilized adult females. The hosts then received single or repeated injections of ecdysterone (=20-hydroxyecdysone) in 10% ethanol in Ringer. In a typical experiment, one group of flies bearing implants received either 7.2 or 720 μg . of ecdysterone/g. fly weight in a single dose, or in six equal installments over a period of eleven days. In none of the singly injected flies did the discs grow significantly or metamorphose. In contrast, multiply treated implants increased in size more than three fold, and most secreted some cuticle, but failed to metamorphose.

To cause metamorphosis, repeated doses of higher concentrations of ecdysterone were necessary. Thus 3600 μg ./g. given over an eleven-day period in three injections caused metamorphosis in thirteen implants. These metamorphosed implants were completely covered with cuticle, and formed bristles, claws, tibial sensory organs, sex combs, and sensilla trichodea.

These results indicate the following: 1) lack of ecdysone in uninjected adult flies accounts for the absence of metamorphosis in implanted discs. 2) Ecdysterone is inactivated rapidly in the adult. 3) To cause either growth or metamorphosis, ecdysone is needed as a sustained stimulant, not merely as a trigger (hence the effectiveness of repeated doses). 4) Low concentrations of ecdysone stimulate the enlargement of discs, whereas, 5) high concentrations stimulate cuticle secretion and metamorphosis.

This discovery of a simple chemical method of regulating the growth and metamorphosis of imaginal discs promises to simplify developmental studies with *Drosophila*.

A comparative analysis of the control of the white chromatophores in the fiddler crab, Uca pugilator, from Woods Hole, Massachusetts, and Panacea, Florida.

K. RANGO RAO AND MILTON FINGERMAN.

Saline extracts of central nervous tissues of *Uca pugilator* from the Woods Hole area were known to evoke white pigment concentration, but it was not known whether they also contained a white pigment-dispersing substance. Subsequent investigations of *Uca pugilator* from Panacea revealed that saline extracts of optic ganglia, sinus glands, supraesophageal ganglia, and thoracic ganglia evoked white pigment dispersion alone. However, acetone fractionation revealed the presence of white pigment-dispersing and -concentrating substances in each of these tissues. To learn whether the corresponding tissues from Woods Hole crabs also have both substances, the tissues were extracted in acetone. In each instance the acetone-soluble fraction contained the white pigment-concentrating substance whereas the acetone-insoluble fraction contained the dispersing substance. However, the circumesophageal connectives of the crabs from Panacea and Woods Hole contain the white pigment-concentrating substance alone. Additional studies were performed with Panacea crabs. Saline extracts and the ethanol-soluble fraction of the eyestalks evoked nearly identical white pigment-dispersing activity, but had no white pigment-concentrating activity. The question arose then whether the white pigment-concentrating substance is absent in these extracts or masked by the antagonistic action of the white pigment-dispersing substance. Gel filtration of the ethanol extracts on a column of Sephadex LH-20 in ethanol revealed the presence of both substances. Exposure of the ethanol-soluble fraction of the eyestalks to heat or isopropyl ether resulted in destruction of the white pigment-dispersing substance. Although the white pigment-dispersing substance in lyophilized aqueous extracts of the eyestalk was thermostable, it was almost completely destroyed by exposure to isopropyl ether. Such treated extracts evoked white pigment concentration, thereby revealing that the white pigment-concentrating substance was present in the aqueous and alcoholic extracts but was masked by its antagonist, the white pigment-dispersing substance.

Supported by Grant GB-7595X from the NSF.

Cell recognition in an echinoderm, Asterias vulgaris. CAROL L. REINISCH AND FREDERIK B. BANG.

Echinoderms, particularly asteroids, are ideal animals for study of cell interaction since they afford easy access to the coelomic cavity for continuous sampling, and clear microscopic observation of the papillae (and hence circulating cells) under the dissecting microscope.

Asterias vulgaris has an average circulating amoebocyte count of 10^6 cells/ml. of coelomic fluid. Amoebocytes of *Arbacia punctulata* were used as a challenge inoculum. Because of their pigmentation, interaction between *Arbacia* and *Asterias* cells was readily observed. After the initial *Arbacia* inoculum, periodic samples for counting were withdrawn from the *Asterias* in warmed N-ethyl maleimide (NEM) which prevents invertebrate cells from sticking to one another.

Initial clearance of *Arbacia* amoebocytes as unbound or non-amoebocyte-associated cells occurred within 120 minutes in 13/16 experiments. Characteristic clumping, in response to the challenge *Arbacia* cells, was seen in the papillae. There was a sharp initial drop in the number of circulating free host amoebocytes which occurred within 1 minute after injection. The clumping, when correlated with the disappearance of free *Arbacia* cells, was interpreted as rapid cell recognition of foreign cellular material.

To see whether NEM might alter cell recognition *in vivo*, two sets of experiments were done. In one set, sea stars were pretreated with 2×10^{-4} M NEM before inoculation with *Arbacia* cells; in the second, *Arbacia* cells were treated with NEM before inoculation. Neither pretreating the host animal nor challenge cells altered normal host cell recognition.

An aspect of NEM's effect was alteration of amoebocyte behavior on glass. Normally these cells sent out pseudopodia within seconds after contact with a glass surface. Both viable amoebocytes from an NEM-treated *Asterias* and amoebocytes which had phagocytosed an *Arbacia* cell remained rounded up to $1\frac{1}{2}$ hours. Normal cells or cells which had phagocytosed a carborundum particle, in comparison, spread normally.

Electron microscopical studies on "myosin" from molluscan smooth muscle. ROBERT V. RICE, G. F. ELLIOTT AND A. STRACHER.

Standard methods requiring several days for preparing purified myosin from striated muscle failed to give recognizable particles when applied to a variety of molluscan smooth muscles. The preparations dissolved in volatile buffers of ammonium acetate were sprayed or streaked onto mica, shadowed with PT and coated with carbon in a vacuum evaporator. These preparations when examined in the electron microscope showed only 50 Å globules in contrast to similar preparations from striated muscle which routinely show 1600 Å rods with bulbous heads 200×50 Å. Isolation of protein in the presence of the antibiotic tetracycline did not change the results. In contrast to the standard methods a new procedure, which uses sucrose-gradient centrifugation, was used. This procedure, originated by Dr. Peter Hardwicke of the Biophysics Department of King's College, London, WC2 (personal communication, to be published shortly), gives purified fractions from muscle in a matter of hours. Fractions from various levels of the sucrose-gradient centrifugation were dialyzed against 0.6 M ammonium acetate and prepared for electron microscopy as described above. Fractions were also prepared by negative staining. The 50 Å globules were found in a band about 1 cm. from the tube top, and were relatively free of other fractions including F-actin filaments and paramyosin ribbons. The pellet contained thin paramyosin ribbons (a sub-structure of the paramyosin filaments) and some F-actin. When the pellet was dissolved in 0.1 M ammonium acetate the paramyosin ribbons were studded with 50 Å globules which were generally absent if the pellet was dissolved in 0.6 M ammonium acetate. From these experiments we tentatively conclude that thick paramyosin filaments, which from x-ray diffraction are known to contain closely packed rods (1400 Å long) of troponomyosin A, also have as part of their structure, associated small (50 Å) globules. The globules are very labile and are easily detached in 0.6 M salt solutions. At the surface of the filament the small globules probably interact with F-actin and represent the "cross bridges" between thick and thin filaments in molluscan smooth muscle.

Activity level and cardioregulation of the northern sea robin, Prionotus carolinus (L.). JOHN L. ROBERTS.

Innervation of teleost hearts by parasympathetic efferent fibers has been suggested recently to be a pathway for cardioregulation only at low levels of activity. This is because the release of vagus "braking" action by bilateral vagotomy and atropine block at the heart does not result

in cardioacceleration at higher activity levels or with steady swimming at moderate velocities. A corollary of this view is that parasympathetic activity is nearly always turned off and therefore of no significance to active species that characteristically are continuous or nearly continuous swimmers. Results that follow bear out these contentions with regard to the bottom-dwelling, inactive sea robin, *P. carolinus*.

Like most bottom fish, standard level heart rates of sea robins (about 70 cpm) as measured from ECG recordings are considerably higher than ventilation rates (5-40 cpm) obtained at 21.5° C. (visual, EMG and branchial cavity pressure recordings). Also, ventilatory pauses and rate and amplitude changes, and an associated transitory bradycardia are pronounced as a result of visual disturbances.

Thermal stress applied by warming (1° C./min.) restrained, lightly anesthetized fish (MS 222, 1/23,300) from acclimation temperatures between 20° and 22° C. until the brain (deep body temp.) reaches 27° to 28° C. results in a hypoxic bradycardia concurrent in time and degree with progressive respiratory failure. Both the visually induced and the hypoxic bradycardia are largely relieved by pericardial injection of 0.2 to 0.3 ml. atropine (10⁻³ M). Atropine injection also elevates standard level heart rates (from 69 to 107 cpm), but seems to have no effect upon cardiac rates of fish subjected to enforced moderate swimming (1 body length/sec.; 9- to 11-inch fish). In fact, heart rate accommodation seems to saturate (about 120 cpm) at a low swimming velocity (undetermined) above which no further rate change occurs.

Partially supported by a Faculty Research Grant from the Graduate School, University of Massachusetts, Amherst.

Do ploidy and cytoplasmic volume influence the cleavage rate of sea urchin eggs?

RONALD C. RUSTAD.

Two different approaches have been used to evaluate the consequences of reducing the amount of nuclear or cytoplasmic material on the cleavage rates of the eggs of *Arbacia punctulata* and *Lytechinus pictus*. In a series of studies, conducted in collaboration with S. Yuyama and L. Rustad, eggs were cut into nucleate and anucleate halves or quarters. After fertilization the diploid half and quarter-cells divided at the same time as the whole cells, and the haploid cells divided significantly later. However, there was no comparable increase in the duration of the second, third, or fourth mitotic cycles of the haploid cells.

The delay found in the first division cycle of the haploid cells might be a special consequence of the fact that the male pronucleus is initially condensed and not in equilibrium with the cytoplasm. Therefore, the "Hertwig effect" was used to prevent the male pronucleus from participating in the normal mitotic cycle of whole eggs. The sperm were exposed to massive doses of γ -rays (200 to 800 Kr.). At the lower doses almost all of the fertilized eggs underwent multipolar mitoses after several hours cleavage delay; however, a small percentage divided into two cells at the same time as the diploid eggs. High doses often immobilized the sperm. In successful experiments with the higher doses from 50% to more than 90% of the eggs which were fertilized with the irradiated sperm completed each of their first four cleavages at the same time as the controls. These experiments demonstrate that a biologically haploid cell containing a female pronucleus, instead of a male pronucleus, divides at the normal rate. Therefore, neither cytoplasmic volume nor ploidy *per se* determines the rate of cleavage of sea urchin eggs.

These studies were supported by the U. S. Atomic Energy Commission and the Office of Naval Research.

Mercenene clam extract: effects on the population kinetics of an experimental carcinoma and a non-neoplastic tissue. SISTER ARLINE CATHERINE SCHMEER, O.P.

The cell population kinetics for the 4-day-old transplantable Krebs 2 solid carcinoma (a non-metastatic tumor) and the cryptal progenitor compartment of duodenum from tumor-bearing CF1 mice has been determined by utilizing pulse label radioautographic techniques reported at these General Scientific Meetings last year. This report describes the effect

of mercenene clam extract on the population kinetics of the above named tissues. The Krebs propagation and transplantation was identical to that reported previously. The tumor and duodenum (a non-neoplastic tissue) were prepared for radioautographic analysis by utilizing standard procedures. A series of experiments consisted of 7 daily injections of mercenene into mice with 4-day-old Krebs carcinoma. Experimental and control animals were sacrificed 24 hours after each daily injection of clam extract, so that the last groups killed had received 7 doses of mercenene, and had tumors 11-days-old. $^3\text{H-TdR}$ was administered 30 minutes before sacrifice. The DNA synthetic index (DSI) (% labeled tumor cells/each 1000 non-labeled tumor cells scored), and mitotic index (MI), were then calculated. Preliminary results indicate that (1) mercenene is inhibiting the tumor cells during the G_1 phase of the cell cycle by preventing the cells from entering S (DNA synthetic period), (2) those cells in G_1 when the clam extract was administered were destined to die approximately 3 days later, and (3) those tumor cells not in G_1 were unaffected until, or unless, they were in G_1 when mercenene was injected, (4) the percentage labeled nuclei declined for a period equal to the time it takes the cells to die, and (5) the percentage of labeled cells increased as the non-DNA synthesizing population was eliminated. There was no significant difference between the MI of the experimental and control groups of Krebs tumor. The DSI and MI for the duodenal cryptal population demonstrated no significant variation between the experimental and control animals for any given time interval. The conclusion, then, is that mercenene, administered daily for 7 days, has no effect on the kinetics of the progenitor compartment of duodenum of tumor-bearing mice. The clam extract does act as an oncolytic agent against the cells of the Krebs solid carcinoma by preventing the cells from entering S. As reported previously, in *in vitro* studies on HeLa and human amnion (non-neoplastic) cell lines, the extract is non-toxic to amnion cells. It is extremely significant to have an anticancer drug that is selective in killing only neoplastic cells both *in vitro* and *in vivo*.

This research has been partially supported by U. S. Public Health Service Fellowship GM-34,903; and grants made to Dr. Gordon E. Stone, University of Colorado Medical Center, Denver, from the American Cancer Society (E-417) and the National Institutes of Health General Medical Science Division, GM-13,752. I wish to acknowledge the excellent technical assistance of Miss Musetta Hanson and Miss Ann Rees.

The control of molting in arthropods by steroid hormones. HOWARD A. SCHNEIDERMAN AND A. KRISHNAKUMARAN.

Insects are the only arthropods in which the control of molting is well-understood. A group of steroids—the ecdysones, thought to be produced by the prothoracic glands—cause molting in almost all insects upon which they have been tested. Ecdysones have also been isolated from a crustacean and have proven active in insects. However, no reports exist of successful efforts to induce molting in arthropods other than insects with ecdysones. This report demonstrates that ecdysones also control molting in four diverse classes of arthropods besides insects.

The ecdysone that was employed in most experiments was ecdysterone (=20-hydroxyecdysone), a steroid which has been isolated both from Crustacea and insects. As experimental animals we selected representatives of the major arthropod subphyla and classes: *Limulus polyphemus*, the horseshoe crab, Class Merostomata; *Araneus cornutus*, a spider, Class Arachnida; *Armadillidium vulgare*, a terrestrial isopod and *Procambarus* sp., a freshwater crayfish, Class Crustacea. Animals were maintained under conditions in which spontaneous molting was minimal. They were usually divided into 3 groups: experimental animals which received an injection of hormone in an appropriate Ringer of 10% ethanol; injected controls which received only the solvent; and uninjected controls. The effects were unambiguous: injection of 3 to 50 μg . of ecdysterone per g. live weight stimulated molting in all species of arthropods examined. In many animals molting occurred almost synchronously. The response was proportional to the dose: the larger the dose, the sooner the molt. Similar results could be obtained by injecting other ecdysones (inokosterone, alpha ecdysone, ponasterone A) and by topical application of a methanolic solution of ecdysterone.

From these data it appears likely that ecdysones were employed by the common ancestor of the major arthropod groups in the late pre-Cambrian, at least 200 million years before

vertebrates evolved. Probably this particular chemical control of molting evolved simultaneously with the development of a chitinous exo-skeleton and may exist in some non-arthropod groups.

Skeletal wall structure of a calcified bryozoan (Phylum Ectoprocta). THOMAS J. M. SCHOPF AND DOROTHY F. TRAVIS.

Knowledge of how the mineralized skeleton of bryozoans forms is essential for the evaluation of phylogenetic schemes derived from examination of calcareous wall layers. Discussion of this topic at the First International Conference on Bryozoa in 1968 showed that it is one of the most important current problems in the study of bryozoa. We have therefore given attention to the manner of calcification in *Schizoporella unicornis*, a representative of the most common group of bryozoans (Order Cheilostomata, Suborder Ascophora Vera).

A model of the calcification of this species can be reconstructed from the following cross-sections.

(1) Sections through the basal, lateral, proximal and distal walls into the main body cavity: cuticle, calcitic layer(s), epidermis, and peritoneum.

(2) A section through the frontal wall and compensation sac into the main body cavity: cuticle, epidermis, peritoneum, coelom, preitoneum, epidermis, aragonitic layer, calcitic layer, cuticle, compensation sac, cuticle, epidermis, peritoneum and coelom of the main body cavity.

The coelomic space above the compensation sac is continuous with that of the main body cavity through the proximal and lateral pores (areoles) of the frontal surface. The prominent pores through the calcitic and aragonitic layers are filled by epidermis and are attached to the cuticle layer that is immediately above the compensation sac. We envisage the function of these tissue plugs to be that of structural supports which help to suspend the compensation sac during its formation by invagination from the frontal wall and during deposition of the calcitic and aragonitic layers of the frontal wall.

A somewhat similar relationship of the calcified skeleton to soft tissues, including the extension of the coelom into the frontal layer above the compensation sac, was illustrated by Calvet (1900: pl. 7) for a species referred to *Microporella*.

The influence of actinomycin-D on distal-hydranth regeneration in Tubularia crocea.

IRWIN SINGER AND JOHN D. PALMER.

The regeneration of new hydranths on isolated hydrocaulus segments has been divided into arbitrary, well-defined morphological stages for the sake of descriptive convenience. The purpose of the study reported here was to determine the role of RNA synthesis in the completion of each of these stages. Actinomycin-D, an inhibitor of DNA-dependent RNA synthesis, was used for this purpose.

Seven- to eight-mm. hydrocaulus segments, removed 3 mm. below the hydranth, were placed in Millipore-filtered sea water containing 0.5, 1.5, 2.5, 5.0, 7.5, or 10.0 $\mu\text{g./ml.}$ actinomycin-D, for 4 hours at 14° C. Between 15 and 30 animals were subjected to each concentration. After this treatment they were washed, placed in fresh sea water, and distal-hydranth regeneration observed at 24-hour intervals for 6-7 days. In control animals, distal-hydranth regeneration was completed by 72 hours. Actinomycin-D had no effect on coenosarc healing or endodermal-ridge regression, but after these stages there was inhibition of further differentiation: inhibition increasing with increasing concentrations. In concentrations up through 2.5 $\mu\text{g./ml.}$, regeneration was retarded, but by 96 hours at least 88% of the individuals had completed regeneration (*i.e.*, reached the emergent stage). At 5.0 to 10.0 $\mu\text{g./ml.}$ never more than 36% of the animals reached the emergence stage, even by 168 hours.

Because of the very pronounced effects starting at 5 $\mu\text{g./ml.}$, the following experiments were undertaken with this concentration. Two to four hours after transection, healing of both ends of the hydrocaulus segments was complete and at this time they were treated with actinomycin for 4 hours; no inhibition resulted. If, however, after healing, the *proximal* end of the hydrocaulus was opened again, then a 4-hour treatment with actinomycin produced an inhibition curve identical with that obtained in the first experiments described above. The inhibition curve remained unchanged until actinomycin was administered through the open

proximal end at 22 hours after transection; a 4-hour treatment at this time produced a mitigated inhibition curve.

This work was supported by National Science Foundation grant GB-5045 to J.D.P.

The effects of temperature and various biological substances on mucociliary function of intact Nudibranchiata. PRANTIKA SOM AND F. B. BANG.

Evidence from our current studies on the mucociliary system of the footpad of intact Nudibranchiata, a shell-less opisthobranchiate mollusc, suggests that the increased viscosity of mucus plays a role in decreasing mucociliary transport rate at higher temperatures. With rise of temperature from 0 to 11° C. there was a concomitant rise in mucociliary transport rate reaching a maximum of 0.78 mm./sec. at 11° C., beyond which a gradual decrease in rate was accompanied by a greater viscosity of the mucus produced. Good ciliary vibration was still apparent up to 31° C. if the overlying sticky mucus was first removed. That the increased viscosity of mucus due to higher temperature decreased the rate of transport seemed to be confirmed by use of a mucolytic agent, N-acetyl cysteine, which increased the rate about 25% at temperatures above 11° C. Below 11° C. little variation from the untreated animal was seen.

A crude preparation of cholera toxin, known to affect mucus secretion in vertebrates, was employed in our system. Exposure of animals for 30 minutes at 21° C. in a concentration of 1 mg./ml. of this material accelerated the transport rate from pre-treatment average of 0.25 mm./sec. to 0.38 mm./sec. over the next 30 to 40 minutes. This was accompanied by increased secretion of mucus. There was a sudden fall to nil rate continuing for 8 to 12 hours followed by an equally sudden increase in rate to levels in excess of those seen before exposure to the toxin and persisting for 5 to 6 days. During this latter period the mucus secreted was very thin. Most animals, treated or normal, did not survive longer, but 2 animals, exposed to toxin, survived 10 days after treatment and showed a return to near normal transport rate, though the mucus continued to be thin.

The use of pilocarpine, a well-known stimulant of mucus secretion, increased the transport rate in normal animals at various temperatures but failed to stimulate toxin-treated animals. Thus, cholera toxin apparently causes a persistent lesion of mucus cells demonstrated by: (a) increased and persistent flow of thin mucus; (b) failure to respond to pilocarpine; and (c) an aberrant increase in rate with increasing temperatures.

Supported by a grant from the Tobacco Research Council.

*Nicotine-induced alteration of DNA and cell membranes in gametes of *Arbacia punctulata* and its bearing on human lung cancer.* CARL CASKEY SPEIDEL AND RALPH HOLT CHENEY.

In an extensive series of experiments, gametes of sea urchins were subjected for from 1 to 24 hours to nicotine solutions of 9 different strengths ranging from 0.5 to 0.00002 molar. Nicotinized gametes were then fertilized with normal gametes of the opposite type. Development was studied through the stages of cleavage, blastula, gastrula and pluteus.

Strong treatments of spermatozoa that still allowed fertilization could cause visible abnormalities and death of the young zygotes in any of the stages of cleavage before motility appeared. *Moderate* treatments allowed development to gastrulation or pluteus formation. Abnormalities were apparent also in these zygotes. Skeletal and ciliary deficiencies with associated deviations in swimming were particularly conspicuous. *Weak* treatments caused no perceptible abnormalities.

Eggs were subjected to similar nicotine treatments and fertilized with normal spermatozoa. Ensuing abnormalities were similar to those indicated above, although less strong treatments were sufficient to induce abnormalities of like degree. Motion picture records were made.

Since abnormalities of similar nature were induced in the two sets of experiments, we believe that the same underlying mechanism has been altered in nicotinized spermatozoa and eggs. The nicotinized spermatozoan head which enters the egg is composed almost entirely of haploid chromosome material. This combines with the normal haploid chromosome material of the egg, thus bringing about an abnormal diploid condition which then is responsible for

the resultant abnormalities exhibited by the progeny. Fundamentally, this indicates to us that DNA, the chief constituent of chromosomes, has been altered by nicotine.

Furthermore, a nicotized egg may suffer alteration of its cell membrane in such a manner as to allow polyspermy. Our observations suggest that nicotine may alter the cell membrane, cilia and DNA of respiratory tract cells in man. Such alterations would certainly favor the development of lung cancer in heavy cigarette smokers.

Supported by Grant RH 00325-02 to C. C. S. from the Environmental Health Division of the USPHS.

Pigment migration within fiddler crab melanophores. JAMES R. STEWART AND JONATHAN P. GREEN.

Hormonally and osmotically induced melanin migration in *Uca pugnax* was studied in a variety of inhibitor or activator solutions. Two bio-assay systems were used: perfusion of isolated legs with the test solution and incubation of isolated abdomens in the test solution. Advantages of the isolated abdomen assay over the isolated leg assay are its convenience and the assurance of uniform distribution of the test solution to all melanophores. Comparable results were obtainable with either assay system. Melanophores were staged according to the scheme of Hogben and Slome.

Ouabain ($1.4 \times 10^{-4} M$), an inhibitor of sodium-dependent active transport, in combination with melanophore-dispersing hormone (MDH) in sea water, had no inhibitory effect on melanin dispersion in either legs or abdomens. Ouabain in the absence of MDH had no dispersal effect on concentrated melanin. Similar results were obtained with 2,4-dinitrophenol ($5.0 \times 10^{-4} M$), an oxidative phosphorylation uncoupling agent.

Mersalyl (0.1 M), a sulfhydryl bond inhibitor, in sea water reduced the rate of MDH-mediated melanophore expansion by 39% over a three-hour period. Mersalyl in 100% and in 80% sea water did not effect pigment dispersion.

Dinitrophenol ($5.3 \times 10^{-4} M$) in 200% sea water caused a 20% decrease in the rate of melanin contraction over a one-hour period. Similar results were obtained with ouabain ($1.4 \times 10^{-4} M$) in 200% sea water.

Calcium chloride and magnesium chloride (0.6 M) increased the rate of melanin contraction over that of 200% sea water by 30% and 28%, respectively, over a one-hour period.

These results suggest that sulfhydryl bond formation may be involved in the action of melanin-dispersing hormone. It is unlikely that oxidative phosphorylation or a sodium-dependent active pump process alone is responsible for melanin dispersion. On the other hand, the available evidence points towards active ion extrusion followed by water efflux as a mechanism for melanin concentration.

Supported by National Science Foundation Grant #GY-4304 to Brown University for undergraduate research, by NSF Grant GB-7261 (to J. P. G.) and by NSF Grant GB-3447 to the Department of Invertebrate Zoology, Marine Biological Laboratory.

ATPase and actin combining properties of "myosin" from molluscan smooth muscle.

A. STRACHER, ROBERT V. RICE AND G. F. ELLIOTT.

Attempts to prepare myosin from the opaque portion of the adductor muscle of *Mercenaria mercenaria* by the standard method used for vertebrate striated muscle have failed to produce a protein with the known enzymatic properties of myosin. For this reason, the method of Hardwicke, utilizing a sucrose gradient, was used in order to localize and characterize the fraction containing the ATPase activity. After completion of the density gradient run, the bottom of the tube was punctured with a needle and ten 2-ml. fractions were collected. Analysis of each tube for enzymatic activity showed that the third fraction from the top of the tube contained the ATPase activity. This fraction was dialyzed against 0.1 M ammonium acetate, pH 7.0, in preparation for shadow casting. Only 50 Å globules were seen in the electron microscope from two separate experiments. Assuming an $E_{280}^{1\%} = 7.0$, the specific activity was equal to approximately 0.5 μ moles Pi/mg./min., comparable to myosin from vertebrate striated muscle. The activity was quite readily lost if the protein was allowed to remain at 4° C. for longer than 6 hours. Dithiothreitol did not protect against this loss in

activity. The dependence of ATPase on both pH and Ca^{2+} was similar to that of myosin from skeletal muscle. If the homogenization and sucrose gradient were carried out in the presence of 0.6 M KCl, the pellets consisting of paramyosin ribbons obtained at the bottom of the tube were devoid of ATPase. The pellet from the 0.1 M KCl run, however, had a significant amount of ATPase. These results are consistent with the findings described in the abstract by Rice *et al.* Preliminary experiments suggest that an interaction between rabbit skeletal F-actin and molluscan smooth muscle "myosin" occurs, as evidenced by a disappearance of the ATPase from tube 3 when F-actin is added to the 0.1 M KCl homogenate before layering on the sucrose gradient.

Studies on the life-history of Distomum pyriforme Linton, 1900. HORACE W. STUNKARD.

Stunkard (1967) reported on the life-cycle and developmental stages of a digenetic trematode whose metacercarial stages occur in medusae. Ophthalmotrichocercous cercariae develop in rediae in *Anachis avara*. They penetrated and persisted as unencysted metacercariae in *Gonionemus vertens* and *Chrysora quinquecirrha*. When fed to fishes, sexually mature worms were recovered from the scup, *Stenotomus chrysops*. Adult worms are very similar morphologically to *Distomum pyriforme* Linton, 1900, described from the pyloric ceca and intestine of the rudderfish, *Palinurichthys perciformis*, and tentatively they are assigned to that species. The studies have been continued; the number and location of the penetration-glands and flame-cells of the cercaria have been determined. Eight pairs of penetration-glands are situated in the lateral areas between the ocelli and the acetabulum. On each side, ducts from three cells pass forward, lateral to the ocellus, and those of five cells pass mediad of the ocellus. All open at the anterior end of the body. Details of the excretory system have been worked out. The pore is terminal, preceded by a sphincter which closes the uroproct. Anterior to the uroproct there is a pulsatile bladder which is continued anteriorly by a tubular reserve vesicle, filled with spherical concretions, that extends to the level of the pharynx. From the anterolateral faces of the bladder, collecting ducts, lined with cilia, pass forward, lateral to the digestive ceca. Anterior to the acetabulum each divides into anterior and posterior branches. Each branch bears three sets of flame-cells, four in each set and the formula is [(4+4+4) + (4+4+4)]. Eggs from the worms have been embryonated; the miracidia develop and emerge in 8 to 10 days at laboratory temperature; they have ocelli with conspicuous lenses, long cilia, and swim rapidly. Life-history data clarify the systematic position of *D. pyriforme*.

Investigation supported by NSF GB 3606, continuation of G 23561.

Localization of bioluminescence in the marine dinoflagellate Pyrocystis lunula Schüitt using an image intensifier. ELIJAH SWIFT AND GEORGE T. REYNOLDS.

Cultures of *P. lunula* were entrained on a 12:12 LD cycle. The non-motile bioluminescent lunate cysts were examined during scotophase in small chambers with water immersion lenses. During scotophase the cyst's plastids migrate to distal areas and facilitate observation of the centrally located nucleus and the perinuclear area, which is *ca.* 25 nm. in diameter and contains obvious starch grains. Macroflashes were stimulated by electrical impulses delivered to electrodes placed near the specimen on the microscope stage. Miniature flashes and glows were associated with chemical and mechanical damage to the cysts. The microscope output was focused on the photosensitive cathode of the image intensifier at magnifications from 20× to 200×. For viewing macroflashes, the image intensifier gain was 9.0×10^6 . For viewing the steady glow, the gain was *ca.* 1×10^6 . By means of dark field illumination, the structural features of the specimen were photographed through the same microscope-intensifier system, so that the structural sources of the luminescence could be identified.

The photographs obtained indicate that the electrically stimulated flash (macroflash) is produced in the entire perinuclear area. Miniature flashes were localized in several different parts of the perinuclear area. These miniature flashes occurred in a protoplast after the introduction of propionic acid to its medium. The steady glow was observed to come from the entire perinuclear area. On the photographs, the nucleus appears as a dark area. The steady glow was presumed to be caused by damage to the cyst since the glow, $>2 \times 10^6$ quanta

sec.⁻¹, is much brighter than spontaneous glow of *P. lunula* cysts. Thus the photographic records provide evidence that the macroflash, miniature flash, and glow are all produced in the small area of the protoplast of *P. lunula* near the nucleus.

Supported in part by AEC contract AT(30-1)-3406 and in part by NSF Grant No. GZ 259.

Fertilization-induced increase of trypsin-like activity in sea urchin "membrane" during embryogenesis. WALTER TROLL, ALBERT GROSSMAN AND SARAH CHASIS.

The mechanism by which protein synthesis and cellular differentiation are initiated by the entry of the sperm into the egg has remained a central problem of biology. Monroy and his co-workers have shown that the ability of ribosomes from unfertilized eggs to engage in protein synthesis is enhanced by treatment with trypsin. This report is concerned with the induction of a trypsin-like enzyme as part of the fertilization process. The enzyme is measured using the synthetic substrates tosyl-arginine methyl ester (TAME) and acetyl lysine methyl ester (ALME). Crystalline trypsin hydrolyzes these substrates, preferring TAME over ALME by eight to one. An enzyme attacking these substrates was isolated from *Arbacia punctulata* eggs by freezing and thawing five times then washing with Millipore-filtered sea water, sonicating and centrifuging. The residue represents the membrane fraction and contains the enzyme. Eggs were fertilized and samples taken at hourly intervals. A 10-fold increase of activity was noted over a 20-hour period. The enzyme differed from trypsin in that it attacked ALME and TAME equally well. Pre-incubation with Actinomycin D (20 µg. per ml.) completely abolished the increase. Tosyl lysine chloromethyl ketone (TLCK), a specific inhibitor of trypsin, inhibited the membrane enzyme. The presence of 0.002 M TLCK prior to and during incubation inhibited the division of *Arbacia*; at 0.0002 M, development stopped at blastula. This suggests two roles for the enzyme: (1) the activation of protein synthesis, and (2) the activation of genome.

Work supported by GM-13728, Core Grant BSS-ES 00014 NCI-CA 06989 and Allied Chemical Corporation. A. G. is a Senior Postdoctoral Fellow of the New York Heart Association.

The responses of granule cells in the cerebellum of Mustelus canis. NAKAAKIRA TSUKAHARA, HELENA TÁBORÍKOVÁ AND JOHN C. ECCLES.

The selachian cerebellum differs from the cerebella of higher vertebrates in that the granule cells are concentrated in two large medial cords. Mossy fibers pass to these cords *via* the stratum fibrosum. The axons of granule cells, the parallel fibers, pass up to the molecular layer and then transversely, so making synapses with the dendrites of Purkyně and stellate cells. Purkyně cells are absent over the medial part of the granule cell cords, so a microelectrode there records selectively field potentials produced by granule cells. A mossy fiber volley in the stratum fibrosum evokes in the granule cell cords an initial diphasic potential followed by a slow negative wave, which are attributable, respectively, to mossy fiber impulses and the excitatory synaptic potential induced thereby in granule cells. With increase in the mossy fiber volley these potentials become larger and granule cell spikes are superimposed upon the slow wave. Evidently spatial summation is required for evoking this discharge. A second stimulus through the same electrode reveals that mossy fibers recover fully from refractoriness in about 5 msec. and produce a large second EPSP. This temporal summation of EPSPs causes impulse discharge when single volleys are ineffective, and such facilitation persists for over 70 msec. When parallel fibers are stimulated directly, impulse propagation occurs along the transverse course of the parallel fibers with a conduction velocity of about 0.2 m./sec. These impulses propagate to the granule cells antidromically, the evoked spike potentials being observed in the granule cell cord. With double stimulation it is found that the recovery of antidromic invasion of granule cells occurs earlier than with Purkyně cells, beginning at 5 msec. and with full recovery by 20 msec. No inhibitory action of presumed Golgi cells has been found, otherwise the investigations summarized in these two papers indicate that physiologically the neurones parallel those of mammalian cerebellum.

This work was supported by the Grass Foundation.

Oxygen-linked dissociation of hemocyanin II. A proposed model. K. E. VAN HOLDE AND HENRY A. DEPHILLIPS, JR.

As has been shown in a previous abstract, the aggregation state of squid hemocyanin is a function of the per cent oxygenation. Any satisfactory model must explain both the dissociation upon partial oxygenation and the reassociation upon complete oxygenation.

Since each 19S subunit contains about sixteen oxygen binding sites (from known stoichiometry), there should exist a distribution of levels of oxygenation among the set of subunits, at any given value of average oxygenation. If noncooperative binding is assumed, as a first approximation, this can be shown to be a binomial distribution. We now further assume that subunits carrying more than a critical number (n_1) but less than a second critical number (n_2) of bound oxygens are incapable of entering into 59S aggregates. It is then possible to calculate the per cent dissociation to be expected at any overall level of oxygenation. With the choice $n_1 = 10$, $n_2 = 15$, excellent agreement is obtained between theory and experiment.

This model further predicts that at low overall oxygenation the 19S component will be most highly oxygenated, whereas at high oxygenation the 59S component will be slightly more oxygenated. These predictions have been checked by use of the scanner attachment to the analytical centrifuge. Measurement of the optical density of the two boundaries at 345 $m\mu$ yields the concentration of oxygenated protein; on the other hand, the total protein concentration in each boundary can be measured from integrated schlieren patterns. The results of experiments at both low and high levels of oxygenation are in accord with the predictions of the model.

This research was supported by Public Health Service Grants Nos. HE-11671 and HE-12326.

Ribosomal RNA cistrons in single and multinucleolate oocytes. W. S. VINCENT, H. O. HALVORSON, H. R. CHEN AND D. SHIN.

The multinucleolate oocyte of amphibians has been shown to contain reiterated ribosomal RNA (rRNA) cistrons to the extent that the germinal vesicle contains more DNA ascribable to these cistrons than to the entire remaining genome. The question arises as to whether this amplification of ribosomal cistrons is a phenomenon only of amphibian oocytes, or whether it exists in other multinucleolate and uninucleolate forms.

We have examined the ribosomal cistron content of DNA isolated from red blood cells and ovarian tissues containing immature oocytes of the striped bass, *Roccus saxatilis*, a form with multinucleolate oocytes; we have also examined DNA from the sperm, eggs and ovarian tissues of the starfish, *Asterias forbesii*, which contains uninucleolate oocytes. Ribosomal cistrons were detected by RNA-DNA hybrid formation on nitro-cellulose filters with appropriate controls of temperature, incubation time, rRNA saturation levels, salt concentration, fixation of DNA to filters and competition by homologous rRNA.

All DNA species examined demonstrate a heavy satellite region detected by rRNA hybridization following equilibration centrifugation in cesium chloride. In the starfish DNA a satellite band was detected with the Model E ultracentrifuge at $d = 1.710$, with a main band at $d = 1.689$. Determination of rRNA binding to DNA gave the following results (in $\mu\mu\text{g}$. rRNA bound / μg . DNA): striped bass rbc DNA: 0.20, ovarian DNA: 2.34; starfish sperm: 0.37, ovarian DNA: 0.35.

From these data we conclude that although the rRNA cistrons are normally redundant in these two forms, amplification of the cistrons is found only in striped bass which has multinucleolate oocytes, not in the uninucleolate starfish oocyte. This suggests that nucleolar number in oocytes may be a function of ribosomal gene amplification.

Supported by Grants from NIH: GM-11480 and AI-1459, from NSF: GB-3273 and B-1750.

The electroretinogram and spectral sensitivity of the common scallop. GEORGE WALD AND EDWARD B. SELDIN.

The eye of the common or bay scallop, *Aequipecten irradians*, contains two retinas, proximal and distal, each sending a branch to the optic nerve. Hartline showed that the

proximal retina discharges impulses in the optic nerve only when light is turned on or up, the distal retina only on turning light down or off.

The electroretinogram (ERG) in response to flashes or continuous light consists potentially of two waves. At low intensities in the dark-adapted eye the first wave appears alone, its latency about 100 msec. and peaking at about 200 msec. At higher intensities and on light adaptation it is accompanied by a second, slow wave, peaking at about 300 msec. In dark-adapted eyes the first wave may be cornea-positive or -negative. In partly light-adapted eyes it is always cornea-negative, as is always the second wave. These on-responses presumably originate in the proximal retina. The ERG displays no off-response, hence apparently no signal from the distal retina.

The spectral sensitivity of dark adapted eyes was measured—the number of photons per 18-msec. flash that at each wavelength evoked a constant amplitude of ERG (20 or 50 μ volts). A simple curve results, with average λ_{\max} 490–495 $m\mu$ (range 480–507 $m\mu$). This is not changed in shape or position in the spectrum by adaptation to bright red or violet light. It seems therefore to be based upon a single visual pigment, resembling in spectrum the rhodopsins earlier extracted from squid and cuttlefish retinas.

This investigation was supported in part by a National Science Foundation grant to G. W.

Optical changes associated with nerve excitation and membrane polarization. A. WATANABE, I. TASAKI AND L. CARNAY.

Nerves from lobster, spider crab and squid fluoresce when stained with such dyes as acridine orange or 8-anilino-naphthalene-1-sulfonic acid. The intensity of fluorescence, detected at a right angle to the incident light, was found to increase transiently when the nerve was stimulated electrically. At the peak of excitation, the increase was within the range of $(1-5) \times 10^{-4}$ times the intensity of the background fluorescent light. The time-course of the increase was roughly parallel to that of the action potential. Similarly, the birefringence and turbidity of the nerve were shown to exhibit transient changes during nerve excitation. (The results of the present birefringence and turbidity studies are consistent with those reported recently by Cohen *et al.*). Optical changes of the nerve could also be produced by altering the ratio of the univalent-divalent cation concentration of the medium, by temperature changes and by electrotonic polarization of the nerve. The sign of the optical signals produced by cathodal polarization was the same as that produced by the action potential; anodal polarization produced optical changes of the reversed polarity. Within a certain range of polarization intensity, the optical effect of polarization and that of the action potential were additive. This finding suggests that the optical changes of the nerve are a rather complex phenomenon, involving various anatomical structures in the vicinity of the axonal membrane.

The effect of high hydrostatic pressure on the mechanical properties of the membrane of the sea urchin egg. LEWIS WOLPERT, DOUGLAS MARSLAND AND MICHAEL HIRSHFIELD.

There is evidence that high hydrostatic pressure can solate and weaken gel-like structures within cells. It is thus of interest to determine its effect on the mechanical properties of the cell membrane since it has been suggested that there is a gel-like cortex beneath the plasma membrane which makes a significant contribution to its mechanical properties. The mechanical property measured was the resistance to deformation when the eggs are compressed. The eggs of *Arbacia* were compressed by placing a small plate made either from a fragment of coverslip (120 μ thick) or Meline plastic sheet (50 μ thick) on a small group of eggs (20 to 100) so that the eggs supported the plate. The deformation of the eggs was determined by measuring their increased diameter. The diameter of the eggs was about 70 μ and a force of about 8×10^{-8} dynes/egg increased this to about 110 μ . Compressed unfertilized eggs from which the jelly coat had been removed by acid sea water showed no change in diameter when subjected to 12,000 lb./in.² for 30 minutes. The mechanical properties of the cell membrane of unfertilized eggs appear to be unaffected by high hydrostatic pressure. When compressed fertilized eggs, from which the fertilization membrane had been removed by urea treatment, were subjected to pressure of about 10,000 lb./in.² there was a significant

increase in cell diameter. This increase was such that it appeared as if the resistance to deformation of the membrane were halved. Surprisingly after a few minutes the diameter decreased again indicating a contraction at the membrane even while under high hydrostatic pressure. Release of the hydrostatic pressure resulted in a further reduction in diameter and thus contraction at the surface, the final diameter sometimes being less than that before the hydrostatic pressure was applied.

The effect of light on the dogfish eye. SEYMOUR ZIGMAN AND SUSAN J. BAGLEY.

The influence of strong light over protein synthesis in dogfish (*Mustelus canis*) ocular tissue was observed *in vitro* using elasmobranch Ringer's medium. Corneas, lenses, and retinas of medium-age dogfish were obtained fresh and incubated at 20° C. under 95% O₂:5% CO₂ in Ringer's solutions containing 1 μc./ml. of C¹⁴ amino acid mixture in the dark or illuminated by a 40W circular cool white fluorescent lamp with 475 and 570 mμ wave-length maxima. The intensity of light in the incubation chamber was 500-600 ft. candles.

Incorporation of C¹⁴-amino acids into the insoluble (1200 g residue), particulate (105,000 g residue), and soluble proteins was measured by counting TCA precipitates of these fractions after removing residual radioactivity by suction filtration. Specific activities were calculated from the results of liquid scintillation counting in Bray's solution of hyamine hydroxide solubilized protein, and the Lowry (phenol) protein analysis.

Amino acid incorporation into the particulate fractions of cornea epithelial cells, lens epithelial cells, lens proper, and whole retina between 7 and 22 hours in the light was reduced to approximately $\frac{1}{2}$ that of the tissues kept in the dark. Soluble protein synthesis was reduced by about 50% also in lens epithelium and retina. Inhibition of amino acid incorporation into soluble proteins was not observed.

When total collagen was extracted from corneal stroma, amino acid incorporation was found to be inhibited by 20% during 24 hours of illumination. No effect was noted upon incorporation into soluble lens proteins. No appreciable conversion of soluble to insoluble proteins took place.

Whole retinal rods isolated by soaking and shaking isolated retinas in ice-cold Ringer's medium (which contains urea at 0.3 M), were incubated up to 24 hours in the light and in the dark. By 24 hours, a loss of 65% of protein synthetic activity was observed in illuminated rods, concomitant with appreciable cell damage. Partial recovery of protein synthesis could be achieved by returning the rods to the dark after 6 hours.

The mechanism of light inhibition of protein synthesis in ocular tissues is under investigation.

Supported by ONR/CNA grant (University of Rochester) and Rochester Eye Bank.

Comments on the extraction of Thalassopsammon from intertidal and subtidal sands.

DONALD J. ZINN.

The removal of the interstitial fauna from intertidal and subtidal sands continues to be a relatively difficult technique in the efficiency and reliability of extraction of a desired taxon or taxa. Although the more commonly used methods of extraction, conveniently categorized as puddling, mechanical, salinity gradient, bubbling and elutriation, elutriation and bubbling are apparently the most satisfactory means for finding maximum group representation (qualitative), these methods as originally described are only occasionally suitable for quantitative extractions. For example, ciliates, tardigrades, turbellarians and gastrotrichs are removed with greatest efficiency by sea-ice (salinity gradient); mystacocarids, copepods, oligochaetes, polychaetes and kinorhynchs are best extracted with bubbling, while ostracods and nematodes may be optimally removed by elutriation.

Recent and present investigations have indicated strongly that (1) extraction efficiencies of different higher and lower taxa exhibit different rates; (2) the presence of excess detritus, non-capillary sediments bacteria, yeasts and algae can alter extraction rates radically to the point of prevention; (3) soft-bodied and contractile forms are most efficiently extracted with modifications of a standard method best for the class; and (4) the most reliable method for testing efficiency is to place a known number of identified individuals into a measured

amount of sterilized natural sand and extract with the developed method of choice. Uhlig, using the sea-ice method he developed, indicated an efficiency of 90% for ciliates and 75% for turbellarians, while Fenchel, also working on ciliates, found a variation of 70% to 90% for the same method depending on the manipulation of the animals.

Special methods must be developed by each investigator for extracting interstitial taxa of special interest. Examples of variants on basic methods include introduction of narcotizing agents (magnesium chloride, propylene phenoxtyol, etc.), bubbling with a gas (carbon dioxide, nitrogen, etc.), elutriation with diluted and/or heated sea water and varying the temperature of the sample before extraction.

Contribution No. 164 of the Systematics-Ecology Program, Marine Biological Laboratory, Woods Hole, Massachusetts. Grateful acknowledgment is made for assistance provided by the University of Rhode Island Grant-in-aid No. 132-151.

ERRATA

The following corrections should be made in the paper by Lofts, Pickford and Atz (1968), *Biol. Bull.*, **134**: 74-86:

Page 75, line 17—*Group III*. 10° C.

Page 76, line 20—Group II.

Page 78, Table I—For II, read III; for III, read II.

Page 85, Rasquin and Hafter. *Anat. Rec.* **111**.

THE BIOLOGICAL BULLETIN

PUBLISHED BY THE MARINE BIOLOGICAL LABORATORY

SEXUAL RESPONSES OF ADULT MALE AEDES AEGYPTI USING THE FORCED-COPULATION TECHNIQUE^{1, 2}

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Our present knowledge concerning the sexual responses of the male Yellow Fever Mosquito *Aedes (Stegomyia) aegypti* Linnaeus is based largely on the studies of Roth (1948), Spielman (1964), Jones and Wheeler (1965a, 1965b), Spielman *et al.* (1967), and Craig (1967). According to Roth (1948), adult males in cages copulate repeatedly and remain in a permanent mating state throughout their lives. He observed that one virgin male caged with 16 virgin females for 30 minutes copulated 30 times and made 10 additional attempts to do so. He found that males would copulate with 50-minute-old females. Jones and Wheeler (1965a), using the forced-copulation technique, reported that males would copulate with 4 to 10 virgins within 10 to 20 minutes. Wheeler and Jones (1960) found that while males could be forced to copulate with newly emerged females, they were not inseminated. Lea (1967, 1968) found this to be true for young caged females, and showed that this was influenced by the corpora allata, and was temperature- and strain-dependent. According to Gwadz (1967), females in a cage copulate repeatedly but do not become inseminated until they are 40 to 48 hours old.

Vandellhey and Craig (1958) reported that multiple insemination occurred in caged groups of *Aedes aegypti*; Craig (1967) subsequently noted this was true for matings which occurred over a short period of time. Jones and Wheeler (1965b) discovered that males did not force-copulate with inseminated females, and Ludlam (1965) confirmed this. George (1967) reported that while males repeatedly copulate in cages, only the first mating was effective. Craig (1966, 1967) showed that this was due to the male accessory gland component of the semen from the first

¹ Supported by N.I.H. Grant GM 6021. One of us (J. C. J.) was additionally supported by N.I.H. Career Development Award K3, GM 21,529. Scientific article number A 1458, contribution number 4089 of the Maryland Agricultural Experiment Station.

² Based on a thesis submitted by the first author to the Faculty of the Graduate School of the University of Maryland in partial fulfillment of the requirements for the degree of Master of Sciences, 1968.

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insemination. After implanting male accessory glands into virgin females, Craig obtained a 25% insemination rate using forced-copulation and no insemination with those placed in a cage with fresh males. Spielman *et al.* (1967) reported that when once-mated females were paired with males in a cage 47% of them copulated again, and those which remated very rapidly ejected the semen from the bursae.

It is evident that much remains to be learned about the sexual responses of *A. aegypti* under both forced and cage mating conditions. This paper is concerned with the details of the sexual responses of *A. aegypti* males to very young virgins or to previously copulated females, using the forced-copulation technique, and also includes tests of male potency.

The specific terms which will be used in this paper need to be defined at the outset. *Copulation* refers to the placement of the aedeagus into the upper vagina of the female. *Precoital time* extends from the initial second of the presentation of the male to the beginning of aedeagal emplacement. *Coital time* is the period from the beginning of aedeagal emplacement to its withdrawal from the vagina. *Insemination* is the placement of sperm and male accessory gland materials in the bursa and is followed by spermathecal filling and retention of the male accessory gland material. *Fertilization* is restricted to the placement of spermatozoa in the egg.

MATERIALS AND METHODS

The mosquitoes of the Bangkok strain were reared at 27° C. and 70% relative humidity. The larvae generally pupated on the sixth and seventh days after hatching. Pupae were sexed by examining their external genitalia with a stereomicroscope at 16×, using the criteria of Christophers (1960). Virgin adults of each sex and of known ages were also obtained by removal and separation of the sexes as they emerged. The adults were maintained on 5% sucrose. Test lots of adults were kept either in 28 cm. square wire cages or in standard U. S. pint cartons with a fine net cover on top, and were supplied with sucrose on cotton pads. The induced copulation techniques of McDaniel and Horsefall (1957) and Wheeler (1962) were employed; however, none of the insects were decapitated. All forced-copulations (forced-matings) were with mosquitoes that had recovered from the nitrogen anesthesia which was used in the pinning procedure. Females caged with males (sex ratio 1:1) for 2 days were assumed to be already inseminated when used for subsequent forced-copulation tests, and will be designated as *cage-copulated* or *cage-mated* females. All males were offered only once to a female and then were discarded except in the potency tests. Sperm checks of females were made by examination of the spermathecae and/or the bursa at 430×.

RESULTS

1. Male responses to young virgin females

Forty-eight-hour-old males were presented to 171 young females of nine precisely known age groups (0, 1, 2, 3, 4, 8, 12, and 24 hours $\pm 1\%$). The young females often interfered with the copulatory attempts of the males by kicking with their legs (chiefly the metathoracic legs), and when this happened the legs were amputated at the midpoint of the femurs.

TABLE I

Coital times of young virgin Aedes aegypti (L.) females which were force-copulated with virgin males 48 hours old. Cases of prolonged coitus are not included.

Age of females	Inseminated females			Uninseminated females		
	Number of copulations	Coital time, seconds		Number of copulations	Coital time, seconds	
		range	mean		range	mean
0	3	21-724	58.3	1	6	6.0
1 hour	3	18-49	34.3	2	4-6	5.0
2 hours	4	33-70	47.8	2	8-10	9.0
3 hours	3	7-120	55.7	0	—	0
4 hours	6	13-200	62.8	2	6-11	8.5
8 hours	11	40-160	85.1	1	120*	120.0
12 hours**	1	40	40.0	1	16	16.0
12 hours	1	115	115.0	1	4	4.0
16 hours	10	10-134	70.4	1	4	4.0
24 hours**	13	30-398	113.8	2	4-12	8.0
24 hours	9	11-99	31.6	1	7	7.0
	64	7-398	71.4 ± 7.7***	14	4-120	7.5 ± 1.0***

* Male ejaculated prematurely, datum excluded from mean of total coital time.

** Test performed at two different times.

*** Mean based on total data.

The male responses to these females were either (1) rapid copulation, (2) seizure of the female's cerci and then not copulating, (3) no visible reaction at all, or (4) withdrawal from the female terminalium.

The only visible response of the females to the males was withdrawal of their

TABLE II

Copulation and insemination rates of young virgin female Aedes aegypti (L.) which were force-mated with virgin males 48 hours old

Age of females	Number used	Copulations		Inseminated		Uninseminated		Prolonged coitus	
		Number	%	Number	%	Number	%	Number	%
0	16	4	25.0	3	75.0	1	25.0	0	0
1 hour	15	5	33.3	3	60.0	2	40.0	0	0
2 hours	15	6	40.0	4	66.7	2	33.3	0	0
3 hours	15	4	26.7	3	75.0	0	0.0	1	25.0
4 hours	21	10	47.6	6	60.0	2	20.0	2	20.0
8 hours	16	14	87.5	11	78.6	1	7.1	2	14.3
12 hours*	10	2	20.0	1	50.0	1	50.0	0	0
12 hours	15	3	20.0	1	33.3	1	33.3	1	33.3
16 hours	15	12	80.0	10	83.3	1	8.3	1	8.3
24 hours*	18	15	83.3	13	86.7	2	12.5	0	0
24 hours	15	10	66.7	9	90.0	1	10.0	0	0
	171	85	49.7%	64	75.3%	14	16.5%	7	8.2%

* Test performed at two different times.

terminalia. Very young females emitted droplets from their ani and these often interfered with the male's clasping of the cerci and prevented copulation. This was observed primarily with 0- to 1-hour-old females, and was seldom seen in females 8 or more hours old.

The individual coital times of inseminated females ranged from 4 to 398 seconds, with a mean of 71.4 ± 7.7 seconds, compared to a range of 4 to 16 seconds and mean of 7.5 ± 1.0 seconds for the unseminated females (Table I). From 20% to 87.5% (85/171 average of 49.7%) of the females force-copulated, and 33.3% to 90% (65/85 average of 75.3%) were successfully inseminated (Table II).

Prolonged forced-copulation was observed with 7 out of 85 pairs of mosquitoes. Most of the males were unable to free themselves at will by even the most strenuous efforts. Only one male, after copulating for 360 seconds, succeeded in freeing himself after struggling for 208 seconds. When the female was dissected 45 minutes later, sperm were seen *only* in the bursa. The other six pairs were *in copula* so tightly that manual stretching tore their abdomens apart rather than separating their terminalia.

TABLE III

Precoital and coital times of force-copulated Aedes aegypti (L.) males in 4 age groups; each male was offered to 15 virgin females of the same age

Age of males	Number used	Number of copulations	Precoital time, seconds		Coital time, seconds	
			range	mean	range	mean
1 Day	10	58	1-171	19.7	3-45	18.5
3 Days	10	68	1-103	16.2	3-66	20.9
5 Days	10	72	1-43	10.2	4-55	20.9
7 Days	10	89	1-105	18.3	4-57	20.3
Means	40	287 7.2 ± 0.5	1-171	$16.0 \pm 1.2^*$	3-66	$20.2 \pm 0.7^*$

* Mean based on total data.

2. Male potency and variability of response to virgin females

Adults of precisely known ages (1-, 3-, 5-, and 7-day-old lots $\pm 4\%$) were used to test the responses of 10 individual virgin males (per lot), each individual being presented in sequence and at known intervals to 15 virgin females, or to as many as would be accepted using the forced-copulation technique. Each of the 4 tests required about 6 hours to complete. The responses of the males were subjectively graded as either (1) feeble, (2) moderate, (3) active, or (4) highly active attempts to copulate, or as (5) withdrawal of their terminalia from those of the females. The first 4 responses were qualitatively assessed by the following activities of the males: (1) the extent of tarsal contact with the female, (2) the degree of flexion of the male's terminalium into the female, (3) the speed and number of attempts to copulate, (4) the forcefulness of cercal clasping, and (5) the extension of the paraprocts with eversion of the aedeagus. Withdrawal of the male's terminalium included a pushing of the female with his tarsi and a flexion of the terminalium away from that of the female.

The precoital times of the 40 males ranged from 1 to 171 seconds, with a mean of 16.0 ± 1.2 seconds. The coital times ranged from 3 to 66 seconds, with a mean of 20.2 ± 0.7 seconds (Table III). The males copulated with 287 out of 600 females (47.8%). The number of females acceptable to individual males ranged from 2 to 15, with a mean of 7.2 ± 0.5 .

Most of the females were not accepted sequentially by the males; only one out of the 40 males used accepted all 15 females, and this individual did not accept 2 additional females. This highly potent male's initial precoital and coital times were 21 and 24 seconds, respectively. His final precoital and coital times were 3 and 4 seconds, respectively. In these experiments, 6 or more females were accepted sequentially by 19 out of 40 males (47.5%), and 8 or more females were accepted sequentially by 7 out of 40 males (17.5%). Only 2 out of 40 males (5%) copulated sequentially with less than 3 or more than 10 females.

The age of the mosquitoes apparently did not influence either the precoital or coital times (Table III). The number of copulations by 5- and 7-day-old males was significantly higher than that by 1- and 3-day-old males (p less than 0.01).

TABLE IV

Responses of Aedes aegypti (L.) males in 4 age groups; each male was offered to 15 virgin females of the same age

Male age	Withdrawal responses	Coital responses					Total responses
		Feeble	Moderate	Active	Highly active	Row totals	
1 Day	5	3	29	15	11	58	63
3 Days	13	20	22	15	11	68	81
5 Days	1	30	24	13	5	72	73
7 Days	3	28	28	23	10	89	92
	22 7.1%	81 28.2%	103 35.9%	66 23.0%	37 12.9%	287 92.9%	309

The increase in the number of copulations of 3-day-old males *versus* 1-day-old males and of 7-day-old males *versus* 5-day-old males was significant (p less than 0.01).

Presentation of a virgin female to a male resulted in definite responses to the female in 309 out of 600 males (52%). In the other 287 (48%), the males showed no recognizable response to the females. Of the responses that did occur, 7.1% (22/309) were withdrawals and 92.9% (287/309) were coital responses. The degree of the coital responses were as follows: feeble, 28.2% (81/287); moderate, 35.9% (103/287); active, 23.0% (66/287); and highly active, 12.9% (37/287) (Table IV).

3. Forced copulation of virgin versus cage-mated females

One hundred virgin and 100 cage-mated females were each individually offered a series of one to 5 males, in 5 lots of 20, using the forced-copulation technique. If copulation occurred, no additional males were presented. All the mosquitoes were 48 hours old ± 3 hours.

Among the virgins, 95% force-mated for 6 to 76 seconds, with means for the 5 replicates ranging from 22.8 to 29.9 seconds, and with an overall mean of 26.7 ± 1.3 seconds: 81 out of 95 virgins (85.3%) accepted the first male offered, and 10 out of 95 accepted the second male (10.5%).

In great contrast to the above, only one out of the 100 cage-mated females could be force-copulated, and this one did so for only 3 seconds.

Fifty of those virgin females which had previously been successfully force-copulated were again presented to virgin males for a second time, 12 hours after the initial force-copulation. Only 5 out of 50 of them (10%) force-copulated a second time. Their coital times ranged from 26 to 55 seconds with a mean of 28.8 ± 6.8 seconds. However, 48 out of 50 concurrent virgin controls (96%) force-copulated. The coital times of these controls ranged from 12 to 58 seconds, with an overall mean of 25.0 ± 1.3 seconds.

DISCUSSION

Aedes aegypti females which are less than 24 hours old can be inseminated by force-mating (this paper), but none are inseminated at this age within a cage (Lea, 1967, 1968; Gwadz, 1967). Spielman *et al.* (1967) suggested that (1) semen is altered during or immediately after insemination of virgin females and that (2) semen is lost, if not so altered. Thus, it is possible that very young females mating in a cage do not yet have the ability to alter semen, and therefore do not retain it. Since the aedeagus blocks the genital orifice during forced-mating about 5 times longer than during cage-mating, it is possible that the retention of the semen in young force-mated females is due to the fact that it has time to become self-altered, whereas the semen in young cage-mated females flows out of the bursae before it has time to become self-altered. It is known that the male accessory gland material alone gels very rapidly in saline *in vitro* (Jones and Wheeler, 1965a). Since Jones (unpublished data) noted that the bursal walls lack the machinery of secretory cells, the bursa probably does not secrete anything which would alter the semen.

It is also conceivable that the bursae of very young females differ in several respects from those of older females. The very young females are swollen by air and fluids in the gut (Christophers, 1960) and this might distort the bursae. If during the relatively prolonged period of forced-mating, the pairs were more firmly united than during the briefer period of cage-mating, then a longer and firmer union might permit the force-copulating male to fill the bursa of a very young female, whereas a shorter and less firm union might not permit the free male to do so adequately.

Further, there may be a difference in the coital mechanics between forced-copulation and cage-mating of the young females. The descriptions of coitus by Jones and Wheeler (1965b) and Spielman (1964) differ enough to suggest that this is a very real possibility. The male may be unable to grasp the young female properly in cage-mating but can do so when force-copulated. If the cage-copulating male does not grasp the young female correctly, then (1) the male may only appear to cage-copulate with the young female but not ejaculate, or (2) the aedeagus of the free male may not be able to attach to the dorsal vaginal valve properly so that the ejaculate is not deposited in the bursa. The phallosome's position must

be exactly within the coital cavity of the inverted upper vagina or the female will not be inseminated properly (Jones and Wheeler, 1965b).

The occurrence of prolonged coitus in cage populations of *Ae. aegypti* has never been reported, yet it was observed in 7 out of 85 (8.2%) force-copulations with young females. This phenomenon may provide a means of determining if the aedeagus blocks the opening to the vestibule (the funnel through which sperm enter the spermathecal ducts). The factors involved in the termination of coitus are not well known. It is believed that the aedeagus overextends and that the connection with the dorsal vaginal valve is easily broken. The aedeagus is then free and pushes against the male's own anal cone. The female then dislodges the male with her legs (Jones and Wheeler, 1965b). In prolonged forced-coitus, the gelation of the male accessory gland material might hold the aedeagus *in situ*.

Very evident in this study was the high degree of variability of the sexual responses of males that had been reared under nearly identical conditions and which were given presumably similar stimuli. Sometimes a male would not copulate with one virgin female but would with a different female. The coital responses shown by individual males appeared to vary between females. Some individual males are certainly more potent in that they are highly active and will copulate sequentially more than 6 times. Although it could be argued that a series of virgin females differ substantially in their attractiveness to a given male, this seems unlikely. Often, in separate tests, females that were actively refused (as indicated by withdrawal of the male's terminalium) by different males were subsequently acceptable to other males. The mean precoital times of 16.0 ± 1.2 seconds for forced-copulation are short compared to the means of 34.1 ± 4.2 , 45.0 ± 4.7 , and 47.5 ± 6.5 seconds reported for mating in lantern chimneys (Spielman *et al.*, 1967). In the latter study, the time required for flight and tarsal contact probably accounts for the longer interval. The coital time (mean 20.2 ± 0.7 seconds) of the males repeatedly force-copulated agrees with the 19.7 ± 6.4 seconds mean reported by Spielman (1964) for cage-copulated males that had been previously mated twice.

The mean coital time of 26.7 ± 1.3 seconds for 48-hour-old virgins that were force-copulated is higher than the means of 13.7 ± 3.2 seconds (Spielman, 1964), 17.6 ± 0.5)Spielman *et al.*, 1967), and 13.2 seconds (Jones and Wheeler, 1965a) for 3- to 7-day-old virgins that were cage-mated.

The mean of 25.3 ± 0.8 seconds of Ludlam (1965) and the mean of 26.7 ± 1.3 seconds in this study for coital times by forced-copulation agree; they differ from the longer mean of 31.3 ± 1.6 seconds reported by Jones and Wheeler (1965a).

This study strikingly supports the hypothesis that in the Bangkok strain of *Ae. aegypti*, cage-mated females cannot be subsequently force-copulated with males. It was not possible to force-copulate 99 out of 100 cage-mated females. This poses the question of whether the male can distinguish an inseminated from a virgin female. Clasping reflexes were generally present but aedeagal contact was never made with the cage-mated females when the forced-copulation technique was used. Ludlam (1965) did not observe even the clasping reflexes with males of this strain. Although it is conceivable that chemical stimulus is involved here, there is no evidence to support this. On the other hand, there is a possible morphological difference between virgin and previously copulated females. In

the latter, both the cerci and the eighth abdominal segment appeared to be more deeply retracted, and this would make it difficult for the male to seize the cerci with his claspers. This would not account for those cases where the male actively withdrew his terminalium from the female. The work of Spielman *et al.* (1967) differed from these findings, since they were able to force-copulate previously caged females.

Since most (95/100) virgin females which are 48 to 60 hours old are readily force-copulated by males of the same age, an optimum standard for evaluating the behavior of controls was established. Although 10.0% (5/50) of those females which had force-copulated 12 hours earlier did so again, it is possible that this seemingly high value is due to the fact that some of these females were not initially inseminated. More data are needed on insemination rates obtained with the forced-copulation technique.

We are grateful to Mrs. Daisy Liu for her technical assistance.

SUMMARY

1. Virgin females which are 24 hours old or less interfered with the copulatory attempts of males by kicking with their legs and emitting fluid from their ani. About 50% of these young females could be force-copulated, and 75% of these were inseminated.

2. The mean forced-coital time of young virgin females which were inseminated was 71.4 seconds. Those which were not inseminated force-copulated for a much shorter time (15.6 seconds).

3. The sexual responses of males to young virgin females were not uniform with the forced-copulation technique. Active refusal by withdrawal of the male terminalium from the female by the male was more definite with young females than with older ones.

4. Prolonged coitus was observed in about 8% of the forced-copulations with young virgin females; this phenomenon has not been reported for cage-matings nor has it been seen in nature.

5. Ninety-five of 100 virgin females force-copulated, whereas only one 3-second copulation was obtained with 100 females that had been previously caged.

6. A force-copulation rate of 10% was recorded for females that had force-copulated 12 hours previously.

7. The cerci and the eighth abdominal segment appeared to be more deeply retracted in inseminated females than in virgin females.

8. While individual males varied in their sexual responses toward similar females, 52% of them definitely responded to them and 92% of these individuals copulated.

9. Using the forced-copulation technique, precoital and coital time means were 16.0 and 20.7 seconds, respectively. The age of the male did not influence these values.

10. The number of copulations by 5- and 7-day-old males was significantly higher than that by 1- and 3-day-old males.

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EFFECTS OF SEMI-STARVATION ON GROWTH AND MORPHOGENESIS DURING THE LARVAL STAGES OF A COMMON MILLIPED, *NARCEUS ANNULARIS* (RAF.)¹

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The life-cycle of the Diplopoda is divided into three phases: the egg, the larva, and the adult. In *Narceus annularis* (Raf.), the common northern spirobolid millipede, the egg is enclosed in a capsule fashioned out of regurgitated leaf litter by the female at the time of egg-laying (Barber, 1915). The first two instars and at least three days of the third instar are spent within the micro-environment of the egg capsule (Berns, 1968). Following emergence from the egg-capsule, the millipede goes through at least seven additional instars before sexual maturity is attained.

In most arthropods, the passage through the larval instars is mainly related to growth in size (Wigglesworth, 1965). The morphogenetic and differentiative processes (with the exception of reproductive structures and regeneration) are mainly confined to the egg, and to the pupal stage in the holometabolous insects. In millipeds, however, passage through the larval instars is not only characterized by growth, but also by a progressive increase in the number of body segments (in *Narceus annularis*, from 7 segments in instar 2, to 51–59 segments in the adult), limbs, and ocelli. Therefore, each instar is a morphogenetic and differentiative phase.

Preliminary observations (Keeton, 1960) suggested to us that environmental factors, particularly food supply, can strongly influence, in a quantitative fashion, the developmental changes occurring in each larval instar. Consequently, we set out to determine the effect of diet on growth, segment addition, and ocelli addition at each instar, as well as the number and duration of the instars. We were particularly interested in discovering which effects of semi-starvation would carry over into the sexually mature adults, and which effects would disappear as a result of developmental compensation. We also hoped to find characteristics that would be reliable in determining the instar of larvae.

MATERIALS AND METHODS

Adult females of *Narceus annularis* were collected from the field and placed in plastic boxes with a maximal supply of leaf litter for food. Nine hundred third instar larvae from eggs laid by these females were used in this study. Four hundred and fifty of these larvae were placed in 15 covered plastic boxes (12" × 4" × 6"), 30 animals per box, with a maximal supply of leaf litter for food. The other 450 larvae were placed in 15 boxes and were given only fecal pellets

¹ Financed in part by grant GB 3150 of the National Science Foundation.

mixed with soil (1:1) for the first six months. After six months, these larvae were given once a month an amount of leaf litter consumable in one week; they were fed only fecal pellets the remaining three weeks of each month. This modification of the original diet was necessary to prevent excessive mortality. Both groups of animals, well-fed (indicated by + elsewhere in this paper) and semi-starved (indicated by -), were maintained in the same room, at the same temperature and humidity.

Larvae from both groups were examined in each instar for body length and width, ocelli number, number of segments, and gonopod development. Records were also kept of the duration of the instars, and the total number of instars to sexual maturity.

RESULTS

Body length and width

Length and width measurements (to the nearest 0.1 mm.) of specimens preserved two to three weeks following molting are presented in Figures 1 and 2. Length measurements were made from the tip of the head to the tip of the anal valve; width measurements were made by measuring the diameter of the fourth body segment.

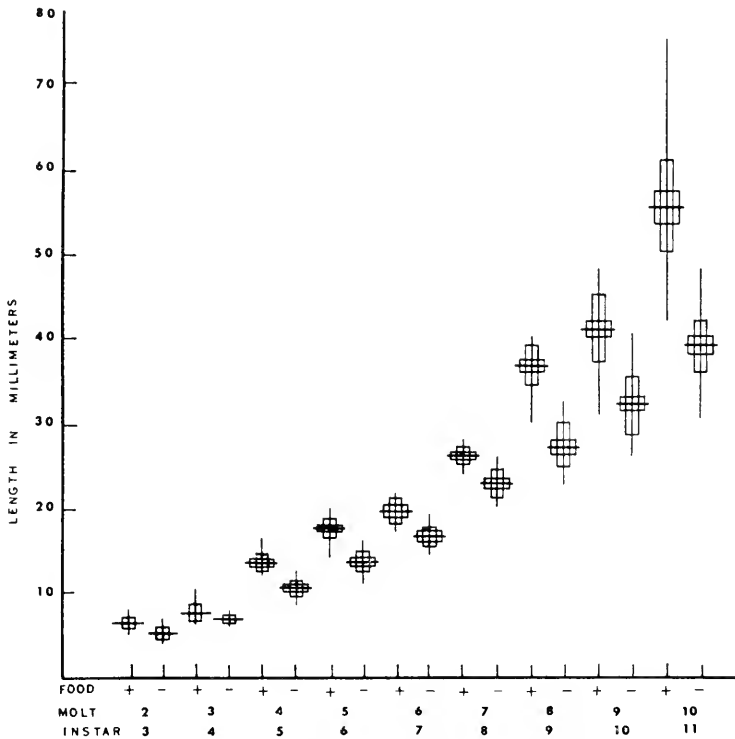


FIGURE 1. Length measurements of well-fed and semi-starved larvae.

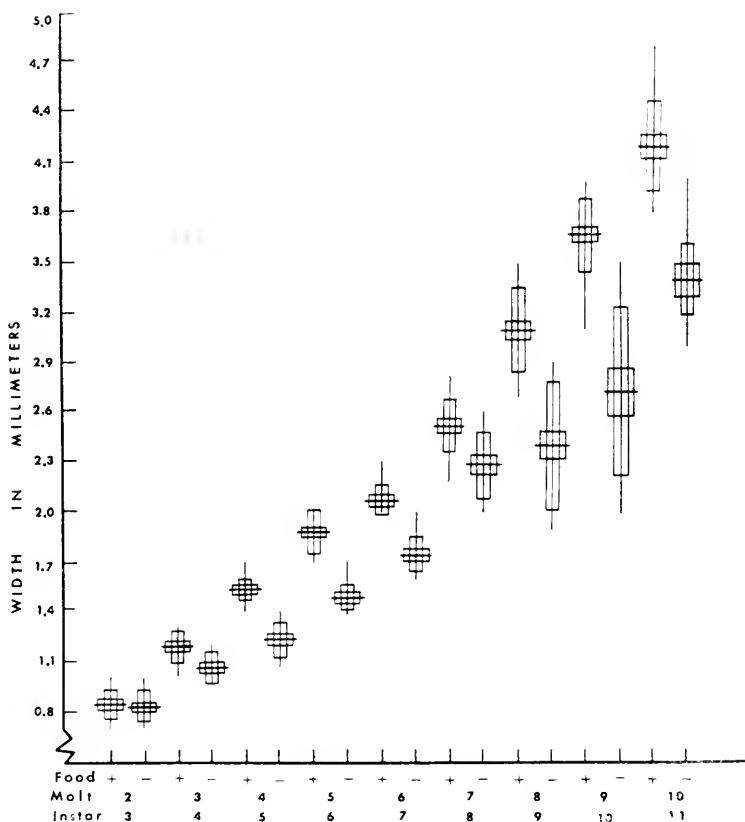


FIGURE 2. Width measurements of well-fed and semi-starved larvae.

The mean values for each instar, both length and width, are consistently lower for the semi-starved animals. In only two instars are the values not statistically different: the mean width of instars 3(+) and 3(-), and instars 4(+) and 4(-). All other means are significantly different at the p 0.001 level (t test), except for length in instars 3(+) and 3(-), and in 4(+) and 4(-), which are significant at the p 0.005 level.

It is clear that diet can affect the overall size of the animal, and therefore these two measurements (length and width) are not satisfactory indices for determining the correct instar. For example, the mean length and the distribution of length values for instar 11(-) are actually less than those for instar 10(+). The mean width of instar 9(-) is less than the mean width of instar 8(+). These are only two examples; other cases of overlap are evident in Figures 1 and 2.

Total number of body segments

The total number of body segments for both well-fed and semi-starved animals is presented in Figure 3 and Table I. The general pattern of segment increase is

similar in both groups of animals. However, there appears to be a difference in the early rate of segment increase from instars 4 through 6; after instar 6 the difference between the two groups does not increase. In fact, there is a slight convergence in instars 9 and 11.

Analysis of the mean number of segments per instar was carried out using the "t" test. Comparison of the means of the two groups demonstrates a statistical difference ($p < 0.001$) in instars 5 through 11. There are instances where the mean segment number for an instar of semi-starved larvae is actually less than the mean for an earlier instar of well-fed larvae; for example, instar 11(-) is lower than 10(+). There is also considerable overlap in the ranges (Table I) of instars 9(-) and 8(+), 8(-) and 7(+), 7(-) and 6(+), and 6(-) and 5(+).

Adults of both the semi-starved and well-fed groups are within the normal range of segment number for the species (51-59; Keeton, 1960) even though the mean of the semi-starved (53.69) is less than the mean of the well-fed (54.91). The mean of the well-fed group is very close to the overall mean for the species, 55.1 (Keeton, 1960).

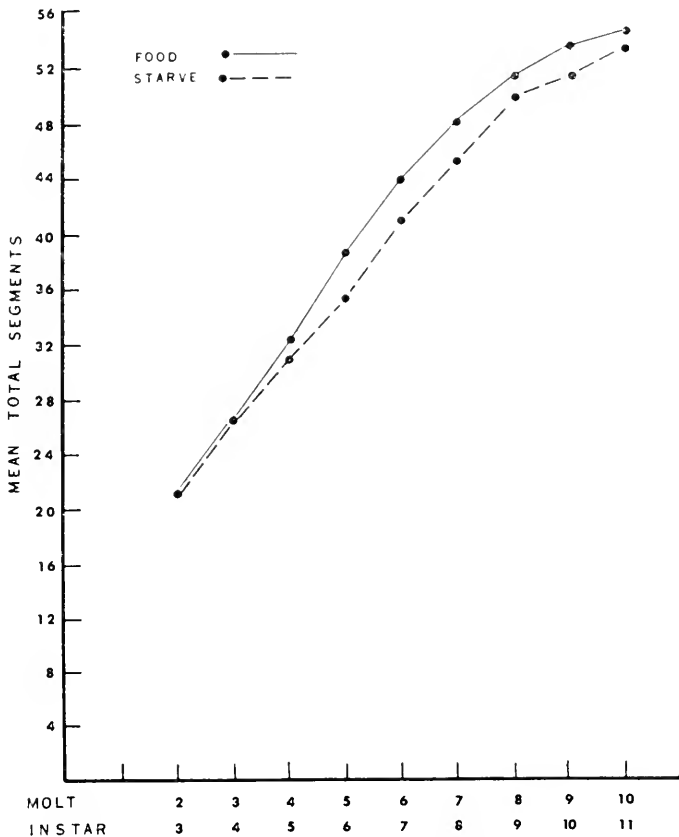


FIGURE 3. Mean total segment number of well-fed and semi-starved larvae.

TABLE I
Segment number per instar (well-fed +, semi-starved -)

Instar	Molt	Mean segment number	Standard deviation	Standard error mean	Range limits	N
3+	2	21.33	0.65	0.19	21-23	12
3-	2	21.00	0.0	0.0	21	13
4+	3	26.67	0.57	0.11	25-27	24
4-	3	26.54	0.80	0.16	25-28	24
5+	4	32.52	0.64	0.13	31-34	25
5-	4	30.92	0.69	0.14	30-32	24
6+	5	38.75	0.78	0.15	37-40	28
6-	5	35.40	1.10	0.25	33-37	20
7+	6	44.10	0.82	0.14	43-46	30
7-	6	41.11	1.40	0.23	38-44	35
8+	7	48.30	0.93	0.19	46-50	23
8-	7	45.52	0.92	0.18	43-48	27
9+	8	51.65	1.40	0.32	49-54	20
9-	8	50.10	1.17	0.26	48-53	19
10+	9	53.77	1.35	0.26	51-57	26
10-	9	51.65	1.74	0.43	49-54	17
11+	10	54.91	1.34	0.28	52-57	22
11-	10	53.69	1.32	0.26	51-55	26

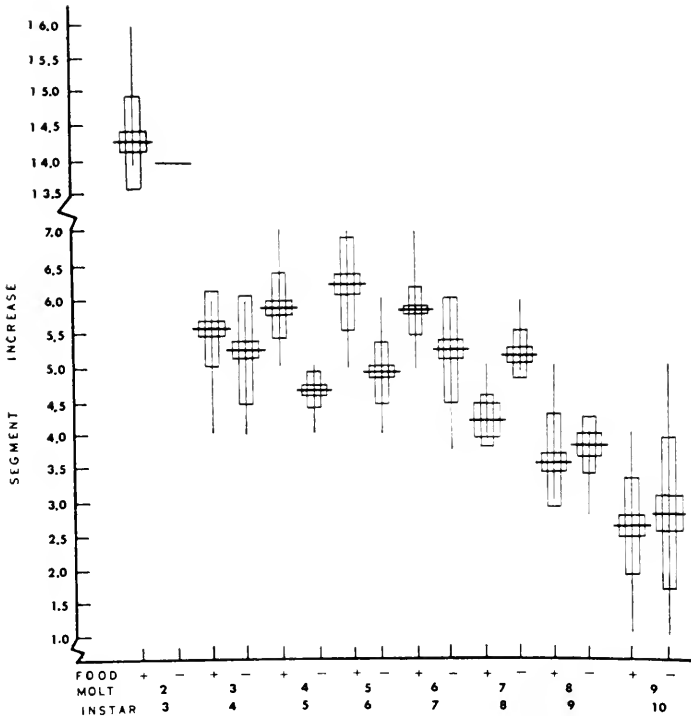


FIGURE 4. Segment increase per instar of well-fed and semi-starved larvae.

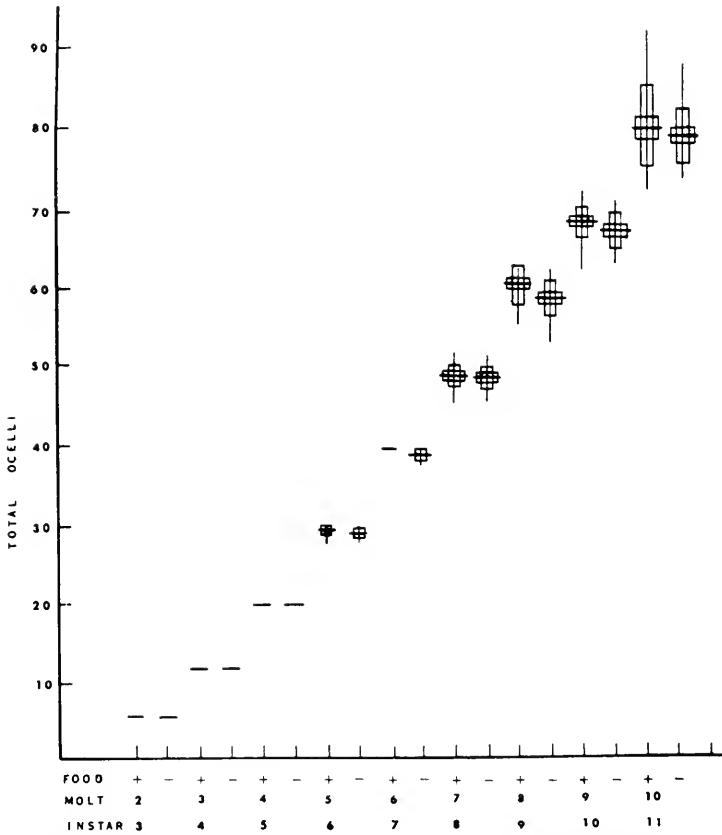


FIGURE 5. Ocelli increase per instar of well-fed and semi-starved larvae.

Segments added per instar

The effect of diet on segment addition is presented in Figure 4. At molts 4, 5, and 6, the well-fed larvae added more segments than the semi-starved (t test, $p < 0.001$). At molt 7, however, the semi-starved larvae added more segments than the well-fed ones ($p < 0.001$); the same trend was apparent at molts 8 and 9, though the difference was not statistically significant ($p < 0.3$ and 0.5).

The new segments develop from a posteriorly located proliferative region (located between the penultimate segment and the anal segment) during the first few days of the instar preceding the molt at which they first become outwardly visible (this was confirmed histologically, and will be the topic of another paper). For example, new segments exposed by the molt (molt 4) to instar 5 actually developed during the first three days of instar 4. Consequently, since the early instar larvae are enclosed within the egg capsule until the fourth day of instar 3, dietary effects on segment addition could not be detected until the molt to instar 5; the segments added at molt 3 (from instar 3 to 4) were originally determined during the first few days of instar 3. At this time both starved and well-fed larvae were still in the egg capsule.

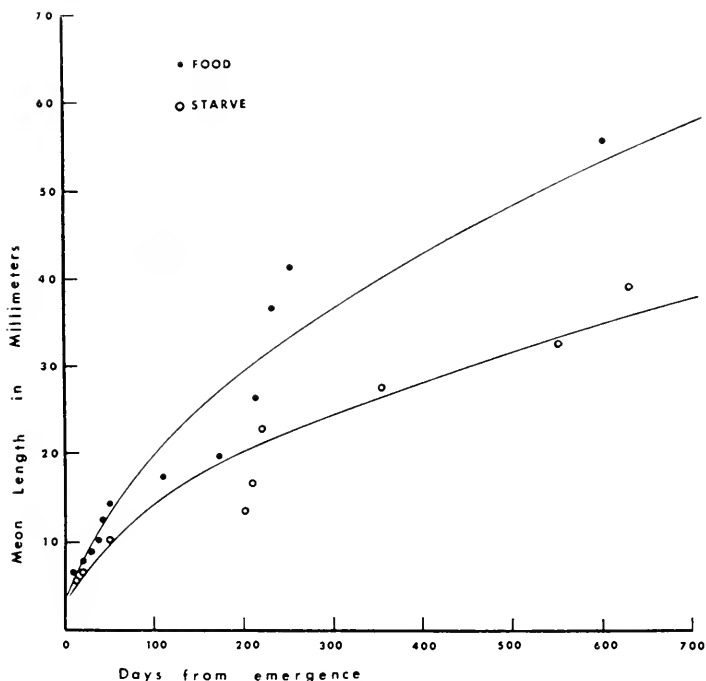


FIGURE 6. Mean body length in time of well-fed and semi-starved larvae. Each point is the mean of at least twenty animals.

Number of ocelli

Data on the number of ocelli characteristic of each instar are presented in Figure 5. In no instar is there a statistical difference between the means of the well-fed and the starved animals. In instars 3, 4, and 5, the means of the two groups are identical; the means of the remaining instars are practically the same. However, it is interesting to note that when there is a difference, the starved animals always have the lower means. Of particular significance is the lack of overlap in the ranges of the ocelli numbers for the different instars. Though the variability of ocelli number increases in the later instars, the ranges remain discrete.

A linear relationship between ocelli and number of molts is evident in Figure 5. It is not implied, however, that the correlation is between time and ocelli. On the graph, the instars are equally spaced; in reality, the time between instars is irregular and often not constant for animals in the same instar.

Number of instars

The several maturational sequences observed for *N. annularis* are shown in Table II. Though the most frequent sequence for both the well-fed and the semi-starved animals was type 3, in which there are 9 segment-building instars, two maturational instars, and maturity following the molt to instar 12, several variations were found. In particular, 25 semi-starved animals went through one

three or more instars. Box No. 20 had animals from instar 7 through instar 10, at 535 days from emergence. Six of 13 boxes of well-fed animals contained larvae in two instars; no boxes had animals in more than two instars. At 535 days, three animals in the well-fed group (box 12) were in instar 11, while one animal in the semi-starved group (box 20) was in instar 7. At this age, then, animals were in any of five different instars, almost half the total number of instars the animal goes through. Similarly, at about 180 days there was a four-instar spread, and equally large spreads were found for other ages.

In an attempt to get some index of development in time, data are presented plotting body length against age in days (Fig. 6). It appears that both groups (semi-starved and well-fed) have similar growth curves even though the starved are always lower than the fed. However, the periods between 100 and 250 days

TABLE IV
Box variability of well-fed millipeds

Box no.	Days from emergence	Instar								
		3	4	5	6	7	8	9	10	11
1	20	0	24							
2	40			24						
3	110				28					
4	173					25				
5	180					5	5			
6	209						7	1		
7	212						8			
8	246						4	9		
9	253							7		
10	265							2	14	
11	533							9		
12	535								10	3
13	549								4	14

are very irregular. In order to make a more meaningful comparison the data have also been plotted on double-log paper (Fig. 7).

The above data are only good enough to point out trends and trend differences between the two groups. The data are not good enough to be useful in predicting size changes.

Maturation of gonopods

The male copulatory organs, the gonopods, undergo definite morphological changes as the millipede passes through the instars. Regardless of the size of the animal, the complexity of the gonopods for a given instar is always the same; it is, therefore, possible to characterize an instar by the morphology of the male copulatory organs (Berns, in press). It must be pointed out, however, that examination of the gonopods of a living larva often results in permanent injury or death. Furthermore, because of the simplicity of the female copulatory apparatus, gonopod staging is only possible in males.

DISCUSSION

From the preceding data it is clear that diet affects both the growth and morphogenetic processes that occur during the larval development of *N. annularis*. The growth effects are reflected in both length and width. However, since an effect on the number of segments is also demonstrated, one cannot fully attribute the length differences to growth alone; one might expect animals with fewer segments to be shorter. Width, however, is a good index of growth.

Diet has an effect on the number of segments built during an instar and on the number of segment-building instars. Semi-starvation results in addition of fewer than the normal number of segments in the early instars, and greater than

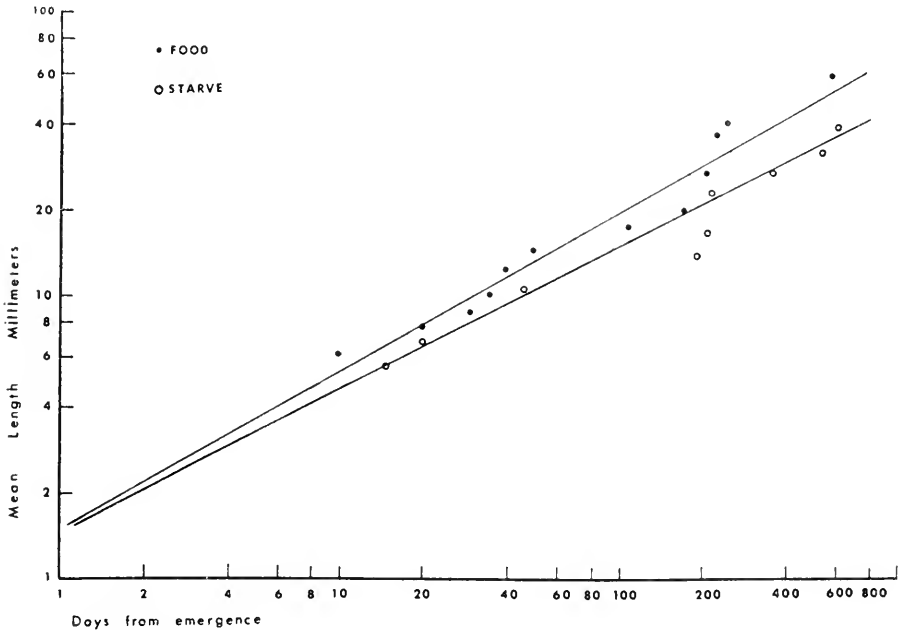


FIGURE 7. Double log plot of mean body length in time of well-fed and semi-starved larvae. These are the same data as plotted in Figure 6.

normal number in later instars. The increase of segment-building in the later instars, even though the animals are still on a semi-starvation diet, reflects the regulatory ability of the milliped segment-building system, which is also seen in the addition of extra segment-building instars by a significant number of semi-starved animals. The result of the segment regulation, whether by increased segment-building in the later instars, or by the addition of extra segment-building instars, was that semi-starved animals achieved a total number within the normal range for the species. However, because of the effects of diet, segment number cannot be used to characterize larval instars, despite the fact that this procedure has been used by most workers in the past (Loomis, 1933; Causey, 1955; Keeton, 1960).

The lack of a definite number of instars is not surprising in view of similar irregularity in other arthropods, but the addition of the extra instars in the milliped appears to be related primarily to prolonging the morphogenetic process of segment-building, rather than to attaining of a required size. In fact the data demonstrate that starved millipeds mature regardless of size, as long as the segment number is within the normal range. Nutrition has been shown to influence the growth rate and the number of ecdyses of several insects: *Blatella* (Seamans and Woodruff, 1939), *Tineola* (Titschack, 1926), and *Bombyx* (Bounhiol, 1938). In these cases, however, size seems to be the critical factor for maturation.

Effects of other environmental factors on the growth and molting of arthropods are well documented (Wigglesworth, 1965). Factors such as temperature have been shown to both increase the number of molts of *Sphodromantis* (Przibram and Megusar, 1912), *Dermestes* (Kreyenberg, 1929), and *Tenebrio* (Ludwig, 1956), or decrease the number of molts in *Melanophus* and *Picris* (Hoskins and Craig, 1935). Humidity, likewise, can effect the number of molts in *Thermobia* (Sweetman, 1934).

Not only is the number of instars variable in the milliped, but the time spent within the instars is even more variable. Semi-starvation merely exposes the limits of this variability; at 535 days of age, animals could be found in any of five different instars. Likewise, an attempt to correlate age with size was unsuccessful.

The only characteristics examined that were not significantly affected by diet were the number of ocelli and the morphological development of the male copulatory apparatus. In *Narceus annularis*, then, ocelli number is the most reliable staging characteristic in live specimens of both sexes.

SUMMARY

1. Growth, as measured by length and width, is substantially affected by diet. The length and width measurements were significantly lower for larvae in the semi-starved group.

2. The total number of body segments was statistically lower in the semi-starved larvae in instars 5 through 11. The difference was manifest at molts 5 and 6, remained constant at molts 7 and 9, and decreased slightly at molts 8 and 10.

3. Even though the mean number of segments was lower for adults of the semi-starved group, the range of adult segment number of the semi-starved animals was the same as the well-fed group and the same as the range given for the species.

4. Starvation caused the addition of fewer than normal (as compared to the well-fed group) segments in the early instars and a greater than normal number of segments in the later instars. Similarly, a significant number of semi-starved animals passed through either one or two extra segment-building instars.

5. The time spent in instars 4 through 11 was so variable that it was impossible to establish an index of development in time. Similarly, growth in time was most variable.

6. Diet did not have a significant effect on the number of ocelli added in each instar, or on the maturation of the male copulatory organs. These two characteristics are suitable characteristics for determining the instar of larvae, though the gonopods are difficult to examine in living specimens.

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EFFECT OF JUVENILE HORMONE ON THE SYNTHESIS AND ACCUMULATION OF A SEX-LIMITED BLOOD PROTEIN IN THE POLYPHEMUS SILKMOTH

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Juvenile hormone (JH) plays a key role in insect development. As long as immature insects produce and release this agent, they do not mature. When they no longer release JH, or release it in small amount, they metamorphose. The effect of JH on the maturation of immature insects is seen when a JH extract (Williams, 1956), or pure JH (Röller *et al.*, 1967; Meyer *et al.*, 1968) is injected into a pupa of the Polyphemus silkmoth, *Antheraea polyphemus*. When a normal pupa is allowed to develop in the absence of juvenile hormone, it molts into an adult moth about 18 days after the first signs of apolysis. However in a JH-injected pupa, adult development is blocked, and the pupa precociously molts into a so-called second pupa 10–12 days after the injection (Gilbert and Schneiderman, 1960). During the pupal-pupal molt, the insect's epidermal cells, instead of secreting adult cuticle, secrete a cuticle which closely resembles the original pupal cuticle. Also the morphogenetic death of many of the epidermal cells of the wings and antennae, which accompanies normal adult development, does not occur.

Because of its effects on metamorphosis, JH might be expected to influence the synthesis, appearance, or disappearance of specific proteins. The present report analyzes the effects of JH on the synthesis, release and accumulation of a specific protein found in the blood of female Polyphemus pupae.

MATERIALS AND METHODS

Female pupae of *Antheraea polyphemus* that had been chilled at 2° C. for six months were removed from the cold and anesthetized with carbon dioxide. Each animal was bled through a cut in the facial region between the right wing and right antenna. About 20 μ l. of blood was squeezed through this cut into a 0.5-ml. polyethylene microcentrifuge tube containing a few crystals of phenylthiourea (PTU) (Schmidt and Williams, 1953). Immediately after the blood was collected, the wound was sealed with melted paraffin. The blood samples were centrifuged for two minutes in a Beckman microfuge at 12,500 *g* and stored at -10° C.

The animals were injected with a JH extract prepared according to the method of Gilbert and Schneiderman (1960). Some of the operated animals were injected with 75 JH units/g. pupal weight (*i. e.*, 150 mg./g. pupal weight of a 1:1

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mixture of JH:peanut oil (PO)). Control animals were injected with 150 mg./g. pupal weight of PO, or were uninjected. Blood samples were collected from the same animals on consecutive days during the pupal-pupal molt and during the pupal-adult transformation. On each succeeding day, 5–10 μ l. of blood was collected through the original wound. After each such operation, the wound was immediately resealed. JH-injected animals were bled every day until they molted into second pupae 8–10 days after the injection. Control animals were bled daily for 17 to 21 days. The fat body and ovaries were removed from all animals at the end of the experiment and were stored at -10° C.

The proteins in these blood samples were separated electrophoretically, and their relative concentrations were estimated by densitometry.

All samples were examined using high pH discontinuous electrophoresis as described by Davis (1964) and Ornstein (1964). Preliminary experiments, in which the gel concentrations and the amount of protein applied to each gel were varied, indicated that the desired proteins were best resolved by electrophoretically separating 0.5 μ l. of blood in 4.5% gel. This quantity of blood was applied to the gel column as 10 μ l. of a 1/20 solution of blood in insect Ringer solution (Ephrussi and Beadle, 1936) and separated electrophoretically.

Upper and lower gel solutions were purchased from Canal Industries, Silver Spring, Maryland. Gels were prepared according to the manufacturer's instructions. The upper and lower buffers were prepared by diluting a stock solution containing 6.0 g. Tris and 28.8 g. glycine per liter of distilled water 1:9 with distilled water.

Electrophoresis was carried out at 22–24 $^{\circ}$ C. at 3 mA/gel until the bromphenol blue tracking dye reached the bottom of the lower gel, usually 50–60 minutes. Following electrophoresis, the gels were removed from their holders and stained for 30 minutes with a solution of 0.5% Buffalo Black in 7.5% acetic acid, and destained electrophoretically in 7.5% acetic acid.

The destained gels were scanned in a Joyce-Loebl densitometer at a 1:9 ratio (1 mm. of gel length = 9 mm. of record). The machine measured and recorded the density of the stained protein bands, and automatically integrated the area under the curve for each band. The densitometric data for *Polyphemus* blood proteins were compared with standard densitometric curves obtained with specific amounts of beef serum albumin, and the amount of protein in a given band was expressed as μ g. of beef serum albumin equivalents (N. G. Patel, unpublished observations). These measurements were accurate to $\pm 5\%$. The blood collected from three experimental animals and three control animals was analyzed in this way.

The fat body and ovaries of experimental and control animals were thawed, suspended in cold Ringer solution in microcentrifuge tubes containing PTU, and sonicated for five minutes in a Bronwill cup sonicator at maximum probe intensity. Microscopic examination revealed that this treatment ruptured all of the cells. The samples were then centrifuged at 12,500 *g* for five minutes. The low density fatty layer was skimmed from the top of each sample, and the supernatants were then separated electrophoretically.

In some experiments, fat body from female *Polyphemus* pupae was implanted into the abdomens of male pupae, which were allowed to develop into adults. Blood was collected from these animals, and analyzed electrophoretically.

RESULTS

1. *Effects of various treatments on development*

The blood sampling procedure had no effect on development. Both experimental and control animals showed the first visible signs of development of apolysis two days after removal from the cold. Control animals completed their metamorphosis 20–22 days after removal from the cold, whereas experimental animals completed their pupal-pupal development 8–10 days after removal from the cold.

The second pupae were covered with a pupal cuticle. Their ovaries showed no signs of development, and as far as could be determined were morphologically and histologically pupal. In contrast, the ovaries of control animals had differentiated in the manner described by King and Aggarwal (1965) although none of the eggs were covered with a chorion.

The fat body of second pupae also appeared pupal by morphological criteria. The cells of pupal fat body are held together in sheets by a connective tissue sheath. Early in adult development, this sheath breaks down and most of the fat body cells become loosened from each other. Later in adult development, the sheath reappears, and the fat body cells are again held together, this time in strands (Madhavan and Schneiderman, 1968). This sheath failed to break down in JH-injected pupae, and was retained when pupae molted into second pupae.

2. *Proteins detected in the blood of female Polyphemus pupae*

Figure 1 illustrates the protein bands that can be detected in the blood of a normal female Polyphemus pupa freshly removed from the cold. In addition to many lighter bands, whose visibility can be increased by applying larger amounts of protein and by varying the conditions of separation, the electrophoretic pattern of female blood contains two conspicuous protein bands. These are designated 4B (R_f with respect to the tracking dye = 0.47 in 4.5% gels) and 5A (R_f with respect to the tracking dye = 0.51 in 4.5% gels). Bands 4B and 5A were not resolved further in experiments which varied the gel concentrations and the amount of protein applied to each gel. For this reason, each band is thought to represent a single species of protein molecule. As seen in Figure 1, protein 4B is sex-limited. It probably corresponds to the sex-limited yolk protein antigen first described by Telfer and Williams (1953) in *Hyalophora cecropia* (their "antigen 7"). Protein 5A is not sex-limited.

3. *Changes in the concentrations of proteins 4B and 5A during development*

Densitometric determinations were made on the blood samples collected from three experimental animals and three control animals and expressed as μg . beef serum albumin equivalents. They are plotted in Figure 2, and reflect the changes in concentration of protein bands 4B and 5A. As Figure 2 shows, the concentration of protein 4B is markedly affected by juvenile hormone whereas the concentration of protein 5A is very little affected. In JH-treated pupae protein 4B may reach twice the concentration it has in control animals. Figure 3 records the ratio (protein 4B)/(protein 5A) and demonstrates that this ratio differs in experimentals and controls largely because of the changing concentration of protein 4B.

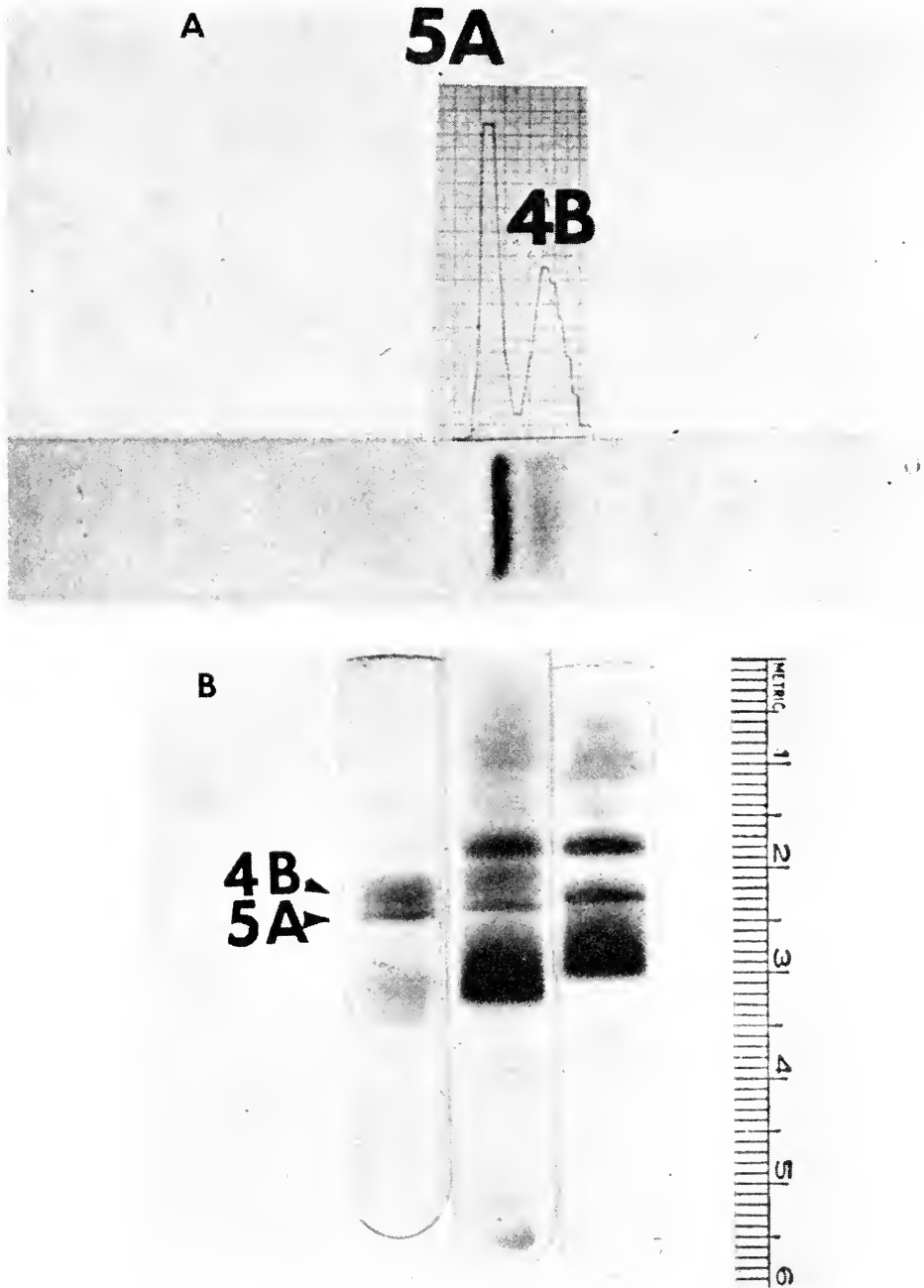
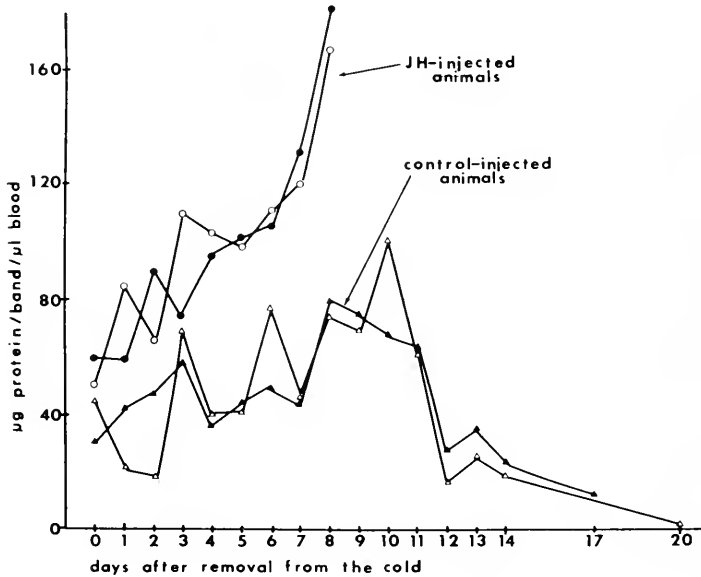
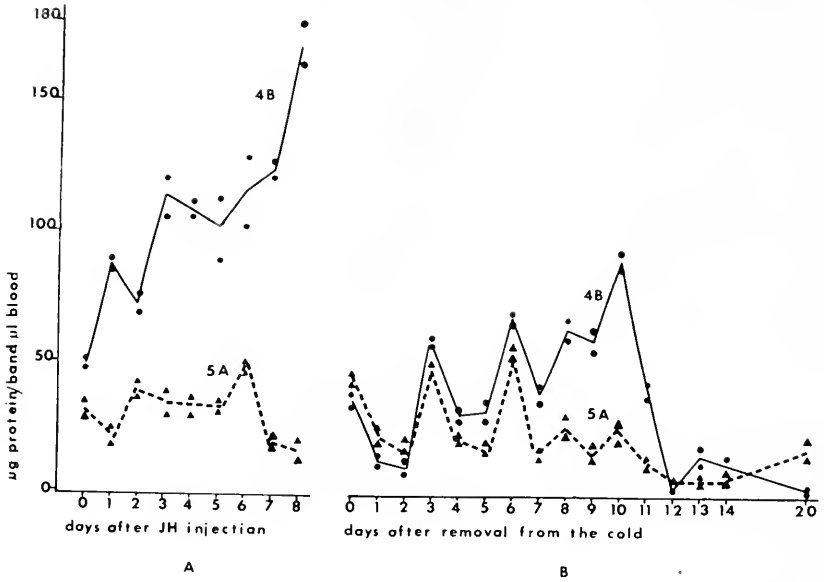


FIGURE 1. Electrophoretic separation of pupal *Polyphemus* blood in 4.5% acrylamide gels. A. Electrophoretic separation and densitometric recording of bands 4B and 5A in female blood. The densitometric values obtained were converted to $\mu\text{g.}$ of beef serum albumin-equivalent by referring to a standard curve. B. Left to right: 0.5 $\mu\text{l.}$ of female blood; 0.5 $\mu\text{l.}$ of female blood + 2.0 $\mu\text{l.}$ of male blood; 2.0 $\mu\text{l.}$ of male blood. Band 4B is not visible in male blood.



C

FIGURE 2. Changes in the concentrations of protein bands 4B and 5A in the blood of females during adult and second pupal development. A. Daily measurements of the concentrations of bands 4B and 5A in a typical animal injected with JH. Each point is based

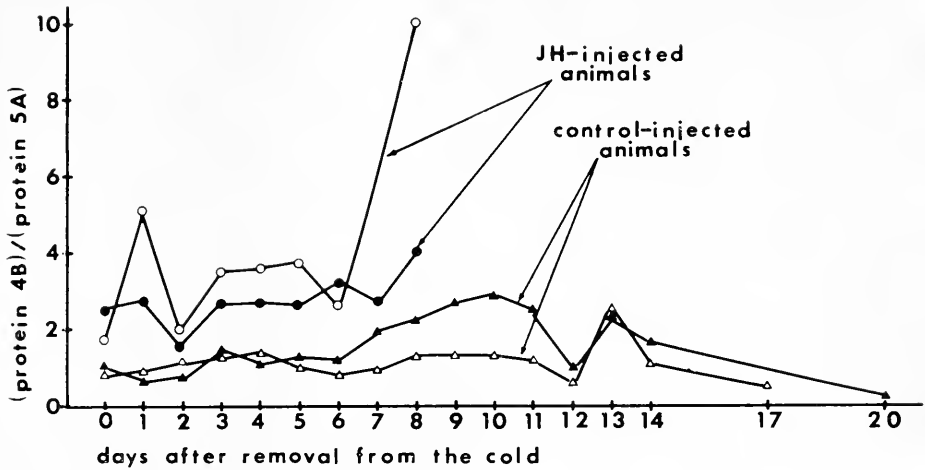


FIGURE 3. Comparison of the daily changes in the ratio of (protein 4B)/(protein 5A) in the blood of JH-injected and control animals. Each line represents one animal. Each point represents the average of the ratios of 4B to 5A.

4. Proteins in the fat body and ovaries

The results of electrophoretic separations of the protein in the fat body and ovaries of experimental animals are summarized in Tables I and II, and indicate the following:

1. The ovaries of adults contain proteins 4B and 5A. The ovaries of pupae and of JH-injected second pupae, in contrast, contain protein 5A, but do not contain protein 4B.

2. The blood of female pupae, 0–17-day developing adults, and JH-injected second pupae contains proteins 4B and 5A. The blood of adult females contains protein 5A, but does not contain protein 4B.

TABLE I

The distribution of protein bands 4B and 5A in the blood, fat body and ovaries of female pupae, developing adults, adults, and JH-injected second pupae

Band number	R_f in 4.5% gel	Blood				Fat body			Ovaries		
		P	DA	A	P ₂	P	A	P ₂	P	A	P ₂
4B	0.47	+	+	0	+	0	0	0	0	+	0
5A	0.51	+	+	+	+	+	+	+	+	+	+

P = pupa; DA = 0-17-day developing adult; A = adult; P₂ = second pupa; + = present; 0 = absent.

on densitometric measurements of one or two separate gels. B. Daily measurements of the concentrations of bands 4B and 5A in a typical control animal. Each point is based on densitometric measurements of one or two separate gels. C. Comparison of the daily changes in the concentrations of protein 4B in the blood of JH-injected and control animals. Each line represents one animal. Each point represents the average of densitometric measurements of two or more separate gels.

3. The fat body of males and females at all stages contains protein 5A, but does not contain protein 4B.

The observation that the female fat body does not contain protein 4B prompted further experiments to establish whether the fat body synthesizes and then releases protein 4B, as has been reported for other blood protein (see Discussion). Telfer (personal communication) has obtained immunological evidence that the fat body of *Cecropia* synthesizes and then releases the yolk protein, antigen 7. To confirm Telfer's observations, female fat body was implanted into the abdomens of male pupae, which were then allowed to develop into adults. Protein 4B was detected in the blood of these developing adults, whereas in normal developing male adults, protein 4B was never detected (Table II). Efforts to detect the synthesis and release of protein 4B by fat body incubated *in vitro* in blood and various media failed (Blumenfeld, 1968).

TABLE II

The distribution of protein bands 4B and 5A in the blood and fat body of male pupae, adults, and JH-injected second pupae

Band number	R _f in 4.5% gel	Blood				Fat body			
		P	A	A _f	P ₂	P	A	A _f	P ₂
4B	0.47	0	0	+	0	0	0	0	0
5A	0.51	+	+	+	+	+	+	+	+

P = pupa; A = adult; A_f = adult derived from male pupa which had received an implant of female pupa's fat body; P₂ = second pupa; + = present; 0 = absent.

DISCUSSION

1. *The fate of protein 4B*

The behavior of blood proteins in *Polyphemus* is best considered in relation to the behavior of the blood proteins of a closely related saturniid silkworm, the *Cecropia* silkworm, which has been the subject of extensive experimental analysis. The blood of the female *Cecropia* pupa contains a high concentration of a specific protein antigen that is present in very low concentration in the blood of the male pupa (Telfer and Williams, 1953; Telfer, 1954). This antigen, which they called "antigen 7," is accumulated by the developing oocytes to a concentration above its level in the blood, and disappears from the blood during metamorphosis. During this time, several other blood protein antigens that appear in the oocytes are not accumulated to concentrations above their level in the blood. Antigen 7 corresponds to an electrophoretically separable protein band (Laufer, 1960). Proteins that correspond to antigen 7 are present in the blood of other saturniid moths (Laufer, 1960). Protein 4B discussed in these experiments is sex-limited, is accumulated by the developing oocytes, and appears to correspond to antigen 7.

At most stages of adult development, a change in the concentration of protein 4B is paralleled by a change in the concentration of protein 5A. Between 0 and 5 days of adult development, the concentrations of proteins 4B and 5A varied from 30 to 50 $\mu\text{g./}\mu\text{l.}$ Between 5 and 7 days of adult development, protein 5A attained

a concentration as high as 65 $\mu\text{g./}\mu\text{l.}$, while protein 4B attained a concentration in the blood of 100 $\mu\text{g./}\mu\text{l.}$ The concentrations of both proteins then decreased as the accelerating demands of the maturing oocytes removed them from circulation. By 10 days of adult development, protein 5A had fallen to 35 $\mu\text{g./}\mu\text{l.}$, while protein 4B had fallen to 25 $\mu\text{g./}\mu\text{l.}$ After this time, they decreased further in concentration.

When one injects JH into the female pupa, one discovers that JH completely inhibits ovarian development, and that second pupal ovaries are morphologically and histologically indistinguishable from those of pupae. In most insects JH either directly or indirectly exerts a trophic effect on the oocytes and is required for both oocyte growth (see review by de Wilde, 1964) and yolk accumulation (see reviews by Telfer, 1965; Engelmann, 1968). Since extirpation of the pupal corpora allata has no obvious effect on the egg development of many non-feeding higher Lepidoptera (Williams, 1959) it is generally believed that the presence of JH during adult development is not necessary for egg development in this group. In these insects egg development occurs during adult differentiation and appears to be coupled to it. Thus, when one blocks adult differentiation with JH, one perforce blocks egg development.

Changes in the blood levels of proteins 4B and 5A are closely linked during normal adult development. When one injects JH, however, this relationship is uncoupled, and the concentrations of the two proteins diverge. In the present experiments this uncoupling was first detected seven days after JH injection. The JH injection results in a 3–4-fold increase in the titre of protein 4B from 40 to about 180 $\mu\text{g./}\mu\text{l.}$ 8 days after injection. During this time, protein 5A remains within its normal range. Telfer (1954) ovariectomized female *Cecropia* pupae, and found that the level of the yolk protein in the blood of such animals rose to more than twice its level in the diapausing pupa. The rising titre of protein 4B in JH-injected female *Polyphemus* pupae strikingly parallels Telfer's results. Indeed, as far as protein 4B is concerned, JH appears to produce the same effects as ovariectomy.

The career of protein 4B consists of periods of synthesis, release into the blood, and accumulation by the growing oocytes. JH does not affect the synthesis or release of protein 4B, but specifically blocks the accumulation of protein 4B by the oocytes. Thus, JH profoundly upsets the career of protein 4B.

2. What tissue synthesizes and releases protein 4B?

While there is conclusive evidence that the fat body synthesizes and releases specific blood proteins (Shigematsu, 1958; Price and Bosman, 1966) it has not been proven that the fat body synthesizes and releases the yolk protein of the female. Telfer (personal communication) has concluded, from immunological analyses, that the female fat body of *Cecropia* was indeed the source of this protein. In the present experiments, the appearance of the yolk protein, protein 4B, in the blood of males that had received implants of female fat body confirms Telfer's findings.

JH injection prevents the breakdown of the connective tissue sheath around the fat body, but does not block the synthesis of protein 4B by the fat body cells themselves. Hence, the data suggest that as far as the synthesis of this

particular protein is concerned the pupal fat body is not a biochemical target of JH at the concentrations we employed. There is also other evidence that JH injection does not prevent an adult-specific protein from appearing in the fat body (Blumenfeld, 1968).

One wonders why the biochemical metamorphosis of the pupal fat body is not blocked by juvenile hormone whereas the metamorphosis of the integument is so profoundly affected. The answer may stem from the fact that many insect cells appear to be most sensitive to juvenile hormone when they are about to synthesize or are synthesizing DNA (*cf.* for example Krishnakumaran *et al.*, 1967, p. 43-44 ff.). Pupal fat body engages in very little DNA synthesis during the pupal-adult transformation whereas epidermal cells do engage in DNA synthesis, hence the insensitivity of pupal fat body to juvenile hormone.

This research was supported in part by research grants from the USPHS. The senior author was supported by a predoctoral traineeship in developmental biology. We wish to thank Dr. Narayan G. Patel for advice on densitometry, Dr. William H. Telfer for several valuable discussions, and Mr. Louis Martonyi for assistance with the photography. Drs. George W. Nace and A. Krishnakumaran provided helpful criticisms on the typescript.

SUMMARY

Injection with juvenile hormone causes pupae of the *Polyphemus* silkmoth to undergo a pupal-pupal molt instead of a pupal-adult molt. Juvenile hormone also affects the concentration in the blood of a sex-limited blood protein, the yolk protein, which is normally synthesized by the fat body of the female pupa, released into the blood, and accumulated from the blood by the developing oocytes. When juvenile hormone is injected into a female pupa, the synthesis and release of the yolk protein are not affected. However, the oocytes do not grow and do not accumulate the yolk protein. Consequently, the concentration of the yolk protein in the blood of developing second pupae increases about four-fold. This effect becomes conspicuous seven days after juvenile hormone injection.

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LIGHT-INHIBITION OF FLASHING IN THE FIREFLY *PHOTURIS MISSOURIENSIS*¹

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Recent investigations of photic inhibition of flashing in the firefly *Luciola* (Magni, 1967; Brunelli *et al.*, 1968a, 1968b) have confirmed the phenomenon of inhibition *via* the visual system reported by Case and Buck (1963), who suggested that the observed inhibition was largely of central origin although they did observe, under conditions of continuous glowing, "unflashes"—dimming of continuous glowing with flash kinetics—upon electrical excitation of the light organ, raising the possibility of a peripheral mechanism. In *Luciola*, however, there appears to be a possibly humorally linked inhibition somehow involving the testis, as evidenced by transferral of inhibition across a saline bridge from a light-inhibited firefly and by disappearance of this phenomenon upon castration of the light-inhibited individual. The inhibitory process in *Luciola* was further shown to have a peripheral element by inhibition of flashes driven by electrical stimulation of the light organ upon illumination of the eyes (Magni, 1967).

Since peripheral inhibition has not readily been demonstrated in insects it is most desirable to determine the extent to which the *Luciola* inhibitory system is present in other species of fireflies.

The present report represents an unsuccessful attempt to detect peripheral inhibition in *Photuris missouriensis*. It provides additional information regarding the nature of the inhibition, originally reported in *Photuris* by Case and Buck (1963), and which seems to be wholly a central nervous system process acting on an exclusively excitatory peripheral light organ innervation.

MATERIALS AND METHODS

Fireflies were obtained principally from collectors in Kansas and Iowa, for whose supervision we are most indebted to Dr. Katherine Smalley of Kansas State College, Emporia. Until use, the fireflies were fed sucrose solution and kept at room temperature or stored at 10° C. Luminescence was detected *via* fiber optic light guides and 931A photomultipliers and recorded on an Offner Dynograph and photographically from a Tektronix 533 oscilloscope. Electrical excitation was delivered from constant-current stimulators or pulse generators. Light stimuli were delivered *via* a fiber optic guide from an incandescent or xenon arc lamp.

Localization of light stimuli was insured by mounting insects with their heads projecting into a hole in a black plastic block receiving the stimulating light guide and sealing the aperture with a mixture of powdered graphite in Vaseline. Light

¹ Supported by ONR grant N00014-67-A-0120-0002.

TABLE I
Time required to photically inhibit spontaneous flashing

Illumination lead time (msec.)	Extent of inhibition of expected following flashes (1.0 = normal; 0.0 = no flash)					
0-40	1.0	1.0	1.0	1.0	1.0	
41-80	1.0	1.0	1.0			
81-120	1.0	0.6	1.0	1.0	0.8	
121-160	0.6	0.0	1.0	0.0	0.2	1.0
161-200	0.0	0.0	0.0	0.0		
201-240	0.0	0.0	0.0	0.0	0.0	0.0
241-280	0.0	0.0	0.0			
281-960	24 expected flashes, all 0.0					

Illumination 3800 lux. Combined data collected from 3 male *Photuris* by random illumination during sequences of regular spontaneous flashing.

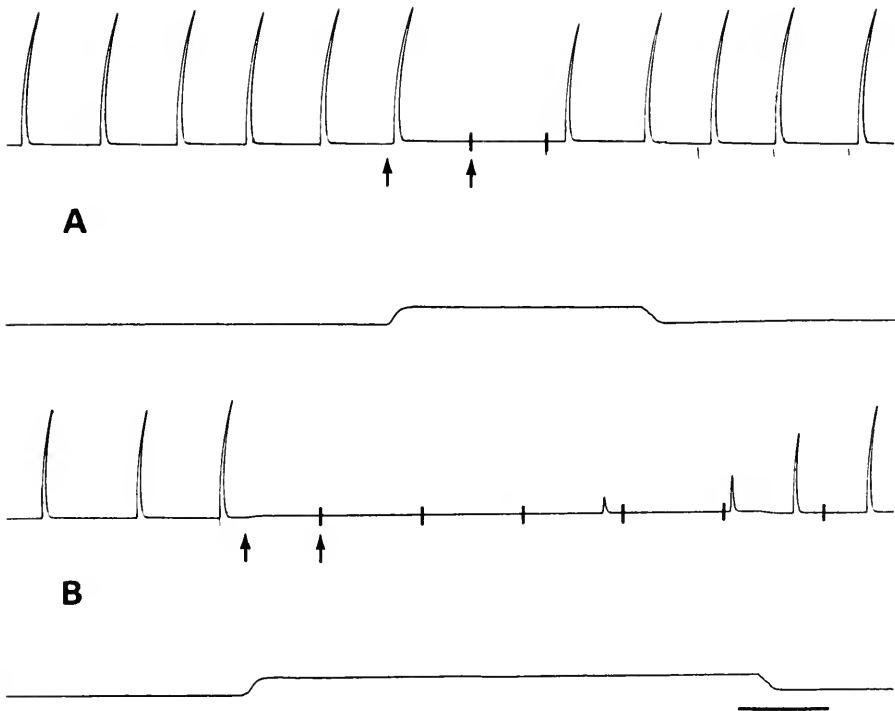


FIGURE 1. Example of records used in determination of latency of photic inhibition *via* eyes of spontaneous flashing *Photuris*. Upper trace in each record registers flashes and lower monitors light source. Latency, the interval between arrows, is measured as time between start of illumination and time of next succeeding expected flash, indicated as vertical bars, as measured from inter-flash interval of several preceding flashes. In 1A, the light goes on insufficiently early to inhibit one flash, completely inhibits the next and possibly partially inhibits and delays a third. Better examples of partial inhibition are shown in 1B. Light stimulus 100 lux, time mark at lower right, 1 second.

sources were calibrated against a Gamma Instruments standard lamp using a Reeder Thermopile and Keithley millimicrovoltmeter. Nor-epinephrine was injected with a Hamilton microliter syringe. The saline of Brunelli *et al.* (1968a) was used for preparing nor-epinephrine solutions and in the saline bridge experiments.

RESULTS

1. Inhibition of spontaneous flashing

Interruption of spontaneous flashing by electrical stimulation of the eye (Case and Buck, 1963) or by illumination of the eye (Magni, 1967) was readily confirmed in *Photuris missouriensis*. In this species the specificity of the interruption appears to be sufficient to permit use of the term inhibition in regard to the

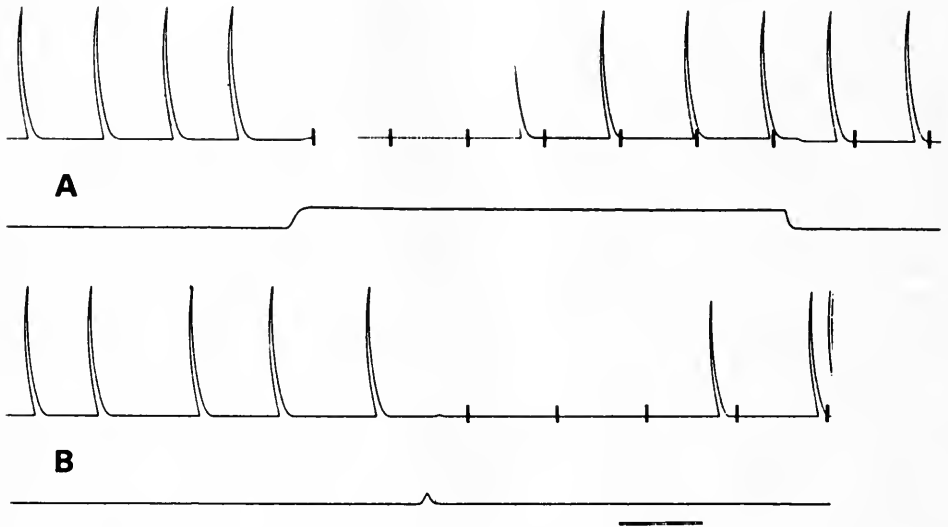


FIGURE 2. Two sequential photic inhibitions showing that a very brief $\frac{1}{2}$ -second, 50-lux flash (2B) can be as effective as a much longer, 100-lux illumination. Upper trace, light organ with some registration of stimulating lamp, which is recorded on lower trace. Time mark, 1 second.

phenomenon. Certainly, the effect is not simply an alarm reaction since *Photuris* responds to injury or disturbance with rapid flashing.

The latency of inhibition is nearly the same for electrical and photic stimulation of the eye. Our most extensive latency measurements were made with light stimuli from either an incandescent or xenon arc lamp. Light stimuli were delivered at random during episodes of regular spontaneous flashing. Latency was estimated as the time from initiation of illumination to the time of the next succeeding modified flash (usually reduced in magnitude) or completely suppressed flash (time of expected occurrence was estimated from the average interflash interval of the preceding series). Results appear in Table I and an example of the records from which the Table was constructed appears as Figure 1. Clearly, illumination at intervals as brief as 160 msec. prior to an expected flash is completely inhibitory. It is

possible to attain inhibition with a lead time of 120 msec. Lead times less than 80 msec. do not inhibit the immediately subsequent flash while they do inhibit the next and following flashes. These observations are not unexpected since Case and Buck (1963) have shown the minimum latency for excitation of luminescence by brain stimulation in *Photuris* to be 120–150 msec.

2. Effects of light intensity and duration on inhibition of spontaneous flashes

These effects are most puzzling since there appears to be a triggering effect of light superimposed upon a graded response. Triggering is illustrated in Figure 2 where an extremely brief illumination of not more than $\frac{1}{3}$ th second with maximum intensity of 50 lux resulted in as much inhibition as continuous illumination of 100 lux for six seconds. Light-dark transitions appear often to be more significant than the total illumination delivered because some *Photuris* can be maintained in the inhibited state for a longer period by a train of approximately

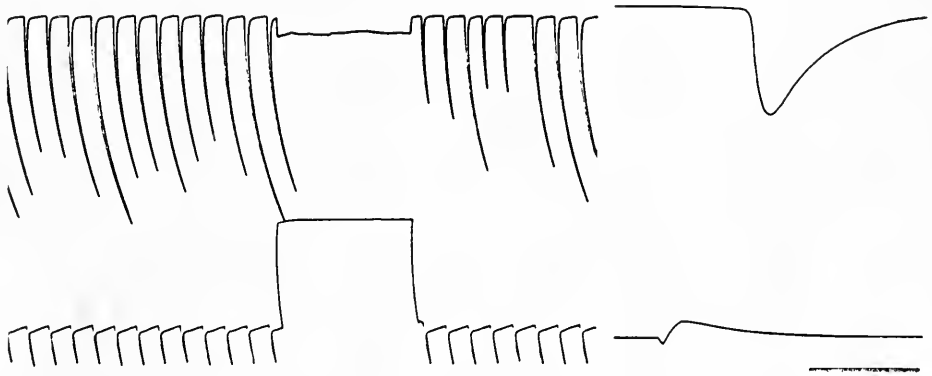


FIGURE 3. Electrical stimulation with paired electrodes in anterior light organ. Flashes recorded as downward deflection of upper trace. Stimulus (5 msec., 5 volts, 1 per second) artifacts and light (1000 lux) on lower trace. Right hand record indicates excitation-flash latency of 136 msec. Time mark at lower right applies to both records: 5 seconds, left; 200 msec. right.

one second on, one second off illuminations than by continuous illumination.

Almost invariably, during all except the most intense illuminations, flashing is reestablished. Typically (Fig. 1b) these escapes are of lower than normal intensity and at a slower frequency than prior to illumination. (The kinetics of the flashes are normal.) There is a marked tendency, however, for flashes subsequent to the first several escapes to occur at the same frequency as prior to illumination and at the same expected time as estimated from pre-illumination flashes, suggesting that illumination does not inhibit a flash pace-making mechanism.

3. Attempts at inhibition of driven flashes

Inhibition of flashes driven at the level of the light organ were unsuccessful in all instances in which the driven flashes were without CNS augmentation. An

example of what would appear to be such inhibition is Figure 3 in which a long series of flashes generated in register with 1/sec. stimulation of the light organ are terminated by illumination. However, the excitation-flash latency (measured osciloscopically) in this sequence amounts to 136 msec. While this is probably insufficient time to allow facilitative effects *via* the brain, there is clearly sufficient time for reflexive facilitation *via* the abdominal ventral nerve cord. Even if the excitation-flash latency were sufficiently short to preclude CNS effects on the immediately following flash, there is no assurance that a long-term facilitatory state built up in the CNS does not affect the flashing sequence. This is illustrated in Figure 4 where a series of voluntary flashes is mixed with driven flashes. Immediate inhibition of the voluntary component occurs upon illumination, along with a gradual diminution of the driven component. We attribute this to an effect of light on the voluntary component, defacilitating the driven flashes. This seems to be confirmed by the events after cessation of illumination in this same experi-

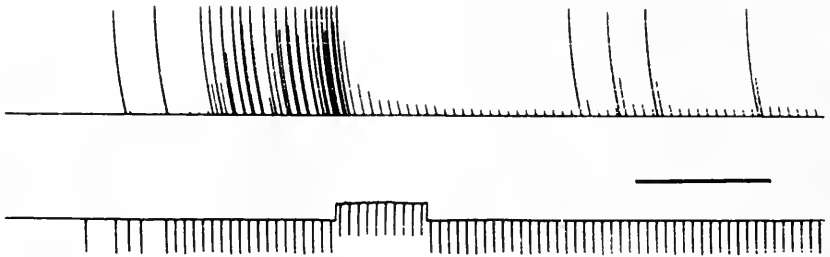


FIGURE 4. Effects of light on voluntary and driven flashes. A series of voluntary flashes, large flashes of uniform intensity, occurs along with electrically driven flashes of irregular intensity. Intensity of driven flashes is seen to depend upon temporal proximity to a previous voluntary flash. Illumination terminates voluntary component with progressive consequent diminution of driven flash magnitude. Upper trace, light organ. Lower trace, stimulus artifact (downward) and illumination artifact (upward). Time mark, 30 secs.

ment in which the driven flashes remain at low ebb until spontaneous flashing resumes. If the level of excitation had been somewhat lower, we presume that the experimental record would have created the impression of complete inhibition of driven flashes by illumination.

4. Attempts to effect transfer of inhibition from one firefly to another

The experimental arrangement used by Brunelli *et al.* (1968a) to demonstrate inhibition transfer between individuals in pairs of *Luciola italica* was duplicated in all essential particulars using males of *Photuris missouriensis*. *Photuris* was used in preference to other American species of fireflies because it most readily flashes spontaneously under experimental conditions. In five experiments with good spontaneous flashing it was not possible to demonstrate transfer of inhibition of spontaneous flashing from one *P. missouriensis* to another upon illumination of the eyes of one animal with light (Fig. 5). Since inhibition transfer occurred in the *Luciola* experiments in 8–15 seconds, we attempted to obtain an estimate of the minimal transfer time of an appropriate chemical agent from one insect to

another in our experimental arrangement. Nor-epinephrine was chosen because it rapidly induces glowing and is probably related to the normal synaptic mediator of the firefly light organ (Smalley, 1965). The time required for transfer was measured as the time between induction of glowing, measured with photomultipliers, in one nor-epinephrine-injected member of a saline-linked pair and its non-injected mate. Even where two insects were in direct physical contact *via* the wounds produced by complete severance of their terminal abdominal segments and when one member of the pair had received an injection of nor-epinephrine sufficiently large to induce instantaneous glowing in its own light organ ($8 \mu\text{l.}$ of $10^{-2} M$), the minimum transfer time was 68 seconds. One of these experiments is illustrated in Figure 6. In another series of experiments, nor-epinephrine was

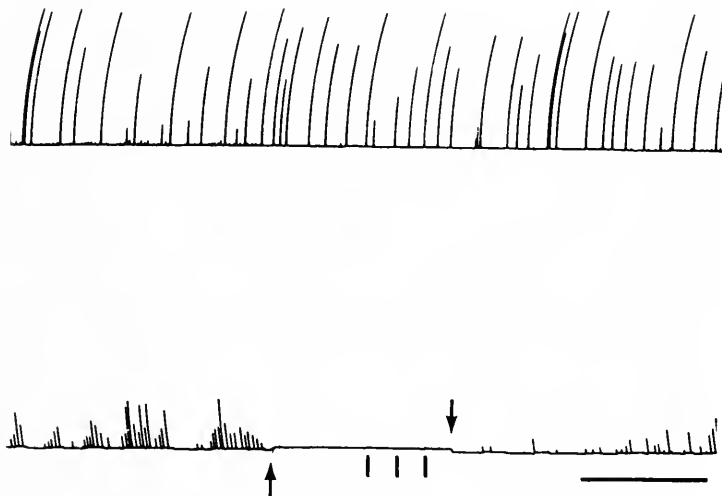


FIGURE 5. Record of an unsuccessful attempt to transfer inhibition from one light-inhibited *Photuris* male, lower trace to another, upper trace, *via* a saline bridge (see text). Light on and off indicated by arrows, 1000 lux. Three minimal escapes, indicated by vertical bars, confirm completeness of inhibition as does the long time required after cessation of illumination to reestablish rapid flashing. Upper trace shows no indication of transfer. Time mark, 30 sec.

introduced into a drop of saline in contact with the hemocoel of the second light organ segment and the time for initiation of glowing in the adjacent segment of the same animal was determined. In these experiments the minimum time required was 30 seconds. Twice this time would, we believe, be comparable to the transfer time between a pair of insects and does, indeed, correspond well with the transfer experiments just described.

5. The role of the testes in flash inhibition

Brunelli *et al.* (1968b) report that castration of the light-exposed member of a saline-linked pair made it more difficult to light-inhibit the castrated insect and made transfer of inhibition to the intact member of the pair impossible. We

attempted to further assess the role of the testis by electrical stimulation of a freshly excised pair of *Photuris* testes in a saline drop directly in contact with the light organ tissue of a spontaneously flashing male *Photuris*. Inhibition of flashing was not observed.

Next a more general experiment was conducted to determine if any structures other than CNS and light organ were involved in photic inhibition. The light organ of a spontaneously flashing male *Photuris* was dissected in such a way that its only connection with the remainder of the animal was by way of the ventral nerve cord. In this state, photic inhibition of spontaneous flashing was still obtainable.

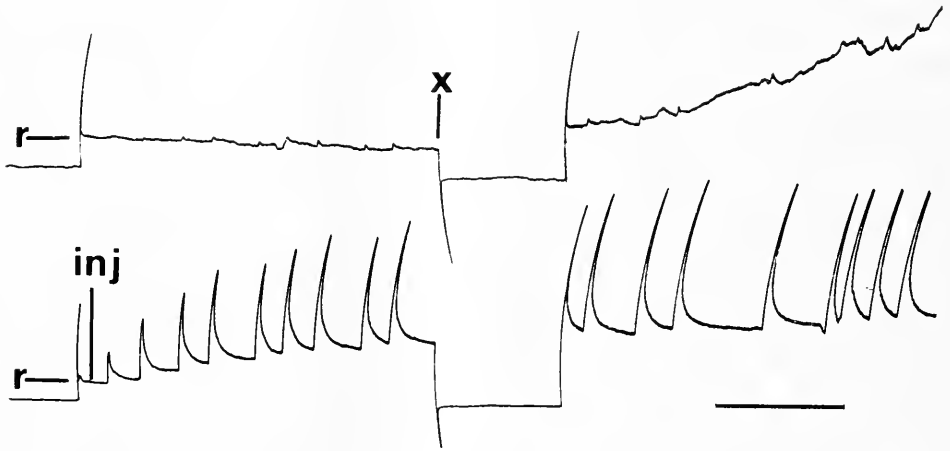


FIGURE 6. Experiment to measure diffusion time of nor-epinephrine between light organs of two fireflies with body cavities joined (see text). Lower trace records rise of luminescence, superimposed upon spontaneous flashing, upon injection at mark from rest luminescent level, r. Upper trace records luminescence from the other member of the pair. Beginning at x, the photomultipliers were turned off during visual confirmation of integrity of saline linkage. By the time recording resumed the second animal had commenced to glow, indicating arrival of nor-epinephrine and leaving the interval terminated at x, 80 seconds, as the minimum diffusion time. Time mark, 30 sec.

DISCUSSION

These data, along with the observations of Case and Buck (1963), Magni (1967) and Brunelli *et al.* (1968a, 1968b), are unfortunately subject to more than one interpretation. When one especially considers the evidence of Carlson (1961) that neural activity exerts a trophic as well as excitatory effect on the lantern, together with the observations concerning the difficulty of "arousing" flashing behavior in fireflies during the day (Case and Buck, 1963), it would seem plausible to propose two flash-suppressing phenomena, both of which are light-initiated. One of these might regulate the circadian cycle of luminescence known to occur at least in *Photinus* (Buck, 1937). The other might effect rapid control of luminescence in response to irregular, brief illuminations during the normally active period. It is plausible to suppose the former might involve a neuro-endocrine link, such as has frequently been observed in insect circadian rhythms while the latter,

because of its rapidity of establishment and of dissipation, might be supposed to be wholly neural in mechanism. The phenomena described in *Luciola* strongly suggest that both mechanisms are present. However, in *Photuris* there does not at present appear to be evidence for any mechanism except the exclusively neural one. In fact, even such phenomena as the "arousal" process and Carlsons' trophic effect might well be simply due to either diminishment of excitation or to a dark-dependent neural excitatory process, rather than peripheral inhibition.

Quenching phenomena, such as described by Case and Buck (1963) have been suggested as possible examples of peripheral inhibition of steady glowing. Since the mechanism of luminescence control *in vivo* remains unexplained, there is little restriction upon the elaboration of theories concerning peripheral processes such as this electrically driven quenching of denervated light organ glows. By analogy with other bioelectrically activated systems (see, for example, Eckert, 1966) it is plausible that light emission is related to depolarization of photocytes and this concept is strengthened by the K^+ dependence of scintillation in *Photuris* (Carlson, 1967). In glowing light organs, such as those showing the quenching effect, it might be assumed that most photocytes are to some extent depolarized. Electrical excitation of such populations with bipolar electrodes would then probably have hyperpolarizing effects in the vicinity of the anode, resulting in quenching, while cathodal effects would be minimal or non-existent, owing to the already depolarized state of the photocytes.

Resolution of the differences which appear in flash inhibitory mechanisms in *Luciola* and *Photuris* is not readily achieved. Certainly unequivocal photic inhibition of flashes driven at the level of the light organ has not yet been attained in *Photuris*. Their reported occurrence in *Luciola* may well represent a fundamental difference in light organ control in the two species. However, a possibly serious restriction to such interpretation is based on the difficulty of eliminating CNS-mediated facilitatory effects, these assuredly being inhibited by light. In the investigation of *Luciola*, this possibility was eliminated from consideration because the maximal excitatory facilitation time (driven flashes in deganglionated light organs) which is observed is 300 msec. (Buonamici and Magni, 1967, p. 332). Most likely, however, what is critical in experiments on insects with intact CNS and normal sensory input, such as specifically concerns us here, is the central excitatory state which may well remain elevated upon excitation for long periods of time. Thus Dethier *et al.* (1965) report persistence of elevations in central excitatory state of *Phormia*, in response to chemosensory input, of as long as 10 minutes. Such may contribute a sufficient level of facilitation to give false appearance of exclusively directly driven flashes in response to light organ stimulation. Abolishment of central contributions during illumination might then give the appearance of photic inhibition of peripherally driven flashing when, in actuality, the only effect might be simply a reduction in the total level of light organ excitation.

Persistence of photogenic volleys in the ventral nerve cord and light organ of *Luciola* is considered to be an argument for the existence of an inhibitory system acting peripherally to the brain (Magni, 1967). While photogenic volleys have not been recorded in our experiments, the reappearance of flashes in time with pre-inhibited flashes after and during photic inhibition is evidence that these

volleys continue to be generated, or at least that the pacemaker producing them continues to function. The difficulty with assessing the significance of photogenic volley persistence during inhibition lies, of course, in the possibility that their effectiveness in producing flashes may well depend upon other CNS-generated excitatory activity which is not recorded. Certainly the low resolution of all recordings so far made of neural activity associated with flashing does not preclude the presence of small fiber tonic activity which might well be light-inhibited.

Our inability to demonstrate humoral mediation of inhibition or involvement of the testis in inhibition is consistent with the thesis that inhibition is purely a central phenomenon acting on excitatory pathways to the light organ. These negative results can, of course, have no direct bearing on the successful inhibition transfer reported in *Luciola*, although the marked discrepancy of our nor-epinephrine transfer times and the time required to effect inhibition transfer in *Luciola* (over a minute as compared with a few seconds) suggests that factors other than simple diffusion of an inhibitory agent are concerned in the *Luciola* experiments.

Since dim, brief illuminations can bring about effective inhibition (Fig. 2), photic inhibition may play a significant role in flash communication. The flash pattern being of critical importance in communication, photic inhibition might serve to prevent the confusion which might ensue if, for example, two males close together commenced signalling slightly out of phase. Inhibition of the late male by the first flash of the other should prevent signal garbling.

SUMMARY

1. Photic inhibition of spontaneous flashing is demonstrated in *Photuris missouriensis*.
2. The minimum latency of inhibition is between 80 and 160 msec. Inhibition is expressed as completely missed or diminished intensity flashes. There is a tendency for the flashes appearing during inhibition to be later than expected but generally flashing returns to pre-inhibitory timing, suggesting continued operation of a flash pacemaker. The extent of inhibition was found to be light intensity dependent above minimal intensities at which on-off triggering effects are dominant.
3. Direct excitation of the light organ could not be inhibited *via* the visual system.
4. Transfer of inhibition from one *Photuris* to another, *via* saline bridging their body cavities, could not be accomplished.
5. These results, arguing for a purely central mechanism of inhibition in *Photuris*, are compared with those of Magni and Brunelli *et al.*, who propose both central and peripheral inhibition of flashing in *Luciola*.

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FINE STRUCTURES OF THE CARBON MONOXIDE SECRETING
TISSUE IN THE FLOAT OF PORTUGUESE MAN-OF-WAR
PHYSALIA PHYSALIS L.)¹

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The Portuguese-man-of-war has been sighted in all seas of the world and has intrigued naturalists for hundreds of years. Much is known about its morphology and general behavior. These aspects have been well summarized in the companion monographs of Totton (1960) and Mackie (1960). Totton carefully outlines the development of the animal which starts as a single hydranth-like form (with one tentacle and a small float) and progresses to a very complex form with modified hydranths budded from but still attached to the original body. He also includes interesting observations on right-hand sailing and left-hand sailing by individuals. His interpretations of the possible significance of the phenomena are at variance with those of Woodcock (1944, 1956).

Despite the long-term interest in *Physalia* very little is known of its physiology. This may be due to the fact that the animal is extremely sensitive to confinement. If placed in an aquarium, it will start degenerating in a day or two. Charles E. Lane, who has made many observations of *Physalia* on the Florida coast off the Institute of Marine Sciences, University of Miami, believes that if the extended tentacles repeatedly touch solid bottom, the animal is adversely affected (personal communication). He feels that a cylindrical sea water tank 50 feet tall, 20 feet wide and with air jets about the periphery of the top to keep the animal centered would probably solve the problem of survival in captivity.

Observations have been made on the gas content of the float by previous workers such as Schloesing and Richard (1896). It remained for Wittenberg (1958, 1960) to first describe significant ratios of carbon dioxide in the gases, ranging up to 8% of the total. This observation has been further explored by Clark and Lane (1961), Wittenberg, Noronha and Silverman (1962), Larimer and Ashby (1962) and Hahn and Copeland (1966).

Another colonial siphonophore, the bathypelagic *Nanomia bijuga* recovered from the deep scattering layers off the California coast, has been reported to possess as much as 90% carbon monoxide in its floats (Pickwell, Barham and Wilton, 1964). This form should receive further attention.

Another instance of carbon monoxide production is in the case of the Pacific bladder kelp (*Nerocystis luetkeana*). This is a kelp that may reach 85 feet in length and have up to 4 liters of gas in the stipe and bulb. Langdon (1917) reports the normal occurrence of an average of 4% (range 1 to 12) carbon monoxide in

¹This investigation was supported by National Science Foundation grant GB-676 and by U. S. Public Health Service grant GM-06836 from the General Medical Sciences Institute.

the gas. The gas cavity is sterile and therefore it was concluded that the monoxide gas is produced by the plant in its normal respiration (Rigg and Henry, 1935).

There are many references to carbon monoxide production in other forms but these deal with special circumstances. Metz and Sjöstrand (1954) recorded small amounts of the gas released by guinea pigs and rabbits, probably accounted for by the decomposition of hemoglobin. Wilks (1959) by grinding up alfalfa and putting it in a flask exposed to direct sunlight obtained traces of the gas. Westlake *et al.* (1961) by exposing moulds to various substrates were able to show that the metabolic pathway, under certain circumstances, may use 1 carbon metabolites. Siegel *et al.* (1962) found that seedlings under reduced oxygen tension (5% or less) could produce measurable amounts of carbon monoxide. Loewus and Delwiche (1963) found no carbon monoxide in the floats of the brown alga *Egrecia*

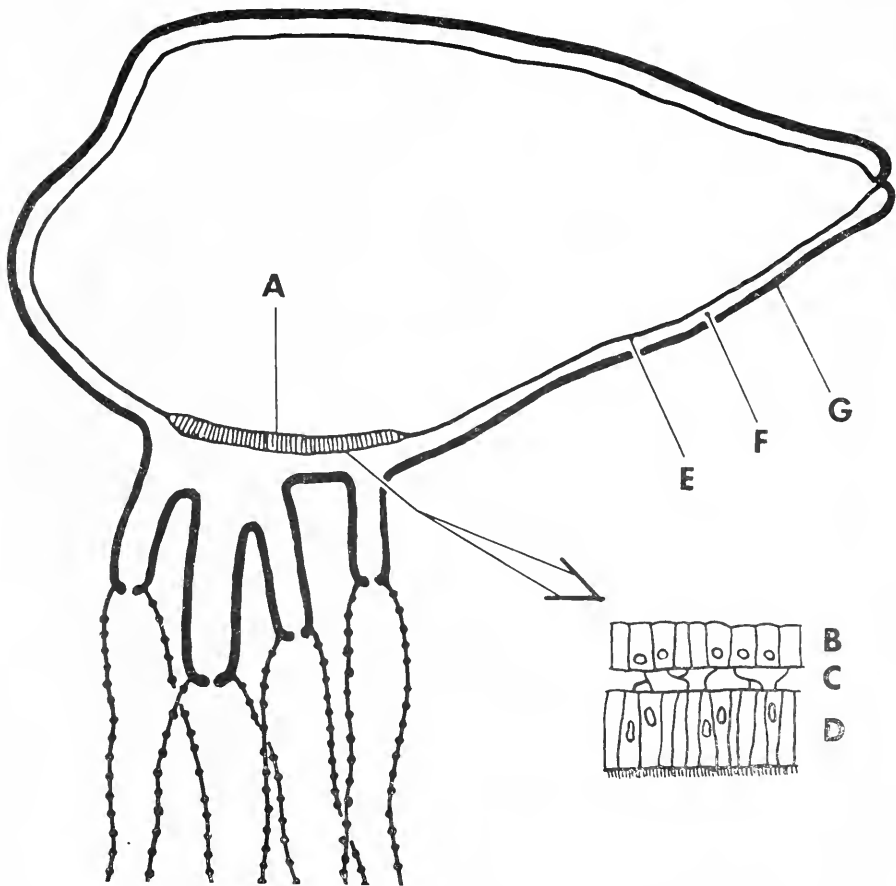


FIGURE 1. Diagram (schematic) of *Physalia* to depict relationship of the gas gland (pneumadena) to the rest of the body. A. The pneumadena (gas gland). B. Ectoderm of pneumadena. C. Mesoglea. D. Gastroderm. E. Pneumatocodon. F. Gastrovascular cavity. G. Pneumatocodon.

menezies but if they homogenized the alga, carbon monoxide could be produced by the dissociated tissues.

It would therefore appear that the siphonophores *Physalia* and *Nenomia* and the bladder kelp *Nerocystis* are the only forms known to date that produce large amounts of carbon monoxide as part of their normal physiology.

The gas is secreted in *Physalia* by a disc of tissue (pneumadena) on the ventral lining surface of the float (Fig. 1). Again, surprisingly little is known of the histology of the gas-secreting tissue. Incomplete descriptions have been given by Okada (1935) and by Mackie (1960). A description by Dahlgren and Kepner (1908) is probably erroneous.

No description of fine structure observed with the electron microscope has been reported except my own preliminary notes (Copeland, 1962, 1966). This report now presents my morphological observations to date.

MATERIALS AND METHODS

Initial observations were made on *Physalia* collected in the open Atlantic off Gay Head Light, Martha's Vineyard, Massachusetts, in August, 1961. A week of prevailing southerly winds drifted the animals into the area from the Gulf Stream in considerable numbers. Although they appeared normal, it was subsequently discovered that they were in a degenerating condition compared with forms collected in the Gulf of Mexico off the Mississippi River Delta.

Investigations of the physiology of carbon monoxide secretion (Hahn and Copeland, 1966) indicated that the gas-secreting system of *Physalia* is quite sensitive to lowered temperatures and, probably, to physical handling. Therefore, fixation was done aboard boat immediately after collection. The best collecting area was 30 to 50 miles off South Pass, beyond the brown to green Mississippi River fresh water overlay and in the blue, open Gulf water. Gas samples were analyzed for each animal (method, Hahn and Copeland, 1966). Tissues from animals with less than 10% carbon monoxide float gas content were discarded. Concentrations as high as 25–28% were not unusual. The highest recorded concentration was 35% in one individual.

As soon as an animal was netted and the streaming tentacles cut off it was placed on a wire gauze frame in a deep ice chest. The entire float of the animal was thus exposed to ice cold air. After 5 to 10 minutes of cooling, dissection was commenced.

The gas-secreting epithelium is a single cell layer in thickness and easily disrupted by direct injection of fixative into the float and on to the surface of the gland. An osmic fume fixation procedure, as used in the study of gas secretion in the teleost swim bladder (Copeland, 1968), was of some aid but the best fixation procedure involved use of 5% glutaraldehyde (Sabatini *et al.*, 1963) buffered to pH 7.4 with S-collidine buffer (Bennett and Luft, 1959). This was followed by post-fixation in 1% osmic acid in the same buffer. Both fixatives and the intermediate buffer rinse were brought to 950 milliosmoles by the addition of sucrose (Caulfield, 1957).

By use of curved, fine-pointed scissors the pneumatocodon was slit for the length of the animal and the intact inner bladder (pneumatossacus) was rolled out. The still inflated pneumatossacus was held with the gas gland (pneumadena)

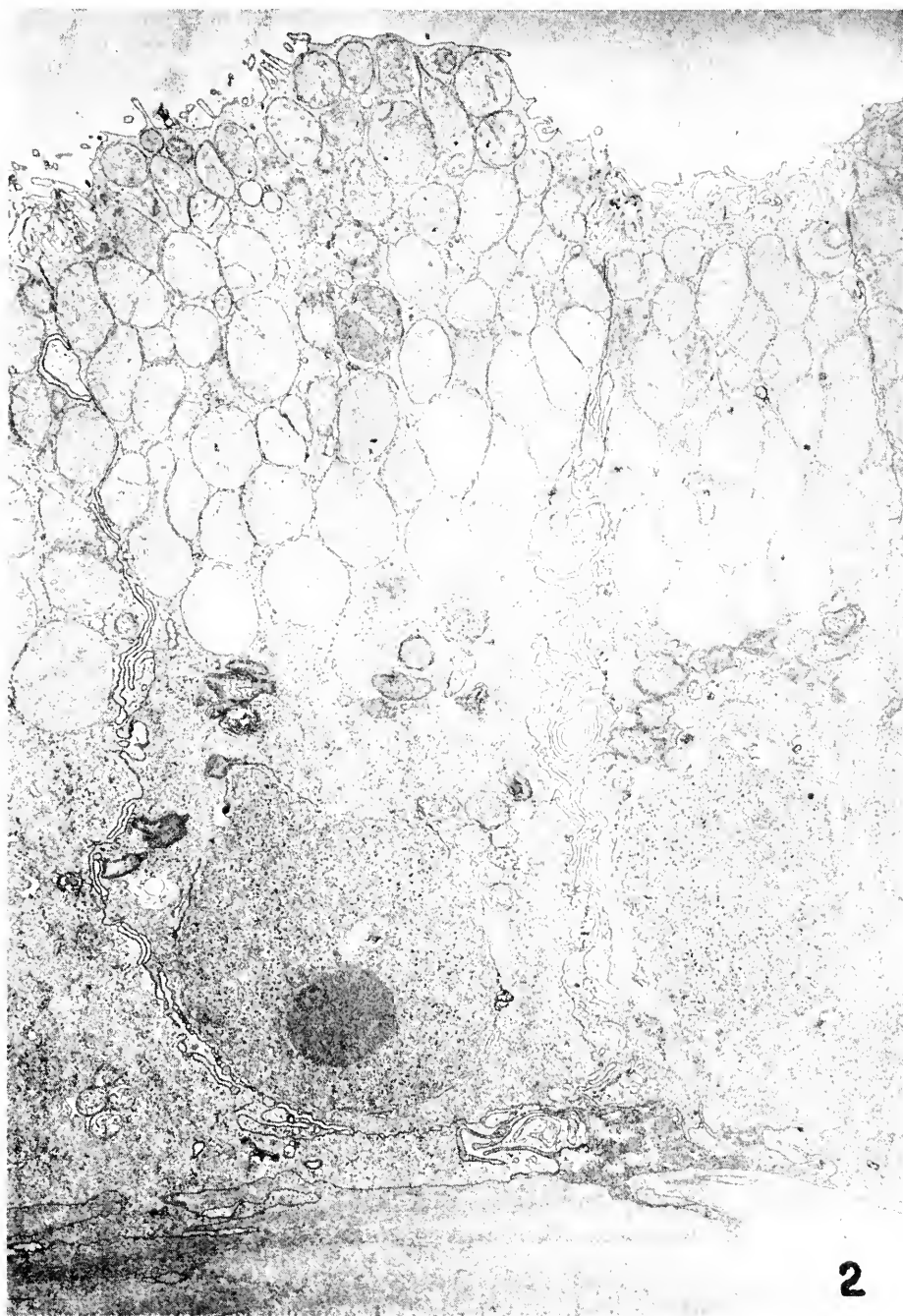
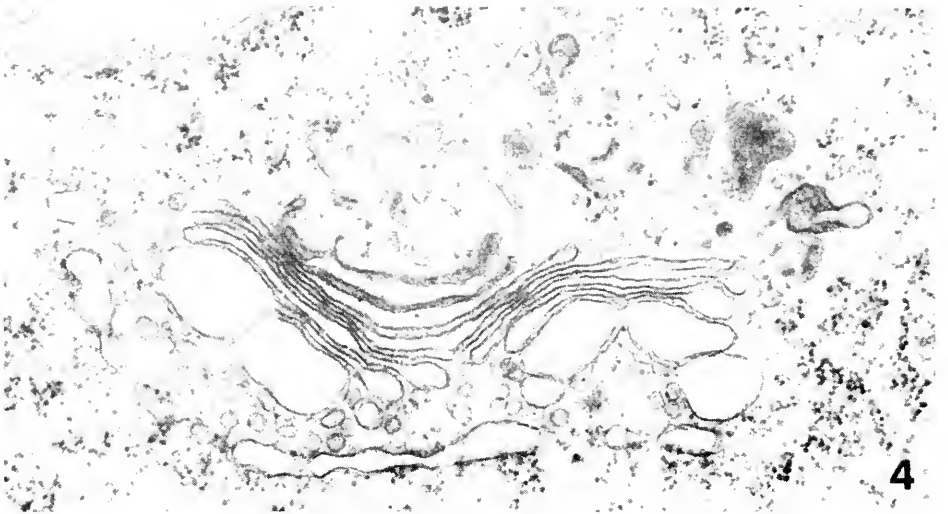
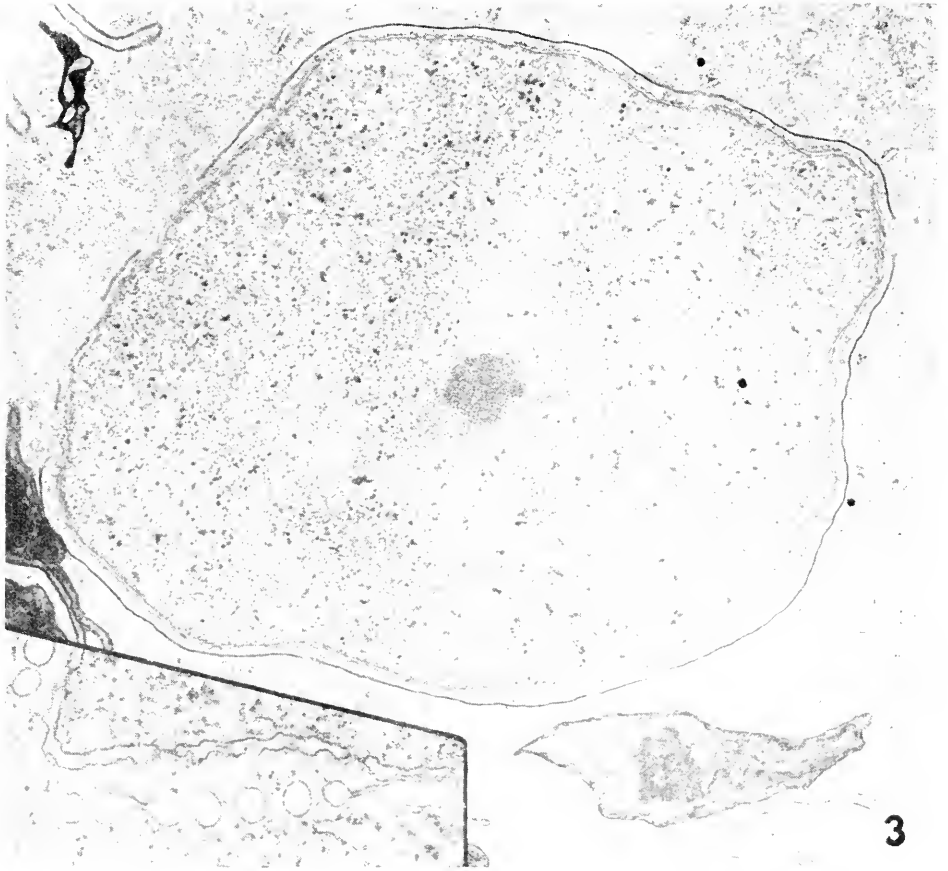


FIGURE 2. Whole cell of pneumadema layer. Gas interface is at top and mesogleal bounding layer is at bottom. See text and other figures for amplification. 4,500 \times .



FIGURES 3-4.

downward and against the surface of the chilled glutaraldehyde fixative. The relatively thick gastroderm and mesoglea were traversed by the fixative before it reached the gas-secreting ectodermal layer. Thus the delicate gas interface surface of the pneumadema was not disturbed by direct contact and the insulating effect of the gastroderm and mesoglea also resulted in more consistent fixation of the ectodermal cells throughout their depth. After 10 minutes of preliminary exposure, some of the same fixative was injected into the cavity of the pneumatosaccus, flooding the ectodermal surface of the pneumadema. After another 10 minutes the pneumatosaccus was lifted from the surface of the fixative, collapsed on dental bite-wax, the pneumadema cut free with a razor blade and placed in a vial to complete a total of 3 hours fixation with glutaraldehyde. Then followed repeated changes of buffer rinse for approximately 2 hours. Post-fixation in 1% osmic acid was for 1 hour. Dehydration was rapid, starting with 50% ethanol and proceeding with constant agitation to absolute in less than an hour. Tissues were then brought to room temperature for final trimming and embedded in Epon 812 by the method of Luft (1961). Thin sections were double stained with uranyl acetate (Watson, 1958) and lead citrate (Reynolds, 1963).

RESULTS

The ectodermal layer of the pneumadema is composed of a single layer of columnar cells with basally located nuclei and large, closely packed mitochondria in the distal halves (Fig. 2). The basal surfaces of the cells frequently have extensions that continue into the mesoglea and interdigitate with similar extensions protruding from the ends of the gastroderm cells.

The nuclei are irregular but tend to be spherical or slightly lobate in contour. In the Gay Head Light material the nuclei were frequently surrounded by an even row of vesicles (Copeland, 1962). In the more healthy material of the Gulf of Mexico vesicles were occasionally seen but they were considered to be due to breakdown of a cisternal space frequently seen adjacent and parallel to the nuclear membrane. The cisternal space itself is not always visible but the membranes are quite evident (Fig. 3).

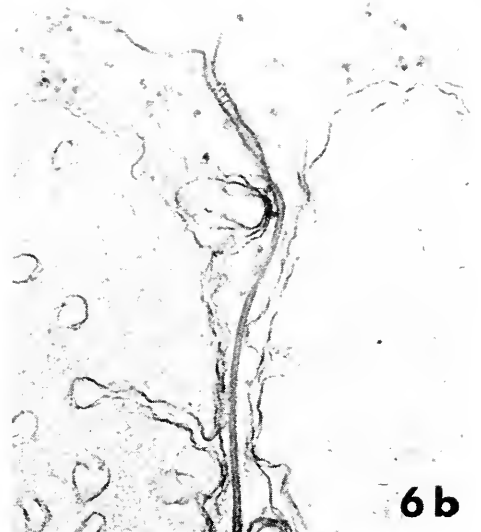
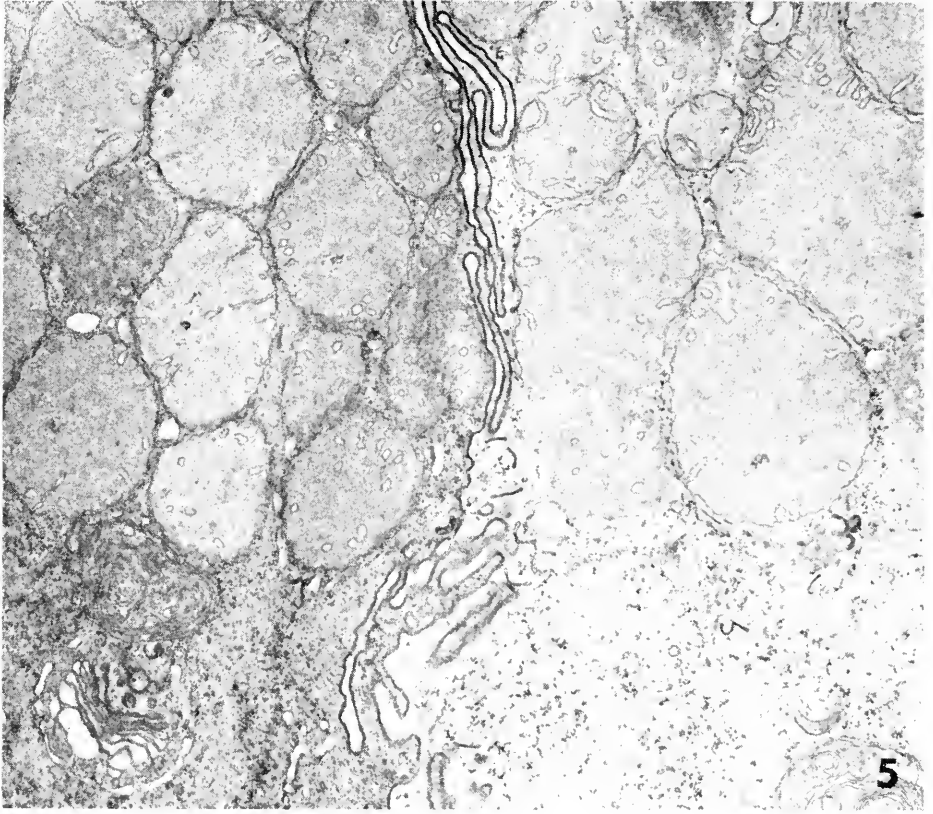
Only scanty endoplasmic reticulum is seen. It is usually of the smooth variety and randomly disposed. A branch of the endoplasmic reticulum is to be observed in association with the Golgi complex when that structure is cut at right angles to its cisternal spaces (Figs. 4 and 5).

The Golgi complexes are usually found just below the mitochondrial zone and have a characteristic configuration (Fig. 4). A flattened unit of the endoplasmic reticulum apparently delivers small vesicles that coalesce into the large cisternal spaces of the Golgi apparatus. The latter then condense and finally release dark, formed bodies at the delivery side of the complex.

A wide range of tinctorial values are to be seen in the cells. Figure 5 illustrates a case of marked contrast. There seems to be no other significant

FIGURE 3. Nucleus surrounded by cisternal membranes. Cisternal space has collapsed. Inset shows cisternal space broken into row of vesicles (Gay Head Light, Massachusetts, material). 12,000 \times .

FIGURE 4. Golgi complex. Endoplasmic reticulum space at bottom. Delivery side of complex is toward top. 40,000 \times .



FIGURES 5-6.

difference between such cells except in the case of extremely dark, obviously moribund cells. Whatever the reason for the tinctorial difference it must be a general one because the entire cell is involved. Figure 5 also well illustrates the mild degree of overlapping interdigitation between the adjoining sides of neighboring cells.

The junction between neighboring cells is plain except for a density between the plasma membranes at the distal ends of the cells near the gas interface (Fig. 6a). Under suitable orientation of sectioning, at least part of the density is seen as a typical septate desmosome (Fig. 6b). Typical zonulae occludentes, zonulae adhaerentes or maculae adhaerentes are not seen.

Immediately below the zone of closely packed mitochondria and sometimes extending down each side of the nucleus is a zone populated by multivesiculate bodies (Fig. 7). These are numerous, usually have multiple encapsulating membranes and internal vesicles that are spherical or oval, sometimes tubular in shape. The multivesicular bodies seem to arise by breakdown of encapsulated mitochondria that become isolated from those above. The initial encapsulation may also trap some of the adjacent cytoplasm (Fig. 7). However, the exact origin and fate of the multivesiculate bodies is not clear at this time.

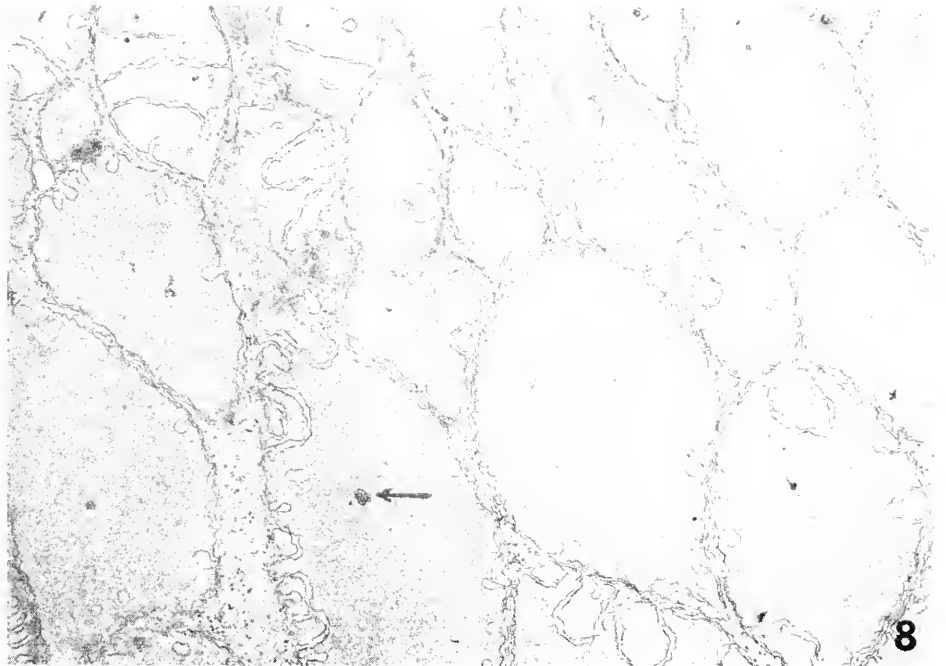
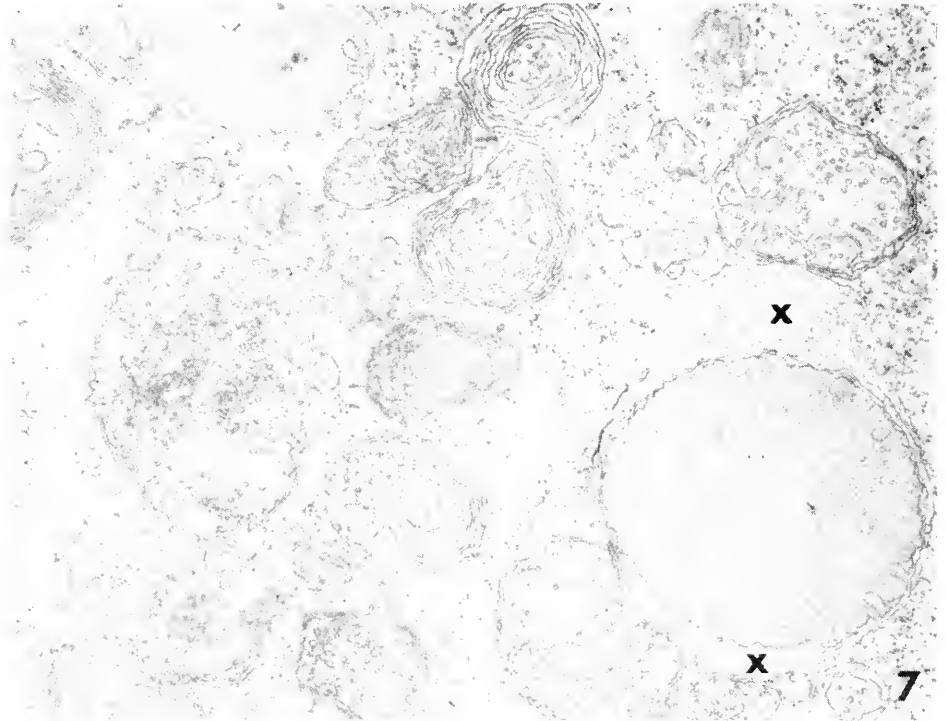
The mitochondria are quite noteworthy. They are oval in form and closely packed in the distal part of the secretory cell (Fig. 8). They differ from most types of mitochondria in having very few cristae. These are short and tubular in the main, but they occasionally form flattened extensions from one side back to the same side or across to the opposite membrane (Fig. 8). Almost the total bulk of the mitochondrion is occupied by a dense granular matrix. Irregular, dark particles are occasionally found in the granular matrix.

In well fixed material the gas surface interface of the secretory cells is seen to have many irregular projections (Fig. 9). These are not regular enough to be called microvilli but there is a resemblance. A few vacuoles or vesicles are seen near the surface but not in sufficient numbers to warrant the certainty that they are occupied with secretion of free gas. Many of them are probably oblique sections of the crypts formed by the bases of the cytoplasmic surface projections.

In a preliminary report (Copeland, 1962) it was stated that the distal end of the secretory cells possessed numerous rows of vesicles possibly devoted to gas release and that the cell was devoid of mitochondria. It was hypothesized that in the presence of high carbon monoxide levels the ectodermal cells had become reliant on respiratory and metabolic support from the gastroderm cells *via* the cytoplasmic bridges across the mesogleal layer. This hypothesis may still have some merit but the interpretation was based on a degenerating condition. The 1962 report was based on material collected in the relatively cold waters off Gay Head Light, Massachusetts. Figure 10 illustrates the morphology characteristic of that material. Figure 11 illustrates an intermediate condition observed in material collected in the Gulf of Mexico which had not degenerated as much.

FIGURE 5. Junction of a light and dark cell. Golgi complex at extreme lower left. Both cells appear normal. Interdigitation clearly shown. 12,000 \times .

FIGURE 6. Cell junctions, seen as parallel plasma membranes with dark substance between them (a). Occasionally the dark substance appears septate (b). Both: 55,000 \times .



FIGURES 7-8.

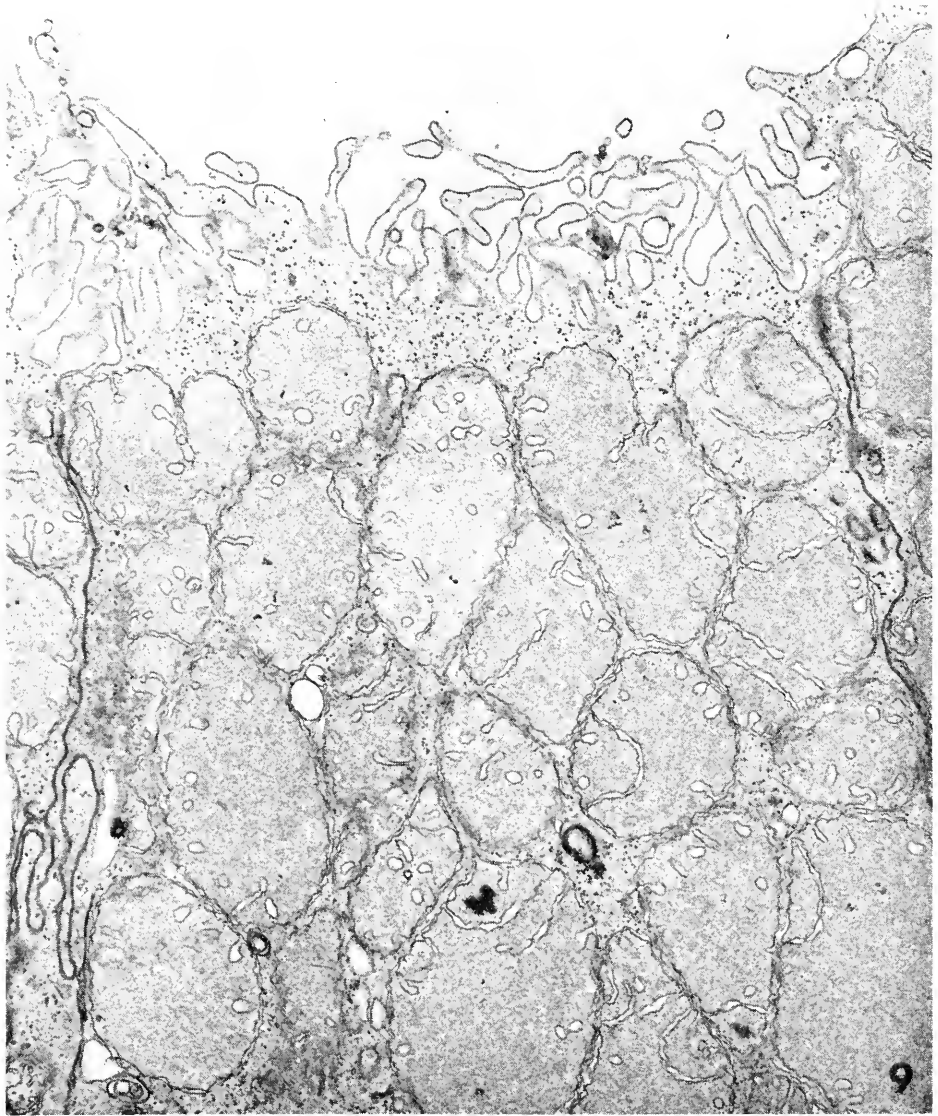
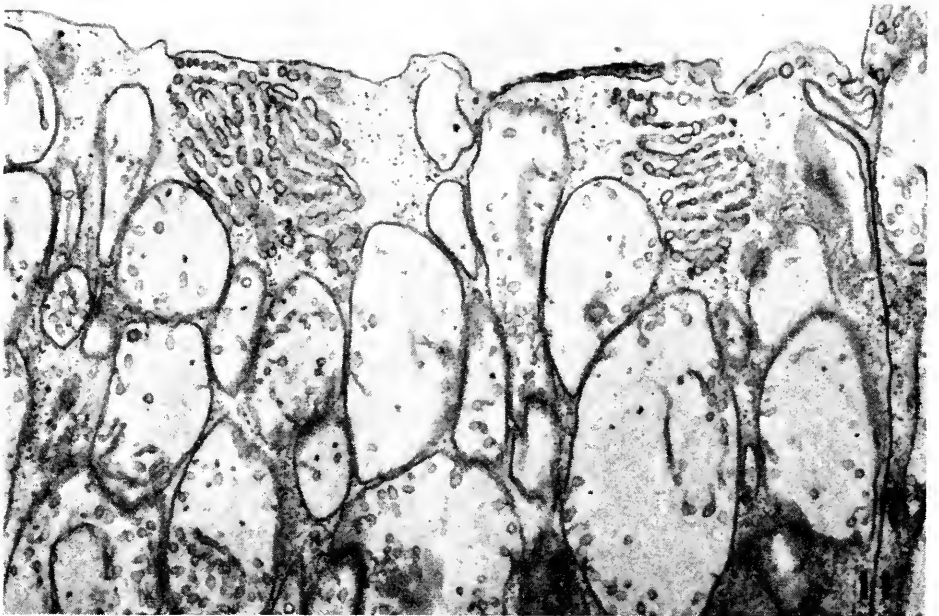
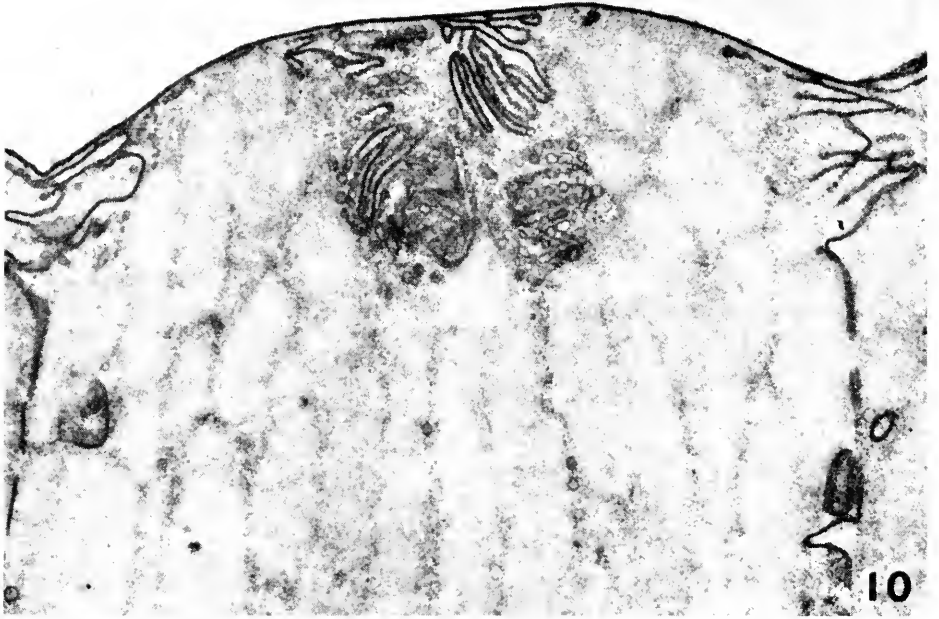


FIGURE 9. Gas surface interface. Note the many irregular projections of the surface. 18,000 \times .

Figure 9 is a comparable view in which no degeneration is visible. It and all the illustrations except Figure 10 are of material collected off the Mississippi River Delta area.

FIGURE 7. Multivesiculate bodies of varying complexity. Edge of normal mitochondria seen above. To right is an encapsulated mitochondrion with some cytoplasm (X's) included in the capsule. 19,000 \times .

FIGURE 8. Mitochondria in detail. Cristae quite sparse and short. They are mainly tubular, though long flattened ones are occasionally seen. Occasional, irregular dark bodies are seen in the matrix (arrow). 18,000 \times .



FIGURES 10-11.

FIGURE 10. "Degenerate" cell type observed in the material fixed off Gay Head Light, Massachusetts. Mitochondria completely fragmented. "Gas release vesicles" seen just below surface of cell. (Compare with Figures 9 and 11.) 11,500 \times .

As a final observation, it should be stated that the external appearance of the *Physalia* is not a reliable indication of the physiological condition of the pneumatadema. The relative percentage of carbon monoxide present in the float gas is, for obvious reasons, a better index.

DISCUSSION

Illustrations of the histological nature of the gas-secreting pneumatadema in *Physalia* are scanty indeed. Dahlgren and Kepner (1908) in their Figure 297 illustrated several secreting cells from *Physalia*. However, from my own observations it is evident that they confused the orientation of the ectodermal and gastrodermal layers in their sections and described instead the much longer columnar cells of the gastroderm with its characteristic brush border. This undoubtedly explains why Mackie (1960) was unable to identify the chromatic vacuoles of Dahlgren and Kepner in his sections of the pneumatadema prepared by similar methods.

Okada (1935) presented drawings of the early developmental morphology of *Physalia* which depicted the gas gland as having tall columnar cells with basal nuclei and distal brush borders. His drawings were diagrammatic and were not intended to present accurate histological detail.

Mackie (1960) gave a description of the histology of the ectodermal layer of the pneumatadema preserved from seven specimens. He described *columnar cells* as being in the majority with occasional *giant cells* scattered among them. In one specimen only he also saw clusters of *islet cells*. Mackie concluded that (p. 391) "The appearance of the cells in the gas gland varies markedly from one specimen to the next, and it is not clear to what extent this variability is due to differing ages of specimens, differing physiological states at the time of fixation or to differing methods of fixation." I have not been able to identify giant cells or islet cells in my own preparations and suspect that Mackie was examining tissues that were in poor condition before fixation.

Carbon monoxide is probably the only gas secreted by the pneumatadema (Hahn and Copeland, 1966). Its survival value to the *Physalia* may exist in the fact that the solubility coefficient, and thereby the diffusivity, of the gas is approximately 30 times less than that of the less toxic carbon dioxide and therefore the carbon monoxide gas is more readily retained by the highly hydrated float tissue layers (Hahn and Copeland, 1966). There is no convincing evidence in the present studies that gas is released in the form of bubbles arising within the cytoplasm of the ectodermal cells. Since only one gas may be involved, it can be released readily by direct diffusion from the cell surface. This differs from the situation in the teleost swim bladder where accumulation of multiple molecular species of gases is dependent on physical phenomena associated with cytoplasmic microbubble formation by at least one of the involved gases (Wittenberg, 1958; Copeland, 1968).

The most striking specialization to be seen in the secretory cell is the peculiar

FIGURE 11. Material fixed in Gulf of Mexico and considered to be transitional between condition in Figures 9 and 10. "Gas release vesicles" similar to those in Figure 10 are seen. Mitochondria, though in poor condition compared with those shown in Figure 9, have not completely degenerated. 11,500 X.

morphological organization of the mitochondria. The cristae are markedly reduced in size and number. Conversely, the matrix is tremendously hypertrophied in comparison with that in most types of mitochondria. It would be interesting to know if the respiratory enzymes of this peculiar mitochondrion are limited to the membranes. It may also be that the granular matrix in some way is associated with the high concentration of folates observed by Wittenberg, Noronha and Silverman (1962) which are thought to be the probable coenzymes involved in the production of the carbon monoxide gas.

The gas-secreting cell is almost devoid of the cytoplasmic morphology characteristic of most secretory cells. Smooth endoplasmic reticulum characteristic of steroid-producing cells is sparse and the rough variety associated with protein production is particularly elusive. The Golgi complexes are reasonably numerous and are well organized. However, the dark bodies or vesicles from the delivery side of the Golgi complex disappear without formation of anything resembling a secretory granule. The multivesicular bodies are numerous enough to suggest some functional role in the gas secretion. That role could well be one of degradation rather than synthesis, *i.e.*, the bodies may be lysosomal in nature. In short, the gas-secreting cell of the *Physalia* pneumadema is highly specialized for its peculiar function and it is difficult to find cytoplasmic homologies in other cell types.

There is no adequate explanation for the cisternal space, with its bounding membranes, frequently observed parallel to the nuclear membrane. It might in some way serve as a barrier between the nuclear contents and the general cytoplasm for an obscure reason. It may equally well be a rather unusual signal of pending degeneration in those particular cells.

In view of the markedly different observations of the Massachusetts forms as compared with the Mississippi Delta forms, I am reluctant to claim that those from the Delta area are completely normal and healthy animals. Presumably they moved from the Sargassum Sea area of the equatorial Atlantic Ocean on the primary Gulf Stream which swings north through the Yucatan Straits, then eastward to the tip of Florida and then up the Eastern Coast. A branch of this current comes almost straight north from the Yucatan Straits into the Mississippi Delta area *via* the old DeSoto Canyon. In terms of time-distance lapse, the Delta forms should be reasonably healthy, especially the smaller (younger) ones. A quick answer could be obtained by analyzing the gas of *Physalia* collected in their rearing grounds. If concentrations of carbon monoxide higher than 35% are observed, then the fine structure of the pneumadema would require further investigation.

There is one puzzling point not answered by the present study. The animals captured off Gay Head Light, Massachusetts, looked normal externally, with well inflated floats. Unless the permeability of the pneumatococcus and pneumatocodon had been markedly reduced, the animals were still secreting some gas, albeit at a reduced rate, from tissue in which the mitochondrial membranes had disappeared. If it is assumed that the substrate-enzyme system for carbon monoxide production is associated with the granular matrix of the mitochondria, then that system might still be intact to some degree in the distal ends of the degenerating cells. In that event, the complex of vesicles seen in Figure 10 might bear in last analysis a

functional relationship to the modified system to produce a lowered level of gas release.

We are indebted to the Freeport Sulphur Company for the occasional use of their seaplane to locate the blue water interface which is usually in the form of a tide-rip. Its location varies tremendously with wind, tide and the general set of Gulf water movement (which is usually from east to west across the end of the Mississippi River Delta). We are also indebted to the Coast Guard for their most welcome help on two separate emergency occasions.

SUMMARY

1. The carbon monoxide gas-secreting tissue (pneumadema) of *Physalia* is a single layer of ectodermal cells that are cuboidal to columnar in morphology. There is only one specific type cell and it is characterized by having the distal part, adjacent to the gas-interface surface, packed with large mitochondria. The mitochondria are unique in having few and small cristae. Most of the internum of the mitochondrion is occupied by a granular matrix. Compared with other secreting cells, there is very little smooth endoplasmic reticulum and less rough endoplasmic reticulum. There are numerous Golgi complexes and complex multi-vesiculate bodies. The nucleus is often enveloped by a cisternal space.

2. The secretory epithelium is sensitive to adverse factors such as lowered temperature and presents changes that, though degenerate in appearance, may still have some functional ability to produce carbon monoxide. There is no evidence to indicate conclusively that gas is secreted by formation of vesicles in the cytoplasm. Therefore, it is assumed that carbon monoxide is secreted by diffusion from the cell surface which is frequently thrown into finger-like projections.

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A STUDY OF THE JELLY ENVELOPES SURROUNDING THE EGG OF THE AMPHIBIAN, *XENOPUS LAEVIS*¹

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The greatest interest in the oviducal jelly material deposited around amphibian eggs has centered on its possible significance in the processes of meiosis (Humphries, 1961) and fertilization (Hughes, 1957; Kambara, 1953; Katagiri, 1963b, 1966; McLaughlin, 1967; Nadamitsu, 1957; Rugh, 1935, 1951; Shaver and Barch, 1960; Shivers and Metz, 1962; Subtelny and Bradt, 1961; Tchou and Wang, 1956). It is generally accepted that the jelly is necessary for successful fertilization of amphibian eggs; however, the particular properties of the jelly which make its presence essential to fertilization are largely unknown. Kambara (1953, p. 84) has suggested that the jelly ". . . plays the role of a mechanical foothold which enables the sperm to penetrate the eggs." In contrast, Katagiri (1966) proposes that the jelly does not act as a foothold but that a smaller moiety in the jelly may facilitate adherence of the spermatozoa to the egg surface. The recent work of Barbieri and Villeco (1966) also suggests that a small molecular weight substance present in the jelly is necessary for fertilization.

It can be expected that additional knowledge of the structure and composition of the jelly would permit more enlightened studies on precisely what properties of the jelly are necessary for fertilization. Chemical analyses of amphibian jelly have indicated that the main components of the jelly are fucose, hexosamines, one or more hexoses, and protein (Bolognani *et al.*, 1966; Folkes *et al.*, 1950; Hiyama, 1949a, b, c; Kusa and Ozu, 1961; Lee, 1967; Masamune and Yosizawa, 1953; Masamune *et al.*, 1951; Minganti, 1955; Minganti and D'Anna, 1957, 1958; Schulz and Becker, 1935). However, not all of the above investigations dealt with each of these components and in every case all the jelly layers surrounding the egg were analyzed together; no attention was given to the possible differences among layers. That these differences do exist has been shown by histochemical studies on oviducts and eggs (Ghiara, 1960; Humphries, 1966; Humphries and Hughes, 1959; Kambara, 1956a, b, 1957; Kelly, 1954; Salthe, 1963; Shaver, 1966).

The present investigation was undertaken to study the deposition, structure, and composition of the egg jelly of the South African clawed toad, *Xenopus laevis* (Daudin). Since amphibian jelly is deposited in discrete layers, it was considered of prime importance in this study to analyze these layers separately.

¹This work was completed while the author was an NSF predoctoral fellow and was part of a dissertation submitted in partial fulfillment of the requirements for the degree of Doctor of Philosophy at Emory University. The work was supported in part by grant GM-09878 from the U. S. Public Health Service.

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Histochemical methods, which are useful for this purpose, have been employed extensively in the present work. In addition, it appeared that it would be possible with this species to separate the jelly layers in small quantities for chemical analyses. To the author's knowledge, such chemical analyses of individual jelly layers have not been reported for any amphibian.

MATERIALS AND METHODS

Ovulation of *Xenopus laevis* was induced by an injection of 500 units of chorionic gonadotropin (Antuitrin-S, Parke, Davis).

Cytochemical methods

Oviducts containing eggs were removed from a pithed animal and cut into small segments for fixation. Uterine eggs and deposited eggs were also used. The material was fixed with 10% neutral formalin, 10% neutral formalin containing 0.01 *M* cetylpyridinium chloride (CPC) (Conklin, 1963) or by freeze-substitution (Humphries, 1966). All material was cleared in benzene and embedded in Tissuemat. For most of the staining procedures, serial sections of 10 μ thickness were cut and placed alternately on each of 3 slides.

Carbohydrates. The periodic acid-Schiff reaction was used with and without prior treatment with malt diastase. Both native and boiled enzyme were used in a concentration of 1% in distilled water for 2 hours at room temperature. Toluidine blue was employed in a concentration of 0.1% in distilled water for 1-2 hours. Coriphosphine O (Gurr) was used in a 0.01% solution in distilled water for 15 minutes and sections were examined in distilled water (Humphries, 1966). Detection of fluorescence in coriphosphine-stained material was by means of a Leitz SM microscope using an HBO 200-watt mercury vapor lamp with Leitz exciter filter UV UG1. Alcian blue was employed routinely as a 0.1% solution in 3% acetic acid (pH 2.4-2.6) or was used in increasing concentration of standardized $MgCl_2$ according to the method of Scott and Dorling (1965).

Bovine testicular hyaluronidase (Sigma Chemical Co.) was used in a concentration of 60 USP units/ml. in 0.1 *M* phosphate buffer at pH 6.0. Sections were incubated for 8 hours at 37° C. Cartilage controls were used. Sialidase (neuraminidase, *Vibrio cholerae*, Calbiochem, 500 units/ml.) was used according to the method of Spicer and Warren (1960). Enzyme-treated sections were stained with toluidine blue or alcian blue.

Proteins. Staining was done according to the bromsulphalein method of Silverman and Glick (1966) and the ninhydrin-Schiff procedure of Yasuma and Itchikawa (1953). The dimethylaminobenzaldehyde (DMAB)-nitrite method (Adams, 1957) for tryptophan was also employed. For the detection of sulfhydryl groups the mercury orange method of Bennett and Watts (1958) was used with and without prior treatment with thioglycolate to reduce disulfides (Barka and Anderson, 1963).

Enzymatic treatment of live, jelly-covered eggs

Eggs were stripped from the female and placed immediately into the various solutions. Treatments were carried out at room temperature in native and boiled

enzyme solutions and in solvent controls. Papain ($2 \times$ crystallized, Sigma Chemical Co.) was used in a concentration of 0.1 mg./ml. in a 0.1 N phosphate buffer (pH 6.3) containing 5 mM cysteine-HCl and 5 mM EDTA. Pepsin (N. F., Merck and Co.) was employed in a concentration of 1 mg./ml. in 0.01 N HCl. Hyaluronidase was prepared as described above. All observations were made on the live eggs.

Chemical methods

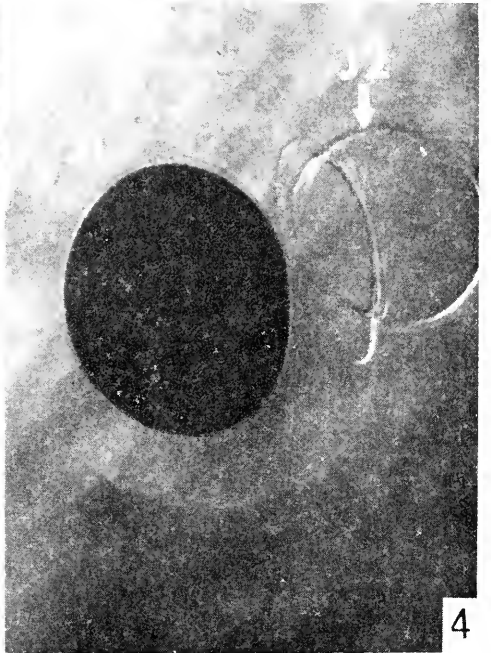
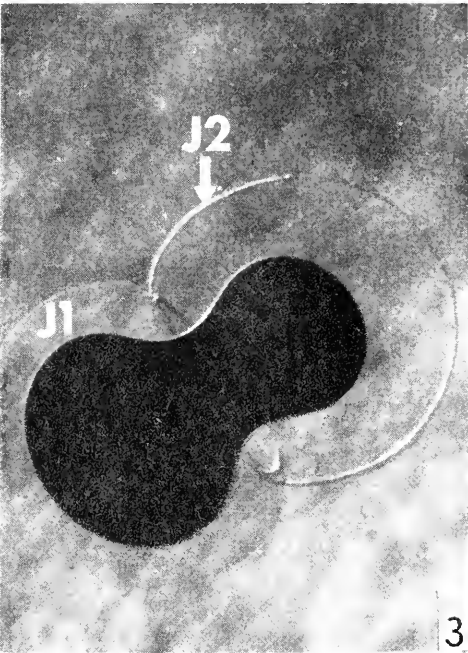
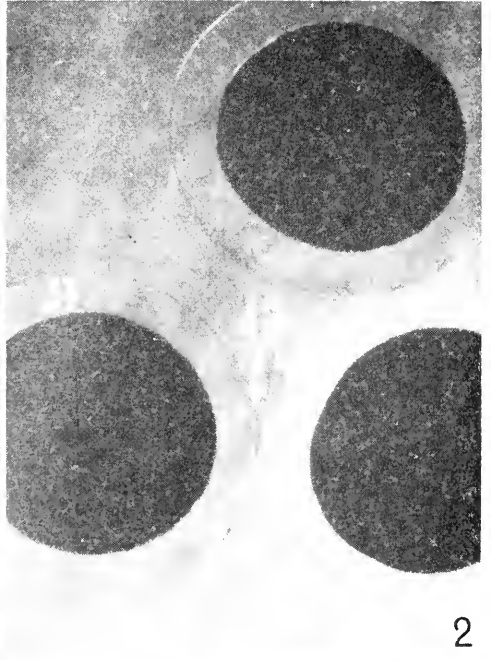
For removal of the jelly, approximately 50 eggs at a time were stripped from a female into distilled water. The jelly layers were removed separately with sharpened watchmaker forceps, placed immediately in vials kept in an ice-salt bath at -10° C. to 0° C. and stored in the frozen state. The jelly coats from the eggs of several females were later pooled and lyophilized.

Nitrogen. Nitrogen content of the jelly was determined by the Nessler method (Lang, 1958) using two different procedures. (1) Equal numbers of coelomic and uterine eggs from the same female were assayed and nitrogen content of the jelly was taken as the difference between the two values. In each case additional eggs from the same female were dried at 100° C. to determine weight. (2) One-mg. samples of lyophilized jelly coats were assayed to determine the distribution of nitrogen among the layers.

Uronic acids. The method of Galambos (1967) was employed. One-mg. samples of each of the separate, lyophilized jelly coats were dissolved in 0.2 ml. of 1 N NaOH, diluted to 2 ml. with distilled water, and 0.8-ml. portions were used for the Galambos procedure. Chondroitin sulfate (20–400 μ g., Nutritional Biochemical Corp.) and glucuronic acid (4–80 μ g., crystalline, Sigma Chemical Co.) were prepared in 0.1 N NaOH and used as standards.

Sialic acid. Total sialic acid was determined by the thiobarbituric acid assay (Warren, 1959, 1960). To study the distribution of sialic acid among the jelly layers, assays were performed on 1–2 mg. of the separate, lyophilized jelly coats. N-acetyl-neuraminic acid (synthetic, Type IV, Sigma Chemical Co.), in concentrations from 0.01 μ mole to 0.05 μ mole, was used as the standard. Readings were made on a Zeiss spectrophotometer in 3-ml. cuvettes (1-cm. light path).

Paper chromatography. The method of Caldwell and Pigman (1965) adapted from Masamune and Yosizawa (1953) was used to identify sugars present in the separate jelly layers. Approximately 2 mg. of each coat were hydrolyzed in 1–2 ml. 2 N H_2SO_4 in capped tubes in a boiling water bath for 5 hours. The solutions were then brought to approximately pH 6.0 by adding saturated $BaOH_2$ and, after centrifugation, the supernatants were evaporated to dryness at room temperature with a stream of cool air. The sugars were redissolved in 2–4 drops of distilled water and 8–12 μ l. were used to spot each chromatogram. Reference sugars (1 mg./ml.), placed on the same paper, included D-glucose, D-glucosamine-HCl, N-acetyl-D-glucosamine, D-galactose, D-galactosamine-HCl, N-acetyl-D-galactosamine, D-mannose, D-mannosamine-HCl, N-acetyl-D-mannosamine and D-fucose (Sigma Chemical Co.). Descending chromatography was carried out in n-butanol-pyridine-water (5:2:2 v/v) on 18.5×22 inch Whatman No. 1 chromatography paper in an equilibrated chromatography chamber. The solvent was allowed to drip off the paper and the total time ranged from 40



FIGURES 1-4.

to 62 hours. Reducing sugars were detected by the method of Trevelyan *et al.* (1950) and amino-sugars by an adaptation of the Elson-Morgan reaction for hexosamines (Smith, 1960).

RESULTS

When a uterine egg is placed in tap water the three transparent jelly coats can be observed easily (Fig. 1). The inner layer of jelly (J1) is approximately 0.2 mm. wide and has a gelatinous consistency. The outermost layer (J3) is about 0.1 mm. wide and because it becomes quite sticky in tap water the deposited eggs tend to be held together in masses. Between these two layers is another very thin layer (J2), approximately 0.02 mm. wide, which appears membranous and fairly tough. With forceps, J3 may be removed easily. If J2 is then punctured with the tip of the forceps, the egg, still surrounded by the inner layer (J1), can be pushed out (Figs. 2, 3, 4).

The jelly layers are deposited as the eggs pass through the long, coiled oviduct. As the eggs travel from the coelom into the oviduct they are collected in a thin-walled ostial region. From here they pass singly into the first secreting region where J1 is deposited around the egg. This section of the oviduct is fairly thick-walled and lined with jelly-secreting glands. Approximately half-way down the oviduct the wall becomes even thicker and the glands in this second secreting region can be distinguished histochemically from those in the anterior half of the oviduct. J2 and J3 are formed around the egg as it travels through this region. The eggs then pass into the thin-walled sac-like uterus where they accumulate before egg deposition begins.

Cytochemical observations

The picture observed in the live jelly-covered egg was best retained when the material was fixed by freeze-substitution. Severe shrinkage of the jelly occurred when 10% formalin or 10% formalin containing CPC was used. Since the histochemical reactions were essentially the same after these three fixation methods the material to be described below is that fixed by freeze-substitution unless indicated otherwise.

Both secreting regions of the oviduct and all jelly layers were distinctly PAS-positive. The second secreting region and J2 and J3 reacted more intensely than did the first secreting region and J1. Diastase did not affect the PAS reactivity of the jelly or the oviducal glands but did remove essentially all positivity from the egg. Toluidine blue-stained material in water showed an intense pink to purple metachromasia only in J1 and the glands of the first secreting region. After

All figures: Live *Xenopus laevis* eggs stripped from a female and immersed in tap water. $\times 30$.

FIGURE 1. All three jelly layers are present.

FIGURE 2. J3 has been removed with forceps.

FIGURE 3. J3 has been removed, J2 has been punctured with forceps, and the egg still surrounded by J1 has been pushed partially out of J2.

FIGURE 4. Same as Figure 3, but egg surrounded by J1 has been pushed completely out of J2. J2 remains on outer edge of J1.

alcohol dehydration of formalin-CPC-fixed material, metachromasia of J1 and the first secreting region was retained although diminished considerably in intensity. In material fixed by freeze-substitution, alcohol dehydration usually abolished any metachromasia. With coriphosphine O, J1 and the glands which secrete it gave a bright orange-red fluorescence. J2, J3, and the glands of the second secreting region were unstained or were only a dull orange. With alcian blue at pH 2.4, J1, and the glands of the first secreting region stained bright blue. The egg, J2, J3, and the second secreting region of the oviduct were unstained.

When material was stained with alcian blue at pH 5.8 in increasing concentrations of $MgCl_2$, J1 and the glands in the first secreting region of the oviduct continued to stain until the $MgCl_2$ concentration exceeded 0.5 *M*–0.7 *M*. Glands of the second secreting region and J2 and J3 did not stain in concentrations above 0.3 *M* $MgCl_2$. Magnesium chloride in excess of 0.2 *M* prevented the egg from staining. Rat rib cartilage staining was abolished in concentrations of $MgCl_2$ above 0.9 *M*.

When any of the carbohydrate stains were used on formalin-CPC-fixed material, the inner portion of J1 gave a more positive reaction than the outer portion of J1. With freeze-substitution J1 appeared essentially homogeneous throughout.

After treatment of sections with hyaluronidase or sialidase, there was no noticeable change in toluidine blue or alcian blue (pH 2.4) staining of the eggs, jelly coats, or oviducal glands.

All four methods employed for the detection of proteins gave the same general staining pattern. With ninhydrin-Schiff the egg was bright magenta. The glands in the first secreting region were essentially unstained, and J1 was only a very pale pink in some sections. J2, J3, and the second secreting region gave a consistent positive reaction although the color was only a pale pink. Somewhat brighter staining was observed after formalin-CPC fixation. Bromsulphalein stained the egg deep purplish-blue. J1 and the first secreting region remained unstained. J2, J3, and the second secreting region were very pale blue. With the DMAB-nitrite method for tryptophan the egg was a positive blue. J1 and the glands which secrete it had only occasional traces of a very pale blue. J3 and the second secreting region were a consistent pale blue. J2 was a deeper blue. When mercury orange was used without prior reduction of disulfides, all three jelly layers and the glands of both secreting regions were negative for sulfhydryl groups. Only the egg stained orange. After thioglycolate treatment, the color in the egg increased and J1 and the first secreting region were pale orange. J2, J3, and the second secreting region were a brighter orange.

Enzymatic treatment of live eggs

After about two hours in papain, J3 had disappeared and J1 had begun to liquefy within the intact J2 layer. After an additional two hours all jelly had been removed. Pepsin appeared to cause dissolution of J3 only. To test further whether J1 and J2 were affected by the enzyme, eggs with only J1 and also separate J2 coats were placed in the pepsin solution. In native enzyme and control solutions J1 shrank but remained around the egg. Separate J2 coats also shrank but, after as long as eight hours, no dissolution of J2 was noticed. Hyaluronidase had no apparent effect on the jelly.

Chemical analyses

When the jelly coats were removed manually from the egg for chemical analyses, it was observed that J3 could be removed without disturbing the other layers and so was probably free of other material. J2 contained a small amount of material from J1 and perhaps also a lesser amount from J3. J1 was thought to be free of contamination from J2 and J3 but, since not all of J1 could be removed without cytolyzing the egg, the innermost part of J1 was not included in the following analyses of the jelly.

Table I gives the values for nitrogen content of: (1) the entire jelly of the eggs of three females (designated A, B, and C); and (2) lyophilized samples of the three separate jelly coats collected from the eggs of several females.

TABLE I
Nitrogen content of eggs and jelly layers

Type of test*	Sample	Mean $\mu\text{g. N/egg}$ (Std. error)	N as % of dry wt. of jelly
(1) Eggs from:	Female A	coelomic eggs	24.62 (± 0.40)
		uterine eggs	27.05 (± 0.64)
	Female B	coelomic eggs	46.91 (± 0.61)
		uterine eggs	51.63 (± 0.22)
	Female C	coelomic eggs	34.63 (± 0.27)
		uterine eggs	38.34 (± 0.74)
(2) Lyophilized jelly coats:	J1		7.1%
	J1		7.9%
	J2		8.1%
	J2		6.0%
	J3		6.8%
	J3		7.4%
	J3		6.8%

* See explanation in "Materials and Methods."

Table II gives the results of sialic acid determinations on separate, lyophilized jelly coats. On a dry weight basis J1 appears lowest in sialic acid with 0.14%. J2 has the highest average value, 0.40%, but substantial variations were found in these samples. J3 consistently contained 0.37%. Hydrolysis for three hours did not result in higher sialic acid values but, in fact, lower values were often obtained.

The test for uronic acid content of the three separate jelly layers was negative.

Chromatography

For each jelly layer three separate chromatography runs were made. Duplicate sheets were included in each run so that the two detection reagents could be used.

TABLE II
Sialic acid content of the separate jelly layers

Sample	Number of samples†	Hydrolysis time	Sialic acid μ g. mg. dry wt.*	Sialic acid % of dry wt.
J1	3	1 hour	1.39(1.17-1.51)	0.14%
J2	3	1 hour	4.02(2.30-6.00)	0.40%
J3	3	1 hour	3.71	0.37%
J1	1	3 hours	0.62	0.06%
J3	1	3 hours	2.78	0.28%

† Number of samples used to get average values listed.

* Equation 2 of Warren (1959) was used for all calculations and 309, the molecular weight of N-acetylneuraminic acid, was used in order to express concentrations in micrograms.

In Table III are listed the sugars present in each jelly coat as found by comparing the R_{glucose} values of the unknown sugars with those of the standards and by comparing the colors obtained after using the Elson-Morgan reagent.

DISCUSSION

The only three descriptions available for the jelly of *Xenopus laevis* eggs are brief and conflicting. Deuchar (1966) reports two jelly coats, Salthe (1963) suggests five coats, and Ghiara (1960) observed three: a thin inner layer, a wide middle layer, and a thin outer coat. He noted metachromasia in the two inner layers after staining live eggs with toluidine blue. The present report presents evidence that there are three coats: a wide, gelatinous inner coat, a very narrow, membranous middle layer, and a fairly wide outer coat.

The inner layer of jelly appears to contain neutral and acidic polysaccharides and possibly protein. Toluidine blue metachromasia, intense alcian blue staining, and a bright orange-red fluorescence with coriphosphine O strongly suggest the presence of some acid groups whose identity remains unknown. The tests for uronic acids were negative. Similar negative results have been reported for the other amphibians investigated (Bolognani *et al.*, 1966; Folkes *et al.*, 1950; Hiyama, 1949a). It is doubtful that the intense alcian blue (pH 2.4) staining of J1 is due to carboxyl groups because at pH 2.6 or lower, carboxyl group staining may be masked as a result of salt links with proteins (Scott and Dorling, 1965). Using alcian blue at pH 5.8 in increasing concentrations of MgCl_2 , Scott and Dorling (1965) found that carboxyl group staining can be abolished at a critical electrolyte

TABLE III
Sugars identified in each jelly layer

J1	Fucose, galactose, glucosamine, galactosamine, glucose?*
J2	Fucose, galactose, glucosamine, galactosamine
J3	Fucose, galactose? glucosamine, galactosamine, mannose? mannosamine?

* Question marks in the table indicate that a spot with an R_f value corresponding to that particular known sugar was found in only one of the three chromatography runs.

concentration (CEC) of approximately 0.3 *M* MgCl₂ and sulfate at 1.0 *M* or greater; a mixture of the two, such as found in heparin, has an intermediate CEC. The CEC for sialic acid-containing structures is approximately 0.2 *M* MgCl₂ (Quintarelli and Dellovo, 1965). Since the CEC of layer J1 is about 0.7 *M*, staining of this layer cannot be accounted for solely by sialic acid. Also, the toluidine blue and the alcian blue stainings were not affected by sialidase treatment. The above evidence and also the persistence of some toluidine blue metachromasia after alcohol dehydration suggest the presence of sulfate in J1 and the glands of the region which secrete it. Past evidence (Minganti, 1955; Lee, 1967) has indicated that amphibian egg jelly has no sulfate. However, other biochemical reports have suggested that sulfate may be present (Minganti and D'Anna, 1958; Bolognani *et al.*, 1966) and, in previous histochemical investigations, several species of amphibians have been found to exhibit metachromasia in the jelly and/or oviducts (Ghiara, 1960; Humphries, 1966; Humphries and Hughes, 1959; Kambara, 1957; Katagiri, 1963a; Kelly, 1954; Shaver, 1966). In no case has the substance responsible for this metachromasia been identified.

The source of the J2 layer is still in question. Although both J2 and J3 appear around the egg in the second secreting region of the oviduct, there has been no way to subdivide further this part of the oviduct and locate precisely the site of deposition of J2. The staining reactions of J2 resemble those of J3 although the reactions for neutral polysaccharides and proteins in J2 are more intense. This may be due in part to the construction of J2, which in sections appears to consist of rather tightly packed fibers arranged parallel to the egg. J2 is not metachromatic and is alcian blue negative at pH 2.4. The alcian blue staining at pH 5.8 is similar to that of J3 and is probably due to amino acids and/or sialic acids, not to uronic acids. Unlike J3, and similar to J1, J2 does not dissolve upon treatment with pepsin. In live material its consistency is quite different from either of the other two layers. It is fairly tough and membranous and not at all sticky as is J3; when J3 is removed, the eggs are no longer held together in masses. Possibly, J2 as such is not deposited by the oviduct but is formed when J3 is deposited on J1. Perhaps some complexing of the components of J1 and J3 occurs, which might change the enzyme susceptibility and staining characteristics of these components. Unless some material is found which is unique to J2 it will be difficult to establish that it is a truly distinct layer and that it is deposited as such by a particular region of the oviduct. There may be a similar situation in the case of *Rana pipiens*, where, although the egg has three jelly layers, the middle part of the oviduct appears immunologically to be an area of overlap between the upper and lower regions (Barch and Shaver, 1963). Katagiri (1963a) describes for the eggs of *Hyla arborea japonica* a "jelly membrane" between the inner and the outer two layers of jelly; as spermatozoa penetrate the jelly they come to a standstill momentarily at this "jelly membrane." Katagiri also suggests that changes in the "jelly membrane" are a prime cause of the decrease in fertilizability of the eggs which have been standing in tap water. Tchou and Wang (1956) noticed that when the eggs of *Bufo bufo asiaticus* remain in water for some time a distinct membrane forms between the outer and inner jelly layers which may act as an obstacle to sperm penetration.

J3 appears to be composed predominantly of neutral polysaccharides and some

protein. The presence of carboxyl groups is indicated and, since there is no uronic acid, these groups could be due to amino acids and/or to the small amount of sialic acid that is present. The actual values obtained for sialic acid are probably much too low because fucose, which has been found in all jelly layers, causes abnormally low readings with the method used (Warren, 1959). Recently, sialic acid has also been found in the oviduct and/or egg jelly of other amphibians (Humphries and Workman, 1966; Humphries *et al.*, 1968; Bolognani *et al.*, 1966; Lee, 1967). Soupart and Noyes (1964) demonstrated histochemically the presence of sialic acid in the zona pellucida of the ova of several mammals, and the possible significance of the compound was investigated by Soupart and Clewe (1965) who treated rabbit ova with neuraminidase and found deformation of the zona pellucida and a definite inhibition of sperm penetration.

In view of the differences observed histochemically in the jelly layers of *Xenopus* eggs it is interesting that, qualitatively, the sugar components of the three layers are nearly identical. From the preliminary work reported here, all layers appear to contain fucose, glucosamine, and galactosamine. Galactose is present in J1 and J2 and possibly in J3. In addition, J3 may contain mannose or mannosamine. The sugars may be joined together in polysaccharides and/or may be attached to proteins. It appears from the histochemical reactions that protein is present in all three jelly layers although these tests are not completely satisfactory since they do not actually indicate protein but are specific for certain reactive sites or for particular amino acids. However, proteolytic digestion of the jelly does suggest that the amino acids are linked together in proteins and, furthermore, the enzymatic treatment has served to demonstrate that some differences do exist among the proteins of the three jelly layers.

In conclusion, it is obvious that the jelly capsule is composed of discrete layers. The separate layers appear to differ chemically and physically and as a result they may have quite different functions. To pool all layers for chemical analyses or for studying fertilization may give misleading results. Information about the macromolecules of the jelly layers is now needed.

I wish to express my sincerest appreciation to Dr. A. A. Humphries, Jr., who suggested this study and who provided encouragement and advice during the period of my graduate studies and the preparation of this manuscript.

SUMMARY

1. The deposition, structure, and composition of the jelly surrounding the egg of *Xenopus laevis* were studied cytochemically and biochemically. During its passage through the oviduct, the egg is invested with three layers of jelly designated J1, J2, and J3, from innermost to outermost. Particular emphasis was placed on analyzing the layers separately.

2. All layers gave positive histochemical tests for neutral polysaccharides. J2 and J3 stained the most intensely. The sugars present in the polysaccharides of each layer were identified chromatographically. J1, J2, and J3 all contain fucose, glucosamine, and galactosamine. Galactose is present in J1 and J2 and possibly in J3. J3 may also contain mannose or mannosamine.

3. On the basis of the histochemical tests it is suggested that J1 contains sulfate. The weak positive reactions for acid polysaccharides in J2 and J3 indicate the presence of carboxyl groups only. Since no uronic acid was found in the jelly it is proposed that the carboxyl groups are part of amino acids and/or sialic acid which was found in small amounts in all layers.

4. The nitrogen values for the three layers are similar. All layers gave positive histochemical tests for protein although the reactions in J1 were weak and irregular. Treatment of live eggs with proteolytic enzymes showed that all jelly layers do contain some protein as an important structural component and that the protein components of the three layers are not identical.

5. It is suggested that future studies should be concerned with characterizing the macromolecules of the jelly and that, for all investigations of the structure and function of the jelly, the differences among layers should be taken into account.

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STUDIES ON THE ENDOCOMMENSAL CILIATE FAUNA OF CARIBBEAN SEA URCHINS

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From a review of the literature, it appears that Jacobs (1914) initiated studies on intestinal ciliates of American sea urchins. During the past 50 years, several investigators have made contributions on the ciliate fauna of sea urchins of the Western Hemisphere; thus almost two dozen species of ciliates have been described from the echinoids. Much of the early work was conducted on sea urchins of Bermuda. Ball (1924) started studies on ciliates of *Diadema* sp. and *Echinometra* sp. of Bermuda and suggested new names for five species of ciliates which she observed; but she never named them. At the request of the Director of the Bermuda Biological Station, Dr. D. H. Wenrich prepared a publication on some of the results of Miss Ball's (now Mrs. Ruth Ball Biggar) study. In the paper of Biggar and Wenrich (1932), the ciliates *Metopus circumlabens* found in *Diadema setosum* and *Echinometra subangularis*, *Cryptochilum bermudensis* in *Toxopneustes variegatus*, *Cryptochilum echinometris* of *Echinometris subangularis*, and *Anophrys elongata* found in both *Toxopneustes variegatus* and *Echinometris subangularis* were described. Lucas (1934) described *Metopus rotundus* found in *Diadema setosum* at Bermuda. Since then, several reports have been given on

TABLE I
Frequency of ciliates in echinoids of Puerto Rico

Host	Ciliates							Total number specimens examined
	<i>Anophrys aglycus</i>	<i>Anophrys elongata</i>	<i>Biggaria bermudensis</i>	<i>Biggaria echinometris</i>	<i>Cohni-lembus caeci</i>	<i>Cyclidium rhabd-lecton</i>	<i>Metopus circum-ladens</i>	
<i>Diadema antillarum</i>		10—	38++				26++	38
<i>Echinometra lucunter</i>		40++	36++	8—		21+	49++	59
<i>Lytechinus variegatus</i>	12+	42++	35++	10—		9++	34+	46
<i>Triplaneustes ventricosus</i>		16—	34++	4—	31+	33++	30+	53

The numbers given in the columns under ciliates in each of the Tables I-V represent the number of hosts found infested with the indicated ciliate; ++ = very abundant, + = many and — = few.

TABLE II
Frequency of ciliates in echinoids of Curacao

Host	Ciliates							Total number specimens examined
	<i>Anophrys aglycus</i>	<i>Anophrys elongata</i>	<i>Biggaria bermudensis</i>	<i>Biggaria echinometris</i>	<i>Cohni-lombus caeci</i>	<i>Cyclidium rhabdotectum</i>	<i>Metopus circumlabens</i>	
<i>Diadema antillarum</i>	4+	1+	6+			6+	8+	8
<i>Echinometra lucunter</i>	4+	3+	2++			2+	2+	6
<i>Lytechinus variegatus</i>	1-	1+	1++					1
<i>Tripneustes ventricosus</i>	8+	8++	3+	1-	3+	1-	3+	12

endocommensal ciliates of echinoids inhabiting the waters of the Atlantic, Gulf, and the Pacific Coasts of North America (Powers, 1933, 1935; Berger, 1960; Berger and Profant, 1961; Beers, 1948, 1954, 1961; Lynch, 1929).

In South America, Urdaneta-Morales and Tengler de McLure (1966) studied the ciliates of *Echinometra lucunter*, *Diadema antillarum*, *Tripneustes ventricosus* and *Eucidaris tribuloides* of the Federal District of Venezuela. It appears that the report of Berger (1961) on the ciliates of *Diadema antillarum*, *Clypeaster rosaceus*, *Echinometra lucunter*, *Lytechinus variegatus*, and *Tripneustes ventricosus* of Bimini Islands, Bahamas, is the only information available on endocommensal ciliates of sea urchins inhabiting between Bermuda and South America. Because there seemed not to have been any major studies on ciliates of sea urchins between Bimini and Venezuela, a survey was commenced in early January, 1968, to compare the ciliate fauna of sea urchins in this area of the Caribbean with the

TABLE III
Frequency of ciliates in echinoids of St. Croix

Host	Ciliates							Total number specimens examined
	<i>Anophrys aglycus</i>	<i>Anophrys elongata</i>	<i>Biggaria bermudensis</i>	<i>Biggaria echinometris</i>	<i>Cohni-lombus caeci</i>	<i>Cyclidium rhabdotectum</i>	<i>Metopus circumlabens</i>	
<i>Diadema antillarum</i>			10++		1-	3+	10++	10
<i>Echinometra lucunter</i>		4+				1+	3++	4
<i>Tripneustes ventricosus</i>		6++	6++		1-	2+	3++	11

TABLE IV
Frequency of ciliates in echinoids of St. Thomas

Host	Ciliates							Total number specimens examined
	<i>Anophrys aglycus</i>	<i>Anophrys elongata</i>	<i>Biggaria bermudensis</i>	<i>Biggaria echinometris</i>	<i>Cohni-lembus caeci</i>	<i>Cyclidium rhabdo-lectum</i>	<i>Metopus circum-labens</i>	
<i>Diadema antillarum</i>		4+	10++		1-	3+	10++	10
<i>Tripneustes ventricosus</i>		6++	6++		1-	2+	3++	11

reports on Bimini and Venezuela. This report concerns the results from studies on the endocommensal ciliates found in four species of sea urchins collected from the Caribbean Sea at St. Thomas, St. Croix of the Virgin Islands, Curacao, Netherlands Antilles, Vieques, and with major emphasis on samples taken from the southwestern coast of Puerto Rico.

MATERIALS AND METHODS

Samples of sea urchins, *Echinometra lucunter*, *Diadema antillarum*, *Lytechinus variegatus*, and *Tripneustes ventricosus*, were collected from the littoral in the Caribbean at five islands. Specimens were taken from Brewers Bay at St. Thomas, St. Croix near Buck Island, the Piscardera Bay at Curacao, Vieques, and the southwestern coast of Puerto Rico. In this survey 282 sea urchins were collected at a depth not greater than four feet.

The specimens were examined for intestinal ciliates immediately, or on an average within eight hours after collecting. The method of examination was essentially as described by Lucas (1934). Studies were made primarily on living ciliates, observed both by bright-field and phase contrast microscopy. In a few cases, a dilute solution (1:10,000) of Lugol's iodine was employed as a supra-

TABLE V
Frequency of ciliates in echinoids of Vieques

Host	Ciliates							Total number specimens examined
	<i>Anophrys aglycus</i>	<i>Anophrys elongata</i>	<i>Biggaria bermudensis</i>	<i>Biggaria echinometris</i>	<i>Cohni-lembus caeci</i>	<i>Cyclidium rhabdo-lectum</i>	<i>Metopus circum-labens</i>	
<i>Echinometra lucunter</i>	1+	4+		4-		2+	2++	5
<i>Lytechinus variegatus</i>	1+	3+	6++			1+		6
<i>Tripneustes ventricosus</i>			2+				1+	2

TABLE VI
Comparative distribution of ciliates

Host	Ciliates						
	<i>Anophrys aglycus</i>	<i>Anophrys elongata</i>	<i>Biggaria bermudensis</i>	<i>Biggaria echinometris</i>	<i>Cohniembus caeci</i>	<i>Cyclidium rhabdolectum</i>	<i>Metopus circumlabens</i>
<i>Diadema antillarum</i>	—	B	B	B	—	—	B
	V	—	V	—	V	V	V
	—	P	P	—	—	—	P
<i>Echinometra lucunter</i>	—	B	B	B	—	—	B
	V	V	V	V	V	V	V
	—	P	P	P	—	P	P
<i>Lytechinus variegatus</i>	—	B	B	B	—	—	—
	P	P	P	P	—	P	P
<i>Tripneustes ventricosus</i>	—	B	B	B	—	—	—
	V	V	V	V	V	V	V
	—	P	P	P	P	P	P

For each host, indication is given for report of individual ciliates at Bimini (Berger, 1961), Venezuela (Urdaneta-Morales and Tengler de McLure, 1966), and Puerto Rico (the present report).

B = Bimini
V = Venezuela
P = Puerto Rico
— = The species was not reported

vital stain. For more detailed morphological studies some of the ciliates were fixed in Schaudinn's fluid, stained in Heidenhain iron hematoxylin, and mounted in Kleermount xylene solution.

RESULTS

Examinations of intestinal samples from the stated echinoid hosts observed in this study revealed that at least seven ciliates live as endocommensals in the Caribbean. The hosts and ciliates are given in Tables I–V. At St. Croix, *Lytechinus variegatus* was not found in the collecting area, while at St. Thomas, *Lytechinus variegatus* and *Echinometra lucunter* were not found in the area. *Diadema antillarum* was not obtained from the collecting area at Vieques.

DISCUSSION

In Table VI, the results of the present study, Puerto Rico only, have been compared with the reports of Berger (1961), and Urdaneta-Morales and Tengler de McLure (1966) on the intestinal ciliates of sea urchins of Bimini and Venezuela, respectively. Because *Lytechinus variegatus* was not included in the study by Urdaneta-Morales and Tengler de McLure (1966), only the ciliates of Bimini and Puerto Rico can be compared for this sea urchin.

It is of interest to note that *Anophrys aglycus* was reported for *Diadema*

antillarum, *Echinometra lucunter* and *Tripneustes ventricosus* of Venezuela but not for the same species at Bimini and Puerto Rico. This ciliate was found in *Lytechinus variegatus* at Puerto Rico, and likewise in *Echinometra lucunter* at Vieques. At Curacao, *Anophrys aglycus* was found in four species of sea urchins. In general, the seven ciliates given in this report are found in two or more species of sea urchins from Bimini, through the Caribbean to South America. There are considerable variations in the ciliate fauna and their abundance at different localities; likewise significant variations in hosts' abundance were observed. Thus it may well be that the diet of the host and the abundance of species living in association may be factors involved in the distribution of endocommensal ciliates.

The authors express thanks to the Staff at the Institute of Marine Biology, University of Puerto Rico, La Parguera, Puerto Rico, for use of their facilities. We are also grateful to the Director of the Caribbean Marine Biological Institute, Dr. F. Creutzberg, for his assistance, and use of the facilities, Curacao, Netherlands Antilles.

SUMMARY

Two hundred and eighty-two sea urchins, *Diadema antillarum*, *Echinometra lucunter*, *Lytechinus variegatus*, and *Tripneustes ventricosus*, collected from the littoral in the Caribbean were examined for intestinal ciliates. The specimens were collected from the islands, St. Thomas, St. Croix, Curacao, Vieques, and with major emphasis, the southwestern coast of Puerto Rico. Studies were made primarily on living ciliates with the exception of a few specimens fixed in Schaudinn's fluid and stained with iron hematoxylin; seven ciliates were found: *Anophrys aglycus*, *Anophrys elongata*, *Biggaria bermudensis*, *Biggaria echinometris*, *Cohnilembus caeci*, *Cyclidium rhabdotectum*, and *Metopus circumlabens*.

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REGENERATING TISSUES FROM THE COCKROACH, *LEUCOPHAEA MADERAE*: NERVE REGENERATION *IN VITRO*¹

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In an earlier study, Marks and Reinecke (1965) reported the migration of cells from the nerve stump of isolated leg regenerates of the Madeira cockroach, *Leucophaea maderae* (F.), and demonstrated that this growth did not follow the cyclical pattern associated with the molting process that was characteristic of epithelial tissues from the regenerating leg. In the same study, an isolated incident was observed and recorded on film in which cell processes grew out from the proximal end of the 5th mesothoracic nerve of an isolated leg regenerate and made contact with a ganglion that had been placed adjacent to it. Wigglesworth (1965) stated that during post-embryonic development, sensory axons *in vivo* normally differentiated and migrated inward to invade the central ganglion while motor axons emerged from the ganglion itself, and Bodenstern (1957) suggested that the same processes occurred during the healing of a severed nerve. Guthrie (1962, 1967) and Jacklet and Cohen (1967) demonstrated that regeneration of transplanted ganglia and the healing of mesothoracic nerves takes place *in vivo* in the adult cockroach. The recent development of a nutrient medium—M-18 (available from Grand Island Biological Co., Grand Island, N. Y. 14072)—that favors nerve growth and development therefore encouraged us to repeat our earlier *in vitro* experiments to study nerve regeneration in detail. In the present study, we attempted to correlate the regenerative growth processes as observed *in vitro* with normal nerve regeneration as found *in vivo* by other workers.

MATERIALS AND METHODS

The technique for preparing leg regenerates was described in detail by Marks (1968). The mesothoracic legs of late instar nymphs were cut off at the trochantero-femoral articulation 24 hours after molting. Regeneration was allowed to proceed for 8 days, at which time the entire coxa was removed from the insect, and the leg regenerate was dissected under sterile conditions; at the same time, the prothoracic ganglion and gland were removed. The explants, two leg regenerates and a single ganglion, were washed in M-18 nutrient medium, placed in a Rose multipurpose tissue chamber under a strip of dialysis membrane, and the explants so oriented that the stump of the 5th mesothoracic nerve projecting proximally from the leg regenerate was adjacent to the stumps of the lateral nerves

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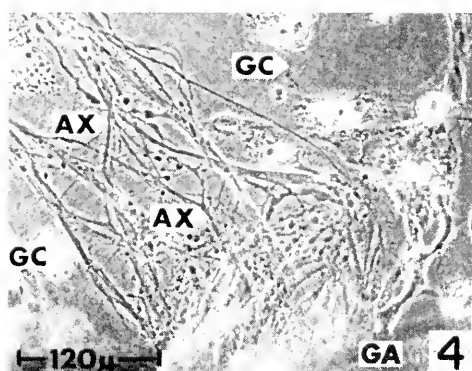
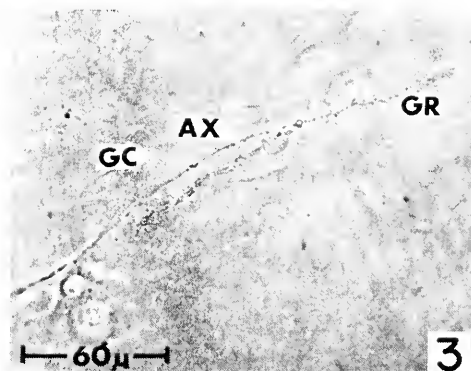
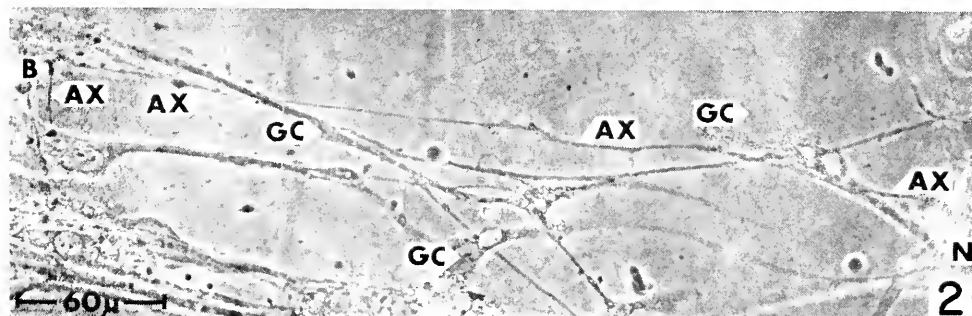
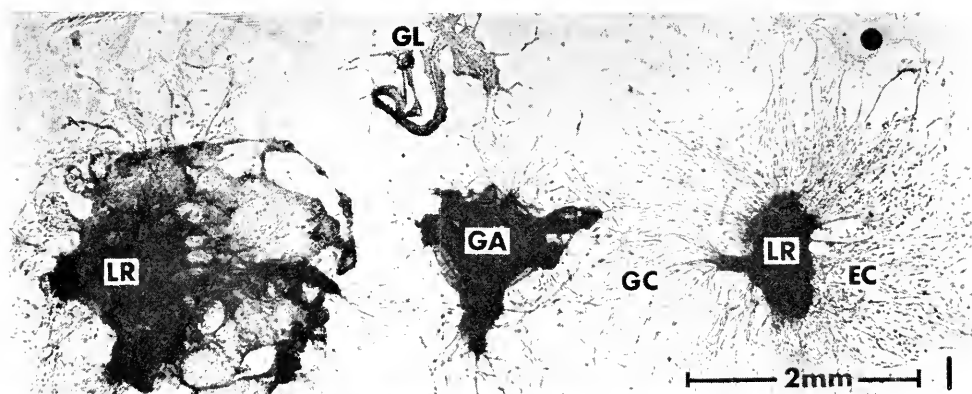


FIGURE 1. Culture containing leg regenerates (lr), prothoracic ganglion (ga), and prothoracic gland (gl). The leg regenerates are surrounded by partial sheets of epithelial cells (ec). Glial cells (gc) are seen emerging from the cut ends of the nerve of both leg regenerates and ganglion and forming a cell bridge. *In vitro* 21 days; transmitted light.

FIGURE 2. Typical growth from the nerve end in a leg regenerate. Sensory axons (ax) emerge from the severed nerve (n) and maintain close contact with several glial cells (gc). *In vitro* 37 days; dark contrast phase.

FIGURE 3. Schwann-like glial cell (gc) with sensory axon (ax) running along the surface. Time-lapse studies show much activity in the region of the growth cone (gr). *In vitro* 60 days; dark contrast phase.

FIGURE 4. Axon (ax) fibers emerging from ganglion explant (ga) migrate through a break in the neurilemma. Glial cells (gc) containing lipid-filled vacuoles are also emerging. *In vitro* 26 days; dark contrast phase.

of the ganglion (Fig. 1). The distance between these tissues was not allowed to exceed 1 mm. In some cases, the prothoracic gland was placed adjacent to the ganglion. The chamber was set up with one glass and one plastic coverslip, and the explants were held against the glass coverslip by the dialysis strip. The completed chamber was filled with M-18 that contained 7.5% fetal calf serum. Another series was prepared with chambers containing only ganglia or leg regenerates. The explants were examined weekly under a phase contrast microscope with a long working distance condenser. If axons and glial cells emerged from the explant, it was scored as positive; if not, it was scored as negative. Only explants that could be clearly scored were included. The results of each experiment were tested against the others by using a test for difference between two sample proportions given by Goldstein (1964). Significance was set at the 95% level of confidence.

To fix and stain these cultures for identification of the cell types, we found it necessary to disassemble the Rose chamber and remove the glass coverslip with its associated dialysis strip. Since the cultures grew between the coverslip and the dialysis strip, extreme care was taken to preserve the delicate structures. First, the 45×50 -mm. coverslip was cut to a 1-in. width with a diamond pen and then the coverslip was attached, together with the undisturbed dialysis strip, to a slide with metal clips. The entire assembly was then fixed in Gendré's fixative for two hours, stained by the periodic acid-Schiff technique, followed by 5 minutes in Hansen's trioxymethanin. Additional explants were pretreated with either alpha-amylase or pepsin before they were stained by the PAS method. After dehydration and clearing, the dialysis strip was so heavily stained that it was necessary to remove it. In most cases, the explant and associated cell outgrowth adhered to the coverslip though occasionally the explant adhered to the dialysis strip and was pulled away with it. When this occurred, the cell outgrowth remained on the coverslip and could be studied after mounting. The trioxymethanin provided a delicate stain that outlined both the nucleus and cytoplasm, so the location of the PAS-positive granules in the cell was easily determined.

RESULTS

The first activity seen in the explanted leg regenerates occurred within 24 hours when the blood cells migrated onto the coverslip; they frequently formed an imperfect monolayer around the leg regenerate. Within 10 days, epithelial cells were migrating from the explant, and a few days later, large granular cells with long attenuated processes migrated from the nerve sheath or neurilemma and often became very long and spindle-shaped. These probably represent glial cells. They could be distinguished from neurons by their granular cytoplasm and by migration of their cell bodies from the explant across the glass. They often joined in long strings that moved over the coverslip and eventually made contact with cells from other explants, either blood cells or other glial cells (Fig. 1). Examination of the cell surfaces frequently revealed fine fibers of darker contrast. When we traced these fibers, we found that they extended from the interior of the nerve stump. The fibers were morphologically distinct from the glial cells and possessed growth cones identified as sensory on the basis of

work done by Pomerat (1967) and Reinecke (personal communication); the apparent location of the cell body in the leg regenerate (*i.e.*, distal to the cut) further identified them as sensory fibers. They sometimes made contact with blood cells as they moved from the explant, but they were usually found closely associated with the glial cells (Fig. 3). In a few cases, they crossed the cell bridge between explants.

The first activity seen in the explanted prothoracic ganglia appeared 12 to 14 days after explantation: blood cells were usually present, but few epithelial cells and no cell sheets were formed around the ganglion. By the 14th day, two types of fibers were migrating from the explant. One emerged from the body of the ganglion through breaks in the neurilemma, and they migrated freely over the glass surface and formed extensive networks. Both blood and glial cells were contacted by these fibers, the growth cones of which were similar to those identified as motor types by Pomerat and Reinecke. The cell bodies of these fibers always remained within the ganglion. Thick ropelike strands often formed and bridged the gap between explants and invaded the tissues of the leg regenerates (Fig. 4).

A second type of fiber migrated from the cut ends of the lateral nerves of the ganglion in close association with the glial cells. They were morphologically indistinguishable from the fibers described earlier from the leg regenerate and were assumed to be axons from association neurons within the ganglion. The location of the cell bodies of these fibers is not known.

In the 14 growth chambers containing one ganglion and two leg regenerates, 81% of the 23 leg regenerates that could be scored and 85% of the 14 ganglia showed fiber growth. Cell bridges between the ganglion and one or both the leg regenerates developed in 64% of the 14 chambers (Fig. 5).

Maximum development of the cultures occurred at 25 to 30 days *in vitro*; after this, the supporting cells began to degenerate. The larger groups of nerve fibers persisted longer and remained as a network that could be identified when freed from the accompanying blood and glial cells (Fig. 6). Axoplasmic traffic was clearly visible in time-lapse photomicrographic studies of these fiber bundles. The makeup of the bundles in older chambers was difficult to ascertain in the living material; sensory, motor, or both types of fibers may be involved. A few glial cells usually remained though no well-defined sheath was observed, and there was no evidence of the secretion of a neurilemma. In some of the chambers, such connecting fibers had formed into compact, cable-like structures that resembled a nerve trunk; in others, they remained diffuse. Such nerve-mimicking structures with recognizable axons and glial cells were found as long as 80 days after explantation.

When a prothoracic gland was present with the ganglion, nerve fibers from the ganglion frequently made contact with the gland. The prothoracic gland remained alive and put out cell migrants, and its presence had no visible effect on the activity of either glial cells or nerve fibers.

Of the leg regenerates set up with the ganglion and prothoracic gland, 87% of 16 produced nerve fibers and support cell migrants; of those set up without the gland, 81% of 23 showed such growth. The ganglion explants had a similar pattern: 86% of 13 showed cell migration in the presence of the gland and 85%

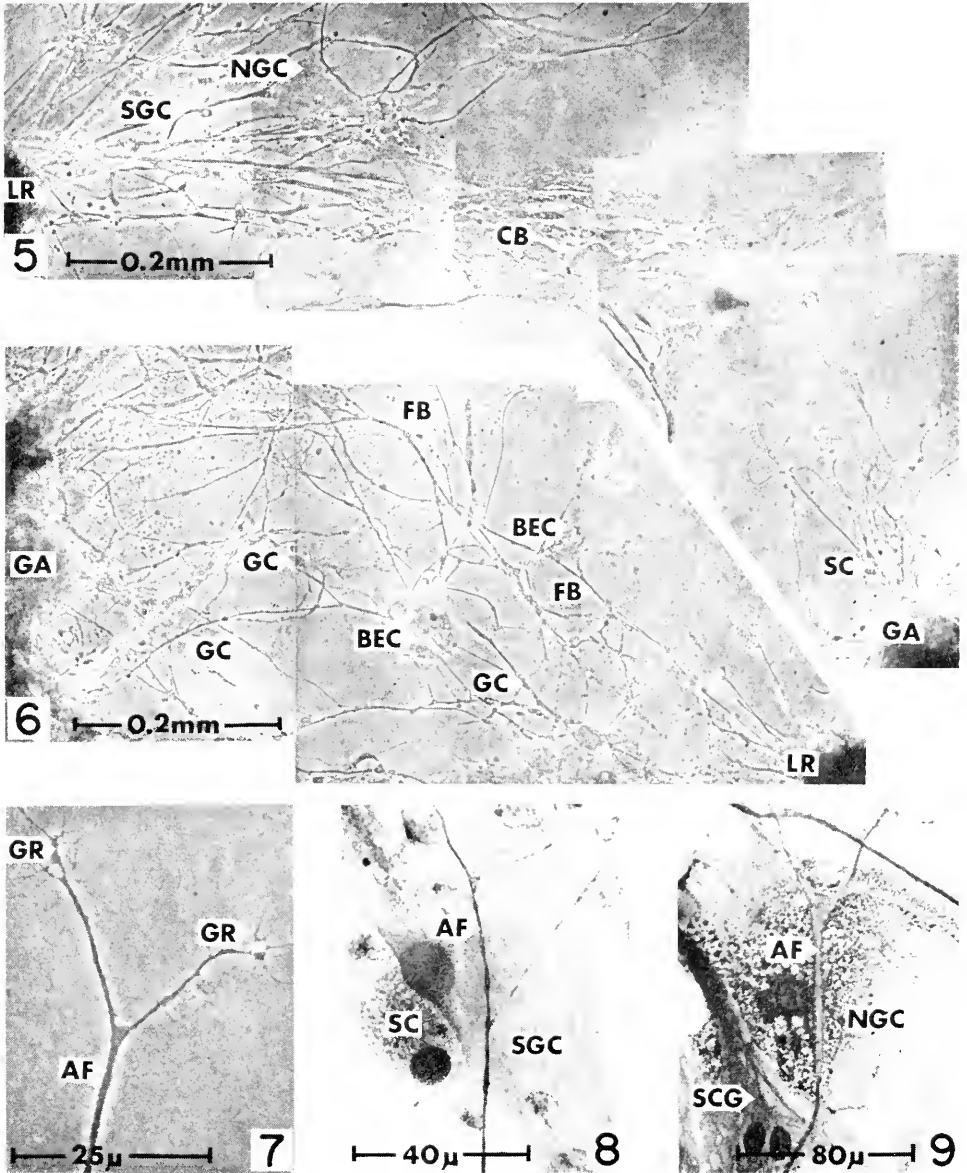


FIGURE 5. Cell bridge between leg regenerate (lr) and ganglion explant (g) is developing. Elongated Schwann-like glial cells (sgc) and granule-filled nutritive type glial cells (ngc) are emerging from the explants. What may be a sheath cell (sc) is seen near the ganglion explant. The center portion of the bridge (cb) appears to be made up of a mixture of blood, glial, and epithelial cells. *In vitro* 34 days; dark contrast phase.

FIGURE 6. Degeneration of blood and epithelial cells (bec) in old cultures exposes the mature nerve net that has formed. A few recognizable glial cells (gc) remain associated with the fiber bundles (fb). *In vitro* 62 days; dark contrast phase.

of 14 in its absence. A summary of these results is given in Table I. Thus, in this test, the presence of the gland had no apparent effect on the occurrence of nerve regeneration.

To check these observations, we made a second series of experiments in which homologous explants from different animals were placed together in a single chamber. Both leg regenerates (four to a chamber) and prothoracic ganglia (three to a chamber) were prepared as described. The chambers contained about the same amount of tissue as those described earlier. Both kinds of tissue grew quite readily. In chambers containing only leg regenerates, the glial cells and fibers migrated in 44% of the 23 explants. Cell bridges of blood and glial cells that carried axons were formed, but they did not reach and penetrate into other explants, and no lasting fiber aggregates were formed.

TABLE I

The effect of the presence of various tissue combinations on nerve regeneration in vitro

Tissue combinations	Test organs:			
	Leg regenerate		Prothoracic ganglion	
	% growth	Total scored	% growth	Total scored
Leg-ganglion-leg	81	23	85	14
Leg-ganglion+gland-leg	87	16	86	13
4 Leg regenerates	44	23	—	—
3 Ganglia	—	—	90	20

The chambers provided an excellent opportunity to study the regeneration of the sensory type fibers since no motor cell bodies were present in the isolated leg regenerate. The long, slender axons were phase dark and could be seen against the lighter cytoplasm of the glial cells. A single axon sometimes varied throughout its length, broadening and then abruptly narrowing again. Irregular branching occurred, and slender filaments frequently extended laterally at irregular intervals. With time-lapse cinephotomicrography, the growth cone portion of the axon was extremely active, and at times, it extended outward from the advancing glial cell. As the growth cone proceeded along the surface of a sheath cell, it formed flattened droplets of cytoplasm that appeared to provide connections

FIGURE 7. Branched axon (af) fiber with two growth cones (gr) that are probably sensory in origin but have lost some of their identifying characteristics. This fiber branched from the single large axon bundle that joined a leg regenerate to the ganglion. *In vitro* 74 days; dark contrast phase.

FIGURE 8. Stained preparation showing: a sheath cell (sc), a Schwann-like glial cell (sgc) with cytoplasmic fibers, typical nuclear morphology, and associated axonal fiber (af).

FIGURE 9. Stained preparation showing two Schwann-like glial cells (sgc) adjacent to a nutritive glial cell (ngc) filled with PAS-positive granules. Nuclear morphology of these two cell types is virtually identical. Axonal fibers (af) are closely associated. *In vitro* 50 days.

with the membranes of the sheath cells. Mitochondria and refractile granules were occasionally seen in the axoplasm.

In chambers containing only ganglia, migration of nerve fibers occurred in 90% of the 20 explants, and both types of nerve fibers were present. The results of these experiments are summarized in Table I. In such chambers, particularly those 40 or more days *in vitro*, fibers presumably from association neurons often migrated independently; however, when a glial cell was encountered, the fiber frequently followed it for a considerable distance. In the older chambers, trunk-like aggregates of fibers were frequently formed, possibly as a consequence of the thigmotropic behavior of these cells.

In preparations of young ganglia (15 to 30 days *in vitro*), fibers that were probably motor fibers, as judged by the structure of their growth cones and by the fact that structures of this type were absent in leg regenerate preparations, were found growing mostly from the body of the ganglion through breaks in the neurilemma. They grew out onto the glass surface where the individual fibers formed angular, jointed patterns and loops, and later, as they matured, fiber bundles. In the older chambers (30 to 45 days), it became more difficult to identify the fibers, and growth cones of intermediate types were frequently found (Fig. 7). Cell bridges formed between individual ganglion explants within the same chamber.

While the phase contrast microscope allowed us to ascertain that the cells involved in forming cell bridges were of more than one type, we could not always distinguish clearly between the different types of cells. Several ganglion cultures were therefore fixed and stained for study.

The nerve fibers and glial cells stained readily with Hansen's trioxihematin, and the neurilemma of the ganglion yielded a positive reaction when it was stained by the PAS technique. The sheath appeared to be lined with small cells containing densely staining nuclei with the chromatin scattered in coarse granules. These cells, often binucleate, occasionally migrated onto the coverslip where they could be identified by their small, dark-staining nuclei (Fig. 8). The cytoplasm was densely packed with PAS-positive granules that were unchanged by digestion with either alpha-amylase or pepsin. The cells occurred in relatively small numbers and fitted Wigglesworth's (1959) description of type I glial cells that belong to the perineurium.

A second type of migrating cell appeared in greater numbers, were much larger, and contained a large, lightly granulated nucleus with one or two prominent nucleoli. The cytoplasm was hyaline around the periphery and frequently included large numbers of granules in the perinuclear region that were also PAS-positive after digestion with alpha-amylase and pepsin (Fig. 9). These cells had an alveolar texture to their cytoplasm, frequently with areas along the periphery that appeared to contain numerous small vacuoles that were also PAS-positive. These cells are one of the components of the cell bridges that form between explants.

A third type of cell present in large numbers varied greatly in shape and was characterized by the presence of numerous fibrils in the cytoplasm and by the absence of PAS-positive granules. They were very large, and some developed long fibrous processes. They were usually the first cells to migrate from a fresh

explant, and they were the main components of the cell bridges (Fig. 8). They were also the cells with which the growth cones of the nerve fibers were most commonly associated as they migrated from the explant. Both the second and third types of cells appeared to belong to the type II category of glial cells (Wigglesworth, 1959) and appeared to be actively involved in the process of nerve regeneration. The second type with its PAS-positive granules may play a nutritive role (Wigglesworth, 1960); the third type with its long, fibrous processes and intimate association with the nerve fibers may play the role of Schwann-like cells. It is likely that these two types have a common origin, the morphological differences being dependent on the physiological role of a particular cell at given time since intermediate types are occasionally found.

DISCUSSION

The nerve cell bridges formed in our *in vitro* studies closely resembled those described by Bodenstern (1957). So well, in fact, does his description of *in vivo* regeneration fit the activities found in the present studies that there can be little question that the *in vitro* findings are representative of the process of regeneration as it normally occurs. In both cases, (a) fibers emerge from the proximal and distal nerve stumps and cross a cell bridge made up of blood and connective tissue cells, (b) fiber bundles can be traced from the proximal to the distal nerve stump, (c) many fibers become lost wandering from the target tissues and sometimes form loops and coils that go nowhere, (d) fibers from ganglia and leg regenerate stumps often reunite with other ganglia or leg stumps instead of forming a union between the distal and proximal stumps, and (e) fibers from the distal and proximal stumps may meet in the middle of the cell bridge and form a reticulum of fibers.

Bodenstern also pointed out that nerve regeneration occurred in adult insects and was independent of the molting cycle and thus of the influence of the prothoracic gland. In our study, the prothoracic gland had no visible effect on the growth of nerve fibers from either the leg regenerate or the prothoracic ganglion. This evidence strongly supports the earlier conclusion of Marks and Reinecke (1965) that nerve regeneration is a noncyclical or wound-healing type of growth rather than the cyclical, molting-dependent type found in nymphal epithelial tissues.

One question was whether observable interactions occurred between the leg regenerate and the ganglion tissues. In the cultures containing both types of tissues, fiber growth occurred in 81% of the leg regenerates and 85% of the ganglia. When these same tissues were cultured in isolation, 44% of the regenerates and 90% of the ganglia showed such growth. The results are similar to those from an earlier study (Marks and Reinecke, 1965) in which fiber growth was found in 64% of 11 regenerates when the ganglion was present and in 29% of 24 regenerates when the ganglion was absent. While neither tissue required the presence of the other in order to initiate regenerative growth, the presence of the ganglion significantly increased the occurrence of nerve growth in leg regenerates. However, the presence of the leg regenerate had no measurable effect on the growth from either the ganglion or the nerve stump of other leg regenerates

in the chamber. This is evident when these results are compared with those of the 1965 studies. In the earlier study, only two leg regenerates were present in the control chambers, while in the present study, there were four. Yet, in both studies, the nerve growth in the chambers containing only leg regenerates was approximately 50% of that in chambers containing both leg regenerates and ganglia.

Another question concerned the directional orientation of the fibers as they emerged from the explants. During the first few days, the sensory fibers emerging from the distal nerve end in the leg regenerates grew parallel to the shaft of the nerve stump so that they appeared to have a directional orientation if the adjacent explants were close together; however this orientation subsided with time, and additional fibers emerged and grew in numerous directions, apparently independent of any directional influences. Both sensory and motor fibers appeared to have a tactile or thigmotropic response that caused them to follow the cell bridges formed by the blood and sheath cells. If the explants were over a millimeter apart or if no cell bridge formed, most fibers became lost and migrated in a random manner over the glass surface.

One additional phenomenon noted by Bodenstern (1957) was confirmed: those fibers that did eventually make contact with another explant appeared to grow in diameter and persisted in old cultures after the surrounding tissues had broken down. This can be partially explained by the thigmotropic response of the fibers causing bundles to form as new fibers migrated along the surface of the old ones. The migration of fibers from both explants over the same bridge would account for the greater size when compared with bundles formed when fibers migrated from one source only and reached a dead end.

CONCLUSIONS

1. The regeneration of nerve tissues which occurs *in vitro* is comparable, within limits, to that which occurs *in vivo*. Both sensory and motor nerve elements take part in the regenerative process. Blood and glial cells play an important role in forming the cell bridge that facilitates the migration of the nerve axons.

2. The presence of ganglion tissue in the growth chamber exerts a demonstrable stimulating influence on the outgrowth of cells from the nerve stump of the leg regenerate. However, the presence of the leg regenerate appears to have no effect on the outgrowth of cells from the ganglion.

3. A tactile or thigmotropic type of behavior by the migrating axons is evident that explains several of the phenomena described.

4. With such similarity between the development *in vivo* and *in vitro*, the advantages of the *in vitro* technique become apparent. High-resolution photography of the activities of single cells is possible, and the activity can be followed for many days by using time-lapse cinephotomicrography. Also, great experimental flexibility is possible because there need be no interference from other tissues. In addition, tissues from several insects may be included in a single chamber which may make it possible to investigate a number of phenomena that are not otherwise accessible for experimentation.

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RHYTHMIC MOVEMENTS OF CONES IN THE RETINA OF BLUEFISH, *POMATOMUS SALTATRIX*, HELD IN CONSTANT DARKNESS

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Rods, cones and retinal pigment in a number of fishes undergo alterations of shape and position in response to light and darkness (Arey, 1916; Parker, 1932; Walls, 1942; Nicol, 1965). Several fresh-water and anadromous fish species kept in continuous darkness exhibited well-defined rhythms of photomechanical changes in the retina, implying an internal control mechanism (Welsh and Osborn, 1937; Wigger, 1941; Arey and Mundt, 1941; Ali, 1961; John and Haut, 1964; John, Segall and Zawatzky, 1967; John and Gring, 1968).

We undertook the following study to determine whether the retina of the bluefish *Pomatomus saltatrix* (Linnaeus), a marine pelagic species, undergoes photomechanical changes ordinarily associated with dark- and light-adaption in the absence of light cues. The facts which prompted the study were: (a) under a natural light regime, the bluefish is diurnal and its swimming speed is associated with daily light changes but not entirely controlled by them; (b) rhythms of bluefish swimming activity are evidently in part controlled by internal mechanisms; and (c) the bluefish seems to depend on vision for capturing prey and is particularly active during morning twilight forage (Olla, 1966a, b).

The above facts led us to hypothesize some internal control for adaption in the retina. To investigate this we measured cone and pigment epithelial movement in the retinae of bluefish which we had sampled periodically while the fish were being kept in constant darkness.

MATERIALS AND METHODS

We used bluefish 12 to 17 cm. long captured in Sandy Hook Bay, New Jersey, for this study. We held experimental fish in two 75-gallon recirculating aquaria with water temperature at $22 \pm 1^\circ$ C. and salinity at 25‰. Each aquarium was illuminated by two 24-inch, 20-watt fluorescent lamps. A frosted plate glass was placed between the lamps and water surface to diffuse light evenly. Each aquarium with its lighting system was enclosed in a light-proof box. A phototimer controlled artificial day length.

Before we could establish the presence or absence of a retinal rhythm, it was necessary to determine the range of photomechanical changes in the bluefish retina under both darkness and light and the approximate time necessary to complete these changes. For dark-adaption we subjected five fish to total darkness and removed one fish at intervals of 15, 30, 60, 120, and 180 minutes after the natural

seasonal time of sunset (1930 hr.). For light-adaption we subjected five fish to a light intensity of 50 ft.-c. and removed one fish at intervals of 15, 30, 60, 120, and 180 minutes after the natural seasonal time of sunrise (0630 hr.).

Our procedure was as follows: we removed a fish at random from an aquarium, immediately decapitated it, punctured the cornea and fixed the head in Held's solution. This procedure took less than one minute. For fish sampled under darkness, we performed the procedure under illumination from a red-tinted, 15-

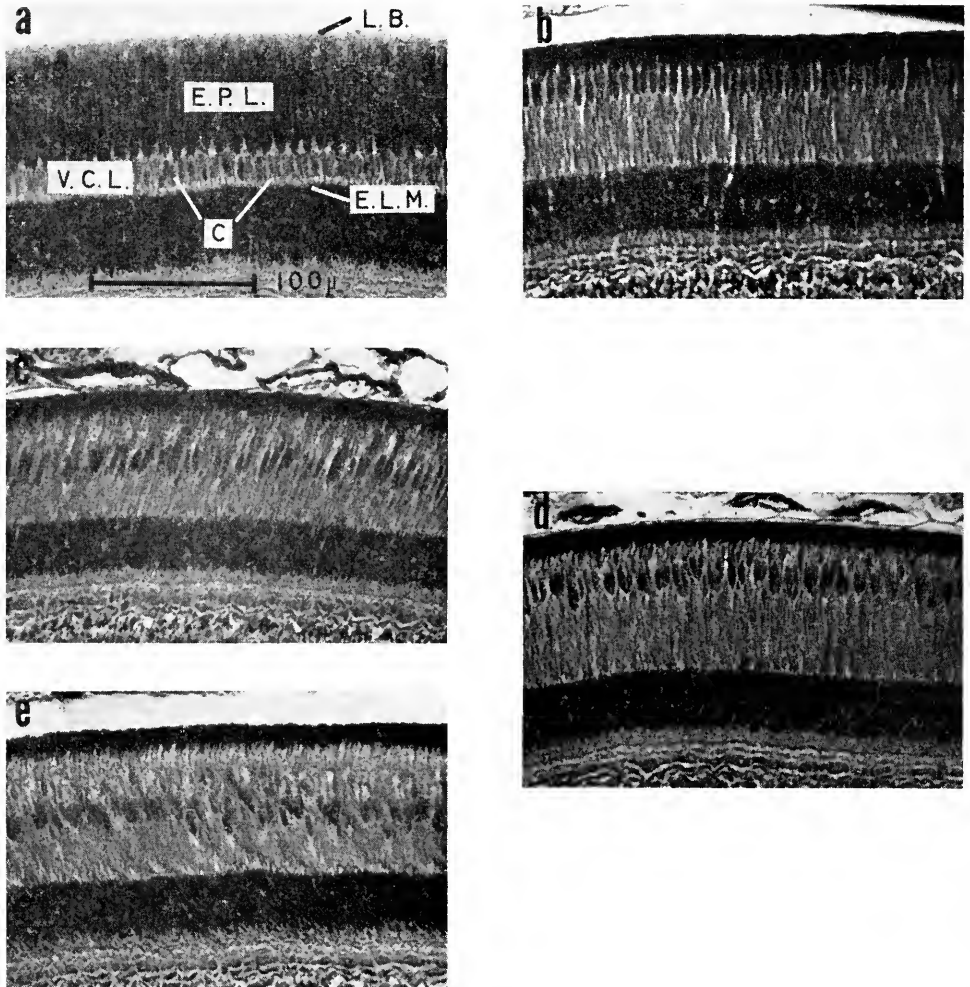


FIGURE 1. Photomicrographs showing the comparison between transverse sections of fully light- and dark-adapted bluefish retinae with those from fish sampled under constant darkness. a. Fully light-adapted retina. b. Fully dark-adapted retina. c. Retina of bluefish sampled at 1200 hours (16½ hr. after light offset). d. Retina of bluefish sampled at 2400 hours (28½ hr. after light offset). e. Retina of bluefish sampled at 1200 hours (40½ hr. after light offset). L.B., lamina basalis; E.P.L., epithelial pigment layer; V.C.L., visual cell layer; E.L.M., external limiting membrane; c, cone ellipsoids,

watt tungsten light bulb. To ensure complete penetration of the fixative, we slit the eyes at the sclerocorneal junction after they had been fixed for 24 hours in darkness. After an additional 24 hours of fixation, we enucleated the heads and removed corneas and lenses. We then dehydrated the eyes in an ethyl alcohol series, cleared them in toluene and embedded them in paraffin (mp 56.5° C.). We sectioned the embedded eyes serially in a transverse plane at 8 μ and stained with Harris' hematoxylin and eosin.

To determine the presence of a retinomotor rhythm under continuous darkness, we held the fish for seven days under an artificial photoperiod of 13 hours light of constant intensity (50 ft.-c.) beginning at 0630 and 11 hours dark beginning at 1930. We assumed that by then the fish had become acclimated to the light cycle. Then beginning at the time the light went off on the seventh

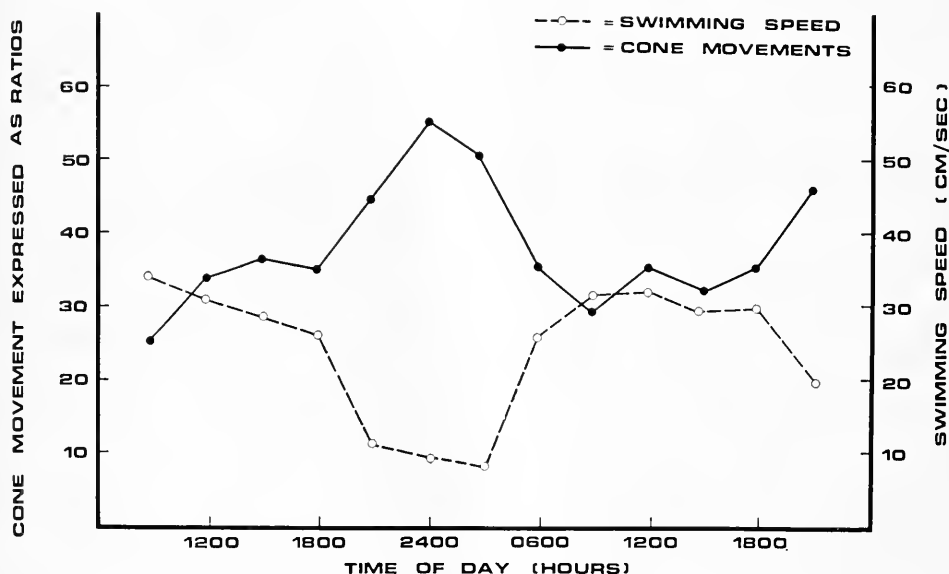


FIGURE 2. Mean values of cone position from fish held in constant darkness compared with swimming activity under constant low illumination (9 ft.-c.).

day (1930) we held the fish in constant darkness for 49½ hours. At 0900 hours of the eighth day and every three hours thereafter for 36 hours, we sampled two fish from the aquaria and prepared the eyes for histological examination in the same way as described above.

With a calibrated ocular micrometer we measured the position of 20 consecutive cone ellipsoids from one eye of each fish beginning 50 μ dorsad and 50 μ ventrad to the optic nerve insertion. We expressed measurements as percentages of the width of the combined pigment epithelial and visual cell layers to compensate for variations in retinal cell layer thickness. We did this by dividing the distance from the external limiting membrane to the proximal end of the cone ellipsoid by the distance from the external limiting membrane to the lamina basalis and multiplying the quotient by 100.

RESULTS

We found that the retinal changes associated with light- and dark-adaption followed the classical pattern for the cones and pigment epithelium (Arey, 1916; Parker, 1932; Walls, 1942; Nicol, 1965). Complete light-adaption of the bluefish retina took between 15 and 30 minutes, complete dark-adaption between 30 and 60 minutes. When the fish had become light-adapted, the cone ellipsoids lay close to the external limiting membrane and the pigment epithelium was expanded and remained in that position until the light phase ended (Fig. 1a). When the fish had become dark-adapted the cones lay close to the pigment epithelium which was in a contracted state (Fig. 1b). Thus under a regime of alternating phases of light and dark, the cones and pigment epithelium moved from one extreme to the other with each change of phase. However, under constant darkness the cones continued to move according to the natural photoperiod from the position normally associated with dark-adaption to that normally associated with light-adaption; but they did this gradually in a rhythmic pattern (Fig. 1c-e; Fig. 2). But whereas the pigment epithelium had expanded and

TABLE I

Medians of retinal cone positions expressed as the ratio of the distance between the ELM and the proximal end of the cone ellipsoid/distance between the ELM and the lamina basalis, $\times 100$; and results of the Tukey-Duckworth End Count Test

Time	Fish A		Fish B		Average of medians	End count
	Sampling location		Sampling location			
	Dorsal	Ventral	Dorsal	Ventral		
	Clock daytime					
0900	23	24	29	26	26	-
1200	31	31	34	34	33	-
1500	34	37	39	38	37	
1800	39	26	33	34	36	
	Clock nighttime					
2100	45	41	43	48	44	+
2400	53	56	56	53	55	+
0300	55	41	50	54	50	+
0600	40	37	32	29	35	
	Clock daytime					
0900	26	23	33	34	29	-
1200	35	35	38	30	35	
1500	40	41	25	24	33	-
1800	35	36	38	32	35	
	Clock nighttime					
2100	43	46	44	45	45	+

+ = Clock nighttime values larger than the largest clock daytime value.

- = Clock daytime values smaller than the smallest clock nighttime value.

Total end count 8
= 0.025*

* = 0.025 signifies that the differences between the clock daytime and the clock nighttime readings are significant at the 2.5% level.

TABLE II

Sign Test, comparing hourly differences of cone position between fish and between location sampled

Hour	Median: Fish B—Fish A		Median: Ventral sample—Dorsal	
0900	Dorsal,	+	Fish A,	+
1200	Day 1	+	Day 1	0
1500		+		+
1800		—		—
2100		—		—
2400	Dorsal,	+	Fish A,	+
0300	Day 2	—	Day 2	—
0600		—		—
0900		+		—
1200		+		0
1500		—		+
1800		+		+
2100		+		+
0900	Ventral,	+	Fish B,	—
1200	Day 1	+	Day 1	0
1500		+		—
1800		—		+
2100		+		+
2400	Ventral,	—	Fish B,	—
0300	Day 2	+	Day 2	+
0600		—		—
0900		+		+
1200		—		—
1500		—		—
1800		—		—
2100		—		+
No. of +		14		11
No. of —		12		12

contracted with change from light to darkness, it remained contracted throughout the whole period of constant darkness.

Variation in the positions of the cones relative to the external limiting membrane in any given part of the retina could result from one or both of at least two causes: (a) differences in the part of the retina sampled, and (b) differences between specimens. To examine these points we subjected the data to the following non-parametric statistical analysis: (1) We tabulated the medians of each set of 20 measurements made dorsad and ventrad to the optic nerve insertion for each fish and each sampling hour (Table I). (2) We then tabulated in Table II the signs of the hourly differences in the position of the cones between specimens and between samples taken either dorsad or ventrad to the optic nerve insertion. Applying the Sign Test of Dixon and Mood (1946), we found no significant differences in the position of the cones between fish and sampling location. (3) Therefore, we added the median cone positions for each sampling hour in Table I and computed the averages of the totals. (4) To compare the positions of the

cones in specimens sampled during daytime and nighttime, we applied the Tukey-Duckworth End Count Test shown in Table I (Tukey, 1959). The results of this test showed the positions of the cones to differ significantly between the two groups, indicating a movement of cones in specimens kept in total darkness. This could be accounted for only by some degree of internal control. (5) We plotted the mean values of cone positions referred to in (3) above (Fig. 2) to examine the pattern of movement over the period of the experiment and for comparison purposes plotted a curve of the activity rhythm of a group of adult bluefish held under constant low illumination of 9 ft.-c. (Olla, 1966a, b).

It is apparent from examination of the two curves in Figure 2 that the dark-adapted position of the cone ellipsoids and low swimming activity of bluefish occur concurrently during the hours of the natural dark phase and conversely, a light-adapted position of the cone ellipsoids and increased swimming activity occur during the time of the natural light phase. These results provide further evidence for the existence of an internal control mechanism which can act independently of light.

DISCUSSION

Swimming activity and photomechanical changes in the retina are both related to light, are diurnal, and are under some degree of internal control. Previous work on bluefish activity rhythms (Olla, 1966a, b) compared with the present results bears out this relationship (Fig. 2). The fact that an overt response such as swimming speed and a covert response such as cone migration in the retina are in part internally controlled suggests a common synchronizing system for phasing rhythmicity. Internal control of retinal adaption for a predator dependent on vision for prey capture has an obvious selective advantage. It would predispose the retina for light and dark. Such a predisposition would effectively lessen the time for light and dark adaption.

We made no attempt to study the longevity of the retinal rhythm. Previous results of bluefish activity rhythms showed persistence of the rhythm for at least five days. In the absence of daily light cues, an eventual desynchronization of the rhythm might occur. Several investigators have described a reduction in adaption as the length of time under constant darkness increased (Welsh and Osborn, 1937; Wigger, 1941; Ali, 1961; John *et al.*, 1967; John and Gring, 1968).

Although Wigger (1941) in his work on goldfish, *Carassius auratus*, Ali (1961) in Atlantic salmon, *Salmo salar*, and John and Gring (1968) in the bluegill, *Lepomis macrochirus*, found an indication of a rhythm in the pigment epithelium of specimens kept in constant darkness, we found no such rhythm in the bluefish. In some species the pigment epithelium may be controlled internally but evidently in the bluefish it responded only to an external stimulus. A predisposition for light and dark in the epithelial pigment layer may not be as functionally critical as in the visual cells.

Nicol (1965) showed differences in responses between double and single cones in the plaice, *Pleuronectes platessa*, merry sole, *Microstomus kitt*, and sole, *Solea solea*, John *et al.* (1967) in the goldfish, *Carassius auratus*, and John and Gring (1968) in the bluegill, *Lepomis macrochirus*. We observed similar differ-

ences between double and single cone movement in the bluefish but did not differentiate between them in our measurements.

Th question of whether retinal rhythmicity in the bluefish becomes changed by manipulation of the light cycle remains for further experiments.

We wish to express our grateful appreciation to Mr. Enoch B. Ferrell for his advice on statistical treatment of the data.

SUMMARY

1. The retina of bluefish, *Pomatomus saltatrix*, undergoes photomechanical changes in response to light and darkness. Complete light-adaption requires between 15 and 30 minutes, complete dark-adaption between 30 and 60 minutes.

2. Retinal cones of juvenile bluefish held under continuous darkness for two days exhibited a diurnal retinomotor rhythm. The epithelial pigment layer remained in a dark-adapted condition throughout the two days and displayed no discernible expansion.

3. Results were related with activity rhythms of bluefish and suggested that internal control of a retinomotor rhythm may predispose the eye to environmental changes in light intensity.

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UREA AND TRIMETHYLAMINE OXIDE LEVELS IN ELASMOBRANCH EMBRYOS

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Urea and trimethylamine oxide (TMAO) have long been recognized as important osmotic constituents of elasmobranch body fluids. Together, these organic solutes make up roughly one-half of the osmotically active constituents of elasmobranchs, allowing them to remain osmotically superior to the surrounding sea water. Eggs of elasmobranchs are nearly isosmotic to sea water (Dakin, 1911), and like the adults, they contain significant quantities of urea (Krukenburg, 1888). Smith (1936), in reviewing urea retention and modes of reproduction in elasmobranchs, concluded that the tendency of elasmobranchs toward intrauterine development may be related to its adaptive advantage in protecting early embryos from urea loss. More recently, Price and Daiber (1967), in their study of the intrauterine environment of ovoviviparous and viviparous forms, have concurred with Smith's conclusion.

In what appears to be the only detailed study of the amounts of urea at various developmental stages of elasmobranchs, Needham and Needham (1930) found that in the dogfish *Scyllium canicula*, low amounts of urea are present when the eggs are laid, and that this urea is added to by the embryos as development proceeds (see also Needham, 1931). Although Needham and Needham did not measure urea concentrations directly, they suggested that urea levels are lower in the undeveloped eggs than in the maternal fluids. They found that only a minimal amount of urea was lost from the developing embryo, and concluded that the increased urea content during development must be due to the excretion of urea by the embryo into the yolk. Their conclusions were rendered uncertain, however, by the lack of a good series of weighings for *Scyllium* embryos, and more importantly by their failure to measure urea in the embryos and yolks separately.

In adult elasmobranchs, the major pathway of urea synthesis appears to be *via* the ornithine-urea cycle (Schooler *et al.*, 1966). Recently (Read, 1968), evidence was presented of an ornithine-urea cycle early in elasmobranch development. Embryos of the oviparous *Raja binoculata* and the ovoviviparous *Squalus suckleyi* were found to contain ornithine carbamoyltransferase and arginase, and the specific activities of these ornithine-urea cycle enzymes underwent marked increases early in development. Neither of these enzymes was found in the yolk, suggesting that any urea formed during development must be supplied by the embryo. This work thus supported the finding of Needham and Needham (1930) that elasmobranch embryos are capable of producing urea.

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In contrast to urea, it is not known whether elasmobranchs are capable of producing TMAO. In what appears to be the only study of TMAO levels in elasmobranch embryos, Goldstein *et al.* (1967) found TMAO in both the yolk and body fluid of embryonic pups, and concluded that active retention of TMAO by the dogfish (*Squalus acanthias*) begins early in development. They were unable to detect synthesis of TMAO by adult elasmobranchs from radioactively labeled compounds (methionine, choline, and trimethylamine), but they did not rule out its synthesis by pathways not involving these compounds. Baker *et al.* (1963) presented evidence supporting a conversion of trimethylamine to TMAO *in vitro* with preparations from elasmobranchs. Cohen *et al.* (1958) found that adult elasmobranchs (*S. acanthias*) starved for up to 41 days were able to retain high blood concentrations of TMAO. Although this finding would indicate that TMAO was being endogenously produced, they suggested that the highly active reabsorption of TMAO by the kidney and its replenishment from TMAO reserves in the muscle could account for its stability.

The present study was undertaken to find out whether either the concentrations or total amounts of urea and TMAO increase during development of the skate *Raja binoculata*. During development, the oviparous elasmobranch embryo appears to be a nutritionally closed system, entirely dependent on its yolk for organic materials. It was felt that if it could be shown that the amount of TMAO is maintained or increases during development, this would indicate that the embryo is capable of producing this compound. In addition, a study was made of urea loss by the different developmental stages of this species.

MATERIALS AND METHODS

Egg cases of the skate *Raja binoculata* were obtained by otter trawl from the region of Bellingham Bay, Washington. They were transported to the Friday Harbor Laboratories, where they were kept in running, aerated sea water. They were collected on two occasions: late December, 1967, and mid-February, 1968. Egg cases of this species are large, being about 30 cm. in length. Each egg case contains four openings, which during the earliest stages are plugged with solidified albumin. Later the albumin plugs dissolve, allowing sea water to enter. Each egg case contains from one to seven embryos. At any given time during the year a complete range of developmental stages may be obtained, thus facilitating their study.

The measurements made using embryos and their yolks were (i) total volume, (ii) urea concentration and urea content, (iii) TMAO concentration and TMAO content, (iv) dry weight and water content, (v) total urea loss per day, and (vi) freezing point depression of the yolks.

The volumes of the embryos and their yolks were measured separately by their displacement of water. For urea determination a homogenate was prepared using nine times their volume of distilled water. Embryos were homogenized in a Waring Blendor, and yolks were shaken thoroughly with water. In a number of cases, embryos were homogenized together with their yolks, so that the urea concentration in the entire system was measured. The homogenates were then centrifuged for 5 minutes at approximately 600 *g* and a sample of the supernatant taken for urea determination. Urea was measured colorimetrically by nessleriza-

tion following treatment with urease. Routine standards were prepared with ammonium sulfate, and the entire method was tested using urea standards.

For TMAO determinations, homogenates were prepared as above using only four volumes of distilled water. The entire homogenate was then extracted with equal its volume of 20% TCA for two hours with constant shaking at room temperature. TMAO was measured by the methods of Dyer (1945) and Dyer *et al.* (1952). Using these methods, the TMAO in the TCA extracts is first reduced using Devarda's alloy to trimethylamine, which is then measured colorimetrically as a picrate salt. It was found necessary to carefully filter the solutions after extraction with TCA and again after reduction with Devarda's alloy to obtain consistent results.

Freezing point depressions of the yolks were measured by the method of Gross (1954). The yolks were first centrifuged at 16,000 *g* for 30 minutes at 4° C. This caused the formation of a small quantity of translucent supernatant, a sample of which could then be used for freezing point determination. Following Krogh (1939), 293 *mM* of NaCl were taken as giving a freezing point depression of 1° C.

Dry weight and water content were determined by placing macerated embryos or yolks in a drying oven at 45° C., and measuring weight loss during consecutive 24-hour periods until no further loss was noted.

In experiments designed to measure urea loss by animals at different developmental stages, embryos were placed separately in covered dishes containing known amounts of Millipore-filtered sea water (usually 250 ml.). Before being placed in the dishes, the embryos were rinsed with additional filtered sea water to reduce the chance of urea loss through the action of diatoms and bacteria. Each dish was well aerated, and after 24 hours urea was measured in samples of sea water taken from the dishes. Urea was measured colorimetrically using 1-phenyl-1,2-propanedione-2-oxime (Archibald, 1945), with urea standards being made up in sea water. Controls included treatment of duplicate samples of the sea water with urease to make certain that urea was indeed being measured. In addition, dishes containing filtered sea water with known quantities of urea were treated exactly as those containing the embryos. There was no significant loss of urea from these dishes during the 24-hour period.

Because of the highly allometric growth pattern of developing skate embryos, length was felt to be an unreliable measure of developmental stage. Values are therefore compared in terms of wet weights of the embryos. The largest of the embryos used in this study still contained yolk in their guts, and were therefore not likely to be taking in nutrients from the environment.

RESULTS

The total amount of urea per embryonic system (embryo plus yolk) was found to increase markedly during development (Fig. 1), confirming a similar finding by Needham and Needham (1930). The concentration of urea in the overall system also increased as a result of a higher urea concentration in the developing embryo than in its yolk (Table I). This overall increase in urea concentration was not, however, great enough to account for the large increase in the total urea content of the system.

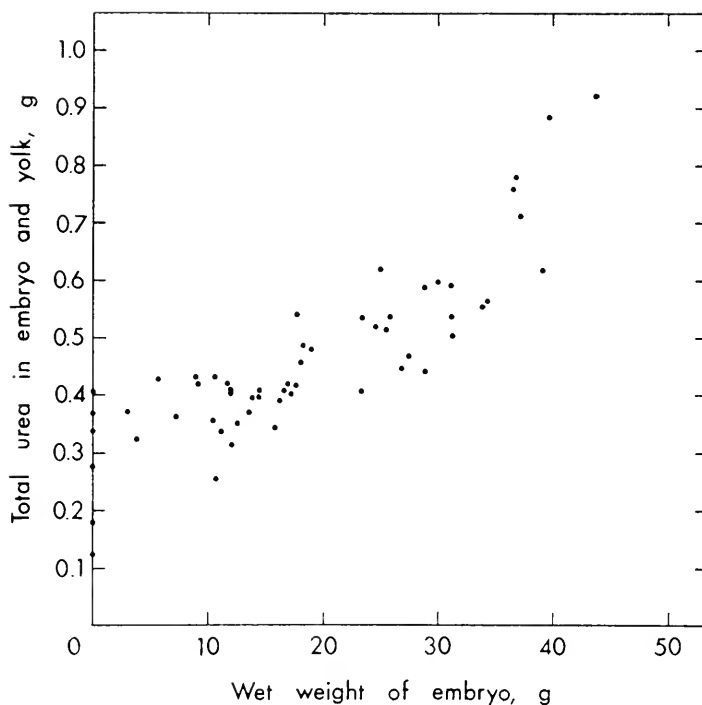


FIGURE 1. Total urea content of the embryonic system (embryo plus yolk) with increasing wet weight (*i.e.*, development) of the embryo. All values in this and remaining figures refer to embryos of the oviparous elasmobranch *Raja binoculata*.

A large increase in the total urea content (Fig. 1) without a similarly large increase in urea concentration can only occur if there is an increase in the total volume of the system during development. This was found to be the case (Fig. 2). As the embryo grows at the expense of its yolk, its volume increases faster than the volume of yolk decreases, leading to a net increase in the total volume. The considerable scatter in the values in Figure 2 reflects variations in the volume of the yolk at any given stage. The final size of the embryos at the time their yolks have been completely absorbed is also highly variable, undoubtedly because of variations in the initial volumes of their yolks.

TABLE I

Concentration of urea and trimethylamine oxide (TMAO) and percentages of water in embryos and yolks of *Raja binoculata*. Data are expressed as mean values \pm standard deviation, and the number of observations is in parentheses

	Urea mg.%	TMAO mg.%	Per cent water
Embryos	1817 \pm 180 (18)	542.4 \pm 119 (28)	77.2 \pm 4.6 (10)
Yolks	1242 \pm 135 (16)	718.4 \pm 107 (28)	51.3 \pm 9.1 (15)

The increase in total volume of the system (Fig. 2) was found to be due to an increase in its water content. This, in turn, was correlated with a higher percentage of water in the developing embryo than in its yolk (Table I). The increase in the total volume was gradual, and dependent on the growth of the embryo. The relative percentages of water in embryos and yolks did not change significantly during development. The present finding of an increase in water content confirms the early work of Ranzi (1932), who found that elasmobranch embryos obtain most of the water they require during development from their environment.

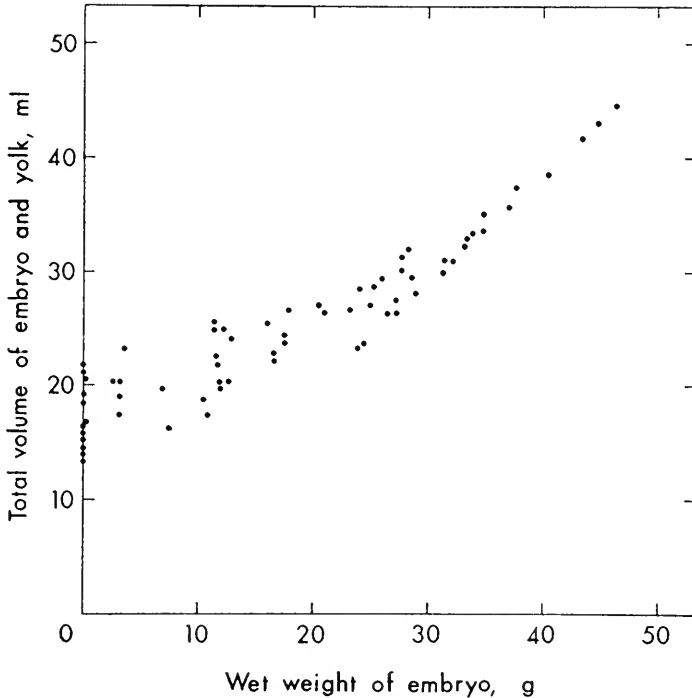


FIGURE 2. Total volume of the embryonic system with increasing wet weight of the embryo.

The difference in urea concentration between embryos and yolks can be correlated with their difference in water content. When the average values for urea concentration in the embryos and yolks are corrected for water content, they are nearly equal (2,354 mg. per 100 ml. water in embryos *versus* 2,420 mg. per 100 ml. water in yolks). Values for urea concentration in the embryos are within the range of urea levels found in various body fluids of adult elasmobranchs (see Bernard *et al.*, 1966). Differences in water content may also help to explain the greater average density of the yolks (1.15 mg./ml.) than of the embryos (1.04 mg./ml.).

The dry weights of embryos plus their yolks ranged between 6.92 and 11.48 g. There did not appear to be a marked change in the dry weight of the system during

development, although only ten complete systems were measured. Ranzi (1932) found that embryos of the oviparous elasmobranch, *Scyllium canicula*, show a gain in inorganic substance but a larger loss in organic substance during development. This led to a net loss of approximately 15% in the overall dry weight of the system. At least part of the loss in organic substance observed by Ranzi was probably due to a loss of urea (see below).

The finding that the relative urea concentrations in embryos and yolks remained constant during development is evidence that urea is not excreted into the yolk, as Needham and Needham (1930) suggested. Values for the total loss of urea

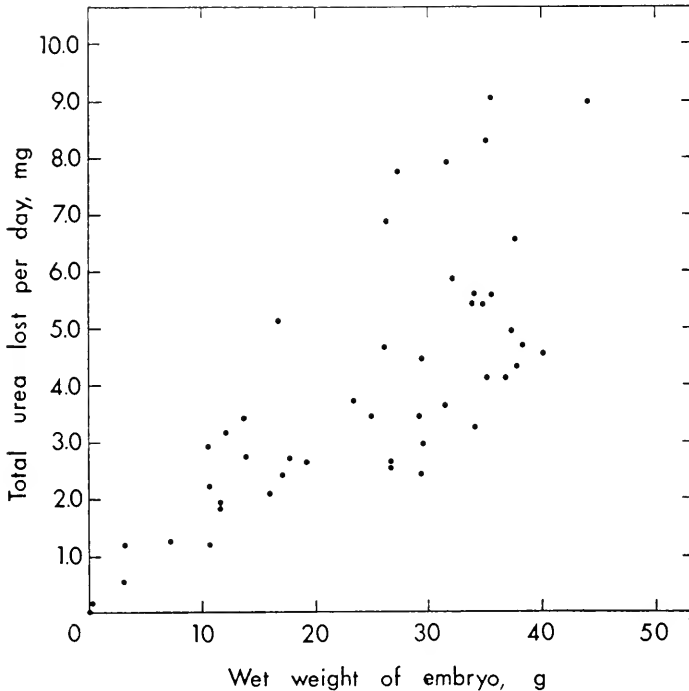


FIGURE 3. Total urea loss per day from the embryonic system with increasing wet weight of the embryo.

from the embryonic system into the surrounding sea water are given in Figure 3. The amount of urea lost per day, although highly variable, increased with the increasing size of the embryo. The amount of variation also became greater as development proceeded. The average rate of urea excretion was 0.24 mg./g. wet weight of embryo per day. No urea, however, was detected in sea water or in the albumin surrounding undeveloped eggs. The extremely delicate membrane of these eggs is remarkable in that it appears to be able to retain urea against a tremendous concentration gradient until such time as the embryo is capable of producing urea. It should be emphasized here that the retention of urea by the undeveloped egg appears to lie solely with its delicate membrane. Needham

and Needham (1930) found that the egg case walls are permeable to urea, so that even in the early stages, when openings in the egg case are plugged with albumin, urea lost by the embryonic system can apparently diffuse to the environment. The enclosure of the eggs in egg cases should therefore not be viewed as a mechanism for urea retention. In the present study it was found that embryos removed from egg cases whose openings were plugged with albumin were able to withstand exposure to sea water for up to a week before losing their integrity and becoming infected with bacteria.

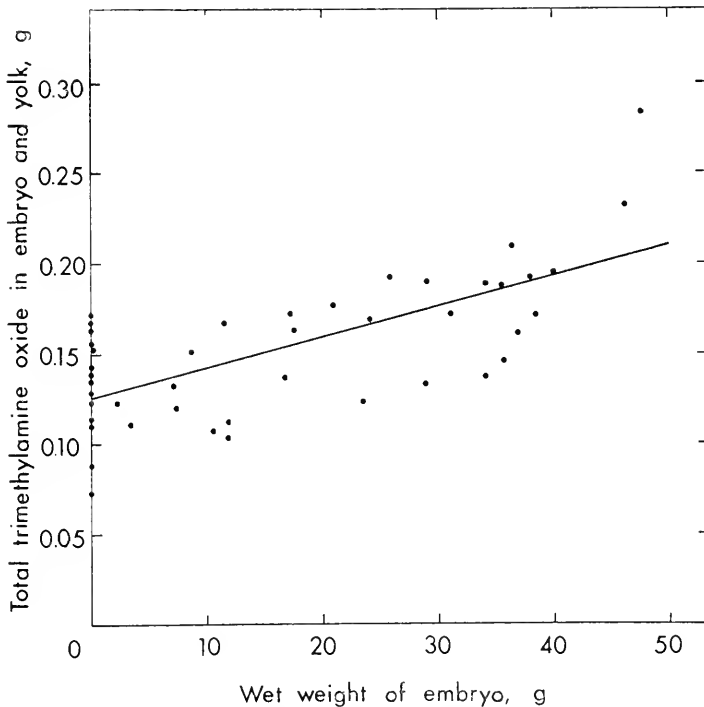


FIGURE 4. Total trimethylamine oxide content of the embryonic system with increasing wet weight of the embryo. The regression line shown has a positive slope of 0.311.

The freezing point depression of the yolks was relatively constant and nearly isosmotic with sea water. The average values for freezing point depression were 1.83 for yolks *versus* 1.82 for sea water.

There was no significant change in the concentrations of TMAO in the embryos and yolks during development (Table I). However, in contrast to urea, TMAO concentrations were found to be consistently higher in yolks than in embryos, even though the difference in the mean values for TMAO (Table I) was not statistically significant. When corrected for water content, the average value for TMAO concentration is twice as high in the yolks as in the embryos (1,410 *versus* 689 mg. per 100 ml. water). The overall concentration of TMAO in the system de-

creased during development. This decrease was correlated with the lower TMAO concentrations in the developing embryos than in their yolks. This, in turn, meant that the total amount of TMAO did not increase as markedly as did urea. However, there did appear to be a definite increase in total TMAO (Fig. 4). A regression line calculated from the values in Figure 4 had a positive slope of 0.311, which, with a standard error of ± 0.069 , differs significantly from zero. (95% confidence interval = 0.202–0.420). The increase in the total volume of the system was greater than the decrease in the TMAO concentration, indicating a net increase in the total amount of TMAO. This increase is evidence that the embryo is capable of producing TMAO, particularly because at these stages the embryo is a closed system with respect to organic substrates. Even if there were no increase in total TMAO, the maintenance of high and relatively stable concentrations of TMAO throughout development would seemingly support the conclusion that TMAO is being produced. Attempts to detect TMAO in the sea water surrounding those embryos used in the experiments designed to measure urea loss were unsuccessful, possibly due to limitations in the sensitivity of the method. The loss of less than about 0.2 mg. of TMAO into the volume of sea water used in these experiments would not have been detected.

DISCUSSION

It seems clear from the present study that embryos of the oviparous elasmobranch *Raja binoculata*, in their ability to retain high urea and TMAO concentrations, have an osmotic relationship to their environment which is similar to that of the adults. The concentration of urea, when measured on the basis of water content, is high and nearly constant throughout development and probably into the adult stage. The concentration of TMAO also remains at a high level, and throughout all stages it is somewhat lower in the embryo than in the yolk. Increases in the total amounts of both urea and TMAO are recognizable mainly through increases in the total volume of the system, since the concentrations of these solutes either decline or remain nearly constant. This increase in volume is in turn due mainly to an increase in water content.

The present report does not in any way invalidate the experimental findings of Needham and Needham (1930), but rather it offers other, and seemingly more reasonable, interpretations of their data. The increase in total urea which they observed during the development of *Scyllium canicula* can be correlated with a correspondingly large increase in water content observed by Ranzi (1932) during development of the same species. Furthermore, Needham and Needham were able to detect urea in the sea water surrounding embryos studied in urea-loss experiments, but they concluded that the amount of urea lost was not significant when compared to the large quantities contained within the embryos. From an osmotic viewpoint, continual loss of even a small amount of urea may be highly significant, particularly if measured over the entire developmental period.

The large variation in the amount of urea lost by embryos of any given developmental stage (Fig. 3), and the relatively stable internal urea concentrations (Table I) may indicate that the production of urea and the utilization of nitrogen-containing compounds are themselves variable, and that the level of urea is con-

trolled at the site or sites of urea loss rather than at the sites of production.² It seems apparent that urea produced during the development of elasmobranch embryos serves as both an osmoregulatory and an end product of metabolism. It should be noted, however, that the conditions under which the urea-loss experiments were run may differ markedly from conditions within the egg case, particularly with regard to the amount of oxygen and carbon dioxide as well as conditions of light and trauma. Possible sites of urea loss were not sought in the present study. Perhaps the most obvious site would be the gill filaments, which are highly developed at an early stage.

The finding of high concentrations of TMAO in even the earliest developmental stages indicates that TMAO, like urea, is retained throughout the elasmobranch life-cycle. The finding that the concentration of TMAO in the yolk is higher than in the embryo, particularly when based on water content, may indicate that this compound is taking the place of osmotic constituents which are at lower levels in the yolk than in the body fluids. However, it is possible that some of the TMAO is bound in the yolk and therefore not osmotically active. The significant increase in total TMAO found in these embryos would appear to be the strongest evidence put forth to date that any elasmobranch is capable of producing this compound.

In reviewing the role of urea in elasmobranchs, Smith (1936) concluded that the tendency of elasmobranchs toward intrauterine development (ovoviviparous and viviparous forms) might be explained by the advantage conferred by this pattern of development in allowing the embryos to better retain concentrations of urea. This view has more recently been supported by Price and Daiber (1967) who concluded (p. 259) that "the inability of the embryo to regulate urea and osmotic pressure in early development has been a selection pressure leading the elasmobranchs toward viviparous reproduction." As seen in the present study, both the embryo and the undeveloped egg are apparently able to retain urea against the diffusion gradient which exists between them and either the albumin or sea water. Furthermore, levels of both TMAO and urea are maintained within relatively narrow limits, which suggests strongly that regulation is occurring throughout development. It seems more likely, therefore, that the initial selective pressure toward intrauterine development resulted from factors other than the need to retain urea (*e.g.*, protection against predation and mechanical injury). Once having adopted an ovoviviparous mode of development there may then have been other selective pressures to maintain closer nutritional ties with the maternal organism.

I would like to thank Dr. Robert L. Fernald, Director of the Friday Harbor Laboratories, for use of the facilities. I am also grateful to Drs. Ingrith Olsen, Aubrey Gorbman, and George W. Brown, Jr., and to Charles Lambert, for their advice and help with various aspects of the study and the preparation of the

² Studies which I have recently completed indicate that, at least in adult elasmobranchs, there may be a number of possible sites of urea production in addition to the liver. Activities of both ornithine carbamoyltransferase and arginase are clearly detectable in a number of tissues, particularly the spleen and kidney.

manuscript. The continual interest and help of Dr. Eugene Kozloff is also greatly appreciated.

The work was supported by N.S.F. grant GB-3386 to the Friday Harbor Laboratories and by a N.A.S.A. predoctoral fellowship.

SUMMARY

1. The total amounts of urea and trimethylamine oxide (TMAO) increase during the development of the oviparous elasmobranch *Raja binoculata*. These increases are not explained by increases in the concentrations, which, when measured on the basis of water content, either decline (TMAO) or remain relatively constant (urea).

2. The increases in urea and TMAO are correlated with an increase in the total volume of the embryonic system during development.

3. The increase in the total volume of the system is due mainly to an increase in its water content. This, in turn, is correlated with a higher percentage of water in the developing embryo than in its yolk.

4. The increase in the total amount of TMAO during development would strongly suggest that the embryo is capable of producing this compound, particularly since the embryo is apparently a closed system with regard to organic substrates.

5. There was a definite, though variable, loss of urea from the embryonic system but not from the undeveloped egg. There was, however, no detectable loss of TMAO. The amount of urea lost per day and the degree of variability increased during development. The undeveloped egg is apparently able to retain urea until such time as the embryo is capable of producing this compound.

6. The maintenance of relatively constant urea and TMAO concentrations implies that regulation occurs throughout development, and that these organic solutes play a similar osmotic role in the embryonic system as in the adult.

7. The ability to retain relatively constant urea levels by even the earliest stages of this oviparous form would seem to make less tenable the suggestion of other authors that the need to retain urea was a selective pressure leading the elasmobranchs toward intrauterine development.

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PEDAL EXPANSION IN THE NATICID SNAILS.
I. INTRODUCTION AND WEIGHING EXPERIMENTS¹

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Possession of an enormously expanded foot, partially enclosing the shell, is characteristic of living naticid snails (or "moon-snails"), and is commonly employed as a diagnostic feature for the group. The adaptive significance of this apparently disproportionate pedal expansion is two-fold. First, the expanded wedge-shaped propodium and the metapodial folds covering the shell obviously contribute to the efficiency of "plowing" locomotion, as the moon-snail moves along on, and through, sandy substrata. Secondly, the naticids are predaceous carnivores with protrusible proboscides, and employ their extensible pedal lobes to manipulate and enfold their molluscan prey. Physiological aspects of this expansion are reported in the present paper, and in the succeeding one (Russell-Hunter and Apley, 1968), the existence of a pedal water-sinus system is demonstrated conclusively, and the functioning of the naticid water-sinus system is discussed.

In his recent, and otherwise excellent, review of locomotion in molluscs, Morton (1964) largely dismisses the possibility of water uptake being involved in naticid expansion, and notes that no modern workers (over the last eighty years) have claimed to demonstrate any aquiferous system of pedal canals. In the last volume of her fine, comprehensive survey of the invertebrates, Hyman (1967) similarly denies the occurrence of internal spaces for water in the naticid foot. Thus, there is clearly a need for an unequivocal demonstration of the use of a pedal water-sinus system in the expansion of naticids, and it is hoped that the present papers will provide it. However, a brief review of why Morton, Hyman, and others were misled by earlier published accounts, and reached their erroneous conclusions seems required.

HISTORICAL INTRODUCTION

In the early nineteenth century, a number of workers on the Mollusca believed that expansion of various structures in a wide variety of marine molluscs was due to the direct uptake of sea water into the blood spaces. Molluscan anatomists from the time of Cuvier onwards had clearly described the large number of protrusible or distensible structures: tentacles, proboscides, ctenidia, penes, and the mantle and foot themselves, which structures in typical molluscs contained only *retractor* musculature. Dilation with fluid to turgescence was early recognized as a characteristic method of protraction. Universal theories involving sea-water

¹ Supported by a research grant, GM 11693, from the National Institutes of Health.

uptake in the mechanisms of pedal dilation in all bivalves and snails were dismissed after a number of workers in the 1880's, but most notably Carrière (1882) and Ray Lankester (1884), had shown that, in many different species in both classes, dilation was achieved by the influx of blood withdrawn from other parts of the body. It remains true that the majority of molluscan organs are dilated by hydraulic mechanisms involving the blood within the spaces of the hemocoel. Carrière (1882) also gives a fairly extensive account of the earlier literature. At almost the same time Schiemenz (1884, 1887) claimed to have demonstrated a system of "aquiferous" tubes in the foot of a naticid snail. His observations were made on the relatively large form, *Polinices josephinus*, and they were undoubtedly accurate as regards both the anatomy and the functioning of the system. Schiemenz had earlier been, with Carrière and Ray Lankester, among the group of malacologists strongly opposed to the more general theories of sea-water uptake, but he chose to work on a representative of what now appears to be the *only* group of gastropods to have a water-sinus system—the family Naticidae. As might be expected, the generalization of hemal distension was widely accepted, and Schiemenz's exceptional case ignored or even denied (see, for example, such general surveys of the Mollusca as those of Lankester, 1883, 1891; Simroth, 1879, 1896–1907; and Pelseneer, 1906).

It should be remembered that earlier workers on the mechanics of movements in soft-bodied animals were sometimes confused as to the physiological capabilities of muscle. Simroth (1879), for example, even invoked "active extension" of muscles in discussing molluscan mechanics. The fact that muscles in *all* animals can exert force *only by contracting* had been established by the beginning of this century. The concept of antagonistic sets of muscles acting around a hydrostatic or hydraulic skeleton in soft-bodied animals became clarified, and an early classic account of the molluscan hemocoel as the "force-transmitting" skeleton acted upon by antagonistic muscles is that of Biederman (1905).

Later workers on snail pedal anatomy and mechanics (among them; Weber, 1924; Rotarides, 1934; Lissmann, 1945a, 1945b) emphasized the importance of the hemal hydraulic skeleton. Perhaps the first worker since Schiemenz to investigate problems of dilation in a naticid was Morris (1950) who, working on the smallish Australian species, *Uber strangei*, produced rather inconclusive evidence of water uptake, and claimed that partial expansion was possible in air. A group of unrelated gastropods where the foot undergoes turgid hypertrophy encompasses the South African species of *Bullia* (family Nassariidae), whose ecology closely parallels that of the naticid genus *Polinices*. Once thought to involve water uptake (Gilchrist, 1916), modern investigation of pedal expansion in *Bullia* (Brown and Turner, 1962; Brown, 1964), using such methods as radiography with the blood loaded with radio-opaque dyes, has shown the dilation to be due to blood movement alone. In the monograph which incorporates their extensive and detailed synthesis of existing knowledge of prosobranch snails, Fretter and Graham (1962) mention the work of Schiemenz, but note that in the smaller British naticid species, *Natica catena* and *N. alderi*, they could find no pedal water pores. In his excellent book on the mechanical and evolutionary aspects of body-cavities in metazoans, Clark (1964) discusses locomotion in naticids, but regards the foot in all gastropods as consisting of erectile tissue, dilated

by blood. Thus Morton (1964) in his review was perhaps justified in dismissing water uptake as a feature of naticid expansion, and in stating, "No worker has verified Schiemenz's canals. . . ."

The present work was carried out at Woods Hole at different times in the summers from 1961 to 1966. The existence of a pedal water-sinus system in the naticid, *Polinices duplicatus*, and the use of inulin-labelled sea water in its investigation, were briefly reported (Russell Hunter and Apley, 1965) in a note on temporary hyperthermia in that species. While the present results were being prepared for publication, Bernard (1968) has independently demonstrated uptake into a pedal water-sinus system in the West-Coast species, *Polinices lewisi*, though with weighing experiments involving small numbers of narcotized snails, without direct records of volumes of water involved, and without labelling experiments. The present paper reports the results of extensive weighing experiments on "trained" un-narcotized snails, including the relation of different degrees of expansion to the water volumes involved, and the construction of "balance sheet" equations involving the mass of the shell and its actual capacity. The succeeding paper (Russell-Hunter and Apley, 1968) describes and discusses the use of inulin-labelled sea water in the investigation of the relation of the various water spaces and of the rates of water exchange. A third paper is being prepared on the micro-anatomy and histology of the naticid pedal water-sinus system.

MATERIALS AND METHODS

On anatomical grounds, the family Naticidae (in the superfamily Naticacea) is placed in the suborder or order Mesogastropoda (formerly the Taenioglossa). It is worth noting that most other familiar families of predaceous gastropods using protrusible proboscides, including the Muricidae (e.g. *Urosalpinx*), Buccinidae, Nassariidae, and Conidae, differ anatomically from the naticids and are classified in the suborder or order Neogastropoda (formerly the Stenoglossa). Most work was done on the two commonest species of naticids at Cape Cod, *Polinices duplicatus* and *Lunatia heros*, collected on tidal flats at West Barnstable Harbor, at Orleans Town Pond, and at Duxbury. *P. duplicatus* survives lowered salinities and high temperatures, is the better adapted littoral species and is abundant in certain localities (Russell Hunter and Grant, 1966). *L. heros* achieves the largest size of local naticids, is less resistant, and is essentially a sublittoral species colonizing the intertidal in certain areas. A few observations have also been made on four smaller species, *P. immaculatus*, *L. triseriata*, *Natica clausa*, and *N. alderi*. All experimental snails were marked with individual numbers (using various colors of model "dopes" and nail varnishes) and specimens of *P. duplicatus* were maintained in healthy condition in laboratory aquaria for over two months.

The present experiments all depend on the fact that it proved possible to habituate individuals of the two larger naticid species to handling. After a few hours of training, a snail would remain expanded on being transferred to a dry aluminum weighing tray to be weighed in air. There was considerable individual variation in responses and in retention of responsiveness even among snails of similar size.

A triple-beam, and subsequently a torsion balance, were used for the weighings (all to the nearest 0.1 g.). An engineer's dial caliper was used to measure the shells—the measurement recorded here as "shell length" being actually the greatest

transverse diameter of the globose shell, from the outer edge of the peristome to the opposite periphery of the last whorl of the shell. The closed volumes of shells were determined by displacement weighings, each cleaned shell being filled with plaster of paris.

In the majority of weighing experiments a numbered snail, of known shell length and contracted weight, was allowed to expand. It was then transferred

TABLE I

Pedal expansion and retraction in Polinices duplicatus: weights in air, and expelled water volumes

Snail #	Shell length mm.	Contracted wt. g.	Estimated degree expansion	Expanded wt. g.	Weight difference g.	Actual water expelled ml.	Expansion Index
d1	61.0	106.2	7	261.6	155.4	136.3	246
d2	35.0	14.7	8	40.0	25.3	22.9	272
d3	40.9	24.7	5	53.8	29.1	29.2	218
d6	26.7	5.9	4	10.9	5.0	5.2	185
d7	52.2	46.0	10	170.2	124.2	111.7	370
d10	44.2	32.1	10	174.2	142.1	134.2	543
d12	36.8	16.9	10	52.8	35.9	34.3	312
d13	36.2	15.8	9	52.6	36.8	35.0	333
d16	33.2	12.0	7	29.7	17.7	17.2	248
d18	36.9	17.7	8	55.8	38.1	35.7	315
d19	33.6	12.8	8	31.7	18.9	18.6	248
d20	41.9	23.6	9	71.8	48.2	45.8	304
d22	35.9	14.4	5	27.9	13.5	13.6	194
d24	16.5	1.8	9	5.2	3.4	2.5	289
d27	49.6	45.1	5	86.3	41.2	40.6	191
d28	46.1	39.8	6	94.6	54.8	54.2	238
d31	43.5	33.7	5	58.9	25.2	25.3	175
d32	41.5	24.6	7	62.8	38.2	38.2	255
d35	44.8	35.0	6	76.8	41.8	41.6	219
d36	46.1	37.7	8	85.7	48.0	46.3	227
d41	41.2	26.6	7	69.0	42.4	41.1	259
d42	39.8	25.0	6	58.0	33.0	33.1	232
d44	52.5	49.7	9	154.3	104.6	104.5	310
d45	47.4	39.6	9	123.7	84.1	84.0	312
d46	46.9	33.4	10	142.8	109.4	109.2	428
d47	43.3	30.5	10	128.1	97.6	92.4	420
d49	46.6	33.4	8	90.4	57.0	57.5	271
d51	41.7	25.3	9	69.6	44.3	44.3	275
d201	22.8	3.9	5	7.4	3.5	3.5	190
d207	30.4	9.8	9	31.2	21.4	21.3	318
d209	48.1	35.6	9	121.9	86.3	86.3	342
d212	46.2	33.1	8	105.4	72.3	73.8	318
d213	43.6	26.9	9	79.2	52.3	52.4	294
d217	37.9	18.4	10	62.5	44.1	43.8	340
d219	41.3	24.3	10	72.5	48.2	47.6	298
d221	43.3	28.6	10	92.3	63.7	62.4	323
d223	29.0	8.6	8	24.3	15.7	15.9	283
d226	34.6	14.5	8	39.8	25.3	25.5	274
d228	41.2	22.5	10	70.7	48.2	46.7	314
d229	36.3	15.0	10	79.8	64.8	64.0	532
d232	44.3	31.9	5	66.5	34.6	34.3	208
d272	43.4	25.0	8	76.7	51.7	49.7	307

TABLE II
 Repeated pedal expansion and contraction in *Polinices duplicatus*

Snail #	Shell length mm.	Dates measured	Contracted wt. g.	Estimated degree expansion	Expanded wt. g.	Weight difference g.	Actual water expelled ml.	Expansion Index
d4	39.8	June 28	21.5	5	39.6	18.1	19.1	184
		June 30	21.5	7	49.6	28.1	23.1	231
		June 30	21.5	8	57.2	35.7	35.6	266
		Aug. 2	20.6	8	54.0	33.4	33.3	262
d5	47.8	June 29	39.9	8	110.7	70.8	67.0	277
		July 1	39.9	6	84.9	45.0	42.8	213
		July 1	39.9	7	85.0	44.9	45.6	213
		Aug. 2	37.3	6	82.2	44.9	44.9	220
d8	47.5	July 3	35.9	10	123.8	87.9	75.9	345
		Aug. 2	34.8	9	128.3	93.5	93.6	369
d50	44.5	July 14	29.8	8	90.4	60.6	58.9	303
		Aug. 1	30.5	10	114.9	84.4	84.3	377
d210	47.5	July 8	36.8	7	101.2	64.4	63.5	275
		Aug. 20	35.0	8	107.6	72.6	71.7	307
d270	51.1	Aug. 21	40.4	9	127.9	87.5	87.2	317
		Aug. 21	40.4	5	93.2	52.8	50.9	231
		Aug. 21	40.4	7	108.0	67.6	66.0	267

to a dry weighing tray, weighed "damp-dried" in air and then forced to contract. The volume of water discharged into the tray was measured, and then the contracted snail weighed again in air. Other experiments involved different sequences.

RESULTS

Preliminary trials soon demonstrated that the weight difference between the snail expanded and contracted was nearly always equal to the volume of water expelled on contraction. This was true for differing degrees of expansion, and for both *Polinices duplicatus* and *Lunatia heros*. Individual variation was less marked in smaller snails, which tended to expand more rapidly after repeated expansions and forced contractions. Medium-sized (shell lengths 30–45 mm.) and larger (45–70 mm.) snails mostly tended to sulk after a third or fourth forced contraction. However, about a third of the medium-sized snails continued to re-expand relatively rapidly, after ten forced contractions within four hours. Such snails are responsible for a disproportionate amount of the data in this paper, and similar snails were deliberately chosen for the inulin-loading experiments (Russell-Hunter and Apley, 1968).

The main series of weighing experiments involved 91 specimens of *Polinices duplicatus* and 22 of *Lunatia heros* in 188 and 47 cycles of weighings, respectively. Table I presents results for 42 typical specimen cycles of *P. duplicatus*, and Table II shows some results of weighings repeated at intervals of up to six weeks.

It proved necessary to set up an arbitrary scale (1–10) for the degrees of expansion achieved. Only the visually estimated degrees of expansion from 4 to 10 were actually employed. Snails remaining considerably less than half-expanded were rated—4, snails half-expanded—5, two-thirds expanded—7, three-quarters expanded—9, and “considerably more than three-quarters expanded” and fully expanded—10. Ratings 6 and 8 were intermediates between the more definable 5, 7 and 9, and the subjectivity of the assessment resulted in relatively more snails being assessed in these two intermediate grades. Such estimates of degrees of expansion are shown in Tables I and II.

The last column of both tables shows a calculated expansion index, which is $M_E/M_C \times 100$ when M_E is the expanded weight and M_C the contracted weight. In Figure 1 expansion indices calculated from 188 cycles (91 specimens) for *Polinices duplicatus* are plotted against the estimates of expansion. Frequencies of index numbers are shown as histograms, at each estimated degree of expansion, and for clarity the calculated index numbers are grouped in class intervals of five units (*i.e.*, index numbers from 243 to 247 inclusive are given the class mark 245). There is fair agreement between the visual assessment and the index calculated from the ratio of weights.

Similar results were obtained with the 22 specimens of *Lunatia heros* which were habituated to handling. “Hypersensitive,” and therefore untrainable, individuals were more common in this species. Specimens of *L. heros* of from 34.2 to 70.0 mm. shell length had contracted weights ranging from 13.6 g. to 133.3 g., and gave expanded weights from 24.5 g. to 319.6 g. The assessed degrees of expansion (over 47 experimental weighing cycles) mostly lay in the 5 to 7 range, and most of the corresponding calculated index numbers lay between 185 and 225, the highest index of expansion for the species being 262. It is interesting that index numbers can be calculated similarly for the 20 specimens of *Polinices lewisi* whose contracted and expanded weights are reported by Bernard (1968). These would average only 180, the highest index number achieved being about 202. The specimens of *P. lewisi* used were mostly large (mean contracted weight = 149 g.) and therefore comparable to the largest *L. heros* reported on above. Thus, to date, the most reproducible results from laboratory weighings—and the largest recorded expansion indices—have been derived from the more euryoecic, and more readily “trainable,” species, *Polinices duplicatus*.

Significantly, observations on smaller naticid species were disappointing. Pedal expansion and retraction have been observed in *Polinices immaculatus*, *Lunatia triseriata* and *Natica clausa* from the Cape Cod area, and in *Natica alderi* from Scotland. In all four species, retraction is a more rapid process than in *P. duplicatus* and *L. heros*, and habituation to handling well-nigh impossible. The only one of these smaller species in which appreciable discharge of water on contraction could be detected was *L. triseriata*. Measurement was difficult: the largest specimens had shell lengths of 12.2 mm., and contracted weights of about 0.41 g. Assessment of the weight (or volume) of water involved in expansion in *L. triseriata* is hampered by proportionately large variations in the amounts carried on the surface of the snail. However, with no claims to accuracy, only about 0.25 ml. uptake is involved in large specimens of *L. triseriata*. These disappointing data may help to explain the inconclusive evidence gained from Morris's

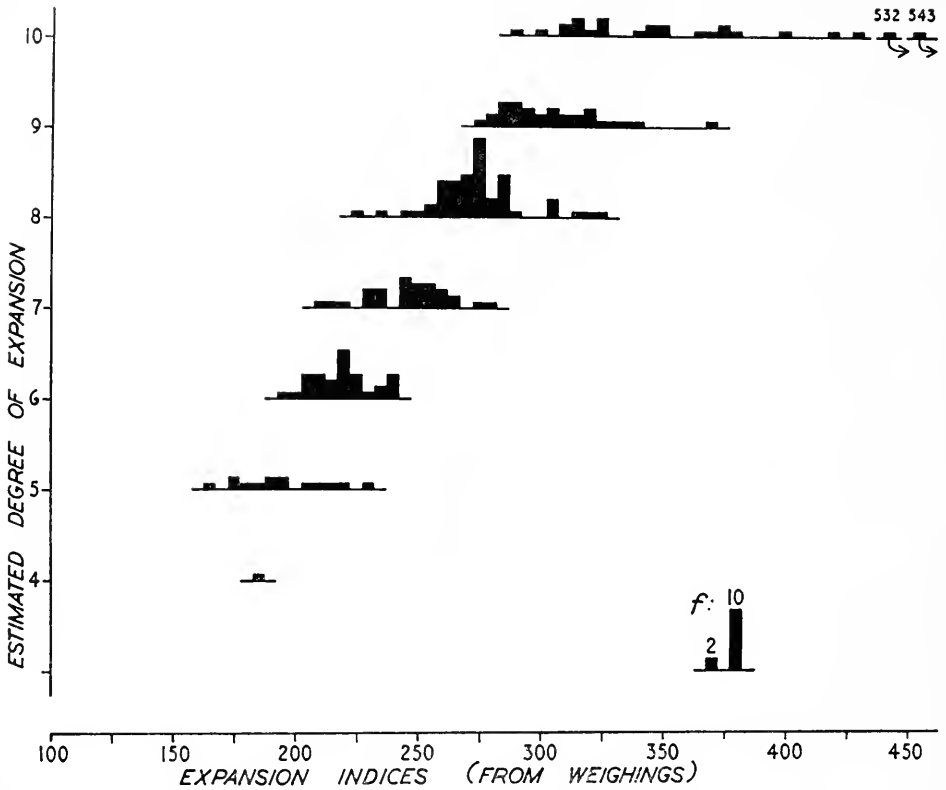


FIGURE 1. The relation of visually estimated degrees of expansion in the naticid, *Polinices duplicatus*, to the expansion indices calculated from the results of weighing contracted and expanded snails. Results derived in 188 weighing cycles from 91 individual snails are shown as histogram frequencies of index numbers. For clarity the calculated index numbers have been grouped in class intervals of five units. For further discussion see text.

(1950) study on the medium-sized Australian species, *Uber strangei*, where water uptake for most specimens must lie in the range 4–7 ml. However, results of weighing experiments on specimens of *Polinices duplicatus* as small as *Uber* were consistent and conclusive (for example, see snails d24 and d201 in Table I).

Some observations on the methods and rates of expansion and contraction in *Polinices duplicatus* are worth noting here. (Extensive records of the times involved in expansion were also made during the inulin-loading work—see Russell-Hunter and Apley, 1968.) Retraction can be rapid, particularly in non-habituated specimens newly brought from the field and, in such cases, can be accomplished in 2.5–4.0 seconds (medium-sized *P. duplicatus*). During such rapid contraction, the characteristic “jets” of sea water from the marginal pores of the water-sinus system along the sole edges of the mesopodium are seen. Experimentally forced contractions in “trained” snails usually take longer, and can take 10–15 seconds in medium-sized specimens that have been habituated to handling. During such slow “reluctant” contractions it is clear that almost all water (excepting that from

the mantle-cavity which is expelled through the *inhalant* siphon) is discharged *via* the mesopodium and its marginal pores; successive waves of muscular contraction pass over both the "plowshare" of the propodium, and the propodial and metapodial folds surrounding the shell, driving the water contained in their water-sinuses into those of the mesopodial system and thence to the exterior. An account of the anatomy and histology of the system is being prepared, but a few notes on general functional morphology may be required here. Naticids retain to a great extent the three-fold division of the primitive molluscan foot. The propodium is large, wedge-shaped when expanded, and is the plowshare of locomotion (cutting the characteristic canal-shaped—rectangular in cross-section—tracks on the surface of sand flats). With the mesopodium, it is also the organ used for enveloping the prey and manipulating it into the correct position for the proboscis to go to work. [In *Polinices duplicatus* and *Lunatia heros*, this does not necessarily mean positioning the bivalve's umbones at the mid-anterior end of the propodium ready for boring, though this is more usual. Both species consume small and medium-sized razor-clams, *Ensis directus*, at Barnstable and elsewhere, without boring, by positioning the anteroventral gape of the razor-clam for proboscis action.] The posterior edge of the propodium is directed upwards in the "propodial shield" which envelopes the anterior third of the shell when expanded. The shield also encloses the head and mantle edge, including the inhalant pallial siphon and exhalant fold, and is responsible for molding the characteristic egg-masses or sand-collars (actually shaped between the propodial shield and the shell). The mesopodium is the largest part of the foot, provides the locomotory sole and is capable of great lateral expansion in enfolding prey and in some kinds of locomotion. There are no upwardly directed folds of the mesopodium. The metapodium is archetypically the upper posterior division of the foot, last to be withdrawn in gastropod veligers after torsion, and bearing the operculum. In naticids, the metapodium when expanded conceals the operculum while its folds, shaped like a thick cup, enclose most of the posterior two-thirds of the shell. The metapodial folds have been termed epipodial, but this term might be better restricted to upwardly directed folds of the mesopodium (which in some archeogastropods give rise to characteristic sensory structures), and it is certainly incorrect (on comparative grounds, see Fretter and Graham, 1962) to term *all* the pedal structures, apart from the propodium, either "metapodium" as does Morris (1950), or "postpodium" as does Bernard (1968).

Thus the great bulk of the water discharged on retraction, and taken up more slowly during expansion, passes through the marginal pores and major water-sinuses of the mesopodium. The experiments using inulin-labelled sea water (Russell-Hunter and Apley, 1968) have shown that only about 5-7% of the volume of sea water taken up on expansion is taken in through the pallial siphon to the mantle-cavity, and only about 2% of the increased weight on expansion is due to superficial water on and between the shell and expanded surfaces. Thus about 90% of the water uptake first enters the mesopodial water-sinuses and then is pumped by contraction of the major mesopodial columnar muscle bundles into the water-sinuses of the propodium and metapodium, the marginal pores being first sealed by contraction of the smaller distal vertical muscles at the mesopodial edges. All the muscle columns are ensheathed by the lining of the

water-sinuses, consisting of a very thin pavement epithelium facing the lumina underlain by a tough layer of collagen-like connective tissue. [The water-sinuses are completely separate from the circulatory system: arterioles and branch sinuses of the hemocoel run with and through the muscle columns. Claims that blood can be involved in the fluids expelled on naticid contraction (Morris, 1950) could only be based on damaged specimens subjected to highly stressed and uncoordinated retraction. Schiemenz (1884, 1887) described the sheaths separating the water-sinuses from the tissues, and his observations are confirmed by recent work (Bernard, 1968; Russell-Hunter, in preparation).]

Normal expansion in healthy specimens of *Polinices duplicatus* takes from 3 to 8 minutes, and somewhat longer in *Lunatia heros*. Initial uptake through the minute marginal pores (about thirty on each side) depends partly on the ciliation of the epithelia around their openings but more on the "recoil" elasticity of the connective tissue sheath system. The latter forces are almost certainly augmented by the temporarily increased blood pressure *within* the muscle columns. The relative times taken in contraction and re-expansion are closely analogous to those for sea-anemones where, similarly, muscles cause contraction, and ciliary and elastic forces are responsible for expansion (Batham and Pantin, 1950). Second and successive experimental expansions within a few hours are possible in *Polinices*, but in habituated specimens the time taken in re-expansion progressively increases (mean times: for first re-expansion in a day's experiments—6.2 minutes, for fourth re-expansion within three hours—21.5 minutes).

In air, on dry surfaces, the larger naticids *cannot* re-expand. The columellar muscle can relax, which allows the operculum to gape, revealing the retracted metapodial folds and the "crumpled" mass of the mesopodium. Only when the latter is in contact with water can real expansion take place. Some experiments with medium-sized, contracted specimens of *P. duplicatus*, set up in shallow water in large weighed petri dishes, showed that expansion was possible with water depths of 3.5–4.0 mm., provided the contracted shell was placed so that the mesopodial margin came in contact with the water when the operculum gaped. Such experiments involved relatively slow expansions (13–26 minutes), and the visually estimated degrees of expansion would be 4 or 5 rather than 9 or 10, but the weight gained by each snail closely matched the water removed from the dish. Only in deeper water does full expansion occur, and only if nearly completely submerged is the mantle-cavity filled and functional. Other experimental investigations of water uptake using dyes such as phenol red were superseded by the experiments with inulin-labelled sea water reported in the next paper (Russell-Hunter and Apley, 1968).

Both in the laboratory and under field conditions, the larger naticids remain continuously expanded for many days (even weeks). The sea water expelled on the first contraction of freshly collected specimens has contained unidentified ciliate protozoans on eight occasions. From two large specimens of *Polinices duplicatus*, first discharged water samples yielded 7 and 3 living harpacticoid copepods, respectively. These were identified as *Tisbe gracilis* by Dr. Harry C. Yeatman, to whom our thanks are due. Harpacticoids have been expelled on five further occasions and have included all ages from nauplii to females with egg-sacs.

The last results to be discussed here are based on the data which were

accumulated on the actual shell weights and contained volumes for *Polinices duplicatus*. The weight of the dry shell plus operculum (W) in 69 individuals of shell length (L) from 6.2 mm. to 60.3 mm. (mean = 27.6 mm.), ranged from 0.057 g to 38.7 g. with a mean of 7.09 g. Ratios of W/L^3 calculated for the 69 snails (W in g., L in cm.) give a mean value of 0.166. The total volume (V) of the filled shell (by displacement) in 61 individuals of shell length from 6.2 mm. to 56.8 mm. (mean = 27.3 mm.), ranged from 0.074 ml. to 58.8 ml. with a mean of 9.96 ml. Ratios of V/L^3 calculated for the 61 snails give a mean value of 0.253. These two sets of figures include smaller snails, while the habituated experimental animals are all medium or large (the 48 snails of Tables I and II have shell lengths ranging from 16.5 to 61.0 mm., with a mean of 41.3 mm.).

It is perhaps worth using such volume figures to dismiss another erroneous theory of naticid expansion. This theory, discussed by Morris (1950) and Brown and Turner (1962), suggests that naticid pedal expansion could result from compensatory water being taken to spaces "around the animal, within the shell." Of course, Brown (1964) has shown that pedal expansion in the nassariid, *Bullia*, is accomplished entirely by movement of blood into foot, sea water flowing in to fill a "free space" between the visceral tissues and the shell. In an expanded *Bullia*, the volume of sea water contained within the shell, but outside the tissues, has been estimated at about twice the volume of water in the mantle-cavity, or up to 80% of the shell's volume (Brown, 1964). Nothing like this is possible in a large naticid. For example, experimental snail d209 (Table I) with a shell length of 48.1 mm. would have a calculated total shell volume of 28.2 ml. This is compatible with the recorded contracted weight of 35.6 g., and implies a (retracted) capacity of about 20 ml. for tissues, blood, and water. This capacity should be contrasted with the volume difference recorded (Table I) between the expanded and contracted states of 86.3 ml. Thus, application of a "free space" theory to the facts of naticid expansion is absurd.

The shell volume and weight figures can be used in "balance sheet" equations.

If T is the mass in g. of the contained tissues and retained water in the contracted snail, then: $T = M_C - W$.

Also, if the tissues have a s.g. near 1.0, T can be regarded as an approximately equivalent volume in ml. and, if V_s is the volume of shell material in ml., then: $T = V - V_s$.

Re-arranging these we have: $M_C = W + V - V_s$. But the s.g. of the shell is approximately 2.57 (calcite s.g. = 2.71), so that: $V_s = W/2.57$ and: $M_C = W + V - (W/2.57)$. Values assessed above of $W = 0.166L^3$ and $V = 0.253L^3$, allow us to predict the contracted weight in terms of the shell length: $M_C = 0.354L^3$. The actual contracted weights of the 91 specimens of *Polinices duplicatus* which provided the data for Figure 1 are fitted very well by this curve. Predicted values can be extended to expanded snails since the expansion index: $I = M_E/M_C \times 100$, and thus: $M_E = I \times 0.354L^3/100$ and, for example, for estimated expansions of 8 with an $I = 270$, then: $M_E = 0.956L^3$. Predicted curves based on shell length for each level of expansion can be made in this way and provide good fits for the 188 actual expanded weights on which Figure 1 is based. There is obviously an element of syllogism in such predictions of M_E , but the above predicted curve of M_C does *not* involve the actual data on contracted weights. Further, the broad

agreement of both sides in such "balance sheets" shows that no major weight factor has been omitted in this consideration of the relevance of weighing experiments to the mechanics of expansion in *Polinices*.

DISCUSSION

The historical introduction to this paper discusses how the mechanism of pedal expansion in naticid snails, correctly explained by Schiemenz (1884, 1887), was ignored or denied in the literature for eighty years. It is hoped that the present papers will provide the needed unequivocal evidence of the functioning of the pedal water-sinus system in naticids. The simple weighing experiments have already been discussed in the perspective of the differing water uptake involved in different degrees of expansion, in relation to the times involved in contraction and expansion, and in the relation to the weight and volume values for different parts of the snail. The essential equivalence of the weight loss on contraction to the sea water expelled, and the fit of actual contracted and expanded weights to values calculated in relation to shell length, are stressed. The relation of various water spaces, and the rates of water exchange are discussed elsewhere (Russell-Hunter and Apley, 1968).

There remain to be discussed certain mechanical and evolutionary aspects of the naticid water-sinus system.

The majority of movements in molluscs involve transmission of forces generated by muscle contractions through the hydraulic skeleton provided by the blood in the hemal meshwork of the molluscan hemocoel. In a large number of cases, molluscan organs are dilated and extended rather slowly by influx of blood, and are withdrawn relatively fast by intrinsic retractor muscles. The underlying anatomical pattern of obvious retractors within each structure without obvious antagonists locally placed is characteristic (Russell-Hunter, 1968). This reliance on distant antagonists, along with the unchanging total blood volume in the hemocoelic hydraulic skeleton, together are responsible for many of the peculiar features in the mechanics of molluscan locomotion and other movements. For example, limitations arise in the number of extensile structures which can be dilated and protruded at one time. If a pulmonate snail is observed in copulation, the sensory and locomotory organs of the head and foot are flaccid and crumpled, a large proportion of the blood volume being involved in dilation of the genital structures. Further, in most gastropods and bivalves the total blood volume is limited to that which can be withdrawn along with the tissues into the closed shell. To a limited extent, these difficulties can be bypassed by compartmentalization of the hemocoel. Certain mechanically efficient molluscan organs where this is the case have been investigated. Capacity to seal off a potentially variable amount of blood in an organ allows the use of intrinsic muscles in local functional antagonism. Examples include the use of radial muscles to extend the siphons (by "thinning" the siphonal walls) in tellinid clams (Chapman and Newell, 1956), and the radial arrangement of antagonists to the main propulsive circular muscles in the mantle wall of cephalopods. In both cases the fluid content of the wall can be isolated from the rest of the molluscan hemocoel and this allows the efficient use of local antagonistic muscles.

The adaptive significance of the great capacity for pedal expansion in naticids is obvious, and has already been discussed in relation to both locomotion and predation. Mechanically it is based on the capacity for water uptake into internal spaces to provide a hydraulic skeleton of variable total volume. Significantly the volume of the hydraulic skeleton of the pedal water-sinus system is not limited (like hemal volume) by the capacity of the closed shell. Other molluscs, notably bivalves with fused mantle edges and massive siphons, make use of contained sea water in a hydrostatic skeleton of variable volume. In forms like *Mya* and *Hiatella*, closure of the mantle openings allows the siphonal musculature to act as the antagonist of the shell adductor muscles around a temporarily constant volume of sea water in the closed mantle-cavity (Russell Hunter, 1949; Chapman and Newell, 1956; Russell Hunter and Grant, 1962). This is the basis of the boring mechanism in *Hiatella* (Russell Hunter, 1949), and of the step-wise process of siphonal extension in *Mya* (Chapman and Newell, 1956). Of course, the potential of the water-sinus system in naticids, where the total capacity of the hydraulic skeleton can be three or four times the capacity of the shell, is much greater than that of the myacean bivalve's mantle-cavity.

Compartmentalization is also important in the naticid hydraulic skeleton. During expansion, the muscular "pumping" which transfers sea water from the mesopodial water-sinuses to those of the propodium can be followed by a period of further uptake through the mesopodial pores. During this process the already dilated and turgid propodium must be sealed off from the flaccid "absorbing" mesopodium. Occasional inefficiencies in unhealthy or tired snails can be revealing. Normally the first stage of retraction is a rapid transfer of most of the propodial water to the mesopodium before columellar contraction. Occasionally, the "sphincter" muscles between propodium and mesopodium remain contracted (as they are phasically during expansion) when the columellar muscle has already begun to retract the head-foot. Rather rarely, a turgid propodium is trapped against the shell edge and injury results. More frequently, metapodial folds are trapped between operculum and shell. In these cases, the initial retraction is interrupted, the columellar muscle relaxes temporarily, the operculum gapes a little, the metapodium is withdrawn properly, and the retraction is completed. Another, more frequent, inefficiency occurs during expansion when the propodium is being filled by contraction of the mesopodial muscles. Normally, all the mesopodial pores are first sealed but, occasionally, some parts of the mesopodial margin remain relaxed and outward jets of water can be detected with dyes or suspended particles during the filling of the propodium. Apart from the overall capacity for propodial isolation, parts of the anterior water-sinus system must themselves be operable as small "closed" hydraulic units with local sets of antagonistic muscles. The evidence for this comes from behavioral, rather than mechanical, observations. Part of the propodium in *Polinices duplicatus* can wrap around and pick up a small bivalve, then transfer it across or back to another temporary propodial fold, rotate the clam, and reverse it end for end. The propodium can then manipulate it into the "correct" position for the proboscis, and form the two folds of the "vice" which holds the clam during boring. The manipulations involved in catching an active medium-sized *Ensis* can be even more elaborate. It should be remembered that the principal sensory organ in the functionally blind naticids is also the propodium.

(The greatly reduced eyes are "buried" behind the propodial shield in the expanded snail.) Observations of a *Polinices* "searching" for prey with the extended propodial tip "scanning" the sand surface like an elephant's trunk, or of the capture and handling of an *Ensis*, emphasize the adaptational significance of the disproportionate pedal expansion. There is even a report of "swimming" using undulating movements of the propodium in *P. josephinus*. Most of the activities of the larger naticids depend on the mechanical properties of the pedal water-sinus system.

As with many biological problems, it is easier to discuss the adaptational significance of the pedal water-sinus system, than to speculate on its evolutionary origins. The latter is made additionally difficult by the phylogenetically isolated position of the Naticidae within the taenioglossan mesogastropods. They represent a line which has evolved the structures and habits of predaceous carnivores (like those of stenoglossans) completely independently. The suggestion by Cox (1960) that naticids could be allied to that aberrant group of archeogastropods, the Neritacea, is not supported by the evidence of functional anatomy (Fretter and Graham, 1962) but does serve to underline the isolated position of the moon-snails. Thus it is almost impossible to set up alleged "relatives" of naticids in which to look for simple versions or precursors of the pedal water-sinus system. Obviously it is worth investigating the elaborate system of the larger naticids as it develops in the young snail, and it may be worth re-investigating the smaller naticids. Some of the latter may prove to possess a simpler, and more than proportionately smaller, water-sinus system.

In spite of the paucity of evidence on evolutionary origins, two hypotheses seem worth advancing. First, in spite of the histological nature of the sinus linings noted above, it seems most likely that the pedal water-sinus system is developed centripetally from an epidermal origin (*i.e.*, is analogous in ontology to nephridia rather than to coelomoducts). Secondly, it seems possible that the system originated as an elaboration of the lumina of anterior or posterior pedal mucous glands like those found in certain mesogastropods, which glands are developed as epidermal invaginations.

We are greatly indebted to Jay Shiro Tashiro and Dr. Martyn L. Apley for checking most of the numerical data and otherwise assisting in the preparation of this paper. We are also glad to record our gratitude to Dr. Clark P. Read who provided the opportunity to begin this work at Woods Hole.

SUMMARY

1. In 1884, Schiemenz published an essentially correct interpretation of the mechanism of pedal expansion in naticid snails. A historical review discusses how this was ignored or denied in the literature for eighty years, and summarizes the need for unequivocal evidence on the problem.

2. Larger naticids, including *Polinices duplicatus* and *Lunatia heros*, have an extensive pedal water-sinus system, and considerable intake of sea water is required for expansion. On dry surfaces, in air, they cannot expand.

3. Naticids can be habituated to handling, and then weighed in air at all degrees of expansion. Extensive weighing experiments show that the sea

water expelled on contraction is always equivalent to the weight difference between the expanded and contracted states. An index of expansion can be calculated relating expanded to contracted weight and this correlates with visual assessments of degree of expansion. Taking contracted weight as 100, half-expanded specimens of *Polinices duplicatus* have index values around 195 and fully expanded ones around 350. For example, this means that a *Polinices* weighing 46 g. contracted, takes in 124 ml. of sea water when it expands fully to a weight of 170 g. Shell weights and capacities can be incorporated in computing predicted curves which fit results from weighing live snails.

4. In healthy medium-sized specimens of *Polinices duplicatus*, retraction takes 2.5–4.0 seconds and expansion takes 3–8 minutes. Retraction is brought about by a sequence of muscle contractions; expansion is largely based on a “recoil” elasticity augmented by local hemal dilation. The larger naticids often remain continuously expanded for many days.

5. The capacity for disproportionate pedal expansion conferred by the water-sinus system is of great adaptive significance to naticid snails—particularly for locomotion in sand and for prey capture. This is discussed in relation to the hemal hydraulic skeleton and the systems of distant antagonists more usual in molluscs. The hypothesis is advanced that the pedal water-sinus system of naticids may have evolved from epidermal invaginations resembling the pedal mucous glands of other mesogastropods.

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PEDAL EXPANSION IN THE NATICID SNAILS. II. LABELLING EXPERIMENTS USING INULIN¹

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The mechanism involved in the hypertrophic dilation of the foot in naticid snails was first correctly explained by Schiemenz (1884, 1887) as being based upon a system of pedal water-sinuses. The peculiar sea-water uptake involved in naticids was then ignored or completely denied by many writers on molluscan mechanics (Lankester, 1883, 1884, 1891; Simroth, 1896–1907) and this continued until relatively recently (Brown, 1964; Morton, 1964; Hyman, 1967). A review of the history of a nineteenth century theory which erroneously attributed distension and protrusion of all kinds of molluscan organs to direct uptake of sea water is provided by Carrière (1882). Russell-Hunter and Russell-Hunter (1968) give a historical survey of published references to Schiemenz's essentially correct conclusions on naticid expansion, review the reasons which contributed to the denial of his work over eighty years, and summarize the present need (in 1968) for an unequivocal demonstration of the use of a pedal water-sinus system in the expansion of naticids.

The work recorded here and in the preceding paper on the naticid water-sinus system was adumbrated in a brief report (Russell Hunter and Apley, 1965) on temporary hyperthermia in *Polinices duplicatus*. Independently, Bernard (1968) and Russell-Hunter and Russell-Hunter (1968) have now reported weighing experiments which demonstrate the nature and extent of the water uptake involved in pedal expansion in naticids. This second paper describes and discusses the use of inulin-labelled sea water in *Polinices duplicatus* in an investigation of the relation of the different water spaces and the rates of water exchange in expanded snails carried out in summer, 1965. A third paper is being prepared on the micro-anatomy and histology of the naticid pedal water-sinus system.

MATERIALS AND METHODS

Brief notes on the systematics and ecology of naticids are given in the preceding paper (Russell-Hunter and Russell-Hunter, 1968). Only one naticid species was used in the inulin-loading experiments, the more euryoecic and more readily "trainable" form, *Polinices duplicatus*. Medium-sized specimens were employed, all collected by hand from a muddy sand flat at Orleans Town Pond, Cape Cod. For each experimental snail, shell size, contracted weight and mean expanded

¹ Supported by a research grant, GM 11693, from the National Institutes of Health.

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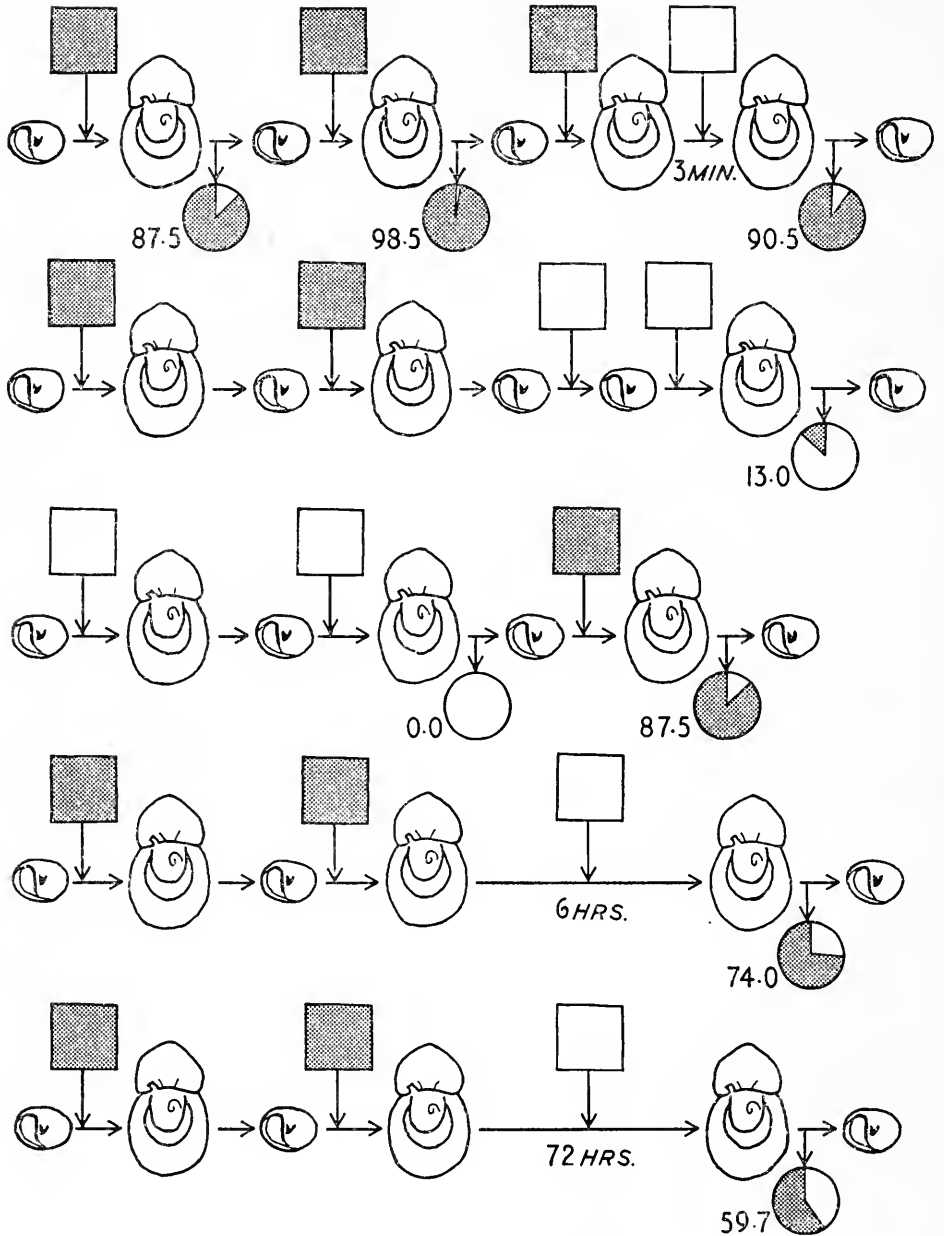


FIGURE 1.

FIGURE 1. Experimental sequences in inulin-labelling of the pedal water-sinus system in the naticid, *Polinices duplicatus*, with average values for some results. The squares above each sequence represent the sea water available for uptake on expansion—shaded squares inulin-labelled, and open squares "clean" sea water; contracted and fully-expanded snails are stylized; and the circles below represent the inulin recovered on forced contraction, expressed as a

weight after training were obtained as in Russell-Hunter and Russell-Hunter (1968). The mean expansion index for each individual allowed a "standard" water uptake to be determined for each, and the actual uptake during an experiment was usually expressed as a percentage of this.

Several sequences were used in inulin-loading, but a typical one would run as follows. A contracted specimen of *Polinices* was placed in a solution of inulin (say 50 mg./l.) in filtered sea water, allowed to expand, forced to contract, allowed to re-expand and then transferred while fully expanded through several washes of clean filtered sea water. It would then be left for a definite time in a known volume of clean sea water, and at the end of the time, weighed in air and forced to contract. Inulin determinations would then be made on the starting solution, the final external sea water, and the internal water expelled at the final forced contraction. This and other sequences are illustrated in Figure 1.

Determinations of inulin concentrations were made by photometric measurement of the red color produced by the reaction of an inulin hydrolysate with resorcinol. Except for minor changes in the preferred range and quantities of samples, this is the method of Schreiner (1950), which in itself is a simplified version of that of Roe *et al.* (1949). Such determinations using resorcinol have been employed by several workers using inulin to determine volumes of extracellular spaces (Ross and Mokotoff, 1951; Cotlove, 1954), and interference from substances likely to be present in biological salines or sea water is minimal. The more cumbersome method using the diphenylamine reaction, employed by some investigators for extracellular volumes (Gaudino and Levitt, 1949), was unsuitable for our purposes.

The analytic procedure (based on Schreiner, 1950) was as follows. Aliquots of 5 ml. of unknown samples or standards were pipetted into 75-ml. Pyrex boiling tubes, each of which already contained 12.5 ml. of 30% hydrochloric acid. Then 5 ml. of an 0.1% solution of resorcinol in 95% alcohol was added to each, and the tubes, loosely stoppered with conical reflux caps, placed in a water bath at 80° C. for 25 minutes. The samples were then cooled in tap water, and the red color determined within 40 minutes. (Trials showed that the absorption values were not significantly changed after 3 hours.) Extinction values were read against a sea-water reagent blank on a Beckman DU spectrophotometer with 1 cm.-path-length cells, mercury lamp and a wave-length setting of 490 m μ . Standards were run with every experiment, as were additional controls (such as samples expelled from the pedal water-sinuses of untreated snails). E-values show a linear relationship to inulin concentrations between 5 mg./l. and 55 mg./l., and experimental procedures were adjusted so that most determinations fell in that range. A stock 0.2% solution of inulin in filtered sea water was prepared at 60° C. (2 g./l.). This was diluted with filtered sea water, immediately before use, to give "exposure"

percentage of the "loading" concentration. These percentages are also shown as figures to the left of each circle and, in three cases, the average elapsed time while expanded in "clean" sea water is also shown. The first sequence is typical of experiments used in investigation of the effects of "multiple loading" and of "flushing" out of the mantle-cavity, the second and third show two methods of investigation of the residual volume of the water-sinus system in contracted snails, and the fourth and fifth typify experiments on the retention of inulin in the water-sinus system over longer elapsed times while expanded. For further explanation, see text and Tables I-V.

solutions in most cases of approximately 50–60 mg. inulin/l. Such high starting concentrations in the experiments with *Polinices* allowed much greater accuracy in the later determinations of inulin in wash waters and in samples from diluted pedal water. Most results are expressed both as concentrations of inulin in mg./l. and as percentages of the (subsequently determined) starting concentration.

RESULTS

In a group of preliminary experiments, numbered and trained specimens of *Polinices duplicatus* (with contracted weights ranging from 17.3 g. to 26.8 g.) were transferred, while contracted, into inulin solutions (46 mg./l. or 97 mg./l.) where they were allowed to expand fully. Thus the water taken into the pedal water-sinuses was inulin-labelled. They were then gently transferred, without much contraction, through successive washes in large volumes of "clean" sea water for periods of from 17 to 34 minutes. After weighing, damp-dried, in air (expanded weights 42.8–67.0 g.), forced contraction yielded water samples (23.5 to 40.6 ml.)

TABLE I

Inulin concentrations in the pedal water-sinuses of Polinices duplicatus immediately after loading (i. e., less than 1 second washing before force contraction and sampling)

Experiment series #	Snail #	Contracted weight (g.)	Inulin concentration mg./l.		Number of loading cycles	Final concentration as percentage of initial uptake
			Initial uptake	Final sample		
III	R2	26.4	60.0	58.3	3L	97.2
VC	RR8	25.6	63.3	63.0	2L	99.5
VC	PP2	23.8	63.3	61.8	2L	97.6
VC	PP4	29.0	63.3	62.7	2L	99.1
VC	PP9	20.3	63.3	62.8	2L	99.2

from the pedal water-sinuses. These had inulin concentrations ranging from 42.4% to 52.6% of the appropriate starting levels.

This reduction to about half of the starting concentration of inulin in the water recovered from the snails on contraction, obviously could involve several diluting factors. These include: first, residual water present in the contracted water-sinuses before exposure to inulin; secondly, the water contained in the mantle-cavity and rapidly exchanged by the ciliary respiratory currents during the washes in clean sea water; and thirdly, water exchanged between washing baths and the water-sinuses because of small partial contractions and re-expansions during the transfer process. Many of the later experiments were designed to distinguish among these three factors and, as might be expected, the first two proved to be relatively constant for each individual *Polinices*, while the third varied greatly (though contributing least dilution to the results in the best habituated snails).

Before turning to these more complex experiments and to those extending over longer periods of time, it is worth noting that even this preliminary series of results from "single loadings" with inulin is important in confirming the existence of closed spaces containing sea water in the expanded naticid. Recent weighing

TABLE II

Multiple inulin loading of the pedal water-sinuses in Polinices duplicatus, followed by thorough flushing of the mantle-cavity

Experiment series #	Snail #	Contracted weight (g.)	Inulin concentration mg./l.		Number of loading cycles	Final concentration as percentage of initial uptake	Maximum mantle-cavity volume as percentage of water-spaces
			Initial uptake	Final sample			
IIA	R1	23.9	58.3	53.1	4L	91.1	8.9
IIA	R5	16.4	58.3	53.8	4L	92.3	7.7
IIA	P3	26.3	58.3	53.7	4L	92.1	7.9
IIIC	R5	16.4	58.3	52.0	4L	89.2	10.8
III	R1	23.9	60.0	57.1	4L	95.2	4.8

experiments on naticids (Bernard, 1968; Russell-Hunter and Russell-Hunter, 1968) merely belatedly confirm and extend Schiemenz's (1884, 1887) findings. The demonstration that a large quantity of inulin-labelled sea water can be carried in the expanded naticid through "clean" wash waters and even through air in a "damp-dried" snail, is a different *kind* of unequivocal evidence for the functioning of the water-sinus system.

In the majority of the 108 successful inulin trials run, the question of the residual volume of the pedal water-sinuses in the contracted snail could be bypassed by multiple loading. In other words the snail was allowed to expand in inulin-labelled sea water, force contracted, then allowed to expand again in the same solution and so on. Initially, three and four successive loadings were carried

TABLE III

Residual volume of the pedal water-sinuses in contracted Polinices duplicatus by two methods

Experiment series #	Snail #	Contracted weight (g.)	Inulin concentration mg./l.		Number of loading cycles	Final concentration as percentage of initial uptake	Residual volume as percentage expanded water-sinus volume
			Initial uptake	Final sample			
VIIC	RR6	36.1	66.8	9.8	2L + 1SW	14.7	14.7*
VIIC	RR1	42.1	66.8	8.8	2L + 1SW	13.2	13.2*
VIIC	PP7	24.8	66.8	8.3	2L + 1SW	12.4	12.4*
VIIC	PP6	24.8	66.8	7.4	2L + 1SW	11.1	11.1*
VIIC	PP2	23.8	66.8	9.0	2L + 1SW	13.5	13.5*
VIIA	PP8	22.5	66.8	59.4	1L	88.9	11.1†
VIIA	PP5	23.5	66.8	57.7	1L	86.4	13.6†
VIIA	PP3	19.5	66.8	59.1	1L	88.5	11.5†
VIIA	PP2	23.8	66.8	57.6	1L	86.2	13.8†

* The first group of five snails were loaded twice with inulin-labelled sea water, force contracted, washed and allowed to re-expand in "clean" sea water before force contracting for the final sample.

† The second group of four snails were taken from clean sea water, force contracted, loaded once with inulin-labelled sea water and rapidly washed in "clean" sea water before force contracting for the final sample.

TABLE IV
Inulin retention in the pedal water-sinuses of Polinices duplicatus after multiple loading and elapsed times from 5 to 20 hours

Experiment series #	Snail #	Contracted weight (g.)	Inulin concentration mg./l.		Number of loading cycles	Elapsed time (hrs. mins.)	Final concentration as percentage of initial uptake
			Initial uptake	Final sample			
VIII	RR1	42.1	61.8	39.8	2L	5.15	64.4
VIII	RR5	35.8	61.8	49.3	2L	5.30	79.8
VIII	RR6	36.1	61.8	50.1	2L	5.31	81.1
VIII	PP8	22.5	61.8	41.5	2L	6.03	67.2
VIII	PP1	28.5	61.8	48.6	2L	6.33	78.6
VB	RR4	36.1	59.6	43.3	3L	12.02	72.7
VB	PP7	24.8	59.6	39.3	3L	12.06	65.9
VB	RR2	41.2	59.6	38.2	2L	12.17	64.1
VA	RR3	30.3	59.9	36.9	3L	20.57	61.6
VA	RR9	24.7	59.9	37.6	4L	20.58	62.8
VA	RR5	35.8	59.9	40.6	4L	21.12	67.8
VA	RR7	33.8	61.1	42.0	4L	21.23	68.7

out, but it became clear that two successive loadings would suffice to label the residual volume (at least within the limits of accuracy imposed by other variables). In Table I are presented five typical results of sampling immediately after multiple loading with only a brief (less than 1 second) wash to remove surface contamination. As can be seen the water expelled from the pedal water-sinuses in these cases has inulin concentrations between 97.2 and 99.5% of the initial uptake. Thus superficial water on the shell and expanded surfaces can only amount to to 0.5–2.8% of the increase in weight of an expanded *Polinices*. In the experiments of Table II, multiple loading was followed by washing, extending over several minutes, in large volumes of sea water. The five cases presented are typical of snails where no contraction or major pedal movement took place during

TABLE V
Inulin retention in the pedal water-sinuses of Polinices duplicatus after longer elapsed times (2–3 days)

Experiment series #	Snail #	Contracted weight (g.)	Inulin concentration mg./l.		Number of loading cycles	Elapsed time (hrs. mins.)	Final concentration as percentage of initial uptake
			Initial uptake	Final sample			
VIB	RR9	24.7	63.1	48.3	2L	58.13	76.5
VIB	PP9	20.3	63.1	40.8	2L	58.11	64.7
VIB	RR8	25.6	63.1	31.2	2L	58.35	49.4
VIB	RR3	30.3	63.1	49.8	2L	58.32	78.9
VIA	PP4	29.0	66.8	33.8	2L	72.15	50.6
VIA	RR7	33.8	66.8	32.8	2L	72.19	49.1
VIA	RR4	36.1	66.8	45.7	2L	72.36	68.5
VIA	RR2	41.2	66.8	47.2	2L	72.34	70.7

the manipulation, and therefore the final sample concentrations reflect dilution by the proportion of water in the mantle-cavity which has had time to become thoroughly flushed out by the ciliary currents. Therefore, the maximum percentage of all water spaces in the expanded snail made up by the pallial cavity lies between 4.8 and 10.8%. Other circumstantial evidence, and an average value of 2% for superficial water, would suggest that for medium-sized *Polinices duplicatus*, mantle-cavity water usually makes from 5 to 7% of the volume taken in on expansion.

The factor of the residual pedal water-sinus volume (when the snail is contracted) is assessed in two ways in Table III. (See also the procedural sequences in Figure 1.) The first group of results in Table III is derived from snails loaded twice with inulin, force contracted and washed while withdrawn, and then allowed to re-expand in "clean" sea water. Thereafter, force contraction yields a water sample from the pedal water-sinuses with an inulin concentration proportional to the residual volume of the water-sinuses when the snail was contracted. The residual volumes thus represent 11.2–14.7% of the volume of the water-sinuses in the fully expanded snail. Another approach was to load snails once only, rapidly wash and then force contract for a water sample. In this case the dilution of the final inulin concentration is proportional to the residual space. Final concentrations of 86.2–88.9% imply residual volumes of 11.1–13.8% by this method—a satisfactory concordance.

As noted above, the majority of experiments began with multiple loading so the potential dilution from the residual volume of the contracted water-sinus (mean value 12.8%) can be ignored. However, all the other experiments on inulin retention in the water-sinuses after various elapsed times can be regarded as showing a basic dilution of up to about 10% resulting from the combined pallial and superficial volumes of water. Superimposed on this are the major differences in water exchange resulting from individual variations in sensitivity and activity which cause different degrees of pedal retraction within the period of the experiment.

Table IV presents some data on inulin retention in snails 6, 12 or 20 hours after transfer to "clean" sea water. Of the water within the pedal water-sinuses, 64.4–81.1% can remain unexchanged for 5–6 hours, and 61.6–68.7% for 21 hours. It should be realized that this set of results represents a biased group, because any snail which has responded to any stimulus by retraction during the 21 hours has been automatically removed from the experimental series.

This bias is even more true of the results on inulin retention over 2–3 days which are presented in Table V. The data were gained from our best habituated snails, and considerable precautions had to be taken to maintain the animals in conditions free from major changes in light, temperature, tactile and vibrational stimuli during the 58 and 72 hours of the experiment. It is remarkable that 49.1–70.7% of the pedal water could remain unexchanged after 72 hours. Other circumstantial evidence, from commensals (Russell-Hunter and Russell-Hunter, 1968), from the occurrence of temporary hyperthermia (Russell Hunter and Apley, 1965), and from other behavioral data, confirms the potentially static nature of the water after it has been taken up into the pedal water-sinuses. It seems likely that, in nature, a large naticid can remain continuously expanded for periods of weeks.

DISCUSSION

Weighing experiments have been used to demonstrate the nature and extent of sea water uptake into the pedal water-sinus system during the expansion of naticids. The experiments with inulin-labelled sea water reported in the present paper provide additional evidence, particularly as regards the relation of the various water spaces and as regards the rates of water exchange. Both topics merit further discussion.

The experiments showed that there are three components in the volume of sea water responsible for the weight increase in expanded snails. Only about 2% is superficial water on the snail's shell and expanded surfaces, and the water rapidly circulating through the mantle-cavity amounts to a further 5-7% of the total. Approximately 90% of the sea-water uptake on expansion enters the pedal water-sinus system. The earlier conclusion that this system is completely separated from the blood in the hemocoelic spaces (Schiemenz, 1887; Russell-Hunter and Russell-Hunter, 1968), is confirmed by the figures for recovery of inulin in many experiments. In other words, there is no dilution of the sea water in the water-sinus system that is not accounted for, and none that could involve exchange with other body fluids. A residual volume of sea water is retained in the contracted water-sinus system and, immediately after contraction, this amounts to 12.8% of the volume of the water-sinuses in the expanded snail. Once again, the evidence from the inulin-loading experiments shows that there is no exchange between the sea water of this residual space and the blood at any time. Similarly, while the snail remains contracted there is no exchange between the residual space and the environmental sea water outside the animal.

Exchange between the environment and the water-sinuses of a fully expanded snail takes place only to the extent that partial contraction followed by re-expansion occurs. (Over a considerable number of "less successful" experiments not fully reported above, there was good correlation between the degree of dilution of the inulin load and the observed frequency and extent of partial contractions.) The data from inulin-labelling prove unequivocally that the pores of the mesopodium remain continuously closed in fully expanded specimens of *Polinices duplicatus*. As noted in the discussion of the preceding paper (Russell-Hunter and Russell-Hunter, 1968), the mechanical value of the pedal sea water to the snail depends on it being closed off from the environment as a hydraulic skeleton (temporarily of constant volume) capable of transmitting forces between antagonistic muscles. It is still remarkable that inulin-labelling showed that 49-71% of the pedal sea water could remain unexchanged after 72 hours.

At first sight, it would seem that surprisingly "stagnant" nature of the water in the pedal water-sinuses would present certain physiological problems to the snail. This does not seem to be so, though questions such as long-term osmotic stress have not yet been investigated. More can be said on the problems of internal temperature and oxygenation. The large static water content, sealed off from the environment, can result in a condition of temporary hyperthermia in *Polinices* (Russell Hunter and Apley, 1965)—a most unusual condition in a marine "poikilothermic" animal. This was detected using thermistor probes in field and laboratory. Snails crawling on exposed tidal flats on a summer's day had internal temperatures ranging from 2.0° C. to above 0.3° C. below the tempera-

ture of the upper 1 cm. of the sand. The temperature of incoming tidal water could be as much as 6.5° C. below the temperature of the sand-flats, and the snails could remain hyperthermic for an appreciable time after tidal submergence. Subjected to lowered temperatures of this order, larger snails (fully expanded weights of 102–159 g.) remained hyperthermic to the extent of 1.8° C. after 5 minutes, 0.9° C. after 10 minutes, and 0.1° C. after 45–60 minutes. Russell Hunter and Apley (1965) note that, empirically, heat is retained about as well as in an equivalent volume of confined but circulating water, but markedly less than in equidimensional foam latex rubber. Such comparison implies some exchange of water *within* the animal during cooling but no exchange with the outside. This is completely in accord with the other evidence on the use of the contained sea water as a hydraulic skeleton. Such a temporary state of hyperthermia has little metabolic importance but is probably of considerable behavioral significance since littoral moon-snails become active immediately after their inundation by the rising tide.

The other physiological question concerns the oxygenation of this "stagnant" mass of internal sea water. There is no evidence to suggest that the water has a lower oxygen tension than that of the snail's blood in the adjacent sinuses. There is circumstantial evidence to the contrary provided by the protistan and harpacticoid commensals which have been recovered from expelled water (Russell-Hunter and Russell-Hunter, 1968). Further, even the most isolated tissues of animals fully expanded for many days remain healthy and responsive. In other words, the ensheathing pavement epithelium and collagen-like connective tissue layers, while mechanically tough, offer no physiological barrier to the diffusion of oxygen.

In broad terms, it seems as if very little energy is expended by a *Polinices* in remaining fully expanded, and perhaps there is a more considerable energy expenditure involved in a complete cycle of contraction and re-expansion. It would seem that the capacity for habituation, which we have exploited in these experiments, must have some adaptive significance. Further, it is clear that sustained contraction and handling over a period of minutes will produce a traumatic change in behavior in *Polinices duplicatus*. "Wild" snails which have experienced such a forced contraction re-expand within 8 minutes, but soon burrow deep into the substratum where they remain immobile but expanded for some time. This behavior was detected as introducing a source of bias when capture-recapture methods were being used to assess population density in *Polinices* (Russell Hunter and Grant, 1966). This study also produced quantitative evidence that recovery from such trauma is complete in just over two tidal cycles, or in about 25.5 hours.

Finally, it should be noted that although inulin-labelling experiments were carried out only on *Polinices duplicatus*, there is reason to believe that the results would be similar with any *large* naticid. All available information suggests that such features as (1) the large proportion (about 90%) of the sea water uptake going into the pedal water-sinuses, and (2) the potentially static nature of the sea water after it has been taken up, would be as true of *Lunatia heros*, *Polinices lewisi* and *P. josephinus* as they are of *P. duplicatus*.

We are greatly indebted to Jay Shiro Tashiro and Myra Russell-Hunter for their help with calculations, and in the preparation of this paper.

SUMMARY

1. Inulin-labelled sea water has been used in an investigation of the pedal water-sinus system, and other water spaces, in *Polinices duplicatus*. Analyses were by a photometric measurement of the reaction of inulin hydrolysate with resorcinol.

2. Of the sea water uptake during expansion: about 90% enters the pedal water-sinus system, about 5-7% is water which rapidly circulates through the mantle-cavity, and about 2% is superficial water on the snail's shell and expanded surfaces.

3. When the snail is in the contracted state a residual volume of sea water is retained in the pedal water-sinus system, and this can amount to 12.8% of the volume of the pedal system in the expanded snail.

4. There is no exchange between the water-sinus system and the blood at any time and, in the fully expanded snail, little or no exchange between the system and the environmental sea water. Labelling showed that 49-71% of the pedal sea water could remain unexchanged after 72 hours.

5. The surprisingly "stagnant" nature of the sea water in the pedal water-sinuses is discussed. Physiological consequences are probably slight, though, under certain ecological conditions, the large static water content is responsible for an unusual condition of temporary hyperthermia. A hypothesis, that little energy is expended by a *Polinices* in remaining fully expanded, is coupled with evidence of traumatic change in behavior resulting from sustained contraction. It seems likely that the features of water spaces and exchange rates demonstrated in *Polinices duplicatus* would be similar in any large naticid.

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THE IDENTIFICATION AND MEASUREMENT OF SUGARS IN THE BLOOD OF THREE SPECIES OF ATLANTIC CRABS¹

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Although considerable attention has been given to the physiology and biochemistry of the Crustacea, their carbohydrate metabolism did not become known until Hu (1958) found evidence that it was not unlike that of other animals. Using paper chromatography Hu found glucose, maltose, maltotriose, maltotetrose and two other oligosaccharides in the blood of the crab, *Hemigrapsus nudus*. Since then these same sugars have been reported in *Cancer magister* (Meenakshi and Scheer, 1961) several other crabs (Dean and Vernberg, 1965a), the crayfish, *Orconectes virilis* (McWhinnie and Saller, 1960) and the lobster, *Homarus americanus* (Telford, 1968a). The non-reducing disaccharide, trehalose, was found in nine species of crustaceans by Fairbairn (1958) and in traces in some crabs (Dean and Vernberg, 1965a). In the American lobster trehalose appears to be a significant minor component of the blood involved in the response to handling stress (Telford, 1968b). Several monosaccharides, galactose, mannose, fucose and fructose were found with sporadic occurrence (above references).

In this paper the results of chromatographic analysis of the blood sugars in the crabs *Carcinus maenas*, *Cancer borealis* and *Cancer irroratus* are related to measurements of blood glucose and total reducing sugar. Some evidence of changes with the molt cycle and reproductive activity is also given. A comparison is then made between these data and the rather scattered observations in the literature.

MATERIALS AND METHODS

Collection of animals and blood samples

Both *C. borealis* and *C. irroratus* were obtained from lobster traps at Port Clyde Maine. Three point five to 4.0 ml. of blood was taken via the articular membrane at the base of the cheliped. Decapod crustaceans become hyperglycemic following the stress of handling (Abramowitz *et al.*, 1944). To avoid this reaction blood samples were obtained at the moment of capture with the animals in as nearly an undisturbed state as possible. *Carcinus maenas* was collected in the intertidal zone at Port Clyde. Only from the largest specimens could 4.0 ml. blood be obtained; normally only about 2.0 ml. were collected. The shore crabs, *C. maenas*, were obtained in several stages of the molt cycle but the other two species were taken only in premolt and late postmolt or intermolt because they do not enter the traps at other times. Collections were made monthly from May

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through October, 1966, and September, 1967. The size range of *C. borealis* used was from 6 cm. up to about 18 cm. across the carapace. *Cancer irroratus* is smaller and the range used was 5–15 cm. The size range of *C. maenas* was 4–11 cm. In all three species males are generally larger than females.

As anticoagulant 4.0 ml. Heller's oxalate (Gradwohl, 1943) was used (2.0 ml. for the smaller blood samples). The anticoagulant was evaporated to dryness before use, thus avoiding dilution of the blood sample. The oxalated blood was stored frozen until use.

Preparation of samples

Large amounts of oxalate are necessary because coagulability varies during the molt cycle, presumably because of fluctuating blood calcium levels (Travis, 1955). Excess oxalate, however, interferes with the determination of glucose. Before deproteinizing the 1.0-ml. aliquots of blood, oxalates were precipitated by addition of 2.0 ml. 1% CaCl_2 . Proteins were precipitated by the Somogyi (1930)

TABLE I

Monthly levels of blood glucose and reducing substances (RS) in Cancer borealis (mg./100 ml.) Port Clyde, Maine, May–October, 1966

Month	♂ N	Glucose	(S.E.)	R.S.	(S.E.)	♀ N	Glucose	(S.E.)	R.S.	(S.E.)
May	48	6.8	(0.49)	10.5	(0.55)	49	8.0	(0.62)	10.0	(0.57)
June	47	7.7	(0.38)	12.2	(0.51)	43	6.6	(0.47)	10.3	(0.54)
July	50	8.0	(0.36)	10.5	(0.45)	43	7.6	(0.25)	9.4	(0.30)
August	51	10.6	(0.47)	15.3	(0.55)	52	8.4	(0.50)	11.1	(0.57)
September	53	10.2	(0.69)	12.4	(0.74)	53	10.2	(0.57)	12.8	(0.63)
October	43	9.9	(0.46)	12.9	(0.52)	52	9.5	(0.45)	12.1	(0.48)
September (1967)	58	8.7	(0.47)	13.1	(0.81)	64	8.8	(0.43)	13.0	(0.67)

method using 2.0 ml. 0.6% NaOH followed by 2.0 ml. 2.2% ZnSO_4 (acidified by drop-wise addition of conc. H_2SO_4 so that 1 vol. of zinc sulfate solution exactly neutralized 1 vol. of 0.6% NaOH). After centrifuging a clear supernatant was obtained, representing a 1:7 dilution of the blood. A 1.0-ml. aliquot of this supernatant was diluted 1:4 with distilled water.

Determination of glucose and reducing substances

Glucose was determined in 1.0-ml. duplicates of the original Somogyi supernatant using glucose oxidase-o-diansidine ("Glucostat," Worthington Biochemical Corp., Freehold, N. J.) dissolved in 0.02 M phosphate buffer, pH 7.0; further details of this procedure have appeared elsewhere (Telford, 1965). Reducing substances were determined in 1.0-ml. duplicates of the diluted Somogyi supernatant (above) using the Folin-Malmros alkaline potassium ferricyanide method (Dische, 1962). In both cases a blank of distilled water and three glucose standards of 5, 10, 20 mg./100 ml. were prepared in exactly the same way as the blood whenever determinations were made.

Paper chromatography

In preparation for chromatography individual blood samples of 1.0 or 2.0 ml. (as available) were coagulated by brief immersion in a boiling water bath (about 45 sec.). The clot was broken up and extracted with three washings of 60% methanol which were then pooled. Deionization on ion exchange columns followed (Telford, 1965) and the eluate was dried at 40° C. The residue was redissolved in 0.1 or 0.2 ml. pyridine (depending on size of original blood sample). Various sugars, alone and in mixtures, were prepared in the same way without showing any changes in chromatographic characteristics. The pyridine solutions were spotted onto Whatman #3 paper and developed for 16 hr. in a descending flow of ethyl acetate:pyridine:water, 8:2:1 (v/v/v) (Jermyn and Isherwood, 1949).

Detection of sugars on the developed chromatograms was by one of three

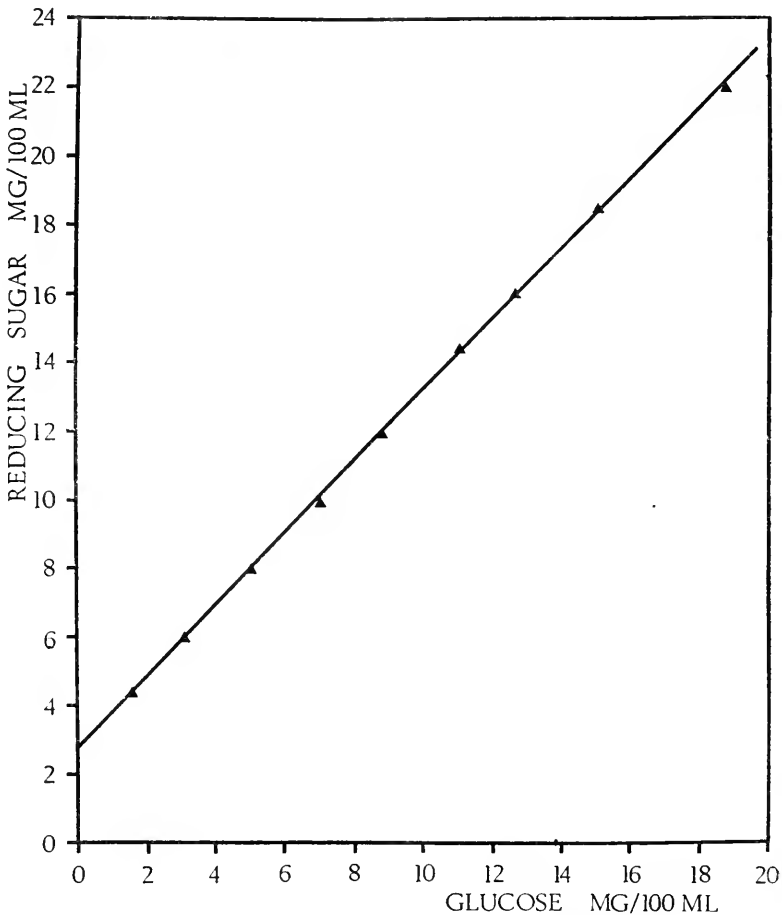


FIGURE 1. Relationship between blood glucose and reducing substances in the crab, *Cancer borealis*.

ways: (1) spray with 3% phthalic acid in 95% ethanol, viewed under UV light, (2) 0.5% benzidine in acetic acid-trichloroacetic acid-ethanol, and (3) AgNO₃ in acetone followed by 10% NaOH in 80% methanol (all methods in Dawson *et al.*, 1959).

Estimates of concentration in the spots were made by visual comparison with spots of known concentration and by chromatographing various sizes of samples to determine threshold of visibility with the detecting agents. Neither method is particularly accurate but estimates of the right order of magnitude are obtainable.

Reagents

The chemicals used were Fisher "Certified" and the carbohydrates for chromatography were obtained from Sigma Chemical Co. Ltd.

RESULTS

Mean levels of glucose and total reducing substances (RS) in the blood of *C. borealis* made at monthly intervals are given in Table I, together with the number of specimens (N) and the standard error of the mean (S.E.).

TABLE II

Blood glucose and reducing substances during the molt cycle of Carcinas maenas (mg./100 ml.)

Stage	Drach (1939)	N	Glucose	S.E.	Reducing sugar	S.E.
Intermolt	C ₄ -D ₀	23	9.9	(0.81)	13.8	(0.72)
Premolt	D ₁ -D ₄	11	12.2	(1.37)	17.1	(1.41)
Very soft	A ₁ -A ₂	5	5.8	—	7.9	—
Early postmolt	B ₁ -B ₂	2	5.7	—	8.0	—
Postmolt	C ₁ -C ₃	19	8.4	(0.90)	13.5	(1.07)

An analysis of variance indicates that the variation in mean monthly glucose levels (1966) is significant at the 0.05 level. The differences in mean monthly reducing substance levels are generally not significant although a t-test between the July and August levels in males gives a significant result. In spite of appearances the difference between the sexes is not significant.

The individual glucose and reducing substance levels varied quite widely. Glucose ranged from 0.4 mg./100 ml. to 29.3 mg./100 ml. and the range for reducing substances was 3.3-29.5 mg./100 ml. The relationship between glucose and reducing substance is approximately constant. Figure 1 is a plot of mean glucose levels within 2 mg./100 ml. classes against the corresponding mean reducing substance levels. The line has a slope of 1.02 and an intercept of 2.9 mg./100 ml. when glucose is nil. The same values for slope and intercept can be obtained directly. The correlation coefficient between glucose and reducing substances, calculated directly from the 584 paired measurements, is highly significant ($R = 0.914$, $t = 54.22$ with 582 degrees of freedom). The analysis of covariance shows that this relationship between glucose and reducing substances

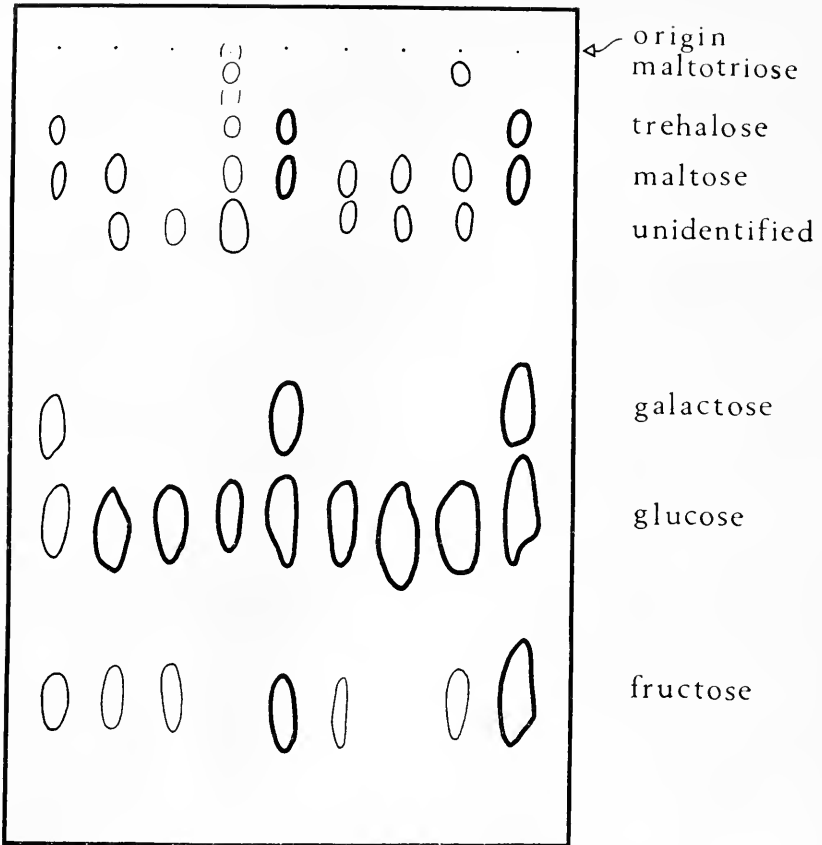


FIGURE 2. Paper chromatogram of blood sugars of *Cancer borealis*. Left, center and right, control mixtures of trehalose, maltose, galactose, glucose and fructose, 1.5 γ , 3.0 γ and 7.5 γ each, respectively (traced from photograph).

does not vary significantly from month to month, nor between the sexes, and it indicates the same level of significance for the variations in monthly mean glucose and reducing substance levels as did the analysis of variance already cited.

The crabs, *C. maenas*, were collected in September, 1967, in various molt stages. Intermolt and very early premolt, $C_1 - D_0$ of Drach (1939), were not separated but were treated together as a single intermolt stage which was recognized by color, hardness of shell, texture, color and consistency of the blood etc. Premolt, stages $D_1 - D_4$ of Drach, was recognized by darker color with blueness of chelipeds, development of new epicuticle and exocuticle and regeneration of damaged appendages. The earliest postmolt stage, stages $A_1 - A_2$ of Drach, was easily recognized by the non-calcified shells and the next postmolt stage, stages $B_1 - B_2$ of Drach, by the partially calcified exoskeleton. Postmolt, $C_1 - C_2$ of Drach, was recognized by the color and texture of the shell and blood. Table II shows blood glucose and reducing sugars in these general molt stages. These dif-

ferences are found by analysis of variance to be significant at the 0.05 level. No attempt was made to determine the significance, if any, of the differences between sexes, because there are too few observations to produce a reliable result.

A total of 67 specimens of *C. irroratus* were taken from lobster traps in May and June (1966), 31 males and 36 females. The mean glucose level for these was 8.1 mg./100 ml. (S.E. 0.23) and reducing substances 11.9 (S.E. 0.39); no difference in sex was apparent. No other determinations were made on this species (except chromatography, below) because *C. borealis* was readily available in much greater numbers in the areas where lobster traps were being set.

Chromatography of deionized blood samples (Figs. 2 and 3) revealed the occurrence of six sugars in the amounts estimated in Table III. Mobility of the sugars is expressed relative to glucose (R_g) because the solvent front was allowed to leave the paper. The bracketed percentage figure indicates the approximate number of samples in which each sugar was found. The estimated quantity is maximal in these samples. The unidentified carbohydrate was com-

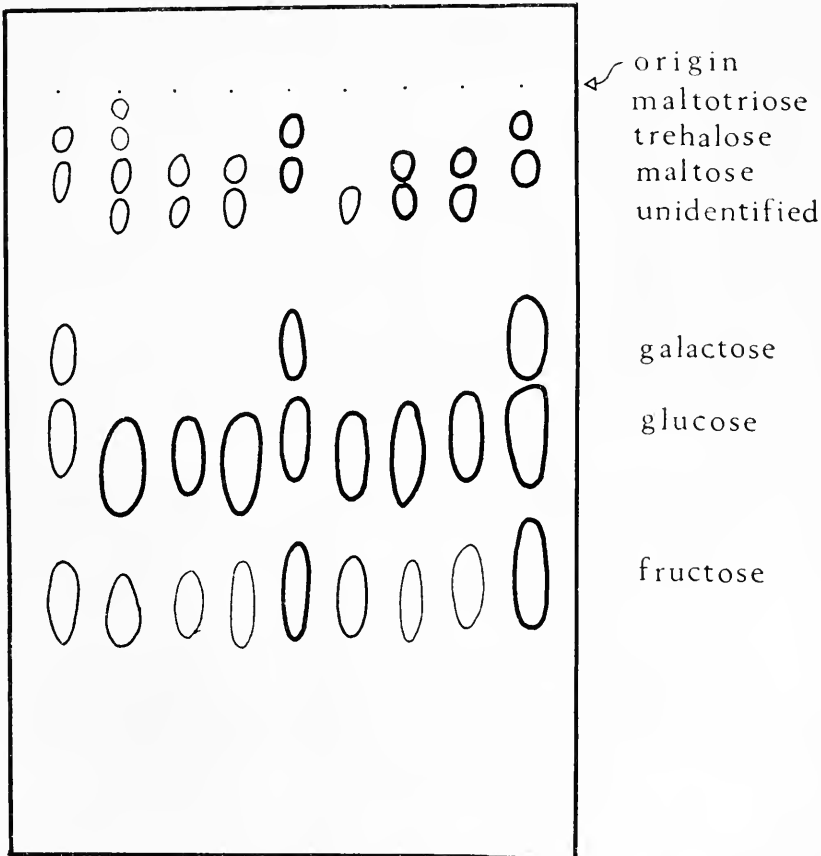


FIGURE 3. Paper chromatogram of blood sugars of *Carcinus maenas*. Controls as in Figure 2 (traced from photograph).

pared to maltose in estimating its concentration. The following criteria were used for identification of the sugars. Maltotriose: plot of $\log_{10} R_g$ against number of hexose units places it on a straight line with maltose and glucose (McWhinnie and Saller, 1960) (Fig. 4). No maltotriose was available for direct comparison. Trehalose: same mobility as control spots, reinforcement of spots when trehalose added to sample, lack of reaction with phthalic acid or benzidine indicates non-reducing nature, appearance of spot with silver nitrate at same time as trehalose (this is due to the very strong alkali used, 10% NaOH). Maltose, glucose and fructose: mobilities, lack of proliferation when these sugars were added to blood samples, corresponding blood sugars show reducing activity and appear in same sequence and time intervals with $AgNO_3$ -NaOH spray (fructose-glucose-maltose). The mobility of the unidentified component suggests a disaccharide and its reaction with both phthalic acid and benzidine indicates a reducing group.

A total of ten female *C. borealis* was obtained carrying egg masses. On one of these the sponge was old; blood glucose and reducing substances levels were

TABLE III
Estimated concentrations of blood sugars found in three species of crabs
with the frequencies of occurrence (%)

Sugar	R _g	<i>C. borealis</i>	<i>C. irroratus</i>	<i>C. maenas</i>
Maltotriose	(0.05)	trace (10%)	trace (30%)	2 mg./100 ml. (30%)
Trehalose	(0.13)	trace (10%)	trace (10%)	1.5 mg./100 ml. (10%)
Maltose	(0.27)	2 mg./100 ml. (70%+)	2 mg./100 ml. (70%)	3 mg./100 ml. (30%)
Unidentified	(0.32)	5 mg./100 ml. (100%)	5 mg./100 ml. (100%)	5 mg./100 ml. (100%)
Glucose	(1.00)	— (100%)	— (100%)	— (100%)
Fructose	(1.40)	1 mg./100 ml. (60%)	1 mg./100 ml. (30%)	1.5 mg./100 ml. (100%)

8.0 and 8.2 mg./100 ml., respectively, in this individual. The respective mean levels for the remaining nine were 12.36 (S.E. 1.18) and 14.47 (S.E. 1.19) mg./100 ml. Twelve females were found with bright orange-red blood and well developed ovaries. These were presumed to be preparing for spawning since the blood color was closely similar to that of the eggs and ovaries and this coloration never appeared in males. Mean glucose and reducing levels were 11.7 (S.E. 1.28) and 12.5 (S.E. 1.17) mg./100 ml. Three pairs of *C. maenas* were captured in copulation and blood was taken from all of the males and two females. No significant difference in blood sugar levels was found despite the ire and pugnacity of the animals (glucose 10.3, reducing substances 13.9 mg./100 ml.).

DISCUSSION

The blood glucose and reducing sugar levels reported here are comparable with those reported previously for decapod crustaceans but some interesting differences emerge. McWhinnie and Saller (1960) found in the crayfish *Orconectes virilis* that glucose levels averaged 3-4 mg./100 ml. and made up about 20-25% of the total reducing substances (by Folin-Wu method). Earlier McWhinnie and Scheer (1958) found the same relationship between glucose and total carbohydrate

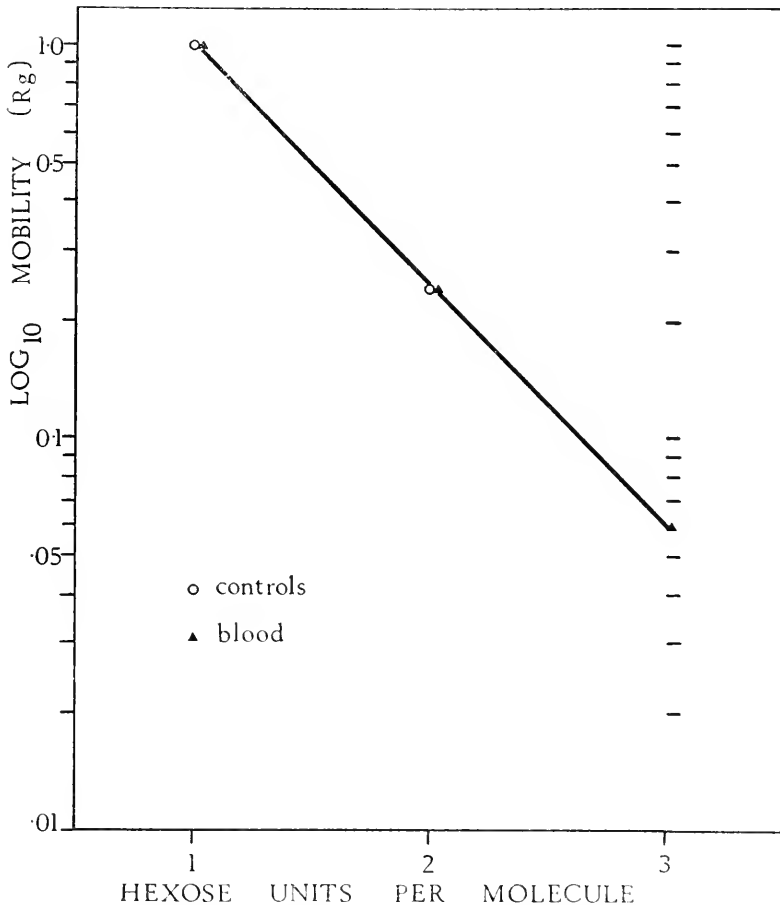


FIGURE 4. Identification of presumed maltotriose by plotting molecular size and \log_{10} mobility (after McWhinnie and Saller, 1960).

(by anthrone method) in the crab *Hemigrapsus nudus*. More recently Dean and Vernberg (1965a, b) using the same methods as McWhinnie and Saller (1960), found the same relationship in several species of crabs and stated specifically that no matter what the glucose level, its ratio to total reducing substances was constant. This is clearly not the case in *C. borealis* where glucose is the variable blood sugar and the non-glucose component remains approximately constant at about 3 mg./100 ml. In the lobster, *Homarus americanus*, this same relationship has been found, variable glucose levels and approximately constant non-glucose around 4.4 mg/100 ml. (Telford, 1968a). In the special case of females carrying eggs there is evidence of significant differences between them and other females (0.01 level by *t*-test). In Table IV these and similar data from Dean and Vernberg (1965b) are compared.

A similar series of changes in glucose levels evidently occurs in the two species but according to the data for *C. sapidus* another reducing substance also undergoes

the same changes. No attempt was made to identify this substance. Dean and Vernberg (1965b) appear to have used a heat-coagulated blood for their reducing substance determination. This method does not completely deproteinize the sample and leaves most or all of the non-carbohydrate reducing matter in the filtrate. These interfering substances may have masked the true relationship between glucose and reducing sugars in the studies with heat-coagulated blood.

The amount of color produced by several sugars with the Folin-Malmros method used in this study was determined and expressed as a percentage of the color given by the same amount of glucose. Trehalose, as expected, gave no reaction; maltose gave 60% of its glucose equivalent, fructose 104%. The estimated amounts of sugars other than glucose found in chromatograms (Table III) are the maximum levels found. Assuming that the unknown reducing disaccharide would react like maltose (lactose, for example, gives about 55%), then the total non-glucose carbohydrate component of the reducing substances would have a maximum value of about 6 mg./100 ml. The estimated normal level is closer to 2 mg./100 ml. or,

TABLE IV
Glucose and reducing substances levels in C. borealis and C. sapidus carrying egg sponges

Species	N	Glucose mg./100 ml.	S.E.	Reducing mg./100 ml.	S.E.	Gluc/Red × 100
<i>Cancer borealis</i>						
no eggs	43	6.59	(0.47)	10.31	(0.55)	64 ^c / _c
new eggs	9	12.36	(1.18)	14.47	(1.19)	85 ^c / _c
old eggs	1	8.00	—	8.20	—	97 ^c / _c
<i>Callinectes sapidus</i>						
no eggs	7	18.47	(2.36)	—	—	20–25 ^c / _c
new eggs	7	37.61	(2.77)	—	—	20–25 ^c / _c
old eggs	9	10.52	(1.03)	—	—	20–25 ^c / _c
<i>C. sapidus</i> data from Dean and Vernberg (1965b)						

about 60–65% of the non-glucose component. In another study of the lobster, *Homarus americanus*, the non-glucose part of the reducing substances was estimated to be about 50% carbohydrate (Telford, 1968a, 1968b).

In the study of the lobster referred to above a cycle of changes in blood sugar levels during the molt cycle was found in a group of 800 animals. The apparent cycle found here in *C. maenas*, although the number of specimens is much lower, is closely similar. Molting of crabs on the coast of Maine occurs principally in the late summer and fall. The seasonal changes in *C. borealis* probably reflect the molting cycle with glucose increasing in premolt (August–September) and dropping in postmolt. The differences here are not clearly defined because the different molt stages were not separated, the monthly samples being made up of a changing proportion of premolt and other animals. In previous studies such a cycle of changes has not been found although expected (McWhinnie and Scheer, 1958; Scheer, 1959; McWhinnie and Saller, 1960).

As a regular component of crustacean blood fructose has not previously been reported. It was reported as occasionally occurring in *H. americanus* (Telford,

1965; 1968a, 1968b) but it was not found by either McWhinnie and Saller (1960) or Dean and Vernberg (1965a). It has been reported in some insects (Levenbrook, 1950). No sign of the curious "galactan derivative" of McWhinnie and Saller (1960) was found, nor of galactose. Maltotetrose has often been reported by other workers but no spot corresponding with its probable position could be found with the detection reagents used here, nor by examination under UV light when tetroses should fluoresce. Following molting the blood of *C. maenas* has only traces of sugars other than glucose; the oligosaccharides are severely depleted as in lobsters at this stage (Telford, 1968a).

This work was supported in part by a National Research Council of Canada studentship made to the writer and a Research Grant to Prof. J. Marsden. The assistance of the fishermen of Port Clyde, Maine, is also gratefully acknowledged.

SUMMARY

1. Blood glucose levels in the crab *Cancer borealis* ranged from 0.4 mg./100 ml. to 29.3 mg./100 ml. with a mean value of 8.6 mg./100 ml. In the same animals blood reducing substances were in the range 3.3–29.5 mg./100 ml. with a mean of 11.6 mg./100 ml. Blood glucose and reducing substances in the other two species of crabs tested, *Cancer irroratus* and *Carcinus maenas*, were in the same ranges.

2. Changes in the blood glucose level account for most of the variations in reducing substances; the other components remain approximately constant, at about 3 mg./100 ml. The relationship between blood glucose and reducing substances is thus a simple straight line one.

3. Variations of blood sugar levels during the molt cycle were found in *Carcinus maenas* and probably occur also in *Cancer borealis*. Qualitative changes in blood sugar composition also occur.

4. Significant changes also occur in *Cancer borealis* females carrying eggs and at this time the relationship between blood glucose and reducing substances changes, the two values converging as the egg mass ages.

5. Paper chromatography of the blood of these three species of crabs shows the presence of glucose, fructose, maltose, an unidentified reducing disaccharide and occasional traces of maltotriose and trehalose.

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THE BIOLOGICAL BULLETIN

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AUGUST, 1968



Printed and Issued by
LANCASTER PRESS, Inc.
PRINCE & LEMON STS.
LANCASTER, PA.

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THE BIOLOGICAL BULLETIN is issued six times a year at the Lancaster Press, Inc., Prince and Lemon Streets, Lancaster, Pennsylvania.

Subscriptions and similar matter should be addressed to The Biological Bulletin, Marine Biological Laboratory, Woods Hole, Massachusetts. Agent for Great Britain: Wheldon and Wesley, Limited, 2, 3 and 4 Arthur Street, New Oxford Street, London, W. C. 2. Single numbers, \$3.75. Subscription per volume (three issues), \$9.00, (this is \$18.00 per year for six issues).

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