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PRACTICAL EXERCISES IN ZOÖLOGICAL MICRO-TECHNIQUE

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President (1916), The American Nerroscopical Society

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REVISED EDITION



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PREFACE TO THE FIRST EDITION

For the past ten years it has been a part of the writer's duties to give instruction in microscopical technique, and it has seemed to him that there is need for a series of practical exercises which will serve to guide the beginner through the mane of present-day methods, with the greatest economy of time, by drilling him in a few which are thoroughly fundamental and standard. The book is intended primarily for the beginner and gives more attention to the details of procedure than to discriminations between reagents or the review of special processes. The student is told what to do with his material, step by step, and why he does it; at what stages he is likely to encounter difficulties and how to avoid them; if his preparation is defective, what the probable cause is and the remody. In short, the book attempts to familiarize the student with the hitle "tricks" of technique which are commonly left out of books on methods but which mean everything in securing good results.

A very brief, non-technical account of the principles of the microscope is inserted (Appendix A) with the idea of giving the student just enough of the theoretical side of microscopy to canble him to get satisfactory results from his microscope. The microscope is so ably treated in the excellent works of Gage (The Microscope) and Carpenier (The Microscope and Its Revelations) that the writer fields himself absolved from any further responsibility in this matter.

The aim of the entire book is to be practical: to omit everything that is not essential; and, above all, to give definite statements above things. Appended to each chapter is a series of memoranda which serve to supply additional information that is more or less pertinent without obscuring the main features of the method under consideration.

In Appendix B the formulae for a number of the most widely used reagents are given with comments upon their uses and manipulation. Following this (Appendix C) is a concise table of a large

number of tissues and organs with directions for properly preparing them for microscopical study.

Insamuch as every experienced worker has his own "hest" method for the preparation of almost any tissue, it is manifestly impossible to give all "best methods" in such a table. The writer believes, however, that the student will find the methods recommended all good ones which will yield satisfactory results.

In Appendix D some directions are given for collecting and preparing material for an elementary course in acology.

It is hoped that the volume will prove of use: (1) as a class textbook; (2) as a guide to the independent individual worker (teacher, physician, college or medical student, or novice); (3) as a reference book for teachers, in the preparation of material for courses in elementary nollogy, histology, or embryology.

In the matter of expressing his obligations the writer is at a loss to know just what to do. Many of the methods in microscopical technique have been handed down tradition-wise from one worker to another until their origin is unknown; they are the accumulated experiences of several generations of workers. Furthermore, many points have been absorbed, as it were, by the writer, from fellowworkers in the Universities of Chicago, Nebraska, and Cincinnati, respectively; consequently the obligation cannot be specifically expressed. Where the name of the originator of a method is known, due credit has been given. The books to which the author is most heavily indebted are the volumes of Gage and Carpenter, already mentioned, Lee's Microtomist's Vade-Mecum, Whitman's Methods in Microscopical Anatomy and Embryology, Hardesty's Neurological Technique, Foster and Balfour's Elements of Embryology, Minot's Laboratory Text-book of Embruology, Huber's translation of the Böhm-Davidoff Text-book of Histology, Stöhr's Text-book of Histology, Mallory and Wright's Pathological Technique, Bausch's Manipulation of the Microscope, and the Journal of Applied Microscopy. Grateful acknowledgment is also made to the various manufacturers of microscopical instruments and appliances for the loan of most of the cuts which have been used in this volume.

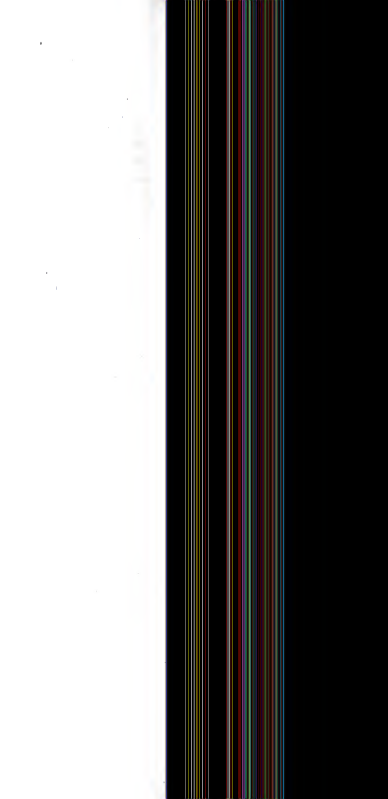
PREFACE TO THE SECOND EDITION

The forwable reception accorded the first ethion of Animal Mirodog has encouraged the author to believe that a second ethion, incorporating some of the many new methods which have appeared during the past ten years, would be equally welcome. The general plan of the book has not been altered (see Preface to the First Edition, on a preveding page), although changes have been made on nearly every page, many sections have been entirely rewritten, and two new chapters, one on "Cytological Methods," the other on "Draving," have been added. The chapter on drawing has been prepared by Dr. Elizabeth A. Smith.

In spite of a determined effort to limit the book to its former size, it has expanded by over fifty pages. For every method dropped there seemed to be a host of good new ones demanding recognition. These in the main, however, have been left to the encyclopedia and the various technical books and journals listed at the end of the values. As in the first edition, the policy has been, not to attempt to give all "best" methods, but rather to select representative good ones which have proved their worth by satisfactory tests in American laboratories.

Whatever merit the new edition may prove to have over that of the earlier one is due in no small measure to the many helpful suggestions of my colleagues in other colleges and universities. I am particularly indebted in this respect to Professors C. E. McChang, R. R. Bensley, H. McE. Knower, F. L. Landaere, F. C. Waite, B. M. Allen, George R. Ia Rue, Edward L. Rize, F. D. Barker, R. M. Strong, and H. L. Wieman, and to Doctors Elizabeth A. Smith and C. H. Hense.

M. F. G.



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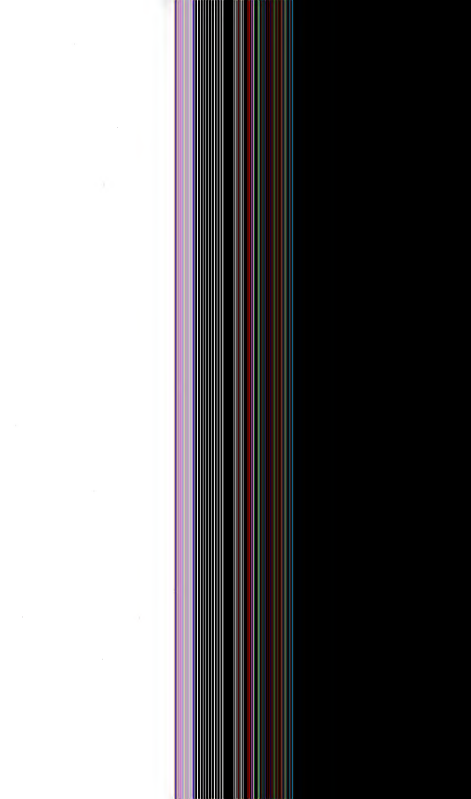
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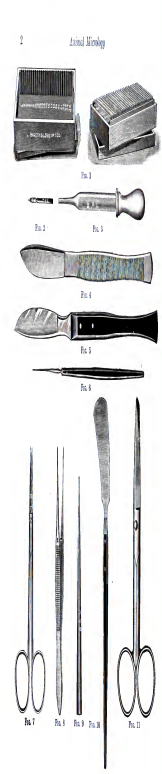


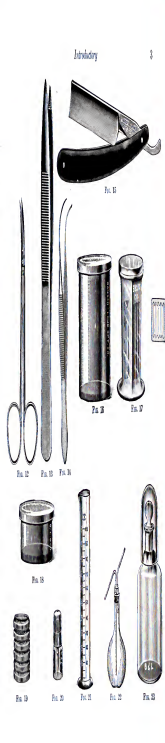
INTRODUCTORY

APPARATUS AND SUPPLIES REQUIRED

The student should provide himself with the following supplies:

One half-gross box best grade glass slides, standard size (25×75 mm.). One-half ounce, 18 mm. or 1/2 in., round cover-glasses, medium thickness, (0.18 mm.), Thirty 25×50 mm. cover-glasses, medium thickness. Two or three Pillsbury slide boxes (Fig. 1). One box of labels for slides. Three to six camel's hair brushes (Fig. 2). Six pipettes (Fig. 3). One set of dissecting instruments as follows: One large scalpel or cartilage knife (Fig. 4). One small scalpel (Fig. 5). Two needles (Fig. 6). One fine straight scissors (Fig. 7). One fine straight disserting forceps, file-cut points (Fig. 8). One blow-pipe (Fig. 9). One section lifter (Fig. 10). To which may well be added: One heavy suissors (Fig. 11). One curved seissors (Fig. 12). One heavy forceps (Fig. 13). One fine forceps, curved tips (Fig. 14). One horn spoon. One desk memorandum calendar. Blank cards (about 75×100 mm.) for keeping records of experiments. The kind of eard used for library card catalogue will do. One section razor (Fig. 15). A piece of moderately heavy copper wire with one end hammered out to a width of 7 to 10 mm. Towels. A glass-marking pencil (wax) or writing diamond will be found useful, Coarse carborundum "engraver's peneil points," which may be purchased for seventy-five cents a dozen, are very satisfactory for marking glass, according to Professor C. E. McClung,





Apparatus ordinarily supplied by the laboratory; Desk with drawers. Locker for microscope. Compound microscope and accessories (Appendix A). Dissecting microscope (Fig. 66). Microtomes (Figs. 27, 28, 29, 32, 33). Paraffin oven (Figs. 24, 25, 26). Tall stenders (about 85 mm. deep). Each student should have at least eight (Fig. 16). Coplin staining jars (Fig. 17). Tall stenders may be used instead. About eight are needed for each student. Flat stenders (Fig. 18); half a dozen for each student. Syracuse watch-glasses (Fig. 19); eight to each student, Balsam bottle (Fig. 20). Graduated cylinders for measuring liquids (Fig. 21). Wash-bottle (Fig. 22). Celloidin bottle (Fig. 23). Turntable (Fig. 36). Injecting apparatus. Reagent bottles and vials. Other apparatus and supplies such as bone-forceps, bone-saws, glass tabing, glass rols, beakers, burners, filter-paper, funnels, evaporating-dishes, sand bath, dropping-bottles, balances, mortar and pestle, etc. For apparatus or supplies not listed in this book the student is referred to the illustrated catalogues of dealers and manufacturers such as: The Bausch & Lomb Optical Co., Rochester, N.Y.; The Ernst Leita Optical Works, Wetzlar, Germany (American branch, 30 E. 18th St., New York City); The Spencer Lens Co., Buffalo, N.Y.; Carl Zeiss Optical Works, Jena, Germany; R. & J. Beck, 68, Comhill, London; The Kny-Scherer Co., New York City; Eimer & Amend, New York City; Whitall, Tatum & Co. (especially for glassware), New York City.

IMPORTANT GENERAL RULES

1. Keep everything clean!

2. Have a definite place in your desk for each piece of apparatus and arrange reagents in order on top of it.

3. Use eards for keeping records of materials. Each card should have a number corresponding to that of each special object or piece of tissue, and should show the name of the preparation, date, reagents used, time left in each reagent—in short, all data concerning the manipulation of the material.

4. Jot down in a blank calendar the various things to be done at future dates, such as elonging of reagent on tissues, etc., and then go over this memorandum earchally each day when you first come into the laboratory.

5. Use only clean vessels in preparing reagents, and clean up all glassware while it is yet moist.

6. Reserve and mark a separate pipette for each of the chief reagents (absolute alcohol, oils, acids, etc.).

7. In making up solutions, 1 gram of a solit in 100 e.e. of liquid is redoned ordinarily as a 1 per cent solution, 3 grams as a 3 per cent solution, etc. But if solutions are to be of 10 per cent strength or over, it is better to weigh out the day material to the desired percentage and then add enough of the liquid to make the whole weigh 100 grams. For example, to make a 25 per cent aqueous solution of eastice potash, add 25 grams of caustic potash hor 75 e.e. of water. A saturated solution contains all of a given substance that the liquid will take up. When a solution is called for without specifying the solvent, an aqueous solution is meant.

8. In weighing salts always first put paper in the scale pans to protect them.

9. In making solutions or mixtures in which only a small amount of one reagent is used, after mixing, pour back some of the mixture into the small vessel and rinse it thoroughly in order to get all of the original contents out.

10. When pouring liquids from bottles keep the label of the bottle turned toward the palm of the hand. Do not lay down stoppers but hold them by their tops between the knneckles.

 Before leaving the laboratory put away your instruments and clean and put in its place whatever laboratory apparents you may have been using.

12. All solid waste materials, acids, stains, etc., should be thrown into stone jars, not into the sink.

CHAPTER I

PREPARATION OF REAGENTS

The following reagents should be prepared by each student. 1. Grades of Alcohol.—To obtain a given percentage of alcohol through dilution of a higher percentage with distilled water, subtract the percentage required from the percentage of the alcohol to be diluted; the difference is the proportion of water that must be added. Thus, if 35 is the percentage required, and 95 the percentage to be diluted; then 95–35=60; hence, 60 pasts of water and 35 parts of 95 per cent alcohol are the proportions for mixing.

This means that in practice one needs only to fill the graduated measuring cylinder to the same number as the perentage required (e.g., 35) with the alcohol to be diluted (e.g., 95) and then fill up to the percentage of the latter with distilled water. In this way one would obtain 95 c.e. of alcohol of the percentage required, if the measuring cylinder is graduated in exbine centimeters.

Prepare about 230 e.c. of 35, 50, 70, and 83 per cent alcohols, respectively, from 95 per cent alcohol and distilled water. The commercial alcohol used, though really about 96 per cent, may be figured on the basis of 95 per cent.

Oving to the differences in the specific gravities of the different percentages of alcohol, the foregoing method gives only approximate results; they are sufficiently accurate, however, for most biological work.

2. Absolute Alcohol.—It is customary in most laboratories to purchase so-called absolute alcohol specifilly prepared for laboratory purposes. Spuibl's absolute alcohol (90.8 per cett) is commonly used. Inasmuch as such alcohol is an expensive reagent, economy sometimes necessitates that the student undertake the more tedious process of making his own absolute alcohol. Crystals of copper suphate are heated until the water of crystallization is driven off and the subplate is fit as white powder. Such anhydrous subplate

is added to a bottle of commercial (96 per cent) alcohol. The water in the alcohol immediately unites with it, turning it blue. Anlydrows subplate should be added until it no longer turns blue. The alcohol is then filtered into a clean, dry bottle which must have a tight-fitting cork or ground-glass stopper. It is well to smear the glass stopper with vaseline, so that when it is placed in the bottle all moisture from the air may be completely excluded. Any laboratory using considerable quantities of absolute alcohol should have its own still.

3. Acid Alcohol.—

8

For sections use the mixture only a few seconds or minutes. For material stained in bulk, add twice as much 70 per cent alcohol and leave the object in it until sufficiently decolorized (2 to 24 hours).

 Ether and Alcohol.—Absolute alcohol and sulphuric ether equal parts. Quantity, 400 e.e. Keep the ether distant from all facues.

5. Normal Saine.—Prepare a 0.75 per cent solution of sodium ehloride in distilled water. This is termed a normal sait solution because it is a solution of about the same density as natural lymph and is much less harmful to living tissues than is distilled water. Quantity, 500 c.e.

6. Formalin (also termed formal, tormol, formolose).—Commercial formalin is a 40 per cent solution of formaldehyde in water. A 4 per cent solution of formalin would be made by taking 4 volumes of commercial formalin and 96 volumes of water. This is, however, only a 1.6 per cent solution of formalidehyde. Make a 10 per cent solution of formalidehyde.

7. Zenker's Fixing Fluid.-

Bichromate of potassium.		
Bichloride of mercury (corrosive sublimate)	5.0) grams
Sodium sulphate	1.0) gram
Water	100	6.6.
Glacial acetic acid	ő	e.e.

Preparation of Reayants

9

Dissolve the bichromate and the sublimate in the water with the aid of heat. Keep the acetic acid in a separate tottle until the fixing fluid is to he used, as it will produce charges in the chrome salt if added at once and allowed to stand.

CAUTION.-In handling corrosive sublimate do not use metal instruments because it corrodes metal. Use a glass or horn spatula.

8. Bouin's Picro-Formol.—

Pictic acid, saturated aqueous solution	75 parts
Formalin	25 parts
Acetic acid (glacial)	5 parts

One gram of picric-acid crystals will saturate about 75 c.c. of

water.

9. MacCallum's Macerating Fluid .-

Nitrie acid.	1 part
Glycerin	2 parts
Water	2 parts
10. Decalcifying Solution	
Nitrie acid (strong)	10 c.c.
Alcohol (70 per cent)	99 c.c.

11. Alum-Cochineal.-

Potassie alum	6 grams
Powdered cochineal	6 grams
Distilled water	90 c.c.

Boil for half an hour; after the fluid has settled, deennt the supernatant liquid, add more water to it, and boil it down until only 90 c.e. of the decordion remains. Fifter when cool, and add a bit of thymol or a little salicylic acid to prevent the growth of mold.

12. Delafield's Hematorylin—Prepare 100 c.c. of a saturated aqueous solution of annonia alum. Dissolve I gram of hematorylin crystals in 10 c.c. of absolute alcohol, and add it, drop by drop, to the first solution. Expose this mixture to air and light for several weeks (two months is not too long) to "ripen." (Ripening consists in an oridation of the hematorylin to form hematein. This may be accomplished at once with some degree of success through the addition of a few cubic continueters of a neutralized solution of percoide of hydrogen or other powerful oxidizing agent.) When ripe, filter the

solution and add 25 c.c. of glyverin and 25 c.c. of methyl alcohol (see memorandum 1). It is well to have a stock solution of this stain already prepared to be used in case the student's preparation is not ready in time.

Most laboratories keep on hand a stock solution of hematorylin made by dissolving 1 parts of hematorylin crystals in 10 parts of absolute alcohol. In the course of several monitos or a year this solution ripens to a dark wine-red color. It may then be used in making up the various hematorylin solutions and, being ripe, will stain at once.

Norm—At this point the student should begin chep, it in order that no time may be lost. The present chapter may then be completed while the tiesees are becoming fixed and hardened.

13. Orange G.—

Orange G (Grübler's)	1	gram
Distilled water	100	e.e.

14. Congo Red.-

Congo red	0.5 gram	
Distilled water	100 c.c.	

15. Lyons Blue.—

Absolute alcohol	100	C.C.
Bleu de Lyon	0.3	gram

16. Eosin.—

Bosin	0.5	i gram
Alcohol, 95 per cent	100	C.C.

17. Iron-Hematorylin (Heidenhain's).--Two solutions are used. They are not to be mixed.

Solution I:

Ferric alum (clear violet crystals)	2.5	gram	
Distilled water.	100	2.0	

Solution II:

Hematoxylin	0.	5 gram	
Distilled water	100	3.3	

Preparation of Roagents

The hematoxylin should ripen (see reagent 12, p. 9) for some three or four weeks. The ferric alum of the histologist is always annonic-ferric sulphote.



Frg. 24.—The Lillie Water-Bath

The bath crucisis of a large character centraling a series of densers of equal size 250 mm long. (When, with, Sham due), Each disearch as expertent as all others where a set stars and have represented in the stream representation of a series of the drawners are separated by performing efficience densities and run on Sales from from the local supports, thus permitting efficience densities and run on Sales from from the local supports in the permitting efficience densities of sum as in the series equal series paraterine in the span show of the hard. Assume and infrastructures have equal solution into the series of the stars are series of the series of th

18. Canada Balsam.—Dry 2 grams of Canada balsam on a smul bath or in a warm chamber until it becomes hard (1 to 2 hours at 65° C). Do not orerheat. When cool add enough xylol to make a very thin, syrapy fluid. Roll a sheet of paper into a cone to serve

as a funnel, and filter the fluid through absorbent cotton. Thicken



Fig. 25 .- Simple Water-Bath

watch-glass incledding or for warming instruments, and tubulatures for gas

regulator and thermometer.

This is a useful bath for individual workers. It is provided with imbed-ding-cups, infiltration vials, a shelf for

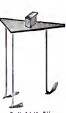
the solution slightly by leaving the cap off the bottle in a place free from dust, and allowing some of the xylol to evaporate. Or, fill your balsam bottle one-third full of the liquid xylol-balsam now on the market and dilute to the proper consistency.

19. Mayer's Albumen Fizative. -Beat the white of an egg with an eggbeater and pour it into a tall cylinder. Let stand until the air brings all suspended matter to the top. Skim off the latter and to the remainder add an equal volume of glycerin, and a bit of salicylate of soda (1 gram to 50 e.c.) or thymol to prevent putrefaction.

20. Celloidin.-Soak 15 grams of dry celloidin overnight in just sufficient absolute alcohol to cover it. Then dissolve it in 200 e.e. of

ether-alcohol (see reagent 4). In a second bottle prepare a thinner solution by taking about one-third of the original and diluting it with its own volume of ether-alcohol. The solutions are best kept in totties with glass stoppers and ground-glass caps. Label the bottles thick and thin celloidin, respectively. bottles with glass stoppers and

melting at about 50° C. (see memo-



response by placing the starts with Restarts the star-prome by placing the time at the 21. Parafin.—In one of the corns of a warm parafin oven (Fig. 21, bits of the placing the starts and 25, or 20), put 75 grams of parafilm history the start of the corns. In the starts and the placing the starts and the starts and a start of the starts and the starts. In the starts and the sta the burner.

randum 5, p. 14). The bath should be kept at a temperature

Preparation of Reagents

13

of some two degrees above the melting-point of the paraffin. A supply of softer and of harler paraffin (e.g., melting at 45° and 55° C.) should also be at hand.

Other Reagents.-Provide yourself with 500 cc. of distilled water, 200 ec. of xylal, 25 cc. of clove oil, 25 cc. of glacial accide acid, 50 cc. of a cedra-wood oil, 25 cc. of a saturated solution of iotime in 70 per cent alcohol, 75 cc. of chlorotorm, 30 cc. of glycerin, and 250 cc. of a headure alcohol if it has not already been prepared. Keep the absolute alcohol and the xylol cancilly corked to exclude moisture. Before measuring out any of these reagents, see that both the graduate and the bottle are perfectly clean and dry.

MEMORANDA

 Rayl Alcohol (grain slookd) is the kind cummonly used in histolegical laboratories. Upon presentation of the proper credentials to the internal revence officers, it may be purchased by the hared from distillers, tar free, by educational institutions. Such commercial alcohol is of adout 96 per send strength. When the strength is unknown, it should be tested by means of an alcoholometer (see 2, below).

Methyl Ababal (called also wood alcohol or wood spirits) is cheaper than ethyl alcohol in case the latter cannot be had tax free, and is fairly satisfactory in most cases. It is poisonous and must be carefully handled. It is of about 90 per cent strength. The "methylated spirits" of English microscopiesti is guin alcohol containing 10 per cent of methyl shedool.

Rectified Spirit is a 91 per cent alcohol (84 per cent in England).

2. The Alcoholimeter is a convenient instrument for determining the strength of alcohol, or the preventage of alsohol in a spiritrows mixture. It is a kind of hydrometer with a scale marked to indicate the preventages of alcohol. Different strengths of alcohol have different specific gavities; consequently the instrument will fact higher or lower in the logitid according to the preventage of alcohol present. The number on the scale just at the surface of the liquid indicates its strength.

3. Rate for Dhrinn of a given strength of a solution with a lower precentage of the same solution. (For where the dilucat is water, i.e., zero per cent, see rule under reagent 1.) Solutest the percentage required from the percentage of the solution to be diluted; also solutest the percentage of the dilucat from that of the strength required. The differences are the relative proportions of the dilucat and the solution to be diluted that must be used. Thus, to prepare a 35 per cent solution from 95 and 20 per cent solutions: 95–35=07; 35–30=15; hence, 60 to 15 or 4 to 1 are the

proportions desired. That is, 4 parts of the 20 per cent and 1 part of the 25 per cent solution must be used to obtain a 35 per cent solution.

4. "To Remove Fined Stoppers, take the hottle in the left hard with the thumb sequiled to one side of the stopper, then tap the other side of the stopper with some heavy instrument, such as the handle of a proket-knife, pressing the thumb against the direction of the tap. Turn the bottle round, gradmally tapping until the stopper locests. Should his device prove of no avail (which is very early), hold the neck of the bottle in a spirit frame, and quickly withdraw the stopper as the glass of the neck expands. This is a somewhat risky procedure, but is very effectual if done smarthy" (Journal of Applied Microsopy, VI, 2116). The glass of the neck and forth until the histin warms the glass.

5. A Simple Parafin Bath, recommended by several workers, may be made by suspending an electric-light bulk (earbox-flauxent) in a tumbler of parafin. The beight of the bulk should be so adjusted that some unmelied parafin remains at the bottom. Tissues will thus come to be where the parafin kip at at the melting-point. Professor F. C. Waite recommends putting a paper core around the entire apparatus if the room is odd. McClendon, in the Biological Bulletin (XV, No. 1) for June, 1912, explains how to make a convenient concrete parafin buth for individual use which is inequentive and effective.

McClung suspends a 50-analle-power helin-wound earlow-flament lamp, provided with a lamp shade, over a backer of parafin. The heat from the lamp is sufficient to keep the top of the parafin melicit. Rapid evoporation of the dealechologing egent is facilitated by such an arrangement and overheating is avoided. The same lamp may be used for spreading the parafin ribbon on slikes, and for drying.

CHAPTER II

GENERAL STATEMENT OF METHODS

Each of the reagents which has been prepared is used for one or more of the purposes to be discussed in this chapter.

All methods of preparation in microscopy are to enable us to learn more of the structure and functions of objects than would otherwise be apparent. We endeavor to study them in as near their natural condition as possible. While the study of living or of fresh material is desirable it can be carried on only to a very limited extent. Most structures of the animal body, though opaque, must be examined largely by transmitted light, hence special preparation is necessary to put them into suitable condition. This is accomplishel—

- 1. By cutting them into thin slices (section method).
- 2. By separating them into their elements (isolation)-

a) Mechanically (teasing), or

b) With the aid of fluids which remove the cement substance (dissociation or moceration).

In most instances the minute structure of a tissue or of an organism can be studied to the best advantage only after the application of certain agents which serve to emphasize the various structural elements. A tissue so prepared is an artificial product in that it is not exactly the same as it was in the living organism, but recent studies of protoplasm in the living condition by competent investigators strengthen the belief that many reagents preserve very faithfully the actual structure of the cell contents. The liquid albuminoids are apparently the materials which suffer the greatest modifications. Since alterations do occur, however, it is char that in our interpretations of prepared material we must reckon carefully both with the original nature of the object and with the factors jutroheed by ourselves.

KILLING, FIXING, AND HARDENING

The first step in the preparation of tissues ordinarily is the employment of some reagent which will kill the tissues and fix their various components in the characteristic stages of their activities. Such material may then be preserved indefinitely for future use.

It is customary to discriminate between killing, fixing, and hardening, although the same reagent may fulfil all three requirements. Killing refers particularly to the destruction of the life of the tissue, a process which may be either slow or instantaneous. In slow killing it is usual to employ narrotice such as ether, chloroform, chloral hydrate, chloretone, carbon dioxide, nicotin, eosain, or weak alcohol. Ice is also used sometimes. Such methods are of particular value with highly contractile animals which are desired in the extended condition. Such forms are narrotized completely or until they are unable to contract, and then frequently fixed and hardened in other or stronger thick. Where practicable, instantaneous killing and fixing is preferable because tissues have then no time to undergo postmortem changes. The same fluid ordinarily is employed for killing and fixing.

The purpose of fixation is-

a) To preserve the actual form of tissue elements.

b) To produce optical differences in structure, or so to affect the tissues that such differences will be brought out through subsequent treatment with stains or other reagents.

To accomplish this the fixing agent must possess the following properties:

 It should kill the tissue so quickly that few structural changes can occur.

2. It should neither shrink nor distend the tissue.

 It should be a good preservative; that is, it must render the tissue elements insoluble and prevent postmortem changes.

4. It should penetrate all parts equally well.

 It should put the tissue in condition to take stains unless it of itself produces sufficient optical differences in the various parts of the tissue.

No ideal single reagent has been discovered which meets all of these requirements, hence it is customary to combine two or more reagents which individually possess certain of these desirable qualities. All of the best fixing reagents are mixtures. For example, anothe add is very generally used in fixing mixtures because it penetrates well, produces good optical differentiation, and counternets the tendency of some reagents (e.g., corresive sublimate) to shrink tissues. Again, osmic add, which is an excellent fixing agent for very small pieces of tissue, penetrates very poorly; consequently for most objects it must be mixed with reagents which penetrate rapidly and thoroughly.

Some firing agents (corresive sublimate, chromic acid, osmic acid, etc.) enter into chemical combination with certain of the tissue elements; others (alcohol, picric acid, nitric acid, hot water, etc.) act by coagulating or precipitating certain constituents of tissues.

The chief object of hardening is to bring tissues to the proper consistency for entring sections. The process, although begun ordinarily by the fixing agent, is usually completed in alcohol. Some objects are not sufficiently hardened until they have remained in alcohol for many hours, or even days. As a rule, tissues should remain in alcohol of at least 70 per cent strength for a minimum of 24 hours after the preliminary operations of fixing, vashing, etc., before they are subjected to further treatment.

WASHING

Fining agents ordinarily, with the exceptions of alcohol and formalin, must be washed out thoroughly or they are likely to interfere with subsequent processes. Append solutions are washed out usually in water or a low percentage of alcohol; alcoholic solutions, with alcohol of about the same strength as that of the fixing agent. Washing usually requires from 10 to 24 hours, with several changes of the liquid. If water is the washing agent, it is best where practicable to use running water.

Chronic acid and its compounds should be washed out in running water. This should be done in the dark in order that presipitation may be avoided.

Pierie acid, or solutions containing it (except piero-formalin mixtures), must be washed in strong alcohol (70 per cent), never in water, because the latter seems to undo the work of fraction.

Corrosite sublimate and mixtures containing it are washed out in water or alcohol. A little functure of iofine should be added to the wash from time to time to insure the removal of all corresivesublimate crystals. Sufficient iodine has been added when it no longer loses its reddish color after being in contact with the preparation for a short time.

Osmic acid and mixtures containing it should be washed in running water.

DEHYDRATING

While under certain circumstances objects may be mounted in aqueous media for examination, in the majority of cases, especially where the preparation is to be a permanent one, it has been found best to remove all water from the tissues, that is, to dehudrate them. This renders preservation more certain, and it is a necessity, moreover, if the object is to be imbedded later in paraffin or celloidin, for neither of these substances is miscible with water. Because of its strong affinity for water and the ease with which it may be manipulated, alcohol has come to be used universally for this purpose. It completes the process of hardening at the same time. The dehydration must be gradual. In tissues transferred from water or aqueous solutions directly to strong alcohol (or vice versa) violent diffusion currents are set up which produce serious distortion of the tissue elements. For this reason a series of alcohols of gradually increasing strength (e.g., 35-50-70-83-95 per cent) is used. The more delicate the object, the closer should be the grades of alcohol.

Professor C. E. McClung recommends a "drop" method for all purposes. If, for example, an object in water is to be earlied up into the higher alcohols, he places a vessel containing 95 per cent alcohol and the vessel containing the object in water under a bell-jar. The vessel containing alcohol is raised above the level of the other vessel and a string or a capillary siphon is set to carrying over the alcohol drop by drop into the water. The amount of 95 per cent alcohol must be apportioned to the amount of water so that the final

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desired strength will be reached by the time the alcohol has all passed over into the water. The ressel containing the specimen should be agitated frequently to scene thorough mixture. Obviously other fluids may be changed by the same method. For a more claborate form of drop apparatus see memorandum 6, p. 182.

PRESERVING

After fining and washing, the process of dehydration is begun ordinarily and tissues are earried as far as 70 per cent alcohol. It is customary to leave them in alcohol of from 70 to 83 per cent strength until they are needed. They may remain here indefinitely. If they are to be preserved for a long time (for months), however, it is better to keep them in a mixture of equal parts of glyverin, distilled water, and strong (commercial) alcohol.

STAINING

A few fixing agents produce sufficient optical differentiation in tissues, but as a rule this must be accomplished through the addition of certain stains. Most of the stains used have more or less of a selective action; that is, they nick out certain elements of the tissue, and thus enable one to see details of structure that would otherwise be invisible. Their action, however, depends largely upon the nature of the fixing agent which has previously been used. The secret of good staining, indeed, has largely in proper fixation.

There are large numbers of stains of very different chemical constitution (acid, neutral, and alkail), and they may act in very different ways upon the material to be stained. For example, some show affinity only for certain elements of the nucleus, others for the cytoplasm of cells, and some are present in tissues only physically as deposits, while others enter into chemical combination with certain of the cell constituents. A few, such as borax-carmine, are general stains, and affect to a greater or less degree practically all the tissue elements.

It is not the purpose of the present book to enter into a prolonged discussion of the theory of staining or to undertake a description and classification of stains. For this the reader is referred to the excellent compendium of Lee (The Microtomia's Vade-Merum).

The stains of widest application are (1) the **Carmines**, (2) the Hematorylins, (3) the Anilins, and (4) Metallic substances.

Commine is a brillant scarlet or purplish coloring matter mode from the holies of the evclineal and kennes scale inserts. The examine stains, including coefficient, have been largely used in the past for all kinds of work, but at present they are used more particularly for staining objects in balk before sectioning, or objects which are not to be sectioned. They are easy to use, and will follow almost any fixing agent. In case of overstaining, weak hydrochlorie acid (0.1 to 1 per cent) is used to decolorine the tissues. For formulae see Appendix B, p. 220.

Henotaglin is a compound containing the coloring matter of logwood. The hematoxylins follow well almost any of the fixing agents; they are especially recommended after fluids containing chromic acid or its salts. According to Mayer, the active agent in these stains is a compound of hematein with alumina. This blue-colored esolution is precipitated in tissues, particularly in model, by certain organic and inorganic salts, such as phosphates. The hematein is produced by the oxidation of hematoxylin. The socalled "ripering" is simply this change, which is brought about by exposing the hematoxylin solution to air. If the pure hematein is used in making the siam, therefore, the latter will be ready for use immediately, because it need not undergo the ippening process (see remarks under 12, p. 9). For formulae see Appendix B, p. 225.

And/a is a colorless coal-tar derivative, and is the base from which many of the numerous coal-tar dyes are made. The callins are brilliant stains of all colors. They are used almost exclusively for staining sections or thin membranes, and are of great service to the microscopist, although, as a rule, they fade in time.

The basic anilin stains, such as methyl green, methyl rolet, gentian violet, methylen blue, safranin, Bismarek brown, tobidin blue, and thionin, are usually nuclear stains. On the other hand, the acid anilin stains, such as acid fuchsin, eosin, erythresin, light green, orange G, bleu de Lyon, nigrosin, benzopuryurin, and aurantia,

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are ranked as cytoplasmic stains. Some of these stains must be made up fresh every two or three months, as they frequently spoil if kept much longer.

The metallic substances used for color differentiation operate principally as improportions rather than as stains. The coloring matter is held physically as a precipitate or reduction product in certain of the tissue elements. The commonest reagents of this class in use are silver nitrate and gold chloride.

The different tissue elements frequently show affinity for different stains; consequently it is a common practice to use more than one stain. Very decided contrasts may thus be produced, such as rod and blue, red and green, green and orange, etc. It is not uncommon, in fact, to have triple and even multiple staining. In such staining the stains are sometimes applied consecutively; in other cases, at different points in the process of general manipulation. Sometimes all the stains may be mixed together, so that immersion of the sections in one liquid is all that is required for double or multiple staining.

A general rule in staining, especially for entire or bulky objects, is that the specimen should be transferred to the stain from a reagent in which the percentage of water is approximately the same as that of the stain. The same is true when the object is removed from the stain. For example, if the stain to be used is an aqueous solution, the object should enter it from an aqueous solution; if the stain is made up in 95 per cent alcohol, the object should enter from 95 per cent alcohol, etc. For reasons see "Dehydrating."

CLEARING

In the vast majority of cases tissues are too opaque for satisfactory examination until they have been treated with certain charifying reagents or *charava* which render them more transported. Such reagents as glyrerin, glyrerin-jelly, etc., are used when the object is to be chared, without alcoholic dehydration, directly from water. Usually, for permanent preparations, the alcoholic dehydration method is employed and it then becomes necessary to use a charifying reagent which will replace the alcohol and facilitate the penetration of the final mounting-medium (balism or damar).

Perhaps the most useful and rapid clearer is xylol. Xylol, however, is very sensitive to moisture, and if the preparation has not been thoroughly dehydrated the final mount will appear milky, Cedar-wood oil, though somewhat slower than xylol, is one of the best clearers. It is also one of the safest, because tissues may be left in it indefinitely. Other good clearers after alcohol are oil of origanum, sandal-wood oil, oil of cloves, cassia oil (cinnamic aldehyde), toluol, oil of bergamot, anilin oil (for watery specimens). carbolic acid (for watery specimens), and beechwood ereosote, Anilin oil will clear from 70 or 80 per cent alcohol. It should be followed by oil of bergamot, cassia, or wintergreen, according to McChung. Clove oil should not be used for celloidin sections because it dissolves celloidin. It is also inapplicable ordinarily after most anilin dyes because of its tendency to extract them. Among the best reagents for celloidin sections are cedar wood oil, oil of origanum, creosote, and Eycleshymer's clearer (memorandum 4, p. 63).

While "clearing" refers especially to the rendering transparent of tissue elements, and *dealcoholication* to the removal of alcohol previous to imbedding in parafin, very frequently the same reagent is used for either purpose and the term "clearing" has come to be used in either sense.

MOUNTING

After tissues have been cleared the final step is to mount them in some suitable medium for preservation and inspection.

If tissues are to be mounted directly from water or aqueous media, glycerin, glyrerin-jelly, or Farnan's solution is used ordinarily. If the alcohole dehydration method is employed, balsam, gun damar, or enparal is the final mounting-medium. The balsam or damar is dissolved commonly in xylol, although turpentine, ebloroform, or benol may be used as the solvent. In my experience xylol-balsam is the most satisfactory for ordinary purposes. However, some of our best American technicians prefer gum damar dissolved in xylol.

IMBEDDING

In order to section tissues or objects satisfactorily it is frequently necessary to imbed them in a suitable matrix. Simple indeeding consists in merely surrounding the object by an appropriate medium to hold it in place while it is being cut. In interstitiol inhelding the object is saturated (infiltrated) with the inhelding-substance which, when all cavities and interstiess are filled, is caused to set; thus it supports all parts of the tissue and holds the components in place when sections are made. Infiltration inhelding is of great importance to microscopists and much of the space of the present book is given up to drilling the student in the details of the two chief infiltration methods, viz, the parafin method and the colloidin method. Infiltration with gun is also not infrequently resorted to, especially for tissues which would be injured by alcohol, or for sectioning by the freezing method.

Parafin is a transhoent, waxy material derived from various sources, one of the commonest of which is crude petroleum. Paraffins of low and of high melting-points, termed respectively soft and hard parafin, should be kept on hand so that mixtures of different degrees of hardness may be made up as necessity demands.

Celloidin is a form of pyroxylin (guncotton or colloilon cotton) specially prepared for interstitial imbedding. It is dissolved in a mixture of ether and alcohol (p. 8, reagent 4) and solutions of two or three strengths are used for infiltration. For details see the method, p. 30. Colloilon instead of celloidin is used by some workers (see memorandum 11, p. 65).

AFFIXING SECTIONS

When mounting sections upon a slide, especially if they are yet to be stained, it is usually necessary to affit them firmly to the slide to prevent later displacement. For parafin sections Mayer's albumen fixative (reagent 19, p. 12), or a combination of this method with the water method, is most widely used. The water method alone often proves adoptate, particularly with thin sections. The slide is flooded with water, or better, albuminized water mode by adding 3 drops of albumen fixative to 30 cc. of distilled water, and the sections are fixated upon its surface. The parafin ribbon should be gently heated until it becomes translatent but not melted, in order to make it spread and flatten properly. As the layer of water eraporates, the sections are slowly drawin down into close contact with the slide. When perfectly dry they are usually so firmly affixed

that they will not become detached even after the removal of parafin from them. It is common, however, and safer to use a thin film of allouren fuxative as a cententing substance between the water and the surface of the file.

In the case of celloidin sections, if only one or a few sections are to be mounted on one slide, it is a common practice to stain the sections and transfer them through the various reagents, even to clearing, before mounting them on the slide. In such cases the sections need not be fixed to the slide. With serial sections, however, the sections must be held in place in some way during their transition through the reagents (see memoranda 12 and 13, p. 165). Unlike parallin, the celloidin is not ordinarily removed from the tissues.

DECOLORIZING

Not infrequently in staining, the tissue becomes overstained and requires that some of the color be extracted from certain of the elements to bring about a proper differentiation. The fact that certain tissue elements retain stain more transionsly than others is sometimes taken advantage of and overstaining followed by decoloriration is practiced intentionally. Alcohol slightly acidulated with hydrochlorine acid (0.1 to 1 per cent) is commonly used for the extraction of surplus color. In special cases other decoloriners are used: for example, iron-alum in the iron-benatoxylin method (reagent 17, p. 10).

Overstaining tissue and then partially developing it is sometimes designated as repressive staining in contradistinction to propressive staining in which the dye, once taken up by the tissue, is not removed. In progressive staining differentiation is accomplished through the selective affinity of dyes for different elements.

BLEACHING

In some cases tissues are observed because of the presence of natural pigments or on account of blackening caused by the fixing reagent. Such tissues must be blacked. Chlorine, percoide of hydrogen, or sulphurous acid are commonly employed. A method is given in memorandum 12, p. 44.

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CORROSION

To obtain skeletal structures, as, for example, the spicales of sponges or the hard parts of insects, various methods of corrosion are employed. Nitrie acid, caustic potash, caustic soda, cau de Javelle are reagents often used for this purpose. Corrosion preporations of injected vessels and cavities may also be made.

DECALCIFICATION AND DESILICIDATION

Tissues impregnated with time solts or with silica must have such hard parts removed usually before they can be sectioned. For detalefication, one of several acids may be used. The details are given in the chapter on hone, tooth, etc. (chap. xi). For detaleifying reagents, see Appendix B, v.

Where desilicitation is necessary hybriduorie acid may be employed, although, because of its property of attacking mucous membranes, its use is attended with more or less danger for the operator. It is added drop by drop to the tissue which has previously been placed in a parsifin-coated vessel (the acid attacks gluss). If the tissue is not too heavily impregnated with silica, it is safer to use an old section rator and try to out sections without previously treating them with hybriduorie acid.

INJECTION METHODS

The injection of colored masses into the blood vessels and other vessels of the body is frequently practiced to aid in determining their distribution and their relation to the surrounding tissues. The dye is termed the coloring mass and the substance to which it is added the reliab.

ISOLATION OF HISTOLOGICAL ELEMENTS

Isolation is one of the most valuable means of forming a correct conception of cells and fibers. It has the advantage over sections that the elements may be inspected in their entirety and from all sides. The separation is accomplished, as already noted, by (1) reagents which dissolve or soften cell cement and interstituil material without seriously affecting the cells (*macerulan or disaciation*), or (2) mechanically by means of dissecting meells (*leasing*),

or both. Harkening and fixing reagents in general if diluted to about one-tenth are efficient for dissociation. Gape recommends normal saline as preferable to water for dilution. The disserting microscope or some kind of lens-holder and lens are valuable aids in isolating tissue elements. For practical methods coasult chap, x; for reagents, Appendix B, iv.

NORMAL OR INDIFFERENT FLUIDS FOR EXAMINING FRESH TISSUES

It is desirable frequently to examine fresh material in as near a natural condition as possible, hence recourse is had to the so-called indifferent fluids. While not wholly indifferent, they ordinarily produce but slight changes in tissues and their elements from the viewpoint of the microscopist. The liquids most commonly used for this purpose are discussed in Appendix B, iii.

GENERAL SCHEME FOR MOUNTING WHOLE OBJECTS (IN 1010) PREPARATIONS) OR SECTIONS

Whole Objects (for balsam mounts) Killing and fixing		Section Methods (paraffin and celloidin) Killing and fixing			
Washing		Washing			
Staining		(Staining, if to be stained in bulk)			
(Decolorizing if necessary)		Hardening and dehydrating			
Dehydrating		Absoutte alcohol			
Clearing	D C. H.J.J		0.R.2. W.d.1		
Mounting	Parafin Method Dealcoholiaation (xylol)		Celloidin Method Ether-alcohol		
	Nelted paraffin		Thin celloidin		
R D	Imbedding		Thick celloidin		
lj not storned in balk	Sectioning	If not etmined in	Imbedding		
Tarough alcohols to stain	Affixing sections	bulk Staining	Sectioning*		
Staining	Removal of paraffir	Washing (and decolorizing if	Dehydrating to 95 per cent alcohol		
Washing	Absolute alcohol	nesessarà)	Clearing		
Dehydrating (and decolorizing if	Clearing		Mounting		
necessary)	Nounting		,		
*If sections are to be arranged serially they are best mounted in sheets or affixed to					

the slide as soon as cut.

CHAPTER III

KILLING AND FIXING

CAUTIONS.-1. Use only fresh tissues and work rapidly so that the tissue elements will not have time to undergo postmartem changes.

2. Remore organs carefully and avoid crushing or pressing the parts to be prepared.

 Tissue should never be allowed to dry from the time they beare the animal walk they are finally mounted for microscopical economation except at one point in the porufin method.

4. Use only small pieces (3 to 6 nm. code) of tissue whenever possible, or penetration of the reagent will be insufficient. Endryge and small dijects up to 4 cm. in size may be placed entire in certain of the fixing fields.

 For fixing and hardening, the bulk of the fluid should be from 10 to 30 times that of the object. Too many pieces should not be placed in the same stal.

6. Use only dean respects. It is well to let the object rest on a bit of extan in the bottom of the vial or have it suspended from the vial mouth so that the respect may penetrate equally from all eides. Penetration is aided by heat.

 When necessary to work fresh tissue, it is usually best to use normal adime, and not voter. Let it from gently over the surface of the object or aboutly twird the latter in the fluid. Do not scrape off foreign matter.

 In many cases the killing and fixing reagent does not harden the tissue sufficiently and the hardening process must be completed in alcohol.

9. Keep the reagents and preparations from direct sunlight.

10. Carefully label each ressel containing tissue. State the contents, the fuild used, and the date. Label on the side.

 Keep on cards a careful record of the reagents used, and the time when changed, for each separate piece of tissue.

PRACTICAL EXERCISE

Kill a frog or other small werebrate by placing it under a bell-jar which exotains a bit of exiton saturated with chloroform. Open the body as soon as possible after death and serure the tissues specified below.

 Alcohol Fizzion.—Remove the dorsal sorts and small pieces of the liver and harden in absolute alcohol (at least, not less than 95 per cent) in a vial or small bottle. The tissue will be ready for further treatment in two days.

Larger pieces of tissue require longer time. The pieces should be thin. Change the alcohol every day for the first three days.

Alcohol is in most instances an unsatisfactory fixing reagent, but it is frequently employed because it is usually at hand and is easily manipulated. Hot absolute alcohol is very often used for insects. Acetic acid (p. 207) is used with alcohol sometimes to increase penetration and to counteract its tendency to shrink tissues. The mixture is usually prefeasible to alcohol akone.

2. Fixing with Zenker's Fluid .-- Place small pieces of liver, kidney, pancreas, spleen, cardiac and pyloric ends of the stomach, bladder, spinal cord, and brain in about ninety times their bulk of Zenker's fluid. Remove a piece of the intestine about 12 mm. long, and after washing it thoroughly in normal saline place it in a vial of the fixing mixture. After fixation, which requires from 6 to 24 hours, wash the objects in running water for from 12 to 24 hours. Then transfer them through 35 and 50 per cent alcohol (20 minutes each) into 70 per cent alcohol. Add sufficient iodine solution to give the alcohol a port-wine color. The iodine will remove any mercuric crystals which may have formed in the tissues. As often as the color disappears the iodine must be renewed. After from 12 to 36 hours of this treatment, the color persists and the objects should then be transferred to fresh 70 or 80 per cent alcohol, which must be renewed until it no longer extracts iodine from the specimens. Too prolonged washing with iodine solution tends to undo the work of fixation and to hinder staining. Many workers prefer to omit the treatment with iodine until the tissue has been cut into sections. In such cases slides bearing sections are treated with dilute iodized

Killing and Fizing

alcohol for 30 minutes and then washed thoroughly in 70 per cent alcohol.

1

Zenker's is one of the best general fixing agents known. Any of a great variety of stains may be used after it, and it fixes satisfatorily almost any kind of tissue. The time during which objects should be left in the fluid varies from 30 minutes for delivete cores to 24 to 36 hours for larger or denser tissues, although many objects may be left a longer time without injury.

3. Fining with Bouir's Finid.—Place small pieces of trachen, tongue, comea, intestine, and testis or ovary in Bouin's finid for from 4 to 16 hours. After fination, wash the tissues in several changes of 50 per cent alcohol, then in 70 per cent alcohol multi the yellow color ceases to be discharged. Preserve in 70 or 80 per cent alcohol. Bouin's finid is an encellent reagent which gives a very delease fination. It is one of the safest for the beginner because it is shnost impossible to go wrong in its use. Objects may be left a considerable time in it without injury.

4. Formalin as a Fixing Reagent .-- Place a piece of spinal cord, sciatic nerve, liver, and fragments of muscle in which nerves terminate in 10 per cent formalin and leave until needed for work later. Formalin in varying percentages is widely used for the preservation and fixation of specimens for dissection. It has been employed especially for the central nervous system. However, Miss Helen D. King (Journal of Comparative Neurology, XXIII, No. 3 [August, 1913]), who has made a careful study of the effects of this fluid on the brain of the white rat, pronounces it unfit for cytological work because of its tendency to swell brain tissue. On the other hand, she finds that nerve tracts are apparently not affected adversely by it. If a formalin-fixed brain is to be used for histological purposes, she advises its transfer to alcohol as soon as it is fixed and hardened. Formalin ordinarily has a slightly acid reaction due to the presence of formic acid. Miss King finds such formalin less harmful than that which has been neutralized. For faithful preservation of cells, however, she prefers some other fixing fluid, such as Bouin's. For simple preservation, solutions ranging from 2 to 5 per cent are adequate, but for fixation the solution should be stronger (10 per cent).

Entire human brains may be fixed and hardened in a 10 per cent solution.

Formalin is much used for fixing and preserving when frozen sections are to be made, and it is particularly serviceable where a stury of fat is desired. It also interferes less with microchemical tests than most other respents.

MEMORANDA

 Tissues Are Preserved in Abcohol of from 70 to 85 per cent strength, but if they are to remain several months; it is better to preserve them in a mixture of equal parts of giverin, distilled water, and 95 per cent alcohol.

2. Hardening .- Read carefully the remarks on hardening in chap. ii.

3. Tissues Should Not Be Left in the Firing Agent longer, ordinarily, than is necessary to get results. Some, however, require a long time to bring out the optical differences of their elements. Experience alone can teach the time required in a given case. Such a reagent as formalin kills, fues, hardnes, and preserves, all at the same time. However, see remarks under 4, p. 29.

4. Firstion by Enjection is highly advantageous with many tissues beeause the fining finit is brought quickly into contact with all parts of the body. The vascular system is first washed out with normal sub solution and then illed with the trar. For finite contaming correstive sublimates a glass syringe and eaunth, instead of a metal one, should be used.

Hollow Organs should be filled with the fixing fluid and then suspended in a vessel of the same.

6. For Transferring Small Objects through respects the method of Walton is an excellent one. For the several reagents he uses shell rials which messure about 10 em. in beight by 3 em. in diameter. Tarough the center of a hat orek which fits the vials a hole is made and a glass tube (about 9 em. X1. about) is inserted so that its lower end dips well into the reagents in the vials. The lower end of the tube is closed with fits-messeled doth and the objects are placed within the tube. To transfer the objects one simply removes the oxic hearing the tube and inserts it in the vial containing the desired reagent. The upper end of the tube past hear of how we have a small hole should he howed in the side of the tube past how the hower level of the larger cork. The vials are supported as indicated in memorandum 7.

Very small objects (e.g., small eggs) may also be made up into little packets in lots of the east-off epidemis of the trog or salamander, according to the method of Professor Boyeri. The epidemial finit is spread over the concrety of a hollow ground side and saturated with adocted of the same strength as that which surrounds the objects. After the latter have been

Killing and Fixing

transferred by means of a pipelite to this sheet of epidemul tissue, two opposite edges of the sheet are folded over the objects, then the other two are brought together, twisted about each other, and pinned with a face pin. The pin, bearing a label, is used as a handle to transfer the little bag of objects from one reagent to another. Since the epidemul tissue outs readily, it need not be removed if the objects are to be imbedied for sectioning.

Professor C. E. McClung places fresh bits of tissue on small strips of paper with the proper index number on the opposite side. The paper is then inverted on the surface of the fixing fluid. The tissue will adhere to the paper through all subsequent processes. For washing, dehydiasting, etc, simply float the paper on the proper fluid. A number of objects may thus be handled together. The plane of section may also be marked by the paper.

Gelatin expandes, such as are used for methines, have been recommended for small objects by various workers. A hole, of sufficient size to let through the reagent but hold back the objects, is pricked in each end of the consule.

7. Shell Vials, Small Bottles, etc., when in use are best supported in shallow anger holes of proper size in thick blocks of wood.

8. Material Which Is to Be Kept Indefinitely should be put in fightly stoppered vials in a place array from strong light. Glass stoppes should be used, since each, besides shrinking and disintegrating with age, may give of extractives which injure deliate issues. It is best to pack the vials in a museum jar on cotton and then seal the jur securely to prevent evaporation. Materiali severa more secure if the museum jur is partly filled with alcoholy in such acuse each small vial should have a label of the content's placed within it. Another way to prevent evaporation from visits or bottles is to "ess"

Amount way to prevent evaporation from vizes of acousts is so exthem with a suitable varnish (see memorandum 9).

9. To Seal Bottles and Preparation Jars ("bottle-capping"), dip the stopper and part of the neck in collodion variable made as follows:

Pyroxylin (e.g., collodion or celloidin)	1 08.
Ether	6 02.
Altohol	8 oz.

When the pyroxylin has completely dissolved add 2.5 drams of camphor. (From Pharmaceutical Era, XXX, 528.)

10. For the Preservation of Anatomical Material for other than cytological or histological dissection, Professo George Wagner of our own laboratory finds nothing as serviceshile as Keller's third. The formula is as follows:

Formalin	1.	5 parts
Carbolic acid	2.	5 parts
Glycerin	10	parts
Water	86	parts

If a good grade of carbolic acid is used, disagreeable odors will largely be avoided. For embalming, a suitable reservoir is filled with the fluid and

suspended some 3 or 4 feet above the body to be preserved. About 6 feet of rabber toting, provided with clamps and a glass example of proper size to fit the femoral artery, is attached to the reservoir. After killing the aximal with illuminating gas or chlorodorm, the field is injected through the feanceal artery. An important pre-aution is to have the column of find in the table and example from air buddles or foreign materials. The pressure should be such at all times as to prevent blood from running lack through the example. The animal should be subjected to this embediming process for two hours. Obviously, by increasing the size of the reservoir and the number of tubes, several azimals may be treated at the same time.

At the conclusion of the embalaning process, if the arteries are to be injected with a colored mass, the rubber tube should be disconnected from the cannels and the latter be left in place in the artery. After the animal has remained for 2H hours in an unright position the hijection may be undertaken. For study of the blood vascular system, however, some workers prefer to inject hesh animals and preserve them in formalin. For a satisfactory starch injection mays see memoranoum 16, p. 42.

In our own laboratory it is the practice to skin the embalmed animal and wap the body in classes/ofth asturated with the embalming fluid. It is kept in a nice box of suitable size which has a close-fitting ful. We find a box 25 index long, 8 inches wile, and 9 inches deep a very good size for ests. For a more detailed account of embalming with Keller's fluid see Bensley, Practical Anatomy of the Rabbit, pp. 194-96.

The following mixture, recommended to the surface have the following mixture, recommended to the surface to realist results. To a mixture of equal parts of givenin and 65 per cent slooled sufficient formalin is solided to make the whole shorts a 2 per cent formalin. Specimens remain perfectly flexible in this mixture, and, indeed, after they have become thoroughly suimated, many forms (rustatese, inserts, etc.) may be removed and lopt as dry specimens which still retain their flexibility.

 Material Which Has Been in Formalin and is to be dissected may be reachered more plickle, and harmless to the skin, by seaking in a 3 per cent solution of earbolic soid.

12. For Weshing out Specimens it is arbitable to have the laboratory water-pipe provided with numerous small cocks about 10 cm. apart, so that each student may have the use of one or more. A piece of rubber tube, long enough to reach to the bottom of the vessel containing the specimen and fitted with a bit of glass tube in the free end, may be standard to each ordet. If the objects are very small they may be placed in performed in individual within may be purchased from instrument dealers. For minute objects the thindles may need to be lined with fine game. Bits of glass tubing with the game, but you go also useful for small objects.

CHAPTER IV

SIMPLE SECTION METHODS

FREE-HAND SECTION CUTTING

This method is important because it requires no costly appliances; although the sections are not as accurately cut as when mechanical aids are used, the method is simple, rapid, and adequate for the more general histological and tachalogical work.

 The section razor is flat on one side (the lower) and hollow ground on the other (Fig. 15). It must be sharp.

2. A shallow glass disk or watch-glass partly filled with water is also necessary. Before making a sertion, dip the ranor flatwise into the liquid, or use a camel's hair brush; see that the upper surface is well flooded.

 Sit in such a way that the forearm may be steadled against the edge of the table.

4. Use a piece of liver which was fixed in formalin, first mixing it in water. Take the tissue between the thanh and forefinger of the left hand, and hold it in such a way that a thin size may be cut by drawing the knife along the surface of the forefinger.

5. Rest the flat surface of the knile upon the forefinger, and, beginning at the heal of the knile, carcially draw the black toward you diagonally through the tissue, slicing off a thin section of as uniform thickness as possible.

 As each section is cut, float it off into the water; if it adheres to the blade, remove it by means of a wet camel's hair brush.

 Practice until very thin sections are obtained, then phase the dish upon a black surface, and with a needle or section-lifter transfer the thinnest and best sections, if only fragments, to a watch-glass containing water.

Norm—In case the tissue has been preserved in alcohol, out the sections under 70 per cent alcohol instead of water, then transfer them to 30 and 35 per cent alcohol successively and faully to water, leaving them in each liquid from 3 to 5 minutes.

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 The section razor is that on one side (the lower) and hollow ground on the other (Fig. 15). It must be sharp.

2. A shallow glass dish or match-glass partly filled with water is also necessary. Before making a section, dip the more factures into the liquid, or use a small's har bruch; see that the upper outline is well finded.

 Sit in such a way that the forearm may be steaded against the edge of the table.

4. Use a piece of liver which was fixed in formalin, first mixing it in water. Take the tissue between the thumb and forefinger of the left hand, and hold it in such a way that a thin give may be out by drawing the knife along the surface of the forefinger.

6. Rest the flat surface of the knille upon the foreinger, and, beginning at the heel of the knille, anticular during the head toward you diagonally through the tissue, diving off a thin section of as uniform thickness as possible.

 As each section is cut, float it off into the water; if it adhenes to the blade, remove it by means of a wet ramel's hair brush.

 Practice until very thin sections are obtained, then place the dish upon a black surface, and with a needle or section-liker transfer the phinnest and hest sections, if only fragments, to a watch-glass containing water.

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4. Use a piece of liver which was fixed in formalin, first in water. Take the tissue between the thrush and foreting left hand, and hold it in such a way that a thin slice may h drawing the kalle along the surface of the forefinger.

5. Rest the flat surface of the knife upon the foreing beginning at the heel of the knife, carefully draw the black you disgonally through the tissue, slicing off a thin seek uniform thickness as possible.

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 Practice until very thin sections are obtained, then p dish upon a black surface, and with a meedle or section-lifter the phinnest and best sections, if only frequents, to a wa containing water.

Nors.—In case the tisses has been powered in advalat, wit the under Toper end takked noted of water, then transfer them to 40 to exert alsohed properties and finally to water, bearing them in each if 8 to 5 minutes.

8. Next, place the sections in about 3 e.e. of Dekheld's hematoryin diluted with an equal volume of water, and leave them for various lengths of time (3, 7, 12, or more minutes) to determine the time for successful staining.

9. Transfer the sections from the stain to tap water, and gently more them about for from 5 to 10 minutes to wash out the excess of the stain. If the sections are still overstained, place them in 5 c.e. of distilled water to which 3 drops of acetic acid have been added. Leave for 5 minutes, or until they become lighter in color, then wash in several changes of tap water until they have again become blue.

 Remove the sections from the water and transfer them through 35, 30, 70, 53, and 95 per cent alcohol successively, leaving them from 3 to 5 minutes in each, and them transfer them to absolute alcohol for 10 minutes, and finally to xylol for 10 minutes, or until clear.

11. Select one or two of the best sections and transfer them to the center of a clean glass slide. After straightening them out properly, drain off the excess of the clearer and before the sections can become dry, add a drop of Canada balsam. Carefully lower a clean cover-glass (for cleaning see memorandum 14, p. 56) on to the balsam. These should be just sufficient balsam to spread evenly under the cover without examing around the edges.

 Label, stating card number, name of the preparation, and other data that it is desired to add (see p. 48, step 10).

 Carry one of the pieces of stomach prepared in Zenker through the same treatment. The sections should be transverse sections of the stomach wall.

14. Clean up all dirty glassware immediately.

MEMORANDA

 The Thinnest Sections are not always the best. For a general view of an organ, large, comparatively thick sections are usually better; for details of structure, thin sections.

2. Small Preses of Tissue may be emented to a orth if too small to bold conveniently between thumb and forefinger. A piece of stout copper wire is heated for a moment in the finne and touched to a bit of parsfirm. As the parsfirm mells, transfer drops of it to the edge of the tissue, which has

been previously placed on the cork. The paraffin cools and holds the tissne fast.

Another and better method of handling a small object is to imbed it in a piece of hardened liver. In sectioning, the liver as well as the object is sliced, but they readily separate when placed in alcohol. Beef liver or dog liver is prepared for such purposes by hardening pieces about 5×2×2 cm. in size in 95 per cent alcohol for 24 hours, and then transferring to fresh 95 per cent alcohol until needed. When much hand sectioning is to be done,

a supply of hardened liver should be kept on hand. Many small objects may be held between pieces of pith and successfully sectioned.



3. Well Microtomes (Fig. 27) are inexpensive instruments which are used for simple sectioning. Such a microtome consists of a tube in which the object is placed, and at one end of which is a plate to guide the razor. The other end is provided with a screw, which, when turned, pushes the contents of the tube above the plate, thus making it possible to cut sections of a uniform thickness. The object to be cut must be firmly fixed in the well. Such tissues as kidney, liver, spleen, hard tumors, cartilage, etc., may be beld sufficiently rigid by wedging small slabs of carrot, turnip, pith, or hardened liver in about them. These supporting substances must, of course, rest squarely against the bottom of the well. Soft tissues, such as soft tumors or brain, must be imbedded. Three parts of paraffin and one part of vaseline melted Pro. 27,-Well Microtome together and thoroughly mixed makes a very good

imbedding-mass for a well microtome. To imbed, warm the microtome slightly and fill the well with the imbedding-mixture. Remove all liquid from the surface of the tissue, and pass it below the surface of the mixture just as it begins to harden around the edges. When the imbedding-mass has become cold the sections are cut in the ordinary way.

4. Temporary Mounts may be made directly from water after staining by using glycerin as a mounting-medium. Transfer the section to the slide, add a drop or two of glycerin, and a clean cover-glass.

CHAPTER V

THE PARAFFIN METHOD: INFILTRATION AND SECTIONING

I. From 70 per cent alcohol take a small piece of intestine (6 mm. long) fixed in Zenker's fluid, and also pieces of other tissues fixed in this fluid, and proceed according to the following schedule. Keep accurate records on your cards.

2. Immerse in 95 per cent alcohol for 30 to 45 minutes. A longer time will do no harm.

 Transfer to absolute alcohol, 1 hour. Before transferring to absolute, remove the excess of 95 per cent alcohol from the object by touching it with a piece of blotting paper or a clean cloth, but do not lebit become dry.

4. Xylol and absolute alcohol equal parts, 30 minutes.

5. Xyld, 1¹/₂ hours, or until the object looks clear. Rapidly remore all excess of xylol before proceeding with step 5, but do not allow the tissue to become dry or dull looking. (For definite objects see p. 53.)

6. Melted pensifin (melting-point about 30°C), 2 hours. The object may be left an hour longer, but it is best to avoid as much as possible subjecting tissues to an elevated temperature. Shift its position in the parafin one or twice to facilitate penetration of the latter.

The duration of the parsfin bath varies according to the size and density of the object. Many objects of from 3 to 5 mm. in thickness are thoroughly saturated within an hour or less; others which are more impervious or which have impendable coverings may require several hours or even days. Lee, assuming that melted paraffin will penetrate as quickly as cold oil, takes as a guide the length of time required to clear the object in cedar oil.

 $C_{\rm AUTIONS,--a})$ Do not have the bath too hot. Cooked tissues are worse than useless,

The Paraffin Method

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b) To keep material clean, it is well to have a false bottom of paper in the vessel containing parafin. Make this by swinging a strip of white paper into the cup so that the loop of the paper is submerged in parafin and the ends attached on either side to the month of the cup.

7. Prepare paper boxes according to the following instructions:

A small restangular block of wood or a stick with a fast cad measuring approximately ISX20 mm. is used. Out a strip of still glaced paper so that it measures about 4X7 cm. Place the fast end of the block in the center of the paper with its long diameter coinciding with the long diameter of the paper. Fold the narrow side margins of the paper up slong the sides of the block first, then do likewise with the ends of the paper. Turn the easy which have been formed at each orner back over what is to be the end of the box, and then fold the long end of the paper back to hold the easy in place, and also to make the end of the box of the same height as the sides. Manifestly, any size of box may be made by raying the size of the block. With a little practice the same kind of lox may be folded without the use of a wooden block. See, however, memorandum 14, p. 45.

8. With a usrm, wide-mouthed pipette transfer sufficient melted parafin to a paper box to eover the bottom, then, with warm forceps, remove the tissue to the box. Next, fill the box with melted parafin. Orient the object with heated medles if necessary. As soon as the parafin has congealed sufficiently for the surface to become opaque, cool it rapidly by plunging it into cold water; otherwise the parafin will crystallize and become unsuited for sectioning. Many workers prefer alcohol instead of water for hardening the parafin block. Waste alcohol may be saved for this purpose.

Carrons.--a) Tissues must be oriented (i.e., placed in proper position for cetting) while the parafin is still in liquid condition. Arrange the tissue so that it will be cut at right angles (transverse) or parallel to the surface of the organ. Avoid oblique sections as they are very puzzling. For present purposes of practice cut transverse sections.

b) If whitisl-looking patches are present in the block after imbedding, they are probably due to xylol which has been carried over into the parafilm. If they occur in the immediate vicinity of the object, the block should be placed in the bath again until melted, and the object should be reimbedded.

c) Be sure that every piece of tissue is marked after it is inhedded. Tissues are sometimes kept in parsifin for months or even years before they are finally sectioned. To mark, sensith the number of the record card in the parsifin or, better, write it on the paper box and leave the box in place.

CUTTING SECTIONS

9. Study the parsfin microtome (e.g., Fig. 28 or Fig. 29); identify the parts and learn how the thickness of sections is controlled.

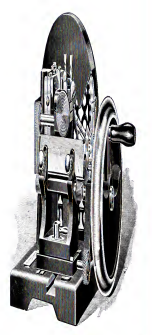


Fig. 28.-Minot Automatic Rotary Microtome

The object-senior is adjustable in three places and is periodry right. The leadscartier is also adjustable and errors have you object. The first is non-robot by an adjustoble can, gring error of any number of minerus in thickness from 1 to 25. By means of an automatically doing optication of minerule is neuronal to the leading position affer in server is full out the enrice leagth. The details of construction are enabled to the neuronal section of the section of the leading of the Barset & Londo Dyticel Oo.

10. Proceed with the block of parafin containing the intestine. Make it has to the carrying disk of the microtone in the following menner: Remove the disk from the machine and, by means of a

The Parafin Method

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heated steel spatula or copper wire flattened at one end, melt a small chip of paraffin on to it. Likewise warm the end of the paraffin block and quickly press it into the melted paraffin on the disk. Cement it finally in place by means of the heated wire or spatula and cool in water.

 With a sharp scalpel trim the free end of the block so that it presents a perfectly rectangular outline (however, see exution d). The length should exceed the breacht by at least on-fourth.



Fig. 29—Spencer Rotary Microtame By means of a wheel at the back the feed may be set for any thickness from 1 to 60 micross.

Caurnovs.—u) In trimming do not cut farther back than the base of the object. This leaves a wide shoulder for support.

b) Leave a margin of about 2 mm. around the object.

e) To avoid reversing sections in mounting, it is frequently advastageous to have the imbedding-mass trimmed unsymmetrically. The edge which first comes in contact with the knife is left longer than the opposite edge. One may thus readily discover when a section or part of a series has been turned over.

 Mount the object firmly in the microtome. It should just clear the knife. The fist end-surface of the parafin block should

be parallel to the edge of the knife, and the block so oriented that, in cutting, the long edge will meet the edge of the knife squarely.

13. Place the knife in position with the handle to the side away from the wheel (if a rotary microtome is used). By means of the adjusting screws tilt the cutting edge slightly toward the object so that the side of the knife will not remain in contact with the paraffin block after a section has been cut. If the knife has a flat under surface it requires more tilt than if the surface is hollow ground. For a flat under surface the tilt should be about 9 degrees from the perpendicular. See that the knife is held firmly in place.

CAUTION.-The knife should be kept in its case when not in the machine. The edge is very easily injured.

14. Set the regulator so that the microtome will out sections about 10 microns thick. A micron is one-thousandth of a mullimeter.

13. Unlose the eatch which locks the wheel and revolve the wheel with the right hand. A few revolutions should bring the block of parafin into contact with the knile. As each new section is cut, it displayes the last one and if the parafin is of the proper consistency units by one edge with the displaced section. Thus a ribbon or chain is formed. When the ribbon becomes of sufficient length, support the free end by means of a hair brush held in the left hand. To prevent breaking the ribbon avoid pulling it toot. Various ribbon-entries have been devised for attachment to the microtume. The best of these is the cylindrial carrier mode by the Spencer Lens Co. according to the suggestions of Dr. C. E. McChung, For an inexpensive, home-made form of this see Hance, Anotonioal Record, X, No. 8 (June, 1946).

Cavrons.—Never bring a needle or other hard object near the edge of the knife. If the parsfin does not ribbon properly, consult the table at the end of this chapter.

16. When a sufficient number of sections have been cut, carefully place the ribbon on a piece of paper. Protect it from droughts of air which will carry array or disarrange the sections.

17. Cut the ribbon into strips of such length that they may be placed in successive rows one above the other under the cover-glass The Parafin Method

that is to be used. Allow one-fourth for the expansion of the ribbon when heated. Mark out on a sheet of paper the exact size of the coverglass so that there can be no mistake in cutting strips of the proper length. A margin of 2 or 3 mm, should be allowed for the cover.

18. Place a small drop of albumen fixative on a clean glass slide (for cleaning see memorandum 14, p. 156), and spread it evenly over the surface, encept the end which is to bear the label (see step 10, p. 49). With a clean finger rub off all of the fixative that can be easily removed so that only a very thin film remains.

19. Flood the slide with a few drops of distilled or alterminized (p. 23) water until the entire surface bearing the fixative is covered by a thin layer of water, but do not put on sufficient to overflow the edge. Some workers use alterminized water alone for affixing sections.

20. Take up the first strip of parafin ribbon with a brush or needle and float it on to the surface of the water. The first section of the series should be in the upper left-hand corner, but back at least 10 mm. from the end of the slide. In case the label is to he placed on the left end of the slide, allowance must be made for it, of course. Add the successive strips of the ribbon in the order of the lines of a printed page until as many rows are in place as will conveniently he under the cover (see step 17, p. 40), allowing for the proper margins. See that each section presents the same aspect to the observer as its predicessor (see step 11, c, p. 39).

21. Warm the slide genity until the parafin fastens out and becomes free from winkles. Be earcful not to melt the parafin, for heat sufficient to do so will render the albumen useless. It is safer to heat the slide by placing it upon the warm parafin oven for a few minutes, instead of bolding it above a flame.

22. Drain off the excess of water and set the slide away to dry after properly numbering it with your glass-narking penell. As the water eraporates, the sections are drawn down tightly into the film of fixative. If, after drying, air is present under the sections it may be seen from the glass side of the slide. The slide is seldom sufficiently dried under 6 hours. It is well to leave it 12 hours; it

may be left indefinitely. The time may be shortened by placing a few thicknesses of blotting paper under the slide and drying it on the paraffin oven or in an incubator. Unless the slide is perfectly dry and the rikbon fully spread, the sections will float off during subsequent treatment. Take precautions to prevent particles of dirt from settling upon the surface of the sections. This is usually accomplished by placing the kildes upon some kind of a rack and covering them with a bell-jar. Prepare sevenal other slides in the same manner as the above if sufficient of the ribbon remains.

Norm—As time permits, out the other sections which are imbedded in parafin. When, as in the present case, it is not necessary to have a complete series of sections, you may place lever sections on a slike and use smaller covers.

Wen a small over is to be used, place the sections at the entire of the side. The center may readily be determined by drawing the orifine of a slide on a card and connecting the opposite corners of the figure by means of diagonal lines. When morning, place a slide over the diagonal; the intersection of the diagonals shows the center.

At this point the student should make a careful study of Appendix A if he is not already tharroughly acquainted with the optical principles involved in microworg.

MEMORANDA

 If Parafin Becomes Dirty it should be melted and filtered through a heated metal funnel.

2. Oil of Cedar, if used for dealookolisation before indexiding, should be followed by at least two changes of parafin are the parafin does not throughly replace the oil and the object is likely to drop out of the sections as they are cut. In my experience this is the commonest difficulty which beginness encounter if they use cedar oil for dealookolination. For this reason ayiel or chlorotom is recommended as prefeable for general work.

 Objects Imbedded in Parafin may be preserved in that form indefnitely. It is one of the most convenient ways, in fact, of preserving material which is to be sectioned in parafin.

4. Small White Objects, if not stained before inhedding, cloudd be tinged with a dilute solution of Congo red to facilitate orientation. For orientation in general see p. 126, memorandum 1.

5. With Delicate Tissues it is necessary that the transition from also had to clearer be gradinal, hence it is best to add the clearer, a little at a time, to the last alcohol, transferring it with a pipette to the bottom of the alcohol. See also "throp" method, pp. 18 and 152.

The Paraffin Method

 The Temperature of the Laboratory must be taken into account when sectioning in parafilm. In summer use a barder, in winter a softer, parafilm.
 For Thin Sections use a hard parafilm, for thick sections, a softer

parafin.

8. For Valuable Tissues Which Crumble in Parafin Alone the following somewhat tellows process (Mark, American Naturalid (1985), p. 628) may be resorted to. Prepare a very fluid collocine in ether-alcohol and coast the exposed surface of the object immediately before entiting each sertion. If the collocine laws a skiny surface or produces a membrane when applied to the parafin, it is not this enough and must be further diluted with ether-alcohol. Apply the collocin with a brush with all excess of the block in which the objects moved. After applying, with a few some of the block in which the objects sequenced. After applying, with a few some of the block in which the objects sequenced. After applying, with a few some only for the solution to due before entiting. See also memorandum 9.

9. Johnson's Pentfin-AsphaleRubber Method for brittle objects is a very useful one. One part of errole India rubber exist into very small pieces is nixed with 90 parts of hard parafin which has previously been methed and targed to a light amber odor with a small amount of asphale ("mineal mither"). The mixture is then subjected to a temperature of 100° C. (not higher) for 24 to 48 hours, or left in a parafin over at 60° C. for several days. Use only the superastant third. It is allowed to cool and remain cold until mended, because the rubber separates out after a time? It hemisture continues methed. Johnson (Journal of Applied Microsopy, VI, 2002) recommends it as even better than parafin for all kinds of work for which parafin is commonly employed. Proceed as in the ordinary method, using xyld (not cedar cold hash for classing for the soften and sky for classing a vertice.)

 Keep All Parts of the Microtome clean and well olled with watch of or pure parafilm oil of 25° C. The instrument should be covered when not in use.

Keep the Microtome Knife Sharp. It should receive frequent stroppings. For sharpening the knife two hones are commonly used.

Horing—If the knife is very dull it is first honed on a Belgian yellow hone, an open-grained stone which cuts the metal of the knife rapidly. The surface of the stone is kept maint with filtered kersene oil or hthered with palm-oil scop. After the nicks and other inequalities of the edge of the knife have been removed, the honing is best finished on a good finegrained blue-water stone.

In homing, the stone is kild flat on the table with its end toward the operator and its surface properly lubricated. A very dull kilde is ground at first on the concarre side only until it develops a fine "where edge" along the full length of the blade. It is then ground on each side alternately until the wire edge has disappeared completely. In graving, the kulte must remain flat on the lone and pass lightly over the full length of the

surface, edge foremost in a diagonal direction from point to heel, although itself remaining at right angles to the long axis of the hone. The honing has been sufficient when all nicks and wire edges have disappeared and the knife, instead of eatching and hanging when the edge is drawn lightly across the ball of the thumb, freely enters the moist epidermis. Finally the blade is wiped clean with a soft cloth, great care being taken not to injure the edge.

Some workers prefer to use first a yellow Belgian hone wetted with a moderately thick scop solution, then an Arkansas stone with a thin oil. Hardesty (Laboratory Guide for Histology, p. 196) uses only a white Arkansas stope with oil.

Stropping .- A broad firm strop of finest calfskin is best. It should be affixed to a solid back so that it will not spring and thus round off the delicate edge of the knife.

In stropping, the motions are the same as in honing (both sides of blade), only the knife passes back foremost and from heel to point. The blade must move lightly over the surface of the strop with very slight pressure on the part of the operator. The stropping is ordinarily considered sufficient when the blade will cut a loose hair freely along every part of the edge. An examination under a low power of the microscope should reveal no nicks in the edge.

12. To Remove Pigments and to Bleach Osmic and Chromic Acid Materials, a 3 per cent solution of peroxide of hydrogen frequently is suff-



cient. Tissues left too long in this liquid macerate.

Mayer's chlorine method is one of the best for bleaching. To several crystals of chlorate of potash in a glass tube a few drops of hydrothloric acid is added. When the greenish fumes of chlorine appear, add from 5 to 10 c.c. of 50 per

Pic. 30.-Metal Us for molding cent alcohol. The object, which in the imbedding-masses. meantime has been standing in 70 per

cent alcohol, is transferred to the tube. From 15 minutes to 24 hours are required for bleaching, depending upon the nature of the material. It is well to suspend the object from the mouth of the bottle. Sections on the slide may be bleached in a few minutes. This method is especially recommended for removing natural pigments and for bleaching osmic material.

13. Large Objects May Be Cut in Paraffin better with a slanting knife than with a square-set one. The block of paraffin must be trimmed to a three-sided prism with its most acute angle farthest from the object. A sliding microtome is used ordinarily and the block of paraffin is so oriented that the knife enters at the sharpest angle of the prism. Each section as cut is removed with a brosh.

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14. Metal or Powelain "US" Are Preparently Used Instead of Paper Bones for molding resultin blocks. The two US (Fig. 30) may be placed together on a small gloss or metal plate in such a way as to mold blocks of any desired size. Bolice pouning the methed parafilm in, the inner walls of the metal pieces should be helded smarred with glycenia so that the block of parafilm will easily separate from them when cool. Many workers, particularly for small objects, indexi in a watch-gloss and harden under alcohol.

DIFFICULTIES LIKELY TO BE ENCOUNTERED IN SECTIONING IN PARAFFIN, AND THE PROBABLE REMEDY

 Crocked Rhbons—o) Usually enseed by wedge-shaped sections. Correct by trimming the block of pacufia so that the edge which strikes the half first and the edge on the opposite side are strictly paculled. See that the block strikes the half evently at right angles.

b) The parafin may be softer at one end of the block than at the other. This can be corrected only by imbebling the object over again in a homogeneous parafin.

 The Object Makes a Scratching Noise on the Knife or Cuts with a Gritty Feeling and the sections perhaps erunable and tear out from the parafilm.

a) This is generally caused by too high heating of the object while in the parafin over. Not only is such an object worthless, but it endangers the edge of the microtone hulfe. Correct by limiting the bath in parafin to the minimum time necessary for a proper presentain of the object and keeping the temperature barely above the melting-point of the parafin.

b) The fixing reagent has formed crystals (e.g., corresive sublimate) which have not been thoroughly washed out.

See also 5 (p. 46).

3. The Sections Winkle or Jam Together; the object itself may be compressed before the knile. This is a serious fault because the arrangement of the parts of a tissue is greatly decanged. It may be due to various causes.

 a) The microtome knife may be dull. Examine the knife and sharpen it if necessary.

b) The paraffin may be too soft. To remedy this defect employ one or more of the following means: (1) could be parafin block in water; (2) out the sections in a couler room; (3) out the sections thicker; (4) embed in harder paraffin. See also menorandum 11, p. 47. If sections are not too hadly winkled they may be flattened out by warming on water as directed in steps 19-21, p. 41.

c) A possible reason is that the tilt of the knife is insufficient (see step 13, p. 40).

d) The edge of the knife may be smeared with a layer of paraffin. Clean the edge with a cloth moistened in xylol.

4. The Sections Roll and Refuse to Röbon.—This is one of the most enspecting of all deletes. If the sections are not tightly curled, they inequality unroll when placed on warm water (step 19, p. 41). Various mechanical devices have been constructed to prevent this eril, but most of them are impractical. Sometimes when a section begins to roll, if the edge is held down by means of a flat-pointed hair brush the euring can be overcome. If a ribion can once be started, the difficulty is frequently corrected. The sections should be out rapidly.

c) The commenset cause of rolling is the hardness of the parafin. This may sometimes he remedial by one or more of the following means: (1) warming the kile with the breath; (2) enting in a warmer room; (3) placing a lamp or burner near the imbedded object; (4) warming the kulle way conclully by holding the back on a warm parafin bath; (5) enting the sections thinner; (6) reimbedding the object in softer parafin; (1) dipping the end of the block in melted softer parafin.

b) The tilt of the knife may be too great (step 13, p. 40).

c) The knife may be dull.

 The Sections Split Longitudinally or Are Crossed by Parallel Scratches.---o) Look for a nick in the edge of the knife. Out in a new place on the knife or sharpen it.

b) A bit of git may have gotten into the object or the parafilm, or there may be a hole in the parafilm. Reinibed after carefully cleaning the object in the cleaning fluid.

c) Tissues may contain hard substances (line subs, silina, crystals preipitated from fixing reagents) which have been imperfectly washed out. It is best to take an entirely new piece of tissue in which these delets do not exist.

d) The tilt of the knife may be too great (step 13, p. 40), or the trouble may be due to loose parafin on the edge of the knife.

e) The object may be too large to cut in parafin. Try smaller pieces of tissue or use the celloidin method.

6. The Knife Scrapes or Rings as It Passes Back over the object after having out a section.

 a) This is sentetimes caused by a kinle with either too great or too little till (step 13, p. 40).

b) The object may be too tough or hard to cut in paraffin without springing the edge of the knife (see 7, b, p. 47).

c) The blade of the knife may be too thin.

 The Sections Vary in Thickness; the machine cuts one thick and one thin or misses a section.

a) This may be caused by the imperfect mechanical construction of the machine. Old machines in which the parts are worn are especially liable The Paraffin Method

to this defect. It may be remeibed to some extent by tightening up the parts of the machine.

b) The object may be too hard for the knife to out and, as a consequence, the edge of the knife springs. When tough or hard objects must be ent, use an old microtome knife or a seriforing mater. See if there is not some means of softening such a tissue without obscuring the microscopical structures sought.

c) Either too great or too little tilt may cause the defect (step 13, p. 40).

d) See that the disk bearing the object is securely clamped in the machine.

8. The Object Crumbles or Drops Out of the Parafin as Cut-1: has probably been insufficiently presentions. Some of the following presentions may prevent the defect: (1) Lave the object in the parafin bath longer. (2) See that it is entirely tree from the dealoohining fluid before placing it into the methed parafin. (Djoets which have been immersed in cedar oil are particularly subject to this defect. For this reason sylal is better than eachs cill for dealoohining in general work. (3) If the object is imperious to parafin or very friable, as are many ora, some other method must be tried. Consult memoranda 8 and 9; see also the celloidin method (memorandrus 8, p. 64).

9. The Ribbon Twists or Carls About or Clings Closely to the Side of the Knite.—This is due to the electrification of sections. If the half is encossive, it is best to postpose the cutting until the atmospheric exactlinos have elanged. The difficulty may be minimized by using a drum ribbon-sarrier (see step 15, p. 40).

10. The Cut Section Catches On and Clings to the Block as it returns instead of remaining on the lande. Probably the laude is dull or its edge is durty; the tilt (step 13, p. 40) is insufficient; the pendfin is too soft, or the room temperature is too high.

11. A Simple Coder for use with the microtome, which facilitates the preparation of thin passifin sertions and is especially useful in a laboratory where the temperature is high, is described by Grave and Glaser in the Biological Bulletin for September, 1910. The apparatus 'is essentially a loalow transated pyramid, open at both ends, and suspended in an inverted position from a standard, so adjusted that the lower end of the short is at a convenient distance above the krite. At the upper read of the short is at a convenient distance above the krite. At the upper end of the short is at a convenient distance above the krite. At the upper end of the short is at a convenient distance above the krite to the escape from the lower end of the ain-channel. At that point a rubber tube connexts the pipe with a suitable receptack." Grave and Glaser recommend the following as a convenient size, base, 12-508-7 in, transated aper, 6-1X-2 in, ; measurements of in-text, 8-503-3 in.

CHAPTER VI

THE PARAFFIN METHOD: STAINING AND MOUNTING

I. STAINING WITH HEMATOXYLIN

Place enough of the following reagents in tall stender dishes or Coglin staining-jars to cover the slides lengthnize, up beyond the sections affixed to them: xylol, absolute; 93, 70, 50, 35 per cent alcohols respectively, clear water, acid alcohol, and, for washing out the acid alcohol in the case of hematoxylin preparations, a separate jar of 70 per cent alcohol to which a few drops of a 0.1 per cent aqueous solution of biearformate of soda has been added. Arrange these reagents in a row in the order named with the exception of the acid alcohol and its accompanying alkaline alcohol wash of 70 per cent alcohol, which may be placed immediately back of the ordinary 70 per cent alcohol. Put a little vaseline along the upper edges of the jars containing alsolute alcohol and xylol and press the cover down tightly to prevent eraporation or the entrance of maisture.

In like nanner place in Coplin staining jars (tail steaders will answer) a supply of Delafield's hematorylin diluted one-half with distilled water, easin, Lyous blar, alum-orchineal, Congo red, and solutions A and B for the iron-hematorylin method. Arrange these stains in a row hock of the alcohol series.

 Remove the parsfin from the sections of intestine (see last lessin) by placing the slides in xylol (tarpentine will answer) for 10 or 15 minutes. The process may be hastened by first gently warming the slide until the parsfin begins to melt.

 Remove the xylol from the sections by transferring the slides to absolute alcohol for 1 minute.

 Pass the slides through the alcohols (95, 70, 50, and 35 per cent), leaving them for a half-minute in each.

 Remove to Delsfield's hematoxylin for 10 to 30 minutes or until stained a pronounced blue.

5. Wash in water for 5 minutes.

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 Pass the slides up through the series of alcohols to 70 per cent, leaving them about half a minute in each alcohol.

7. Dip each slide for from 30 seconds to 5 minutes into the acid alcohol until the sections are of a reddish hue, then rinse them in 70 per cent alkaline alcohol until the blue color is restored. This last alcohol must be kept very slightly alkaline through the occasional addition of a few drops of a 0.1 per cent solution of bicarbonate of coda (see memorandum 10, p. 55). The alkaline alcohol may be omitted when other than hematoxylin stains are used, as its purpose is merely to restore the blue color of the latter.

 Pass the shfess through 95 per cent alcohol (3 minutes), absolute alcohol (5 minutes), into xylol for 10 minutes or until clear (see memorandum 3, p. 55).

9. Carefully drain off all excess of the charter from a slide, wipe the under side, and lay it down fast with the sections uppermost. Put a few dops of thin halsam on the sections near one end. Take up a clean cover-glass and, holding it by the edges between the thumb and first finger of one hand, lower it upon the balsam by bringing one end into contact with the slide near the balsam and supporting the other end by means of a needle held in the free hand. Lower the cover slowly so that as the balsam spreads, no air bubbles will be inclosed under the glass. If a slide is tilted a little and allowed to remain in that position small bubbles will frequently work out unaided. They may sometimes be removed by pressing gently above them with the handle of a needle and gradually working them to the edge of the cover-glass. Keep the slide in a boinontal position until the balsam harders.

CAUTION .-- Do not allow the sections to become dry before adding the balsam and cover.

10. Attach the permanent label. It should contain at least the following data: the number of the record card (p. 5); the name of the tissue; the kind of section (plane of section, thickness, etc.); if one of a series, the number of the slide in the series and the number of the first and last section on the slide; the date, and if desired the name or the initials of the preparator. It is well to add the thickness of the cover-glass (see also memorandum 23, p. 58).

It is best to have the label on the left end of the slide, as it will then not be in position to obscure the scale of the detachable mechanical stage so widely used on microscopes today.

Norn—Prepare four shiles each of the other objects which have been imbedded. Skin and mount one of each kind as you did the intestine, and also one of each kind in the same way, only substitute a lam-sochined for the beamtoryin. The slam-sochined may require 12 hours or more for staining. Preserve the others for double staining.

As time permits, prepare and section the other tissues which were fixed in alsohol and in Bouin's fluid. After you have had the preliminary practice in double staining, stain and mount these as you prefer.

II. DOUBLE STAINING IN HEMATOXYLIN AND EOSIN

 Proceed according to the regular schedule with one each of the slides reserved above, and stain in Delafield's hematoxylin.

2. Wash the sections in water, and proceed farther according to the regular schedule to 95 per cent alcohol.

 Transfer the slide to the cosin stain for 30 to 60 seconds, and after rinsing again in 95 per cent alcohol, place it in absolute alcohol.
 Clear in xylol and mount in balsam.

Norm.—The sections should show both the blue stain (in nuclei) and the red stain (in cytoplasm) when examined under the microscope. If either is too dense or too light, make a note of the last and vary the time accordingly when staining other sections by this method.

III. DOUBLE STAINING IN COCHINEAL AND LYONS BLUE

 Pass the remaining reserved slides through xylol and the alcohols, descending to 35 per cent alcohol.

 Stain in alum-cochineal for from 6 to 12 hours, or until the sections are well colored.

3. Rinse in water or 35 per cent alcohol, and pass the sections up through the alcohols to 95 per cent. If the sections are deeply stained, however, remove the excess of stain with acid alcohol (a few seconds) when the sections are in 70 per cent alcohol.

 Stain for 10 to 20 seconds in Lyons blue. It is very easy to overstain with this dye.

5. Rinse in 95 per cent alcohol, and transfer the sections to absolute alcohol (5 minutes), clear in xylol, and mount in balsam.

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IV. STAINING WITH HEIDENHAIN'S IRON-HEMATOXYLIN

This stain is very valuable in the study of cell division and in determining the finer structure of the nucleus. The iron-alum acts as a mordant, werearing the tissue for the action of the hematoxylin.

 Prepare two sets of sections of intestine, testis or ovary, bladder, pancreas, and stomach. The sections should not be over 6 or 7 microus in thickness. Preserve one set for double staining.

Pass the other set through xylol, absolute alcohol, 95 per cent alcohol, and thence directly into water.

 Transfer from water to the iron-alum, and allow this solution to act for from 30 minutes to 1 hour.

4. Rinse in water 5 minutes.

 Stain in the 0.5 per cent benatorylin 1 hour. If a trace of the iron-alum remains in the sections the hematorylin will turn black. This, however, does not impair its power of staining.

6. Rinse in water 5 minutes.

7. Place the sections into iron-alum again, which will now extract the excess of stain. The time required for proper differentiation varies with the kind of tissue and the hing agent that has been used. From 10 to 30 minutes is usually sufficient, though no definite time limit can be set. Remove the side from the non-alum from time to time and inspect it. When the sections become of a dull-grayish hue the decolorization is usually sufficient. If very accurate results are necessary, the slide should be removed from the inor-alum frequently, rinsed in water, and examined under the microscope. When in a dividing cell the chromesomes become sharply defined, the decolorization should be stopped.

 Wash in running water for 20 minutes or in several changes of water for 2 hours. If any of the inn-alum is left in the sections the color will inde later.

 Wipe off the excess of water, transfer the slide to 95 per cent alcohol, followed by absolute alcohol and xylol.

10, Mount in balsam.

Norm—hen-henstoxylin is perhaps the one most important stain in use today. The student should practice the method until he has mastered it. It is better though not absolutely essential that the stain be "ripe."

For demonstration of extrasources and finer systellogical details, the time of staining may require to be lengthered. In my own systellogical work I find that 2 to 4 hours in invo-stam followed, after rinsing in water, by hemotoryin for 12 to 24 hours, yields better results than does the shorter method.

V. IRON-HEMATOXYLIN WITH OTHER STAINS

Use the sections which were reserved for this method. The method is identical with the one just outlined, except that between step 8 and step 9 the following directions should be inserted: 80, transfer the sections from water to Congo red for a minute, or to orange G for 2 hours, then wash them in water and proceed to step 9.

Norz.-Before proceeding farther, kill a female out or robbit to secure tissues for the colloidin method and to correct failures in the parafin method. In addition to the tissues specified before, prepare (fiz in Zenker's fluid) pieces of tendom, cartilase, spleen, lumph gland, pancress, and sativary glands. (If the respents are at head and time permits, the student, indeed, might advantageously prepare a number of tissues according to the methods indicated in Appendix C.) Fix the every in Bosin's fluid, and reserve it for the parafin method for delivate objects. Fiz parts of the brain and cord in Zeaker's and in formalin as previously indicated, and place bits of muscle in which nerves terminate plentifully (e.g., intercostate) in formalin. Larger pieces (up to 2 cm.) may be used of such tissues as are to be imbedded in celloidin. Bour in mind that the larger the fiscue the larger must it be left in the different reagents. Select the necessary parts of the digestive tract to prepare longitudinal sections in celloidin from esophagus to stomach and from stomach to intestine. As soon as possible begin the preliminary steps in the celloidin method (chap, vii) so that there may be no loss of time. Prepare a piece of intestine for staining in balk (see vi). It should be placed in the stain after thoroughly weshing out the fizing reagent. Preserve parts of it to cut in celloidin. Remore the lower jaw and prepare it for decalcification of teeth (as indicated in chap. xi). Likewise prepare pieces of femar and of tarsal bone for sectioning (chap. xi).

VI. STAINING IN BULK BEFORE SECTIONING

It is sometimes desirable to stain objects before sectioning. The method is a slow one, and requires stains which penetrate evenly and thoroughly. Various preparations of earnine and orchineal give the best satisfaction, although several hematoxylin stains are also frequently used in this way. It is best to stain immediately after fixing and witsing out, before the object has been earried into higher alcohols. In general, it is advisable to section tissues and

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stain on the slides, because the staining can be controlled more effectually. Use the piece of intestine already prepared (see note, p. 52).

 After fixing in Zenker's fluid and thoroughly weshing according to the directions on p. 28, place the tissue in Dekafield's hematoxylin for 24 hours.

 Wash in 35 per cent alcohol for 5 minutes, followed by 50 and 70 per cent alcohol, 20 minutes each.

 Decolorize in acid alcohol 20 to 30 minutes, or until the color eases to come away freely. Restore the bluish-purple tint by treatment with alkaline alcohol.

4. From this point proceed through 95 per cent alcohol, absolute alcohol, xylol, and imbedding, sectioning, and moanting precisely as in the general parafilm method, except that after the sections have been freed from parafilm in xylol, do not mount immediately in balsam, but first transfer the silde back into absolute alcohol, and thoroughly wash it in order to remove the glyverin from the finative and so prevent cloudiness of the final mount. From alcohol the silde is passed through xylol, or carbol-xylol, and mountied in the usual way.

Norz.-When exists of the carnines or hematoxylins are used as stains for estire objects, the preparations usually need to be decolorized with acid alcohol. This may be deferred, however, until after the objects are sectioned.

VII. PARAFFIN METHOD FOR DELICATE OBJECTS

To prevent the distortion of delieste objects which are to be sectioned in parafin, the transition of the material from one reagent to the other must be very gradual and the heat he minimized. Observe the following modifications of the general method and prepare pieces of ovary which have been fixed in Bouin's fluid.

 Pass the object in the usual manner up through the series of alcohols to absolute. It is sometimes necessary to use a more closely graded series of alcohols if the object be very definite. See also the "droy" method (memorandum 6, p. 152).

 From the absolute alcohol pass to a mixture of absolute alcohol two-thirds and chloroform one-third; gradually add more chloroform until at the end of an hour the mixture is at least two-thirds chloroform.

3. Transfer to pure chloroform for 30 minutes.

 Add melted parsfin little by little during the course of an hour or two (24 hours will do no harm), until the chloroform will hold no more in solution.

 Transfer the object to pure melted parafin in a small vessel on the parafin oven for 10 to 20 minutes, changing the parafin once. Imboti in the usual way.

6. Cut the sections about 7 micrors thick. Mount and stain some in Delafield's hematoxylin and cosin and others in ironhematoxylin and Congo red or orange G, according to the directions already given for these methods.

Norm—For very sensitive objects Schult's delytizing apparatus (to be chained from dealers) may be used. It consists of a tube within a tube, each having the lower cal covered by an animal membrane. The tubes are suspected in the seek of a mark henger bottle which contains 95 per cent alcohol. The object is placed in the inner tube and host tubes are filled with water. When suspended in the alcohol, a very gradeal hardening or delyticities of the object tubes place as the alcohol skowy diffuses through the membrane. Smeetings it is necessary to use only one tube, and in such a case the hardening proceeds more rapidly.

VIII, EUPARAL AS A MOUNTING- AND PRESERVATION-MEDIUM

This reagent, introduced by Gilson, has been highly extelled by various workers as a final mounting-medium, although most still prefer belsam or danax for general work. Two forms, colorless and green, are obtainable from Grühker. The green is used only with hematorylin stains, which it intensifies. One of the merits of expanal is that delicate tissues may be mounted in it directly from 95 per cent should, thus arveiding the expense and the risks of passage through absolute alsohol and the essential oil. Another value lies in the fact that, because of its lower index of refraction (1.433), unstained or faintly stained elements which are invisible in balsam (index, 1.335) are rendered visible. It thus becomes particularly serviceable in the study of spindle fibers and other such delicate cytological elements. It hardress rapidly, hence preparations can be used within 24 hours. It is well for the beginner to try it along with other methods.

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MEMORANDA

 In Passing from One Liquid to Another, one corner of the side being sections should first be touched by blotting paper to remove any encess of the liquid last used. This is expectably necessary in transferring from absolute alcohol to xylol, or from 50 per cent to alcohole alcohol.

2. Sections Once Placed in Turpentine or Xylol for the removal of parafin must merer in any subsequent step be allowed to become dry. Particular case must be taken to prevent sections from drying out after removing them from xylol to mount in balsam because the xylol ersopentes rapidly. If the sections become dry the preparation is usually rendered valueless.

 Xylol Used for Removing Parafin should be kept in a jar separate from that which contains xylol for elearing before mounting, and it should be changed occasionally because it tends to become saturated with parafin.

4. Sections Not over to Micross Thick may be plunged directly from 95 per cent alcohol into an opports melium and vice versa. It sections are over 10 micross thick it is better to put them through the complete series of alcohols. With thick sections diffusion is less rapid, and too alrupt a change from one fluid to another may produce distortions or wrench the sections loose from the fluide.

5. To Avoid Rubbing Sections of the Slide, hold the slide with one end toward the light before wiping it and game obliquely along the surface. The shiny side is the one to wipe.

6. The Series of Alcohols and Stains colinarily may be used a number of times without replenishing. When the alcohols become very much discohored or the stains cloudy, they should be reneved. Alcohols should not be used too often, however, as they soon accumulate particles of dirt which settle upon the sections and render preparations unsightly.

7. Absolute Alsohol must be kept free from water. It may be tested from time to time by mixing a few drops with a little targentine. If the mixture appears milky the alcohol contains a harmful smount of water and should be reserved.

8. Two Sildes Placed Back to Back can be handled as realily as a single shide in passing through the various liquids. Various kinds of sild-backets or slid-backets, which enable one to transfer a number of sildes through liquids at one time, may be obtained from dealers.

 Gentle Agitation of a Slide in any liquid facilitates the action of the Iquid. Observe this precaution especially with absolute alcohol.

10. For Washing Sections after Staining in Hematorylin tap water is preferable to distilled water because it is usually sliphtly alkaline. When and alcohol is used to devolutive sections stained in hematorylin, the sections should be washed in 70 per cent alcohol rendered alkaline by the addition of a lew drops of 0.1 per cent solution of binarborate of sola. The alkali of a lew drops of 0.1 per cent solution of binarborate of sola. The alkali neutralizes the acid and restores the bluish-purple color to the section; it also renders the blue color more permanent. If too much of the soda is added the color will be a hary disagreeable blue.

11. To Obtain a More Precise Stain with Delafield's hometoxylin, it is well to dilute it with three or four times its bulk of distilled water. The sections must be left in this solution a correspondingly longer time. Sections stained in this way may not require treatment with acid alcohol. Must workers, however, prefer to overstain and decolorize.

12. The Length of Time Required for Staining Different Tissues is exceedingly variable. Upon removal from the stain after rinsing, if the sections are inselficiently colored, put them back into the stain and examine them from time to time unfil they are properly stained (30 minutes to 24 hours).

13. H Objects Refuse to Skin, it is usually due to one of the following causes: (a) The fixing agent has not been sufficiently washed out. This is a frequent cause of poor staining. (b) The fixing has been poor. The success of a preparation depends largely upon proper function in most cases. (c) The stain is at fault. Hematoxylin will not stain properly until ripe (see "Hematoxylin"); n.9). Many stains especially the anilins, deteriorate and must be replaced. (d) Certain stains will not follow some fixing agents. This can be remedial only by using a different stain or by fixing fixesses in a different fluid. The hematoxylins and carmines are applicable after a very large variety of fixing agents. (e) The parafin has been insufficiently removed from the sections. This may be corrected by disselving off the corrected by disselving off the corrected by disselving and and, after tharoughly removing all parafin, restaining and mounting the sections gain in the ordinary vay.

14. Use Only Clean Slides and Covers --Movys grasp a slide or a cover by its edges to avoid scaling its surface. All cloudiness (seen by looking through the glass toward scane dark object) must be removed. For wriging slides and covers, a piece of club which does not readily form lint should be used. A well-masted linen towel is good, as is also bleached cheesedout out into pieces the size of a handkerchief. Slides may often be cleaned after simply dipping them into alcohol or into alcohol followed by water. If this treatment is insufficient, place them for several hours into equal parts of hydrochloric acid and 85 per cent alcohol, keying them well separated so that the liquid may act on the entire surface of each. Then rines them in water and place them in ether alcohol. It is well to keep a stock supply of such slides and cover glasses in ether alcohol.

To clean a coverglass, grasp it by the edges in one hand, cover the thumb and first finger of the other hand with the cleaning-lobb, and rub both surfaces of the glass at the same time. To avoid breaking the cover, keep the thumb and finger each directly opposite the other. A large coverThe Paraffin Method

glass may be cleaned by rubbing it between two flat blocks which have been wrapped with cleaning-cloths.

To clean sides which have been used, if balsam mounts, warm and place in sylal or targentine to dissolve off the overs. Put the sides and covers into separate gass or porcelain vessels and leave them for a lew days in the following cleaning mixture:

Potassium bichromate	10 parts
Hot water	50 parts
Sulpharie seid	50 parts

In making, sold the soid very eautionaly after the biohomete solution cools. When the shifes are freed from balsam, wash them in veter, rince in a ditute solution of eautite socia, again in water, and finally place them in ether-alcolud until mechel.

15. If Sections Appear Milky or Hary under a melium power of the microscope, when finally mounted, the effect is probably due to one of the following sauses: (a) The clearer is poor and needs replenishing or correcting. (b) The absolute about contains water (see memorandum 7, p. 55). (c) The over hore moisture. Passing a covar-glass quickly through a finane before putting it on to the object will remove moisture. (a) The add has not been entirely removed from the sections. (c) Too much allowen hastive has been used. (f) The giverin of the allowmen hustive has not been removed by passing sections of object stained in bulk (see vi, 4, p. 53) hack into absolute about allow paradin from them.

The defect may be remelied frequently by disadving off the over in rylal or turpentine, descending through the series of reagents to the point where the fault lies, correcting and ascending again according to the regular method. To remove water, for example, it is only necessary to go back as for as absolute alcohol which has a great afinity for water.

16. Dry or Dull-Looking Areas under the Cover-Glass indicate that the sections were allowed to get dry after the removal from the clearer, or that insufficient talksam was applied.

 Delicate Black Pins in sections of tissue fixed with a mercuric fixer are due to deposition of mercury. Remove with iodized alcohol before staining.

18. Balsam Which Endes from under the Cover may be scaped off with an old kulle after it knowns. Bemore the last taxes by means of a brish or a doth dipped in turpetitine or xylol. Balsam may be removed from the surface of a cover by means of a kreach dipped in xylol.

 If Sections Wash off the Slide the defect is probably due to one of the following eccess: (a) The slide was solled or oily. Remely by chaning slides theroughly (see 14, p. 36). (b) The albumen fixative is too old.

(c) The transitions in the alcohol have been too great. This is true sometimes of thick sections. (d) The paraffin ribbon was not thoroughly spread when mounted (see p. 41).

Thick sections are more likely to come off the slide than thin ones. To avoid this, the sections may be collodionized by placing them, after the parafin has been removed, in a thin solution of collection or celloidin in ether-alcohol (? gram in 100 c.c. of ether-alcohol; see 4, p. 8) for a few minutes and then transferring them to 70 per cent alcohol. If carmine dves are to be used, this method is not satisfactory, as carmine stains collection.

20. Flooding Sections with the Dye by means of a pipette, especially in case of stains which act rapidly (e.g., cosin, acid fuchsin, Lyons blue, picite acid, etc.), is sometimes more convenient than immersing the sections in a jar of the staining fluid. Small bottles with combination rubber stopper and pipette (Fig. 31) are now provided for this purpose by dealers.

21. Balsam Mounts in Which the Stain Has Faded may frequently be restained, either with the original or with other stains. All that is necessary is to dissolve off the cover in xylol (2 to 3 days) and pass the preparation down through the alcohols to the stain in the usual manner.

22. Ink for Writing on Glass (Hubbert, Journal of Applied Microscopy, V, 1680) .- Mix drop by drop 3 parts of a 13 per cent alcoholic solution of shellac with 5 parts of a 13 per cent Fig. 31 aqueous solution of borax. If a presipitate forms, heat the solu-Dropping-bonde tion until it clears. Add enough methylen blue to color the mass deep blue.

Professor Robert F. Griggs uses common water-glass (an aqueous solution of sodium silicate or potassium silicate) with an ordinary steel pen. After marking, the slide is heated until the water-glass decomposes, leaving behind a rough, sandy surface, which when rubbed away shows the written characters etched on the slide.

"Diamond ink" obtainable from Eimer & Amend is useful for writing on glass. When not in use it is kept sealed with paraffin. See also p. 1, "Carborandam Points."

23. More Detailed Labeling than that indicated on p. 49 is sometimes desirable. For well-devised schemes see Richard E. Seammon, "A Method of Recording Embryological Material," Kanans University Science Bulletin, IV, No. 5 (March, 1997); also Robert T. Hance, "A System for Recording Cytological Material, Slides and Locations on the Slides," Transactions of the American Microscopical Society, XXXV, No. 1 (January, 1916).

24. For Orientation of Objects in the Imbedding-Mass, see p. 126.

CHAPTER VII

THE CELLOIDIN METHOD

Use the tissues which were prepared (p. 52) for this method, including pieces of the brain and spinal cord which were fixed in Zenker's fluid. Reserve a piece of spleen for the freezing method (p. 67).

 Fixing, washing, and dehydrating are the same as usual (chap.ii). If the object is in 70 per cent alcohol, complete the dehydration by using successively 95 per cent and absolute alcohol. It should remain in the absolute alcohol for from 12 to 24 hours.

 From absolute alcohol transfer the object to equal parts of absolute alcohol and ether 12 to 24 hours.

 Next to thin celloidin (reagent 20, p. 12) for from 36 hours to several days or weeks.

 Thence to thick celloidin for from 24 hours to several days. It may be left for weeks.

Thorough delaydration and thorough infiltration are the great essentials for success with the celloidin method. Some workers prefer to use a graded series of celloidins, such as 11, 3, 6, and 8 per cent, leaving the object in each for from 24 hours to several days or even weeks.

5. Prepare a woolen block (see memorandum 3, p. 63) in such a manner that it will have surface enough to accommodate the object, leaving a small margin, and length enough to be readily clamped into the earlier on the microtome (Fig. 32). Dip the end of the block to which the object is to be stracted into ether-alcohol for a minute and then into thick celloidin. Let it dry so that later air bubbles will not work up out of the wood into the imbedding mass.

6. Oil one side of a strip of stiff paper by rubbing on a very little vaseline, and wrap it, olled surface in, about the prepared end of the block in such a way that it will project beyond the end of the block, forming a collar high enough to extend a little beyond the object



in volaption for collect or parama cutture. These as A-Automate Collectin Alterations in work of a transfer of the second second and a second and a second and a second as a second second second as a the second second second again when the second form is been for our a second with a formation as a second second the second second again when the second form is been for our a second way for unreased in a second as the second

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which is to be placed within it. The the paper in place by means of a thread.

7. Pour a small amount of thick celloidin into the paper cap thus formed and with forceps remove the piece of tissue and place it in celloidin. Add more thick celloidin until the cup is full. By means of needles which have been moistened in other-alcohol arrange the object so that it will be cut in the desired plane.

Norm-Instead of being mounted in blocks, objects may, after saturation with thick edition, be placed in proper position in a glues disk and covered with thick edition. The dish is then lossely covered and placed under a bell-jur so that the other will gradually eraporate, heaving the mass, in a day or two, of proper consistency for certiting. Each specimen is then cut out in a block of suitable size for sectiming. For fastening to base, see memoratolm 5, p. 68.

8. Into a small stender dish put chloroform to the depth of 3 mm. When a film has formed over the exposed surface of the celloidin place it in the chloroform to harden. It need not be submerged. Keep the vessel tightly covered. The object may be left for a day or two, but 1 to 3 hours usually suffices.

9. Transfer the block to 70-83 per cent alcohol, where it may remain indefinitely.

10. Make a careful study of the microtome used for cutting celloidin (Fig. 32).

11. Place the block in the object-carrier of the mirrotome at the proper level and arrange the microtome knile obliquely, so that it will slive through the object with a long drawing cut for at least half the length of the blade. If the object is oblong it is advantageous to have the long diameter parallel to the edge of the knile.

 Keep both the knife and the object flooded with 70 per cent alcohol, preferably from an overhanging drop-bottle.

 Draw the knife through the object with a straight steady pull; avoid pulling down on or lifting the knife-carrier.

14. If the feed is not automatic push the knife back to position always before turning the screw which raises the object. Out the sections about 15 or 20 micross thick. If they end they are best unrolled on the surface of the knife with a camel's hair brush just before they are wholly cut free from the block.

15. As the sections are cut, transfer them by means of a small soft brush or a paper spatial to a flat stender or a watch-glass containing 70 per cent alcohol.

 Transfer some of the sections through 50 and 35 per cent alcohol, 2 minutes each, into alum-coeffined for from 20 to 30 minutes, or until stained (12 to 24 hours).

17. Wash successively in 33, 50, and 70 per cent alcohols, leaving the sections from 2 to 3 minutes in each.

 Transfer the sections to 95 per cent alsohol for 3 to 5 minutes. Absolute alsohol is not to be used with celloidin becaue it dissolves the celloidin.

19. Clear in eedar oil or beechwood crossote for from 10 to 20 minutes.

20. Mount in balsam (see step 9, p. 49).

STAINING CELLOIDIN SECTIONS IN HEMATOXYLIN AND EOSIN

The objects are killed, fixed, and preserved as usual in 70 per cent alcohol, and sectioned as in the foregoing method.

1. Fifty and 35 per cent alcohol, each 3 to 5 minutes.

2. Delafield's hematoxylin, 20 to 30 minutes, or until stained.

3. Water, 5 minutes.

4. Thirty-five, 50, and 70 per cent alcohol, each 3 to 5 minutes.

 Acid alcohol, until the celloidin which surrounds the object shows but little of the stain.

 Seventy per cent alcohol, barely alkaline (see memorandum 10, p. 55), until the red color caused by the acid is replaced by bluish purple.

7. Alcoholic eosin, 30 seconds to 1 minute.

 Ninety-five per cent alcohol, 2 to 5 minutes. Clear in cedar of or beechwood cressote and mount in balsam.

Norm.—As time permits, section other tissues by the celloidin method and stain as above.

MEMORANDA

 H Chloroform Is Not at Hand, 80 per cent alcohol will harden the celloidin, although more slowly.

The Length of Time that objects should be left in ether-alcohol and the celloidin minitures depends upon the size and density of the objects. When time permits, it is always best to leave them several days or even weeks in the mixtures of cellolin. For large objects such as the modulla of a large brain this is a necessity. For an embryo of large size months may be required.

3. Blocks for Celledin Mounting may be of white pine, gloss, whennied fiber, or even a very hard parafin. Cark should not be used because it is liable to give or bend. The vulnamized fiber is the most satisfactory. It may be purchased from dealers in the form of strips which may easily be sawn to the necessary dimension. It is well to saw sevent) parallel cuts into the upper edge of the block to revoide points of statement for the reliability.

4. Other Cleares may be substituted for order oil or creasive. One which clears from 95 per cent and which does not dissolve cellular must be chosen. Other good cleares are: (1) origanum oil; (2) a minture of 3 parts of all of thyme and 1 part of cestor oil; (3) Eveloshymer's clearing fluid, which is a mixture of equal parts of berganot oil, celar oil, and anhydrous earbolic adil.

5. Imbedding a Number of Objects in one mass is frequently covernient. Fold a stiff paper into a box of the proper size (step 7, p. 37) or use metal 1's (Fig. 30). Pour in thick celloidin, put the objects in place, and orient them properly for exiting. Leave a space of about 5 mm between adjacent objects. Full the box with thick celloidin and set it in a disk outstaining a little chloroform, or leave it in 30 per cent alcohol to harden. When ready to proceed, cut the large blocks into smaller cores each containing a piece of tissue. To fasten it to the wood, thin the small celloidin block to the proper dimensions, suften for a few minutes in ether-alcohol the side to be attached, then dip it into thick celloidin and apply to the end of a wooden block which likewise has been dipped into the ether-alcohol and the think celloidin. Press the two together and place them in chloroform or 80 per cent alcohol to backen (see also note under step 7, p. 61).

6. Amin Dyes are usually avoided in the edition method because they stain the celloidin intensely and are not removed in subsequent treatment. When necessary, however, some (e.g., eosin) may be used. Subtanin, for example, may be removed satisfactorily from the celloidin by means of acid alcohal without extracting all the stain from the tissue. If acilin dyes have been used, it is sometimes better to remove the editoilin by treating the setting with absolute alcohol or with ether before the final charing and mounting (see memorandum 15, p. 66).

7. Relative Merits of the Parafin and the Celoidin Methods.—Celloidin is good for large objects, for Initite or fraible objects, and for delivate objects which heat would injure. It does not require zenoral from the tissues ordinarily, hence it holds delivate structures together permanently. Some tissues are not rendered so hard and so difficult to ent as in parafin. Horever, very thin sections exampt be obtained except by great skill. The

method, moreover, is extremely slow. The peculith method is comparatively rapid, serial sections may be cut and mounted with ease, and very thin sections may be obtained. Large objects do not section as subdistenity as in colloidin, although up to 10 mm. or even considerably greater diameter they ext readily. The rule is to use the paradin method when you can.

8. For Britle Objects, a Combination of Califoldin and Paraffin Intitration semetimes proves successful (see, however, menurandum 9, p. 43). The method is too tedious for ordinary use, although it must sometimes be resorted to with finishe or delivate objects such as eggs.

According to Apathy's method, as reported by Koralauser (*Keinea*, July 14, 1916), fixed material is dehydrated as usual, finally passing through three changes of absolute alcohol into e ther and alcohol, where it is left 5 hours. It is next put into 2 per cent celloidin for 24 hours, then 4 per cent celloidin for 24 hours, and ultimately imbedded in 4 per cent celloidin and hardened in chlorotom vapor for 12 hours. The block is then quickly trimmed, leaving a margin beyond the object of a few millimeters on every sile, and put into liquid chlorotom for 12 hours. It is next transferred to an oil mixture mole up by weight instead of volume as follows:

Chloroform	4 parts
Origanum oil	4 parts
Cedar-wood oil	4 parts
Absolute alcohol	1 part
Carbolic-aeid crystals	1 part

Anhydrous sodium sulphate should be kept in the bottom of the tube to take up any water brought in, in the celloidin.

The block must remain in this oil mixture until it clears and sinks; this may take from 3 days to a week. It is next washed in three or more changes of benul to remove oils and slookel, then influctued in parafin, imbedded, sectioned, and mounted in the usual way. In subsequent handling, slobes should not be left for any great length of time in alsolute alcohol, as it will dissolve out the celloidin.

9. To Transfer Celloidin Sections from the Knife, it is an excellent plan to use a paper spatzle; a bit of postal eard held in the defit end of a small stick answers very well. Press the paper down evenly on the section and then shife it of the edge of the knife. The section adheres to the paper. In carrying losse sections from one fluid to another an ordinary section-lifter may be used, or a glass red around which the section is allowed to earl answers very well.

10. Objects Stained in Bulk May Be Cleared While Yet in the Block, then sectioned, and mounted without passing back into the alcohols. After the block of celloidin has hardened sufficiently in chlorothern it is transferred directly to the clearer (redar oil, or a mixture of oil of thyme 3 parts and easter oil 1 part). In certing objects trace cleared the kmile must be theoded with the clearer instead of alrohol. Do not allow the sections to become dry. If it is desired to use this method for a celloifin block which has already been preserved in 70 to 83 per cent alcohod, the block must pass through 45 per cent alcohol (1 to 2 hours) before it is placed in the clearer.

 Calladian instead of Calladian, is used by some workers. Celladian, in fact, is a patent preparation of collodian, which is a solution of guarcettom (pyroxylin) in other and strong alcolol. Thin and thick solutions are employed and the method is in every respect similar to the reladian method.

12. Fining Celoidin Sections to the Slide is accomplished (1) by evvening the sections, when mounted in proper order, with a strip of tissue paper, which is then bound fast by wrapping thread around it. Lee (Microtonia's Vade-Merow, 7th ed., p. 125) recommends (2) the allounen method for each of the sections as well as for parafilm. (3) If the sections on the slide are easefully flooled with 95 per cent alcohol two or times times, this chained off and followed by a small amount of ether-alcohol or ether times until the edges of the sections begin to soften perceptibly (10 to 20 seconds), the sections will generally adhere to the slide sufficiently when the celloid heremas hard again upon exposure to the slide sufficiently when the related alcohol has been drained off; they must then be immessed in 95 per cent alcohol hefure are further stores are taken.

13. For Serial Sections in Celloidin some one of the so-called ploting or ober methods will be found most satisfactory. That of Linstacht (Andonical Record, November, 1922) is among the best: It is a modification of the celloidin sheet method suggested by Hibber for parafin sections and of Weigert's method for serial sections in cellulin.

Plates of glass, thoroughly clean and of suitable size (e.g., 5X7 inches), are coated on one side with the following solution and allowed to dry:

Saothartee.	3 grams
Dertrin	3 grams
Distilled water	100 c.c.
To which as a preservative is added a bit of th	iymel.

When the sugary layer is thoroughly dry, cost it with a 4 per cent solution of celluloid in sectione. A number of such sheets may be made up and kept dried if desired.

As settions are cut, place them in the desired order on the celluloid sheets, meistening from time to time to prevent drying out. Friling may be prevented by brief treatment with alsolute alcohol. When the plate is filled with sections, blot with a smooth-surfaced tailet-paper, then, in order to far the sections to the celluloid, spary by means of an atomizer with a 1 to 2

per cent solution of celloutin in ether and skohol. When it is partially dry immerse the plate in 70 per cent skohol, then in water will disadre the sugary solution and the celluloid sheet bearing the sections will fost off. The sheet may be preserved indefinitely in 70 or 60 per cent alcohol, or stained, elsared, and mounted at any time by any of the methods suitable for oximary cellulin sections. Strips may be cut and mounted secilly on properly numbered slides.

For another "Sheet" method see Böhm, Davidoff, and Huber, A Textbook of Histology, 1910, p. 40.

14. Gilson's Rapid Celloidin Process (Lee, Tie Microtonicle Yade-Meum, 7th ed., p. 112) is valuable under some circumstances because of the great saving of time. After delyrisation the object is saturated with ether and finally placed into a test-tube containing thin redshiftin. The lower and of the tube is then dipped into melled parafin and allowed to remain there mult the cellulin solution has bouled down to about one-third of its original volume. The mass is then mounted in the ordinary way, hardened for an hour or more in chloroform, and eleared in celar oil. Sections are ent as directed under memorandum 10 above.

 Celloidin May Be Removed from Sections when necessary by passing them through absolute skohol into ether and alcohol or oil of cloves for from 5 to 10 minutes, then back through absolute to ordinary alcohol.

16. For Orientation of Objects in the Celloidin Mass see p. 126.

CHAPTER VIII

THE FREEZING METHOD

 Use a piece of sphere which has been properly fixed and later preserved in 70 per cent alcohol. Transfer it through 50 and 35 per cent alcohol successively to water, and wash it for 12 hours in running water.

2. Plose it into a gum and syrup mass for 24 hours (a saturated solution of loaf sugar in 30 e.c. of distilled water, added to 50 c.e. of gum muckage. Prepare a supply of gum muckage by dissolving 60 grans of best gum acada in 80 c.e. of distilled water).

3. Examine the freezing microtome carefully (Fig. 33).

4. Bemore the gum and syrup mixture from the outside of the tissue with a cloth, put a little gum muckage (not gum and syrup) on the freeing disk of the microtome, and place the tissue in it in such a way that longitudinal sections through the hilum may be out. Surround the object with gum muckage and set the freeing apparatus to guing. If earlon dioxide is used, open the valve very cautiously, and let only a small quantity of the gas escape.

Norm—Curbon dimitle is commonly used for charging solar water and beer. It may be purchased in iron optimales containing about 20 pounds of the logadied gas. The optimale, when empty, is exchanged for a charged one, so that the purchaser pays only for the curters. The Roudeen microtome (Fig. 33) may be servered directly upon the carbon-dimitle optimale when the latter is in a horizontal position, or, if desired, the optimale may be placed vertically and the microtome attached by means of an 1-shaped piece of heavy taking. This microtome has the schwartage over the common forms of freezing microtomes of wasting less gas and of greater freedom from obgring. It is schwartageous to have an extra lung handle to the key which is used for opening the escape valve of the exchan-discule optimiler.

Small tubes of compressed earlier discribe, sufficient for one or two freezings, may sometimes be obtained from stores earrying automobile supplies. These can be utilized in operations where immediate diagnosis by means of sections is required.

 As soon as the gum is frozen, continue to add more until the tissue is completely covered and frozen.

6. Work the microtome screw with one hand and plane off sections (15 to 20 microns thick) with the other. The well-sharpened blade of a carpenter's plane is the best instrument for cutting. It must be frequently stropped.

Norra-The blade should be mounted in a short, broad handle, which may be grasped easily and firmly with one hand. In cutting, the bevel edge of the knife should set squarely on the glass ways of the microtome so that the handle



of the knife is inclined toward the operator at about an angle of 45 degrees from the perpendicular. The hand guiding the knife should be firmly supported against the chest while pressing the outting edge steadily against the glass ways of the microtome. The cutting stroke is made by bending the body forward from the waist and thus forcing the blade squarely across the surface of the tissue.

The blade must be kept cold to prevent sections from sticking to it. If the sections fly off or roll, the tissue is probably frozen too hard. The same defect may arise if there is insufficient syrup in the gum with which the tissue has been saturated. To correct, let the tissue thaw a little, and if it is still at fault,

F10, 33.—Banicen Carbon-Diaxide Freezing Microtome

The freezing chamber contains a feed is 20 microns.

soak it again in a mixture which contains a greater proportion of spiral passage through which the er-spiral passage through which the er-the maximum freezing power. The hubbility og gaus gause gause gauses. The facest Several sections may be allowed to

collect on the blade before they need be removed.

7. Transfer the sections to distilled water. The water should be changed several times to dissolve out the gum. Reserve a few sections in water for later use (step 11, p. 69).

8. Immerse a few of the sections for 10 to 30 minutes, or until stained, in Delafield's hematoxylin, then wash them in several changes of tap water.

The Freezing Method

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9. Transfer the sections through the successive grades of alcohol (decohoring with acid alcohol if necessary) up to absolute alcohol, leaving them 2 minutes in each, after which remove them to xylol for 5 minutes or until clear. If desired, stain with easin (30 to 60 seconds) after 70 per cent alcohol.

 Remove one or two of the best to a slide, drain off the excess of xylel, add a lew drops of balsam and a cover-glues of suitable size, and label.



Fig. 34.—Ether or Rhigolene Preezing Attachment

11. Remore the sections reserved in step 7 (p. 68) to a test-tube containing a small amount of water, and shake the test-tube vigorously for a minute or two. This removes the lymphocytes from the sections, and exposes the refordar connective tissue so that it may be examined. Dehydrate the sections and mount in holisam.

MEMORANDA

 Presh Tissues Are Preparently Sectioned by the freezing method. The tissue may be transferred directly to the disk of a microtome without previous imbedding, and sectioned after freezing. This affords a ready means

of rapidly determining the nature of a given tissue, and is very servicedale, especially to the pathologist. The principal objection is that erystals of ice form in the cells and distart them badly. This is avoided when syrup and gem are used for imbedding.

2. Fired Tissues may be sectioned by the freezing method if they are first washed in running water for some hours. Even tissue first in tormalin perhaps the best fixing fluid for tissues which are to be frozen—is better for a washing of at least 30 minutes before being frozen.

3. Sections May Be Preserved in alcohol in the usual way after being out by the freeing method. All trace of gum should be washed out and the sections passed through the guides of alcohol to 63 per cent, where they may remain indefinitely.

4. Sections of Fresh Tissue May Be Fired and vashel out after cutting if desired. This requires but little time, and the sections will take stain much more satisfactorily after having been subjected to a fining reagent.

5. Objects Which Alcohol Would Injure may be sectioned by the freezing method and mounted in aqueous media.

6. Ether or Rhigulene Is Sometimes Used for Freezing, although the method is more expressive and less satisfactory on the whole than the embondioxide method. Fig. 34 shows a common form of freezing attachment used for either of these liquids.

Organs or Parts Varying Greatly in Density may sometimes be out more successfully by the freezing than by any other infiltration method.

8. To Fin Frozen Sections to the Side, after treatment with absolute alcohol, flow a little three-fourths of 1 per cent solution of celloidin in ether and alcohol over the sections and drain off at once. After a few seconds of exposure to the sin place in 80 per cent alcohol for a minute. The film of celloidin should be very thin. If it turns white upon immersing in the alcohol, the original solution of celloidin was too thick; it should be thinned by adding more other and alcohol.

CHAPTER IX

METALLIC SUBSTANCES FOR COLOR DIFFERENTIATION

I. A GOLGI METHOD FOR NERVE CELLS AND THEIR RAMIFICATIONS

The Golgi chron-silver method is one widely used for the demonstration of nerve cells together with their various processes. There are many modifications of the method, all of which are more or less inconstant in their results. In a successful preparation the various cells and nerve processes are not equally blackened, a fact which allows of discrimination between the different elements. Sometimes the ganglion cells and fibers remain unstained while the neurogfla cells are impregnated, or occusionally other elements than nervous tissue (e.g., blood vessels) are affected.

The following method is applied to material preserved in 10 per cent formalin and is a so-called "rapid method."

1. From the brain and spinal cord, which have previously (see note, p. 52) been subdivided and placed in at least 10 times their volume of 10 per cent formalin (3 days to an indefinite time), cert out small pieces 4 to 5 mm. thick from the region desired for study and transfer them to a vessel containing from 15 to 20 times their volume of a 3.5 per cent acproxe solution of prtassium bickromate. They should remain in this solution for from 2 to 5 days. Renew the fluid at the end of 12 hours. Keep the different pieces of tissue in separate vessels so as to avoid confusion.

2. For impregnation, transfer the tissues to a silver-nitrate solution made as follows:

Silver nitrate (erystals)	1.5	grams	
Distilled water	200	C.C.	
Concentrated formic acid.	1	drop	

3. Rock the tissues gently in a small amount of this fluid until the brown precipitate of silver chromate ceases to appear, then

transfer them into from 20 to 40 times their bulk of fresh silvernitate solution and leave them in the dark for from 3 to 6 days. Change the fluid after the first 12 hours.

4. Transfer a few of the brown pieces of tissue to 95 per cent sloohol for half an hour, renewing it once or twice during this time. Leave the rest of the tissue in the silver-ainsite solution for future use in case the first attempt proves unsuccessful.

 Remove the pieves from 95 per cent to absolute alcohol for 20 minutes, changing the latter once. Then transfer them to etheralcohol for 20 minutes.

 Imbed in celloidin without waiting for infiltration to occur (thin celloidin 30 minutes, thick celloidin 10 minutes). Mount directly on a block and harden in ellocoform for 20 minutes.

7. From chlorotorn transfer directly to the clearing third (e.g., ecdar oil), and as soon as clear (30 to 60 minutes) cut sections 50 to 100 mirorns thick, but keep the knife flooded with the clearing fluid instead of alcohol. Cut sections of cortex so that they will be perpendicular to the surface of the brain.

8. When the sections are throughly cleared, transfer them to a slide flocided with the clearing fluid, select such as prove desirable upon microscopic inspection, and discard the remainder.

 Replace the oil with xylol, then remove the xylol by pressing upon the sections with blotting paper. Add enough thick Canada halsan to cover the sections.

Carriex.—Do not put on a over-glass; moisture must eraporate from the section. If this is prevented, the metal deposits break up and the sections become worthless.

10. Keep the preparations level and put them away in a dry place free from dust. If the bolsam runs off the sections, more balsam nurst be added at once. Do not attempt to examine under a high power until the balsam is thoroughly hardened.

MEMORANDA

 A Fuller Account of the Golgi Methods will be found in Hardesty's Neurological Technique (pp. 55-61), or in Lee's Microtomia's Vade-Mecum (pp. 419-57).

Metallic Substances for Color Differentiation 73

2. An Osmium-Bichromate Minture is frequently used instead of formain for fixing fresh tissues. To 85 parts of a 3.5 per cent solution of potassium hielenmate add 15 parts of a 1 per cent solution of comize avid. Small pixes (4 to 6 nm. thick) of fresh tissue are placed in 40 times their volume of this mixture and kept in the dark for from 12 to 24 hours. This fixing fluid is then replaced by a 3.5 per cent solution of potassium hielenmate, as in the case of material funct in formalin (see above). From this point the method is identical with the one given above.

3. The Determination of the Elements That Will Be Imprepated appears to depend upon the length of time the tissue is left in the 3. Spec cent solution of potassium bichromate. Hardesty gives the following lengths of time for different structures: neurogita, 2 to 3 days; contial cells, 3 to 4 days; Purkinje cells, spinal cord, peripheral ganglina cells, 4 to 5 days; nerre fibers of the spinal cord, 5 to 7 days. Avones are impregnated onlinanity only in so far as they are not michilated.

4. Mounting the Sections upon a Cover-Glass is preferred by some workness. The cover-slip is then fastened over the opening of a perforsited slide with the section downward.

5. For Permanently Mounting Golgi Preparations under a Cover-Glass Huber recommends the following method: The sections are removed from xylol to the shike and the xylol then removed by pressing blotting paper over the sections. A large drop of xylol-laskum is then quickly aspliced and the shike is carefully heated over a fame from 3 to 5 munths. A large coverglass is warmed and put in place before the taksun cods.

6. The Cor Modification of Golg's correstive-sublimate method is widely used. It is likely to impregrate nearly all of the cells in the section. This may prove to be disadvantageous rather than otherwise, however, where cells are numerous and close together. Small pieces of nervous tissue are placed for from 1 month in summer to 2 or 3 months in winter in the following solution:

The later treatment is the same as for ordinary Golgi preparations.

 Trachese of Insects, Bile Capillaries, and Gland Durts may also be studied by the Galgi chrom-silver method. A bit of the wing muscle of a bundle bee is a good object in which to demonstrate the facer ramifestions of trachese.

IL OTHER SILVER-NITRATE METHODS

a) For Nerves (after Hardesty)

 The fresh nerve, or better, a spinal nerveroot, may be dotained from a freg which has just been killed. Without stretching the nerve, carefully insert beneath it the end of a strip of postal eard or similar eard which has been trimmed to the width of 50 mm. The nerve when cut off at each side of the eard will adhee to it and remain straight and at approximately normal tension.

 Clip off the end of the eard bearing the nerve into a clean vial which contains 0.75 per cent aqueous solution of silver nitrate. Place the vial in the dark for from 12 to 24 hours.

 Transfer the nerve to puze glycerin on a slide and tease the fibers apart thoroughly under the dissecting microscope.

 Add a cover-glass and expose the fibers to sunlight until they become brown (30 minutes).

5. To make the preparation permanent, take off the ovver and remove the glycenin by means of filter paper, add a few drops of warm glycenin-jelly (n. 96), put on a clean ovver-glass, and press it down. Wipe away the exoded jelly, and when the preparation has could seal the ovver with gold size, followed by Bell's cement (see steps 5 and 6, p. 95).

The preparation should show the "cross of Ranvier" and the "lines of Fronman."

b) For the Cornea

 Quickly rub a piece of silver nitrate over the comes of an eye which has been removed from a recently killed frog.

 Slice off the cornea and place in distilled water. Brush the surface with a camel's bair brush to remove the epithelium (conjunctivum).

 Expose to the action of sunlight or strong daylight until the tissue turns brown.

 Wash in distilled water and mount in glycerin, or mount in belsam, after proper dehydration.

If the preparation is successful the cells should be strongly outlined by the precipitated silver. If desired, after washing,

Metallic Substances for Color Differentiation

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the nuclei of such cells may be stained in hematoxylin according to the usual method.

MEMORANDA

1. Fresh Membranes are also commonly treated with silver nitrate to outline cells. The membrane should be stretched over some smooth surface, or, better, by means of two small vubanite rings which fit one into the other in such a way as to stretche bits of membrane files a durn-head, and hold them fast. Such stretched membrane is first washed with distilled water, then agitated in a 1.300 silver-nitrate solution, in direct sullight, until it dacknes. It is then washed in distilled water, then agitated in a 1.300 silver-nitrate solution, in direct sullight, until it dacknes. It is then washed in distilled water, removed from the rings, and mounted in giverin, giverin-jelly (a. 90), or, after dehydration, in balanm. II preferred, after washing it may be stained in benatorylin to krine out the myels, and then mounted.

2. Cajal's Method for Neurofitriks is widely used. Small pieces of nervous fissue are fixed in formalin for 6 hours, washed in water 4 hours, and transferred to 40 per cent alcohol for 6 hours. They are next kept for 24 hours in 40 per cent alcohol to which annorms has been added in the proportion of 5 drops of annomism hybrite to 50 e.e. of the alcohol. The tissues are then placed in an inclustor in a 1.5 per cent sliver-nitrate solution and kept for 5 drops at temperature of 38° C. Next they are placed in a mixture of 100 parts of water, 15 parts of formalin, and 1 part of programs and they are placed in a mixture of hybridinous for 24 hours, after which they are ready to be passed through graded alcohols into parafin or celloùlin and sertioned in the usual way.

An excellent application of the Cajal method to serial sections has been derived by Malone (Lontonical Room), IX [1915], 701), who also describes how to obtain satisfactory Cajal preparations from sections previously stained by the Nesl method.

3. The Pyridne-Silver Method, a modification of the Cajd method, deviced as a differential stain for non-methallatel nerve fibers, has come into wide use in American laboratories in the study of various other problems. It is aften used in the preparation of sectors of spinal gaugia, sympathetic gaugia, and spinal cord and is the most reliable of the silver stains. Relatively large pieces of tissue can be successfully stained. Ranson's technique (American Journal of Antony, XII [1911], (09) is as follows: "The nerve or gauginon is placed in 100 per cent alcohol, with 1 per cent annuonis for 45 hours (95 per cent alcohol with 5 per cent annuonis will give much the same results, but scense more likely to bring out the neurlinnua nuclei). The pieces are then washed for from 3 to 3 minutes (according to their size) in distilled water and transferred to gyridine for 24 hours. They are then

placed in the dark for 3 days in a 2 per cent aqueous solution of silver mixeds at 35° C, then rinsed in distilled water and placed for 1 to 2 days in a 4 per cent solution of pyrogallic axid in 5 per cent formalin. Sections are made in parsfin, and, then mounting are ready for examination."

For use of the method in staining and sectioning the entire head of a small azimal or entrys, after desalizition, see Huber and Guild, Anatomizal Record, VII (1913), 233 and 331. For a discussion of the method with hilfography, see Rasson, Review of Neurology and Popularity, November, 1914.

III. GOLD-CHLORIDE METHOD FOR NERVE-ENDINGS

 Trace some of the motor nerves of a reptile or mammal to where they enter the muscles (intercostals are best), and clip out small pieves of the muscle. Use material that has been preserved in 10 per cent formalin (see note, p. 32).

 Place the bits of muscle in 10 or 12 times their volume of a 10 per cent solution of formic acid in distilled water and leave them for from 30 to 40 minutes.

 Transfer the tissue into from 8 to 10 times its volume of a 1 per cent solution of gold chloride in distilled water for from 30 to 40 minutes. Avoid direct sunlight. The moscle should become yellow in color.

4. Remove the tissue without washing it to about 25 volumes of a 2 per cent formic-acid solution and keep it in the dark until it assumes a purple color (24 to 48 hours). When the fibers appear reddish violet in color the reduction has gone for enough; if they show a decidedly bluish inge the process has gone too far.

5. Wash the tissue in several changes of distilled water for an hour and transfer a small piece to a slide. Tease the fibers apart very carefully under a dissecting lens. Great care must be exercised to avoid tearing the nerve fiber from its endings. Examine from time to time under a low power of the compound microscope, and when a nerve fiber with its termination is found, carefully separate it as much as possible from the other fibers.

 Add glycerin-jelly and a cover-glass. Seal in the ordinary way (p. 95).

Norz.—Tissues may be dehydrated in the ordinary way and mounted in balsam or imbedded in parafin or celloidin and sectioned.

CHAPTER X

ISOLATION OF HISTOLOGICAL ELEMENTS. MINUTE DISSECTIONS

I. ISOLATION

A. Dissociation by Means of Formaldehyde; ellisted and cohuman epithelium.—I. Kill a frog and secure the hinder part of the roof of the mouth, bits of the brain, and a small piece of the intestine. Slit open the laster. Leave the objects for 24 hours in a dissociating finid made by adding 0.5 e.e. of formalin to 250 c.e. of normal saline solution.

2. Scrape the roof of the mouth after removal from the fluid and mount the ellisted cells thus obtained on a slide. Similarly remove some columnsr epithelium from the internal surface of the stomach and mount on another slide.

 Add a coverglass and examine. If the cells eling together in elumps, separate them by drumning gently upon the coverglass with the handle of a needle.

4. Stain by placing a drop of alum-cochineal on the slide just at the edge of the cover and applying a bit of filter paper to the opposite edge of the cover. The filter paper absents the fluid from under the cover and the sisin replaces it. Keep the preparations under a bell-jar or other cover to prevent evaporation of the staining fluid.

5. After a few hours replace the stain by glycerin in a similar manner.

6. If a permanent preparation is desired, the cover-glass must be scaled (p. 65), or, after staining, the tissue must be dehydrated and mounted in balsam in the usual manner.

B. Isolation of Muscle Fibers by Maceration and Teasing.— 1. Place small fragments of volnatary muscle, of the root of the tangue, and of heart muscle of the frog into separate vials containing MacCallum's macerating third (respect 80, p. 238). After 2 days

pour off the fluid, fill the vials about half full of water, and separate the faseicles by shaking each vial. Further isolate the fibers by teasing.

Tenning—In trasing, the important thing to remember is that the elements of the tissue are to be separated, not breken up. Both painner and sharp clean needles are indispensible. The process is best earried on under the less of a disserting microscope or a timorthar dissection, although it may be done without such aid. A background which enables the tissue to be seen distinctly should be belevied, black for cohores or white for cohored objects. Back-and-white porcleain slabs are made for this purpose and are very convenient. A good disserting microscope has attached beneath the stage a reversible plate one side of which is black, the other white. Use a small piece of tissue and begin tassing at one end of it.

2. With the aid of a disserting microscope carefully tense out in water a number of fibers. Use a small piece and, beginning at one end, with both needles separate the piece along its entire length into two; likewise further subdivide these until the ultimate fibers are isolated.

 Transfer some of the fibers through the alcohols and xylol and mount in belsam. Stain others in alum-tochineal for some hours and mount in glycerin as above.

C. Maceration by Means of Hertwig's Fluid (Hydra. Testis).— 1. The solution consists of:

Prepare the ingredients for this mixture by diluting the stock solution (1 per cent) in each case with distilled water. Make a separate 0.1 per cent solution of acetic acid also.

2. Treat a hydra with the osmic and acetic acid mixture for 3 minutes and then transfer it to the 0.1 per cent solution of acetic acid. Wash in several changes of this fluid to remove all osmic acid and let the hydra remain in the acetic acid for 12 hours.

 Wash in water, stain in alun-cochineal or in acid carmine (reagent 88, p. 222) and mount in glycerin as above. If the cells are not sufficiently separated, gently tap on the cover-glass.

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 Submit small bits of the testis of some animal to the same treatment. Scain with methyl green (reagent 60, p. 231) or acid carmine (reagent 38, p. 222).

D. Mall's Differential Method for Retiruham.—1. Cut services of fresh spleen or lymph gland 40 to 80 microws thick by the freezing method and digest for 24 hours in the following solution:

Pancreatin (Park, Davis & Co.)	5 grams
Bicarbonate of soda.	10 grams
Water	100 e.e.

 Wash thoroughly in water, then, in order to remove cellular débris, shake for some minutes in a test-tube half full of water. Spread out on a slide and allow to dry.

 Apply a few drops of a 3.5 per cent solution of piceic acid in 11 per cent alcohol and allow it to dry on the preparation.

 Stain for about half an hour in a 10 per cent solution of acid fuchsin in 35 per cent alcohol.

5. Wash in the picric-acid solution (step 3) for a moment, then pass through alcohol and xylol and mount in balsam.

II. MINUTE DISSECTIONS

A. Alimentary Canal and Nervous System of Inserts.—1. Curveluly dissect out the alimentary scala and the central nervous system of a orekrouch with the ali of the dissecting microscope or lens. Wash each by gently flooding it with distilled water from a pipette, and then over it with Bouin's thait or corresive sublimate (reagent H.p. 122) for 30 minutes.

 Wash in several changes of water during the course of half an hour and stain for 40 minutes or more in borax-earmine.

 Wash in 50 per cent alcohol and decolorize in 70 per cent acid alcohol until the objects become bright scarlet in color.

4. Wash in 95 per cent alcohol for 5 minutes and then transfer to absolute alcohol for 5 minutes, xylol or targentine 10 minutes; and mount in balsam. Apply cover and label.

B. Guzard of Cricket or Katydid.—Pull off the head of a cricket or katydid. The ginard usually remains attached to the head part. Cut it open lengthwise, wash out the contents and mount as alove, but omit the staining. The inside should be turned uppermest.

C. Sing of Wasp or Bee. – 1. Place a wasp or bee in water, cover to keep out dust, and let it stand for two or three days until the smell becomes unpleasant.

 Wash in clear water and spaceae the abdomen gently until the sting protrules. With forceps pull it out earthily. The poison gland and duct should come array with it.

3. Place the parts removed on a shife and under a lens draw the sting out of its sheath by means of a smllb field which should be drawn over the outer surface of the sheath from the base to the apex of the sting.

4. Stain and follow out the same subsequent treatment as for II, A, above, or mount without staining. It is advisable to compress the object between two slides as soon as the acid alcohol is washed out. The slides should be tied together and left in 95 per cent sloubol several hours. Then proceed in the onlinary way.

D. Salirary Gland of Cockroach or Cricket—Let the animal soak in water as for preparation of sing. When sufficiently decayed pull off the head cardrally with forceps. The ecoplages, the salirary glands, and erop usually once along with it. Stain and mount as for sting. For preparation of fresh salirary gland use the "salirary gland of a Chiconomous larsa."

E. Month Parts of Insect.—1. Place the head of a bee or cocknoch in 95 per cent alcohol for 2 or 3 hours. Transfer to absolute alcohol for 30 nimetes, and then to cedar oil for 30 minutes to an hour.

2. Resurve the head to a slate and in a drop of the oil dissect out the mouth parts. Transfer them to a clean slate, remove the excess of oil, and arrange them in their relative positions in sufficient balaxm to hold them in place, then set the slate scale in a place free from dust until the balaxm hardens enough to keep the parts from slating. Make any necessary rearrangement. Add more balaxm and a cover.

MEMORANDA

 The Cover-Glass May Be Supported by means of small was feet, bits of broken cover-glass, or fine glass threads when the tissue is too bulky to allow the cover-glass to fit down closely to the slide.

2. A General Rule for Dissociating Tissues is to use small pieces of the tissue and not a very great amount of the fluid.

 For Minute Dissections clove oil is often a convenient medium. It tends to form very conven drops, cleass well, and renders the object brittle; any or all of which properties may be useful in such dissections.

 The Fixation of Pieces of Macented Tissue (e.g., macented epithelium) in 0.5 to 1 per cert osmic acid for an hour or so eften proves advantageous.

5. Congo Gyreeni is recommended by Gage as especially good for isolated preparations, particularly nerve cells. It is made by dissolving § gram of Congo red in glyreeni. It acts both as a stain and as a mounting-melium. The preparation may be scaled (p. 93) if desired.

CHAPTER XI

TOOTH, BONE, AND OTHER HARD OBJECTS

Sectioning Deckifiel Tooth.-I. Kill a cat and remove the lower jaw (p. 52). With a fine saw cut out about a quarter of an inch of the bone bearing a tooth (e.g., exame), remove as much of the surrounding tissue as possible, and place the object in Zenker's fluid for 1 or 2 days. Wash thoroughly in water and place in alcohol for at least 24 hours. Transfer to nitric-acid decalcitying fluid (reagent 10, p. 9). Use a relatively large quantity of the fluid and change it each day until the tooth is decalchied (2 to 6 days). It is sufficiently soft to cut when a needle can be thrust into it easily. Use this test sparingly, however, as it injures the tissue.

2. Wash it in repeated changes of 70 per cent alcohol until all trace of the acid is removed.

Transfer the object through 50 and 35 per cent alcohol successively to running water and wash for 24 hours.

4. Cut sections by means of the freezing microtome as directed under that method (p. 67). If a freezing microtome is not available use the celloidin method.

5. After disselving out all of the gun from the sections in distilled water, stain in atom-cochineal and Lyon's blue (see method, p. 50). Dehydrate. Remove one or two of the best sections (through the center of the tooth) to a slick, clear, and mount in the usual way in balsam.

6. Stain other sections in 1 per cent centic acid for 24 hours and mount in glycerin-jelly. When the jelly has hardened, seal the cover with gold size, and when this is dry add a thin cost of Bell's cement (see p. 95). If preferred, dehydrate and mount in balsam instead of glyrerin-jelly.

Sectioning Decalcified Bone.—Saw out a short piece from the femur of a cat (p. 52). Prepare transverse sections by decalcifying

and sectioning in the same manner as for teeth. Do not destroy the periosteum. Prepare likewise longitudinal sections of a tarsal hone.

Sectioning Bone by Grinding.—1. With a fine saw cut a thin transverse section of the femur of a cat. Let it macerate in water until quite clean, then dry it carefully.

 Grind the disk of hone between two hones, keeping the hones parallel in order to avoid wedge-shaped sections. The section is not thin enough until fine print can readily be distinguished through it.

3. Wash the section thoroughly in water, transfer it to absolute alcohol for 10 minutes, then to pure ether for half an hour.

 After removal from the ether, clamp it between two shdes by means of a string or a rubber band and let it dry thoroughly.

5. Place some xylol-lalsam in the center of a slide and heat it for a few minutes to drive off the xylol, then press the section of bone down firmly into it and put on the cover-glass. The air in the spaces of the bone makes then stand out black. The balsam should not be thin enough to enter these spaces.

MEMORANDA

1. Failure to Stain Properly is due ordinarily to insufficient washing out of the acid.

2. Teeth and Other Hard Objects may be prepared by grinding in the same way as bone.

3. For Other Decaktifying Fluids than nitric acid, see Appendix B, v.

CHAPTER XII

INJECTION OF BLOOD AND LYMPH VESSELS

Red Injection Mass.—1. Rub up 4 grams of earnine thoroughly with 8 c. of distilled water in a mortar and add annowinn hydrate drop by drop until a transparent red color results.

2. After quickly washing it to remove dust, etc., sock 10 grams of best French gelatin in distilled water until it is scollen and soft (18 hours), then remove it to a porcelain erraporating-tish and melt it at a temperature of about 45° C.

 While the gelatin is yet fluid, slowly add the coloring matter, stirring constantly until a homogeneous mixture is obtained.

4. Before the mass cools add also some 25 per cent acetic-acid solution drop by drop, stirring thoroughly until the mass becomes slightly opaque and the odor of ammonia gives place to a faint acid smell. Watch for this change closely, for a few drops too much of the acid will spoil the entire mass by precipitating the carmine. If the ammonia is not completely neutralized, on the other hand, the coloring matter will diffuse through the walls of the injected vessels and stain the surrounding tissues. Walker (American Journal of Anatomy [1905], p. 74) makes results more certain by mixing 1 part of the laboratory ammonia with 4 parts of distilled water, and then determining the exact amount of the laboratory acetic acid which will neutralize it. Knowing this, it is easy to determine the total amount of acetic acid which must be added for the amount of ammonia which has been used in any quantity of the gelatin mass. Just before using, the mass should be heated and strained through clean flannel wrung out of hot water.

With a large animal it is advisable to keep animal and apparatus submerged in warm normal saline during the operation of injection, but with a small animal this is unnecessary if the operator works rapidly.

Bue Injection Mass.—Prepare a gelatin mass as directed above. To the warm mass add sufficient quantity of saturated aspects solution of Berlin blue to give the desired blue color. If the blue does not disaclive, add a little oxalic acid to the mixture. The blue mass need not be made for the present practical exercise unless the student visitus to undertake a double injection as indicated in memorandrm 2, p. S7.

Yellow Injection Mass.—Prepare a gelatin vehicle consisting of 1 parts of gelatin to 4 parts of distilled water. Take equal volumes of the gelatin mass, a cold, estimated solution of hieleromate of potassium, and a cold, saturated solution of lead averate. Add the hieleromate solution to the gelatin and heat almost to bulking: then add slowly, while stirring, the solution of lead averate.

INJECTING WITH A SYRINGE; SINGLE INJECTION

A common method of injection, and one which proves satisfactory in many instances, is by means of a metal or glass syringe. Although not as desirable in the main as the method of continuous air pressure, many good injections may be made by means of the syringe. The apparatus consists of a syringe fitted with a stop-tock in the notale, and a separate tube, known as the example, which fits on to the end of the notale. The syringes are made in different sizes, and each is provided with an assortment of cannulae to fit vessels of different ealiber.

 Provide yourself with several strong threads about four inches in length for ligating blood vessels. Have the red injection mass melted and beated to about 50° C. Also have ready some hot water to warm the syringe.

2. Kill a cat or a rabbit by means of chloroform or illuminating gas. The latter acts more rapidly and causes less struggle on the part of the animal. Work rapidly so that the entire animal may be injected while yet warm. Stretch it out in a dissecting pan or the it out on to a board, or, better, keep it inmersed in a vessel of warm normal saline solution.

 Slit the skin along the ventral surface of the body to the middle of the neck and reflect it to the right and left side. Pin it back out of the way.

 Snip a small hole through the body wall just posterior to the ensitorm cartilage. Insert the index finger of the left hand to guide

Injection of Blood and Lymph Vessels

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the scisons and pievent injury to the underlying organs, and cut the costosternal cartilages of the right side up to the first rib. In like manner cut the cartilages of the left side up to the first rib.

5. Ligate the sternum tightly as close to the first ribs as possible to prevent leakage from cut blood vessels.

6. Cut off the apex of the heart and expose the ventricles. The left ventricle is seen as a round opening, the right as a slit.

7. With a sponge wrung out of warm water rapidly absorb the blood from the thorax.

 Choose the largest cannula that the sorta will admit and thrust it through the left ventricle into the sorta.

9. With a pair of fine-pointed forcess (preferably with curved points) pick up one end of a thread for ligating and cantully work it through inder the aorts (do not mistake the rens curva superior for the aorta). The the thread around the aorts over the cannula, making a double or surgeor's knot. Draw it tightly on the cannula so that the latter will be held firmly in place. Run another thread through under the aorta and have it in readiness to ligate the aorta when the cannula is withdrawn.

 Warm the syringe by sucking hot water into it repeatedly, then fill it and the cannula with the warm injecting fluid.

 Force out a little of the fluid from the syringe to expel all air, and connect it carefully with the cannula.

12. Force the injecting mass into the blood vessels by a slow steady pressure. Begin with a very low pressure, so that the large vessels will be thoroughly filled before the mass entres the explicites. The pressure should be gradually increased. Avoid sudden increase of pressure should be gradually increased. Avoid sudden increase of pressure or too strong pressure, for either may cause a rupture of the blood vessels and consequent extravisation. From 8 to 10 minutes is about the time required to make a good injection of the ext.

13. Examine the intestines and the guns from time to time and also the inside of the thigh (from which the skin has been reflected); they should be deeply colored by the mass before the injection is complete. If the mass begins early to flow from the right ventricle, the ventricle should be ligated. In any event, it is well to the the

ventricle a few minutes before completion of the injection, to insure filling of all blood vessels.

Norm—If the guns remain uncolored, the examples has probably been forced past the arteries which lead to the lead. In such a case, complete the injection of the trunk and then, if injected tissue from the head region is desired, our obliquely into one side of the innominate artery, the a cannot in place, and inject toward the head as in the case of the arts.

14. When the injection is complete, shut the stop-cock, ligate the sorta, or chump it with pressure forceps beyond the end of the cannuls and then remove the latter.

15. Place the animal in cold water or cold alcohol for half an hour, then remove pleos of liver, spleen, panetras, stomach, intestine, salivary glands, kidneys, and voluntary muscles and harden in strong alcohol or in 10 per cent formalin.

16. When sufficiently hardened transfer the objects to etheralcohol and proceed to imbed and cut in celludin according to the method already given. Make longitudinal sections of the bidney parallel to its flat surface. Cut transverse sections of liver, stomach, and intestine, longitudinal ones of the muscle, and sections passing longitudinally through the bilam of the salivary glands and spleen. The sections should not be under 30 microus thick. Mount some unstained; stain others in diluted Delafield's hematoxylin or in hemalam.

MEMORANDA

1. Apparatus for Continuous Air-Pressure Injections is now provided in many kaloratories. If a regular cylinder for air pressure is not present, however, anyone with all'the ingentity can reality fit up a suitable appartus. A carboy or large-monthed bottle which can be tightly corbital will answer as a chanker for compressed air, a water tap, or a tank of water elevated to the height of 7 or 8 feet will provide sufficient pressure. By making the proper connections by means of rabber and glass tuling a steady stream of compressed air may finally be conducted to a flask containing the injection mass; the flask works in the same way as an collinary wash-dottle [Fig. 35]. All corks and fittings must be tightly secured with wire or storag corel. If desired, by adding on extra perfection to the ork in the air chamber, a mercury manometer may be added to register the amount of air pressure. If a metal cannula is not at hard a glass one may be made as indicated under memorandum 9, p. 90. Inline of a stop-cock, us a pinel-cock on the rubber delivery tube.

Injection of Blood and Lymph Vessels

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2. A Double Lightion of the Vascular System may be made by first injecting the blue mass until it is seen to flow from the right ventricle, then detaching the tabe which conveying a two from the right ventricle, then detaching the tabe which conveying a role mass. This second mass should be in a bottle or fishs connected with the pressure bottle by means of an additional table through the control with the tabe from the pressure bottle by means of a V-true. Each must be provided with a pinel-cock or champ to hold back its contents while the other is in operation. If a syring is used, it is better to have a second syring for the second mass, although one will assere if it is rised out with how sater before being illed with the second mass.



Pra. 35.-Apparatus for Continuous Air-Pressure Injection (after B. G. Smith)

constant nined with it, so that when it reaches the capilaries they will become completely plugged. Walker uses the red gelatin mass first, then follows with a gelatin colored with ultramarine blue. The granules of the atter are too large to center the capilaries, hence a double injection with reins red and arteries blue is obtained.

It should be home in mind that the larger veins cannot be injected in a direction contrary to their flow because of the valves they contain.

3. The Langs, Liver, and Kidneys are readily injected through their larger blood vessels with two masses, and afford very instructive material when thus prepared. A triple injection of the liver may be made by injecting the begative artery and the begative and portal veins. The third mass may be colored with China ink. Withman (Mehods in Microsophical Andony and Endrydoy) recommends first injecting the hepstir artery

and afterward the two veins. The blood should be washed out of the organ to be injected, with warm salt solution.

4. To Inject Lymphatics, the puncture method is commanly employed. For example, as appears solution of Berlin blue is sharen into a hypotermic syringe, the sharp point of the cannula is thrust into the issue, and the syringe emptied by slight, steady pressure. For practice, thrust the examula into the pad of a cut's foot, and force in some of the injection mass. If the leg is rubbed upward, the finit will flow along the lymph channels and into the glands of the grain. Instead of this haphaneri method, however, much better results will be insured by the use of the needle-and-elamp device of W. S. Miller (Johns Hopking Hopking Bulletia, XVI, No. 173 [1936]).

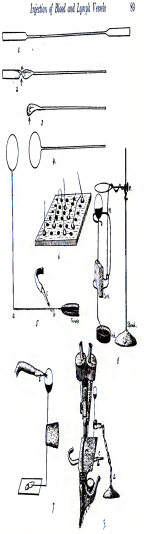
5. Micro-Injection of Embryonic Vessels has been much resorted to in recent years for the study of early stages of lymph and blood vessels, and some very delastic and effective methods have been devised. Infla ink is the medium ordinarily used. Embryos of medium size are generally injected with a hypoternic syringe, but smaller embryos require a more delastic procedure in which glass tubes with the finest possible capillary points are used.

layone who has seen the beautifully injected specimens of Dr. H. McE. Knower will concele the success of his method. The general scheme of his apparatus is shown in Fig. 30, and the accompanying legend is selfexplanatory.

He expresses the essence of his method as follows: "If a gentle transfit is applied to a glass bulk blown on the end of a capillary tube, while the fine point of the tube is held beneath the surface of some fluid, such as India, ink, air will be driven out of the bulk and ink will run up to replace it as the bulk cools. When the system has come to equilibrium, the point of the tube is inserted into the desired blood vessel under a dissecting microscope (binocular if possible), the tube boing exaried on a holder to avoid varning it or the bulk. India ink is row injected, as desired, by warning the bulk when ready." The method is explained in minute detail in the *Audomical Roosd* for Angust, 1908 (Vol. II, No. 3), together with applications to fish, amplifue, prefiles, binds, and mammals.

Instead of using a glass bulb, as does Knower, Heaser inserts a simply made resistance cell of fine German-silver wire into an ordinary sub-morth bottle. The earls of the wire extend out through the cort ior electrical connection. When a current is passed through the wire the air surrounding it in the bottle, becoming heated, expands and affords a steady and prolonged pressure.

Dr. Emily Ray Gregory uses direct pressure from a good goale of De Vilhist' atomizer bolls which lies on the floor. The bulk, kept from rolling by a erecheted net cover, is openeded by the foot. Pressure is transmitted through a f-inde red-valuer tube to the short glass injection tube



Pro. 36.—Apparatus for the Injection of Small Embryon, etc., under the Microscope (stree Knower).

1. Glass tube drawn out after moderate heating unor middle (see arrow). Tube should be turned while heating. The capillary take will usually be three or four times

shardle terms while horing. The explicitly take will usually be three or how times the length of havin in the figure. 2. Superforms the horizopher fames. The shored he transf will have into the figure of the length of the start of the horizon of the figure of the length of the le

holder. Size reduced.

more: solar curva. 8. Wire support w, for holding halls while alling. Reduced size. 9. Method of using holded holder a, with bimodiar. The small gas-jet c lies in front ready for heating the bulb. Size greatly reduced.

which has the outer end drawn into a capillary top at right angles to itself. The method is given in detail in the August number of the Anatomical Record, 1916 (Vol. XI, No. 1).

For other suggestions regarding micro-injections see Hoyer, Zathehrift für unsenchaftliche Mitrokopie, Band XXV (1988); Evans, American Jaural of Anatamy, IX (1910); and Salin, Contributions to Endergology, III, No. 7 (1913), Carnegie Institution of Weshington.

 To Keep Getain Injection Masses let them congred, then cover the surface with 95 per cent alcohol, and have in a well-stoppered vessel until meeted.

 Injection through the Penneral Artery is frequently practiced, and is preferred to injection through the sorts by some workers. An oblique cut is made in one side of the artery and the cannols inserted pointing toward the heart. Others prefer to cut into the dorsal sorts and inject both asteriorly and posteriorly.

8. The Injecting Syringe must work without jerking or catching along the wall of the burrel. It should always be exercitely cleaned after using. If the piston does not fit the barrel tightly enough it should be wrapped with gauge.

9. Glass Cannulae may be made by grasping the ends of a short piece of soft glass tubing and heating the middle in a flame until the glass becomes soft, which is indicated by the pellow color of the flame. The tubing should be constantly rotated, so that all sides heat equally. When the glass becomes soft, draw the tube out steadily until the diameter of the soft portion becomes as small as desired. When the glass has cooled, the tube should be cut with a fla at the proper place to make two enmalae of it.

 If the Blue Color Fades in the gelatin mass in the tissues, it may frequently be restored by treating the tissue or section with oil of cloves or turpentine.

11. A Gild Fluid Gelatin Mass hashen used successfully by Tandler (see abstract by A. M. C. in *Journal of Applied Microscopy*, V, 1023). To prepare the mass, dissolve 5 grams of finest gelatin in 100 e.c. of tepid distilled water. Color to the desired shale with Berlin blue, and then add slowly 5 to 6 grams of potassium indule. The mass remains fluid at ordinary temperatures, but when injected objects are placed in 5 per cent formalin it sets completely and is thereafter mathematic by reagents. The minutest vessels are injected, and sectors may be stained in the usual ways. Subjection to strong acids, such as subjuurie or hydrochlorie, does not affect the mass; hence it may be used for injecting specimes that are to be decalcified afterward. To preserve the fresh mass, add a low crystals of thymod and here in a stoppenel bottle.

 Corrosion of Injected Vessels or Cavities is sometimes practiced. A mass must be employed which will not be attacked by the reagent used

Injection of Blood and Lymph Vessels 91

for destroying the surrounding tissues. One of the best masses consists of white was 5 parts and rusin 6 parts, melted together at a temperature of about 15° C. For fine vessels increase the proportions of was, for larger ones add more rusin. Vermilion, Prussian blue, or chromate of lead may be used for coloring. The part to be injected should be placed in warm water and the mass injected at a temperature of from 30° to 60° C. The injected part is left in cold water for from 1 to 2 hours, and is then corroled in pure hydrochherie acid for from 6 to 48 hours, according to the resistance of the tissue. Finally, wash the preparation theoroghy in running water. For bibliography and more detailed directions see Technique de injection, by Hermana Joris, Université Libre de Brunelles, 1908.

13. Wood's Metal is one of the commonst injection media used for comsisten preparations. It is a fushle alloy consisting of 1 or 2 parts of eachnium, 2 parts of tim, 4 of lead, with 7 or 8 parts of bismuth. It melts at from 66° to 71° C.

14. Cellulid Dissolved in Acetme was found prefendle by Fint (American Journal of Antomy, VI [1966]), in his study of long development, to cellulian or Wood's metal for corresion preparations. He injected from aspiration bottles into the longs through the trackee, and used hydrochlaric soil for corresion.

15. Air Injection of Minute Vessels was found to be an indispensable method by Locy (American Journal of Anatomy, XIX [May, 1916], 3) in his work on the hung and air passages of the chick: "In stages subsequent to 96 hours, the lungs and air sats were dissected out of the previously fixed and hardened specimens, then cleared in cedar oil, after which the organs were placed in a mixture of 1 part cedar oil and 2 parts chloroform. On becoming permeated with this fluid, the preparation was removed from the mixture and placed on a filter paper until the chloroform might evaporate. The evaporation of the chloroform served to draw the order oil from the luming of the various branches of the bronchial tree into the lung tissue and to fill the spaces thus made with air. When this preparation was replaced in ours cedar oil, the difference between the refractive index of the imprisoned air and the surrounding medium gave the lung tubes the appearance of being filled with a metallic cast. Thus the minute air passages that could not be injected by other means were made clear. The finer details would disappear after a few minutes as the cedar oil percolated into them, but the same specimen, if carefully manipulated, can be treated repeatedly without apparent injury, and a complete picture could finally be obtained."

For later stages Loey also used celloidin and Wood's metal injections followed by corrosions. He has obtained some beautiful Wood's metal

casts of the adult lung. The lungs of the freshly killed lowl were distended under pressure with 50 per cent alcohol until the air save were fully expanded. The entire bord was then immersed in alcohol for 24 hours or more before metallic injection was attempted.

16. An Excellent Injection Mass for other than histological purposes as used in our own laboratories is made as follows (Wagner):

Water	100 c.e.
Glyceria	20 e.e.
Strong formalin	20 c.e.
Cornstarch, powdered	75 grams

Mix by gradually adding water and glyverin to the starch, rubbing out all humps. For yellow color add 10 grams of chrome yellow; for green, 10 grans of chrome green; for red, 10 grams of vermilion. Strain through chrosedoth and add the formalin. If the mass is too thick to strain, add the formalin fast. Constarch is vasily superior to humdry starch and the colors recommended diffuse less into tissues than camine or Berlin blue.

17. For Clearing of Injected Organs or Embryos "in Toto," see memoranda 15, 16, and 17 (pp. 102-104).

CHAPTER XIII

OBJECTS OF GENERAL INTEREST: CELL-MAKING, FLUID MOUNTS, "IN TOTO" PREPARATIONS, DRY MOUNTS, OPAQUE MOUNTS

When objects of considerable thickness are to be mounted, it is sometimes necessary to resort to cells which will contain the object. and support the cover-glass. Fluid mounts and aqueous media must occasionally be used for delicate objects which would be injuriously affected by alcohol, or which are unsuitable for mounting in balsam. When such mounts are used, whether in a cell or not, the cover-glass must ordinarily be sealed with a cement if the preparation is to be permanent. In all cases where it is at all practicable, balsam mounts are to be preferred for permanent preparations. Glycerin is a convenient mounting-medium for many objects, especially for temporary mounts. It is often used where such media as balsam would render the preparation too transparent; it is much more favorable, moreover, to the preservation of color than are resinous media. For making cells and sealing circular covers, a turntable (Fig. 37) is desirable, although the work may be done by following a guide ring drawn on paper and placed under the slide.

I. TURNING CELLS

Prepare 12 or 15 slides as follows: 1. Place a slide on a turntable and adjust it so that its center lies over the center of the turntable.

 Dip 3 small camel's hair peeril into gold size, but do not take up enough of the fluid to drop (see also memorandum 13, p. 101).

3. Choose a guide ring on the turntable which is of slightly smaller diameter than the coverglass to be used, which the table and hold the pencil lightly over the guide ring. The ring which

has been spun should be even. If it is not, practice turning rings until satisfactory ones are made. If the gold size is old it is probably too thick to make suitable rings. Pure linseed oil may be used to dilute it, but it is advisable to use only fresh gold size if it is obtainable.

4. The slide must be set aside to dry before it can be used for mounting. A gentle heat will aid in drying.



5. To some of the cells add successive coats of gold size as the previous one dries, so that you will have cells of varying depth.

IL MOUNTING IN GLYCERIN

A. Water Mites and Transparent Larvae .-- 1. Kill several small, colored water mites or transparent larvae of insects by means of chloroform (a few drops in water) and place them for half an hour (two or three hours for larger objects) into a mixture of water and glycerin equal parts, after which transfer them to pure glycerin.

2. Apply a thin coat of gold size to the upper edge of a cell which is of sufficient depth to accommodate the object.

3. Breathe into the cell to moisten it so that the glycerin will adhere throughout and prevent the formation of air bubbles.

Objects of General Interest

 Fill the cell flush with glycerin and put the object into it, carefully spreading out all parts.

5. Breathe on the lower surface of a clean coverglass, put one edge down on the edge of the cell, and then gradually lower the cover so as to avoid bubbles of air. When in place, press the cover down gently with the handle of a needle and see that it adheres all around. Wash off the emodel giverin and carefully wipe the slide with a cloth.

 Turn a comparatively broad ring of gold size around the edge of the cover to seal it, and when this is dry add a very thin coat of Bell's cement. Label and put away in a horizontal position until dry.

Carmox.—It is indipensable that the edges of the coverglass be perfectly dry before attempting to seal the preparation; otherwise the cement will not adhere.

B. Killing and Mounting Hydra.—1. With a disping-tube (memorandum 10, p. 101) remove a hydra to a warm watch-glass and leave it in only a few drops of water. Have ready some hot Bonin's fluid or corresive acetic, and when the hydra sends out its testacles and expands its body, apply the reagent by suddenly squirting it into the watch-glass so that it sweeps over the hydra from alocal to oral extremity and carries the testacles out straight. Then fill the watch-glass with the hot fluid.

 After 10 minutes pour off the fixing fluid and wash the animal thoroughly in 70 per cent alcohol.

3. Replace the alcohol with alum-cochineal or dilute hematorylin and stain for from 30 minutes to several hours.

4. Remove the stain with a pipette and replace it with a mixture of equal parts of glycerin and water for half an hour, followed by pune glycerin. Proceed farther as in the preceding exercise.

Norn.-After removal from the stain, if necessary, decolorize in acidulated water or alcohol (0.5 per cent hydrochloric acid), then wash out the acid thoroughly in tap water.

Hydra may also be dehydrated, cleared, and mounted in bakam (see also "Hydra," p. 264).

III. MOUNTING IN GLYCERIN-JELLY

Glycerin-jelly is frequently preferable to pure glycerin for mounting because it is a solid at ordinary temperatures. One formula for making it is as follows:

Water	42 e.e.
Gelatin	6 grams
Glycerin	50 e.c.
Carbolic-aeid erystals	2 grams

Let the gelatin stak in the water for half an hour, then dissuive with gentle heat. Add about 5 c.c. of white of egg and heat (not over 75° C.) for half an hour. The egg albumen gradually precipitates and earries down all fine particles of dust, etc., so that the gelatin is left perfectly clear.

Filter through moist, fine hot hannel and add the glyverin and the earbolic axial. Use only clean gelatin of the best quality. Warm for 10 or 15 minutes, stirring continuelly until the mixture is homogeneous. If heated above 75° C, the gelatin may be transformed into metagelatin, which will not harden at ordinary temperatures.

A. Small Crustacea.—1. By means of a dipping-tube isolate such small creatures as Cyclops, Daphnia, or Cypris.

2. Kill by warming slowly in a drop of water on a slide.

 Place them in a cell of proper depth, draw off all water with a pipette, and gently warm the slide.

 Place the bottle of glycerin-jelly into a vessel containing warm water until the jelly becomes fiquid, but do not let it get any warmer.

 Fill the cell flush with the warm jelly and arrange the objects in suitable positions.

 Breathe upon the lower surface of a clean cover-glass and put it in place in the usual way.

7. Wash away any trace of the jelly from the outside of the cell and when the slide is dry run a ring of gold-size cement around the elge of the correr. After this drise, vanish with Bell's cement. It is not an absolute necessity to seal glycerin-jelly mounts, but the write has always found it a wise precaution.

Objects of General Interest

B. Muscle of Insect.-1. Out off the head of an insect and bisect the trunk so as to expose the interior. Observe two kinds of muscular tissue, that of grayish color belonging to the legs, the yellowish to the wings.

 Take a shred of muscle and on a dry slide eardfully separate pieces of muscle fiber and stretch them out, while keeping them moist by breathing on them.

 Mount in glycerin-jelly as directed in the previous exercise (see also p. 251).

IV. MOUNTING IN BALSAM

A. Flat Worms.—1. Obtain specimens of Planaria from the under surface of flat rocks in the edge of streams (see "Planaria," p. 266).

2 Place the animal in a little tepid water. Watch until it is extended full length, then flood it quickly with corresive sublimate to which 1 to 3 per cent of acetic acid has been added. The animal may be removed after 30 minutes or an hour and washed thoroughly in 50 per cent alcohol to which a little fineture of iodine has been added.

3. Stain for 24 hours in alum-cochineal or in Delafield's hematoxylin diluted one-half with water.

4. Wash in water followed by 35 and 50 per cent alcohol each 15 minutes.

 Decolorize in acid alcohol until the color ceases to come away freely (10 to 30 minutes).

 Wash out the acid in 70 per cent alcohol, using the alkaline alcohol if hematoxylin was used in staining.

 Flatten the animal by compressing it between two slides by means of a rubber hand, and place it for 24 hours in 95 per cent alcohol.

8. Transfer to absolute alcohol for 1 hour, and to xylol until clear.

9. Mount in balsam in a thin cell or without a cell at pleasure. If on examination the separate organs of the animal are not seen distinctly, it probably has not been compressed sufficiently. This difficulty may sometimes be avoided in a measure by letting a cover-glass rest upon the live planarian to fatten it out slightly, and then running the fixing fluid under the cover. Specimens which have been in the laboratory for some weeks or months make better preparations than those fresh from the stream.

B. Mosquito, Gnat, or Aphid.—1. Kill a mosquito with eyanide or ehlowform and place it in cedar oil or turpentine for an hour.

2. Remove, and place it on its back on filter-paper. Carefully spread the legs of the insect, put a drop of thick balsam on a slide, invert the slide, and bring the balsam in contact with the thorar of the mospito. Spread the wings and the legs of the insect and gently press it down into the balsam.

 Add thinner balsam, see that the proboses and antennae are floated out properly, then add more balsam, and put on a coverglass.

V. OPAQUE MOUNTS

Some objects are mounted to be viewed by reflected instead of transnitted light. They may be mounted in the ordinary way, and when they are examined as opaque objects, the light from the mirror should be turned away and, if necessary, a strip of dark paper placed under the slide to shot of all light from below.

A. Beetles.—Choose a shallow cell for mounting the wing cases and legs of one of the Curculionidae, preferably Curculio impendia, the South Ameriean diamond beetle.

 Stak the part in colar oil or turgentine for half an hour, then place it in the cell in the proper position, the outer side of the case toward the observer.

2. Fill up the cell with balsam and add the cover.

B. Wings of Moths or Batterflies.—Prepare parts of the wings of moths or batterflies as in A. The wing of the clothes moth makes a good opaque mount.

C. Head of a Fly.-1. Secure the specimes (preferably one having colored eyes, as one of the galfies) and choose a cell of the proper size for it. The cell should be of such a depth that the over will rest lightly upon the object and retain it in the center of the cell. The head should present the front view when mounted.

Spin a very thin cost of gold size on to the dry edge of the cell so that the cover will adhere.

 Soak the head of the fly for a couple of hours in equal parts of glycerin and water.

 Moisten the cell by breathing into it, fill it with glycerin, and transfer the object to it.

 Breathe on the over-glass and apply it very earchilly to avoid air bubbles. When the cover settles into place, press it down gently to make it adhere to the censent.

 Set it saids to harden. When hard, seal on the turntable with gold size followed by Bell's cement when the gold size is dry.

Objects of General Interest

D. Foreleg of Dytiscus; the Great Water Beetle.—1. Detach the foreleg of a male, and seak it in 10 per cent potash solution (see reagent 86, p. 287) for a day or two.

Wash it in water, run it up to 95 per cent alcohol, and leave it there for 24 hours.

 Pass it through absolute alcohol and clear in cedar oil, turpentine, or xylol.

4. Lay the leg, disk side uppermost, in a drop of balsam on a Sink, add another drop of balsam, and carefully cover with a clean cover-glass. Pure a small weight (e.g., half of a bullet) on top of the cover to hald it down until the balsam hadrens.

VI. DRY MOUNTS

A. Stales—Propere a very skallow cell and let it dry. Throughly dry the scales from a motif's wing by gently heating them on a slide over a frame. Place the scales in a cell, warm the slide with the cell will becomes stoky, put on the overe and press it down until it adheres all around, and finally scal as in previous exercises.

B. Eggs of Butterflies, Small Feathers, Antennae of Inserts, etc., may be mounted as dry objects. Care must be taken to have them perfectly dry, or they will in time cloud the cover with moisture from within.

MEMORANDA

 Smill or Soft Insects or Their Larvae may frequently be mounted directly in glycerin, or they may be dehydrated and mounted in balsam. A method often used is to kill them in strong carbolic acid and mount them directly in balsam. The earbolic acid both dehydrates and clears. It is better, however, to clear the preparation further by immersion in celar of or style before adding the balsam.

2. Insects Having Hard Shells must first to scale in 10 per cent potable to soften them and render them transported if they are to be examined by transmitted light. The softer parts of insects so treated are destroyed and only the external parts remain. Such insects may be movated in given or givenin-jelly, or they may be dehydrated, elsered, and mounted in halson.

 Delicate Insects which are too frail to withstand much handling may be placed at once in colar oil or turpentine and after an hour mounted in balsam (see "Mosquito," p. 98).

4. Wings, Legs, Antennae, Mouth-Parts, etc., of Such Forms as Files and Bees, which have been preserved in alreaded, should be completely debythsted, eleared, and mounted in balsam in cells of the proper depth.

 Transparent and Soft Insets may be stained in alum-ordineal or hexatorylin in the ordinary way and mounted as whole objects, if desired.
 They will stain better if they have been fixed peeriously in some corresive-

sublimate mixture and then washed properly (see reagent 14, p. 212). To stain, follow the method outlined in IV, A, p. 97.

6. To Center an Object in a Cell (the head of an insect, for example), thread a fine needle with a hair and rm it through the object. Remove the needle and inshed the ends of the hair in the ement on opposite sides of the cell. When the coverglax is put in place the object may be adjusted by pulling the hair. After the side is finished and dry, the ends of the hair should be cut off at the edge of the cell.

Another method which will frequently asswer for an object to be mounted in balsam is to place the object (after clearing) in the center of the cell, cost it with balsam, stjust it propeely, and then set the slide away in a place free from dust till the balsam thickens. Finally fill the cell with balsam and soil the cover.

7. The Rabuka or Lingual Ribbon of the Samil or Sing should be dissected out and socked for a day or two in a 10 per cent solution of potask. If the animal is a small one, out off the head including the bureal mass and sack it in a solution of potash until the soft tissues are distroyed and only the radult remains. From the potash the radult is transferred to water and washed for some hours. With a strip of paper on each side to prevent crushing it, it should be placed between two sides, and the solids bound together by means of string or rubber banks. While held in this position, dehythete and clear it. Finally remove one shole and the paper and month the object in balaxm on the other shole. A shallow cell may be used if desired.

8. Plakes and Tapeworms are prepared in the same manner as Planaria (p. 97). The time of immersion in the various fluids should be lengthened in proportion as the object is larger than the planarian. See also p. 207.

For in two staining. Mayer's parametrize and Mayer's hematum are highly recommended by nearly all specialists on these forms. The animals should be much overstained and then very rapidly and completely destained in strongly arishabed (2 to 4 per cent HCI) 70 per cent alabed (Cort, *Transactions of the American Microscopical Society*, XXXIV, No. 4 (Detober, 1916).

To prevent the curling up of flat worms which are to be infiltrated for sectioning, Peaslee binds them by wrappings of thread to a bit of bristol board which is not removed until the animal is to be imbedded.

 Spirogyra, Protococcus, Volvor, Desmids, etc., may be mounted in a cell in the following copper solution:

Acetate of copper	l gram
Camphor water	240 e.c.
Glycerin	240 e.e.
Glacial acetic acid	0.3 e.e.
Corrosive sublimate, saturated aqueous solution	0.1 e.e.

Objects of General Interest

Mix throughly, filter, and keep in a glass-stoppered both. The green color of the plant may frequently be preserved for some time in this medium. The speaimen is washed in water, transferred to the cell, then the solution is added. The cell is overed and scaled in the usual way.

10. A Dipping-Tube is a simple glass table. To openate it, hold the tip of the forefinger over the upper call and dip the lower end into the water until it causes just above the object desired; lift the funger and let the air out of the tube, and the water will rush in at the lower end carrying the object with it. Replace the funger over the top of the tube and remove it; the water will rush in it as long as the funger is held fundly over the upper call. When the funger is removed the water and the object pass out. The object may sometimes be more readily likelunged if the tube is rotated. A pipetter made of a large-love glass table and atomizer bulk is also yeary envircedable.

11. To Keep Water from Evaporating from a Cell Too Freely, use a round cell and over it with a square cover-glass. Apply a lowsh well with water to the slide beneath one of the projecting corners of the overer from time to time. Capillary attraction will draw in the water and will keep the cell full. It a continuous supply of fresh water is necessary, one end of a lowely trained cotton thread may be kind long one side of the overe and the other end of the thread immersed in a small vessel of water which stands within half or thread-parters of an inch of the cell. A reservoir made from the bottom of a shell vial or homeopathie vial answers very well; it may be emented to the shile.

Protozoa and other small forms may be kept alive on a side for a number of hours by simply mounting them in water under a overe in a cell of blotting paper which has been assumated with water. For somaria for studying microscopic organisms, Walton (Ohio State University Bulletin, XIX, No. 5 [915]) used ring-like pieces of less paper cet somewhat smaller than the overegass. Such squaria keep for several hours. They may be made more permanent by letting them stand 15 to 30 minutes in order to allow the outside water to evaporate, and then running parafin oil around the margin of the over-glass.

12. Deep Cells are nade frequently by eating out rings of paper, lead, or block-in with gun purches and remarking them to the shife. Glass and hard-tubber rings of various sizes may be purchased from dealers. To support coverglasses Barker uses circular doth patches with a hole in the center. These may be bought of a stationer.

 Filtered Shellar is recommended by McChang as excellent both for making and for scaling cells. It may be colored with Bismarck lowen and similar dyes. Barker uses any good quality of enamel paint.

14. For a Method of Preserving Fine Dissections for microscopic study, as opaque objects, see memorandum 15, p. 134.

15. The Clearing of Total Spetimens as developed by Spalteholz (Ueber das Durchsichtigmachen von menschlichen und tierschen Pröparaten, 21 ed., 1914, S. Hirzel, Leipzig), whereby relatively large anatomical and embryological preparations can be made transparent, is an extremely useful method, particularly with injected objects. The following account of the method together with modifications introduced by herself is taken from Miss Sabin's article in Contributions to Embryology, III, No. 7, Carnegie Institution of Washington, 1915: "In general the essentials of the method are, first, fixation in formalin: second, a thorough bleaching of the tissues with hydrogen peroxide to remove the hemaglobin and other pigments; third, dehydration; and, fourth, clearing the specimens in an oil which has the same index of refraction as the tissues. As applied to embryonic tissues, the method, developed by Professor Spalteholz, to whom I am very much indebted, is as follows: The specimens which have been injected with India ink are fixed for 24 to 48 hours in 5 and 10 per cent formalin. Commercial formalin is slightly acid, which is an advantage for the India-ink injections, since the ink diffuses in an alkaline solution. Specimens which have been injected with silver nitrate are ruined by fixation in formalin, because the silver salt is changed to a white precipitate which obscures the vessels. If injections of hone are desired, the formalin may be made slightly alkaline and the diffusion of the ink prevented as much as possible by tying off all vessels before firstion. For large fetuses, which are to be cleared in toto, Dr. P. G. Shipley has found that the subsequent bleaching is made easier by washing the specimen in running water before fixation, thus removing much of the hemaglohin. After fixation, the specimens are washed in running tap water from 12 to 24 hours, followed by distilled water to remove the formalin, -The bleaching is done in hydrogen peroxide. Spalteholz adds a few drops of ammonia to precipitate the barium salts. This is not necessary with harium-free oxide. For adult tissues, Spalteholz uses undiluted peroxide; for the embryonic tissues about 2 to 3 per cent is the best strength. The small embryos with ink injections take about 20 minutes to bleach; for the silver specimens, 2 to 3 minutes suffice, and they must be watched constantly and the bleaching stopped before the silver is affected. Following the bleaching, the specimens must be washed thoroughly in running water and in distilled water. The dehydration may be begun with 50 per cent alcohol and the percentage increased successively by five points or less. After two changes of a good grade of absolute alcohol, the specimens are passed through changes of benzene into the synthetic oil of wintergreen. The small amount of benzene which is carried over evaporates quickly, and the few bubbles which develop in the bleaching process can be removed with needles. The oil of wintergreen should be entirely colorless, but both the specimens and the oil will gradually become brown with age. This is especially true of the

shee-ainste specimes, but they will keep for six months or a year in cil. They can be returned to alcohol for storage and recleared when desired, or they may be made permanent in balaam. The advantage of keeping the total specimens in cill earber than in balaam is that they can be disserted. On the other band, they are made more permanent in balaam. The cill of wintergreen makes the tissues tough, so that it is possible to obtain minute dissertions of the injected specimens.

"The Spaltehols method as applied to embryos can be very much simplified by changing the fixative. For mammalian embryos the best fixative is Carnov's mixture. This is absolute alcohol 60 parts, chloroform 30 parts, and glacial acetic acid 10 parts. In this mixture the acid is sufficiently strong to bleach the hemaglobin so that the peroxide is unnecessary. The penetrating power of the fixative is very great, which is of importance, since no injected specimen can be cut into until it is thoroughly fixed. The relations of the tissues are well maintained and the swelling due to the acetic acid tends to counteract the shrinkage that always takes place in the oil of wintergreen. The fixative does not affect any of the injection fluids. The process after fixation in the Carnoy's mixture is simple; specimens remain in the fixstive from 2 to 12 hours and are then placed directly into 70 per cent alcohol, dehydrated in graded alcohols, and cleared as before. The specimens can then be studied in toto, or dissected or imbedded in paraffin and sectioned. They should be imbedded through a mixture of the oil of wintergreen and paraffin. They do not become brittle in the oil, so that they may be sectioned after staying in the oil for many weeks. The shrinkage in the oil, however, seems to increase on long standing. The advantages of the fixation in Carnoy's mixture are that the specimens are even clearer than after bleaching with peroxide, there are no bubbles formed to damage the fissues, the time of the procedure is shortened, and the fixation is much better should it be desired to section the specimens after studying the vessels in whole embryos. Specimens which are strongly pigmented, however, must be bleached with hydrogen peroxide before they can be cleared."

For other methods of clearing in toto preparations see memoranda 16 and 17.

16. The Potsh Clearing Method (or Modifications of II) for "In Toto" Preparations is, seconding to Thurlow C. Nelson of our own laboratories, once of the best as well as one of the simplest methods for the demonstration of schedul and cartilaginous structures. Unless nervous or other structures are to be stained, the animal should be put into a 1 per cent potash solution immediately after killing. Tweaty-four hours in this medium should be sufficient to clear the overlying tissues to such an extent that the skeledal elements are clearly visible. After removal to glyrerin the specimen will keep indefinitely.

For Staining Nerve Tissue "In Toto," Nelson combines the Charles Shille hematorylin stain with the potash method. Three solutions are required: 1

Potassium hydroxide, 1 per cent aqueous solution

2	
Glacial acetic acid	1 part
Glyverin	1 part
Chloral hydrate, 1 per cent solution	6 parts
3	
Glycerin	1 part
Ehrlieh's acid hematoxylin	1 part
Chloral hydrate, 1 per cent solution	6 parts

A minuor, for example, is killed in 95 per cent alcohol and left for 45 hours. After the viscena are removed, it is next transferred to the potash solution for from 1 to 3 days. When transported, the specimen is put into solution No. 2 for 72 hours, then into solution No. 3 for a week. It is then destanded in solution No. 2 for 18 hours and cleared in glycerin. The nervous tissue should show up dark purple in the semi-transport muscular tissue of the body.

17. A Benzaldehyde Clearing and Fixing Method for "in Toto" Preparations has also been devised by Nelson (memorandum 16) as follows:

Small objects, such as eggs and small fish, may be put into the aldehyde directly as it acts as a dehydrating agent to a slight extent, being missible with 30 parts of water. Larger objects are dehydrated by running up through the alcohols and then clearing in the benzaldehyde. If a specimen of the fresh-water mussel, for example, Anodon, is treated with 50 per cent alcohol for 2 hours, 70 per cent for 5 hours, 80 per cent for 15 hours, and 95 per cent for 2 hours, and is then put into benzaldehyde, the mantle begins to clear at once, the whole preparation being almost transparent in 12 hours. As benzaldehyde is very unstable, being oxidized to form benzoie acid, it must be removed before the tissue is exposed to the air for any long period. Likewise vessels containing it must be kept tightly stoppered. Objects may be kept in the fluid permanently by sealing the container in such a way as to exclude all air. This method of clearing finds its chief use in clearing up the entire bodies of small animals so as to show injected circulatory systems and such structures. Small objects may be mounted in balsam from benzeldehyde if preferred.

To infiltrate with parafilm or to stain in 600, the object should first be passed through a mixture of 6 parts of xylol to 1 part of absolute alcohol. A satisfactory stain may be prepared by dissolving methylex-blue crystals in 2 co. of absolute alcohol and adding 6 co. of benares.

CHAPTER XIV

BLOOD

I. EXAMINATION OF FRESH BLOOD

c) General.—1. Throughly clean a slide and cover, hathe one thumb in ether-alcohol (reagent 4, p. 8), steriline a sharp needle by heating it in a fame, and then prick the back of the thumb with the needle.

 Place a small drop of the resulting blood on a slide and quickly put on a cover-glass. To prevent eraporation, the edges of the over may be surrounded by olive oil or vaseline.

Living corpuseles may also be studied in a drop of normal saline or in Ringer's solution (p. 239).

b) Effect of request.—When it is desired to study the effects of requests on fresh blood (e.g., distilled water, 1 per cent tamic acid, etc.), a dwy of fresh blood is placed on a shie, the over is put on, and then the blood is "irrigated" with the reagent. That is, a dwp of the reagent is placed at the edge of the over to be drawn under by capillary action. The process may be basened by applying the edge of a bit of blotting paper to the opposite edge of the over.

c) To demonstrate block-platieta.—Place a small drop of a 1 per cent solution of methyl violet (reagent 61, p. 20) in normal selt solution, on the back of a thomb which has been cleaned by washing it in ether-alcohol. With a sterilized needle prick the thumb through the stain and mount a drop of the blood which enclose. Examine it under a high power. Both platelets and white encyuseles are stained.

d) Statised preparation of Shrin. - Mount a dray of blood on a shife as in a moist chamber for from 20 to 30 minutes to cougalate. Lossen the cover with a few drays of water and then theroughly irrigate the preparation with water. Drain of the water, bloot the preparation with blotting paper, and add immediately a dray of a 1 per cert approx solution of event (pregent 43, p. 224). Remove this site 3 minutes, rines the preparation is water, then treat it 3 minutes with a 1 per cent approas Solution of methyl violet (reagent 61, p. 231). Kines the preparation in water, let it dry, and finally mount in balsam.

e) Crystals of the blood.-

 Henoglobin Crystak—Allow a drop of blood to dry on the slide without covering it. Long rhombic prisms of a red color crystallize out. The blood of a rat is best for demonstration. A more certain method is as

follows: To 5 cc. of blood in a test-tube add a few drops of ether and shake the mixture rigorously until the blood becomes laky. Place a drop or two of the laked blood on a slide and allow it to dry in the cold.

 Hematoidin Crystals; reddisk-yellow crystals (rhombie plates).— They can be obtained from old blood extravasticus (e.g., cerebral hemorrhage, corpora lutes, etc.) by tessing. Mount in Canada balsam.

3. Henin or Peichmant's Crystals—To a small drop of blood on a slide or a bit of eloth which has been previously saturated with blood, add a few crystals of common sult. Heat over a flame until the mixture has become dry, leaving a redifish-town residue. Apply a covergless and fixed the preparation with as much aretic axid as will remain in place under the over. Heat the preparation until the acetic axid balls. After the axid has evaporated, the preparation may be made permanent by adding Canada halsam. The crystals are very small, narrow rhomine plates of dark-hown color. They vary in size and may lie singly, across one another, or in stellate groups.

The presence of these crystals is positive evidence of the presence of blood, hence their demonstration is of great importance in stains or fluid suspected of containing blood.

II. COVER-GLASS PREPARATIONS

a) Dry preparations (Ehrlich's method).—1. In this method the preparation is "fitted" by means of heat. Under one end of a copper has or copper triangle (Fig. 26) place a fame. After 15 or 20 minutes a given point on the bar will have a practically constant temperature. Throcogdby dean the bar, run a stream of water along the toy of it toward the fame, and locate the point farthest from the fame at which the water bolls. The blood smears when prepared are to be placed film side up in a row across the bar about three-fourths of an inch neurer the fame than the point at which the water just bolled. This will subject them to a temperature of about 129° C.

2. Thereughly clean and dry two error glasses, touch one to a small drop of perfectly fresh blood as it comes from the farger or toke of the ear and instantly drop it on to the second cover. The blood should spread in a thin film between the coversy, if it does not, it has begun to complete and the preparation will be inferior. Rapidly separate the covers by shing them spart, wave them in the air a minute to dry the films, then place them down with the same side uppermost. Do not prove the corror topford to spread the blood because this runs the corpusels. If the red corpusels are to retain their shape the films, then place the most bay and uniformly thin. Practice until you have prepared such a film.

Some workers prefer to make smears on slides instead of on cover-slips. A small drop of blood placed near one end of a perfectly clean slide is spread by drawing an edge of the end of another slide through the drop and along the surface of the first slide to its opposite end. Crushing is thus avoided.

Norm — In the clinical examination of blood great care must be exercised to have it alsolutely read; furthermore, the over-glasses should be handled with forceps inclead of by means of the fugers. It is recommended that me pair of the furceps be Connect or spring forceps of some kind (Fig. 39). The bloc of the are is perlays the best regim from which to obtain the blood. The needle with which the puncture is made should always be sterilized. Wipe array the first drop of blood that appears. The drop finally chosen should be not little larger than a pin-head. The whole operation cannot be performed too rapidly. To shorten the time it is well to have an assistant to prick and manipulate the ear while the operator strends to the preparation of the film.

 When several satisfactory films have been prepared, place them on the heated har, as indicated in step 1. Cover them to keep out dust and leave them for from 30 to 60 minutes.

4. Remove the covers and stain the preparations 15 or 20 minutes with Ehrlich's triple stain (reagent 42, p. 223) by flowing the film with the stain by means of a pipette. Rinse of the surplus stain with water, blot the film with blotting paper, and dry it by holding it with the edge downward high above the fame. When dry, mount in balsam on a skile.

Norz.-Instead of heating the preparation, much the same results may be obtained by subjecting films (prepared as in step 2) to other and alcoloi (p. 8) for from 1 to 12 hours, drying them again in the air, and then staining as above.

b) Royid method.—1. Prepare a film as above (a, 2), but before it bas dried treat it for 30 minutes with a saturated aqueous solution of cornesive sublimate (reagent 14, p. 212).

2. Wash the preparation thoroughly in water or in 50 per cent alcohol.

 Stain for 10 minutes in Delaheld's or Ehrlied's benatorylin (reagent 52 or 53, p. 226), rinse in 70 per cent alcohol, and stain for 20 seconds in easin (0.5 per cent solution in 70 or 50 per cent alcohol).

 Rinse in 95 per cent and in absolute about each for 2 minutes, pass through xylol, and mount in balsam.

After rinsing following staining, some workers simply blot the preparation with blotting paper, dry it in the air, and mount it in halsam.

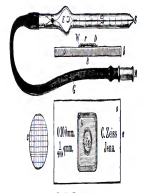
III. ENUMERATION OF BLOOD CORPUSCIES

The instrument used is the hemosytometer (Fig. 38). It consists of a special slide for counting and two graduated pipettes for diluting and messuring the blood.

Obtain a drop of blood from the lobe of the ear or from the finger. Fill the smaller pipette (with the 101 mark just above the chamber containing

beal) of the hemosytemeter to the mark 1 by earchil surtion. The tip of the torgue placed irmly over the hole in the monthpiece will keep the blood from dropping tock. If the blood is drawn beyond the 1 mark, blow it out immediately, clean the tube, and repeat the operation.

Wipe the blood from the outside of the pipette quickly and draw in sufficient Toisso's solution to make the level of the combined liquids stand precisely at the mark 101. Close the ends of the pipette with thumb and



Pio. 38.—Hemocytometee a, view of slide from shorre; h, view of slide from one skile; c, counting-ilsk wideh lies at the center of B; E, bead for mixing; M, monthpiece.

middle finger and mix the blood thoroughly with the solution by shaking the tube for 2 minutes. The blood is thus diluted 100 times.

Totizon's solution:		
Sodium sulphate	8.0	grams
Sodium ekloride	1.0	gram
Neutral glycerin.	30	0.0
Methyl violet, 5b.	0.02	i gram
Distilled water		

Blow out a drop of the liquid to remove the manifed solution remaining in the expillary tabe. Have the counting-disk and coverglass perfectly clean. Allow a drop of the diluted blood to flow on to the disk and place the coverglass over the drop. The cell of the disk must be entirely filled by the drop of blood. Let the corpusels settle a minute or two before beginning to count. Examine the field under a low-power objective to see that the corpusels are evenly distributed. If they are not, the blood was not thoroughly mixed and the whole operation should be repeated after thoroughly mixed and the whole operation should be repeated after throughly mixed and the whole operation should be repeated after

Examine the preparation under a high power of the microscope, and count the number of red corposels in 20 to 40 small separes; of those corposelss which happen to lie on the boundary line, count the ones that lie only in the upper and on the left sides of each square. Take the average number in a square and calculate the number of corpuscies in a cubic millimeter of blood.

The depth of the entire cell is 0.1 mm, the area of each small separe is $\tau_{\rm b7}$ sq. mm, consequently the volume of blood in each square column is $\tau_{\rm b7}$ cm, mm, or 1 cm, mm, of diluted blood would contain 4,000 times the average number in a square. One cubic millimeter of unidated blood contains 100 times as many, or 400,000 times the number in one square. What result do you obtain? For accuracy, three squarts counts should be made and the average taken.

After finishing the count, clean the pipette by successively deaving into and expelling from it water, alcohol, and finally ether. Do not blow through it, but cause the ether to evaporate by sucking air through the tube. For counting the white corpuscies use the large pipette and dilute the blood 10 times with one-third of 1 per cent glarial sectio acid. The acid destroys the red corpuscies, and times the white corpuscies are more readily seen. Proceed in the same manner as for red corpuscies.

Use water instead of alcohol for cleaning the counting-tilde, as alcohol dissolves the cement of the slide.

IV. OBSERVATION OF THE BLOOD CURRENT

a) Gradution in the web of a ford's foot-Wind a long strip of cheesechith around a frog stretched out upon a narrow piece of thin board, leaving one hind host exposed. Such the dubt in water in order to keep the animal's skin maist. Fin the extended foot in such a way that the web between the tors is stretched over a notch or hole in the end of the board. Examine under the microscope. If the preparation is favorable, hencoytes may perhaps be seen penetrating the walk of the vessel (dispodents) and passing into the surrounding fissues.

b) Circulation in the meentary. Information—Immobilize a frog (the male is better) by injecting a few drops of a 1 per cent solution of curve into one of the docsal lymph asis. Curve paulytes the nerve-codings. After waiting 20 minutes for the curve to be absorbed into the invalation, end open the abdominal wall for a short distance along the left site and draw out several loops of the intestine. Pin out a favorable area of mesentery over a cork ring and, after covering it with a cover-glass, examine under the microscope. Keep the parts moistened with normal sub solution. Such a proparation is especially furveable for studying the migrations of lencorytes through the walls of the vessels. Do not have the mesentery stretched too tightly or the circulation will cease. After a time the phonomenon of inflammation may nealiby the observed. It is hastened if some irritant (e.g., a drop of rewords it is amind to the mesentery.

MEMORANDA

 For Demonstration of the Different Granules of Leucocytes, etc., see p. 240, under the general topic of blood.

2. To Study Blood in Sections, light a small vessel in two places to keep in the corposelss, then remove the piece so prepared and first in Bonin's fluid or osmic acid. Indeel in parafilm and cut thin sections. Stain material find in Bonin or corresive-sublimate reagents by the henatoxylin-techn method (p. 50) or with the Earlich-Biomil stain (41, p. 222). The blood fixed in oscile acid may be stained by the sufficient-gentian violet method (33, p. 234).

3. Ametoid Morements in Leucortes may readily be observed in blood (preferably amphilizat) which has been mounted on a solie in very slightly marked memal salme. Place a hair under the coverglass and seal the edges of the latter with vaseline or melted parafin. For continuous study of the white corpuseles of warm-blooded animals a warm stage of some kind is necessary to keep the temperature of the blood near the temperature of the body.

4. Ingestion by Lewceytes.—Rub up sufficient India ink in a few drops of normal soline to make a grayish fluid. With fine sensors make an indiation one of the dorsal lymph sars of a chloroformel frog (nuclei to and else beside the mustyle). Introduce a capillary pipetie into the wound and obtain a small droy of lymph. Min it on a slick with a drop or two of the prepared ink. After placing a hair across the field, put on a over-glass and scal the edges with vachine or methed parafla. Under a high power of the microscope the cells may be seen englishing the colored particles.

Gage (The Microsop) resommends a mixture of lamp-black, 2 grans; sodium édiočile, 1 gran; gran Antóie, 1 gran; distilled water, 100 cc. Mix thoroughly is a mortar and filter through one layer of grane and one of lens paper. When injected into an animal the lencorytes will ingest the particles of earbon.

5. Wright's Stain for Blood.—This is a molification of Leishmark Romanowsky stain. To prepare the stain make a 0.5 per cent solution of solium hearboarke in distilled water and add to it 1 per cent of methylen blue (B.X., or "medicinally pure"). Subject the mixture to live steam in an ordinary steam sterilizer (e.g., Arnold; not a pressure sterilizer or a water-lath) for one hour. The container should be of such size that the liquid forms a layer not more than 6 cm. deep. When the mixture is cool, filter to remove any precipitate. To each 100 e.e. of the filtered mixture add, with constant storing, 500 e.e. of a 0.1 per cent acproas solution of "yellow, water-soluble" cesin. Collect the resulting precipitate on a filter, dry it theorogith, and, rubhing up in a porceluin dish or motar if messary, make a 5 per cent solution in pure methyle alcohol. To prevent the alcohol

from eraporating, keep the bottle containing the solution tightly stoppered. Should precipitation occur, filter the stain and add a small quantity of methyl alcohol.

Mallory and Wright in their Pathological Technique, p. 333, give the following summary of the method for staining blood films:

 Make films of the blood, spread thinly, and allow them to dry in the air.

Cover the preparation with a measured quantity of the staining fluid for one minute.

3. Add to the staining fluid on the preparation the same quantity of distilled water as there was of the stain. Allow this mixture to remain on the preparation for two or three minutes, according to the intensity of the staining desired. Examplifie granules are best brought out by briefer staining.

 Wash in water, preferably in distilled water, until the film has a pinklish tint in its thinner or better-spread portions and the red corpuscles acquire a yellow or pink color.

5. Dry between filter-paper and mount in balsam. The preparations retain their colors as long as any preparations stained with anilin dyes. Fresh films stain better than those which are several hours old.

Engineering when stained by Wright's stim should appear orange or pink in color (with deep-blue nuclei, when nucleated); long/horstes should show purplish-due muchs and cytoplasm of tobin k-eng blue with occusional dark-blue or purplish grandles; polynodear neutrophilo leasongles should have blue or dark like-voltered nuclei with cytoplasmic grantles of reddishlike color; essionplithe leasongles should show blue or dark like-voltered muchs; and blue cytoplasm with grandles the color of essin, large neuronologr leasongles should show blue or dark like-colored muchs; which explasm pale blue in one from and blue with dark like or deep purple-colored grandles in the other; mot-blue should exhibit incolor, are indeelded; myeloopte have dark-blue or dark like-colored muchs and blue extraplesm containing numerous dark-blue or reddish-like-colored grandles; blood-platelet are stained blue.

6. For Malarial Penseites Wright's stain (memorandum 5) is everlient. It yields the so-called Romanowsky stain; the color of the elementan warks from like to very dark red, while the body of the parasite stains blue. A full account of the method will be found in Mallory and Wright's Pathological Technique, p. 417.

7. Ehrlich's Triple Stain for blood is given on p. 223.

CHAPTER XV

BACTERIA

No strengt is male here to give even an elementary account of bateriological technique. Only such places of the work as are encerned with the immediate microscopical examination of bacteria are touched upon, and these chiefly to afford some practice in this kind of manipulation. For special technique, identification, or obscriptions of apparatus and arcessories, the student is referred to standard testbooks.

BACTERIAL EXAMINATION

Bacteria when prepared for microscopical examination are in the form of A. Cover-glass preparations, B. Bacteria in discuss (section method), or C. Hanging-drop preparations.

A. Cover-Glass Preparations

I. Killing and fizing .-

1. From Fluid Media (e.g., locallon, milk, water, saliva, blood, pas, etc.).—Sterline a platinum wire loop by heating it red hor in a flame. When cool, touch the loop to the culture and spread the adherent bateria in a thin film over the surface of a covergluss which has been sterlined in a flame. After the film has drived in the sit, kill and far the bateria to the cover by passing it three times, film sile uppermash, through the space of a flame.



Fig. 39.-Cornet's Cover-Glass Forceps

Each time should not exceed half a second. Prepare several films from a given material. Convnet or similar forcess (Fig. 30) should be used for handling such films, because the cover-glass can be left in them through the entire operation of fixing and staining.

If a platinum loop is not at hand a second cover-glass may be used to spread the smear. The first cover-glass is held in a pair of cover-glass 112 Bacteria

forceps and the second coverglass is dropped on to it. The glasses are then rapidly drawn apart with a sliding motion by means of forceps. The glasses should not be pressed tightly together. Professory in making such preparations is gained only after considerable practice. The chief secret in making a good preparation is to get the films extremely thin and evenly distributed.

2. Prom Solid Media (gelatin, agor, meat, polato, aximal issues and organs, etc.)—The procedure is the same as for 1, except that a drop of sterilized water or bouildne is put on the cover-glass to doubtate the spreading of the bacteria in a film over the over.

II. Staining and mounting.-

 Gentian violet (memorandum 3, 4, p. 116), 5 minutes. The coverglass is left in the foreps, film side up, and the film flooded with the staining fluid.

2. Rinse in water.

 Gram's solution (memorandum 3, f, p. 117) until the color becomes black (2 to 3 minutes).

 Ninety-five per cent alcohol until the violet color has almost completely disappeared.

5. Rinse in water and examine by placing the cover-glass film side downward on a slide. Only a thin film of water should remain between the slide and the cover. Remore surplus water by means of histing paper. If a prelonged examination is to be made, water lost by eroparation must be replaced by occessionally placing a small drop of water at the edge of the cover. In ordinary work the final inspection is frequently modes at his slage. If a permanent preparation is desired, however, proceed with the following steps:

6. If the batteria are well stained, a counterstain of Bismarck brown (memerandum 3, d, solution 2, p. 116) may be added (5 to 10 seconds). This step may be omitted.

7. Absolute alcohol, 10 to 15 seconds.

8. Xyol.

9. Xylol-balsam.

Norm—In stating, if the cover glass is warmed over a flame some 15 or 20 seconds until the stain steams, the action of the stain is usually more intense and more rapid. Builing, however, must be avoided.

B. Bacteria in Tissues

Tissues may be fixed and hardened (e.g., Gilson's fluid, Appendix B, p. 203, reagent 16; or Zenker's, reagent 6; or formalin, reagent 15) in the endinary way and sections made by the usual methods. Where provinsible, parafin sections are preferable to celloidin sections, because the celloidin

tends to hold the stain and thus obscure the bacteria. Sections should be fixed to the slide (parsfin by alloumen fixative, celloidin by ether vapor).

Batteria which do not stain by the Gram method (memorandum 3, 4, p. 117) are difficult to demonstrate, because it is hard to stain them so as to differentiate them from the tissues in which they lie, intrhermore, most of them easily loss whetever stain they may have taken up. Löffler's alkaline methyten blue (memorandum 3, b, p. 116) is, perhaps, the most useful stain for these organisms.

Methyleo-Blue Stain for Bacteria in Tissues.—1. Siain sections (parafilm) 30 minutes to 24 hours.

2. Acetic acid (1 to 1,000 of water), 10 to 20 seconds.

3. Rinse in absolute alcohol 20 to 30 seconds.

4. Xylol.

5. Xylol-balsam.

With celloidin sections substitute 65 per cent alcohol for absolute (step 8), then treat with creases or colar oil until sections are clear. Mount in xylol-balsam.

Anilin gentian violet, methyl blue, methyl violet, or fuchsin (memorandum 3, o, p. 116), also carbol-fuchsin (memorandum 3, c, p. 116) may be used in the same way.

Gram's Method for Bacteria in Tissues (Weigert's modification).--1. Stain sections (any kind) in lithium carmine 2 to 5 minutes.

Lithium Carmine (Orth's):

 Anilia gestian violet, 5 to 20 minutes (celluidin sertions should first be delyufusted in 95 per cent alcohol and afficed to the shife with ether rapor).

3. Rinse in normal saline.

4. Gram's solution (memorandum 3, f, p. 117), 1 to 2 minutes.

5. Rinse in water.

6. Blot sections with filter-paper to remove as much water as possible.

7. Anilin oil, several changes. The oil dehydrates, and at the same time decolorizes the celloidin.

8. Xylol, several changes.

9. Xylol-balsam.

Bacteria

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C. Hanging-Drop Preparations

 A shile with a concave center is used (Fig. 40). With a fine-pointed brash paint a narrow strip of veseline around the margin of the concervity. The vaseline makes the cover-glass stick to the shile and also prevents eraporation.

 Place a small drop of the fluid containing bacteria in the center of the cover-glass. If the bacteria to be examined are on a solid medium, the "drop" should be made by mixing

a small perion of the growth with a drop of boullon, normal saline, or serum. Place the cover-glass,

drop downward, over the depression in the slide and press it down well into the vaseline.

3. Use only a small opening in the displangm when examining the bacteria, in order to get as much contrast by refraction as possible. Focus first with a medium-power dry objective on the edge of the drop, then employ the oil immersion. Such medianel organisms are frequently difficult to find and there is great danger of breaking the cover-glass with the objective.

Hanging-drop preparations are used mainly in determining the motility of bacteria, or in the study of spore formation. For the latter purpose the shole and cover-glass must be carefully sterilized and the scaling with vaseline complete. The preparation may then be placed on a warm stage or in an inerbator and examined from time to time.

MEMORANDA

 The Mein Paints to Be Observed in the Microscopical Examination of Bacteria are as follows: (1) form of the individual, whether spherical (neural, spinal (apirallaw), or realike (bacillus) with end square, pointel, or rounded; (2) uniformity in size; (3) the arrangements of individuals whether single (microscet, etc.), in pairs (e.g., alphoneci), no chains (e.g., draptoscet), groups of from (e.g., ketwore), onbical groups of eight or more (survivae), or small grapelike bunches of various-sized corei (doupdiococol); (4) presence or absence of cell wall, gelatinous capsule, etc.; (5) matility in bring forms (do not confuse with Bownian movement); (6) reaction to stains; (7) presence of spores which are recognizable as bright, highly refractive numbel bodits.

2. Material for the Demonstration of Bacteria (secons, healbur, spirilhum, and begrates forms) will be found in abundance in foul water, expeeatily when contaminated with seconge. By scraping the inside of the check such forms as Leptothrix may often be found. Make a cover glass preparation; kill and far in the flame in the collinary way; stain in methyl violet,

genian violet, or fucksin (basic) and, if desired, counterstain lightly with Bismarck forwar; example in water or delyrchate in absolute alcohol, clear in sylol, and mount in talsam.

To demonstrate bacteria in tissues, a mouse may be inconlated with anthrar, and parafilm sections of the spleen prepared. Shin by the gentianviolet method.

3. Some of the Most Important Stains for Bacteria are as follows:

a) Anilia water solution of gentian ricket (Koch-Ehrlich's).-

Gentian violet, saturated alcoholic solution...... 10 c.e. Anilin water (see reagent 30, p. 218)............ 100 c.e.

After chains, the mixture should be set asile for 24 hours because of the prepiritation which takes place soon after making. Solutions of fucksin (tack) and methyl blue are made in the same way. These solutions begin to decompose after shout 10 days and must then be freshly prepared. They yield good results with many species of batteria. The gentian which, particularly, is wilely used in connection with Gran's method (see f).

b) Alkaline methylen blue (Löffler's).-

Methylen blue, saturated alcoholic solution...... 30 e.c. Caustic potash, aqueous solution (1:10,000)...... 100 e.c.

This stain keeps well and is one of the most widely used of the general stains. It is especially serviceable in staining the bacillus of diphtheria or of glanders.

e) Carbol-fuchsin (Ziehl-Neelson's) .-

This stain keeps well, stains powerfully, and can be used on many forms of bacteria.

d) Neisser's method for the diagnosis of diphtheria.-

Solution 1:

Methylen blue (Grübler's)	l gram
Alcohol, 96 per cent	20 c.c.
Distilled water (add after the methylen blue has	
dissolved in the alcohol)	950 c.c.
Glacial scetie acid	50 e.c.

Solution II:

Coverglass preparations are stained for from 2 to 3 seconds in Solution 1, tinsed in distilled water, placed in Solution II for from 3 to 5 seconds, rinsed again in water, and examined in the ordinary way. The bacteria of virulent diphthetis should appear as patchrown rods, some of which show at one or both ends bluids-black oval bodies of greater diameter than the rod. Such dark bodies will not be seen in the persolo-diphthetis bacili.

The bacilli must have been grown for from 12 to 18 hours on Littler's blood-serum which is a mixture of ghrose bouillon 1 part and beef-blood serum 3 parts. The mixture is run into test-tubes and coognized at 100° C; the tube should be tilted to one side to give a slanting surface for enhure purposes. The formula for ghrose bouillon is as follows: dry glucose, 10 grams; Liebig's extract of beef, 3 grams; peptone, 10 grams; softum esharde, 5 grams; water 1,000 c.e.

e) Gabbel's solution for demonstrating tubercle bacilli.-

Methylen blue	1 to 2 grams
Distilled water	75 e.c.
Concentrated sulphuric acid.	25 c.c.

The acid devolutines, while the methylen blue serves as a contrast stain. The solution sets isopilly. A modification of the method to be commended is first to stain the preparation with carbol-fachsin (see c) by warming the stain on the shife multi it steams, rinsing in water, and then proceeding with the methylen-blue solution. Smegma and leprosy bacill, and the trepowerms of sphills are also stained by this method. Tuberels bacill, and the trepowerms of sphills are also stained by this method. Tuberels bacill are also stained by Gran's method (see f). To examine spottom for tuberels bacill, the spottom is earchally inspected for small yellowish-white cheesy masses varying in size from the diameter of a spin-head to that of a small pea-Very thin smear preparations (see A, p. 112) are made from such masses.

f) Gram's method -

The preparations are first stained in anilin gentian violet (memorandum 3, d), and then immersed in Gran's solution for from 1 to 2 minutes. They are then rised in alcohol until the violet color is no longer visible to the naked eye. To devolate them sufficiently, it may be necessary to treat them again with the indue solution. Finally rises in valet and examine, or, if a permanent preparation is desired, rises in adsolute alcohol, transfer to rylol, and mount in balaan. If the prevantions are from enhances, it should be

borne in mind that the method works well only when applied to bacteria from actively growing cultures; old cultures seldem yield satisfactory results.

PATEOVENIC BACTERIA STAINED BY GRAM'S METHOD	PATHOGENIC BACTERIA DECOLORIZED RT GRAR'S METHOD
Bacillus aerogenes capsulatus	Bacillus of bubonic plague
Bacillus of anthrax	Bacillus of chancroid
Baccilus diphtheriae	Bacillus coli communis
Bacillus of malignant edema	Bacillus of dysentery
Bacillus of tetanus	Bacillus of glanders
Bacillus tuberculosis	Bacillus of influenza
Micrococcus tetragenus	Bacillus mucesus capsulatus
Pneumoooecus	Bacillus proteus
Staphylococcus pyogenes aureus	Bacillus pyocyaneus
Staphylocorcus pyogenes albus	Bacillus of typhoid
Streptococcus pyogenes	Diplococcus intra cellularis meningitidis
Streptocorcus capsulatus	Gonococcus
	Spirillum of Asiatic cholera

4. Staining Spores (Akbott's method).—Prepare a cover-glass surear in the usual way. Apply the stain (e.g., methylen blae) and hold the coverglass over a fame until the liquid steams. Repeat the basing several times, but do not bail continuously. Hence the cover-glass in water and then devolvine the preparation in a 0.3 per cent solution of hydrochloric usid in 05 per cent alcohol, until all color visible to the tasked eye has disappeared. Wash in water. If a counterstain is desired, stain for from 8 to 10 seconds in antih-furthin solution. Russ in water and mount in the usual way. The spores are stained blue.

5. Staining Flagella (Bange's modification of Littler's method)—The locomotor organs of motile barteria are long, hairlike prolongations (1 to many) termed flagella. Special methods of staining are necessary for their demonstration.

Make thin cover-glass smears of an 18-hour culture which contains motile forms. Dry and fix in the ordinary way.

The mordant.-

Ferric chloride, aqueous solution (1:20)	25 e.c.	
Alum, saturated aqueous solution.	75 e.e.	
Shake well and add		
Fuchsin (basic), saturated aqueous solution	10 c.c.	

Filter and allow to stand for some time before using. Treat the smear for 5 minutes with this preparation, gently warning by holding it high above a fame. The fluid must not boil. Rinse in water, then stain faintly with eachof-uchsin. Repeat the process until a successful vsultis obtained. Mount in the usual way.

CHAPTER XVI

SOME EMBRYOLOGICAL METHODS; SECTIONS AND "IN TOTO" MOUNTS OF FROG AND CHICK; AMPHIBIA; FISH; MAMMALS; OTHER FORMS

THE FROG

Frog eggs and tadpoles are best fixed in Tellyesnicky's fluid (5, p. 209). Eggs in early clearage stages should, after such fixation, be preserved in 2 per cent formalin, but later stages and tadpoles are better preserved in 70 to 80 per cent alcohol. Before eggs can be seetioned, the thick albuminous costs which surround them must be removed (4, p. 121). In addition to the ordinary clearage and yolk-polog stages, I find 3, 5, 7, and 9 mm. tadpoles, both as whole mounts and sectioned, the most neful stages for a course in endryology. Older stages are also necessary for the study of external features of later development.

Allen (op. cit., p. 122) finds that clearage furrows show with greater distinctness if the eggs are bleached. The fixed and hardened specimens are placed in ordinary commercial hydrogen peroxide for a week or more until the pigmented area is of a light-hrown color. If formalin-bardened material is used, the formalin should be washed out before the objects are placed in the peroxide, otherwise the tissue will be distorted by the rapid liberation of oxygen. Tadpoles may be bleached white by this method and mounted entire (see B., 120).

A. Section Method

 Select several 7 mm. tadpoles which have been fixed in Tellyesnicky's fluid (5, p. 209) and stain for from 12 to 24 hours in alumexchineal.

2. Run the stained specimens up through the grades of alcohol into absolute alcohol.

 Transfer to absolute alcohol and chloroform, equal parts, for an hour, then to pure chloroform. After an hour add melted parafin

to the chloroform from time to time until the latter contains all the paraffin it will hold in solution. Leave the objects in this mixture for at least 24 hours.

4. Transfer to melted parsifia (melting-point about 48° C.) and keep for 2 or 3 hours at a temperature just high enough to liquefy the parsifia. Indeed after reading step 5.

5. Prepare at least three sets of sections, one set in each of the three different planes of the body. The sections should be cut some 20 or 30 micross thick. Read carefully the directions under memorandum 1, p. 126, before imbedding, so that the sections will be properly criented. Read also caution under step 10 on p. 125.

 Carefully following directions under memorandum 1, p. 126, meant the sections with albumen fixitive and albuminined water as usual (steps 17-21, pp. 40–41).

 Dissolve out the parsifin from the sections in the usual way after the latter are throughly dry. If not completely dry and tightly stuck to the lide some of the sections will float off.

 Pass the slides back into absolute alcohol for a few minutes, then into fresh xylol until clear. Add balsam and the cover-slip.

B. Whole Mounts (after Allen, op. cit., p. 122)

 Select 5 to 7 mm. tadpoles which have been fixed in Tellyesnicky's fluid (5, p. 2009) and bleached in hydrogen percende for one week, or until white.

 Stain for 12 to 24 hours in alum-cochineal diluted with distilled water until the stain shows but a faint tinge of color.

 Run the tadpoles up through the increasing grades of alcohol into celar oil, ereasete, or synthetic oil of wintergreen and leave until clear.

4. Using Canada balsam or damar which has been heated in an oven for some days until it will harden immediately upon cooling, prepare several shdes by dropping such heated balsam upon them until drops 3 or 4 mm. deep and as wide as the cover-slip are formed. Let these harden. If small bubbles appear in such drops, place the slides in an incubator or a parafilm oven for some hours; the bubbles rise to the surface and may be skimmed off or burst with a fame.

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5. To mount a hadpole in such a drop, beat the drop by holding it inverted over a fame for a moment, then with a scalpel previously dipped in xybel make a groove in it of suitable size and shape to fit the tadpole. The sides of the groove should press against the object in such a way as to hold it in the desired position. Some tadpoles should be mounted to present a dorsal, others a lateral, aspect.

6. When the object is in place, fill in the space about it with a drop of soft balaam and gently but firmly press a heated cover-slip down upon the surface of the drop. Guard against including air bubbles.

 Examine the preparations after a week or more and, by pushing the cover-silp toward one side or the other, adjust any of the objects which may have shifted from the desired position.

MEMORANDA ON AMPHIBIAN MATERIAL

1. To Study Amphibian Eggs Entire, use a hand lens or dissecting microscope. Flue the eggs on a bit of absorbent cotton under 70 per cent alcohol in suit cellors. The eggs are fragile, consequently, to manipulate them, use a soft-hair pencil or a current from a pipette. Use the same egg for surface view and for sectioning when possible.

 Special Egg Pipettrs for handling definete objects should be prepared by breaking off the tip of an ordinary pipette to enlarge the ordine. After rounding up the broken edges in a flame, cover the broken end with a small piece of suft-robber tubing.

3. Amphibine Eggs in General may be fixed (in masses of 15 or 20) in Tellysainchy's fluid (5, p. 209) or in Woresster's aceto-formal-sublimate mixture (20), p. 214). Chromie acid (11, p. 211) brings out surface views well, but the material becomes very brittle and does not take stains readily. H surface views alone are desired, formalia-preserved material will answer.

4. To Remove the Gelations Costs of Eggs, rull them over and over on a bit of blotting paper. Either fresh or preserved eggs may be handled in this way. To prevent very saft eggs from during down and allering tightly to the blotting paper, roll them oft on to a paper of hardler testure just before the last trace of gelatinors film has been removed.

Whitman (American Naturalisi, XXII, 857) recommends putting the fixed eggs into a 10 per cent solution of sodum hypothetic diluted with 5 α 6 volumes of water and leaving them until they can be staken free. This requires only a few minutes. Rinse the eggs in 35 per cent should. It is advisable to remove the allouninous scats before handening in alsohol.

Child (Zeitschrift für wissenschaftliche Mikroskopie, XVII [1900], 205) states that the albumen which surrounds many ova becomes transparent and dissolves if after fixation (in any way except with chromic acid) the ova are passed up through the grades of alcohol to 80 per cent, hardened, and then passed down again through the alcohols into water which has been slightly modified with any acid except chromic.

5. Amphibian Eggs Are So Friable that they are ordinarily sectioned in celloidin. If they are cleared from 95 per cent alcohol (avoiding absolute) into cedar oil or oil of wintergreen they are less brittle. However, good sections are obtainable with the Johnson asphalt-rubber method (memorandum 9, p. 43) or by the more tedious parafin-celloidin method (memorandum 8, p. 64). Older embryos are readily sectioned in paraffin according to the method already given (p. 119).

6. Whole Eggs May Readily Be Cut into Halves with a safety-ranor blade. They are often more serviceable for study along with observations

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Ser.		1. Section

on the external changes of cleavage, blastula formation, gastrulation, etc., than the most elaborate serial sections. 7. The Germinal Layers are much

more distinctly seen in sections of young Fro. 41.-Method of Mount-ing Endryos (after B. M. Allen). the frog.

8. Very Thick Sections are sometimes useful. A 7 to 9 mm. tadpole, for example, cut into three sagittal sections is excellent for studying the general topography of organs. Also older tadpoles with the skin removed from one side before mounting are serviceable.

9. Eggs Mounted Entire or in Halves, according to Allen's gelatin method (Kansus University Bulletin, IX, No. 8 [December, 1914]), are very successfully studied as opaque objects. The method, also useful for preserving free-hand sections of various kinds, fine dissections, etc., is as follows:

a) Dissolve thymolin distilled water with the aid of heat until a saturated solution is obtained. Filter until clear.

b) Soak gelatin in the thymol water until it has absorbed all it can hold, then drain off the excess of water. The gelatin, now ready for melting, may be kept in corked test-tubes.

c) Prepare cells for mounting large objects by placing strips of glass upon a slide as indicated by the shading in the diagram (Fig. 41).

d) Liquefy some of the gelatin by placing a test-tube full into a warm water-bath, then pour it into the newly made cell.

e) When the gelatin has set, melt small areas in it with a hot needle and insert the objects (e.g., a series of frog eggs in different stages of segmentation) to be mounted. Each of these must be held in proper position with the needle until the gelatin solidifies,

f) Flood the cell with gelatin heated just enough to be a fiquid and place a slightly warmed side on top as a cover. Avoid air bubbles, and see that there is a complete film of gelatin between the cover-slide and the glass strips. The cover should be held firmly in place till the gelatin solidifies.

g) With a toothpick or similar object thoroughly clear out every trace of gelatin from the grooves formed by the projecting edges of the slides and the strips between them. Dry well.

b) Bun some emant such as gold size into the groove and set the proparation asile to harden. Add more emant from time to time and the groove is completely filled up. Keep the preparation out of direct sought or away from radiators, as it must not be subjected to heat.

If the objects have previously been hardened in formalin, so much the better, as the formalin will gradually diffuse out into the gelatin and harden it.

Hollow-ground slides designed for use with a hanging drop may be used if preferred. It is sometimes desirable to sublify the gelatin more rapidly after the object is mounted, by placing it on ice.

10. For Artificial Fecundation of Amphibian Eggs see 16, b, p. 135.

THE CHICK

A setting hen or an artificial incubator is necessary. In many ways the latter is more convenient as it may be kept in the laboratory and is ready at all seasons of the year. There are many kinds of good incubators on the market at present which may be had for a small sum.

Whatever method of inculation is employed, the eggs must be fresh and must not have been subjected to rough handling. The date and hour at which inculation is to begin should be written on the shell of each egg in ink. If late stages of development are desired, the egg must be turned every few days. All products of combustion from the lamp or burner should be kept from the eggs and the supply of fresh air and moisture earefully maintained. The temperature should be maintained at 38° C. (102° F.). Should it rise above 40° C., embryos will be destroyed.

Prepare at least 5 embryos as directed in the practical exercise, 2 for in toto preparations and 3 for sections.

 Place an egg which has been incubated for between 46 and 54 hours, while it is yet warm, in a vessel which contains sufficient normal saline warmed to 38°C. to cover the egg. In the ford the

embryo always makes its appearance as a germinal disk or ciostricula, as it is termed, situated on one side of the yolk, which is the real egg of the hen, the white being simply a natritive mass added in the oviduet. This disk or *blastolerw* in the early stages of ineuhation always turns uppernost no matter in what position the egg may be placed. Moreover, it has been found that the embryo in nearly every instance hes in such a position that when the blant end of the egg is toward the left, the head of the chick is directed away from the operator. This fact affords a very reliable means of orienting the embryo, especially in the very young stages when the anterior and posterior ends are not easily recognized by the observer.

Break through the shell at the broad end over the air chamber by tapping it sharply, and let out the air, or the broad end will tilt up.

3. Begin at the hole made in the end and with blunt forceps remove the shell and shell membrane bit by bit from the upper surface of the egg until the endryo oranes plainly into view. Remove with a pipette the thin layer of albumen which lies above the blastoderm.

4. With as little agistion of the liquid in the vessel as possible by means of fine scissors cut rapidly around the blastoderm well outside of the vascular area.

5. Carefully float the blastoderm into a thin watch-glass, keeping it as flat as possible. Shake it gently to remove the piece of vitelline membrane covering it, or any yolk which may adhere. The aid of a needle may be necessary to remove the vitelline overeing.

6. With a pipette remove all excess of fluid from the watch-glass but do not let the embryo become dry. In order to keep the elges from ourling up and obscuring the embryo, touch small, rectangular pieces of dry filter-paper to the blastoderm around its periphery (Miller's method: Anatomical Record, V, No. 8 [August, 1911]), and when they adhere use them for spreading the tissues flat. Stick the free ends of the paper strips to the bottom of the watch-glass to hold the membranes in this spread condition.

 Carefully add piero-sulphanie fazer (reagent 26, p. 217) until the embryo is completely immersed. The third should be allowed to act for from 2 to 3 hours.

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The paper strips may be left in place through subsequent treatment to 95 per cent alcohol for embryos which are to be sectioned, or to xylol for those which are to be mounted whole.

Norm—Some prefer to fir the embryo before removing it from the egg. After some of the abbumen is drawn off, the fining agent is spainted on to the blassoferm. As son as it is opaque the latter is then removed to a vessel containing the fining agent. Andrews (Zeitherbif) for viscourschiftliche Mitradopie, XXII (1994, 1971) injects picro-schiburic acid (1) between the vitedline membrane and the blastoferm and (2) between the blastoferm and the yolk by means of a pipette which has a fine upcurvel point. The blastoferm and the yolk by fired from the yolk. This operation should be performed before the egg has been subjected to the action of any reagents.

8. Wash in repeated changes of 70 per cent alcohol. Pass down through 50 and 35 per cent alcohol to water. Stain in alum-occhineal for 24 hours (Conklin's hematoxylin may be used if preferred).

9. Wash the object in water and transfer it through 35 and 50 per cent alcohol, leaving it 30 minutes in each. Decolorize the embryo slightly in weak acid alcohol, then wash in 70 per cent alcohol, and leave it there until ready to proceed.

10. Transfer the object through 95 per cent (1 hour), absolute alcohol (2 hours) to xylol, where it should remain about 2 hours or until it ceases to appear opaque.

Mount two embryos entire, one with the ventral, the other with the dorsal, side uppermost. Put bits of broken cover-glass or threads of glass under the edges of the cover to avoid erushing them.

The three remaining embryos are to be so sectioned (steps 11 ff.) that the student will have a complete series of sections in each of the three different planes of the body with reference to the axis of the spinal cord: viz., transverse, sugittal, and frontal. Read carefully memorandum 1 on orienting serial sections.

CATTION—Before sectioning any embryo always make an outline drawing of the entire embryo; then rule lines across the drawing parallel to the plane of section. Unless this is done great difficulty will be experienced frequently in understanding the sections.

 Infiltrate the embryo with parsfin in the usual manner by leaving it in melted parsfin for 2 or 3 hours. A parsfin melting at

about 45° C. should be employed and sections should be cut 20 to 30 microns thick.

12. Imbed and cut in the usual way (chap. y). Mount the entire series.

MEMORANDA

1. Directions for Orienting Sorial Sections.—o) In mounting broasense sections (sections across the main axis of the object), the sections beginning at the anterior and of the object are laid on the shole in the same sequence as the reading on the page of a book. In order to have right and left sides and dorsal and ventral surfaces in proper relation to the observer, mount the object in such a way that, in conting, the handle will enter it on the left sides and at the anterior end. Leave room at one end of the skide (see p. 49) for a label and also a small margin at the opposite side.

b) To get proper crientation of frontal sections (sections lengthrese of the object in a plane including right and left sides), arrange the object so that the hule will enter it on the right side and since of the doesnl surface first. Mount sections, with their posterior ends toward the upper edge of the side, placing the first section of the series to the left end of the upper row. This throws left and right, dorsal and ventral, into their proper position as viewed through the compound microscope, and the observe locks from the dorsal toward the ventral spect of the object.

c) To mount sopiilal serious (sections lengthesise of the object in a phase including ventral and doesal sides), arrange the object in such a position that the latter state ventral surface and slices off the right side first. Mount with the posterior end toward the upper edge of the side, placing the first section of the series at the left end of the upper row. Through the compound microscope the diserver views the object from the right toward the left. The head will appear to be toward the upper end of the slide, the doesal surface toward the left.

It is frequently advantageous to have the imbedding-mass trianment unsymmetrically by leaving the edge which first comes in contact with the huide larger than the opposite edge. One may thus readily discover if a section or part of a series has been accidentally turned over.

 Orientation of Objects in the Imbedding-Mass so that sections can be cut accurately in definite planes is trequently difficult to accomplish. The following methods are useful in meany instances:

I. For prooffs excison.—With a soft penel rule the strip of paper which is to be used for making the imbedding-box into small separes or restangles. After imbedding, upon removal of the paper a copy of the penel marks will be formal upon the block of parafilis. If the object has been arranged in the methed parafilm with reference to these lines, it is easy so to arrange the block in the microtome as to ent the object along any desired plane.

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It is frequently an aid to orientation by this method to have one of the central ruled lines broader than the others, or double.

Small objects which cannot conveniently be criteried in method parafilm may be properly oriented and fixed to a small strip of paper ruled as above, before they are placed in the parafilm both, by a mixture of clove ed and collidica of about the consistency of thick melasses, as in Pattor's method (Zeitzchrift far wisenschoffliche Mikrostopic, XI [1894], 13). One or a number of small objects which have previously been cleared in oil of bergrand or cloves are monthed in small separate droplets of the reagent and oriented under a dissecting lens with reference to the ruled lines. The paper is then placed in turpeatine which washes out the olor oil and fixes the object in place. The paper which objects attached is then passed through melted parafin and imbedded in the ordinary way. Upon removal of the paper from the hardneed block a sufficient number of pencil marks Patton employed ribbel paper.

II. For celloidin sections.-

Eycleshymer's Methods --- a) For imbedding, metal boxes made of two L's (Fig. 30) are used. The L's are held together by overlapping strips. The ends and sides of the box are periorated at regular intervals by small holes which have been drilled opposite one another in such a way that threads drawn through them are parallel. Threads of silk are run through the holes from side to side, drawn taut, and cemented to the outside of the box with a drop of celloidin. Each piece of thread should have an end two or three inches long hanging outside the box. A piece of heavy blotting paper is used as a bottom for the box. The object is oriented on the parallel threads and the imbedding-mass poured in and hardened. The loose ends of the threads are then soaked in a solution of thin celloidin which contains lamp-black, the celloidin drops holding the threads tant are dissolved by a drop of ether-alcohol, and the blackened ends are drawn through the block of celloidin. The lamp-black leaves distinct black lines through the mass which will serve for properly orienting the celloidin block on the microtome.

This method is valuable also in reconstructions from sections (see chap, xvii). In such work it is very desirable to establish "treamstruction points" to guide in fitting the wax plates together properly. The black rings of hamp-black left in the sections answer admirably for this partyses.

b) For small objects in which reconstruction points are not required Byelskymer uses fine insert pins from which the heads have been elipped and the headless ends loosely inserted in handles. The objects are momited on the points of the pins and ericated in the desired position. Each pin is then removed from its handle, and the free end is inserted from below into a small perforation which has been made by passing a somewhat larger pin.

lengthvise through a cock. A number of pins may be mounted on the same cock. To prevent the dojects from besoning dry, the cock must frequently be inserted into the month of a vial full of alcohol in such a way that the objects are immersed. If desired, the dojects may be sketched is advander alcohol by weighting the cock with lead and placing it in a beaker of alcohol. To pass the objects through the various goales of alcohol, etc., simply transfer the cock bearing them to successive vials of proper size containing the different fluids. For imbedding in celluidin uses the method given on p. 39, steps 2 ft. When the celluidin mass hardened, the paper is removed and the pins are drawn out through the cock, thus leaving the objects in place ready for sectioning.

3. In Measuring the Length of Embryos some embryologists (e.g., Minut) measure the greatest length of the embryo along a straight line (finds not included) when the embryo is in its normal attitude; consequently in some early stages where the embryo is greatly flaved the neek-lend would be the paint to which to measure instead of the tip of the head, because it is themost anterior regime; in stages where the embryo is straight, the head would be included. Other embryologists (e.g., His and German authors in general) make use embryologists (e.g., His and German authors in general) make use embryologists (e.g., His and German authors in general) make use embryologists (e.g., His and German authors in general) in the taxe of human embryos, use the so-called "sitting beight" and "standing height."

 For the Embryology of Teleosts the following are the most useful mounted stages:

I. Whole mounts—The 2, 4, 8, 16, 32, and 64-cell stages (only the blastodisk segments); early periblest; late periblest; early germ-ing; embryour shield; various stages of early embryos, such as embryos of 45, 50, and 60 hours.

II. Sotions (parafin)—Of 4, 16, and 32 cells (vertical sections parallel to the first place of charage); late charage (vertical sections); early, mid, and hat periblast (vertical sections); transverse and sagittal sections of early gene-ring, embryonic shield, early embryo, hate gene-ring, and closing of blastopore, respectively.

All stags may be fixed in piero-acetic (reagent 24, p. 216) or Bonin's fluid for 30 to 40 ninutes. The eggs are finally preserved in 83 per cent alcohol. Child finds that fixation for about a minute in 10 per cent acetic acid saturated with corresive sublimate, followed by 10 per cent formalin, gives good results without the yolk becoming hard. The owa of the Salmonidae must be removed (after fixing and hardwing) from their envelopes before the endryro can be studied.

Before the preserved material can be momented in *toto* or sectioned, the essential part (the blastcolerm) must ordinarily be dissected off under a dissecting lens by means of sharp needles. If the blastcolerus are to be

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moundel entire they may be passed down through the alcohol (see Walton's device, memoranhum 4, p. 30), stained in Conkin's hematoxylin (rengent 54, p. 225), then dehydrated and mounted in the usual way. To avoid crusting the objects, the cover glass should be supported by means of bits of broken cover or glass threads. Material which is to be sectioned may be stained in two or the sections may be stained on the Side. In the latter event, to facilitate orientation, it is measured to the glass the facility with Booleaux read or some other cytuplismic stain makes the fixing respect has already done so. For the same reason it is best to imbed the material in a wathelgas, arranging it near the bottom of the parafinemass so that one can see with a microscope how to shape the parafin block in order to cent sections in the proper plane. The immension in the melted parafin should not be longer than 5 or 10 minutes. The parafin is best howhened under 95 per cent alcohol. The sections may be stained by any of the hematorylin methods; iron-leanatorylin (5, 51) yields excellent results.

5. To Study Living Eggs of Telessts, a thin, flexible piece of sheet cellukid or mize should be used instead of a coree glass. The egg must be rotated from time to time, and this is easily accomplished with such a flexible corer.

6. For Artificial Fecundation of Teleost Eggs, see p. 135.

7. Thing the Microscope into a Horizontal Position and examining the egg in its normal medium by direct light is an encellent method of studying blackdisk formation in such forms as Coredebrus, for instance. Inasmuch as the blackdisk forms on the lower side of the egg it appears to be on top when viewed through the compound microscope.

8. To Preserve Teleost Eggs in Corresient Form for Demonstrating disorial elearage, embryonic shield, germ-ring, etc., Smith (Pronoutions of the American Microscopical Society, XXXIII, No. 1 [Journary, 1914]) scales pieces of j-incl glass tubing stone and by holding in a frame. A series of eggs find in corresive-active mixture and preserved in formalin is placed in each tubeard the opening plugged with outcom. The tube may be held in the hand and examined with a lens or dropped into a watch-glass filled with water and examined under a lens or binoreliar microscope.

 For the Average Course in Embryology of the Chick the following mounted stages are the most useful:

I. Montel in toto.-Approximately, 48 hours rivered from shore and below 36 a a a a a 30 a a a a 24 a a a a 18 a a a 18 a a a 12 a a a 64-72° a a a

96 hours (studied in alcohol under the dissecting microscope)

II. Sections.—

48 hours, transverse, sagittal, and frontal 36 " 4 4 30 " u 4 24 4 ų, 4 18 " 4 10 " W. 72^{-4} и и и и 96 " a 8

The number of embryos needed for the above-mentioned preparations is as follows:

> 5 embryos of 48 hours (71-29 sumites) 4 a a 36 a (15-18 a) 3 a a 30 a (10-14 a) 4 a a 24 a (4-6 a) 2 a a 18 a 1 a a 12 a 1 a a 60 a 3 a a 60 a 3 a a 30 a (10-14 a) 4 a a 24 b (4-6 a) 5 embryos (errival fleure formed) 3 a a 30 a (10-14 a) 6 a (10-14 a) 6 a (10-14 a) 7 a (10-14 a) 8 a (10-14 a) 8 a (10-14 a) 8 a (10-14 a) 9 a (10-14 a) 8 a (10-14 a)

10. To Mark Anterior and Posterior Ends of Young Chick Embryos in blastoderms which still have a homogeneous aspert, Duval's esnie-acid method is very useful. With a strip of paper 5 mm. wide by 50 mm. long a triangular bottomless box with narrow base is constructed. This is placed on the yolk inclosing the blastoderm in such a position that the base of the triangle corresponds to what will be the anterior region of the embryo (for orientation of embryo in the egg, see step 1 of the practical exercise). Press the box down against the yolk and fill it with a 0.3 per cent aqueous solution of osmic acid. In a short time the preparation begins to darken and the osmic acid should be removed. The blastoderm may then be removed in the ordinary manner and fixed as desired (Duval used chromic acid for fixing). However, it is very difficult to separate the blastoderm from the egg during the first 24 hours of incubation, and it is advisable, therefore, to fix and harden both together and to remove the blastoderm later (see note under 7, p. 125). The blackened area affords a convenient means of orienting the preparation for sectioning.

11. For the Stages of Maturation, Pertilization, and Segmentation in Mammals while mise will prove most useful because these processes are better known in them than in other mammals; furthermore, an abundance of material may be proceed. The ovan, however, is extremely small, mesaming only about 50 micross in disarder. It is surrounded by a very thin area pelluvids (1.2 micross). Long and Mark find a modified Zenker's full the most satisfactory for function. They make up two solutions: (1) a

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4 per cent aqueous solution of polassium bichromate; (2) a 4 per cent aqueous solution of corresive solutimate and 20 per cent acetic acid. The two solutions are mixed in equal proportions when needed for fixing.

Fenale mise are in heat soon after parturition. They tend to orulate every 21 days during the spring months. The maturation process requires from 4 to 15 hours and usually occurs between 13 and 29 hours after parturition. The second polar spindle usually forms immediately before orulation, but the second polar body may not be extruded unless the egg is fertilized. Orulation occurs between 14 and 29 hours after parturition. The eggs are easily visible at first in a fold of the ordinet sear the orary. Insemination is most successful when it occurs between 15 and 30 hours after parturition. The spermatoona reach the eggs in the upper end of the Fallopian tube in from 4 to 7 hours.

For details and hibitography see (1) Kritham, Biological Balletia, XII, No. 4 (1907); also Transactions Connectical Academy of Aris and Science, XIII; (2) Long and Mack, "The Maturation of the Egg of the Mouse," Publications of the Carnegic Institute of Washington, D.C., 1911; also, "The Living Eggs of Bats and Mite with a Description of Apparatus for Othiming and Observing Them," University of Colifernia Publications in Zology, IX, No. 3 (Feb. 23, 1912); (3) Dainels, "Mice, Their Breeding and Rearing for Scientific Purposes," American Naturalist, October, 1912; (4) see also Danforth, Anatomical Record, X, No. 4 (February, 1916), for some very practical suggestions regarding use in classes.

The phenomena of naturation and fertilization in the altino rat are desenthed in a paper by Solotta and Burchlandt in Asatonische Hele, XLII, 1911 (summarised in Huber's paper cited on p. 133). See also Krisham and Bur, "The Breeding Habris, Maturation of the Eggs and Ovulation of the Ahtino Rat," American Journal of Anatomy, XV, 1918.

12. For Early Stages of the Mammalian Entryp relidits are commonly employed because they breed readily, especially in the syring of the year, and the observer can note the exact time when the female is covered if she has been kept separate from the book until she comes into heat. The period of gestation is 30 drys and imporpation takes place again immeliately after litering. The two uteri of neabbit diverge as two anterine hours from the single median ranging and each terminates in front in a narrow, colled tube, theorishnet or Fallopian tole. To obtain the early stages the abdomen is slit open from publis to steman, the intestinal tract is cut oray or publied to one side, and each terms and oxidute cardially removed and stretched out along a glass plate. The segmenting one are found in the oxidute up to nearly 70 hours from the time of copulation. After that period of time they must be looked for in the uteres. Formulation takes place about 9 hours after colding. Which in the oxidutet, with the aid of a lens they may sometimes become through its wells. A segmenting orum once located, a transverse

ent is made to one side of it through the wall of the oriduct, and the orum, which is very small, is greatly squeezed out by compressing the ordnet behind it. With a spar-banded meelle or the point of a scalpel the orum is conveyed to the funing fluid. In case segmenting oru are not visible from the exterior of the ordnet, the latter must be slit open carefully with a pair of fine-pointed subsers, and the eggs sought for by means of a lens. In case, no red corpore latten are visible on the surface of the orany, indicating a recent discharge of one from the Gradian follows, for the search is useless.

Bablit ora of 18 hours show 4 blastmeres; 36 to 18 hours, advanced segmentation; 72 hours (about 0.6 mm. in diameter; in anterior end of uterus) show the fully segmented orum—an outer layer of clear, endical cells, an inner mass of irregular granular cells; 72 to 00 hours show enlarged blastodermic vesicle and establishment of embryonic area; 5th and 6th days (0.8 to 4 mm.) show geminal layers; 7th day, primitive streak; 8th day, melullary folds.

The earlier stages (up to 70 hours) may be fixed for from 5 to 8 minutes in a 0.3 per cent aqueous solution of comic acid, stained in piorw-earning, and transferred to a minimum of givenia and water, equal parts. They should extrain in this fluid for a week nucler a kell-jur so that the water gradually eraporates. The object may them be mounted in formir-giverin (formine acid 1 part, giverin 99 parts). To avoid pressure of the cover-glass, the object should be mounted in a cell or between two slips of paper or pieces of cover-glass. If the preparation is to be permanent the cover-glass should be scaled (see p. 95).

To render the cell outlines distinct, stages of from 70 to 50 hours are best treated, after rinsing in distilled water, with a 1 per cent aqueous solution of silven nitrate for 3 minutes and then expected to light in a dish of distilled water multi they become known. They are then treated with water and glycein and mounted in fremine glycein as in the case of younger stages.

For sections the embryos should be placed in Bouin's or Zenker's fluid for one or two hours, then washed in the customary way for these methods, stained in alam-cordineal, and sectioned in parafilm.

In opening the uterus, the invision should always be made along the middle of the free side, opposite the insertion of the peritoneal fold, because this line of insertion marks the region of attachment of the endryo within the oriduct. By the 7th or 8th day the developing ova have taken up positions at intervals along the inner walls of the uterus and have become so firmly attached to the unerous membrane that they can no longer be detached unmultisted. For further particulars regarding the embryology of the rabit, the reader is referred to E. Van Benelen and Charles Julin's "Recherches our la formation des amenes foetals chen les mannifers," Arbino de Biologi, V (1884), 378. See sko Assheton, "A Reinvestigation

into the Early Stages of the Development of the Babbit," *Quarterly Journal Microscopical Science*, XXXVII (1995).

With the skit of Huber's paper (Journal of Marphology, XXVI, No. 2 [1915]; sko Menoira of the Wistar Institute of Automy and Biology, No. 3), which ovvers the development of the allow rat from the promother stage to the end of the 9th day, it is now feasible to use the rat for early embryonic stages. Huber found Carnoy's fining fluid (reagent 2, 0, p. 207) the most satisfactory. He fixed issues for several hours, washed them in serveral changes of absolute alcohol, and stored them in the latter. He found that sectioning orary, oviduet, and uterus on more was more satisfactory than isolating and sectioning the separate ora, aldough he used both methods. For staining he employed moinly hemalum, followed by Congo red. His methods are described at some length in the paper.

13. For Older Stages of the Mammalian Endryto pig endryts are commonly employed. They may often be protocol in large numbers and with little trouble at the larger port-posing establishments. The most valuable stage for study is an endryto of from 10 to 13 mm, in length. In most laboratories it is reasonary to make a detailed study of an endryto of about this stage and then a more general survey of both smaller and larger sizes.

Early stages are much more difficult to obtain than advanced stages. Endryss of 6 mm, length and over may usually be readily located by the endrygements which they cause in the uterine walls. The others should be handled carefully and opened as soon as possible. The endryo is best removed by means of fine forceps and a horn spoon. It is very delicate and should not be handled roughly. The chances are that in removing the embryo the membranes will be ruptured and the annihole and shouth fluids will escape. Larger endryos should here the body eavily punctured to should the line fluid.

Submerge the embryo without removing the membranes in a bountiful apply of Kleinenberg's picro-subhanic acid (respect 29, p. 217), moving it about gently to rines off any congulum that may form on the surface. Lardwersky's fluid (19, p. 214) is also a good fixing agent for pig embryos and is to be preferred for the other ones.

Leave embryos of 6 to 9 mm. 24 hours; 12 to 15 mm., 4 hours; 20 to 25 mm., 6 to 8 hours.

For washing and subsequent treatment see reagent 26, p. 217. Embryos may be stained in toto in alum-tochineal or borax-carmine.

For studying the uterus, placentation (diffuse in the pig), and embryonic membranes in place, formalm-hardened material may be used after first thoroughly washing it in water.

For gross dissection of embryos the specimen should be studied in alcohol under the dissecting microscope.

Because of the asymmetry of young embryos it is impossible to secure strictly towarce, ouglid, and found sections. Minot resummends, there, fore, that for practical purposes the plane of section be taken with regard to the head above invespective of hour it may set the other parts of the body, and suggests the floor of the fourth ventricle of the levin as the guide for orientation. In his Laboratory Tort-Book of Entryology the especially recommends that each student prepare sections of the following stages of pig embryos: 9 mm, transverse and sugittal, frontal of the head; 6 mm, transverse, frontal of the head; 17 mm, transverse and sugittal, frontal of the head; 20 mm, transverse and sugittal, frontal of the head; 24 mm, frontal of the head.

14. Human Embryos of all ages are very valuable material for scientific purposes. Physicans and surgeons are urged to preserve such material properly and turn it over to some competent embryologist. Very young human embryos are enceedingly distrable. Fill the containing vessel completely with fluid in order to avoid staking.

An excellent fixing reagent, the ingredients of which a physician can usually readily provine, is Lawlowsky's mixture (reagent 19, p. 214). The embryo should remain in this fixial from 1210 Shours, succeiling to size, and then be preserved in 80 per cent solvhol (or enumerical solvhol to which has been added about one-fifth its volume of distilled water). Use a widemonthed bottle with tightly fitting stopper.

Zenker's fluid (reagent 6, p. 200) is better for larger-sized embryon. Material should be left in it from 15 hours to several days. For washing and preserving follow the directions given under the description of the fluid. For feluxes use a bruit-jur of such a size that the embryo can be kept in about 10 times its volume of fluid.

In case the above-mentioned finits are not available, the material may be placed in 10 per cert formalin (1 part of commercial formalin to 9 parts of distitled voter) and left indefinitely. As a last resert, if no other fining reagent is available, the embryo may be placed in the strongest alcohol which can be secured and later transferred to 80 per cent alcohol for preservation.

The specimen should not be handled nor allowed to be in water. When the proper reagents are not at hand, excefully wrap the object in cloth and keep it on ice if possible until they can be secured. Very small embryos may be fixed and preserved with membranes ruptured. To secure the best function of fetures (2 months and beyond), the specimen should be diriched, or at least the body enviry should be opened.

15. For Mice-Dissection of Small Embryos, effort frustion Heuser stains for 24 hours in alum-cochineal diluted with 5 times its volume of water. He then fixes the early to a small square of thin ground glass with ediolith ement (alour) 0.75 per cent solution of ediolith) and dissects in alcohol

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under the bioscular microscope. The dissected specimen may be preserved at any stage by placing it, still attached to its gloss support, in alcohed in a shell vial of suitable size. Necessary data may be written with a penel on the ground surface.

Steedar (American Journal of Anatomy, IV, No. 1, p. 87), stire dehydrating in alcohute alcohot stackes the embryo with a drop of thick cellodin to a smaked kinglass strip coated with thin cellodin and places it in 80 per cent alcohol. The black serves as a good background for the embryo and may be written upon. During dissection the kinglass strip is clamped to a gass side which has been cenceried with balsam to one faces of a set-glass polyhedral paper-weight. The object can thus be placed in any desired plane and dissected in alcohol under a binerska microscope.

10. Artificial Perudation when it can be practiced is the must convenient means of securing early stages of development. This is possible with many worms, coelenterates, echinodrums, cyclostanes, teleosts, and annua amphilia.

a) In exhibitering (e.g., sea urchin) the fenck is out open and a number of the firing eggs transferred to a watch-glass which contains firsh sea watcr. The testes of a muck are tessed out in sea watcr and a drop of the mixture is conveyed by means of a pipotie into the disk containing eggs. Inmediately upon fertilization a membrane forms around each fertilized egg. In about 90 to 30 minutes after fertilization the signs of the first charage should appear. The blasting forms in about 60 hours, and the gastruk in about 90 to 50 minutes after fertilization, etc., the following stages should appear. The blasting forms in about 60 hours, and the gastruk in about 10 hours. For the study of fertilization, etc., the following stages should be fixed in Bounds fluid (p. 29) for 30 minutes and stained in iron-hematorylin (p. 51); 5 minutes after fertilization, nucleus giving off polar bolies; 30 minutes after fertilization, approaching promotely; 30 to 55 minutes after fertilization, approaching ponnolei; 30 to 55 minutes after genes (mitotic figure) in the first charage.

b) In emphasize (e.g., freg) both male and female are cut open, the rasa deferentia or testes are tensed out in a watch-glass full of water, and the ora are then removed from the lower ends of the orbitest and placed in this water. After fertilization the eggs should he placed in glass disks in not over 4 inches of water. Many eggs should not be placed in one disk. See also memoranda pp. 121-123.

c) In bloots the eggs are obtained by stripping the female when she is in sparning condition. At such times the eggs are loose in the body cavity and may be pressed out by gettly manipulating the belly of the fish. The head of the fish should be bid in one hand, the tail in the other, and the themh or the thumb and forefinger used to press out the ova into a chan, dry finger-loosel. The milt of the male is obtained in the same manner in a dish containing a little firsh or sea water, depending upon the habitat of the fish. When the water becomes milky with sperm pour the mixture over the eggs. Eggs and sperm are then gently stirred about by means of a feather

to insure through mixing. However, in some telessis (e.g., stickleback, Fundulus) it is necessary to kill the nade and tease out the testes. In the examer (Chendahrus) 10 minutes after fertilization the formation of blastodisk and polar bodies may be observed; 30 to 33 minutes after fertilization the two promobil may be found in close approximation.

If other than the very early stages are required, the fertilined eggs must be transferred to a hoteling-box or jar, depending upon the kind of egg. This is best done by means of a horn spoon and a feather. Dead eggs, recognizable by their opacity, should be removed at least once a day. The conditions under which the eggs of different species theirs are so varied that the reader must be referred for details to such special publications as those of the United States Bureau of Fisheries or the fish commissions of the various states.

17. For the Study of Early Clearage in Living Material the eggs of some of the water scale safered an abundance of encodent material. By watching asparine which contain scales the fresh material can easily be obtained during the spring and summer. Twigs and bits of based to which the egg-masses may be stretched should be placed in the organia.

If one is at the seashere, the sea urchin, starksh, squid, various marine annebis, moltoxis, and evelenterates afford an abundance of material. Of the marine fishes Fundulus and Ctendelorus are excellent. Of feesh-water fishes the whitefish (spanning in November or Desember) and the packerel (m April) show elserage well, although in the whitefish it is very show.

18. Por Quick Preparation of Clearage Stages for study in 606, in forms where there is considerable yold, Sparth finds useful often a miniture of equal parts of global acetic axid, glycerin, and water, to which enough Delabeld's hematoryfin is added to make it a light tan color.

 Chinese Black added to the water on the slide in which eggs with very transporent jelly (e.g., Nervis) are being examined outlines the egg distinct y and shows the path of the spermatoroon through the jelly.

20. For the Study of the Formation of Polar Bodies, Fertilization, and Early Charage in Sections radius surpasses the eggs of Assaris. The Assaris (1. magalorephalo) from the hore is preferable, addough A. hubbioide from the pig will answer.

The orisaes, two in number, are very long convoluted tubes. Diffeent regions contain eggs in different stages of development. The thicker tubes toward the anterior end of the animal contain clearage stages; back of these are cells showing extrusion of the polar bodies and fertilization stages. The material must be fresh; either hring the live Ascar's to the hovestory or take the firing fluid to the place to obtaining the material. Sit open the abdominal wall of the worm and remove the orisans and after separating the runnerous convolutions somewhat, fix them entice for 24 hours in a mixture of absolute should 4 parts, glacial acetie sci1, 1 part, or for

Is to 25 minutes in actival odd-chlordorm (reagent 2, b, p. 307) saturated with cornsive sublimate. Preserve in 80 per cent alcohol. To locateegys of the desired stage tease out egys at intervals along the ovisors, stain with ach camime (reagent 38, p. 222), and examine. The proper region corelocated, out out small lengths of the table, inheid? It in parafilm, and make thin transverse sections. In order to keep the egys from shriveling, the bath in hot parafilm must be curtailed. Use the method for debrate objects (p. 53). Stain by the iror-hematorylin method (p. 51). Accuris eggs when smaared on a slide in thick albumen frastive, which is then coagulated with formalin, will go on developing if put into an involutor.

21. The Cultivation of Removed Endrymic Tissues in elotted lymph, plasma, netrient agar, bouillen, and in various suit solutions is an important phase of embryological technique which has been developed largely in the past ten years, but the subject is too extensive to treat in detail in limited space. A general method is given and the reader is left to look up molifications and other methods in the papers listed at the end of this memorandum.

All dissections must be carried on under asceptic conditions. Sterilize all instruments, pipettes, shides, covers, and vaseline in a Bausen fame. For entirution of tissues use a Locke's solution to which destrin has been added. It is made as follows:

To 100 e.c of distilled water add:

NaCl	0.900 per cent
CaCl ₁ ,	0.025 per cent
KCl	0.042 per cent
NaHCO ₂	
Dextrin	0.250 per cent

Renove an S- or 10-day click embryo, under aseptic conditions, to about 10 or 20 cc, of the sterilized solution heated to 30° C. Cut out bits of intestine, kidney, live, heart, or spleen a few millimeters in diameter and place into another disk which contains 10 to 20 cc, of the solution at 30° C. Cut each stall piece up into smaller pieces a fraction of a millimeter thick. Draw these up into a sterilized flue pipette, one at a time, with some of the solution and make hanging-drop preparations (Fig. 40) of them on sterilize ourse-dips which are thoroughly clean and free from every trate of grease. Invert each cover-stip on to a raseline-ringed, hollow-ground slike which has been sterilized. For rings use a vaseline melting at about 45° C. A bit of parafin may be added to ordinary vaseline to stiffen it.

Incluste the exharts at about 30° to 40° C. Growth begins within 10 to 30 hours and, as indicated by the number of mitotic figures, reaches its maximum on the second or third day. When it is desired to examine the living tissue do so on a warm stage. The margins of the growing regions are the best points to examine because these the cells are only one or two layes thick. (Method of Lewis and Lewis)

When permanent preparations are desired the corre-slip is removed from the vaseline fing and the film of tissue is fixed, on the cover-slip, by means of comiscial vapor. After fraction the denser contral piece of tissue is torn savey, leaving only the thin film of new growth which is treated on the cover-slip as one treats sections on a slide. Sain in inc-hematoryfin or in Earlich's hematoryfin and exain.

Further details of tissue eulture in ratio and hiblographics will be found in the following papers: Harrison, Anatomical Record, I (1907); Journal of Experimental Zology, IX (1910); Carrel and Buryons, Journal of Experimental Medicine, XIII (1911); Lewis and Lewis, Johns Hopkins Hospital Bulletin, XXII (April, 1911); 241; Anatomical Record, VI, Nos. 1 and 5 (1912); American Journal of Anatomy, XVII, No. 3 (March, 1915); Anatomic cal Record, X, No. 4 (February, 1916).

22. The Living Embryo of the Chick may be kept under observation for some hours while still in the egg by employing one of the so-called "window" methods. The simplest method is to cut out a disk of the shell on one side under as mently ascritic conditions as possible, so that the embryo is exposed. A bit of the white is removed and a film of celloidin is hold over the opening to form the window. It must adhree firmly at every point around the margin. The embryo will continue to develop for some time if the egg is put back into the inculator.

CHAPTER XVII

SOME CYTOLOGICAL METHODS

In the very many cytological methods which have been in vogue during the past few years two fixing fluids, Flemming's strong and Bonin's, and two stains, iron-hematoxylin and safranin, stand preeminent as of general utility. Iron-hematoxylin with or without a conntenstain may be used successfully after either of the fixing fluids mentioned. The safranin is more likely to prove successful after Flemming's, although in some materials good preparations can be made with it after Bonin's fluid. Two other fixing fluids, Gilson's mercuro-nitrie and the acetic-sublimate mixture of Cannoy and Lebran, are also of wide application, especially when followed by iron-hematoxylin as a stain.

A resent method of fixation developed by Dr. Ears Allen gives promise of equaling if not surpassing any of the foregoing, particularly in the study of mammalian tissues. The method is given on p. 149.

In preparing tissues for cytological work it is imperative that pieces should be small, not more than 3 or 4 mm. thick where practicable, to insure thorough and even fixation. To severe prenetation from all sides, it is well to place a few layers of filter-paper in the bottom of the vessels in which tissues are fixed and to shake the tissues about a little from time to time. Mechanical injury may be avoided by binding a bit of clean linen on the ends of the forceps with which tissues are handled.

I. MITOSIS

For general study of cell structures, and particularly cell division, I have found nothing which can readily be obtained in quantities sufficient for class use that surgasses the crayfish testis; the blastolisk of the whitefish, the epidemis and testis of Ambystoma and Necturus, and the maturation and clarage stages of Assaris. Assaris material has already been discussed (p. 136).

Robertson (Journal of Marphology, XVII [June, 1916]) finds that in grasshoppers of the family Tettigidae, taken before the last moult, cells dividing nitrotically may be found in large numbers in the mesenteron, proctoclasma, fat-body, hypodernis, and the follicles of the gonals. The columnar epithelium of the mesenteron seems to be the most favorable region for finding such divisions. Inasmuch as the members of this family have only thirteen or fourteen chromosomes, the material should prove to be exceptionally valuable for purposes of class demonstration

Testis of Crayfish

The testicular cells of the erayfish (Candorna virilia) will be found in active proliferation from the middle of June to the middle of July. The chromosomes are too small and too numerous for satisfactory individual study, but the spindles and centrosomes are distinct and the general pictures of representative stages are clear-cut and easily found.

Section Method.—1. Fir small bits of the testes, 3 to 4 mm. thick, in Flemming's fluid (12, p. 211) for 24 hours. Wash in running water 6 to 12 hours, dehydrate, and indeel in paraffin according to the methods for deficate objects (p. 33) or the "drop" method (6, p. 152).

 Cut sections 5 to 7 microns thick and mount several slides by the water-albumen method.

3. After removing the parafilm with xylel and running the slides down through the slooks, stain some of the sections by the ironhematoxylin method (p. 51) and counterstain with acid fucksin or orange G. Bun the slides up through the alcohols, clear in xylel, and mount in a thin balsam. Use a No. 1 cover-slip if the preparation is to be studied with an cell-immersion lens.

4. Place others of the slides in safranin (72, p. 244) for 24 hours, then rinse in water and run up through the alcohols to 95 per cent. Counterstain for 30 seconds in a 0.5 per cent solution of light green (Liddyrin S.F.) in 95 per cent alcohol. If the sections are left too long in the green stain the safranin will be washed out. Pass the slides through alsohote alcohol into clove oil for a few minutes, rinse in xylol, and mount in thin halsam under a No. 1 over-slip.

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Smear Preparations.—Remore the fresh testis to a slide and tease somewhat with needles in order to rupture the cysts which inclose the germ cells. Spread the mass evenly over the slide with the end of another slide and then fastten it between the two slides in a very thin film. Avoid any considerable pressure. Separate the slide by slipping them apart. Each should hear a very thin coating of the material. Plunge them into Flemming's fluid and leave for 24 hours. Wash in running water for 6 to 12 hours, then with forceps pick or sense off all humps of tissue which might later keep the over-slip from fitting closely. Stain and momt as if the films were sections.

If preferred, some of the shifts can be fixed in Bouin's fluid for an hour or two, washed in 50 per cast alcohol, then stained in ironhematorylin and counterstained in acid furths in or orange G.

Blastodisk of Whitefish (Coregonus)

 Spawn the females and fertilize the eggs (in early December) as directed in memorandum 16, p. 135.

2. Select eggs in the 32- to 64-cell stage of cleavage (40 to 60 hours after fertilization) and fix for 6 or 8 hours in Bouin's fluid.

 Wash in repeated changes of 50 per cent alcohol, then in 70 per cent alcohol until the yellow color ccases to come from the eggs.

 With needles carefully dissect off the blastodisks under a binocular or other dissecting microscope.

 Dehydrate, section in parafin (method, p. 36), stain in ironhematoxylin with or without a counterstain, and mount as usual. Sections should be about 7 micross thick.

The eggs of the pickerel, obtainable in April, may be handled with equal case. They cleave much more rapidly than do those of the whitefish.

Testis of Necturus

The cells of the testes will ordinarily be found undergoing rapid problemations in late July and early August. Those toward the posterior end of the testis show the most advanced stages of spermatogenesis, those toward the anterior end the least advanced stages.

Both cells and chromosomes are very large. The spindle usually shows up well and the chromosomes exhibit considerable variety in shape and size.

1. From different regions of the testis fix some bits of testis in Bouin's fluid (6 to 8 hours) and others in Flemming's (24 to 36 hours). Wash out the Bouin as in 3, p. 141, and the Flemming according to directions in step 1, p. 140.

2. Dehydrate, imbed, and section according to the usual paraffin method.

Iron-Hematoxylin Preparations .--- 3. Stain sections of each kind of material according to the ordinary iron-hematoxylin method with or without a counterstain (pp. 51-52.).

Safranin-Gentian-Violet Preparations .--- 4. Also stain some of the Flemming material according to the safranin and gentian-violet method (73, p. 234).

Safranin-Gentian-Orange Preparations .-- Use saturated aqueous solutions of safranin, gentian violet, and orange G respectively. Rinse and stain in gentian violet 2 to 5 minutes (time determined by trial). Pipette absolute alcohol over the sections until the violet is out of the cytoplasm, then follow with orange G, pipetting it on and removing it again almost instantly. Wash off with absolute alcohol, dip in oil of cloves, clear in xylol, and mount in thin balsam under a No. 1 cover.

Somatic Cells of Ambystoma

Epidermal Cells .- Cut off the tails of several one-month-old Ambystoma larvae into Flemming's fluid. At the end of 2 to 4 hours strip off bits of the epidermis from the tails and fix these strips for some 20 hours longer in the fluid. Wash in running water 6 to 8 hours, stain some according to the iron-hematoxylin method (p. 51) and others with safranin and light green (step 4, p. 140). Dehydrate and mount as usual.

If Bouin's is used instead of Flemming's fluid, the peeling off of the epidermis need not be done until the end of fixation (6 to 8 hours). Peritoneal Cells .- Parmenter recommends larger larvae than

those used for epidermis. He cuts away the side walls of the body

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eavity, pulls out the intestine, and fixes the remaining tissue in the region of the spinal column in Flemming's or in Borin's fluid as above. Bits of the peritoneum on either side of the dorsal nubline are stripped off and prepared as was the epidermis.

Either of these kinds of preparations shows splendid polar views of cell-division stages, though little or nothing of lateral views. They are especially favorable for showing longitudinal splitting of chromosomes.

Living Cells.--Cursive Ambystoms or other young samphibian larvae by adding (seconding to size of larva) 5 to 10 drops of a 0.5 per cent solution of cursare in equal parts of glyverin and water to a watch-glassful of water. After 40 minutes remove for half an hour to a 1 per cent solution of solium chloride in water. Wrap in histing paper and examine the tail fin on a sible under the microscope. Cell divisions may be seen in progress.

If replaced in fresh water such larvae recover after some hours. After cuarinsion some workers prefer to cut the larva in two parts in front of the hind limbs, studying only the tail. The gills also show interesting cell activities. If cursre is not at hand a 3 per cent alcohol or ether may be used, although not so successfully.

II. MITOCHONDRIA

Recent studies tend to show that mitochondria occur more or less extensively in nearly all kinds of tissue. They were largely overlocked in the past because many of the fixing fluids in use contain strong organic acids, such as acetic, and these dissolve mitochondria. They are sometimes stained with great sharpness by iron-hematoxylin, following fixation in Flemming's strong mixture (12, p. 211) in which, instead of 1 c.c., only 3 to 6 drops of glacial agetic acid are used for every 15 c.c. of chromic acid. For their careful study, however, cytologists are using special methods. These are too numerous and complex to be reviewed in an elementary guide to technique. Bibliographies and discussion of the technical details will be found in the publications of Bensley and particularly of Cowdry. See Bensley, American Journal of Anatomy, XII (1911), 297-388; Condry, Internationale Monatsschrift für Anatomie und Physiologie, XXIX (1912); American Journal of Anatomy, XVII, No. 1 (November, 1914); ibid., XIX, No. 3 (May, 1916); Contributions to Embryology, No. 11, Carnegie Institution of Washington,

Three methods of wide application are as follows: Bende's Method.—1. For for eight days in a modified Flemming fluid made as follows:

Chromie acid, 1 per cent.	15 e.c.
Osmic aeid, 2 per cent	4 e.e.
Glavial acetic acid	3 drops

2. Wash in water for 1 hour, then for 24 hours in a mixture of 100 parts pyroligneous acid and 1 part chromic acid.

 Transfer to a 2 per cent potassium bichromate solution for 24 hours, run up through the grades of alcohol to xylol, and finally infiltrate with parafin and section. Sections should be about 5 microns thick.

 After removal of parafin from sections run them down to distilled water and place them in a 4 per cent iron-alum solution for 24 hours.

5. Wesh thoroughly in water and transfer to a solution of Kahlkauni's sulplainarinete of soda (made by taking 1 part of a saturated express solution of the stain to from 80 to 100 parts of distilled water) for 24 homs.

6. Raws the slide in distilled water and flood it with a crystal violet aulin-rater solution (equal parts of anilm water and a 3 per cent solution of the dye in 16 per cent solution). Warm until the solution steams, keeping it heated for about 3 minutes.

 Wash in distilled water, transfer to 30 per cent acetic acid for 1 or 2 minutes, then wash in running water for 5 or 10 minutes.

 Dry the slide with filter paper, dip it for a minute into absolute alcohol, place in oil of bergamot until eleared, then transfer it through xylod and mount in balsam in the usual way.

A successful preparation should show chromatic elements a deep purple and the cytoplasm a light red with mitochondria violet.

Wilman (Journal of Marphday, XXIV, No. 3 [1013] modifies the method by transferring slides from the alian'n solution, after rissing, into a 3 per cent solution of crystal violet (3 c.e. of anim stain in 100 c.e. of distilled water) for 10 minutes; rinsing and passing into 80 per cent alcohol for 5 seconds; passing through 55 per cent and alsofute alcohol; and, when properly differentiated, elearing and mounting in the usual way.

Bensley's Acid-Fuchsin, Methyl-Green Methods.--Fix tissues for 24 hours in the following:

Osmie acid, 2 per cent	2 e.e.
Potassium biehromate, 2.5 per cent	8 e.e.
Glacial scetic acid	1 drop

Sections should be 4 microns or less in thickness. Mount by the water method (p. 23), remove parafin with tolood, then pass through absolute alsohol to water. Treat for from 30 seconds to 1 minute (determined by

trial) with a 1 per cent solution of potaxium permanganate, then for the same length of time with a 5 per cent solution of onale acid. The permanganate extracts the morelanting elements of fraction and the onalle acid removes the permanganate. Thoroughly wash in water.

Stain for 5 minutes in Altmant's will feelsin (acid fuckin: 20 grans, aniin water 100 e.c.) which has previously been warmed to 60° C. Wash theroughly in distilled water, dip for an instant into a 1 per cent solution of methyl green, then wash, rapidly delythate in alsochute alcohol (avoiding alcohols of intermediate strengths), clear in tothol, and mount in balsam. Tohinin blue can be abstituted for methyl green.

If the material does not stain well with the and fuelsin, or if the methyl green or toluidin blue obliterates it, treat the sections with a 2.5 per cent superus solution of potassium bieleromate for shout half a minute and rinse in water just before staining in and furthsin.

In spinal ganglion cells, for example, mitochondria should appear bright red; Nissl substance, green (or blue); neurofibrils in the anon kalleek, light brown; and the canalirular systems should be revealed.

Bensley's Copper-Chrome-Hennsteryfin Method.—Fire materials in asetic-semic-bichemante mixture and prepare for staining as in the preceding method. Wash for I hour in distilled water, then thercoughly dehydrate. Leave in absolute alcohol (1 hour) into pure hergemot oil for 3 hours, followed by equal parts of bergamot oil and pantifin (1 hour), then by parafin mething at 60° C. (2 to 3 hours). Out sections 4 microas thick and fir to the slinks by the water method. Remove parafin with toked and pass down through the alcohols to distilled water.

Place sections for 5 minutes in a saturated appears solution of copper acetate, wash in several changes of water, and transfer for a minute to a 0.5 per cent appears solution of a well-inpend hematorylin. Wash in water and transfer for 1 minute to a Sper cent appears solution of neutral potassium elevante. The sections should turn a blue-black color. If they are douby a light-blue should, place them again in the copper acetate and repeat the operations from there on.

Wash several minutes in water, then differentiate under the microscope in Weigert's borar-ferrityanide miniture (boras, 2 parts; ferrityanide of potassium, 2.5 parts; water, 200 parts) dibutel with 2 volumes of water. Wash 6 to 8 hours in tap water, then dehythete, elser in tobul, and mount in balsam. Ditorbordrin should appear a blush black against a clear background.

Mitschondria in Living Cells stain specificulty with James green. There are several James greens, but of these Condry reports that only James green B (dischyschrainnanelimethylandin ethorike) of the Fartbwerke Horehst Co. (obtainable from L. A. Mett & Co., New York) will give the desired

reaction. He finds that mittochoodnia will stain in human lymphocytes in a dilution of Janus green in normal saline solution of 1:300,000. Ordinarily, for bring tissue, a dilution of 1:15,000 or 1:20,000 in normal saline, in Lock's solution, or in Ringer's solution, is employed. Janus blue, G and R, may also be used as a vital stain for mitochoodia.

While firsh tissues may be stained by innersion in the dye, much better results are obtained by injection through the blood vessels, in normal saline, after the vessels have been thoroughly fitsched out with normal saline solution.

Matchandra in Tissue Grown "In Vitro" (see p. 137) may be studied reality, according to M. R. and W. H. Levis (American Journal of Anatomy, XVII, No. 8 (1915)), who observed their changes, growth, and division in embryonic tissues of the chick.

III. STAINING OF LIVING OR FRESH TISSUES

Into-viton staining, so called, has come more and more into prominence during the past fer years. It is questionable if a staining of really "vitol" elements ever occurs, although undoubtedly various grantlas in cells may be stained while the cells are yet living. Certain stains also may be used with fair success with fresh or lightly fined cells.

For inter-sites staining, neutral red, lisenarch levers, Janus grean, and methylen blue are the dyes most commonly employed. They are used in the proportion of about 1 part of the dye to 10,000 or 20,000 parts of some normal finit, such as normal saline, Ringer's solution, or Loeke's solution.

For lightly fixing and staining fresh cells methyl green axialulated to about 0.75 per cent with ascelic axid is in common use. Also for the study of fresh cells tissues are teased in a solution of Ripert and Petit (p. 211) to which 0.1 per cent osnic axid has been added, and then stained in methyl green. Acid carmine is frequently used for the study of chromosomes in fresh cells (see memorandum 2, p. 131). For the use of Sudan III with hiring animals see p. 147. Trypas blow will make certain parts of the Bring body take on an interse blue color, bot the color sense to be due whally to the enguliment of the colored particles by certain plaqueytic cells, particularly in the connective tissues of the body (Erans and Schulemann, Scione, XXXXIX [1914], 443-54). Thus, 1 e.c. of a 0.5 per cent solution injected into the peritoceal cervity of a muses will rapidly blue it from east to tail without noticeably interfering with its normal activities.

Equal parts of glyrein, 95 per cent alcohol, and distilled water is a useful examining medium in which fresh tissues may be kept for a long time without deterioration.

IV. TESTS FOR CERTAIN CELLULAR STRUCTURES

The following tests while not always specific are serviceable in helping to identify some of the more usual cellular contents:

Archoplasm stains intensely with acid fuchsin or light green.

Calification may usually be detected by means of 3 to 5 per cent hydrochloric acid. When treated with this solution carbonate of lime emits bubbles of earbon dioxile, while thoughate of lime simuly dissolves.

Cell Walls are usually well defined by acid furths in when used as a counterstain with some of the bematoxylins.

Centrosomes are best shown by the iron-henatoxylin long method (p. 32). Holdenhain finds that they are more sharply defined if the sections, previous to nonclasting in iron-alum, are stained for 24 hours in a weak solution of Bordeaux red.

Chromatin—In fresh tells, nellyl græn stains only ehrematin when it eulers any part of the cell. Alsence of coloration with this dye does not necessarily mean absence of ehrematin. Moderate digetion with gastine juite (at about 40°C) will remove alloumins and heave ehrematin. Prolonged treatment with 1 per cent caustic potash or with faming hybrokheire arith will remove all ehrematin from a nucleus. A 10 per cent solution of softme charite swells chromatin and may dissolve it.

After sublimate finations, thin (3-mirron) sections stained by the Ehrlich-Bandi method (41, p. 222) or American's furthin-methyl-green method (46, p. 224) should show "sective" chromatin or chromosomes green, limin, and plasmosomes red.

Chromosomes.—The best single stain in fined material is ofther ironhematoxylin or sufmain, although chromotoid bolies and mitochoodria, when present, may also stain by these reagents. In fresh tissue, acid carmine (80, p. 222) rokors chromosomes, as does also methyl green (80, p. 231).

Fat—As the fat of issues is dissolved by xylol, slockel, and other reagents used in the parafin and reliabilit methods, only travel, free-hand, or frozen-sectioned, fresh material, or material fixed in some non-fat solvent first such as formalin or Miller's fluid, can be used.

Ownie Acid (21, p. 218) is the communest test for fat. It stains most but not all fasty bodies bown or black. Osmizatel fats are rendered sufficiently insulable to permit of dehythetion and mornting in balsam, if absolute alrehol is seconded and codur of it used instead of xyhi for cheming. However, expand (p. 54) may be used.

Subox III is a specific stuit for fat (see 76, p. 253). Large fat drops stain from a builliant red to an orange; small cons may be yellowish red. The fat of animals feel with the days will become intensity colored by it. The fat in the hypers of yeak haid down in hers? eggs while the lowis are fed on the days (Riddle, Soizoo, XXVII (1908, 945) is stained red, and eggs to colored (Gage) horea into elicks with the body fat colored pink.

Solution B is superseding Solan III as a stain for ist. It stains for onego to red. If a permanent month is desired, frame sections may be fixed 10 minutes in formalia vapor, stained for 12 hours in a saturated filtered solution of the dyp in 30 per cent alcohol, washed in water, contrastained in adm-hematorylin, washed and monted in gybrein or glypenin-jelly. However, see 74, p. 255.

Fat may be removed from tissues ordinarily by treatment with alcohol, ether, or chloroform.

Free Add in tissues may be detected by Congo red, the solutions of which become blue in presence of free acid. Neutral red is turned bright red by acid, yellow by alkalias.

Lecthin may be distinguished from fat by its less solubility in ether and its greater expanity for status. Formalin-fund material if hroughs into accounce has the fat dissolved, but not its lectricin. The latter may be stained by somic acid, heratorylin, orange G, acid fachsin, methyl green, or bolnikin blue, although the tissue should be dehydrated in acotone and left as listle as possible in alcohol.

Glyogen—Readly soluble in aqueous media, hence tissues should be fixed and hardened in 95 per cent alcohol. Gage (*The Microscope*, p. 278) states that a Lugel's solution made of 1.5 grams of iodine or systals, 8 grams of iodile of potssium, 1.5 grams of solium chloride, and 300 e.e. of water gives a differential stain (a malogany red) for glycogen in sections. For vary solidie glycogen he recommends that 30 per cent slocked be substituted for the water in the stain. He deparadins with rylol, mounts in yellow wassline, and scals with shelter or taksm.

Hemoglobin, after proper fixation, stains a characteristic, clear, deep-red color with essin. For crystals see p. 105.

Intra-cellular Reduction Processes may be detected by Janus green used as an intra-robust stain (p. 146). With reduction the color changes from blue or green to red.

Mitochondria (see pp. 143-146),-The most nearly specific single stain for mitochondria in fresh tissue is probably Janus green (p. 145),

Marin in cells, after schlimate fixation, stains with loade but not with orid anilin dyes. Ether thiosin or tolaidin blue stains mucin redshis, surrounding elements blue. Methylen blue and skiranin are also good stains for mucin. See also muci-carmine and muci-kensatin (62 and 63, p. 231).

Nissi's Granules (tigroid substance).-For the methylen-blue method see p. 230. See also pp. 145 and 232.

Oxidase Reaction (Schultze's).-The presence of an oxidizing ferment in cells may be disclosed by the following method:

Solution 1.—Boil 1 gram of a-raphthalin 100 c.e. of distilled water until it metrs. Add pure potassium bydrate (about 1 c.e.) until the raphthal is completely dissolved. The solution should pass from yellow to yellowish brown.

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Solution 2.— Make's 1 per cent squeous solution of dimethyl-p-phenylendiamin (Merck) at room temperature. Filter.

Use frozen sections of formalin-thron material or cover glass preparations fined in vapor of formalidelyse. More the preparations genity back and forth in sahtrion 1 for shout 3 minutes, then do the same in solution 2. Wash in distilled water and mount and examine in water or in giverin-jelly. Oxidase granules are stained drep blue.

Plasmosmus remain unstained in fresh material treated with acid methyl green (60, p. 231) which stains chromatin. With the Ehrich-Bondi stain they stain red, sometimes enouge, with salmain, gentian riolet, and other basic dress, in regressive staining (p. 24), the plasmosmes retain the stain more tenakously than resting chromatin does. Because of greater refractivity they are frequently demonstrable in unstained preparations and sometimes in bring cells.

Secretion Antecedents.—See Bensley, American Journal of Anatomy, XII, No. 3 (1911); XIX, No. 1 (1916).

Spindle Fibers are frequently well stained by add furthsin when used as a counterstain after such fixers as Flemming's, Gilson's, or Bonin's fluids. See also "Euganal," p. 54.

V. ALLEN'S METHOD (Anatomical Record, X, No. 9 [July, 1916])

Fix tissues in Allen's B-15 fluid, which is made up as follows:

Pictic acid, saturated aqueous solution	75 e.e.	
Formalin (e.p.)	25 c.c.	
Glacial acetic acid	5 e.e.	

Just before using heat to 37°C, and add 1.5 grams of chronic add erystals, agitating the mixture riporcustly until the crystals are dissolved. Then add 2 grams of urea crystals. During fixetion keep the Equiti heated to 37 or 35°C.

Pieces of brain 0.5 c.c. in volume fix in 1 hour. Bits of young mature testes require a little longer and pieces of older testes 2 to 3 hours.

Let the find and the contained tissue cool to none temperature and dehydrate gradually by the drop method (menorandum 6, p. 132) with alcold up to 75 per cent alcold, then finish dehydration with anilm oil. Regulate to about one drop per second, or less if the quantity of fining fluid is small. Bring pixes of soft tissue, some 0.5 c.e. in volume, up to 75 per cent alcold in about 1 hour. Harder tissues require more time.

To wish out all pierie acid throughly, replace the ordinary 75 per cent alcohol on the object with 75 per cent alcohol containing a lew drops of a saturated squeens solution of libitum earlonate. Keep up aglasion with a very slow current of air and continue the washing until the yellow color

ceases to appear in the fluid. As soon as possible after washing to avoid strinkage in alcohol, start replacement by andin oil, letting if doop in skorly. To insure rapid mixing a storager current of air may be required; when nearly pure andin is reacted leave the tissue in it until it is clear like amber.

Replace the anilin with bergamot oil or synthetic oil of wintergreen, following the same method. Change the oil one after the tissue has arrived in pure oil.

Warm the oil and tissue slightly and add to it every 10 minutes a few drops of melled parsfin, which must be thoroughly mixed with the oil by means of a pipetite. When the minute is about So or 90 per cent parafin, transfer the object to pure parafin with a melling-point of 52° to 53° C. H bergamot oil has been used, make at least four changes of parafin. Leave the tissue in each about 30 minutes and in a fifth parafin about an hour. Testis material requires longer time. Imbed and section in the usual way.

VI. PHOTOGRAPHING CELLULAR STRUCTURES

 Select a slide in which the part to be photographed is well stained in iron-hematoxylin. The background should be unstained, as a sharp contrast in the slide gives better results in the picture.

 Using an aporhromatic lens and a compensating or projection ocular, bring the part selected for the picture into sharp focus under the microscope.

3. Place the microscope containing the object in focus under the samera. Adjust the hellows to any desired length. Remember that a lower extlar with a longer bellows, at the same magnification, produces a better picture than a higher oxular with a shorter bellows. Also remember that in using a camera mounted in a vertical plane three is loss likelihood of jarring the object out of focus when the microscope is placed under the camera than when it is adjusted in a horizontal plane.

 Focus the object under the microscope upon the ground-glass screen of the eamera. To insure sharp focus, a focusing glass should be used. The picture may now be taken.

5. Fort and Strockell (*Leitschrift für vissenschaftlich Hiltwebupie*, XVIII (1901), 421-30) derelayed the following method, which obvisites forusing upon the ground glass of the eamers every time a picture is taken. Prours from an outlist several concare spectacle keess (-1 to -10 diapters). When steps 1 to 4 here been completed, remove the microscope from under the camera without the least shift of focus. Do not touch the microscher or fue objecture leaves until one is found through which the outlier one after another of the spectacle leaves until one is found through which the object appears in as sharp focus as it did upon the ground-glass sereen. Note the number of the particular leave und objective, the same object is to be photographed with the same ordur and objective, the same the and the same ordur and objective, the same the and spectra the same ordure and objective, the same the and spectra the same ordure and objective, the same the and spectra the same ordure and objective, the same the and spectra the same ordure and objective.

bellow length, one heed only place the dispirie lens over the oralar, from until the points desired in the histhet photograph stand out sharply, remove the dispirie lens, place the microscope under the camera without shift of froms, and take the picture. Since the dispirite lens corrects the focus for that bellows length, the necessity of refronsing upon the ground glass is removed. Whenevers different combination of oralar objective and bellows length is used, the proper dispirite lens must be found.

6. It is good practice to allow the microscope to stand for a time after the object has been focused through the dioptic lens in order to be sure that the focus does not shift. After the picture has been taken, by replacing the lens over the coular one can see if the focus has been held throughout the fine. Even with the utmost care the focus will sometimes charge.

MEMORANDA

1. Accessory Chromosomes (see chromosomes, X-elements) are perhaps best demonstrated in the tests of some species of the short-homed gassboyees taken about the time of the last model. In these forms minote figures are large and chromosomes usually distinct. Among the Hemipten, the spaceh long (Anose tristil) for single X-element, and the stink lung Euclidea, for X- and Y-elements, are recommended. Flemming's or Bouin's fluid may be used for fixing, and iron-hematorylin or safrain for staining.

2. For Quick Determination of the Chromosomal Condition of cells astro-carmine preparations (35, p. 222) are useful. The realize tests of an insert is put on the solver, or smears from the tests of larger animals are made and fineded with the stain, after which a correscip is added. The edges of the over should be earled with vascine to prevare exponsion. Chromosomes are stained in a few minutes. Such preparation should be carefully tested by observations on welf-find and stained materials, however, since the great amount of acetie acid in the acid carmine swells chromosomes and is likely to lead to errorewers conclusions regarding details.

 Protoplasmic Currents in cells may be seen to good advantage in Rhinopods, in the plasmolia of Myxomycetes, and in the stamens of Tradescantia.

4. Celluidin Instead of Parafin is being used none and more by various epidogists for making sections, the purpose being to avoid the had efforts of hot parafin. For example, Danchakoff (Leitelnijt for wisenschöftliche Mikwakoje, XXV [1960] starts with very thin celluidin, changes to somewhat thicker, then to still thicker (about 3 per cent), and finally, having arranged the tissues, lets the solvent evaporate vary slowly throughout 4 or 5 days to a week. The mass should become equiline, hanogeneous, and

about as hard as vulcanized rubber. It is stored in 80 per cent alcohol. Thin sections can be cut if the mass is sufficiently hard.

5. Une is being used very successfully in such thing thisks as Flemming's and Bonin's by Professor C. E. McClung and his pupils (see Aller's method, p. 199). From 1 to 3 or more grants per 100 e.c. of their is the quantity used. The exact proportions for any particular tissue can be determined only by trial.

6. Very Gradual Changing of Fluids is recommended in the treatment of tissues to be used for cytological studies. This may be accomplished very

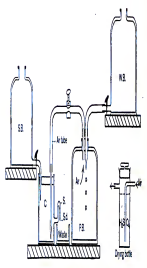


Fig. 42 .- Apparatus for Gradual Change of Liquids (after Ezra Allen)

successfully according to the drop method described by Allen in the Anatomical Record for July, 1916. The apparatus he designed for the purpose is shown in Fig. 42.

A 2 or 3-liter sopirator boths (*IF.B.*) is filled with water and its stopcock opened slightly until the water begins to drop into the tightly eached pressure boths (*P.B.*). As the air is compressed in this boths bubbles will begin to issue from the air tube through the liquid in the small container (*C*) in which the tissue liss. To secure a steady stream of air through *C*, the end of the air tube is drawn out into almost a capillary and the rubber tube connexing it with *P.B.* is champed nearly shut. The purpose of the eurent of air is to insure quick and thorough mixing of liquids in *C.*

The alcohol or other replacing fluid in the supply bottle (S.B.) is started to dropping at the desired rate, the flow being regulated by a stop-cock or

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champ. A siphon on the side of C removes the excess of fluid into a waste jar, so that the concentration of the liquid which is being added steadily rises.

For higher abouts or oils the air should be dried by passing it through a tube containing calcium chloride or through subhurie acid (see figure). Two liters of water in W.B. should last all night at the rate of a drop a second.

7. The Use of Amin Olin Place of the Higher Alcohols for completing dehydration is strongly advocated by Allen (op. cl., p. 149) because dehate fascus are less likely to skrink in it. Strore smill oil does not mix with parafin it must be followed by some dearing oil. Allen prefers oil of bergemot or symbelic oil of wintergreen.

8. Coding Tissues by placing them on ice for 15 or 20 minutes before function may smeetimes prove helpful in preventing the champing or sciencing together of chromosomes. With difficult material, such as that of biols or mammals, it is well, at least, to try this method along with others.

 The Mature Testes of Young Mammals or Birds are better for studies in spermatogenesis than those of older animals.

10. For Dissection of Living Cells an appendix designed by Barber is being used very successfully by several investigators. It is described in full in the Known University Science Bulletin, IV (March, 1907) and in the Pullippine Journal of Science, IX (1914). Some workers employ a double form of it.

11. For the Estimation of Very Minute Quantities of Carbon Dioxide, Tashino has designed several pieces of accurate and delicate appacatus which are described in full in the American Journal of Physiology, XXXIII, No. 2 (1918), 107-45, and the Journal of Biological Chemistry, XVI, No. 4 (1914), 455-94. These various types of apparatus may be obtained from the Einer & Ameral Co., of New York City.

 Holmgren's Canals may be demonstrated in various cells by faxing in Bensley's formal-bichromate-cublinate mixture (*Biological Bulletin*, XIX, No. 3 (August, 1900). The composition of the final is as follows:

Neutral formalin (freshly distilled)	10 e.c.
Water	90 e.c.
Potassium bichromate	2.5 grans
Mercurie chloride	5 grams

Use the solution soon after making up.

 Buparal (VIII, p. 54) is of great value in the study of certain achromatic cellular elements.

CHAPTER XVIII

RECONSTRUCTION OF OBJECTS FROM SECTIONS

In investigating objects which possess complex internal cavities or complicated structures it is frequently very difficult to gain an adequate idea from the direct study of serial sections, or by means of macented or teased preparations; consequently rarious methods of plastic or geometrical reconstruction from the sections are resorted to. For such reconstruction, sections must be of uniform thickness, serial, and they must possess similar orientation.

RECONSTRUCTION IN WAX

Bon's method of constructing wax models of objects from serial sections is widely used for both embryological and anatomical subjorts. The thickness of the sections, the magnification of the microscope, and the plane of section must be known.

Wax plates are prepared as many times thicker than the actual sections as the latter will be magnified in diameters. For example, if the serial sections are $\frac{1}{34}$ of a millimeter thick (33 $\frac{1}{34}$ microns), and they are to be magnified (0) diameters, then the wax plates must be made 60 times as thick as the sections, or 2 mm, thick. This is a thickness commonly used. Count the number of sections to be reconstructed, and prepare an equal number of plates.

Preparation of the Wax Plates

 a) The hot-water method.—1. Prepare the wax according to the following formula:

Beeswax	6 parts
Paraffin (melting-point 56° C.)	4 parts
White lump (not powdered) rosin	2 parts
Melt together and thoroughly mix.	

2. To prepare plates of the proper thiskness (2 mm.), use shallow studget-walled rectangular its pass which will alloed a water surface of 3):4 feet; 2,000 gauns of the prepared wax pourced on very hot water in 154

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such as pan will give a plote 2 mm, thick. Air bubbles which form in the wax may be driven of before it cools by playing a Bunsen flame over the surface. The wax should spread evenly over the surface of the waite if both wax and water are sufficiently hot. If gaps remain, elsee them by drawing a glass file over the surface of the wax. To prevent the plate from splitting while cooling, after it has sufficient somewhat, out the edges free from the walls of the pan. When the water has become tepld, remove the wax plate to a flat support and leave it to harden.

b) The complete machine method.—Several instruments have been devised for making the plates more rapidly and more accurately than by the hot-water method. Hoter's appearing, for instance, consists of a heavy cast-icon plate with movable side pieces which can be adjusted to a height corresponding to the desired thickness of the war plates. The whole instrument is supported upon three adjustable legs, by means of which it can be made easely level. Melted war slightly in encess of the quartity necessary for a war plate is poured on to the iron plate in an even layer, and rolled out with a hot roller until the roller comes to run directly on the side pieces of the instrument. When the war plate is cont encode to handle, it may be placed in a pan of cold water to harden.

Practical Exercise.-When possible an outline drawing of the part to be reconstructed should be made before it is sectioned.

 Reconstruct the heart of a chick at the end of the third day of incubation, under a magnification of 60 diameters. For this magnification, if it is desired to use a wax plate 2 mm. thick, the original sections should have been 33.3 microns thick.

2. Place a sheet of blue tracing-paper on the wax plate with the colored side toward it. Over the tracing-paper place a sheet of ordinary drawing-paper. With the aid of a camera lucida or other projection apparatus, outline on the drawing-paper the part to be reconstructed. In doing this the outline is also traced in blue on the wax. Number each drawing, and also indicate the number of the section on the slide to which it corresponds; also number the wax plates with reference to the drawings.

3. Lay the wax plate on a suitable fat surface, and cut out the outlined parts with a sharp, narrow-bladed knife. Leave bridges of wax to hold in place the parts that would otherwise be separate pieces. Pile up the successive sections in proper sequence as they are cut out.

4. In finally putting the model together, accurately adjust the parts (for reconstruction points see memorandum 13, II, a, p. 127), and build up the model in blocks of five sections each (Bardeen's suggestion). If necessary, unite the essential parts by means of parts or fine nails. Remove all temporary wax bridges (see 3) by means of a bot knife.

When all blocks are properly adjusted and united, smooth over the surface by means of a hot spatula.

MEMORANDA

1. Geometrical Reconstructions, first described by Professor His, are often all that is necessary to give one the desired information about internal organs. Before sectioning, an outline drawing of the object is made in a plane at right angles to the intended plane of section, and under the same magnification that will be used for the reconstructed drawing. For example, if the sections are to be transverse, the outline drawing of the object would be a profile view from the side. After sectioning the object, each section is charm under the same magnification as was used for the outline drawing.

To reconstruct any special part of the object, show a median line on the outline drawing corresponding to the long axis of the object. At right angles to this line draw a series of equidistant pecallel lines corresponding in positions to the sections that have been made. For example, if the magnification is 100 diameters and the sections 10 micross thick, then the parallel lines must be 1 mm, apart. Then, beginning with the first section, indicate by dots in the proper plane in the sections having the relative distances of the parallel lines. All of the sections having thus been plotted, connect the dots of corresponding parts in the successive nons. It is frequently sufficient to reconstruct only every fifth or even every tenth section. When the plane of section is not quite at right angles to the axis of the object, an equal alterction of angle must be made between the median line of the outline drawing and the parallel lines.

Such a reconstruction as that above would give lateral views of the various internal parts. To get their asperts as seen from above or below, the original outline drawing of the specimen as a whole should have been made from this point of view instead of from the sile. In actual work one should make reconstructions in both planes.

For a molification of Weber's method of graphic reconstruction, see Scanman, Andonical Roard, IX, No. 3 (March, 1915). For suggestions on profile reconstructions see Streeter, American Journal of Andony, IV, No. 1 (1914), St.

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 A Special Drawing-Table for rapid and convenient drawing of sections for reconstruction has been devised by Barbeen. For details, see Johns Horking Bulletin, XII. 148.

3. Starts of Biothing Paper instead of wax are recommended by Mrs. Gage (Antomical Roord, I, No. 7 [November, 1997]). Models are insided by coating with parafilm. The advantages claimed for this method are lightness, durability, and ease and cleantiness of production. The method is also given in Gage, The Microscope, pp. 335–32.

4. Pattes with the Paper of the Drawing Rolled into the War, following directions in Karl Peter, Meldoen de Rebonstruktion (Gastar Fischer, Jeau), have been found very satisfactory by Rive. A thin drawing-paper, smooth on the drawing side, provus on the other, is used and drawings are duplicated by means of a carbon sheet. One copy is kept for reference, the other is pressed into the plate. To accomplish the latter, the slab on which the wax is realed out is smeared throughly with turpentine sol the drawing kail face down on it. The melted wars is one-ord whose these both the drawing and, when begianing to harden, is overed with tissue paper, which is then plates. The thicker paper with the drawing out it gives a fixed contour which is a helpful guide when it ownes to smoothing down the model.

Twisted wires in short lengths are used for supports. The earls are spread to afford anthonge where they lie between the plates. Two longer wires should be twisted together, then out into proper lengths.

5. Patagraphy of Sections upon Large Platis has been resorted to by Warren H. Lewis (Automini Record, IX, No. 9 Stytember, 1915), (19-20) as a substitute for the laboritors and time-moreorming method of draving each section on paper. He uses line humaide or ano G hard (matte) prints. He considers the photographs for superior to dravings and maintains that, when time is taken into account, the method is loss expressive than the old method of traving. Lewis' paper is full of valuable suggestions and should be read by everyone who contemplates doing much work in reconstruction. The duif points emphasized are: the use of photographs; the use of series of guide-lines which coincide with planes that are sit right angles to each other and perpendicular to the plane of the sections; and the use of plaster-of-Paris easts.

6. For Rapidly Cotting Out the Wax Plates, Chester H. Heuser, of the Wastar Institute of Anatomy and Biology, has devised a series of nortal styli of varied design which when electrically heated are handled much as one would handle a pen in writing. The cotting instrument proper (opper, inn, or brass wire, pointed or fastneed at one end according to need) is could enough for about 15 mm. at the working end, with a thin have of an insulating askestos paste. A piece of No. 32 German sliver wire about.

20 cm, long is then wrapped around the coated surface, with the earls kopt well insulated from each other, and finally covered with the poste. The earls of the German silver wire are attached to small copper wires which run to a lamp-band bearing several incandescent bulls with sockets connected in parallel. The lamp-band serves as a theostat, so that any desired temperature can be obtained in the stylus by altering the number of lights and thus regulating the current which passes through the German silver wire. Styli of different kinds of metal attached in common to the same lamp-band after inserting small heostats in the system to regulate the temperatures of the individual instruments more acoustlely.

Professor Mark (Proceedings of the American Academy of Arts and Science, XLII, No. 23 (1907)) uses an electrically heated wire moved rapidly by a molified serving machine for cutting out the wax plate.

CHAPTER XIX

DRAWING

I should make it absolutely newsary for everybody, for a longer or shorter period, to learn to draw. It gives you the means of training the young in attention and accuracy, which are the two things in which all manifold are more deficient than in any other meetial quality whatever—Hucky.

Drawing is an important part of the work in most biological sciences. The essential phases of a subject can be condensed into a few pages if the drawings accurately represent the dissections or microscopical preparations studied. The following simple directions are written mainly to aid the student in preparing his notebook, but it is hopped that they may also be useful to individuals preparing manuscripts for publication.

Materials for Class Work—All the materials needed for ordinary class work can be selected from the list here given with the approximate price attached.

Pencila-one-HI, use HB, one 2B, 10 cents each. Pens-Gillot's lidiographic perpoint, No. 201, 5 cents each. Ish-waterproof India ink made by Charles Higgins & Co., retails at 25 cents per bottle. Buler-celluloid, 10 cents. Buly enser-10 cents. Cropon pencila-reci, blue, and yellow, 5 cents each. Low-kaf publich with bond paper, 40 cents.

The pencils may be of any standard make; the 4H is a hard pencil for line work; the HB, a medium pencil, is useful in placing outlines and shading. The 2B, for black shading, should not be used unless the drawings are afterward fixed to prevent rubbing. The ruler graded in centimeters on one edge and inches on the other is indispensible. The enser is for ensing pencil lines; for ensing ink a sharp haife is best. A larger assortment of colored pencils will often prove useful, but the three primary colors will answer most purposes. Two-phy bristol loard at 2 cents per sheet may be used

in the notebook instead of the bond paper. Where classes are large, bookstores will make up bound notebooks for 35 cents each, containing a good grade of paper upon which drawings can be made. Where pend drawings only are required, pens and ink may be omitted from this list. On account of the small amount of locker or drawer space usually available for one student, an elaborate drawing outfit should be avoided. The excellence of student drawings is judged by the exactness with which the preparations are depicted.

L METHODS OF REPRESENTATION

Outline.-In beginning a drawing, the field which the picture will occupy should be marked off with dots. The placing of two faint lines which cross at right angles in the middle of the drawing-field is a great help to beginners, especially in drawing bilaterally symmetrical objects. Next determine the relation of the length of the object to the breadth; then calculate the size of the drawing. If the object is large, a reduction will be necessary; if small, it can be represented better 5 or 10 times its original size. The important points can be indicated in the drawing-field by dots which, when connected by light lines, roughly block in the object in correct proportion and size. This crude picture may then be worked over until angles are removed and a neat outline results. For the preliminary mapping of the object an HB pencil is best, as the lines are easily crased. The outline when finished must consist of a continuous line of uniform thickness with no overlapping edges where the pencil has been removed from the paper and put down again. Outline is the most important part of the drawing, for "a good outline may redeem bad finish, but no amount of excellence in finish can save a picture that has been incorrectly outlined." What details to include depends upon the purposes of the drawing. The principal points of an anatomical drawing stand out more clearly when they are not obscured by unnecessary details. In histological drawings, details are essential, but they should still be kept subordinate to the general effect of the picture.

Depth.-Usually the third dimension, depth, is not considered in drawings made from sections. When drawings of reconstructions

or whole mounts are inade, however, all three dimensions must be indicated in the drawing. Likewise, drawings of such things as digestive canal, nerve cord, heart, lungs, and kidneys stand out better when depth is represented. This can usually be done by indicating degrees of light and shade.

Ink Drawings—In drawings which are to be inked, the outline should be carefully drawn in pencil and as many corrections as possible made before ink is applied. Place the ink upon the pen by means of the quill attached to the cork of the ink-lottle. If the original outline is even, the inking can be done readily; a line uniform in thickness results from the applications of firm, steady pressure. The pen will give a ragged line if held so that one mb bears more heavily upon the paper than does the other, or if it becomes sticky with dried ink. A smoother line will be obtained if the penholder is held at a wide angle to the paper and only the very point of the pen is allowed to touch.

Shading.—Where differentiation of parts is desired, shading may be used. This can be done either by stippling or by lines. In making the dots in a stippled drawing, the pen nust be grasped firmly and only the point placed upon the paper. If the pen strikes the paper at an acute angle, three-sided instead of round dots result. The dots must all be of the same size. To indicate degrees of shade, vary the number of dots, not their size. A heavy shading can be accomplished by placing the dots close together, whereas dots farther apart give the impression of light shading. Lines can be used with good effect upon large drawings. Let the lines, placed an even distance apart, follow the shape of the shadows. To indicate heavy shadows, lines in an opposite direction can be placed across the first set. Be caceful not to cross-hatch until the first lines have dried, otherwise blots will occur.

Pencil Drawings.—When correctly executed, pencil drawings are more artistic and permit of more subtle differentiation in detail. Minute points can be shown by the use of stippling. In stippling with a pencil, fellow the same procedure as in the use of a pen. The pencilpoint should be sharp and rounded on all sides. In large drawings, lines can be evenly placed to outline shadows, but they are not as

effective as shadows put in by blending graphite, obtained by rebbing the pendi over the paper. For the latter method a stub is necessary. This can be made from a strip of paper 1 inch wide and 5 indees long in the following manner: Begin to roll the paper at one end and let each turn overlap the preceding turn slightly, until an elongated coil results. The pointed end of this can be used in spreading graphite evenly over a surface. The graphite is placed on the part of the drawing where the darkest shadows occur with an HB or 2B penell, and is worked over with the end of the stub until the sharp edges of the shadow gradually grade out into the lighter parts. The 2B penell outinarily should not be used as the source of the graphite, as shadows can be darkened by reveating the application of the HB penell.

Shadows—In most pictures of biological subjects, one cannot stand off and observe where the hight falls upon the object and what part is in shadow. For that reason a knowledge of where the shadows occur if an object is illuminated from any one direction is necessary. To gain such knowledge from a description is impossible; it is therefore advissible for students wishing to shade their drawings to consult an artist who can give usable information in the form of demonstrations. A caveful study of textbook illustrations will aid materially. Shading requires practice, and even then it may not be successful. In most cases where not imperative it had better be left out entirely.

Fixing penal drawings.—Where penal drawings are made with soft penals which are liable to rub, they must be fixed. This is done with a fixing solution and a special atomizer which can be longht at any art store. To prepare the fixative, make a saturated solution of white shellact in alcohol. Allow this to stand for a day or so; dilute one-half; then filter off the liquid. To prevent eraporation when not in use, this must be kept in a tightly stoppered bottle.

The drawing should be placed in an upright position, about 2 feet from the spray. In order to avoid a glossy surface spray lightly.

Wash-Drawings.—After a faint online of the section or object has been made with a hard penell, fasten the paper to a load with thumb tacks, and with a large brush dampen the entire surface, removing the surplus water with the brush or a blotter. Mix the

wash as follows: With a wet brush remove some pigment from a eake of Winsor & Newton's Charcoal Grey or Ivory Black, and put in the water in the mixing-pan. Repeat this process until the wash is slightly darker than the desired background tint. Next put a wash of this over the entire background; it will dry lighter. Allow the paper to dry partly before darker tones are put in. Where a very dark portion is confined to a small area, the paper should be quited dry, otherwise the wash will run into the surrounding part of the drawing. If details are to be put in by stippling or linework, make a wash (from the same eake) the color and consistency of ink. This can be applied with a pen or brush after the paper drives.

Some artists use a dry paper which, however, requires more skill in applying the wash. The darkest tones can be applied first, gradually working up to the lightest. After experimenting, use that which gives the best results for the purpose in hand.

Where serveral wash-drawings are to be made with the same toxes it will be found simpler to put in all the tackgrounds first. Mix up plenty of wash for this purpose, as it is not easy to duplicate the exact shade at another mixing. Wash allowed to stand becomes darker upon erroporation of the water; hence, if after the backgrounds are put in the work must be deferred until later; the same wash will do for darker tints. Do not redsmpen the whole surface, as that will lighten the background. The darker tones can be blended into the background with a clean wet bruch.

MEMORANDA

1. Cleanthess.—Even where the dravings are correctly made as to size and proportion, the notebook will not present a good appearance if the pages contain fugges marks and blots to mar their whiteness. With sufficient diligence fugge-marks can be eased, but the best way is not to make them in the beginning. Where laboratory work requires dissection, rough sketches can be made upon scrap paper and later expired into the notebook. Blots can be swelled if care is used in playing the ink upon the pay, a spall is stached to the stopper of the ink-bottle for this purpose.

Size and Arrangement of Drawings.—Uniformity in size and arrangement of drawings should be preserved wherever possible. For example, in the development of the troy's egg, no increase in the size of the egg takes

place from the single-cell stage to the end of gestrulation, hence all drawings representing this series of development must be the same size. The neural tube, largest in circumforcese in the head, decreases gradually toward the posterior end of the lody, and must be drawn corneelly in cross-sections, otherwise one will have an erroneous idea of its structure. The drawings must not be crowled upon the page. Exact margins and spaces equidistant between drawings, though not artistic, give the impression of matness desirable in scientific work.

Labeling.-The results will not equal expectation if the labeling is poorly done. Too few students consider this item in making a notebook. The peer-layoint must be rubbed upon has sandpaper until it is smooth and exoinal. The beginner should draw 3 parallel lines, 2 mm. spart, upon which to place the letters. The letters may be placel close together or further apart, but in either case the space between the letters must be kept the same. Larger spaces are left between works. Letters may be straight

ABCDEFGHIJKLMNOPQRSTUVWXYZ abcdefghijklmnopqrstuvwxyz 0 1 2 3 4 5 6 7 8 9

 $\rm Fra.43.-Simplified Goldic or "Shop Skeleton" Letters and Figures Used in Labeling Drawings.$

or sharled to suit the individual taste. If one has never done any holding, time should be taken to practice upon a piece of paper heltore finishing a drawing. Many letters like h, d, and q contain an q contained with a straight line, hence it is necessary to learn to make an q properly. In making h, d, p, and q, be earchil that the up or down stroke is straight and joins the eurre smoothy. The h, d, f, h, k, and l, extend above the other letters to the height of capitals, while g, f, q, g, and g entrail just as far below the line. The *t* falls between the stem letters and the short letters. The Cothie style given in Fig. 43 is easy to learn and will answer all purposes.

II. PREFERABLE MODES OF REPRESENTATION FOR SPECIAL COURSES

General.—Methols of representation vary as the subject-matter in each instance requires different handling. The modes here described have been found practical in different biological courses; however, each may be altered to suit individual requirements. In

Drawing

elementary courses in general noilogy unshaded ink-drawings are best. They are more accurately executed by elementary students because defects are so obvious that they do not pass unnoticed; moreover, the student exercises more care in making the drawing because of the greater difficulty of changing it after it is once drawn. Such drawings may be made more or less diagrammatic, depending upon the shallity of the student and his previous training. Students with no former experience in representing upon paper what they see meet not be discouraged, for clear-cut ontline drawings can be made by any one, if due consideration is given to the points enumerated under the foregoing paragraphs.

Embryology .- The first step in embryological drawing is a careful outline with a 4H pencil. As an aid in drawing complex sections -a cross-section of an old tadpole, for example-a cover-glass ruled into squares can be fastened into the ocular. The drawingpaper is ruled into the same number of squares as the cover-glass; each square as many times the size of one square of the cover-glass as the intended magnification. Parts of the object under the squares in the ocular can be located in corresponding squares upon the paper (Isaacs, Anatomical Record, IX, 711-13). In such drawings cells are not usually indicated. It is a decided advantage to have the parts colored, especially if organs from the same germ layer are colored alike. If three colors are chosen, one for each of the germ layers, ectoderm (blue), endoderm (vellow), and mesoderm (red), and if these are consistently used throughout the sections, one can see at a glance the development of the organs. Crayon pencils work up rapidly and give good results if the color is put on lightly, so that in the finished drawing the colors blend. To spread crayon evenly, use a blunt, rounded point and make long strokes with the side of the crayon. For example, in a cross-section of a frog embryo, the neural tube with its optic cups is ectodermal in origin and the blue color immediately indicates its relationship to the ectoderm which is similarly colored; the mesoblastic somites and mesenchyme are red; the lining of the archenteron yellow. In older embryos where many parts develop from the mesoderm, other colors may be used for special parts-as brown for kidneys. The shade of red may likewise oe

The Street of the second se

varied, a dark red being used for blood ressels, obtained by putting on a heavy layer of crayon; while a light red may be used for mesenclyme. By this method a student has constantly before him the layers from which various organs develop, while the instructor can immediately see that the student has or has not a clear conception of the manner in which the organism is built up. Water colores can be used instead of crayons; but in the hands of most students they are less satisfactory. Where cellular structure is put in, colored inks may be employed. However, drawing the individual cells in an embryo is too time-consuming a process for ordinary elses work in embryology.

Histology.—Histological drawings are best executed in penell. Details must be shown. In general, these, especially nuclear differences, can be put in by stippling. In intercellular matrices, such as connective tissue, the texture of the tissue should be represented by irregular lines. Light lines give the effect of fibers and fibrils very well. Where different tissues of an entire organ are to be distinguished, the whole drawing may be covered with a light groundsubstance of blended graphite and the details worked up with pen and link. This combination is effective and has the added advantage of quick execution.

Cytology—Cytology requires even more detail than histology. Stippling or wash is used chiefly. A different arrangement of the dots gives the gramular, alwohar, or reticular appearance of cytoplasm. Solid lines should be avoided as much as possible; fibers can be represented by dotsplaced close together in a linear row. Chromosomes can be stippled, made solid, or blended with graphite over their entire area. Where the cell cytoplasm is homogeneous, a light ground-oast of graphite can be placed over the entire cell and the cell parts stippled upon this with pen or pencil. A wash can be substituted for the graphite; but as it requires more careful application it is not recommended for class work.

Crayns used judiciously in both histological and cytological drawings are often effective. Secretory granules, where present, can be put in with color if the crayous are sharpened to a fine point. Ground-substance of tissues can be depicted with crayon and the

characteristic features of the tissue added with pen or pencil. Granules of white blood corpuseles have definite color reactions upon the basis of which they are elassified; such granules should therefore be colored in a drawing.

III. DRAWINGS FOR PUBLICATION

To make illustrations for publication in a book or scientific journal, one must not only understand form, color, perspective, and composition, but also know something about the science of reproducing drawings in printed form. Manuscripts which contain drawings that can be cheaply reproduced are more readily accepted for publication than those which require expensive plates. Before undertaking a series of drawings for publication make a careful study of similar work in standard journals, particularly in the journal in which you expect to publish.

Materials for Manuscript Drawings—In general, drawings for publication should be made in black and white, because this style can be reproduced most cheaply by publishers. For working with ink, a water-proof India ink, such as Higgins', is best. This can be applied with pen or brush. Gillott's penpoints are satisfactory. Insamuch as each person not only handles a pen differently, but uses different degrees of pressure in working, he must determine by experience the number of the pen best suited to his needs. For fine linework and stipping, the writer has found that Gillott's lithographic pen No. 290 gives the best results. Fine red-sable brushes may also be used for this work, although for line-process reproduction (p. 168) the pen drawing is likely to prove nore succesful.

Drawings should be made upon a good quality of paper. Bristol board, either 2- or 4-ply, or Whatman's hot-pressed (smooth) watercolor paper, either of which can be obtained at any store carrying a complete line of stationery materials, can be used. Whatman's paper is of the same texture throughout; moreover, it can be used for ink, wash, or pencil-work. Anvil drawing-paper, No. 105, is excellent for large illustrations or charts. This is a cloth-backed paper which can be rolled without injury. A disadvantage is the

large size and expense of a roll, which contains 10 yards of paper 36 inches wide. It is part out by the Keuffel & Esser Co., of New York. Cloth-lacked papers can be obtained from other firms. A stipple-board (Ross board), manufactured by the Charles J. Ross Co., consists of a chalk surface upon a paper back. The advantage of this paper is the rapidity with which drawings can be made. A stipple effect is obtained simply by rubbing the flat sile of a penelpoint back and forth over the chalk surface. Different stipple effects are obtained by using different grades of the paper. For all general uses No. 8 is best.

Water colors may be preemed in a variety of forms and makes for wash-charrings. On account of the cost of reproduction, however, in papers which are to be published colored drawings should be avoided wherever possible. Winsor & Newton's Ivory Block or Charcoal Grey, which comes in a cake, is best for black-and-white wash-drawings. Never use blue in a drawing to be photographed, as it does not take. Benember also that yellow and brown appear as black. Likewise keep in mind that the results are in no appreciable degree dependent upon the number or kind of tools used, but upon the skill shown in execution.

Camera Lacida—As an aid in making drawings of microscopic objects, an instrument known as the camera bucida is often employed. With such an instrument the image of an object under the microscope can be projected upon the drawing-paper (see p. 189).

Reduction of Drawings.—In making drawings for publication, it is advisable to make them larger than they will appear in the finished cut, as in the reduction many irregularities are lessenel. But under no circumstances make a crude drawing with the idea that in the print it will appear perfect, for while reduction minimizes, it does not obditerate deflects. The original drawing should not ordinarily be more than twire the size of the intended cut, while a reduction of one-fourth or one-third will probably give a better result. If one is not careful about the spacing of lines and dots used in the drawing, reduction tends to make them run together.

Line-Process.-Line-process is not only the cheapest form of reproduction used by journals, but likewise the most accurate.

Prints reproduced in this manner contain contrast not gained by any other method. The drawing is photographed directly upon a zinc plate. Ink is then applied to this plate with an ordinary ink-roller. The sticky ink adheres only to the lines of the photograph. The inked plate is now placed in a both of acid which eats away the uninked portions. The metal plate, containing the design in relief, mounted upon wood, type high, is set up and printed with the type upon text-paper if the details are not too fine and if no colors are required.

Lines.—For this type of reproduction use ink as the working medium and apply it with pen upon smooth white bristol board or water-color paper. Lines should not be extremely fine, and eare should be taken that they are far enough apart to reproduce well. Likewise, cross-hatching, if employed, must be coarse enough to stand the reduction, otherwise it will appear as a solid black mass. While ink thinned to a gray gives a difference in tone in the original drawing, it must be borne in mind that this behaves erratically in reproduction. Gray lines often appear in the print as broken black lines, or they may be entirely lost. To avoid graying black lines the pen must be wiped frequently and filled with a fresh supply of ink. To get effects in tone vary either the thickness of the lines or the distance between the lines. Lines in the foreground should be farther apart and heavier than those in the background.

Data—If drawings are stippled, the dots should be made with the amount of reduction in mind. Too fine stippling necessitates etching upon copper, a telious process, which doubles the cost of production. To secure degrees of light and shade, vary the distance between the dots, not the size of the dots. Lithographic erayon used upon stipple-board (see p. 168) will reproduce by line-process if the drawing is coarse enough to stand a reduction to one-half its size.

Grapht—Graphs for line reproduction should be made upon co-ordinate paper in which the lines are blue. As blue does not photograph, the co-ordinates, both perpendicular and horizontal, which are to appear in the cut must be inked.

Color — A separate block and printing are required for each color used in the engraving. In a drawing in which several colors are to appear, the original drawing in black and white must be accompanied by a colored sketch. From a process-plate of the black-and-white drawing a print is taken. For each color used in the sketch, the printer then makes separate blocks which are printed one over the other upon the print from the first plate. Avoid fine gradsitons in color, as thereby the expense of reproduction is increased.

Half-Tone.—Shaded and wash-drawings and photographs are onlinearly reproduced by the half-tone process. A screen containing 133 to 115 meshes per square inch is photographed. This is done to break up the flat tones into dots, otherwise they would print as black masses. The photograph thus made is transferred to a rine or copper plate and etched in the same way as a line-process plate. Where the details of the picture are not too delicate, the print can be made upon text-paper. For proper reproduction, drawings which require a fine screen must be carefully printed upon special coated paper.

It must be remembered that the serven on account of its fine dots introduces chadow into the lighter parts of the picture; while the white spaces between the dots render the darker areas lighter. These changes demand stronger contrasts in the original drawings and photographs which are to be reproduced by half-tone than those desired in the print.

Wost-drawings—Wast-drawings (p. 162) for reproduction should not contain solid black lines, as the screen breaks up the lines into dots. Be careful that no penel lines remain in the drawing and avoid sundges and finger-marks. If clear white spaces are desired in the print, the shadows produced by the screen in these places must be tooled out of the plate to give the required effect—a process which is slow and expensive.

Combination with wesh.—Drawings can be made in which certain parts are put in with pen or brash lines upon a wash-background. This makes a satisfactory combination for half-tone prints. On the other hand, fine penell strike upon a wash-background are usually last

in reproduction, since they offer insufficient contrast with the background.

Pencil—Pencil-work does not reproduce satisfactorily by halftone. Where blended graphite is used for a background instead of wash, a reproduction can be made if the print is made upon good coated paper. However, the results are so inferior to the originals that it pays to take the time to learn how to make good washdravings.

Color.—Color can be applied upon half-tone prints, although a separate block and printing are necessary for each color. Plates are sometimes made as described under "Line-Process," and these printed in color upon a half-tone background (see p. 170).

Photographs.—Photographs are usually reproduced by halftones. A hard-finish, glossy paper which brings out strong contrast between the blocks and whites of the picture is best for prints. Ano hard X, glossy white Velox paper made by the Eastman Kotek Co., and solio paper of a brownish tinge reproduce well. Photographs will stand some reduction, but on the whole it is better to have prints the same size as the intended cut. Prints should be squeegeed. This is done with a roller which, passed over the wet print, removes the moisture and gives a hard finish to the gelatin film when it drive.

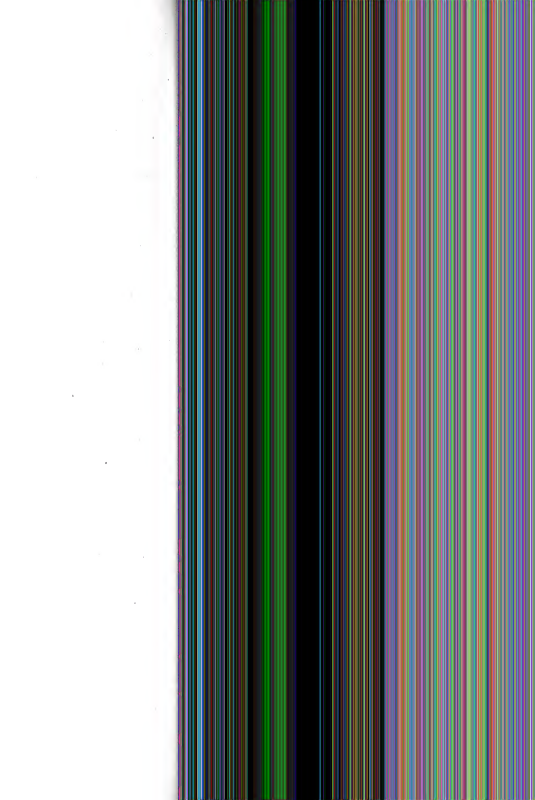
Lithography — Lithography is the most expensive form of reproduction. While line-process and half-tone plates are made by mechanical means, lithographic work is all done by hand. A picture to be reproduced by this method must be transferred to a stone and the parts cut in with a graver in the hand of an expert. Undoubtedly it is the most artistic form of reproduction, but most journals will not accept drawings which have to be engraved unless the author or artist pays the extra cost of reproduction. Intricate drawings of many colors which cannot be reproduced by hne-process or half-tone come out well by this method, if owst is not an item to be considered. Likewise stippled penel drawings make excellent illustrations with this sort of reproduction. Heliotype (gelatin plate) is a somewhat less expensive method for reproducing colored drawings and photographs.

Arrangement of Drawings for Reductions.—Line-trawings where not used as text-figures should be arranged in the form of plates. Half-tones are usually so arranged, since they generally require a special costed paper. To arrange drawings in the form of a plate, one must know the exact amount they are to be reduced when printed. If one-shall, for instance, they must be arranged as a plate twice the size of the journal page. The drawings can either be made directly upon bristol board in the order in which they are to be printed, or they may be posted upon the board in such order. The latter method is usually practiced. Due allowance must be made for lettering and the margins of the page.

Lettering.—All the original drawing schould be so lettered that the letters will be of the proper size when reduced. Letters can either be pasted on or printed by hand (p. 164). A drawing presents a neater appearance if the lettering is parallel to the base line. A cut is more legible if, instead of abbreviations, the names printed in full are connected to the proper part of the drawing by leaders. Gummel sheets containing letters, numerals, and such works as "Plate" and "Fig." in several sizes can be lought for use in this work. Likewise publishers of some journals will print letters and works which can be pasted on.

For a good elapter on laboratory drawing see A Laboratory Guide for Histology, by Irring Hardesty (P. Blakiston's Son & Co., Philadelphia).

APPENDIXES



APPENDIX A

THE MICROSCOPE AND ITS OPTICAL PRINCIPLES

For an understanding of the optical principles involved in microscopy, four things must be borne in mind with regard to a ray of ordinary daylight:

1. It has an appreciable breadth.

2. It travels in a straight line in a homogeneous medium.

 It is bent (refracted) in passing obliquely from one medium into another of different density.

 It is in reality a composite of a number of different colored rays, ranging from violet to red, and each of these has a different refrangibility.

The amount of refraction undergone by light in a given case depends upon the A//difference in density of the two media which A'the light traverses. Thus, glass is denser than air; hence, in passing from air



obliquely through a glass plate (Fig. 44), a ray of light AB would be bent out of its original course. On reaching the air again, however, it would resume its original direction, although it would be displaced to an amount equal to the distance between A and A'.



It is on account of such displacement that an object in water, for example, appears to be at a different point from where it really is.

On the other hand, after traversing a prism, a ray does not resume its former direction, but takes a new

course upon leaving as well as upon entering the prism (Fig. 45). This new direction is always toward the have of the prism, and the amount of deviation depends upon the shape and density of the

prism. If the base is down, then the ray is bent downward; if the aper is down, the ray still deviates toward the base, that is, it is bent upward.

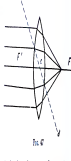
Lenses.-Each of the two principal forms of lenses is in effect practically two prisms, (1) with the bases placed together (Fig. 46, a,

> convex level, or (2) with the apexes together (Fig. 46, b, concare level). In the convex lens, since rays of light are refracted toward the bases of the respective prisms, they will converge; in the concare lens, for the same reason, they will diverge. The terms converging lens and diverging lens, b therefore, are used frequently as synonymous with the terms "convex lens" and "concare

lens." All lenses are modifications or combinations of these two types.

If parallel rays of light pass through a convex lens (Fig. 47) they are so refracted as to meet in one point *P*, which is termed, in consequence, the focal point or principal focus. If, on the other hand, the source of light he placed at

the focal point, then, after traversing the lens, the rays of light will emerge parallel. If parallel rays of light eame from the opposite side of the lens, manifestly there would be a second focal point at P'. The two principal foci are termed onjugate foci, and will be equidistant from the center



of the lens when both sides of the lens have equal curvature.

The ray which passes through the center of the lens (Fig. 47, ϵ) and the focal point traverses what is termed the principal axis of the lens. The optical center of the lens is a point on the principal axis through which rays pass without angular derivation. It may be within or outside the lens, depending upon the form of the latter.

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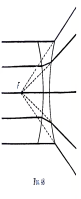
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Any line (ef), other than the principal axis, which passes through the optical center of the lens is termed a *secondary axis*.

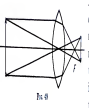
In the case of a concave lens, pacallel rays will be caused to diverge (Fig. 48) and the principal focus, *P*, of the lens is determined by the extension of the divergent rays till they meet at a point which the categories of the divergent rays till they meet at a point which

lies on the same side of the lens as the source of light. Such a point has no actual existence, and is known, consequently, as a virtual focus. The focus of a convex lens, on the other hand, is real, and may be determined readily by allowing the sun's rays, which are practically parallel, to pass through it on to a screen.



By moving the lens backward and forward, the spot of projected light varies in size and brightness. When smallest and brightest the spot is at the focal point of the lens.

Images.—In Fig. 49 the object, represented by an arrow, lies beyond the principal focus of a convex less as in a photographic camera, for example, or the objective of a compound microscope.



Light rays pass out in all directions from any luminous point. Hence, one ray from any point on the arrow, the tip, for instance, will pass through the focal point, *P*, and

one will pass through the optical center of the lens. From what was determined above, manifestly the ray through *P* will emerge as one of the parallel rays upon leaving the lens, and the one through the optical center of the lens, since it traverses a secondary axis, will not be refracted, hence the two rays must cross. Their point of intersection is the point at which the image of the arrow-inp will

be formed. The same fact may be determined, likewise, for any other point of the arrow, for example, the opposite end. Thus the distance from the lens at which the image is formed may readily be determined. In focusing a photographic camera, for example, the image comes sharply into view on the ground-glass plate at the back of the eamera when the plate is brought into the plane in which these rays through the focus and the optical center intersect beyond the lens. It will be observed from the figure that the image is reversed. The size of the image diminishes as the object like farther beyond F.

In case the object lies between the lens and the principal focus, as in Fig. 50, parallel rays from the object would converge to meet at the conjugate focus F', and an eye at this point would see the

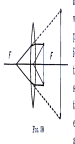


image projected and enlarged without being reversed. The plane in which the image is formed is determined by finding the points of intersection of the secondary axes through points of the object with the imaginary elongation of the refracted rays as shown in the figure. The

image is magnifed because the observer judges of the size of an object by the visual angle which it subtends. The greater the convenity of the lens the shorter the forus, and also, since the mays are bent more, the greater the magnification.

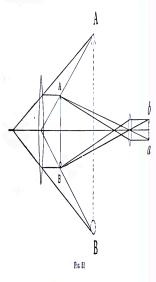
The Simple Microscope.—The simple microscope (the ordinary so-called magnifier, etc.) operates upon this principle; the image of an object is projected and enlarged but not inverted (Fig. 50).

The question arises as to why there is a bot distance to hold the simple microscope from an object. Why will not any point answer so long as it is within the focal point? As a matter of fact, the object may be placed at any point within the focus, and it will be found that the nearer it is brought to the lens the less it is magnified. There is one most favorable point for observation, however, which is neither

at the point of highest nor of lowest magnification, but an intermediate point, where the lens is freest from chromatic and spherical aberrations.

In reality the eye forms an integral part of the optical arrangement when the microscope is being used, but in our elementary exposition of the subject it is disregarded.

The Compound Microscope.—The general principle of the compound microscope is represented in Figs. 51 and 58. The object of (Fig. 51) his beyond the principal focus of the first lens or *objective*



(really a system of lenses), hence the image AB is reversed. This image, in turn, is viewed through a lens, the equiver or order situated nearer the eye of the observer. The ornhar acts as a simple magnifier, projecting and enlarging the image but not reversing it again. As a matter of fact, the ordinary order of a compound microscope cannot be taken from the instrument and used as a simple magnifier because it is marke of two planocourse lenses which are so adjusted that the image from the objective of the compound microscope is not brought to focus until it has traversed the larger or field-lens of the expeisee (Figs 55, 58). The image is really examined, therefore, at a point between the two lenses of the expeiser. Such an evepice is

termed a negative eventice or outhar and is which used today for microscopical work. The commonst form, the Huggenian, is an adaptation of an ocular designed originally by Huygens for the telescope. By contracting the area of the real image, the fieldlens of a negative ocular not only brightens the image but also



|-lach Objective |-lach Objective]-lach Ol-Immersion Objective Fiz. 32.—Lons Systems of Varius Objectives Bausch & Lomb [-lach, l-jach, and j-lach ol-Immersion objectives, respectively

increases the size of the field that can be examined. It is usually also designed, in conjunction with the eye-lens, to help render the image achromatic.

Positive evenieses are also made. An inverted image of the object is formed below the system of ocular lenses. Such an ocular operates as a simple

Fin. 33

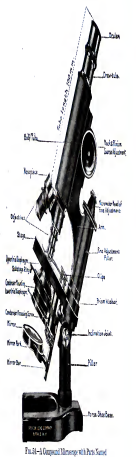
microscope. A good objective is made up of from two to five systems of leases, as shown in Fig. 32. A singlesystem in turn may be a *doublet* (Fig. 57) or

a tripld, each made of different kinds and shapes of glass. A good objective is a very delicate piece of apparatus and must be handled with great care. Each component is very accurately ground and the systems distanced with extreme precision in order to get a clear image. If not already familiar with the parts of the com-

pound microscope; the student should study Figs. 54, 55, and 58, with a microscope before him.

DEFECTS IN THE IMAGE

Spherical Aberration.—A simple convex lens, unless corrected, will not give a sharply defined image because it does not refract



to the same degree all rays passing through it. Those which traverse its edges are brought to a focus nearer the lens (Fig. 53). This results not only in an indistinct image but in a distortion of shape as

Fieldlers

-Draw Tube

Body Tube

well. Straight lines, for example, appear curved, and when the parts of the object are in focus in the center of the field, those nearer the margin are hazy and indistinct. This defect is greatest in strongly curved lenses, that is, since magnification increases with increased curvature, in high powers. Spherical aberration is corrected by one or more of the following processes: 1. Cutting off the margi-

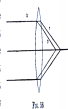
 1. Cutting off the marginal rays.
 2. Changing the shape of

the surface of the lens. 3. Combining several lenses equivalent to a single lens.

Chromatic Aberration.— As with a prism, ordinary light in passing through a lens is broken up into its component colors. This process is

technically termed dispersion. Since the colors are not all bent to

the same ertent, the result is that each color has a different focus; the ones which are bent most (violet rays) come to a focus nearest the lens, and those which are least affected (red rays) meet at a point farther away (Fig. 56). This failure of the color rays to meet in



one focal point is termed chromatic aberration, and if uncorrected

Draw Tube Diaphragm

houric

Baceliens Middle lens Front lens (Object ive

Fig. 55.—Sectional View of Microscope Tube Including Ocular and Objective.



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Length

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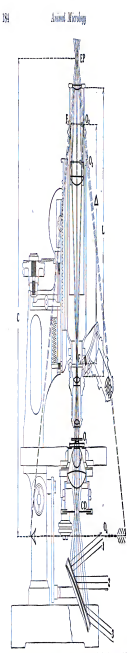
Option

causes the image of an object viewed through such a lens to be bordered by a colored halo.

The defect is corrected by properly combining glasses of different dispersive powers but of kindred refractive powers. Fint glass (silicate of potassium and lead), for example, has a dispersive power equal to about twice that of crown glass (silicate of potassium and lime), although their refractive powers are nearly the same. By combining a biconvex lens of crown glass with a coneave lens of flint glass so constructed that its dispersive c r power will just equal that of the crown glass (Fig. 57), the error may in large measure be corrected. Such an arrange-Pro 57 ment does not interfere seriously with the refractive powers of the lens so constructed. Unfortunately no two kinds of glass have been found which have proportional dispersive powers for all colors, so that in the ordinary achromatic objective only two of the different colors of the spectrum have been accurately corrected and brought to one focus. The colors left outstanding form the defect known as a secondary spectrum. In the apochromatic objectives (p. 188) three rays are brought to one focus, leaving only a slight tertiory spectrum.

NOMENCLATURE OR RATING OF OBJECTIVES AND OCULARS

Oculars—Different makers, unfortunately, use different systems in marking their lenses to indicate relative powers of magnification. In the case of lettering the system is wholly arbitrary; the only rule is that the nearer to A the letter is the lower the magnification. When the objective bears afgure it is usually indicative of the magnifying power of the part marked. Thus a $\frac{1}{2}$ sinch objective magnifies approximately 120 diameters; a $\frac{1}{2}$ inch, 80 diameters; a $\frac{1}{2}$ inch, 20 diameters; a 1-inch, 10 diameters; a $\frac{1}{2}$ inch, 80 diameters; a $\frac{1}{2}$ inch, 20 diameters; a 1-inch, 10 diameters; a 2-inch, $\frac{1}{2}$ diameters; a $\frac{1}{2}$ inch d



For No-Diagram Showing Path of Light Roys through the Compound Micro-scope Topother with Images from Dausch and Londo stabilization. A, upper ford plane of objective. F, however ford have of exploses 4, optical table length - distance between P, and PF: (0, object: 0, real Image 1 P, transposed by the objective length - distance of the projection of the projection distance of the projection distance of the projection distance of the PM and the projection distance of the PM and t



Fig. 58.—The Spencer No. 15 Compound Microscope



Fig. 60.—The Bausch & Lomb CAS Compound Microscope

and so on. Such magnification is termed the initial magnifying power of the objective.

The objectives of most manufacturers are now rated in millimeters and the conventional distance of vision taken as 250 mm. An objective of 3 mm. focus, therefore, yields an initial magnification of 83.3 diameters ($\frac{1}{2}$ X20=83.3). Compensating oculars (see below) hear numbers which indicate the number of times the eyepiece, when used at a given tube-length, increases the initial magnification. Ocular 12, for example, with a 3-mm. objective would yield a magnification of 83.3X12=1,000 diameters, with a standard length of tube. Unfortunately this simple system does not apply to most onlinary oculars, which are more or less arbitrarily lettered or numbered.

SOME COMMON MICROSCOPICAL TERMS AND APPLIANCES

(Alphabetically Arranged)

Achromatic Objective.—An objective corrected for chromatic aberration (p. 182). The correction is not absolute.

Achromatism.-Freedom from chromatic aberration.

Angular Aperture.—The angle (measured in degrees) formed at the point of forus (F, Fig. 62) by the outernosit rays (AF, IF) which traverse the objective to form an image. This angle is an important consideration because on it depends in large measure the defining or resolving power of the objective. It is evident that the large the angle is the grater the number of rays of fight that will be a larger the angle is the grater the number of rays of fight that will be a larger the angle is the grater the number of rays of fight that will be a larger the angle is the grater the number of rays of fight that will be a larger the angle is the grater the number of rays of fight that the larger the same to be the same to the same transmitten and the same angular aperture, will have the same brief have it they are of the same angular aperture, on the other hand, if they have the same magnifying power, but differ in angular aperture, the ballhangy is reduced in the one of smaller angle. In immersion haves the liquid used between the lens and the object, by reducing refraction, has the effect of increasing the angle of aperture. New "Immersion Objective," also "Numerical Aperture" (pp. 196, 199).

Apertometer.-An instrument for measuring both the angular and the numerical aperture of objectives. It is fitted to the stage of the microscope.

Aplanatism — Freedom from spherical aberration (p. 181). The result is a flat field as viewed through the microscope. Aplanatic lenses are usually also achromatic.

Appriment: Objective—In improved form of objective which is more easily achievable than the collinary objective because it is corrected for rays of three colors instead of two, and this correction is equally good in all parts of the field. In the ordinary schematic objective after correction there is a residue of color which is known as the secondary spectrum. In the appelermentic leaves correction is made for a third color, and usually



Fig. 61.—The Bauch & Loub KA Bincenlar Disserting Microscope, Greenough Type.

only a slight tertiary spectrum is left uncorrected. Spherical aberration is also more fully corrected. Furthermore, in these objectives the foci of the optical and the chemical rays are identical, hence the lenses are well adapted to photography. In the glasses of the appelmenties, silicon is replaced by boron in the first series and by phosphorus in the coven series. Fluorite was used in compaction with the glasses in the earlier forms of appelmentation lenses with the result that the lenses frequently deteriorated

in warm, moist climates. Several makers are now able to construct apochromatic objectives without the use of fluorite. Both dry and immersion apochromatics are made,

Binocular Microscope .- A microscope adapted to vision with both eves at once. One of the most important advances in microscopy during the past ten years has been the development of binocular microscopes with erecting prisms which enable one to carry on dissections, to study thick injected preparations, and to perform other manipulations under higher power and otherwise more advantageously than formerly. In general, they consist (Fig. 61) of two optically distinct tubes so combined that the objectives focus on the same point from different angles. A magnified, stereoscopic vision is thereby provided, so that objects which have depth stand out in pronounced relief. The upper parts of the tubes may be rotated so as to adjust the eye-points of the oculars to the width between the pupils of the observer's eyes. If, upon closing one eye and then the other, an image is not seen by each eye without moving the head, the eye-points are

too close together or too far anart. The oculars should be separated or approximated accordingly. When they are correctly adjusted one should get a distinct, stereoscopic alappearance. Other adjustments are provided to compensate for differences of focus in the right and the left eye.



A very simple form of binocular magnifier, known as F the Hardy Binocular Loop, may be obtained from F. A. Hardy & Co., of Chicago, Illinois. It is worn like a pair of Fig. 62 spectacles. A more elaborate binocular magnifier, to be worn with an elastic headband, may be secured from the Bausch & Lomb Optical Co., of Rochester, New York,

Binocular Compound Microscopes with but a single objective are also obtainable. By means of a prism which extends partly across the field about half of the light is directed into the left eye, the rest passing unobstructed to the right. While permitting of higher magnification, this type of binocular is not as universally serviceable as the other form described. The image is not erected.

Brownian Movement or Pedesis.-An oscillating or dancing motion observable in small particles in a liquid when seen under the microscope. Calibration of Microscope.-See "Micrometer" (p. 197).

Camera Lucida .- An apparatus containing a glass prism or thin glass plate so arranged that when it is placed over the eveniece of the microscope, the observer may see the image of the object under the microscope projected on to his drawing-paper on the table. The point of the pencil is also visible; consequently the outline of the object may be readily traced on the paper. In the simpler earnera Incidas a thin neutral-tint glass slip is so arranged that it is in alignment with the evelens of the ocular, except that it sets at

an angle of 55 degrees to it. When the microscope is tilted into a borizontal position the observer sees the image of the object reflected from the upper side of the glass slip, but, since the latter is semerical transparent, he also sees the while paper spread below on the table (Fig. 63).

Another form of simple camera lucida is the Wollaston. To use it the microscope must be inclined. The essential part of the camera consists



of a quadrangular prism. The eye of the observer is so placed over the edge of the prism as to receive rays of light from the object with one portion of the pupil, and from the drawing-paper with the remainder.

Some form of the Abbe camera lucida, however, is used by most workers. It consists of a cap which is fitted immediately above the evenues and which contains two right-angle prisms cemented together to form a cube

(Fig 64). The lower one of the prisms is shvered along its cemented surface, although a small central opening is left through which the object under the microscope may be viewed; connected with the cap is an arm which bears a mirror, and his mirror may be so adjusted as to reflect the image of the durving-paper on the table on to the prisms from one side. The prisms



are so set that the silvered surface of the lower one reflects this image upward to the eye of the observe which also, coincidentially, is viewing the magnified image of the object through the hole in the silvering. When proper adjustment of the hight received from object and paper respectively is made, a pencil-point may be distinctly seen when brought into the field of vision over the paper, consequently the outline of the object may be accurately travel.

The secret of success in working with a camera lucida is to have the illumination in the two fields properly balanced. Small screens of tinted glass are provided with the instrument for such regulation. With the Abbe camera huida the microscope may be used in a vertical or in an inclined position. If the microscope stand is inclined, the drawing-board upon which the paper rests must have the same inclination, or the outline when drawn will be distorted. Likewise, if the mirror of the camera is at any other angle than 45 degrees, an adjustment of the drawing surface must be made; in short, the axial ray of the image and the drawing-surface must always be at right angles to prevent distortion. This means that if the mirror is depressed below 45 degrees the drawing surface must be tilted toward the microscope trice as much as the mirror is depressed. For example, if the mirror is depressed to 37 degrees (8 below 45 degrees), the drawing-board must be tilted (raised) 16 degrees. See also remarks under "Micrometer" (p. 197). When the camera is in proper position the field of the microscope should appear at about the same size as without the camera. If the field is reduced or unevenly lighted, the camera is too near or too far from the ocular, or it is tilted, or the prism is not properly centered.

Compensating Order.-A specially designed eyepices for use with appehrmatic lenses. It was found advantageous to undercorrect the objective and then to restify the abstration by overcorrecting the order. The so-called searching order is a low-power compensating coular used for the first finding of objects. The object once located in the field, the higher working coulars are used in observation.

Condenser.—A less or a series of lenses mounted in a substage attachment for the purpose of conventrating light upon the object to be examined. They are made in various guides of excellence, non-achromatic, achromatic, and appedromatic. Some wild-endpe condensers are used as immersion condensers; the immersion fluid is placed between the upper surface of the oundenser and the lower surface of the object-sible. Condensers are especially valuable with high-porce objectives and of-immersion lenses. For the best results the condenser must be accurately centered and the object must he at the apex of the some of light formed by it. Unintelligent use of the condenser is a very common fault. Condensers are constructed to receive parallel rays of light, hence the plane mirror only should be used with them if the illumination is from daylight. See "Ellumination" (p. 194).

Correction Collar.-A device for adjusting the distance between the leas systems of objectives so that the proper corrections may be made for different thicknesses of cover-glass. Low-power objectives are not so sensitives of high power to the influence of the cover-glass. Orthoury objectives, however, are mounted in a rigid setting and corrected for a specific tube-length and a standard cover-glass (about 0.18 mm, thick; i.e., a No. 2). With a cover-glass of different thickness correction should be made by

altering the tube-length of the microscope, lengthening it for a thinner cover and shortening it for a thicker one. With a 4 mm. (\$ in.) or a 3 mm. (\$ in.) dry objective a deviation of as little as 0.05 mm. in the thickness of the coverglass, if uncorrected, is sufficient to obliterate fine details of the object. With homogeneous immersion lenses the defect caused by different thicknesses of cover-glass disappears (see "Immersion Objective," p. 195). See also "Tube-Length" (p. 201).

Cover-Glass Correction, Cover-Glass Thickness .- See "Correction Collar" (p. 191).

Daylite Glass .- A specially constructed glass which, when used as a screen with a nitrogen-filled tangsten lamp, yields a light almost exactly like daylight. It gives very nearly true color values. The light is soft like that from a white cloud, and is more comfortable to work with in micros-



copy than any other form of artificial illumination. See Gage, Science, XLII (October, 1915), 534, for a fuller description.

Dark-Ground Illumination .- See "Ultramicroscopy" (p. 201). Demonstration Microscope.-A microscope designed to be passed around a class with specimen in place. Most of the compound types are in the form of adjustable tubes

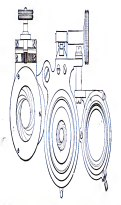
Fro. 65.-Inconfescent Electric Lamp which, when in use, are pointed for Use with Microscope. toward a window or a lamp.

Demonstration or Pointer Ocular .- An ocular provided; at the point where the real image of the object is produced, with a delicate rod of some kind which may be rotated to point out objects in the field. A simple type may be made by cementing a hair across the opening of the ocular diaphragm with balsam. When the balsam is hard the hair is cut at the center of the opening and one end is removed. It is necessary to have both ends of the hair supported until the balsam hardens, otherwise the free ends will sag and not be in focus. To use, rotate the ocular.

A double demonstration eyepiece, by means of which the image formed by the objective can be viewed simultaneously by two observers, has also been devised.

Diaphragm .- Opaque plates with openings of various sizes for regulating the illumination of the object to be examined. The iris disphragm (Fig. 66) is the best type. It consists of a series of overlapping plates placed around a central opening the size of which may be varied by means of a lever. Revolving diaphragms are commonly used on the cheap grades of microscopes. They consist of round disks perforated by openings of various sizes

which may be rotated between the mirror and the object. The maner to the object the diaghangm is placed the better the intensity of the illumination can be regulated. Most of the better class of microscopes are provided with two inis diaphragms, one beneath the condenser to be employed when the latter is in use, the other flush with the stage to be used only when the condenser is out. If this second it is diaphragm is lacking, its place is taken by means of a cap-diaphragm which may be titted into the substage in the place of the condenser. A control-keep diaphrapm is one with an opaque center and a sit around the edge, so arranged that a follow cone of light, consisting of rays of great obliquity, will be produced.



F1a, 66.—Top View of a Substage Attachment with Condenset and Lower Iris Diaphragm Thrown out of Optical Axis.

Dissecting Microscope—In instrument to constructed as to enable an operator to earry on minute dissections under magnification. Ordinarily they are simple microscops mounted on a stand of some kind. The best instruments (Fig. 67) are provided with well-corrected lenses, with glass stage, mirror, black-and-white substage plate, and rests for the hands. See also Figs. 68 and 69 for moldined forms. The binocular type, Fig. 64, is indispensable for the finer modern technique. See "Binocular Upe, Fig. 64, is

Embryograph — A form of camera huida for drawing at slight magnifractions small objects, such as embryos. A camera huida attached to a simple microscope is frequently used for this purpose.

By-Point.—The point above an order or lens at which the largest number of rays from the instrument enter the eye. The largest field of the microscope is visible from this point.

Flatness of Field .- See "Aplanatism" (p. 187).

Homogeneous Immersion Objective.—See "Immersion Objective" (p. 195).

Huygenian Ocular.-See p. 180.

Illumination — hay means employed to direct light upon the object under observation. Light which traverses the object is said to be transmitted light. Most microscopical work in biology is done by means of transmitted light, hence the object must be readered more or less transparset if not naturally so. If the object is symmetrically lighted, the lighting is designated as acrial or control illumination. If one side is lighted more than another, the term obligat illumination is employed. In the case of transmitted light, the light which traverses the object is usually hight reflected from a micror because it is generally incorrected to impossible to hold the instrument directly toward the source of light. Makers of microscopical



Fig. 67.-Dissecting Microscope

appliances, however, now supply admirable miniature electric lamps which may be used with the mirror or in place of it (Fig. 65).

Light which falls upon the object and is reflected from it to the eye, either directly or through a microscope, is termed reflected light. Such illumination is employed but little in ordinary histological work, but it is useful in the examination of opaque objects such as metals, inserts, etc. The illumination may be increased by means of a bull'scope condensor or a mirror. In some microscopes the mirror can be swang above the stage for the surpose of illumining an object which is to be studied by reflected light.

The best light for microscopical work is light reflected from while clouds. Direct smilpht is never used. The light should some from in front of the observer of from one side. Various kinds of artificial light are used for microscopical work, such as an ordinary hanp with flat wick, the Welshach, or the ordinary electric light. Some of the never electric lights especially designed for microscopy are excellent. Whatever the source, the rays must be standy and buillant. If a lump with flat wick is used, practice builliancy is seemed when the edge of the flame is turned toward the microscope; the object should be lighted directly by the image of the flame. To do this with low powes, the lump may have to be turned so that the flame is oblique to the microscope.

In artificial light the rays are diregent, not paullel, as in the case of sunlight, hence they will not come to focus at the same point when reflected from the mirror as the latter do. This should be corrected by using a large bulk-eye condenser between the source of light and the mirror, or by shiling



Fro. 68.—Lens Holder with Flexible Arm

the mirror slung the mirror-her farther away from the stage so that the coneave mirror will have a longer distance in which to bring the mays to focus. If a substage condenser is used, the same results may be obtained by depressing the condenser somewhat below the level of the stage. Lamps made for the microscope often have a metal chimney with a bull-seye in one side.

The objectionable yellowness of most artificial light may be eliminated by interposing a piece of green signal-glass between the hump and the mirroscope. With most mirroscopes, round slips of blue glass which fit into the employ as a screen an annonia subplate of copper solution in a globular field. To make the solution, dissolve a small annount of copper solution fask. To make the solution, dissolve a small annount of copper solution water and add annonia. At first a precipitate appears, but if an eccess of annonia is added this is dissolved and a transparent deep-blue liquid results. This should be diduced with water softiciently to get a blue of just the paper depth to render the transmitted light white as seen through the microscope. The globular hash also acts as a condenser.

Immersion Objective.--A kind of objective in which a liquid is used between the front lens and the cover-glass. Cedar oil is the most widely

used medium. Inasmuch as the optical properties of evaluar of (refraction and dispersion) are almost the same as envorm gives, it is often termed a homogeneous immersion fault. A homogeneous immersion heat, therefore, would be one intended for use with such a fluid: The advantage of an immersion over a dry less loss in the fact that, other things being equal, after learning the onver-glass rays which would be so refracted in a rare medium.

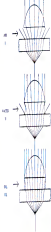


Fra. 40.—Eigh-Power Dissorting Lons This leas, represented as in use on the stand of a disserting microscope, is provided with two double reflecting from persons which even the image so that the operator works as with a simple lons.

like air as to miss the front less of the objective reach this less in the case of immersions and traverse the objective. With homogeneous immersions the rays of light are varied without deflection through cover-glass and finid and into the glass of the front lens. Water has a greater density than air and less than glass; hence, with a *voiter* immersion more rays of light reach the front less than with a dry lens and less than with a homogeneous immer-

sion lens (Fig. 70). . The effect of an immersion is practically to widen the angle of the lens (see "Angular Aperture," p. 187). The value of the immersion objective is enhanced if the immersion fluid is placed between the upper lens of the condenser and the slide as well as between the objective and the cover-slip.

to multiply the apparent dimensions of an object viewed through it. It should be expressed in diameters, not in areas. While magnifying power is very important, it is only so in connection with resolving power. If high power were the only essential, a series of single lenses might be used. The impossibility of using such a series for high magnifiestion is due to the fact that proper correction of aberrations cannot be made, and, consequently, a distinct image cannot be obtained. Immession, and Homogeneous For determination of magnification see Oil Immersion Objectives Ba spectively (after Bausch). "Micrometer" (p. 197).



Fre. 70.-Diagram to Illus-

Mechanical Stage .- A stage attachment (Fig. 71) for the more accurate manipulation of an object or a series of objects which must be moved about



under the objective. The best mechanical stages are provided with scales and verniers so that an object once recorded may be easily found again. They are often very serviceable, especially with high powers. Micrometer.-A scale for

measuring objects under the microscope. The stoge micrometer consists of a finely divided scale (1; and 1; mm.) ruled on glass or metal. It is commonly mounted on a glass slide of standard size. To determine the actual size of an object with the stage micrometer, it is most convenient to use a camera lucida. The outline of the object to be

Pro. 71.—Attachable Mechanical Stage for Microscope.

measured is projected on to a sheet of drawing-paper and marked off. The object is then replaced under the microscope by the micrometer and the micrometer scale is projected on to the paper. Knowing the actual distance between the lines on the micrometer scale, the magnification as well as the real size of the object is readily calculated.

The size of the image projected by a camera lucida on to a piece of drawing-paper at the level of the table, however, does not represent the true magnifying power of the microscope. The latter is really considerably smaller if the microscope is in a vertical position because the magnification of a lens or a system of lenses is calculated in terms of the conventional distance of vision (250 mm., see p. 183) while the distance from the ocular to the table is considerably more than 250 mm. Since the rays of light diverge after leaving the ocular, manifestly the projected image will be larger (possibly by as much as 60 per cent) at the level of the table than at a level just 250 mm. from the point of emergence of the rays from the ocular. To determine the actual magnification of the microscope, therefore, one would have to bring the drawing surface to within 250 mm. of this point of emergence, sketch the projected scale of the stage micrometer on the paper, and then, by means of an ordinary metric rule, compute the number of times the divisions of the micrometer scale have been magnified. The standard distance of 250 mm., if the Abbe camera lucida is used (with camera mirror at 45 degrees), includes the distance along the mirror-bar from the optical axis of the ocular to the mirror, plus the distance from the mirror to the drawing surface.

In practical work it is not necessary to make drawings or measurements exactly at this standard distance; one needs only to have a scale made out for the distance from the exames heids at which the drawings are astrally to be made, although it must be earcfully borne in mind that any variation in the elevation of the drawing-surface will alter the size of the projected image. A series of earcfully prepared scales for various combinations of objectives and oculars should be made and kept for hotone use. On each should be recorded the tub-length used, the number of the dojuetive and of the coulter, the length of the eamers miror-bar, and the angle of the mirror, for if any one of these is changed the scale is no longer accurate.

When much measuring is to be done an ocular micrometer is used. It consists of a circular glass disk with a scale ruled on it and is inserted in the ocular between the eye-leas and the field-lens. By means of a stage micrometer the value of the divisions of the ocular micrometer is determined for a known tube-length and every combination of leaves it is desired to use in the work of measurement. Suppose that it takes four divisions of the ocular micrometer to correspond to one of the finer divisions of the stage micrometer, then since the divisions of the latter are equal to γ_{17} mm, each space in the

ocular micrometer must be equal to 717 mm., that is, 0.0025 mm. A flor or acress micrometer is a more convenient form of ocular micrometer, which is provided with delicate movable spider lines that can be adjusted to the space to be measured by means of a fine screw with very accurately cut threads (Fig. 72). At the end of the screw is a graduated disk which gives the value of the distance between the spider lines. The pitch of the screw is either the inch or 0.5 mm. When once the valuation of this ocular micrometer has been determined by means of a stage micrometer, measurements can be made rapidly and with great precision.

The step micrometer, in which the intervals are arranged in groups of ten, each group being conspicuously marked by a black, stairlike notching along one side, is one of the most desirable types of ocular micrometers.

Micron.-The one-thousandth part of a millimeter; expressed briefly

by the Greek letter u It is the unit of measurement in microscopy.

Mirror .-- The compound microscope is usually provided with both concave and plane mirrors, which may be rotated or swing in any direction. The plane mirror is used with the condenser; the concave, whenever it is of advantage to have light concentrated upon the object, with the con-



denser out. The mirror should be capable of being moved up or down the mirror bar so that it can be accurately focused upon the object. See also "Illumination" (p. 194).

Muscae Volitantes.-Small filaments or speeks which float across the field of vision. They are really small opacities in the vitreous humor of the eve.

Numerical Aperture .- A system which expresses the efficiency of an objective by indicating the relative proportion of light rays which traverse it to form an image. With the introduction of immersion objectives, it became evident that angular aperture alone is not sufficient to indicate the real capacity of an objective. For instance, an immersion and a dry lens may be of precisely the same angular aperture, and yet the immersion lens is more efficient because it sends more rays of light through the objective (see "Immersion Lens," p. 195). It was found necessary to take cognizance

of the medium which intervenes between the cover-glass and the front lens of the objective.

Professor Abbe, in 1873, proposed the name numerical querture and introduced the formula N.A.= u sin u, in which a signifies the refractive index of the medium between cover-glass and objective, and u equals half the angle of aperture. That is, by multiplying the refractive index of the medium by the size of half the angle of aperture the numerical aperture is obtained. For example, suppose that one had an oil-immersion less of 90 degrees angular aperture, then half the angle of aperture is 55 degrees, and by turning to a table of natural sizes, the size of 45 degrees is found to be 0.707. The refractive index of eachs of is 1.2. Then N.A.= 1.32X 0.707=1.053. Suppose that the lease were a dry instead of an immersion less; then since the refractive index of an is 1, the formula would read N.A.=120.707=0.707. Thus the two products 1.075 and 0.707, respectively, represent the relative capacities of an illimmersion and a dry dojective of 90 degrees angular aperture.

Partoral—A term ordinarily applied to evolvess of different powers that may be exchanged in the microscope without very materially affecting the lows of the instrument. The term is also applied to objective statehed to a revolving userpiece if each is approximately in focus when turned into place.

Pedesis.-Same as Brownian movement,

Penetration.—The quality of an objective that permits of "hoking into" an object having sensible thickness. It is greatest with low powers and nervor angles and is natagonistic to resolving power. It is the natural consequence of certain conditions in the making of lenses and is reduned of secondary importance, begave practically the same results are obtained by manipulating the fine adjustment.

Photonicrography.—Photography of small or microscopic objects. The subject, although of great importance, is too extensive to enter into in the brief space that could be albitted to it in an elementary treatise such as this. (See, however, p. 150.) An excellent chapter on photomicrography and a bidilography will be found in Gage's *The Microscope*.

Pointer Ocular.-See "Demonstration Ocular" (p. 192).

Polarisope.-As used in microscopy the polarisope consists of two parts, each composed of a Nicel prism of lockand spar, one, the polarisor, fits into the substage, and the other, the outlare, is inserted between the objective and the tithe of the microscope or, in some forms, just above the coultr. The polarisope is used more in chemical and in geological than in histological work. Some of the uses are as follows: determining whether an object is singly or doubly refractive; detecting the presence of minute crystaks, determining the composition of nexts; ceamining sections of bane,

hoof and horn, hairs and fibers of animals and plants, starch, etc., for certain , characteristic and striking effects.

Projection Ocular.—An order specially designed for projecting a nicroscopic object on to a screen or for use in nicrophotography. While ordinarily used with appendence objectives, they may be used with ordinary objectives of large numerical aperture. The systems is movable so that a sharp focus (molinated by a distinct image of the disphragm) may be obtained at different screen distances.

Reading Power-The quality of an objective which eachies the observer to make out fine details of structure. It is the most essential property for precision in observation, and determines largely the excellence of an objective. Readving power depends upon earchil correction of alcentations, general accuracy in the mechanical construction of the microscope, and upon the aperture of the objective (see "langular Aperture," "Numerical Aperture," pp. 187, 189). Readving power is tested by the resolution of the parallel lines ruled on glass or the strine on the surface of diatoms. The test is to determine how many lines to the inch or continuetier may be distinguished, and whether the objective simply glimposes the ratings or whether it resolves them clearly. The where the angle of aperture the better the resolving power, provided the width is not so great as to interfere with the correction of the leaves. The increased resolution of immersion leases is due to the that the immersion fluid practically whiless the angle of aperture (see "Immersion Objective," p. 185).

Tub-Length—The distance between the places of insertion of orchar and objective into the table of the microscope. There are two standard tube-lengths; the short standard is 160 mm. (b); inclus), the long standard, 216 mm. (b); inclus). Some makers, however, do not adhere to the standards. The optical efficiency of the instrument is the same in either case. The short length is more abrantageous in that it is more compact. The lenses must be corrected for the length of tube with which they are to be used. The short standard is in use in must American laboratories.

Utramicroscopy.—A system of microscopical inspection in which objects are examined by reflected light. The object appears to be self-luminous against a dark field, hence the term dark-pround illumination is often used as descriptive of the method. Objects to be studied in this way are usually semi-transparent or consist of fine particles such as occur in colloidal suspensions. In highting rays of great obliquity are used so that only such traverse the objective as are deflected from some object in the field. The great value of the method lise in the first that particles may be realised which are which invisible with the microscope as originarily used.

For low powers without a condenser, the diaphragm must be wide open and the mirror so tilted that the object is lighted by oblique rays which

cannot get directly into the front less of the objective. With a condenser, a control-stop diophrapm is used which admits only marginal rays. By making an ordinary displaragm eccentric, somewhat the same effect may be second. For practice, place a drop of 10 per cent solution of salveylic acid in 55 per cent slowhol on a side and leave it until the slowhol evaporates. Examine the residue of crystals in the ordinary way and then by darkground illumination. By the latter method the crystal should appear bellliantly lighted on a dark lookground. Add a small drop of the solution to the crystals and watch crystallization under dark-ground illumination.

For kiah powers, according to one system, very wide apertures (greater than 1.00 N.A.; see p. 199) are nexessary in the condensers. Some makers (e.g., Leitz, Reichert) use specially modified condensers. Others (e.g., Beck, Sedentonf) substitute a parabolic reflector for the condenser. In the method of Siedentopi and Zsigmondy, the field is lighted from one side, at right angles to the axis of the microscope, by a wedge or cone of bright light. In another method, useful for both high and low powers, an objective of wide aperture and a condenser of moderate aperture are employed. The field's lighted, as in ordinary microscopy, by a cone of light from the condenser, but a diaphragm or stop of the right size to cut out the central rays of light is placed on the back lens of the objective. In this way only those rays which have entered the marginal zones of the objective pass on to form an image, and among these are the rays which have been deflected by objects in the field. A somewhat similar result may be attained by using a stop in the eve-point (p. 193). For a fuller discussion of ultramicroscopy and darkground illumination see A. E. Wright, Principles of Microscopy, chap. xiv.

Overcorrection and Undercorrection—In correcting for character shermition, if the concave lens is stronger than is necessary to neutraline the absention of the concave lens, the blue rays are locupit to focus beyond the true principal focus of the objective, and the latter is said to be overcorrected; if the concave lens is not strong enough, the result is what is known as undercorrection. In case of overcorrection, the object takes on an ownge tim if, after focusing, the distance between object and objective is slightly increased; or it becomes of bluish olor if the distance is decreased. In case of undercorrection, just the reverse is true. In some instances the objective is purposely undercorrected, and the oppiece (e.g., compensating coulter) is equally overcorrected.

Working Distance.—The distance between the front lens of the object tive and the object when the latter is in focus. With high powers it is very small, so that with some oblimmersion objectives, if a thick cover is used, it is impossible to focus upon the object. For this reason thin over-glasses (No. 1) should be used on preparations which are to be used with high-power immersion lenses. For examination under high-power dry lenses, however, see remarks under "Correction Collar" (n. 191).

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MANIPULATION OF THE COMPOUND MICROSCOPE

 Always handle the instrument continuely; it is a deficate mechanism. Lift it by the base or by a handle specially provided, never by the tube.

2. The work-table should be of such a height that the observer can sit at it confortably without compressing the class or tiring the neck. Sit as upright as possible. If the instrument is inclined, it should set farther in on the table than if it is in the upright position.

 With a piece of old linen, a chamois skin, or a bit of lenspaper, carefully clean the cycpiece to be used and put it in place. Always use the low-power cycpiece first.

4. Likewise clean and attach the objective (low power first) after elevating the tube far enough above the stage for this purpose. Guard particularly against screwing the objective in crocked, as this will injure the threads. It is best to swing the objective between the first and second fingers of one hand and bring the screw squarely into contact with the screw of the tube (or nosepice); with the thumb and forefinger of the other hand it is then screwel into place.

5. Bring the draw-tube to the standard length (see "Tube-Length," p. 201) for which the lenses are corrected. If a nosepicee is used, allowance must be made for its height. In some of the more recently made microscopes, however, the scale on the back of the draw-tube includes the nosepicee. In poshing in or drawing out the draw-tube includes the nosepicee. In poshing in or drawing out the draw-tube includes the nosepicee. In poshing in or drawing out the draw-tube includes the nosepicee. In poshing in or drawing out the draw-tube includes the nosepicee. In poshing in or drawing out the draw-tube always grasp the milled head of the coarse adjustment also, so that the tube as a whole will not be shifted.

6. Place the slide which bears the object on the stage with the object over the central opening of the latter, and elamp it in place by means of the spring elips. While looking at the object from one side, turn the mirror until a flood of light slines up through the center of the stage.

7. Lower the tube until the objective nearly touches the overglass, then look through the eveptice and slowly raise the tube by means of the owner adjustment until the specimen to be examined is planly visible. Focus accurately by means of the fine adjustment. If a high-power objective is being used, since it must owne very near the overe, the operator should lower his head to the level of the

stage, and look toward the light between objective and cover-glass in order to prevent actual contact. This is of great importance, for otherwise the objective or the object is liable to injury. Remember that in focusing up the lowest part of the object comes into view first, the highest part last. It is often easier to locate the object if the preparation is moved about slightly while focusing.

8. The higher the power the more difficult it is to find an object or a particular part of it. For this reason the finding is usually done by means of a low-power objective, or a low-power ocular, or both, and after accurately centering the object in the field, the high power is attached. In case a revolving nosepiece is used, great care should be used in turning in the high power not to strike the shide with the objective. This is very likely to happen if the objectives are not particel. When objectives are not parfocal they may usually be made so by putting a paper or bristol-kcond collar on them.

9. After the object is in focus give any further attention to the illumination that is necessary (see "Illumination" and "Mirror," pp. 194, 199). If intensified illumination is desired, use the concave mirror, or use the substage condenser and the plane mirror. For ordinary purposes the field should be evenly illuminated, although oblique light is frequently useful. Manipulate the diaphragm until the structure to be studied shows with the greatest distinctness. Too much light "thourse" the object, and is hard on the eyes. (To determine the proper distance at which the concave mirror should stand below the stage, let direct sunlight shine upon the mirror, and then adjust the latter so that the apert of the one of light comes just at the top of the stage where the object will rest.)

If particles of dust or cloudness appear in the field, determine, by moving the slide and rotating the coular one after the other, whether slide, objective, or ocular requires further cleaning. A camel's hair brush is often more effective than lens-paper or cloth in removing bits of dust.

10. In using off-immersion objectives, a small drop of cedar of (specially prepared by the maker of the lens) is applied to the front lens by means of a small rod or brush. It is very important to keep the oil free from dust, and to see that it does not contain air bubbles

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when applied to the lens. Cardully lower the tube until the od on the objective comes in contact with the cover-glass. The operator should lower his head to the level of the stage to observe this properly. Focus up as with a dry objective. For critical work innersion of should also be placed between the condenser and the lower side of the slide. With a piece of lens-paper or a soft cloth clean the innersion lens immediately after you have finished using it. Likewise remove the of from the cover-glass. Oil which has hardened on the coverglass should be removed with lens-paper wet with xylol.

If the immersion oil becomes too dense, as is likely after some months, it may be diluted with pure cedar oil.

11. The range of the fine adjustment is limited. Keep it as near the middle point as possible. If the tube does not respond to the movement of the screw, you have probably gone beyond the range of the fine adjustment.

12. In working with the microscope keep both eyes open. The eye which is not in use soon becomes accustomed to ignoring objects in the field of vision. To avoid faigure it is well to use first one eye and then the other for observation. The eye should be placed at the eye-point (p. 193) of the lens. This is some distance from the eye-lens in low-power eyepiexes, close to it in high-power eyepiexes.

While observing, with one hand keep the fine adjustment moving. This relieves the eye of the strain of attempting to focus on different depths of the object.

13. Put the microscope in its case when you have finished using it, or at least cover it with a cloth or cone of paper. For further details regarding the use or care of the microscope consult one of the following books: The Microscope, by Gage; Principles of Microscopy, by Wright; The Microscope and Its Revelations (1,20) pages), by Carpenter and Dallinger.

14. Do not apply alcohol to any part of the instrument. The leases may be cleaned ordinarily by breathing upon them and wining them with a rotary motion on leas-paper or a piece of soft old linen. In case a solvent must be used for balsam or oil, bearene is the one commonly recommended. It must be quickly wiped away so that it will not affect the setting of the lens. 15. Read carefully in the catalogue of the maker of your instrument what is said about its construction.

16. Determine the magnifications of your various combinations of lenses as described under the heading of "Micrometer" (p. 197).

The beginner in microscopy should acquaint himself with various common objects that are liable to get into his preparations in the form of dust, etc., so that he may not mistake them for essential parts of his specimen. Such objects are hairs, fibers of silk, wool, linen, cotton, and the like, and particularly air bubbles. Air bubbles are usually circular with black borders and bright centers; they may show tinges of color. Examine a drop of salirs for examples.

APPENDIX B

SOME STANDARD REAGENTS AND THEIR USES

I. FIXING AND HARDENING AGENTS

1. Acetic Acid.—Acetic acid is more commonly used in mixtures or in diluted form than pure. It is rahable because it tends to produce good optical differentiation and facilitates preateration. When employed alone it causes some tissues to swell and disintegrate. Inazonech as most fixing agents give the best results when they have an acid reaction, from 1 to 5 per cent of acetic acid is generally added to acidify them in cases they are not naturally acid. Any reagent containing a very large proportion of acetic acid should be allowed to act for only a short time. Acetic acid is also of great value in mixtures because it counteracts the shrinking action of certain reagents. Ordinary acetic acid is of alore 30 per cent strength; glacial acetic, of about 99.5 per cent strength. The latter is meant when mentioned in this book unless otherwise specified.

A strength of from 0.2 to 1 per cent is recommended by Flemming for work on cell nuclei. Strong glacial accine acid is sometimes used for highly contractile animals, such as Coelentersta, Mollusca, and Vernes. The animal is rapidly flooded with the acid and remains immersed until it is thoroughly penetrated (6 to 10 minute). It is then washed in repeated changes of 50 or 70 per cent alcohol and left to harden in 70 to 83 per cent alcohol. The pure acid, if allowed to act for more than a few minutes, swells and softens the tissues. Acetic acid should not be used when connective tissue or celicate calcaneous structures are to be preserved.

2. Acetic Alcohol-Carnoy recommends each of the following formulae:

0)	Glacial acetic acid	1 pert	
	Absolute alcobol	3 parts	
b)	Glacial acetic acid		
	Absolute alechol		
	Chloroform	3 parts	

The chloroform is said to hasten the action of the mixture. Either of these reagents penetrates well and acts rapidly. Solution b is especially good for glandular or lymphatic tissue. Almost any stain will follow them. Even such difficult objects as the eggs of Assaris may be fixed by the second mixture. The reagent should be washed out in alsolute or at least in strong alcohol.

Absolute alcohol to which 20 per cent acetic acid has been added is also in use in Boveri's laboratory for Asearis. Material is left overnight in it.

A mixture of absolute alcohol, glacial acetic acid, and chloroform, equal parts, saturated with corresive sublimate (formula of Carnoy and Lebram), becomes even more valuable for the function of difficult objects. According to Lee, isolated ova of Ascaris are fixed in 30 seconds, entire orihorts in 10 minutes, in this liquid. It is good for cytological work in general.

 Alcohol.—Alcohol is used especially for gland cells and for preserving the brain and spinal cord for Niss's method of staining nerve cells. See "Alcohol Fixation," p. 25; also reagents 1 and 2, p. 7, and memoranda on p. 13.

Alcohol and Chloroform.-See 2, b.

Bensley's Formol-Bichromate-Sublimate Mixture.—See p. 153. Bichloride of Mercury.—See "Corrosive Sublimate."

4. Bichromate of Potassium.—Bichromate of potash is one of the olisist and best known fixing reagents. At present it is more commonly used in mixtures than alone. It is widely used in hardening nerrous tissue. Its fixation of nuclei is unsatisfactory unless it is properly corrected through the addition of aceite axid. It acts very slowly, about three weeks being necessary to harden properly a sheep's eye, and from three to six months for a good-sized learn. A weak solution (2 per cent) should be used at first, to be replaced gradually by stronger solutions up to 5 per cent. When hardening is completed the object should be theoroughly washed in running water and them put into alcohol; begin with low percentages of alcohol and gradually increase the strength up to 70 or 80 per cent. Change the alcohol as often as it becomes yellow. After the object has been placed in alcohol, keep it in the dark in order to prevent a

precipitate forming on the surface. Either earnine or hematoxylin may be used as a stain after bichromate of potach. In ease carmine is used, the staining is best done before the object is placed in alcohol. Tissues which do not stain well should be placed for 3 hours in avrid alcohol and then wached in alcohol before staining.

5. Bichromate of Potassium and Acetic Acid (Tellysnicky's fluid).-

Bichromate of potassium	3 grams
Glacial acetic acid	5 e.e.
Water	100 e.e.

It is best not to add the acetic acid until just before using. This is a good general reagent. It is valuable for embryos. Objects should remain in some 20 volumes of the fluid from 24 to 45 hours, according to size. It is well to change the fluid croce after a few hours. After firstion, tissues should be washed thoroughly in running water (6 to 12 hours) and passed through alcohols of increasing strength beginning with 15 per cent.

6. Bichromate of Potassium and Corrosive Sublimate (Zenker's fluid) .-- For formula, see p. 8, reagent 7.

Zenker's is a valuable respect for both histological and embryological material (embryos up to 25 mm.). Several hours are required for fraction: 2 to 4 hours for a 2-day chick; 8 to 10 hours for objects or embryos of 6 to 8 mm.; 24 hours for embryos of 12 to 14 mm., etc. For washing, running water is employed for from 12 to 24 hours. The object is then transferred to gradually increasing strengths of alcohol up to 70 per cent, leaving it according to sine from 1 to 3 hours in each alcohol. To remove the excess of corrosive sublimate, see 14, "Caution" 1. Almost any stain follows this reagent well. Both nucker and cytoplasmic structures are properly fined.

7. Bichromate of Potassium, Corrosive Sublimate, and Formalin (Zenker formalin mixtures).--

A. Helly's Fluid:

Prepare a Zenker's fluid, but instead of acetic acid add formalin in the same proportion and in the same way. Good for tissues in which it is desired to examine the granular cytoplasmic contents.

B. Danchakoff's Mixture:

Corrosive sublimate	50 parts
Potassium bichromate	25 parts
Sulphate of soda 10	or 12 parts
Water	1,000 parts

Boil to dissolve. Just before using add sufficient formalin to make the solution contain 5 per cent for soft tissues or 10 per cent for dense tissues. Fix for from 2 to 4 hours, never more than 6 hours. Keep the mixture at about 37° C. during fixation. This fluid is useful for some kinds of eptdoletical work.

Bichromate of Potassium and Cupric Sulphate (Erlicki's fluid).—

Bichromate of potesh	5 grams
Sulphate of copper	2 grams
Distilled water,	220 e.e.

Pulverize the crystals before adding the water.

Ericki's fuil is an excellent reagent for general use, and is especially valuable for voluminous objects such as advanced embryos. Its principal dravback is the length of time required property to harden objects (five days to three weeks). The process may be hastened by beeping the fuil containing the tissue at the temperature of an inculator (30° C). At the end of this time transfer the object to 35 per cent alcohol, keeping it in the dark for two hours to avoid precipitation. The alcohol should be changed occasionally during this time. Repeat the process, using 50 per cent alcohol, and finally preserve the material in 70 per cent alcohol.

9. Bichromate of Potassium and Sodium Sulphate (Müller's fluid).--

Bichromate of potassium	to 25 grams
Sodium sulphate	10 grams
Water	1,000 e.e.

Müller's fluid is an old and widely used reagent. It is especially valuable for the nervous system. It acts very slowly. Specimens require immersion in a large quantity of the fluid from three to ten weeks, according to size. The solution should be changed every two

days for the first ten days, and later about once a week. If a serum appears at any time, the fluid should be changed. In washing, the tissues are placed in running water for a number of hours and are then treated with gradually increasing strengths of alcohol in the usual manner. For some purposes, however, the tissue is transferred directly from the fluid to 70 per cent alcohol. In any event, the material should always be kept in the dark to prevent precipitation.

Bouin's Pierce-Formal—See pp. 9 and 29. Carnoy's Acetic Alcohol—See 2. 10. Chloride and Acetate of Copper (liquid of Ripart and Petit).—

Camphor water	75.0 grams
Crystallized acetic acid	1.0 gram
Distilled water.	75.0 e.e.
Acetate of copper	0.30 gram
Chloride of copper	

This is a good reagent for cytological work where objects are to be studied in as fresh a condition as possible. Methyl green (reagent 60) should be used for staining. Only aqueous media are employed with such material.

11. Chromic Acid — Aqueous solutions of from 0.2 to 1 per cent are used. The acid is best kept in the form of a 1 per cent stock solution. Tissues are left in at least fifty times their volume of the acid for from 24 hours for small pieces to one or more weeks for larger ones. The objects are then washed in running water for several hours, after which they are treated with gradually increasing strengths of alcohol. Do the washing and dehydrating in the dark. If sections of chromic-acid material do not stain readily, they should be treated for three hours with acid alcohol, washed out with ordinary alcohol, and then stained. Hematoxylin or some of the anilins are the best stains for chromic material. Chromic acid hardens much more rapidly than bichromate of potach. It makes tissues extremely britte.

12. Chrom-Acetic-Osmic Acid (Flemming's solution) .--

This is the so-called "strong" solution of Flemming. The mixture should not be made until immediately before using, because it deteriorates if allowed to stand for any considerable length of time. The fluid is valuable for cyclological work, especially for the study of karyokinetic figures. Only small pieces of tissue should be used, as the reagent penetrates poorly. They should remain in the fluid for from 24 to 43 hours and then be washed in running water for from 6 to 24 hours. From water they are transferred to gradually increasing strengths of alcohol. Particles of ist are blackmed by the mixture. Sections stain well with safranin or bematosylin. Read the remarks on osmic acid, 21.

13. Chrom-Acetic-Formalin Mixture .--

Chromic acid, 1 per cent solution	16 parts
Glacial acetic acid	1 part

Just before using add to two volumes of this mixture one volume of formalin. This is a good fixing fluid for general embryological work.

14. Corrosive Sublimate (mercuric chloride, thichloride of mercury).—Corrosive sublimate is ordinarily used as a saturated solution in distilled water (about a 7 per cent solution) or in normal submer. The latter keeps better and contains a greater percentage of the sublimate. Corrosive sublimate is an excellent and rapid fixing fluid for many objects (glands, epithelia, etc.). Objects should remain in the fluid only long enough to become thoroughly fixed; this has been accomplished when they have become opupue throughout. Only a few minutes or even seconds are required to fix very delicate objects, but denser tissues may require from 4 to 24 hours. The value of the fluid is usually enhanced by the addition of 5 per cent of glacial acetic acid. Small pieces of tissue (not over 0.6 em. in diameter) should be used where practicable. Washing may be done in running water (several hours) or in 50 to 70 per cent alcohol.

CATHONS.--(1) With corresive sublimate or mixtures containing it, the mercuric salt is often not wholly removed in washing. If the tissues are to remain several days or weeks in alcohol, the alcohol will gradually extract it. If they are to be used within a few days,

however, it is necessary to remove the excess of sublimate by adding a few drops of a 10 per cent alcoholic solution of iodine to the 70 per cent alcohol. Sufficient of the solution is added to give the alcohol a port-wine color; as often as the color disappears the iodine must be renewed. After from 12 to 45 hours of this treatment, the iodine color persists, and the object should then be transferred to fresh 70 or 80 per cent alcohol, which must be renewed until it no longer extracts iodine from the specimen. Some workers prefer not to treat tissues fixed in a mervarie fixer with the iodined alcohol until they are sectioned and on slides. The treatment then requires only about 30 minutes.

(2) In handling corrosive sublimate, a glass or horn spoon should be used instead of a metal instrument, because it corrodes metal.

(3) Use distilled water, not tap water, in making an aqueous solution.

15. Corrosive Sublimate and Acetic Acid .--

Corrosive sublimate, saturated aqueous solu-	
tion	100 parts
Glacial acetic acid	

This is an excellent reagent for embryonic tissues and for organs which do not contain a very great amount of connective tissue. See remarks under 14.

16. Corrosive Sublimate, Nitric-Acid Mixture (Gilson's mercuronitrie mixture).--

Corrosive sublimate.	5 grams
Nitric acid (approximately 80 per cent)	4 e.e.
Glacial acetic acid.	1 e.e.
Alcohol (70 per cent)	25 e.e.
Distilled water	220 e.e.
Filter after three days.	

Gilson's is an excellent general reagent and gives a very delivate fraction. Objects should be left in the fluid from 15 to 30 minutes for delivate ones to 6 hours for those which are larger or denser, although many tissues may be left for 24 hours without injury. This is one of the most satisfactory killing and fixing reagents that the beginner can use. Danchakoff's Mixture.—See 7 B. Erlicki's Fluid.—See 8.

17. Ether-Alcohol.—Equal parts of sulphuric ether and absolute alcohol.

Flemming's Solution.-See 12.

18. Formalin.—See reagent 6, p. 8, and reagent 4, p. 29. It should be borne in mind that formalm is a reducing agent and will rapidly decompose such reagents as osmic acid or chromic acid if mixed with them. It preserves fat and myelin, so that they may be stained by the standard methods, and various substances, such as anyloid and hemosiderin, to which it may be desirable to apply chemical tests.

Commercial formalin is always slightly acid. This is not objectionable for ordinary fixation. If neutral formalin is required, add magnesium carbonate to the commercial variety, keeping a deposit of the carbonate on the bottom of the formalin container.

19. Formalin, Alcohol, and Acetic Acid (Lavdowsky's mixture).-

Formalin, commercial	10 parts	
Alcohol, 95 per cent	50 parts	
Glacial acetie acid	2 parts	
Distilled water	40 parts	

This mixture is recommended in some cases for the treatment of embryos, especially when the nervous system is to be studied. It penetrates well and preserves faithfully; the alcohol counteracts the swelling effects of the acetic acid and the formalin. Material may remain in it without injury for several days. The fluid should scorer or later be replaced by 70 per cent alcohol. No preliminary washing is necessary.

Formalin-Zenker.-See 7.

20. Formal Schlimate (Worcester's fluid)—a) Make a saturated solution of corresive sublimate in 10 per cent formalin. This reagent is recommended by Baymond Pearl (Journal of Applied Microscopy, VI, 2451) as "extremely satisfactory" for killing and fluing protonon. Washing may be done in water or 4 per cent formalin. The material may be preserved in 4 per cent formalin or carried up the grades of alcohol to 70 per cent alcohol.

b) If to 9 parts of this formol-sublimate mixture 1 part of glacial acetic acid is added, Worester's formol-sublimate-acetic mixture is obtained. Pearl recommends this highly for telesst eggs and for embryological material in general. It will not produce coagulations and cloudiness in the gelatinous envelopes of amphibian eggs, if thoroughly washed out after fixing. Preservation is the same as for (a). Johnson (Journal of Applied Microcopy, VI, 2652) also recommends this reagent very highly for general work except in the case of nervous tissue.

Personally. I have found it advisable not to prepare either of the above mixtures until needed because the formalin, which is a reducing agent, causes much of the mercuric salt to pass over into the insoluble mercurous salt.

Gilson's Mercuro-Nitric Mixture.—See 16. Helly's Fluid.—See 7 A. Hermann's Fluid.—See 27. Kleinenberg's Picro-Sulphuric.—See 26. Lardowsky's Mixture.—See 19. Müller's Fluid.—See 9.

21. Osmic Acid (really the tetroxide of osmium OsO,),-Osmic acid kills quickly and fixes well. It is exceedingly volatile. The chief objections to it, aside from its extremely poisonous nature, are its poor powers of penetration, and the fact that it becomes reduced in the presence of the least amount of dust containing organic particles. The substance must be handled with the greatest care, as even the vapors are dangerous. It is usually put up in small quantities (0.1 to 1 gram) in hermetically sealed glass tubes. In making up solutions, the wrappings are removed from such a tube, and the tube is dropped into a reagent bottle, where it may then be broken by means of a glass rod. Aside from its use in mixtures (see reagents 12 and 27), the vapor or a 0.05 to a 1 per cent aqueous solution is commonly used. A stock solution of 1 per cent is usually kept on hand. It must be kept free from dust. As the most practical way of preventing reduction, Lee recommends that the osmic acid for ordinary work be kept as a solution in chromic acid (a 2 per cent solution of osmic acid in a 1 per cent aqueous solution of chromic

acid). This solution may be employed in making up Flemming's solution or for the purpose of function by means of commun rapor. For rapor function, however, many workers prefer the vapor from the solid crystals.

To fix by means of the vapor, the tissue is pinned to the lower end of a cork which fits tightly into the bottle containing the oscile acid, or it is suspended by a thread. Objects which will adhere to a slide are fixed by simply inverting the slide over the mouth of the bottle. The time required for such fixation varies from thirty seconds or a few minutes for isolated cells to several hours for thicker objects, such as the retina. For fixing in the solution, 24 hours are required ordinarily. Objects are then washed in running water for the same length of time. The yapor. The stains which follow osnie and best are hematoxylin, methyl green (for study in squeous methal), ahm-carmine, piero-earnine, and safranin.

22. Pieric Acid.—A cold saturated aqueous solution (about 1.2 per cent) of pierie acid is commonly used. Small objects are fixed in from a few minutes (minuscia) to 6 hours; objects up to 1 em. in sinc in from 24 to 36 hours. They may be left a much longer time, however, without injury. Large objects may require weeks for proper fixation. After fixing, tissues should be washed in 70 per cent alcohol until the alcohol is no longer colored by the pierie acid. The tissue should not pass, during subsequent treatment (with a few exceptions in case of staining), into an aqueous medium or into an alcohol of less than 70 per cent strength, because such media seem to undo the work of fization.

23. Pictic Alcohol.—Gage recommends a 0.2 per cent solution of pictic acid in 50 per cent alcohol as an excellent fixer and hardener for almost any tissue or organ. Time required, 1 to 3 days. Entite adjects which have been fixed in pictic acid or in pictic alcohol stain readily in board camine or paracarmine.

24. Puro-Acetic.—Saturate a 1 per cent aqueous solution of asetic and with pierie axid. This liquid is whely used as a general reagent, and is to be preferred for most purposes to pierie axid alone. For washing, etc., see remarks under 22.

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Picro-Acetic-Formalin.-See Bouin's Picro-Formol, pp. 9, 29. 25. Picro-Sublimate .--

Rolls:

Pierie acid, saturated aqueous solution	1 vol.
Corrosive sublimate, saturated aqueous solution	1 vol.
Distilled water	2 vols.

This mixture has been especially recommended for embryos. They are left in the fluid for 12 hours, then washed in weak alcohol and transferred to gradually increasing strengths of alcohol.

O. nom Rath's:

Pieric acid, cold saturated solution.	1 ml.
Corrosive sublimate, hot saturated solution	1 vol.
Glacial acetic acid	1 vol.

After fixing for several hours, transfer the material directly into alcohol.

26. Picro-Sulphuric (Kleinenberg's) -

Pierie acid, saturated aqueous solution	98 vols.
Sulphurie acid	2 vols.
Water	200 vols.

This is an excellent reagent for embryos, either for entire mounts or for sectioning. Chick embryos of 24 to 48 hours should remain in the liquid for from 2 to 4 hours; older embryos for from 3 to 6 hours. For washing, 70 per cent alcohol is used. It should be changed (frequently at first) until the color ceases to come out of the embryos. Preserve in about 80 per cent alcohol.

Lillie recommends the addition of glacial acetic acid sufficient to make a 5 per cent solution of acetic acid.

27. Platino-Aceto-Osmic Mixture (Hermann's fluid) --

Platinum chloride, 1 per cent aqueous solution... 60 c.e.

Hermann's fluid is one of the most valuable cytological reagents. Only small pieces of tissue should be used. The washing and subsequent treatment are the same as for Flemming's solution (12).

For subsequent treatment with pyrogallol, see 70. Read, also, remarks on osmic acid (21).

Rabi's Piero-Sublimate.—See 25. Rath's (O. vom) Piero-Sublimate.—See 25. Ripart and Petit, Liquid of.—See 10. Tellyennichy's Fluid.—See 5. Van Gehochten's Fluid.—Same as 2, b). Worcester's Fluid.—See 0. Zenker's Fluid.—See 6.

II. STAINS

Read the general statement about stains in chap. ii.

28. Alum-Cochineal—For formula see p. 9. Alum-cochineal is one of the best stains for entire objects. It is easy to work with, and does not overstain. The time required for staining is from 24 to 36 hours ordinarily. After staining, the object should be washed in water for 15 or 20 minutes to extract the alum, which would otherwise crystallize when the preparation is placed in alcohol. Too long an immersion in water may extract the stain to too great an extent. From water the object should be passed upward through the grades of alcohol, remaining about an hour in each. The writer has found alum-cochineal especially valuable for fattyouns (taperarms, flakes, etc.) and enforces. If it is desired to use a counterstain with it, Lyon's blue, piece

29. Alum-Carmine.—

Boil for 20 minutes, and filter when cool. The uses and manipulation are the same as for reagent 28. These stains affect calcareous structures injuriously.

 Anilia Stains.—Read the general remarks about anilin stains in chap, ii (p. 20). The formulae for some of the most important are given separately in this list in their proper alphabetical position.

The dyes are dissolved in water, in alcohol of any desired strength, or in anilin water, according as they are soluble in these media, or as they meet the needs of the operator. Some workers even use some

of them as counterstains dissolved in the clearing fluid. For the study of nuclei, after Hermann's or Flemming's fluid has been used for firing, the writer has found a weakly alcoholic anilin-water solution to be the most satisfactory. As cytoplasmic contrast stains alcoholic solutions (in 70 to 95 per cent alcohol) have given the best results. Anilin water is made by shaking up 4 e.e. of anilin oil in 90 e.e. of distilled water and filtering the mixture through a wet filter. Enough alcohol may be added to make it a 20 per cent alcohol, if a weakly alcoholic solution is desired.

The length of time which sections should be immersed in the stain varies from a few seconds or minutes for some of the dyes (especially when used for cytoplasm) to 24 to 36 hours for others (especially nuclear). Sections usually overstain, in which case they are differentiated by means of alcohol, either pure or slightly acclutated with hydrochloric acid. The color is thus extracted rapidly; devolution should be stopped immediately after the color ceases to come from the tissue in clouds (20 seconds to 3 minutes). If acidnated alcohol is employed, it must be in much weaker solution than that used for extracting earnines or hematorylins. One part of hydrochlorie acid to 1,000 of water or alcohol is about the correct proportion. When one desires to study the karyokinetie figures of nuclei, the acid-alcohol differentiation should be employed, but if resting nuclei are to be studied, only neutral alcohol should be used.

31. Anilin Blue, Orange G, and Acid Fuchsin (Mallory's triple connective-tissue stain).--

Solution I:

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Acid fuchsin. Distilled water.	
Solution II:	
Anilin blue (Grübler's water soluble)	0.5 gram
Orange G (Grübler)	2.0 grams
Phosphomolybdic acid, 1 per cent aqueous	
solution	100 e.e.

The tissue should have been fixed in Zenker's fluid. Stain either paraffin or celloidin sections in the acid-fuchsin solution for

5 minutes or longer, depending upon the freshness of the tissue. Transfer directly to solution II and stain for from 10 to 20 minutes or longer. Wash and dehydrate in several changes of 95 per cent alcohol. Pass paraffin sections through absolute alcohol, clear in xylol, and mount in balsam. Clear cellokin sections from 95 per cent alcohol in crosset or other cellokin clearer, or blot and clear in xylol, inally mounting in balsam.

Connetive-tissue reticulum, collagen filmis, mucus, anyloid, and various other hyalme substances stain in different shades of blue; muclei, eytoplasu, axis-cylinders, neuroglis fibers; filoreglis fibrits, and fibrin stain red; elastic fibers, pele pink or yellow; and red blood corpuscles and neyelin sheaths, yellow. If the add fuelsin is omitted, muclei and protoplasm stain yellow and the connectivetissue fibrillae and reticulum stand out sharply in deep blue. This is an excellent stain for developing hone, inasmuch as eartilage stains light blue and hone dark blue.

Bensley's Acid Fuchsin-Methyl Green Method.—See p. 144. Bensley's Copper Chrome Hematoxylin Method.—See p. 145.

32. Bismarck Brown.—Boil 1 gram of the stain in 100 e.e. of water, filter, and add 30 e.e. of strong alcohol. Bismarck brown is a nuclear stain which does not overstain, although it acts rapidly. After staining, wash in 95 per cent or absolute alcohol. This stain is also used in aqueous solution for intro-witow staining; the nucleus of the living cell may thus be colored. It has been used as an introvitow stain mostly in the study of infusoria. The stain may be fixed by means of a 0.2 per cent chronic-with solution, but this, of course, distroys the life of the cells.

33. Borax-Carmine (Grenacher's) .---

Borax (4 per cent aqueous solution)	100 e.e.
Carmine.	1 gram

Boil until the earmine dissolves, then add 100 e.e. of 70 per cent alcohol. Filter after 24 hours.

This is a stain much used in the past for staining in balk. Objects must be left in it for from 24 hours to serveral days. They are then transferred, without washing, to acid alcohol and left until the color no longer comes array in clouds. Objects should become bright

searlet in color. Finally they should be washed and hardened in neutral alcohol.

34. Bordeaux Red.-

Bordeaux red	l gram
Distilled water	100 e.e.

This is a good plasma stain. Recommended by Heidenlain as a contrast stain for iron-lematorylin in the demonstration of centrosomes (p. 147). Stain for 12 to 24 hours.

35. Carmalum (Mayer's).—

Carminie acid	l gram
Alun	10 grams
Distilled water	200 e.c.

Dissolve with heat and filter the solution when cold. Add a few crystals of thymol or a little salarylic acid to prevent the formation of mold. Carmalum is one of the best stains for staining objects in bulk and will follow almost any fixing reagent, even osmic acid. If the object has an alkaline reaction it does not stain so well. Washing is done in water.

36. Carmine (Beale's) .--

Powdered carmine	l gram
Ammonia	3 e.e.
Pure glycerin	96 e.e.
Distilled water	96 e.e.
Alcohol, 95 per cent	24 e.e.

The ammonia and part of the water are first mixed and the carmine dissolved in the mixture. The remaining water is added and the solution is left in an open dish until the ammonia has almost evaporated. The alcohol and glyverin are then added. For staining, equal parts of the stain and glyverin are used. The staining is carried on for 24 hours under a hell-jarin an uncovered dish. A second open dish containing acetic acid is placed under the bell-jar. After staining, the sections are washed in water, then in weak hydrochloric acid (1 to 500 of water), and again in water. Minot recommends this stain and method of treatment especially for the placenta and for the central nervous system of endryos.

37. Carmine, Picric Acid, and Indigo Carmine (Calleja's staining fluid).—

Solution I:		
Carmine	2	grams
Lithium carbonate, saturated aqueous solu-		Ū
tion	100	e.e.
Solution II:		
Indigo-carmine	0.25	gram
Pieric acid, saturated accesus solution	100	6.6.

Place sections in solution I for from 5 to 10 minutes, then into acid alcohol until they become pale ref (20 to 30 seconds); wash well in water. Next place the sections in solution II for 5 to 10 minutes, then into acetie acid (0.2 to 0.5 per cent) for a few seconds, and wash well in water. Dehydrate rapidly and clear in xylol. The method is useful for epithelial cells and connective tissue.

38. Carmine, Add (Schneider's).—Add earnine to builing acetic acid of 45 per cent strength until no more will dissolve. Filter the solution when coal. This is a valuable reagent for the study of the nuclei of fresh cells. It is very penetrating and gives a brillant stain. The strong acetic acid ultimately destroys the cell.

30. Congo Red.—For formula, see p. 10. This is a good counterstain for hematoxylin, especially when applied to fetal and young tissues. Its solutions become blue in the presence of free add, hence it is useful in determining the existence of free add in tissues.

40. Cyanin (Chinolin Blue; Quinoline Blue).—Dissolve 1 gram of cyanin (prepared by H. A. Meiz & Co., of New York) in 100 e.e. of 95 per cent alcohol, and add 100 e.e. of distilled water. This is a good cytological stain. Sections are stained for 5 to 10 minutes. Chromosomes stain a deep blue. I have found cyanin followed by erythrosin (0.5 per cent alcoholic solution) especially valuable for spermatozoa.

 Enrich-Bondi Triple Stain (Heidenham) -- The ingredients should be obtained from Grübler and Hollborn, Baiersche Strasse 63, Leipzig, or from their agents.

The solution of orange should be prepared first, and the solutions of fuchsin and methyl green added to it with continual stirring. Each solution must be thoroughly saturated; it takes several days for this to occur. The above-mentioned mixture constitutes a stock solution which should be diluted with about 50 or 100 times its volume of water before using. According to Lee (Microtomist's Vade Mecum), "if a drop be placed on blotting paper it should form a spot bluish green in the center, orange at the periphery. If the orange zone is surrounded by a broader red zone, the mixture contains too much fuchsin." For use with this method, tissues should be fixed in pure corrosive-sublimate solution. Sections should be thin (3 to 5 microns) and must remain in the stain from 12 to 24 hours. They should then be rapidly washed in 95 per cent alcohol, placed for a short time in absolute alcohol, and cleared in xylol. If the sections remain in the alcohols any considerable length of time, the methyl green will be extracted. The stain is very uncertain in its action, but when it is successfully applied the results are excellent. It is used chiefly in evtological studies, especially in connection with gland cells. Grübler prepares a dry powder for this three-color mixture, but the results are usually not as satisfactory as when the mixture is properly made fresh. To prepare the stain from the powder, a 0.4 per cent solution of the latter in distilled water is made, and to 100 c.e. of this solution 7 c.c. of a 0.5 per cent aqueous solution of acid fuchsin is added.

42. Ehrlich's Triple Stain.—For blood films Ehrlich's so-called triacid mixture is a serviceable stain which is widely used.

Orange G, saturated aqueous solution	14 c.c.
Acid fuelisin, seturated aqueous solution	7 e.e.
Distilled water	15 c.c.
Absolute alcohol	25 c.c.
Methyl green, saturated aqueous solution	12 c.c.
Glycerin	10 e.e.

Each solution must be thoroughly saturated (several days). Add the ingredients in the order named, shaking the nixture well before each addition. It is best for the stain to stand several weeks before it is used. Neutrophil granules stain violet, oxyphil granules a brownish red. The mixture stains in from 5 to 15 minutes.

43. Bosin.—See reagent 16, p. 10. This suffin due is often used after hematoxylin as a contrast stain. It is specific for certain granules of leucocytes and for red blood corpuseles, giving to the latter a very characteristic coppery-red tinge. Some workers prefer to dissolve it in water or in some cases in the clearer.

 Erythrosin.—An eosin; properties and manipulation much the same as ordinary eosin (see reagent 43).

45. Fuchsin, Acid (Rubin S, Acid Magenta, Magenta S).-

Acid fuebsin	0.5	gram
Distilled water	100	C.C.

This is an excellent anifin stain for cytoplasmic structures. It is also used in some instances as a specific stain for nerve fissue. Acid Inchsin should not be contounded with host fuckain, which is a nuclear stain. It too is used in aqueous solution. When fucksin alone is mentioned by writers, without specifying whether it is acid or basic, the basic fucksin is ordinarily meant.

46. Fuchsin-Methyl-Green Stain (Ainerhardts).—Keep in separate bottlas 0.1 per cent aqueous solutions of acid fuchsin and methyl green respectively. When ready to use, mix 2 parts of the acidiuchsin solution with 3 or 4 parts of the methyl green, after acidulating every 50 e.c. of the former with 1 drop of a 10 per cent solution of active acid.

This stain works best after a sublimate fiver. Chromosomes stain green, finin and plasmosomes red. Sections should not be over 3 or 4 microns thick. Stain for 15 minutes and transfer directly to 95 per cent alcohol. As soon as the green stain ceases to leave the sections in clouds, pass the sidds rapidly through absolute alcohol and xybol and mount in balsam.

47. Fuchsin (Acid) and Picric Acid (Van Giesen's stain) .--

Acid fuelsin, 1 per cent aqueous solution...... 10 c.e. Pierie acid, saturated aqueous solution....... 90 c.e.

This stain is frequently used in conjunction with hematoxylin in the study of fibrous or of nerve fissue. Small bits of fissue should be fixed in corresive stulin ate or its mixtures. Sections are slightly overstained with hematoxylin, rimsed in water, and then stained

5 minutes in the piece-fuchsin mixture. To avoid extracting toe much of the yellow color in dehydrating and clearing, the alcohols and clearer should each have a few crystals of pieric acid added to them. The result should be: nuclei and epithelia hown; white fibrous connective tissue red; elastic tissue and muscle yellow.

 Gentian Wielet—This is one of the best of the nuclear amin stains. It is best made up in anilin water and weak alcohol (see reagent 30).

Géntian violet	l gram
Anilin water	80 c.c.
Alcohol, 95 per cent	20 c.e.

The stain works well with thin sections. It is also widely used in the study of bacteria. For differentiation, Gram's method is used.

Gram's solution:

Iodine	l gram
Iodide of potassium.	2 grams
Water	300 e.e.

After staining, the sections are placed in this solution until they are black (2 to 3 minutes) and are then decolorized in absolute alcohol until they appear gray. See also reagent 73.

49. Gold Chloride.—The gold-thloride method is used chiefly in the study of nerve-fiber terminations, both motor and sensory, although it is sometimes used for the coloration of other tissue elements (capsules of eartilage, etc.). The process is really an impregration; through the agency of sunlight and of certain rengents (acctin, eitric, formic, or oxalic acid) the gold is deposited in the tissues in the form of very fine particles. There are numerous modifications of the method, one of which is given in chap ix.

50. Golgi's Chrome-Silver Method .- See chap. ix, p. 71.

Hematoryin.—For general statement see chap. ii, p. 20, and the remarks under 12, p. 9.

51. Hematoxylin, Conklin's Picro .--

Add one drop of Kleineaberg's picro-subhuris (20) to each cubic centimeter of the solution. This is a good stain (1 to 3 hours) for embryos which are to be mounted entire. If the embryos are to be sectioned they should be stained for 12 hours.

52. Hematoxylin, Delafield's.-See reagent 12, p. 9.

53. Hematoxylin, Ehrlich's Acid.—

Hematoxylin	2 grams
Absolute alcohol	100 e.c.
Glacial acetic acid	10 e.e.
Glycerin	100 c.c.
Distilled water	100 c.c.
Potassium alum	10 grams

Dissolve the hematoxylin in the actic acid with 25 c.c. of the alcohol; then add the glycerin and the remaining alcohol. Dissolve the alum in the water by the aid of heat and slowly pour the warm solution while stirring into the solution of hematoxylin. The solution must be exposed to light and air at least 3 weeks to ripen. It is not ready for use until it acquires a deep red color. This solution is an excellent nuclear stain and will keep for years.

Mone's acid hemotein is the same as this, except that hematein (Grühler's) is substituted for the hematorylin. This solution should stain without having to ripen.

54. Hematoryin, Heidenhair's Iour.--See reagent 17, p. 11, for formula; p. 51 for method; chap. xvii for cytological uses. This stain is much used in the study of cell structures such as centrosomes, chromosomes, etc. Tissues are best fixed in Bonin's, in some of the sublimate solutions, or in acetic about, athrough it will follow liquid of Flemming or Hermann. Sections should not be over 6 microns thick. The ferric solution must be renewed oceasionally as it soon spoils.

55. Hematoxylin, Mallory's Phosphotungstic Acid .--

Hematein ammonium	0.1	gram	
Water	100	66	
Phosphotungstic acid crystals (Merek)	2	grams	

Dissolve the hematein by heating it in a little water. When cool add it to the rest of the mixture. If the stain is too weak at

first, it may be ripened by adding 5 c.c. of a 0.25 per cent aqueous solution of permanganate of potassium or by allowing it to stand two or three weeks. Hematoxylin may be used instead of hematein ammonium if 10 c.c. of the potassiun-permanganate solution is added to ripen it.

The stain is recommended for the demonstration of neuroglia, myoglia, and fibroglia fibrils, for fibrin, and for centrosomes and spindles of mitotic figures.

Janus Green.-See pp. 145, 146.

56. Light Green (Lichtgrün S.F.).-This is a beautiful cytoplasmie anilin stain which is frequently used after safraain as a counterstain. Not more than 0.5 per cent solution should be used as it stains very rapidly and very deeply. It may be used either as an aqueous or as an alcoholic solution. The writer has found a 0.5 per cent solution in 95 per cent alcohol very satisfactory. Sections should remain in it only a few seconds. Do not confuse it with methyl green, which is sometimes called light green by dealers.

57. Logol's Solution.—A solution of iodine in water containing iodide of potassium. It is used in various strengths. One of the commonest formulae is that of Gram (p. 223), although some prefer a solution with only one-third the amount of water used by him.

58. Lpons Blue (Bleu de Lyon).—This is one of the best of the numerous anifm blues. It is a good contrast stain when used after such nuclear stains as safrarin and earnine. See reagent 15, p. 10; also p. 50.

Magenta, Acid.-See 45.

Mallory's Triple Connective-Tissue Stain .- See 31.

59. Methylen Blae.—This reagent is an extremely useful one; it is of great value in the study of the nervous system, and it can be made to give results with intercellular cement substance, lymph spaces, etc., as satisfactory and with greater certainty than impregnations obtained with gold chloride or silver nitrate. It is also serviceable as an introvitom stain. Furthermore, methylen blue (saturated solution in 70 per cent alcohol) followed by each is sometimes used for the double staining of blood corpuseles. Methylen blue should not be confounded with methyl blue.

Ordinary commercial methylen blue usually contains, in addition to the blue dye, a small quantity of a reddish-riolet dye. Such methylen blue is termed polychromotic and is especially serviceable in staining certain cell granules. Only the pure methylen blue, however, should be used for nerve staining and other intra-itam work.

a) "Intra-Vitam" Skin for Snall, Comparatively Transparent Aquatic Organisms.—Add sufficient methylen blue to the water containing the organisms to tinge it a light blue. Different tissues will take up the color after different intervals of time. A given tissue after having attained a maximum degree of coloration will rapidly lose its color again. It is necessary, therefore, to watch the organisms closely for the maximum of color in the tissue desired. If the observer wishes, the stain may be fixed for more prolonged study by following the processes indicated under (i). The order in which various tissues take the stain seems to vary in different organisms. Usually gland cells stain first, then with more or less derivation, other epithetial cells, fat cells, blood and lymph cells, elastic fibres, smooth muscle, and striated muscle. Nerve cells and nerve fibres do not ordinarily take the stain when the ertine animal is immersed.

b) Ehrlich's Method for Nerve-Terminations and the Relations of Nerve Cells and Fibers to the Central Nervous System .-- The stain should be Grübler's methylen blue (rectificiert nach Ekrlick). A 1 per cent solution in normal saline is used. Warm the solution till it steams, stir it thoroughly, and, when cool, filter. The tissue must be perfectly fresh. Chloroform the animal and immediately inject the stain into the main artery of the part to be investigated. If the animal is small, the entire body may be injected. The vessels should be filled full, but care must be taken not to rupture them. The part should become decidedly blue in color. It is well after 10 or 15 minutes to inject more stain. At the expiration of half an hour after the second injection remove small pieces of tissue containing the nerve elements desired, and expose them freely to the air on a slide wet with normal saline. Examine every two minutes under the microscope (without cover-glass) until the particular element to be investigated (cell, axone, termination) has developed a well-marked

blue color. It is important to eatch the color at the proper stage and fix it because it soon begins to fade.

Firing the Stain.--When the desired element has developed a satisfactory blue color, the tissue is transferred immediately to a saturated aqueous solution of Annowium picrote (Dogiel's method) and left for from 6 to 24 hours. For final mounting the tissue should be teased out sufficiently to show the proper elements and then mounted in a few drops of a mixture of pure glycein (free from acid) and annowium picrote (saturated aqueous solution), equal parts. It is well to let the tissue stand in 20 to 30 rohmes of this glycerinpicrote mixture for a day or two before mounting it. If the preparation is to be hept the cover glass should be scaled (n. 95).

Sections—If it is desired to make paraffin sections and mount them in bakam, after treatment with the annomium picrate (10 to 15 minutes), the tissue must be placed into 20 or 30 volumes of Bethe's fluid, which renders the color insoluble in alcohol.

Bethe's fluid:

Molybdate of ammonia	l gram
Chromic acid, 2 per cent aqueous solution	
Hydrochlorie acid, concentrated C.P	1 drop
Distilled water	

The tissue is left in this mixture for from 45 to 60 minutes (for small objects) and then washed 1 to 2 hours in distilled water. Dehydrate directly in alsolute alcohol; follow this with xylol, inded in parafilm, and section in the ordinary manner. Sections may be counterstained in alum-carmine or alum-ordineal.

c) Immersion Method—Material which earnot be readily injected or which has failed to stain may be stained by immersion. A 0.1 per cent solution of the stain is used (dilute 1 volume of the solution used for injection with 9 volumes of normal saline). To small pieces (2 to 3 mm, thick) of the tassue add a few drops of the stain at intervals of about three minutes. The tissue should always be moist, but never covered sufficiently by the solution to exclude air. Examine the preparation from time to time under the microscope and when the nerve elements are well stained, fix in animoxium pierate and proceed as in (b). In case of the central nervous system, fairly

good results may sometimes be obtained by dusting the methylen-blue powder over the freshly cut surface of the part to be studied. The development and fixing of the color is the same as in (b).

d) Nissl's Method of Staining Basophil (Tigroid) Substance in Nerve Cells.—

Nethylen blue	3.75 grams	
Venetian scap (white castile scap)	1.75 grams	
Water	1,000 e.c.	

It is best to keep the stain for some months before using.

Ganglia should befined in alcohol, formalin, or cornsire sublimate and sectioned in parafin. Fix the sections to the side, dissolve out the parafin with xylol, and run the preparation down to the squeous stain in the ordinary way. In a test-tube heat a few eulaic centimeters of the stain until it steams, then apply it while still warm to the sections on the slike, which has been placed flat on the desk. It takes about 6 minutes for the stain to act. Pour off the surphs stain and rinse the slike in distilled water. Lay it flat on the desk again and flood the sections with endlowedowd (55 per cent alcohol, 9 parts; amilin oil, 1 part). Let the sections devolutine (20 to 30 seconds) until they are a pale blue; then drain off the anilm-alcohol and transfer the preparation to also balo the alcohol. Clear in xylol and mount in blasm. The basephile randows should appear deep blue in color. They are arranged for the most part concentrically around the nucleus.

e) Unna's Method of Staining Unstriated Muscle in Sections.— Stain in a 1 per cent equeous solution of polychomatic methylen blue, risse in water, and then leave for 10 minutes in a 1 per cent aqueous solution of polassium ferricyanide. Transfer to acid alcohol until sufficiently decolorized, then complete the dehydration, and mount in the usual way.

f) For Ordinary Section Staining where a nuclear stain is desired methylen blue answers very well. It is usually used (2 to 24 hours) in aqueous solution. The treatment is the same as for safranin.

g) Impregnation of Epithelia, etc.—Place the fresh tissue, preferably a thin memorane, into a 4 per cent solution of methylen blue in normal saline. To demonstrate the outline of cells, leave the

tissue in the stain not longer than 10 minutes. To get a negative image of lymph spaces, canals, etc., in contrast to the ground substance which becomes deeply impregnated, leave the tissue in the stain 20 to 30 minutes. For this purpose it is advisable to remove any membranous covering which invests the organ. In either case, after staining, far the tissue for 30 to 40 minutes in a saturated aqueous solution of ammonium picrate, changing it once or twice, and examine in dilute glycerin. To preserve the preparation permanently, proceed as in (b). To do away with the macerating action of the ammonium picrate, add 2 per cent of a 1 per cent osmic-acid solution to the fixing bath.

60. Methyl Green.—This is one of the best of the unckar anilin stains. It is particularly valuable because it often instantly stains the chromatin of nuckei in fresh tissues. Use in strong aqueous solutions, adultated to about 1 per cent with accile acid. It does not give a satisfactory chromatin stain if the tissue has been fixed in acetle acid or mixtures containing it. It follows pure corresivesublimate solution admirably.

61. Methyl Violet.—This stain is commonly used in 0.5 to 2 per cent aqueous solutions for staining harteria, nuclei, and anyloid. It may often be substituted for gentian violet.

62. Muci-Carmine (Mayer) .--

Carmine	1.() gram
Aluminium ehloride	0.8	5 gram
Distilled water		
Aleohol, 50 per cent.	100	6.6.

Mix in the order given; heat gently till the fluid darkens (about 2 minutes); filter after 24 hours. To use, dilute with 5 to 10 volumes of water. The stain (3 to 10 minutes) is specific for mucus-containing cells.

63. Muci-Hematein (Mayer).-

Hematein	0.2 gram
Glycerin.	40 e.e.
Aluminium ehloride.	0.1 gram
Distilled water	60 c.c.

Rub up the hematein in a mostar with the glyeerin and the almminime chloride, then add the water. It stains in from 3 to 10 minutes, much appearing blue. If a drop or two of nitrie add is added, its nuclear staining capacity is enhanced.

64. Neutral Red.—Neutral red is used widely as an intro-ritom stain. It is a good stain for cytoplasmic granules, and in some cases for moons-cells. For intro-riton staining it may be used in the same way as methylen blue (with the omission of fixation). For staining fixed material, a 1 per cent or stronger aqueous solution is employed. Granules are stained orange red (hright red in acid medium, yellow in alkaline medium). Rosin finds that in nerve cells stained in neutral red (followed by water, acid-free alcohols, xylol, and balsam) modeoli and Niss's granules are stained red, the rest of the cell yellow.

63. Orange G.—This is an excellent cytoplasmic stain and is often used on sections as a contrast to earmine, hematoxylin, and satramin. It is especially good as a contrestain in tissues of vertebrate embryos. Grübler's orange G is the most reliable. It should be used in saturated aqueous solution. The solution does not keep very well.

66. Orcein (Unna's method for elastic fibers) .-

Oreein (Grübler's)	l gram
Hydrochlorie acid	lea
Absolute alcohol	100 e.c.

Sections are stained in a watch-glass or porcelain disk. The dish is warmed over a flame or in an oven until the stain becomes thick through the evaporation of the alcohol. Rinse the stained sections throughly in 70 per cent alcohol, wash in water, run up through the alcohols, clear in xybl, and mount in balsam. Elastie fibers should appear dark brown, connective fissue a pale brown. Nuclei may be brought out by staining in Unna's polyehrome methylen-blue solution after washing the sections in water.

67. Paracarmine (Mayer's).-

Carminie acid	1.0 gram
Aluminium chloride	0.5 gram
Calcium chloride	4.0 grams
Alcohol, 70 per cent	100 e.e.

Paracarmine is an excellent stain for large objects. It does not overstain ordinarily. The stained tissue is washed in 70 per cent alcohol. In case overstaining occurs add 2.5 per cent glucial acctic acid or 0.5 per cent aluminium chloride to the alcohol used for washing. Objects to be stained should not have an alkaline reaction nor contain limy materials.

68. Pieric Acid—Pierie acid is widely used as a contrast stain with earnine, hematoxylin, etc. It is best manipulated as a stain by adding a little to each of the alcohols used in dehydrating, after application of the nuclear stain. However, if acid alcohol is to be used, the pierie acid should be used only in the grades above the acid alcohol. It may be employed in staining entire objects as well as sections. See also remarks on washing under 22. 69. Piero-Carmine.—

Ammonium hydrate	õce.
Distilled water	50 e.e.
Carmine	l gram
When dissolved add pierie acid (saturated	
aqueous solution)	50 c.c.

Expose to air and light for two days, then filter. A few crystals of pieric and should be added to the alcohols used for dehydration after staming.

Picro-Fuchsin.-See 47.

Pyridine-Silver Method (Ranson's Cajal) .- See p. 75.

70. Pyrogalol.—Tissues which have been faced in Hermann's or in Flemming's fluid for 24 to 36 hours may be treated (without periodes washing) with a weak solution of pyrogalol or with erade pyroligneous acid. Lee (*Microtomial's Vale-Meaun*) recommends the pyrogalol as much preferable. Tissues should remain in the fluid from 1 to 24 hours, depending upon size. The result is a black stain which colors both nucleus and cytoplasm. If desired, an additional chrometin stain may be employed. Sufranin (72) for 24 hours is recommended; decolorine slightly with very dilute acid alcohol. The stain is excellent for cytological work (for "sphere," etc.).

71. Resorcin-Fuchsin (Weigert's elastic tissue stain).-

Basic fuchsin, 1 per cent aqueous solution...... 100 e.c. Resorcin, 2 per cent aqueous solution....... 100 e.c.

Heat the mixture in a porcelain dish and while builting add 25 e.e. of hypor ferri asequichlorati; stir and keep builting for 2 to 5 minutes. Cool and filter. Throw away the liquid. Dry the precipitate which remains on the filter-paper throughly in a porcelain dish over a water or sand bath. Return the dried precipitate together with the filter-paper to the first porcelain dish, add 200 e.c. of 55 per cent alcohol, and boil. Remove the paper when the precipitate is dissolved off. Cool, filter, and replace the alcohol lost through eraporation, up to 200 e.e. Add 4 e.e. of hydrochloric acid. The stain works best after formalin-fused material.

Stain the section 20 minutes to an hour in this solution. Wash in alcohol, dehydrate in absolute alcohol, clear, and mount in halsam. Elastic fibers should appear dark blue on a clear background. If desired, the sections may also be stained, either before or after the staining of the elastic fibers, with one of the earnine or hematorylin stains.

 Safranin.—Safranin is one of the most important of the basic anilin dyes. Read earefully the remarks on anilin stains under 30.

Safranin	l gram
Anilin water (see 30).	90 c.c.
Alcohol, 95 per cent	10 e.e.

Filter before using. Grübler's "Safnanin O" is the most reliable dye. Sections of tissue fixed in Hermann's or Flemming's solution are left in the stains for from 24 to 48 hours. Decolorine as directed under 30.

73. Seframin and Gentian Violet.—This is a combination that is almost indispensable in the study of cell problems, especially spermatogenesis. For formulae of stains see 48 and 72. Tissues are best fined in Flemming's or Hermann's solutions. Stain thin sections for 36 to 48 hours in the safranin; differentiate in alcohol very slightly addulated (see 30), then stain for 5 to 10 minutes in the gentian solution and transfer the sections to Grant's solution (see under 45) for

1 to 3 hours. Finally differentiate in absolute alcohol. As soon as purple clouds have eeased to come from the sections in absolute alcohol, they should be transferred to clove oil for a few minutes and thence to xylol. The clove oil seems to intensity the suframin in the chromatic granules, but too prolonged an immersion in clove oil extracts the gentian violet.

74. Scharlach R—A saturated solution of the dye in equal parts of 70 per cent alcohol and acctone is used. This is a specific stain for fat. For example, cover-slip preparations are faced in formalin vapor for 5 or 10 minutes, stained 5 minutes in the Scharlach R solution, rinsed in 70 per cent alcohol, washed in water, counterstained with alum-hematoxylin or methylen blac, washed in water, and mounted in giverin-jelly. Frozen sections of formalin-fixed material may be treated in much the same way.

With any evaporation of the alcohol a precipitate forms, hence staining should be done in a tightly closed vessel. After staining with alum-hematoxylin, if the sections are put into a 1 per cent aqueous solution of acetic acid for 3 minutes, the contrast is sharper.

75. Silver Nitrate.—The nitrate-of-silver method is used largely as an impregnation method for work on nerve tissue and for denonstrating intercellular substances and outlining boundaries of cells in the epithelial coverings of membranes, etc. Wash the fresh tissue in distilled water, then place it for 2 to 5 minutes in 0.5 to 1 per cent aqueous solution of silver nitrate. Rinse in distilled water, then expose the tissue to height smlight in water or giverin (or in 70 per cent alcohol, if it be mounted in balsam) until a brown exlocation appears. Temporary mounts should be made in giverin. For applications see pp. 71–74.

76. Sudan III.—This is a specific stain for fat. See remarks on p. 147. A saturated alcoholic solution is used (5 to 10 minutes). Wash rapidly in alcohol. Since alcohol is a solvrant of fat, too long an immersion will destroy the preparation. Mount in glycerin or glycerin-jelly. The tissue should have been fixed previously in Miller's third (9) or other medium which does not dissolve fat.

 Thionin.—This is an excellent stain for chromosomes when used in saturated aqueous solution for about 5 minutes. After corresive-sublimate fixation it is, when used dilute for 10 to 15 minutes, a specific stain for maxin (muxin red, everything else blue).

Van Giesen's Stain.-See 47.

78. Weigert-Pal Stain for Medullated Nerve Fibers .-

Solution	111
1233 8849	11.

Hematoxylin	1	gram
Absolute alcohol.	10	6.6.
Distilled water	99	0.0.
Lithium carbonate (1 part of a saturated aqueous solution to 80 of distilled water)	1	6.6.
Solution II:		
Potassium permanganate Distilled water.		gram e.e.
	100	40
Solution III; mix just prior to using: Oxalic acid, 1 per cent aqueous solution Potassium sulphate, 1 per cent aqueous	50	88
solution.	9 ·	C.C.

Tissue should have been fixed in Müller's fluid or in 10 per cent formalin, hardened in alcohol, and sectioned. Stain sections until black (6 to 24 hours) in solution I. Wash well in water to which a few drops of lithium carbonate have been added. Transfer to solution II and lave until the gray matter of the nervous tissue becomes brown (‡ to 2 minutes). Rinse in water and decolorise in solution III until the gray matter of the tissue becomes a light brown and the white natiter a steel blue (‡ to 1 minute). Each section must be earchilly watched to get a satisfactory result. Wash in running water or in several charges of water, dehydrate, clear, and mount in halsam. If desired, a counterstain of ahumcochineal may be given before the final dehydration.

79. Wright's Stain (for blood and for the malarial parasite).— See memoranda 5 and 6, pp. 110-111.

III. NORMAL OR INDIFFERENT FLUIDS

(For Fresh Tissues)

80. Aqueous Humor.—Obtained by puncturing the comea of a freshly excised beeff seye. A small amount may readily be obtained by means of a capillary inpetter from the eye of a freshly killed frog.

Amniotic fluid from pig or cow fetuses is a serviceable fluid for the examination of fresh tissues.

81. Blood Serum.—Blood is allowed to clot and after 24 hours the serum is poured off. If necessary it may be further freed of blood cells by means of a centrifuge. The serum will keep for only a day or two. Schultz's indized serum make by saturating blood serum with holine is sometimes classed as an indifferent fluid, but it is really a dissociating fluid.

82. Locke's Solution.-See p. 137.

83. Normal Saline.--

Sodium chloride.	0.7 to 0.9 gram
Distilled water	100 c.c.

84. Ringer's Solution.-

Sodium ehloride	0.80	part
Caleium chloride (anhydrous)	0.02	part
Potassium chloride	, 0.02	part
Sodium bicarbonate	0.02	part
Distilled water	100	parts
Dextrose (may be left out)	0.10	part

The following formula is probably better adjusted to tissues of warm-blooded animals:

Sodium chloride	0.900 part
Calcium chloride (anhydrous)	0.024 part
Potassium ehloride	0.042 part
Potassium bicarbonate	0.020 part
Distilled water	100 parts

Ringer's solution corresponds more nearly to normal blood serum than does normal saline and is therefore less likely to produce distortions in tissue elements.

IV. DISSOCIATING FLUIDS

 Bichromate of Potassium—A 0.2 per cent aqueous solution is commonly used. Nerve cells of the spinal cord and also various epithelia dissociate well in it (2 to 3 days).

 Caustic Potash.—A solution of 35 parts in 100 parts of water is often used for isolating fibers of smooth muselé or heart fibers.

It acts by rapidly destroying the connective tissue (30 to 30 minutes). Examination of the tissue is made by moniting it in the dissociating fluid. If water is added, the tissue will be destroyed. Usually only temperary preparations are made in this fluid, but tissues may be made permanent by neutralizing the alkali by means of acetic acid.

Digestion Method.-See pp. 79, 268.

87. Gage's Formaldehyde Dissociator.-

Formalia		
Normal saline solution.	250	C.C.

Good for epithelia and for nerve cells.

- 88. Hertwig's Macerating Fluid.-See p. 78.
- 89. MacCallum's Macerating Fluid.-

Nitrie acid	1 part
Glyverin	2 parts
Water	

This finit is recommended for heart nussele of adults or embryos. Hearts should remain in it from 8 hours to 3 days, according to size. The method is radiable for showing the arrangement of cardiae muscle fibers.

90. Rarvier's One-Third Alcohol.—This is one of the commonest as well as one of the best macersting fluids. It is simply a 30 per cent alcohol. Equithelia will macerstic in it sufficiently in 24 hours. A still weaker alcohol (20 to 25 per cent) is used for isolating the nerve fibers of the retina.

91. Sodium Caloride.— A 10 per cent solution of sodium chloride is excellent for tendon, etc. It dissolves the cement substance of epithelial cells and of connective tissue. As a stain, a saturated aqueous solution of pierie acid (stain for 24 to 36 hours) followed, after thorough washing in water, by a dilute alcoholie solution of acid fuelsin gives excellent results.

V. DECALCIFYING FLUIDS

Tissues are fixed in Zanker's or other fluid, thoroughly washed, and hardened for at least 24 hours in alcohol before decalification. 92. Chromic Acid.--Chromie acid diluted to 1 per cent or in comhimstion with other fluids is frequently used for decalification.

Some Standard Reagents and Their Uses 239

Chronic acid, I gran; water, 200 c.c.; nitric acid, 2 c.c., is a mixture widely used. It decaleties well but acts more slowly than the 10 per cent mixin-acid mixture. Bone should first be hardened in Mülke's find (9).

93. Nitric Acid—A 10 per cent solution of micric acid in 70 per cent alcohol may be used. If mitric acid is used for young or letal bones, it is advisable to use only 1 part of the acid to 99 parts of the alcohol. After washing out in 70 per cent alcohol, the decalefied bone may be kept in 95 per cent alcohol.

94. Phioreglucin Method.—This is a rapid method. Young bones may be decalcified in half an hour and old and hard ones in a few hours. Testh require a somewhat longer time. Phioroglucin itself does not decalcify, but protects the tissue from the action of the strong nitrie acid. One gram of phioroghecin is dissolved in 100 e.e. of pure non-luming nitrie acid with the aid of gentle heat. Ten c.e. of nitrie acid in 100 c.e. of water is added to the mixture. Wash thoroughly and stain in Delaheld's houstorryin. After staining leave sections in tap water for 12 hours.

93. Pinte Acid.—A solution kept fully saturated is useful for delivate bones. It stains and decalefies the tissue at the same time. Wash in 70 per cent alcohol.

96. Von Ebner's Fluid.—

Alcohol, 95 per cent	500 e.e.
Water	
Sodium chloride	2.5 grams
Hydrochloric acid	5.5 c.c.

This is an excellent fluid for bone because in it the ground substance of the bone does not swell up. Sections are best examined in a 10 per cent solution of sodium chloride.

 TABLE OF TISSUES AND ORGANS WITH METHODS OF PREPARATION

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General Remarks		Wright's stain, chup. xiv. See also II, p. 106 Houndorythe coops (2): The propagation should be washed with water or 433. or Physical (2): Macobol before staining.	See I. e. p. 105 Technique same as for cover-glass preparation	See I, d, p. 105	blood		24 hours as all represent the second representation of the second repre	Eat. ad. aol. of methylen Rinee in water, bloc, dry, and mount in balaam. Baseo, gran., blue
Staining, etc.		Wright's stain, chup. ziv. See also II, p. 106 minu.to'x/inte costa (52. The preparation of 43) or Ehrbleh (42)				Ehrlich's hematoxy lin (53), To every 100 c.6. add 0.5 g. com, The 2 to	24 hours Equal parts, sat, gives in sol, of indulin, mapbelly, stain y cillow, and some	Bat, ad sol, of methylen
Person or head a- teron Method - Person - Collordin H - Free Hand								
Plaing and Hardening. or Other Proliminy. Treatment		Dry Corrosive sublimate (14) for 15 minutes			Examine fresh	lither-alcohol (17) 1 to 2 hours	De	Ď
Animal Organ, or Tisaue Trising and Endamine. Resemptation of Or Or Presence of		Proc.			Frog; pigeon; man	Leucocytes of man	Leucocytes of guines-pig. rabbit, nr pigeon, Not in man	Mononuclear leucocytes of
Object or Blement	I. PUSANING DRCANS	Hicod. cover-glass props. From	Crystals Erythytees (red cor- pueden)	Bibeim	Fresh blood Frog; pigeon; man	Granulen, seid ophil (cosmophil, oxyphil)	Granuissi, amphophil Isucceptes of guines-pig. Gadulinophil, ar physics. Not in man	Granules, basephil Mononuclear leucocytes of

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Animal Micrology

Granules, mast cells .	Erradi numbers in normal Erradi and normal bloud. Large numbers in feukeanio blood	For Do. for blood For bismon was strong	P. Thin sections. Mucous mem- branoof mouth intestine. etc.)	Blain (for 24 hours in alum- carruine dabila (dabila) I. R. Aba alu, 25 cool burst gives alu, 25 cool burst gives alum (10 con 25 cool, of alum-entruine)	Differentiate in abs. Als. for 24 hours and finally mount in balasm. "Nuclei, red; mass granules dark blue
Granules, neutrophil.	Polymolear leucocytes of man, sume transitional cells, pus cells, and my- clocytes	Ether-alcohol (17) 1 to 2 hours		Iduction's triple (42)	Rines in water, blot, dry, and mount in acido. scanties, brownish red framines, violeti balaam.
day and a subsection of the su	Mesenteric glands of kit-	Acetic alcohol (2a)	Ú Å	Hematoxylin, cosin (52. 43)	Technictue anno as for cover-class proparation. Longer distant sectors busined business for a sector business for the liness for the business of the part of the p
Marrow, red.	Coversions of sumea-pis- from rit of sumea-pis- fact, or young rabbit Serapp the out surface of a	Ether-alcohol (17) 1 to 2 hours		Ehrlich-Biondi (41) 24 hours, ar hematokylin, erosin (82, 43)	After 41 these ducidy in strong alcohol, clear in clove off collowed by sylol, and mount in balaam See 1, c, other, sit, andms (st)
Spleen, section	Cat	Gilaon (16) or Zenker P. or C. (6)	P. or C.	Hematoxylin, seein (52, 13) or, beinger, bematoxy- lin, and Van Ciench	Fix the spieen entire and later out nut segments of the proper size for sectioning
Thymus SYS-	New-born infant	Acetic alcohol (2a)	P. or C.	-Henrickylin, cosin (53,	
Artery (medium size)	Man; dus; cut Do	Absolute alcohol (3) Absolute alcohol (3) Absolute alcohol (3) or Zonkar (0)	ΰΰ	Hematoxylin, acid fuchain (52, 45) Do	Rematesylin, acid fuchain Make both transverse and Inagtudinal sections (52, 42) Do
cessels Elastic elements	Pleees of bia mater from base of brain Aorta, artery, etc.	Zenker (6) for 1 hour Teased prepara- Carmalum (35) Absolute alcohol (3) C.	Tonsed propara- tion C.	Carmalum (35) Orgein (66) ar Resorciu-	Mount in giveerin or debudgete and mount in for discribution of explicates see in- jection method, chap. xii
Vessels of blood	Vessels from the measurery			Fuchain (71) Silver pitrate (75) 1 p.c. solution injected into the blood vessels	Capillatics in the examined entire; larger vea- sels should be slit open and spread out

Chlote or Element Arbair Organ, or Termen O'Firing and Homomention of Termen O'Termen Heart Dog oth man Arbair Heart I In the fit recould heart or Zonder Ch	Anitotal CORRANACT Thurse Pristing and Anticidual Sciences and the second second second second based on Sciences (1) (1) the fat measured based on Zenderse (0) (1)	Fisting and Hardening, or Oper Presentation Presentation (3) Absolute alcohol (3) Zenker (6)	Election ar Isola- tion Methoda P Dentifin D Free linad H Free linad C.	Staining, etc. Brankerzyin, acid fuchato Dibe, 230000 Bites (33, Methylen hine (50), soeih	Fatalular, etc. Goneral Remarks Hermiters/International Conternal Remarks Description of the Remarks Performance Internation Proceedings of the Remarks Proceeding International International International International Proceedings of the Remarks Proceedings of
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		Fix as above and de- caloity in von Eb- caloity in von Eb- nitrie acid (90) or	C, or F.	Piero-carmine (09)	liee also shap. xi
endochondral)	Extremities of fetal pige, cuts, or burners fetures	Zenker (0)	ť	Mallory's stain (31)	In advanced fetumes the boxe should be decalci- fied (see 93)
none, development of	Parietal, frontal, or lower	Do	ő	De De	
Bone, fibers of Sharpey	Vault of a fowl's alcult	Decalcify in yon Eb- H. Thick see-	H. Thick sec-		Tesse lamellae aparti fine tapering fibers and aperioures from which he visible fibers have been
Hone, Haversian canals					See method for grinding bone, chap, si
puectes sectation of cor-	Thin fragments of bone	Strong nitrio acid for			Cover on a slide and drum upon the cover-glass with the handle of a dissecting needle
		Fix in a misture of Multiple (3) b party (3) 1 party 1 your strand and	Tensed prepara-		Reamine in 10 p.e. aq. sol. of sodium objection
Cartilage (in general)		Zeologi (6) or corrol H. or C.	Н. от С.	Hematoxylin, piorie acid	Hematoxylin, piorie acid Thin sections of costal cartilage may be cut and (52, 68) (52, 68) reactions of readily without any provide propa- ration. Finer details of arcuceure vanish in
Cardiage, capcule of	Small pieces	P'rout.		Treat with I P.c. ad, sol.	zylol-balaam mounts. Try suparal.
diseue and elustic fibers		Freshimate (14)	÷.	Piero-carmine (69)	Examine in giveerin. Elastic fibers, yellow: connective-tesus fibers, pink
fortuage, elastic (yellow	Epiglottis: vooal process of hrytenovi externations	Zenker (6) or sorro- sive sublimite (14)	1, e. 0.	Hematexylin, cosin (52, 43)	Thin sections of fresh cartilage should be exam- ined in normal suline (33)
Cartilage, glycogen in	car (maau, rabbit)	Fresh	H.	Lugol's inde-jodide of po-	Glycogen. M present, is stained a mahogany brown: clastic fibers are stained a different
Cartilage hyaline.	Head of femur: costal car- tilage; ensiferm process sive sublimate (14)	Zenker (6) or corror sive sublimate (14)	H. or C.	Hematogylin, pierio acid	This sections of the fresh cartings also should be examined in pormal saline (83)
Cartilage, white fibro- Intervertebral disks	Intervertebral disks	De	H. or C.	Do	â

PRICIARATION Continued
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METHODS
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TABLE

Object or Element	Animul. Organi, of Themes Fisting and Therdening. Experiments for of Theorem and Theorem and	Fixing and Hardening. or Other Treindinary	Bection or Isola- tion Method F Colloutin H Free Hand	Staining, etc.	General Itemarks
Cells of fibrillar connec-	Internuscular connective		Spread out the	Spread out the Piero carmine (09)	
Gonnie e e e e e e e e e e e e e e e e e e	Bections of any organ Internuscular or subcuta- meous connective tissue	Zenker (0) or corro- nive aublimate (14) Irrigate with acetic Tease acid	Ŭ Å	Mallory (31), or Calleja (37)	
Elastic fibers (coarse) Encircling fibers	Ligamentum nuchae of a See Preparation for fine elastic fibers	P'reab.	Tenso	Picro-carmine (69)	Transforms on and (1 1 ref shound in givenin The constrictions sature (7.4): mount in givenin The constrictions seems at intervals actions the
Pat cella	Fatty areas of mesentery Muller (9) or forma- H.	Muller (9) nr forma-	н.	Scharlach R (74)	Do not subject to any treatment with alcohol occupt as directed under the their occup.
ted membrane	Tenestrated membrane Basilar artery, cut open Strong causiic potash Scrape a way Acid fuchain (48) tong theory andoar acidan and area causa duan and area causa a solution. 6 hours	Strong caustic potesh solution, 6 hours	Scrape away	Acid fuchain (45)	Wish well in water before stating p. 147
Tibrillar (white fibroun)	_	Prest	T'ease		Examine in normal saline (74). Bee also "Ten- don" (p. 245)
Jelly of Wharton	Unbulled bord by young (2 to 3 months) burnen embryo, nr nf h 3 to	Zenker (6) or Lav- H. or P. dowsky (19)	H. or P.	Hematoxylin, cosin (52, 43)	
Ligament	Ligamentum teres	Zenker (0)	H. or C.	Hematorylin, sold fuchain	Hemitorylin, sold fuebain Crees and longitudinal sections
Mucold connective tis-	Boof	Zenker (6)	Do	(02. 40) Do	Do
Retoutar (adonoid) con-		Freak	F. or H. Thin		Bee p. 79

Synovial villi	Proving conjunction intermeters have bordes of patella. Tendon from tail of rat in Tendor-Ashilla of from	Fresh Dry Fresh Zentior (0) or Gilson	H. Henne	Methylen blue (39 f.) Flore-esrmine Sokin in 7015 in bores geottunine, in plore sold	With the provide and with with the functional strains (12). Say also "Prioritian connective formation in givenin formula in environ	. 41 t
IV. DYOESTIYE ORGANS Distriction of Panels of Cells of Panels	Gat or dog an or redents Read present in caratyons	A mixture of potan- (3.5 Per. ar. ar. 2012) (3.6 Per. ar. 2012) (3.6 Per. ar. 2012) (4.6 Per. 2012) (4.6 Per. 2012)	P. Thin sections	(85) Iron-hematozylin (54)	Carraius injection method, shep. sil	Table of Ti
Competence of a contract Competence of a contract Contraction of a contract Contraction of a contract Contraction of a contraction		Zanitas (2) or Flam. Zanitas (2) or Flam. Zanitas (2) or Glam. (10) 33 Assetta n. Assetta n. Assett	0 0 8 8 8 6 9 6	Hennikovskih, sesin (62, 150 150 150 150 150 150 150 160 100 100 100 100 100 100 100 100 10	Poor instantions men Ma, 175,000 00 Doc That definition that a Doc That definition that a boot of the fit with a docentration defin The start and a docentration for the fit was a docen-	ssues and Organs
Contric glanda (fresh)	. Itabbić Large intentine	Frent Press and Press of moderna manifestate Correstives sublimate 1 C(14)	ä	Picrosoftanting (30) Mallory (31)	Tease in D.5 p.e. solution of sodium chloride	245

Constraint of	
NOLLVAYAAAA	
SUCHUSIN	
OBCANAS	
224	
SHIISSIL	
ÅQ.	
TARLE	

4	6					A	nit	na	1	lic	roi	ay	y							
	General Romacka	Technique same as for "Cella of Paneth"	Food the aginal on fat bacon for a couple of days	Perfore follows	ž	membrane should be made	Inject a, ognoputrated ag, sol. of Borlin blue	duct. Avoid too great pressure		Make sections both parallel and vertical to the	entface.	After stations, shake out cells by shaking sections					water to mounted spectmens. To see gramples teams fresh pareress in normal saline (83)			To prevent the shripkage of connactive tissue of vill away from the spithalium, allow the its testing to good before fixing.
	Staining, etc.		Bufranin (72)	Hematoxylin, sosin (52,	Hematowylin, cosin (52,	4.3)	Mallory (31)		d	(02, 40) Do		100	Thionin (2 to 3 drops of sub. aq. sol. to 5 c.c. of	water), 10 munites	Gold chloride (49)	Hematoxylin, Congo red	Dieges in boyax carmine	De	Hematoxylin, cosin (52, 43)	Do. Also stain pleces in borax-carmine before sectioning (33)
	Bestion or Isola- tion Nethod P Paradiu F Paradiu H Freeshand		P. or tease in nor-	P. or C.	ó		5: or P.		Thin sections	P. or C.		F. or H.	P.		T consec	Å.		Å.	P. or C.	P. er C.
	Fishing and Hardening, or Other Freihninery Treatment		Osmie acid, 1 p.e. (21)	Zenker (0) or Gilson P. or C. and B. anto.	Zenker (6), or Tellyen-		Alcohol (3)		Zenker (6) or Gilson P. This sections	Acetic alcohol (3)		Muller (9) or Zenker F. or H.	Corrout ve sublimate	IOL 2 DO 2 HOULS		Zenker (6)		Å	Å	Po
	Autronit, Orsam, or Thenue Frishing, and Hardoning. Demunatration or Or Presenting unary		Frog; rat		Lower hp of man or dog		Guinen-pig		Man, pig. or dog	Pig		å	Glanda (sections and thin membranes)		Stomach or duodenum of	Man or dog		Do	Small intention at its point of opening into the large	Kitten or puppy
	Object or Element	Grandies of salivary	fat From Prom	Large intestine	Lip or dog	Liver, amyloid infiltra-	Liver, bile capillaries Guinen-pig		1.5	Liver, hepatic lobules	Liver, interlobular con-	Teethe tissite	Mucin	Norve plexuses (all-	montary canal)	Panoreas		Parotid gland	mated modules)	time!! Intesting

Stomach, cardiac end	Dog	Acctic alcohol (2)	0 8 4	133), or boyas comin (52,	
Stomach, fundus and	ÅÅ	ÅÅ	00 55 54	20	Cut some longitudinal sectors to show transition
Sublinguel glands Taste-budary glands	Foliate papila of rabbit	Flomming (12)	P. Thin sections	Bafranin, gentian-violet (73)	Technique same as for D Panerease Orient the buds excelly for sectioning. Both Dongludinal and cross sections are instructive
Tongue (monoral)	(onSuch	Acctle alcohol (2) or D. or C.	P. or G.	Hematoxylin, cosin (52.	Cross-sections
Tongue, papillae and Mucous membrane of up- follicuit finguates per surface (rabble, mau)	Mucous membrane of up- persurface (rabbit, mau)	å	р. ег О.	Å	For fungitions and the of tensure. For full- values patillase, react of tensure. For follouth, react of tensure.
Tonell	Cat or rabbit	Auction alcohol (2), or P. or C.	P. or C.	Å	
Tooth, decalcified	Caritor tooth of dog. cat. Sheep of his embryu	Fis Tellygenicky	P. or F. Serial	Fig. in Tellygenicky P. or F. Serial Bornz-carmine, pierio acid	Technique mune se for decalcified boue. See also First pieze, 7 cm. embryo. Second stage, 10 cm.
Tooth, enamel prisms		P.o. mitrio herd (93)		Î	Tome out bits of engined from teeth prepared i a
Tooth, ground hasts	New-born child or other Müller (9), 6 to 8 days young animal	Müller (9), 6 to 8 days	Withdraw pulp	Withdraw pulp Piero-carmine (69)	From Conditional and the second secon
V. EAR			CONTINUE OF LC		
Coructionus glands	Guines-pig or rabbit. Cut away the lower Jaw and prove the three ar the dispeting three ar the	Alcohol (3) File 10 Flow minus (13) G. or P. C ut prof. dismutive mini prof. dismutive mini	G: or P, C ut from have to	Iron-hematoxylin (54) Do	In the grounswarpik decomposition is secondulated two to the tunno of the conduct conductor to a second to priority 1, do, thus under the lutuid so that air
Eustachian tube		Zenker (6) or Gilson	ő	Hematoxylin, cosin (52.	Sections should be transverse and should include the carfillant
Middle car		Zooker (0)	00 55 64	22	
endings of cochies	Young fetuses; new-born	Colsi method (50), or methylen blue (56, 5)			

Object or Element	Autori, Orgent, or Thouse Frising and Princhening. Recommendation	or Other Presiment	Focus or foolar from Mathod F Paraffin F Pressing H Free Hand	Staining, etc.	General Remarks
Otolithe	Young dog: skate	Flemming (12). Also decade ffy in guar of four "control".	, d	Bufranin (72)	Chinel into the secondula repove hits of the manual.
Culated epithelium	Roof of frog's mouthing ill plate of mussel, clam. or				Scrape the surface with a control and examine (53) in material thus obtained in normal eaches (53).
Ciliated columnar opi-	Trachea: epididymus	Gilson (76) or Zenker P. or C.	P. or C.	Hematoxylin, cosin (52, 43)	
Columnar and glandu-	Αã	åå	P. er C.	åå	
Endothelial cells	Shit open a medium-sized yein or artery from a freshly killed eat or dog	Subject the inner aur- field of the inner aur- nitration 175, unlyer high field of 15	To be mounted	Expose to surfight until a brownish-red color is visible, then at an in hematoxylin (52)	The venued should be planed out fist, endothelium obleve medical solution in the solution line ob- velues metery is to inject the solution into the
Intercellular bridges	Epidermia of larval sala- mander	Flemming (12)	P. Thin	Irgn-bematerylin. aoid fuchain (54, 45)	Also mount bits of spidermis flat withput section-
	Presh spithelia	Formaldehyde diago-		Tease, or shake Piero-carmine (69)	Examine in the dissociating fluid or in dilute glycerin
Mesothelial cells	Central tendon of dia- perioardium peritoneum, or	н		Expose to surfight until a brown other of any lightly	Rinne in distilled water before placing in the silver nitrate and again upon removal from it. Mount in giveerin, or dehydrate and mount in
Pigmented epithelium	Pigment layer of eye	Formalin, preserved To be mounted	To be mounted	Carmalum (35)	Dashmara

1			-		See "Cornea," "Bkin," "Reophagus," etc.	
		Gilson (16) for 20 min.	To be mounted	Hematoxylin (52)	Examine also scrapings from inside your cheek	
Terminal bars.	Smail pieces of intestine Corrective sublimate P. Thin Iron-hematoxylin 100, or Flamming	Corrosive sublimate (14) in normal sa-	P. This	Fon-hematoxylin (34)		
Transitioual epithe-	Transitiousi epithe Bladder of frog. guines-	30 p.c. alcohol (90) for 24 to 16 hours. Dis- tend the bladder with some of the		Plore-earmine (69). Stain to toto for 24 hours	Pierce-earmine (00). Stain Serape of some of the mucoue membrane and in toto for 24 hours	
VIII. BUYER Vessels of eye-	Injected eye (chap. sii) of albino rabbit or rab	Alcohol (3) or Muller Bisect into ante- (6) (3) or Muller Fior hod roste-	Bisect into ante-		The haives of the rat's cychall may be dehydrated	Table
Cboroid Cornea	Frenh	Flemming (12)	Tomo Thin	Iron-bematozylin (54)	Examine in glycerin. Por sections see "Eyeball"	d I
		Use the gold-chloride method (chap. xi), the method (39, b) blue method (39, b)			, seauce	Tissues
Constitution areas and Fresh cyo	Fresh cye	Scrape the epithelium from the cornea and then the cornea and then with ablok all-		Place in water for 2 to 3, days	Tangential acc. Place in water for 3 to 3 Spaces and canalicul show dark upon a light to the four with a resort	and A
Eyeball General)	Bisset the eyeball into an-	Muller (93) or Flem-	ó	Borna carmine, pierie acid (33, 05)	After fixing, the part desired may be sectioned and monoted. Small eyes may be fixed entire.	TOO
Illyelid	Evelid of an infant	Tellyesnicky (5)	P. or C.	Hengatoxylin, cosin (52.		8
Harder's glands. Tris	Rabbit (median angle of Sec. Eveball.	Zenker (6) or Gilson (16)		å	Do not confuse with lachrymal glands Meridianal metions of the actorian half of the ane	
Lachrymal gland	Man or rabbit	Zenker (6) or Gilson P.	, d	Hematoxylin, coain (52.	will show it. In case of the rabbit do not confuse with Harder's	
Lens	Bad or sheep	Muller (9)	ö	Boras corraine, pierie acid	Make an artero-posterior sections, also other sec- tions, at right angles to these	
	Å	Flemming (12)	÷.	Hematosylia, cosin (52, 43)	Fix the lens entire, then peel off the anterior cap- sule for sectioning	24
						0

			Animal I	lierology						
General Remarks	Junctional lines of lens fibers will be sevo. Peel off lamina and tease	Wash in distilled water and examine in giveerin or dehydrate and mount in balaam. See also	Four sections about Plane through the forces con- training and the section plane of Nethine from "Sec Colling 150% or markiveling morthod (26, 5)"	Becomines in non-mail solution (htt). If the propagate will appear purplish reacting the propagate propagate will appear purplish reacting the mark become provide the propagate of the propagate propagate propagate provide propagate propagate propagate propagate propagate provide propagate propagate propagate propagate propagate propagate propagate propagate propag			Examine in giveerin and mount in giveerin-jelly, or dehydrate and mount in water	Longitudinal and transverse sections		Tease on a slide and examine in the dissociating fluid
Staining, ata.		Expose under surface to sunlight until it becomes brownish red	Hematexylin, orange G			Henneoxylin (52)	Picro-carmine (69)	Aeld hematorylin (53)		
Section of Isola- tion Nethod P. – Paralla F. – Celloidin F. – Freezibin H. – Freezibin	Tenne	Cut out cornes	4			P. Thin	Tenne, or shake	P. or O.		
Fixing and Hardening, or Other Preliminary	0.2 p.c. sol. potamium biohromate for a	For 2 to 3 days Draw off squeous hu- pro- allyer hitrate	Fluxon to the fully of the second sec	Rapidly cut out a set third of the poster third of the poster ball, will of the poster	out the return		Macerate in 20 p.c. Tenne, nr shake Picro-carmine (69)	Corrective sublimate	MacCallum (89)	Dissociate in 35 p.c. caustie potanh for 15 min. (80)
Automic Organ, or Tiasue Fraine, and Hardonine. Dismonstructure	Beef or sheep	Rabbiti fowl	Beth mammal (human if a number and amphili- tain and amphili- tain and antipation and an and antipation and an and an and an an a			Man; lingual muscle of Flemming (12)	Tongue of from	Man; dog; sheep	Adult or embryo	Gastroensmius of frog
Object or Element		Memory Descention of Descention Rabbits fowl	R. estimation		VIII. MUSCULAR TISSUE	Areas of Cohnheim	Branched striated Tongue of from	Cardiac muscle	Cardiac muscle, iso- lated fibers Adult or embryo	Eads of striated fibers Gestroenemius of fros

TABLE OF TISSUES AND ORGANS WITH METHODS OF FREPARATION-Continued

					Tal	ile q	f Tu	isues and	Organs					25	1
	Ends of teased fibers show fibrillae. To make a perinshop thereading the propagation wash thereading.	Treases on a slide and examine in the dissociating		3		Within 5 or 10 minutes the sareolemma separates from the muscle substance. Examine in nor-	On of the thin muscles from the leg of Hydrophi- line (a water beside) is the chaste symmic of insect muscle. Examine without adding fluid:	Curavia and entrepretion rate observable of the local second areas and wash in discilled water before to second areas and wash in discilled	Make loogitudinal and tranverse sections. In local control of the section of the	See also "Corebral cortex," "Cerebellar cortex," etc.	Tease in normal saline (83) and examine under		Also try Golgi method (70). See ohap. ix. Failures with the Gelgi method are more	Also try Golgi method (50). See ohap. ix	
			Iron-hematoxylin, aeid fuchim (54, 45)	Piero-carmine (69)				Hematorylin (52), Need not be statned, hewever	P. Very this see fat, sq. sol. of add fuchain tions for a days) benefits also for a days) benefits also.	Beals's carmine (36) di-	Che mascrated haid	Borax-carmine (33), or	Bergy-contraste, pierie acid	Do	
	Tease in water		Á	Tenne, or shake		Tease		Tenne	P. Very this sec-	Tenne	Tenne	ú	P. er C.	P. or C.	
	Maccento in 0.1 p.c. Tease in water	Dissociate in 35 P.0. Cautie potent (86)	Flemming (12)	Macorate in 20 p.c.	(0/00)	Cold asturated solu- tion of annionium	Isolate by teasing io normal saline (83)	Stretch the muscle by Tenne cwterthy in set 0.5 trenity in set 0.5 trenity in set 0.5 trenity in the by the		Malcobel for 24 bours		Muller (9)	Muller (9) or Erlicki (3)	De	
~	Frog	Small muscle with its ten- fon (e.g., sartorius of	Bladder; intestine	Intestine, stomach, or	See " Purkinje fibers" un-	Fresh strinted muscle	Free: manual: wioz (the-	Â	Small nerve (lay it along a toothinck or amali appr- ter without atrefoling.	Small pieces of brain	Finest body	Brain and cord			
	Fibrillae in stristed	Muscle to tendon	Cootions)	Nex-striated fbers (wolated)	Purkinje fibera	Sarcolemma	Striated fibers (fresh)	Striated fibers (fixed).	AKONS (akis cylinder).	Brain cells	Brain sand	Central nervous sys- tem (general topography) Brain and cord	Corobellar cortex	Cerebral cortex	
									ř.						

52		Anim	ıl Mie	rolog	y		
General Remarks		defined the match have the control conduct. This is a second second seco	Mount under (197) stain In sectioning, cut from corium toward epithelium In carentium (33)	Do:	Remove all fat before freating the tissue with any of the solutions. Leads for the tissue with any conserving the events of the solution.		Finally examine in a drop of normal saline (89).
Staining, etc.	Hematowylia, eosia (53, 43) Methylen blue (59, b or c)	Fron-hommeorylin (54) di- Bieale's comment (36) di- futed with 1 voit of the maccerating guide of		Hermatoxylin, cosin (52,	43) Gold chloride (49)	Boras cacmine, piorio acid	
Section or Isola- tion Method C: Calicidin H Free Hand	To be mounted flat	F. Thin Teace	II. Iver hardened	P. Thin hardened	C. Very thin sec-	P. or C.	Tease loose and
Fixing and Hardening, or Other Preliminary	Flood with active al- control (2) for 30 milling then wash well in 50 p.o. alco- hol	Macoratolimates (25), Macoratolimates (25), Adama (27) in Gase a	Treat thin pieces with II. In hardened (21) "(1) the name of the land of the l	Muller (9) or Zenker		Multer (0) for 8 days Number 3 parter and 1 pot owner action Multer 2 parter and 1 pot owner action Multer (0) of Ernoki	Press an end of the prove throw you by the the throw you by our the prove the throw the provements of the prove
Animal, Organ, or Tissue Recommended for Demonstration	Epread out piecese under shoore one which abows blood vessels well abows Fresh seleral conjunctive	cous membrane See "Marchi's method" Generation of the sec- ficient correlation from Films correlation from	Waxy skin from lateral edges of duck's or goose's upper braket	, Po	Shein from freebly ampu- tated toe or finger (volar surface)	Nerve or cord, 2 to 4 weeks after the lesion has heen made	Small freeb nerve
Object or Element	Choroid plexus	Degenerating fibers Ganglion canalicul Ganglion cells	Chrandry's corpuscies.	Herbst's corpussies	Intra-opithelial nerve	degenerating fibers . Medula objongata	Meduliary sheath

Medullated fibers of		Muller (9) or Erlicki		Weigert's hematoxylin (78)	
Medullated nerve	Small nerve	I p.c. cemic acid (21) Tease	Tease	(Dehydrate. On a slide tesse and at the same time clear small pieces in clove oil. Mount
Meissner's corpusales	Papillas of corium on volar purface of haod, inger- tip, or foot			Gold chloride (19) or methylen blue (39, 5	
	Frog vabbit	Camic acid (1 p.c. for frog; 2 p.e. for rab-		Dehydrate. Clear with bergamot oil (45 hours)	The off dissoftware out the myelin and thus renders Visible the neurokesatin
Nerve-endings (n		The sold chieride (49). (56, 5 of 5), or the chief silver (50)			The methods are enumerated in order of their inusels see chap. is
Nerve-fiber bundles (transverse sections)	Scintic	Zepker (6) or Gilson C. or P.	0. 01 9.	Carmalum, Lyons blue	
Neuroglia	Spinsi cord	Muller (9) or Erlicki	Å	Bafranin (72) for 24 hours	Safrabia (72) for 24 hours Differentiate in absolute alcohol. See also Goldi
Wiest's granules)	Lumbar colargement of apinal cord	Corrosive sublimate (14), formalin (18),		Methylen blue (59. d)	See also neutral red (64)
Nodes of Ranvier	Small nerve		Towns	Silver nitrate (75)	
one medulisers	Sympathetic nerve; vagua 0.5 p.c. camic acid Tease	0.5 p.e. camie neid	Tense	Picro-carmine (69) 24 to	Piero-carmine (69) 24 to Medullated fibers (myellu) are stained black; 43 hours
		from fat		appears brownish	bodies between the strands of fat
	ter or new-born guines-				
transverse sections)	Cat	Corrosive sublimate C. or P. (14)	C. er P.	Hematoxylin, cosin (52, 43)	This parafic sections may be stained in Ehrlich- Biondi (41)
Spinal cord; azones and cells (transverse sections)	Cat or dog	Müller (9) for 4 weeks. Transfer to stain without washing	C. State before sectioning	1 p.e. ac. sol. sodium car- minate (3 days)	Maller (9) for 4 weeks. C. Static before 1 p.c. net. nod, acdium car-Before dehydrating wash the staticed bieves for victority washing accioning a

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			Anin	val I	licrolog	I		
General Remarks	Hermatoxylic. cosin (52, Spinal ganglia of higher vartebrates are difficult 43) usually be substituted for them		Oversiaio with hemutoxy- Differentiate in alsoholis sol. of potassium ferri- oyanide	Scrape off a little of the spithelial covering and examine in glycerin			¢.	
Staining, etc.	Hematoxylin, cosin (52, 43)	De	Overstaio with hematoxy-	Piero-earmine (69)	30 for 5 with the summer of the state of the		Hematoxylin, cosin (52, 1.00-hematoxylin (54)	Hematosylin, cosin (52, 43) Do
Beetion or Isolar tion Method D:	4	Å			P. Make section perpendicular to sprface of membrane		0 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
Pixion and Hardening. or Other Prelimituary	Zenker (6)	Do	0.33 p.c. ad, ad, of	30 p.c. alcohol (90) 24 hours, followed by 1 p.c. conformed	30 p.e. 5 minutes followed by 1 p.e. followed by 1 p.e. bours starden in	Golgi method (50), or method (50), or	Gilson (16) or Zenker P. nr C. (6) Dn P.	Acetic alcohol (2) Gilaon (16)
Auffinal, Organo, or Theurs Pratos, and Hardening. Decommendation	Cat or dos	Frog, mammal (first tho-	Second to the second se	Small pieces of officiery mucous membrane	Rabbit (divide head long) todinally: mand mucous membrane is of brown-	Nerve processes of ol- factory cells	Human; monkey Rabbit	Ttabbit Child dos, eat. or rabbit Azeara dos neu. 11. chur. XVI, men. 30.
Object or Element	Spinal ganglia (sections) Cat or dog	Sympathetic gaugin	Tactile cerpuseles	NOSE Solated olfactory cells Small pieces of olfactory membrane	Mucous membrane .	Nerve processes of ol-		uters (trans-
				×			i x	

PREPARATION-Continued
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256			lam	11	lien	logy				
pontenued	Ganeral Romarks	Before sectioning wash the preparation in dis- tilled water and transfer it to 70 p.c. alcohol		Carmine or Barlin-blue injection through pul- monary artery. Nee chap. xii	(66), or resorain Also, treat pieces of fresh lung with about 15 p.c. potassium hydrate	Hematoxylin, sosin (52. Fill the lung with the fixing fluid as well as km-	The stain should differentiate the chief from the	(52. If very accurate results are desired the mucous membrane should be removed and sectioned	atora and the second alteredes with draw (rout, the Manual and remain attaction to the epidemia	The hair of the mouse and of most bats is peculiar
TABLE OF TISSUES AND ORGANS WITH METHODS OF PREPARATION-Continued	ŝtaining, etc.	Safranin (73)	Hematoxylin, cosin (52, 43)		Organ (56), or resortin	Hematoxylin, sosin (52, 43)	Ehrlich-Biondi (41)	Hematoxylin, cosin (52, 43)	Acid hematoxylin (53)	
SCIOHTEM HTIV	Bection or Isola- Fion Mathod P. Callodin P. Callodin H. Freeshor	e sit	C, or P,		, A	P. or G.	4 5 04	C. or P.	Mount flat with under aurface upward	
TES AND ORGANS 1	Fixing and Hardening. or Other Pretiminary	Flemming (12) Fill the lung with 0.5 Fill the lung with 0.5 polution advertant fion for acvertant	Gibon (16) or Zenker (6) Trebnique same as	HONOWALL LOA	Alcohol (3)	Acetic alcohol (2)	Plenning (12)	Acctic alcohol (2) or Florming (12)	Maccerate in 0. 25 p.0. accerate in 0. 25 p.0. accerate accerate accerate	from dermis
TABLE OF TISSU	Animal, Organ, of Tissue Fixing and Hardening. Recommended for Other Freinmany. Demonstration	Kalabit Kittean					New-born child	Child; ent	BERTY AVD. TT: A.P. Sterry and sevents an	Examine in water (under cover-glass) or mount dry in balsam
	Object or Element	KII. RESPIRATORY OR	Fetal lung	Lung, blood vessels of	Lung, clastic tissue of	toposraphy)	Thyroid gland	Traches Child; ont		Hair
		***							*00.*	

Rate, elemente: Weight, filt with brinding and the state from the state of t					I	ahli	e of 1	Fissue	8 QX	d ()rga	118			25
Weight 10. Methods	Tensee if necessary and examine under a cover-		The orientation should be precise, so that exactly longitudinal or eross sections result.		Make vertical sections through nipple and gland		Add a drop of piero-carmine (99) and examine collectum in normal saline (83). Avoid pres- ente of the cover-glass	Myke longitudinal sections of the satire piece of finger	Transfer to a slide without staining and examine in the dissociating fluid	To section in paraffin, take only small pieces. Examine in giveerin		Inless with Berlin blue or mirnios mass through ulone artery. See chap. Kit		For spithelial cells see "Transitional spithelium"	First inject the fresh kidney (through renal artery) with carming mass. See chap. Xil
Weight for substruction and substr			Do	Å	Å	Bafraoin (72)		Alum-carmine (29)		Unstained	Iron-hematozylin 54)	Unstaioed		Iroa-hematozylia (54)	Acid bematoxylin (53)
 M. Martin, R. M. Martin, Parkins and Martin Martin, S. M. Martin, Markins, S. M. Martin, M. M. Martin, S. M. Martin, S. M. Martin, S. M. Martin, M. Martin, S. M. Martin, S. M. Martin, M			ö	ď	Ď	C. or P.					ő				ö
			Zenker (6)	Tellyesnicky (5)	Zenker (6)	Flemming (12)	Examine milk in nor- mainaine (83) with-	Multer (9); theo de- calcity (92); hardeo a peccod time in al-	Heatin strong saustic	Flemming (13)	Zenker (16)			Discend the bladder	Alcohol (3) or Zenker
Rate, alcontenter Rate, contenter Rate, contenter Rate	Warn in sulphurie acid	Skin from forehead (not	Preferably the upper hp of	Evend of new-born child.	Preferably human. Nipple	Mammal during gestation	Obtain colourum from a program parturition	First finger-joint of little finger		Stratum Malpighii	See sections of hair follicie Volar surface of finger or	Haud of child. Forefoot Bog cat of data, also in the uncher-surface view of epidermia			Rabbit or eat
	ir, elements	air, development	air folliele	air, renewal	country wind	pecial) ary gland	filk and colostrum	all (sections)	all, elomonts	rickle cells	ebaceous glands	kin, blood vessels	SINARY ORGANS	adder	idney, blood vessels

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ontinued	General Remarks	Hematoxylin, cosin (53. Make radial horizontal sections embracing the whole organ			Bealin 24, hours in henne. Examine in givenrin, tensing further if necessary					For male urothra see " Penis" under XII
TABLE OF TISSUES AND ORGANS WITH METHODS OF PREPARATION Continued	Staining, etc.	Hematosylin, cosin (53, 43)		Ivon-hematosylin (54)	Stain 24 hours in hema- toxylin (53)			Hematoxylin, cosin (52, 43)	Iron-hematosylia (54)	De
WITH METHODS	Bection or Inola- tion Method C. Paraffin F. Freeshin H. Free Hand	ť	Tenne	P. Thia	After staining. tense in dis- titied in dis- titied sherify alkeline with alkeline with	drate		Å.	Å	P. or C.
JES AND ORGANS	rixing and Hardening. or Other Dreitminary.	Acetic alcohol (2) or Gilson (16)	5 p.c. aq. sol. of neu- tral ammonium	Zepker (6) or Gilson P. Thia	After 15 min. remove, purses 1 ptd. recove, purses 1 ptd. recove, more the fill of the fourthy becoughly	Treat as for "Cortex	Golgi method (50), or methylen-blue	Zenker (6)	Gilson (16)	Ď
TABLE OF TISSI	Animaal, Orman, or Therus Remonstration.	Small manmal			A freehly killed animal, If and atrong (75 p.c.) If 1 into high veses ately high pressure	Young animals				Human; dog
3	Object or Element	Kidney, cortex, and	tubules of uriniferous	Midney; giomerulus and its capsule	Kidney: isolation of uriniferous tubules	Kidney: medullary rays (vertical sections) Young sairnels	Eldney, nerves of	Suprarenal gland	Ureter	Urethra (female) Human; dog

APPENDIX D

PREPARATION OF MICROSCOPICAL MATERIAL FOR A GENERAL COURSE IN ZOOLOGY

(In addition to the methods enumerated here, see also II, chap. x, and ehap. xiii.)

PROTOZOA

a) Cultures .- Amebae, etc., may usually be obtained in quantities sufficient for class use by the following method recommended by H. S. Jennings (Journal of Applied Microscopy, VI, No. 7, p. 2406). A number of glass dishes measuring 8 or 9 inches in diameter by 3 inches deep are crowded full of water plants (especially Ceratophyllum and Elodea), filled with water, and the plants allowed to decay. Keep the dishes in warm, light places. In two or three weeks the layers of plants at the surface of the water will be covered with a brown slime which should be examined occasionally under the microscope for the desired forms. The scum that appears on the surface of the water consists mainly of bacteria upon which amebae largely feed. They will be found most frequently in the slime that immediately surrounds the plant tissue. Since they frequently last only two or three days in a culture, to insure material for class work, a number of cultures must be made at different dates and from different localities. Other protozoa such as Arcella, Diffugia, Carchesium, Stentor, etc., will also be found in the cultures.

As soon as the anchoe appear in such cultures, several days before they are desired for use, Smith (*American Naturalist*, XXXIX [1996], 467) ekins off the brown scum and puts it in small bacteria dishes (4X1] inches) with enough water to fill the dish about 1 inch deep. He adds a little of the decaying regetable matter from near the surface of the original culture, covers the dishes and keeps them in a worm place but out of direct sunlight. In this way numerous large active specimens may often be obtained.

Pieces of frog or mussel allowed to decay for about 10 days in pond water will usually afford an abundance of a small species of ameba.

Barker keeps amelae and parameria from dying out by adding a sheet of fish food whenever the culture begins to be depleted. For a pure culture method see Kofoid, *Transactions of the American Microscopical Society*, XXXIV, No. 4 (October, 1915).

Chilononas and Infusoria usually appear in a few days in cultures of hornwort and partly decayed water-lily leaves packed in batteria dishes as for amelese, but with proportionately more water.

Parametium may be kept from dying out by keeping bits of stale bread in caltures. A culture of pond water and bread will usually develop large numbers of parametia in from a week to ten days. See also last paragraph under Euglena.

Euglena will be found in some of the cultures, but usually not in any quantities before the end of four or five weeks. They appear along the side of the dish toward the light. Stephensen finds that a few grams of pulveined rice covered with pond water provides an abundance of Eugleon in from ten days to two weeks.

Turner balls 20 grams of dry quince seed for half an hour in 1½ liters of distilled water, then passes the thick exulate which is given off, together with the water, through a wire sierce (Einer and Amend, No. 80). He then makes up the volume to 2½ liters with distilled water, sterilizes it and places it in a stoppered bottle. The medium will keep for months. Cultures made by inovalisting tubes or fasks of the medium with Eugleon will keep for a year, and specimens can be obtained in considerable numbers at any time after four weeks. Cultures will keep longer in a thicker medium but will not reproduce rapidly.

Standard masses for use in experimental work may be prepared by evaporating the exolute to dryness and making up solutions with distilled water. A 0.2 per cent solution of the dried exolute seems to furnish the optimum density. Mold frequently invades the cultures, but will not grow in a density of 0.2 per cent or less. Mold growths are less Ekely to occur in cultures which have been rendered slightly alkaline. Cultures should be kept at room temperature and in a moderately lighted place.

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Parameeia and other infusoria will live and reproduce for about two months in the medium, feeding upon Euglena and bacteria.

Carchesium and Vorticella are frequently found on decaying duckneed (Lenna) and hornwort (Contophyllum). To scene a culture, have a more plentiful supply of water than for anelase. Professor Walton tells me that he always finds a supply of Epidylis on the shells of fresh-water scalls.

Difinium, a form which feeds largely on parametia, is highly recommended by Mast (*Science*, December 20, 1912, p. 871) as of great value in biological study. It is easy to obtain (in parametia cultures), shows the phenomena of fission and encystement with particular clearness, and has a remarkable method of feeding. Didnia can be kept indefinitely in the encysted state, and when wanted for study will appear within 24 hours to a few days in active form if introduced into a vigorous culture of parametia.

Optima may be obtained readily by killing a frog with chloroform and slitting open the large intestine. Examine scrapings of the epithelial wall in normal saline (reagent 83, Appendix B).

Sportona.—Gregarina may be found in the alimentary canal of the meal worm or the excitoach and Monocystis in the male reproductive organs of the earthwarm. They are best studied in normal saline. If it is desired to stain and mount specimens, they may be fixed in corresire-acetic (reagent 15, Appendix B) for 5 minutes, washed thoroughly in 35 per cent alcohol, to which a little instruct of iodine has been added, and stained with Ehrlich's triple stain (reagent 42), or hematorylin and acid fuchsin (reagents 52 and 45).

Herpetomonas may be obtained from the intestine of the fly and of the squash bug, and Trypanosoma from the blood of the rat.

Volvez.—Volvez globater, the form commonly described in textbooks, is found in the early spring, often in great abundance, in small permanent pools which outsin duckweed and Riccia. A smaller and less desirable form, Volvez ourcus, may be found later in the spring and throughout the summer in the same pools. When water from such pools, together with a small amount of the water plants, is placed in bacteria dishes, so arranged that one side is strongly exposed to light, the volvex present will collect after a few hours at the edge of the water on the lighted side of the dish. If the contents of the vessels which contain volves are kept in as near the natural condition of the pool as possible, the organisms may be kept alive for some weeks in the laboratory. Tap water is injurious to them. Avoid having too much decying material in the water, although some is essential. Keep in glass-covered dishes near windows (out of direct sunlight) in as cool a place as possible. Any considerable rise in temperature beyond that of the original pond will result in their death. Small crustacea feed upon volves and will, if present in any considerable numbers, soon externing the them. The second stages are more likely to be found in the cone at the bottom of cultures.

Because of the uncertainty of obtaining living volvon at any stated time, it is well to have an abundance of the material preserved in 5 per cent formalin. Such preserved specimens show the flagella more distinctly than do living ones. See Smith, *The American* Notorolid, XLI, No. 451 (1907), p. 31.

b) Quieting infournin.—1. Let sufficient water eraporate from under the cover to permit the latter to press lightly upon the animals. Guard against too great eraporation of water or the infusoria will be erashed.

 Entanglement in fibers of cotton, etc., sometimes proves efficacious.

3. A small amount of gelatin, or, better, cherry-tree gum, dissolved in water makes a viscous mass which is often useful in retarding their motions. A bit of white of egg may be used in the same way.

4. Animals may be narrotined by means of a small drop of very dilute should (reeferably methyl should) or ebloretone (about one drop of a 1 per cent solution to 10 drops of water). (Chloretone is manufactured by Parke, Davis & Co., of Detroit, Mich. For its use as an anasethetic in biological work see Journal of Applied Microscopy, V, 2051.)

c) Feeding--Place finely pulverized earnine or indigo under the cover-glass. The colored powder rapidly accumulates in the food vacandes. In such a preparation the action of the cilia of

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infusoria is also indicated by the rapid movement of the particles in the vicinity of the animal. See also memorandum 4, p. 110.

 d) Staining.—For intra-tilam staining see reagents 59a, 32, and 64, Appendix B.

To see cilia of infusoria treat the animal with very dilute iodine solution or a drop of a dilute solution of tannin.

To see the macronucleus and the micronucleus use a drop of a 2 per cent solution of acetic acid, or, better, methyl green (reagent 60, Appendix B).

e) Permanent mounted preparations.-Benedict's method is as follows:

"Snear a glass slide with allouren fixative, as in preparing for the mounting of parafin sections. Then place on the surface of the film of fixative a drop or two of water containing the form which it is desired to stain. Let nearly all the water evaporate by exposure to the air of the room until only the film of fixative remains moist. The slide can now be immersed in Gilsen or any other fixing reagent, and then passed through the alcohols, stains, etc., in the same way that mounted sections are handled.

"I have had no difficulty in getting preparations of parametrium by this method, with very little distortion of the body and any kind of staining desired. By this method students can prepare in ten minutes very satisfactory preparations of protonos for demonstration of nuclei, etc."-Journal of Applied Microscopy, VI, 2047.

For firstion of protonos Calkins (Journal of Experimental Zoldoyy, I, No. 3, 1904) uses estimated aqueous solution of corresive sublimate to which 10 per cent of glacial acetic acid is added. See also reagent 20 (formal sublimate), p. 214. Barker fives, washes, stains, destains, dehydrates, clears, and mixes protonos with balsam, all in homeopathic vials. After each operation material is allowed to settle well. Reagents are pipetted off.

Plankton, in general, are well fixed if passed directly into fresh Zenker's fluid to which a few drops of 1 per cent solution of osmic acid have been added. Under such treatment eilistes remain expanded.

SPONGES

To isolate the spicules of calcareous sponges, boil a bit of the sponge in 5 per cent solution of caustic potash for a few minutes.

Fairly thick transverse, longitudinal, and tangential sections of Groutia showing spicules in the tissues are useful. Make these with an old ranor or sharp scalpel. To hold the object while sectioning, place it between two pieces of pith or cork. For a careful study of the relations of the two systems of canals in the body wall, thinner sections are necessary. To prepare these it is best to decalefy (2 per cent chronic acid, 24 to 36 hours) the sponge and cut cellohifm or parsfin sections on the microtome, although fairly good sections may be made by hand. They should be dehydrated and mounted in balsam if permanent preparations are desired; if not, they may be examined in giverin.

To color the collar cells use an aqueous solution of anilin blue.

Spicules of *eliceous* sponges are isolated by treating bits of the sponge with strong nitrie acid or a mixture of nitrie and hydrochloric acid.

COELENTERATES

Hydra should be sought for in spring-fei pools. In the autumn they are found most frequently on mooth deal leaves which are completely submerged. Material should be collected and placed in hottery jars or larger glass jars, which are then filled with fresh, elear water, and placed in a fairly light place, but not too near a window. Put a small amount of hornvort or Chars in each jar. In a few hours (12 to 30) the hydra will be found attached to the sides of the vessel and to the plants. They may readily be kept in the laboratory throughout the winter if glass plates are placed over the jars to prevent excessive eraporation and the temperature is not allowed togo below freezing. Fresh water should be added from time to time to make up for evaporation. In ease their supply of food (Cydops, Duphnin, and other small erastace) is exhausted it should be renewed by skimming out from other aquasin the small forms upon which the animal feeds and putting them in the hydra jars.

Keeping hydra in the dark at somewhat lower temperature for several days favors the formation of spermaries and ovaries.

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For staining and mounting entire see p. 95. Kill in the same way for sectioning. The most instructive sections are (1) transverse sections, (2) longitudinal sections through the mouth and a bud, and (3) sections showing the secual organs. Stain in bulk with hematorylin (reagent 52, Appendix B), inded in paraffin, using the method for delivate objects (p. 53), and after the paraffin has been removed from the sections, stain them for a few seconds in acid fuclsin. Dehydrate and mount in the usual way.

The sections are much more satisfactory if the hydra have been placed in small stender dishes filled with filtered water (not distilled) and kept from ford for a week or ten days before killing. This eliminates the metabolic products and oil globules which ordinarily obscure the details of structure.

To Stain the Nematorysis of Living Hydra, place several of the animals in a small stender dish of water which has been introl a sky blue through the addition of methyler-blue solution made up as follows:

Methylen blue	1.0	gram
Castile scap	0.5	gram
Water	300	C.C.

After two hours the hydra may be transferred to fresh water; the memotoryst cells are stained a deep blue. (Method of Little, Journal of Applied Microsiopy, VI, 2216.)

To Discharge Nematocysts drum on the covergless gently with a pencil. By using a very small opening to the disphragm they are usually sufficiently distinct without staining.

For Other Polypoid Forms the methods given for hydra will answer in most cases.

For Collecting Free-Swimming Medusoid Forms full directions will be found in Brook's Invertebrate Zoology.

Compound Hydrozoa should be placed affive into the cells which they are to occupy when mounted; 1 per cent formic acid is then added drop by drop to the sea water. After the animals have been killed, the fluid is replaced by glyceric jelly and the cover-glass is put in place. Another method is to kill the animals slowly by adding a

few crystals of chloral hydrate from time to time to the small vessel of sea water containing them.

Small Jellyfish may be fixed and hardened in 1 per cent cemie acid and, stained or unstained, mounted in cells.

Anenones, Medusae, and other definite marine forms may usually be killed in the expanded condition by means of magnesium sublate. Success lies in securing a quick diffusion of a quantity of the subplate through the water without eausing mechanical disturbance of the animal to be anaesthetized. Griffin accomplishes this by tying a considerable quantity of the magnesium subplate in a piece of chesseoloth and hanging it over the dish of sea water containing the animals in such a way that the bottom of the hag hearly dips into the water. Mayer's method of anaesthetizing mechanese by earloan dimide is also often applicable to other sensitive contractile forms.

PLANARIA

Look for planarians on the under sides of stones in small streams of running water. They are usually examined alive. To see them thrust out the protosets, keep them from food for a few days and then feed them on dead files. Planaria which have been kept in the laboratory for months display the internal organs much more clearly than freshly exptured ones.

If it is desired to study stained specimens, for preparation see p. 97.

To Kill Planaria with Pharyux Protruded, Cole (Journal of Applied Microscopy, VI, 2125) recommends covering them in a watchglass with a 1 per cent appecus solution of chloretone until they are immobilized and then rapidly transferring them to 5 per cent formalin. Other fixing agents than formalin can be used.

TREMATODES

The most easily obtained forms are those found in the hungs, intestine, or bladder of frogs. A good form for study is occasionally found in the liver of the ext. Search for it in the bile passages. Fix trematodes in corresive sublimate, wash out with alcohol to which tincture of iodine has been added, and stain for 24 hours in shum-coefined (reagent 28, Appendix B) or camalum (reagent 35).

As with planaria, they should be compressed between two glass slides (see p. 97). To kill trematodes in a distended condition, Barker flattens out each individual on a glass slide or in a watchglass with a camel's hair brash and floods it with the killing agent.

If the large liver fluke of the sheep (Fascida hepatica) can be obtained, both the alimentary canal and the excretory system may be injected with India ink or with finely powdered earnine in water. For injection a very fine-pointed eannals with rubber cap is used, or the manipulator may operate the earnula by simply blowing through it. The excretory system is injected through an incision made with a sharp-pointed scalpel in the median line near the hinder end of the animal. For the alimentary canal the incision should be made about 1 mm. to one side of the median line. When the injection is completed, fasten the animal somewhat between two slides (see p. 97), handrain 95 per cent alcohol for 12 to 24 hours, then dehythate, clear, and mount in helsam.

Larval Stages may frequently be found in the so-called "liver" of pond snails.

CESTODES

Near large cities an unlimited supply of the sheep tapeworm (Monico) can usually be secured from slaughter houses. Ample supples can ordinarily be obtained from dogs, or, less frequently, from cats. Taperroms can be kept alive for a considerable length of time in tepid water. The most instructive portions to mount are sceler, securally mature, and terminal predictids. For fixing and staining use the same methods as for distances. La Rue fixis that carmine stains are better for trematoles and hematorylins for cestoles. The scelexishould not be compressed. To kill cestoles in an extended confiltin, Barker wraps the living worm around a glass shile, than immerses the slide in the killing reagent. The worm is removed as soon as killed.

To Find Cysthereri, open the body eavily of a rabbit and look for large whitish bodies imbedded in the peritoneum or liver (the cysticercus of *T. corotal)*. Likewise, the cysticercus of *T. crossicallis* may be found in the liver of the mouse. If a cysticercus is found, its outer wall should be slit open in order to show the reversed scolex.

NEMATODES

See memorandum 20, p. 136. Nematodes occur frequently in the intestines of pigs, dogs, cats, and rabbits.

Trichinella.-The simplest way to obtain it is to apply for infected park to the government inspector whose headquarters are to be



found near all large slaughter houses in cities. Bits of the infected nuscle should be teased and fattened out in a compressor (Figs. 73 and 74) until a favorable area has been found. The fattened

Fig. 73.-Compressor

tissue may then he dehydrated and monited unstained or it may be stained in hematoxyfin (reagent 52, Appendix B). Better results will be obtained if the material is fuxed for from 4 to 6 hours in Carnoy's fluid (reagent 2) before dehydrating or staining. If desired, the tissue may be sectioned in celloidin or parafin.



Fig. 74.-Compressor Used by the Government Bureaus for Meat Inspection

To Demonstrate Living Trictinellae, Barress (American Monkly Microcopical Journal, XIV, 104) subjects small hits of trictinined muscle to a mixture of 3 grains of pepsin, 2 drams of water, and 2 minans of hydrochlorie acid for about three hours at body temperature with occasional shaking. When the firsh and cysts are dissolved, the liquid is poured into a narrow glass vessel and allowed to

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settle. The live trichinae may be withdrawn with a pipette from the bottom of the fluid and examined on a warm stage.

ROTIFERS

Rotifers will usually be found in abundance in some of the lalouratory aquaria on the lighted side of the vessel. For ordinary class work they are best studied alive. They are difficult to preserve properly. Full directions for killing and preserving will be found in Jenning's paper, "Rotatoria of the United States," U.S. Fish Commission Bulletis (1992), p. 277.

To Quiet Rotifers, Cole (Journal of Applied Microscopy, VI, 2179) anaesthetizes them by adding from time to time a drop of 1 yer cent aqueous solution of elloretone to the water on the slide in which the animals are being examined.

BRYOZOA

They may be treated in the same way as compound hydraoa. Planudello may frequently be found in shallow fresh-rester streams on the under side of flat rocks; Peritoatello, in rivers and streams on the upper surface of massel shells, etc.

STARFISH

Barker's technique for Policellaria is as follows: Boil the aboral part of a ray from a formalin-preserved startish in 5 per cent caustic soda for from 3 to 5 minotes. Wash quickly and thoroughly in water, stain in water-coluble cosin, wash in acid (acetic) alcohol, dehydrate, and mount.

EARTHWORMS

Earthwards are best collected on warm, rainy nights when they may be found extended on the surface of the ground near their burrows. They are most plentiful in old gardens or rich lawns. A lantern and a peil are the only implements necessary. Earthwarms may frequently be found, however, in large numbers on the surface of the ground on cloudy days immediately after prolonged hard rain. In winter living ones can nearly always be found under manure plac.

To Prepare Earthwoms for Class Work, secure good-sized specimeus, wash them in water, and place them in a vessel containing moist filter-paper. Put only a few worms in each dish and adjust the cover so as to admit a little air. After 12-to 24 hours it is well to remove any dead or injured specimens and to change the filter-paper. The dish should be kept from direct sunlight in a cool place. After two or three days the grit and dirt in the alimentary canal will have been passed out and its place taken by paper which the worms have eaten. They are then ready to kill and preserve or section.

Place the worns in a flat vessel and pour on sufficient water to cover them. During the next two hours add a little alcohol from time to time until the storegth of the liquid is increased to about 8 or 10 per cent. Then wash all mneus from the body of the worns and replace them in 10 per cent alcohol until they no longer respond to pricking or pinching with forceps. Transfer them to 50 per cent alcohol for several hours, herping them straightened out as much as possible; then to 70 per cent alcohol for 12 hours, followed by 95 per cent alcohol for 24 hours. Preserve finally in 70 per cent alcohol.

Chronic Acid Method—Although requiring considerable more work in preparation, specimens hardened in chromie acid are somuch superior to alcoholic ones for general dissection purposes that the extra trouble is well worth while. The worms are anaesthetimed as in the preceding method, but from 10 per cent alcohol they are injected with 1 per cent apneous solution of chromic acid and then immersed in it for 4 hours. While working with the chromic acid the hands and wrists solutib to conted with vaseline.

Keeping the worms extended and submerged in 1 per cent chromic acid in a large shallow dish, inject the acid into the body eavily slowly, about half an inch behind the editellum, and again near the posterior end of the body. Avoid pieving the alimentary canal. The injection is not complete until the worm is turgid along its entire length. The worms must be kept straight and untwisted while in the chronic acid. Remove them at the end of 4 hours (a longer time in the acid will make them heitile) and wash thoroughly in running water until the yellow color is gone (12 to 16 hours). Remove them to 50 per cent alcohol for 2 days, then to 70 per

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eent alcohol for 2 or 3 days, and finally preserve in fresh 70 per cent alcohol.

For injection a water-pressure apparatus (Fig. 33) is best. The reservoir A should be placed about 4 feet above the compression chamber B. The cannula should be made of a piece of quarter-inch glass tuding with one end drawn out to a very fine hore and so broken as to kare a sharp point and edge for piercing the body wall of the worm.

For actioning, the preliminary steps are the same as in the alcohol method, but from 10 per cent alcohol the worms should be placed into Zeaker's fluid (reagent 6, Appendix B) for 4 to 6 hours. For washing, etc., follow the directions given in the discussion of the reagent. To facilitate penetration of the fluid, it is well to shi open the body earity of the worm in places that are not to be sectioned. The most instructive sections are cross-sections of the middle of the body and sagittal sections of the anterior end which include the plarynx. The worms may be stained in bulk (24 to 36 hours) in boarx-carnine (reagent 33) or hematorylin (reagent 52) before sectioning.

Entire neptridu together with a small part of the septum which they traverse should be carefully dissected out, stained in boraxcarmine (reagent 33), dehydrated, eleared, and mounted in halsam.

An overy should be removed entire, stained with borax-earmine, dehydrated, cleared, and mounted in balsam.

A testis should be treated in the same way as an ovary. Tease it in the balsam before adding the cover-glass.

To Keep Earthworms Alive in Winter, Jennings (Journal of Applied Microsopp, VI, 2412) places them, immediately after collection, into bacteria diskes (9 in. in diameter by 3 in, deep) between folds of muslin which is kept damp but not dripping wet. Not more than a donen worms should be placed in one disk and the cloth should be changed or washed at least every two weeks. The worms may be fed on haves, etc., from time to time.

To Immobilize Earthworms for study of eirenlation of the blood under the microscope or projection lantern, Cole (Journal of Applied Microscopy, VI, 2123) places them in a 0.2 per cent aqueous solution

of chloretone for 3 or 4 minutes. Such worms may be slightly compressed between two slides.

To Examine Corpuscles of the Coelonic Pluid, expose the worms for a minute or two to the vapor of chlorotorm. The coelonie fluid exudes through dorsal pores. Touch a cover-glass to the fluid and mount.

The Setae Can Be Isolated by boiling a bit of the tissue containing them in a solution of caustic potash. When isolated, dry them and mount in belsam.

LEECH

Levelos are obtained from fresh-water pools, streams, and marshes, but to get sufficient numbers for class use it is manily necessary to purchase them from dealers. Live leveloss intended for dissection may be killed with eblecoform. Cross-sections prepared in the same way as for earthworms are very instructive.

ARTHROPODS

For Mounting Small Crustacea see III, A, chap, xiii. To Quiet Small Crustacea for Microscopical Examination (Cole, Journal of Applied Microscopy, VI, 2180) place them in a watchgluss containing 2 parts of 1 per cent chloretone and 5 parts of water. The same treatment is useful for the karvae of insects. Some, such as the normsh of the dragon fly, will require more chloretone.

For Various Dissections and Parts of Insects see II, chap. x.

For Mounting Inserts Entire (teetles, mospitoes, grats, aphile, larvae, etc.) as microscopic preparations, and for mouting nursele, wings, heads, legs, scales, antennae, etc., see chap, siii.

Live nymphs of the dragon fly are especially valuable for study under the compound microscope because they show very clearly the valvakr action of the heart, the tracheal gills and tracheae, and the brain and its relation to the eyes. The heart is located well toward the posterior end of the abdomen between the main tracheal trunks. Cole (Journel of Applied Microscopy, VI, 2274) recommends that the animals be anaesthetized by subjecting them to a 1 per cent appends solution of chloretone.

MOLLUSKS

Gills of the Fresh-Water Mussel may be fixed in corresive sublimate (reagent 14, Appendix B) for from 20 to 30 minutes, washed out in water and then in dilute alcohol to which instrue of iodine has been added. Make cross-sections in parsfin, stain in dilute hematarylin (reagent 52), and mount in the ordinary way.

Cross-Sections of the Entire Mussel are valuable to show the relations of the galls, kidneys, and heart. Wedge the values apart slightly and immerse the animal for 24 hours in 1 per cent chronic acid (reagent 11). Wash out thoroughly in running water and transfer the specimens to 70 per cent alcohol for two or three days or until needed. To section, remove both values, place the animal on a board, and with a more cent transverse sections. These are to be examined with the naked eye or with a dissecting lens.

To KII Stails in an Expanded Condition, put them into a vessel of cold water, then run a layer of hot water on to the surface of the cold water. See that the vessel is full of water and cover it with a glass plate to exclude the air.

For Lingual Ribbon of the Snail see memorandum 7, p. 100.

AMPHIOXUS

Specimens must ordinarily he secured from dealers. The animals should be stained entire in borax-carnine (reagent 33, Appendix B) and sectioned in celloidin. The most instructive sections are crosssections of a female with well-developed gunads and longitudinal sections of anall individuals. Mounts of entire small specimens should also be made.

VERTEBRATA

For any of the tissues of vertebrates which teachers may desire to prepare, ample directions are given in Appendix C.

For Demonstration of Circulation of the Blood in the frog, see ehsp. xiv.

APPENDIX E

TABLE OF EQUIVALENT WEIGHTS AND MEASURES

WEIGHTS, METRIC AND AVOIRDUPOIS

1 kb=1,000 grans=1 liter of water at its maximum density=2.2 pounds. 1 gran=1 cubic continueter of water at its maximum density=15.43 grains=0.003 curve. 1 pound=433.59 grans. 1 curve=23.35 grans. 1 grain (Troy)=0.065 gran. 1 dran=1.77 grams.

WEIGHTS, METRIC AND APOTHECARY'S

1 kilo=1,000 grans. 1 gran=15.43 grains=0.002 ounce. 1 point=373.24 grans. 1 ounce=31.10 grans. 1 dran=3.89 grans. 1 serupk=1.30 grans. 1 grain=0.005 gran.

MEASURES OF LENGTH, METRIC AND ENGLISH

1 meter = 1,000 millimeters = 30.37 inches. 1 centimeter = 0.394 inch. 1 millimeter = 0.024 inch. 1 yard = 0.914 meter. 1 foot = 30.45 centimeters. 1 inch = 2.54 centimeters.

LIQUID MEASURES, METRIC AND APOTHECARY'S

1 liter=1,00 cubic centimeters=2 11 pints. 1 cubic continenter=0.034 fluid ounce=16.23 minims. 1 guilen=128 ounces=3.79 liters. 1 pint=16 ounces=473.18 cubic centimeters. 1 fluid ounce=8 fluid denms=29.57 cubic centimeters. 1 fluid denm=60 minims=3.70 cubic centimeters. 274

Equivalent Weights and Measures

THERMOMETERS

To reduce degress Fahrenheit to degress Cartigrade use the formula, C=5/9(Γ =32). For example, if the number of degress Fahrenheit is 77, then C=5/9(Γ =32)=25 degrees. Or, for instance, to reduce =31 degrees. Fahrenheit to Cartigrade, C=5/9(-31-32)=5/9(-63=-35 degrees. To reduce degrees of Cartigrade to degrees of Fahrenheit use the formula F=9/5C+32. For example, if the number of degrees Cartigrade is 25, then F=(3/5)(25)+32=77 degrees. Or, to reduce -35 degrees Cartigrade to Fahrenheit, F=(9/5)(-33)+32=-31 degrees.

APPENDIX F

REFERENCES

Only a very limited habiligraphy is given. A full one will be found in Gage's *The Microscope*. Above all, for desirable special methods the student is advised to look through the articles in **cur**rent journals which cover the field of his own researches.

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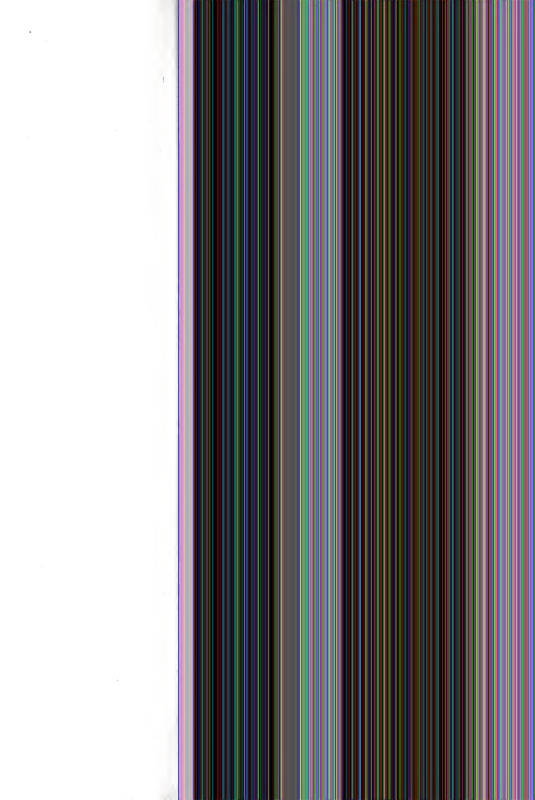
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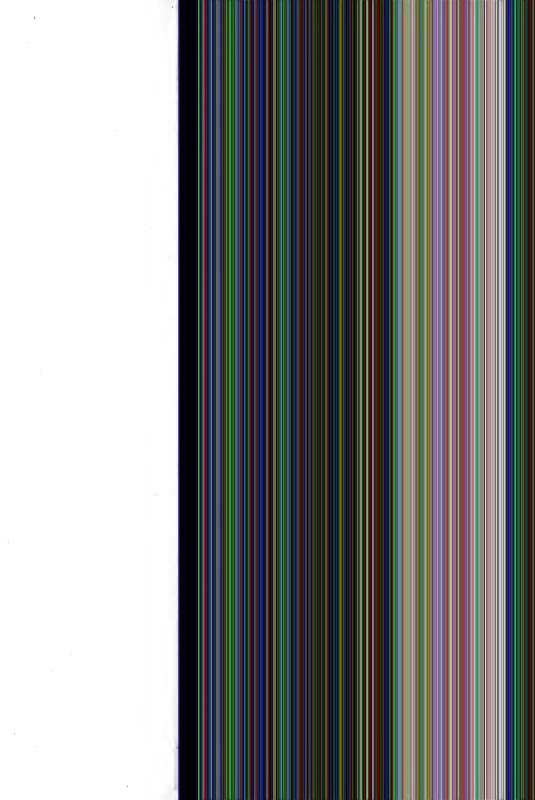
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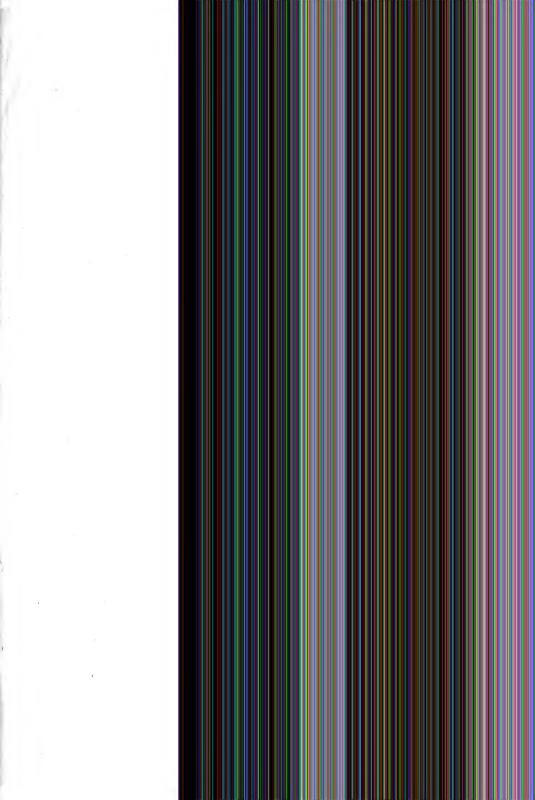
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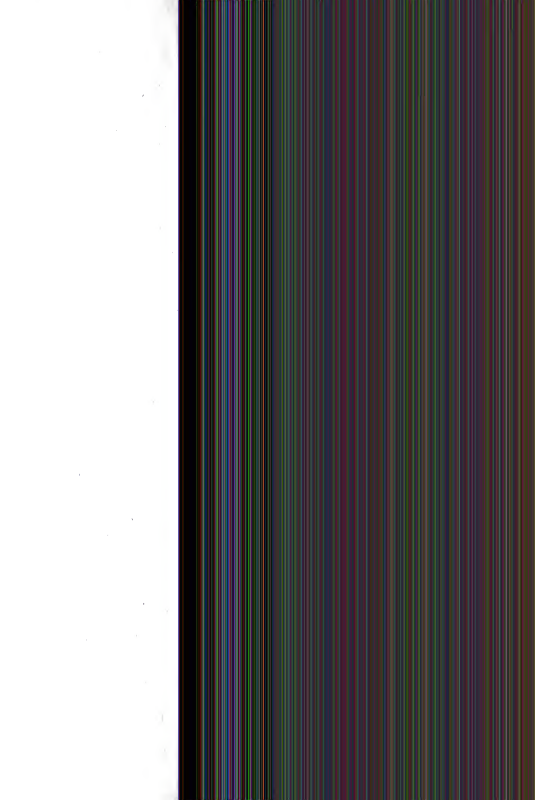
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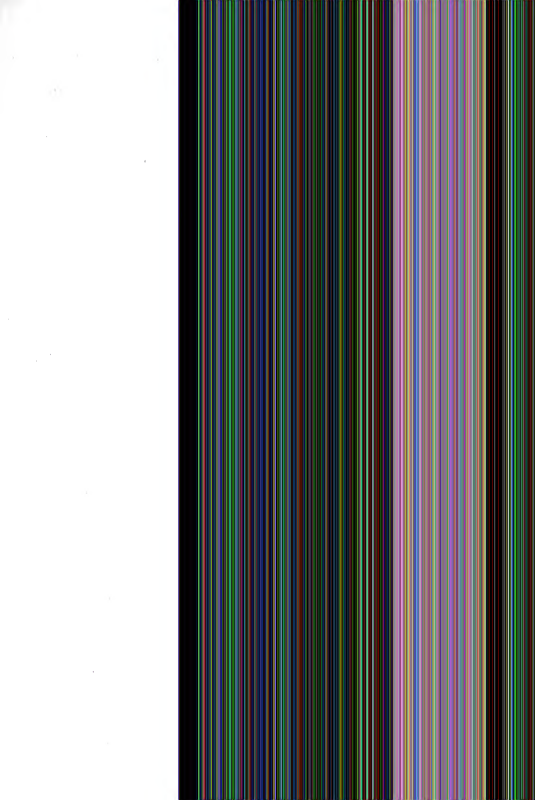
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