

X-RAY STRUCTURAL STUDIES OF SOME GROUP VIII
COMPOUNDS WITH CATALYTIC IMPLICATIONS

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To Jeanie

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KEY TO ABBREVIATIONS

LIPS	ligand-induced proton shift
H ₂ dmg	dimethylglyoxime
dmg	dimethylglyoxime dianion
Hdmg	dimethylglyoxime monoanion
H ₂ dmg ₂	bis(dimethylglyoximate) with relative proton positions unspecified
sulfa	sulfanilamide
dhph	1,4-dihydrzinophthalazine
dhphpy	1,4-dihydrzinophthalazinebis(2-pyridinecarboxaldimine)
pyca	2-pyridinecarboxaldehyde
clan	4-chloroaniline
H ₂ dph	diphenylglyoxime
H ₂ mpg	methylphenylglyoxime
fph	pentafluorophenyl
cp	cyclopentadienyl anion
tpp	triphenylphosphine
an	aniline
4-FPYTSC	4-formylpyridinethiosemicarbazone

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X-RAY STRUCTURAL STUDIES OF SOME GROUP VIII
COMPOUNDS WITH CATALYTIC IMPLICATIONS

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X-ray structural investigations of compounds containing Group VIII metal atoms are presented. The compounds studied illustrate interatomic interactions which may be of importance in catalytic processes. The structures of metal-containing compounds were solved by locating the heavy atoms in Patterson functions and locating the remaining atoms in Fourier syntheses. The direct method of symbolic addition was used in the one, all light-atom case presented. Trial structures were refined by the method of least-squares.

The crystal structure of trans-chloro(dimethylglyoximate)(dimethylglyoxime)(4-chloroaniline)cobalt(III) illustrates an unusual ligand-induced proton shift. Both neutral and dianionic dimethylglyoxime groups are found in the complex and the 4-chloroaniline ligand is oriented over the dianionic dimethylglyoxime. The structure of trans-bis(dimethyl-

glyoximato)bis(4-chloroaniline)cobalt(III) chloride shows that complex to contain two monoatomic dimethylglyoxime ligands and the 4-chloroaniline ligands to be skewed relative to the diglyoxime ligands. The crystal structure of trans-chlorobis(diphenylglyoximato)(4-chloroaniline)cobalt(III) is described. Trends in the structures of these compounds and in the previously reported structures of similar compounds are discussed. Ultraviolet and infrared spectra of these compounds are given.

The synthesis of a novel chelating ligand capable of binding two metal ions is described. The characterizations, including crystal structures, of its protonated form, 1,4-dihydrzinophthalazinebis(2-pyridiniumcarboxaldimine) nitrate dihydrate, and of a nickel complex, μ -chlorotetraqua[1,4-dihydrzinophthalazinebis(2-pyridinecarboxaldimine)]dinickel(II) chloride dihydrate, are presented. The planar ligand is shown to bind two nickel ions with a separation of 3.603 (1) \AA . A chloride ion occupies a bridging site in the plane of the nickel atoms and the ligand. The magnetic moment per nickel atom of the chloride bridged complex was determined to be 2.74 B.M. at 40°C. The plausibility of structurally similar complexes mimicking the nitrogen-fixing enzyme nitrogenase is also discussed.

The X-ray crystal structures of 1-(π -cyclopentadienyl)-1-triphenylphosphine-2,3,4,5-tetrakis(pentafluorophenyl)cobaltole and 1-(π -cyclopentadienyl)-1-triphenylphosphine-2,3,4,5-tetrakis(pentafluorophenyl)rhodole are reported.

These compounds are viewed as stabilized intermediates in the catalyzed cyclization of acetylenes. In each case the metal atom forms a metallocycle by σ -bonding to the terminal carbons of a butadiene-like fragment. The π -bonding in the metallocycle appears to be delocalized.

CHAPTER 1 INTRODUCTION

Western civilization has demonstrated the efficiency-oriented phenomenon of expending large amounts of energy to find ways of requiring less human energy. This is evident in the evolution from animal trails to freeways and from muscle to sophisticated, high-energy machinery. On the molecular scale the more efficient path is provided by catalysts. As alchemists searched for the "philosopher's stone" many chemists have been seeking catalysts. The application of catalysis is now advancing through the development of an understanding of the mechanisms of catalytic processes.

Life processes are dependent upon chemical reactions controlled by enzymes. "It is not generally appreciated how little is understood about the mechanisms by which enzymes bring about their extraordinary and specific rate acceleration."¹ Investigation of enzymes should not only be fundamental in the understanding and maintenance of life processes but also should contribute to developing more efficient industrial processes.

Much of the investigation of enzymes has concerned the use of model compounds. "Model building and the application of material analogues are becoming increasingly important for the elucidation of fundamental problems of biochemical

structure and reactivity."² X-ray structural studies of enzyme models are important for the exploration of structure-activity relationships. Solid state studies of enzyme model compounds are of particular relevance because of the high degree of order the macromolecular enzymes themselves possess.

While electrostatic and hydrogen-bonding forces are usually considered the major binding forces in enzyme-substrate interactions, the strong charge-solvating and hydrogen-bonding ability of water tends to reduce the possibility of obtaining large binding energies from these forces. To explain the large binding energies found, "hydrophobic forces" are presumed to exist in these intermolecular interactions in aqueous solution.³ The enthalpies of mixing of aromatic liquids with aliphatic liquids indicate that aromatic molecules prefer an aromatic environment.^{4,5} "Stacking interactions" involving the π -systems of aromatic groups within the enzyme's protein structure may account for part of the "hydrophobic forces" and contribute to the orientation of the enzyme-substrate interaction.³ The ligand-induced proton shift (LIPS) observed in $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{sulfa})$ [the key to abbreviations is given on page x] is an indication of the importance of this π -type interaction. A further examination of LIPS was undertaken and is presented in this work.

The design of enzyme models is often based on sparse structural information about the prosthetic group of the enzyme. Efforts to mimic the nitrogen-fixing enzyme nitrogenase

have been concerned with the metal to nitrogen bond. The probable binuclear nature of the enzyme's active site^{6,7} has largely been ignored. The structures of a novel binucleating ligand and its nickel(II) complex are presented here as a first step in the construction of a new generation of models for nitrogenase.

When the mechanism of a chemical process is believed to be understood, stable compounds similar to the intermediates of the reaction may be prepared and examined to support the proposed mechanism. One proposed mechanism for the catalyzed cyclization of acetylenes would have a five-membered ring containing a metal atom and a cyclobutadiene fragment as one of the intermediates.⁸⁻¹³ The first structure of such a stabilized intermediate containing a cobalt atom and the structure of the rhodium analog are presented in this study.

CHAPTER 2
SYNTHESIS AND CHARACTERIZATION

Synthesis

Crystals of all cobaloxime compounds were generously provided by R. C. Palenik* and were used without recrystallization.

M. D. Rausch and R. H. Gastinger synthesized the metallocycles containing cobalt¹⁴ and rhodium.¹⁵ They supplied well-formed crystals of those metallocycles for X-ray structural studies.

Unless otherwise indicated all solvents were reagent grade and were used without further purification. All preparations were carried out in air. All melting points were taken on a Mel-temp apparatus in open capillaries and are uncorrected.

The published method¹⁶ was used to prepare dph for succeeding experiments. To 6.40g (49.0 mmoles) 1,2-dicyanobenzene (98%; Aldrich Chemical Company, Milwaukee, Wisc.) in 12.5 ml 1,4-dioxane was added a mixture of 15.0 ml (ca. 250 mmoles) hydrazine hydrate (85%; Fisher Scientific Company, Fair Lawn, N. Y.) and 4.0 ml glacial acetic acid (reagent; Baker and Adamson, Morristown, N. J.). After being heated

*These complexes were prepared using standard procedures¹⁷ with synthetic details to be published at a later date.

for three hours the mixture was cooled and the red product was collected (yield, ca. 40%). The decomposition temperature of 193°C was in agreement with the reported value.

A solution of 0.0955g (0.50 mmoles) of the previously prepared dhph in 40 ml absolute ethanol was added to a solution of 0.237g (1.0 mmoles) $\text{NiCl}_2 \cdot 6\text{H}_2\text{O}$ (reagent; Matheson, Coleman and Bell, Norwood, Ohio) and 0.095 ml (0.99 mmoles) pyca (99%; Aldrich) in 40 ml absolute ethanol. Upon slow, almost complete, evaporation in air of that solution olive green crystals of $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhphpy})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$ formed.

Analogous procedures were carried out replacing $\text{NiCl}_2 \cdot \text{H}_2\text{O}$ with $\text{CoCl}_2 \cdot 6\text{H}_2\text{O}$, $\text{CuCl}_2 \cdot 2\text{H}_2\text{O}$ (reagent; Fisher), ZnCl_2 (reagent; Mallinckrodt Chemical Works, St. Louis, Mo.) and $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ (reagent; Matheson, Coleman and Bell) without success in obtaining a crystalline product. Similar procedures were followed with the addition of ca. 0.2 ml of 12 M hydrochloric acid (reagent, 38%; Baker and Adamson) to solutions of $\text{CuCl}_2 \cdot 2\text{H}_2\text{O}$ and $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$. Again, no suitable products were formed. Attempts to separate and recrystallize reaction products from water, water-ethanol, methanol and pyridine failed to give a crystalline product. When CuCl_2 was present, gas evolved from the reaction mixture.

Additional attempts were made to isolate complexes similar to $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhphpy})]\text{Cl}_3$ using dhph obtained by recrystallization from hot water of $\text{H}_2\text{dhphSO}_4$ (ICN-K and K Laboratories, Inc., Plainview, N. Y.) to which an equivalent

amount of KOH (certified A.C.S.; Fisher) had been added. Those attempts were unsuccessful.

The red-orange plates of $H_2\text{dhphpy}(NO_3)_2 \cdot 2H_2O$ used in crystallographic studies had been recrystallized from water. The crude product formed upon cooling a solution made by adding 0.190g (1.0 mmole) dhph in 20 ml warm water to a solution containing 0.583g (2.0 mmoles) $Ni(NO_3)_2 \cdot 6H_2O$ (reagent; Mallinckrodt) and 0.89 ml (9.4 mmoles) pyca in 10 ml warm water followed by drop-wise addition of nitric acid (reagent, 71%; Baker and Adamson) to a pH less than 1.

Also, $H_2\text{dhphpy}(NO_3)_2$ was prepared by first adding 1.90 ml (20.0 mmoles) pyca to a suspension of 2.878g (10.0 mmoles) $H_2\text{dhphSO}_4$ in 100 ml water. A brick-red solid formed upon addition of 1.11g (ca. 17 mmoles) KOH. After washing with water and drying in air, the brick-red solid was suspended in 100 ml of 95% ethanol and 1.30 ml (21 mmoles) of nitric acid were added. Small red-orange needles of $H_2\text{dhphpy}(NO_3)_2$ which decompose at 126°C were filtered, washed with ethanol, and then ether and air dried (yield 4.0g, 75%).

Freshly prepared hydrated metal hydroxides were reacted with $H_2\text{dhphpy}(NO_3)_2$ in methanol. Each of the metal hydroxides was filtered after adding 1 M KOH to aqueous solutions of $Ni(NO_3)_2 \cdot 6H_2O$, $Cu(NO_3)_2 \cdot 3H_2O$ (reagent; J. T. Baker Chemical Company, Phillipsburg, N. J.), $Fe(ClO_4)_2 \cdot 6H_2O$ (reagent; G. Frederick Smith Chemical Company, Columbus, Ohio) and $Zn(NO_3)_2 \cdot 6H_2O$ (reagent; Matheson, Coleman and Bell). After

the reaction mixtures were stirred until there was no further change in color, they were filtered and the filtrates were allowed to evaporate. Only the reaction with nickel(II) hydroxide produced a crystalline product. Attempts to recrystallize that maroon product from methanol, ethanol, ethanol-water, and 2-propanol did not yield crystals suitable for crystallographic studies.

Discussion of Characterization

The microanalyses recorded in Table 1 were performed by Galbraith Laboratories, Inc., Knoxville, Tennessee, for the dhphpy compounds and by Atlantic Microlab, Inc., Atlanta, Georgia, for the cobaloxime complexes. The calculated percentages of carbon, hydrogen, and nitrogen for the dhphpy compounds correlate well with the measured percentage. Two water molecules per molecule of dhphpy in each are indicated by the elemental analysis. This is confirmed in the structural determination. Similarly, the elemental analysis of $\text{ClCo}(\text{H}_2\text{dmg})(4\text{-nitroaniline})$ is in agreement with the expected formula with two water molecules present. Based on the measured density and crystallographic data the molecular weight of $[\text{Co}(\text{H}_2\text{dmg}_2)(4\text{-methylaniline})]\text{Cl}$ should be 596. This is greater than its formula weight of 538.9 and the presence of molecules of solvation is expected. Three water molecules or one molecule of the ethanol solvent per formula could account for the difference. Neither of these possi-

Elemental Analyses of Selected Compounds

Table 1

	%C		%H		%N	
	found	calc.	found	calc.	found	calc.
$\text{C}_\ell\text{Co}(\text{H}_2\text{dmg}_2)(4\text{-nitroaniline}) \cdot 2\text{H}_2\text{O}$	33.87	33.71	4.87	4.85	16.90	16.85
$[\text{Co}(\text{H}_2\text{dmg}_2)(4\text{-methylaniline})_2]\text{Cl} \cdot 3\text{H}_2\text{O}$	48.03	49.03	6.23	5.99	14.27	15.59
$\cdot \text{C}_2\text{H}_5\text{OH}$	44.56		6.46		14.17	
	49.28		6.55		14.37	
$\text{H}_2\text{dphphy}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$	45.36	45.29	4.12	4.18	26.10	26.40
$[\text{Ni}_2\text{C}\ell(\text{H}_2\text{O})_4(\text{dphphy})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$	32.39	32.65	3.84	3.84	15.22	15.23

bilities is confirmed by the CHN analysis (see Table 1).

IR spectra of samples as mineral oil mulls between polished plates of fused sodium chloride were recorded on a Beckman Model IR10 grating spectrophotometer from 4000 to 500 cm^{-1} . The spectra were calibrated using the 1601.0 cm^{-1} absorption of a polystyrene film. IR spectra of selected compounds are reported in Table 2. The IR spectra of the bis(diglyoxime)cobalt(III) complexes with aniline derivatives exhibit many features of similar cobalt complexes with nitriles and isonitriles described by Batyr *et al.*¹⁸ The spectra of the cobaloximes show the absorption assigned¹⁸ to the C=N stretch between 1550 cm^{-1} and 1580 cm^{-1} . The absorptions associated¹⁸ with the N-O band at ca. 1245 cm^{-1} and ca. 1095 cm^{-1} are present also. A weak absorption in the $1700\text{-}1800\text{ cm}^{-1}$ range appears in some of the spectra but with low resolution. Peaks in this region have been assigned¹⁹ to the O...H-O bridge between the dioximate ligands. The presence of a symmetrical bridge has been suggested²⁰ to rationalize this low frequency.

Absorption spectra in the ultraviolet region were recorded on a Cary Model 15 spectrophotometer. Spectra of solutions were measured from 26.7 kK ($375\text{ m}\mu$) to 47.6 kK ($210\text{ m}\mu$) using the double beam method with the pure solvent as the reference. Solutions of the cobaloxime complexes in methanol (spectroquality; Matheson, Coleman and Bell) and solutions of the dihydpy compounds in 0.1 M hydrochloric

Table 2
Infrared Spectra^a of Selected Compounds

$\text{C}\ell\text{Co}(\text{H}_2\text{dmg})^-$ (dmg) (Clan)	$\text{C}\ell\text{Co}(\text{H}_2\text{dmg}_2)^-$ (4-nitroaniline)	$\text{C}\ell\text{Co}(\text{H}_2\text{dpq}_2)^-$ (Clan)	$\text{C}\ell\text{Co}(\text{H}_2\text{mpg}_2)^-$ (Clan)
3525 (s)	3565 (s)	3535 (s)	3480 (m, b)
3425 (s)		3410 (s)	3360 (m)
	3195 (s)		3165 (m)
	3105 (s)		3065 (m)
		2405 (w, b)	
		2305 (b, w)	1897 (w)
		1930 (w)	
	1778 (b, w)	1730 (b, w)	
			1610 (w)
		1598 (s)	1580 (w)
		1563 (s)	1530 (m)
		1530 (s)	1490 (s)
			1445 (s)
		1323 (s)	1343 (s)
		1244 (s)	1292 (m)
		1203 (s)	1200 (m)
		1186 (w)	1168 (w)
			1156 (w)

Table 2 - continued

$\text{C}\ell\text{Co}(\text{H}_2\text{dmg}) - (\text{dmg})(\text{Clan})$	$\text{C}\ell\text{Co}(\text{H}_2\text{dmg}) - (\text{dmg})(\text{sulfa})$	$\text{C}\ell\text{Co}(\text{H}_2\text{dmg}_2) - (\text{4-nitroaniline})$	$\text{C}\ell\text{Co}(\text{H}_2\text{dmg}_2) - (\text{clan})$	$\text{C}\ell\text{Co}(\text{H}_2\text{mpg}_2) - (\text{clan})$
1152 (s)				
1085 (s)	1084 (s)	1088 (s)	1130 (s)	1138 (m)
				1085 (w)
			1013 (m)	1007 (s)
973 (b)	972 (m)	971 (m)		958 (m)
	922 (m)		920 (w)	
	837 (m)	858 (s)	885 (s)	
825 (m)	824 (m)	818 (w)	823 (w)	825 (m)
		798 (w)		
			757 (w)	780 (m)
742 (m)			743 (m)	733 (s)
705 (m)			686 (m)	685 (s)
645 (w)		670 (m)		

^aEach column contains the respective absorption peaks (cm^{-1}) and the relative intensity (s, strong; m, moderate; w, weak; b, broad).

Table 2 - extended

$[\text{Co}(\text{Hdmg})_2^2^- \text{Cl}_2^-]$	$[\text{Co}(\text{H}_2\text{dmg}_2)^{-} (\text{4-methylaniline})_2] \text{Cl}$	$[\text{Ni}_2\text{Cl}_2(\text{H}_2\text{O})_4^{-} (\text{dhpipy})] \text{Cl}_3$	$\text{H}_2\text{dhpipy}^- (\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$
3125 (s)	3420 (m, b)	3280 (s, b)	3460 (s, b)
2415 (w)	2400 (w, b)	1620 (m)	2050 (w, b)
2380 (w)		1517 (s)	1750 (w, b)
1892 (w)		1465 (s)	1609 (s)
1785 (b, w)		1380 (s)	1552 (s)
		1296 (w)	1290 (s)
		1285 (w)	
1612 (m)	1600 (s)	1260 (w)	1168 (w)
1582 (s)	1570 (s)	1224 (m)	
1493 (s)	1506 (s)	1137 (s)	1141 (m)
		1115 (s)	
		1096 (w)	
1234 (s)	1228 (s)		1057 (m)
1205 (s)	1197 (m)		950 (w)
	1158 (w)		914 (w)

Table 2 - extended - continued

$[\text{Co}(\text{Hdmg})_2^- (\text{Clan})_2]\text{Cl}_3^2-$	$[\text{Co}(\text{H}_2\text{dmg}_2)^- (\text{4-methylaniline})_2]\text{Cl}_3^2-$	$[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4^- (\text{dhpipy})]\text{Cl}_3^2-$	$\text{H}_2\text{dhpipy}^- (\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$
1083 (s)	1076 (s)	865 (w)	870 (w)
1008 (m)	1013 (m)	824 (w)	775 (m)
967 (m)	968 (m)	768 (m)	758 (s)
219 (m)			
805 (m)	808 (m)		
735 (m)		743 (m)	
700 (m)		701 (w)	
647 (m)			

acid were used. The UV spectra are reported in Table 3.

The UV spectra of all these compounds are dominated by intense charge transfer bands. Yamano *et al.*²¹ report three bands in this region for compounds of the formula $[\text{Co}(\text{H}_2\text{dmg})_2 - \text{A}_2]$ where A is an aniline derivative. These three bands are present in $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}$ and $[\text{Co}(\text{H}_2\text{dmg})_2(4\text{-methyl-aniline})_2]\text{Cl}$. The band between 25.0 and 27.5 KK (400 to 360 m μ) was assigned²¹ to the charge transfer from the aniline ligand to the cobalt ion. In agreement with this assignment the band for the complex of the more basic 4-methylaniline at 27.6 KK is lower in frequency than that for the analogous complex of clan at 28.9 KK. The band near 33.0 KK (300 m μ) was assigned²¹ to the charge transfer from the cobalt ion to the dioximate ligand. The band near 40.0 KK (250 m μ) was assigned²¹ to the intra-Hdmg $\pi + \pi^*$ transition.

The UV spectra of cobaloxime complexes with a chloride ligand trans to a substituted aniline show three bands, also. One band is between 27.0 and 33.0 KK (370 to 300 m μ). The other bands lie near 39.0 KK (255 m μ) and 43.0 KK (230 m μ). No assignments have been made for these three bands.

The charge transfer spectrum of a solution of $[\text{Ni}_2\text{Cl}_2(\text{H}_2\text{O})_4(\text{dhphpy})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$ in 0.1 M HCl exhibits the same absorptions as that of a solution of $\text{H}_2\text{dhphpy}(\text{NO}_3)_2$ in 0.1 M HCl. The intense bands at 25.4, 32.7, and 37.3 KK (395, 305, and 268 m μ) are presumably due to the aromatic system of the ligand.

Ultraviolet Spectra^{a,b} of Selected Compounds

[Co(Hdmg) ₂ (clan) ₂]Cl	28.9(16000)	[32.7]	39.7(21000)
[Co(H ₂ dmg) ₂ (4-methylaniline) ₂]Cl	27.6(11000)	32.8(7400)	39.7(16000)
ClCo(H ₂ dmg)(dmg)(clan)	32.7(9200)	39.5(24000)	44.8(24000)
ClCo(H ₂ mpy) ₂ (clan)	31.1(7300)	39.8(27000)	[44.1]
ClCo(H ₂ dpq) ₂ (clan)	29.7(12000)	37.6(43000)	42.4(42000)
ClCo(H ₂ dmg) ₂ (4-nitroaniline)	27.2(20000)	[39.6]	42.9(33000)
H ₂ dhphpy(NO ₃) ₂ ·2H ₂ O	25.4(5600)	32.8(4600)	37.3(4600)
[Ni ₂ Cl(H ₂ O) ₄ (dhphpy)]Cl ₃ ·2H ₂ O	25.4(25000)	32.7(20000)	37.3(20000)

^aThe compound name is followed by the absorption frequencies (KK) with the extinction coefficients in parentheses.

^bFrequencies listed in square brackets are for poorly resolved peaks.

The magnetic moment per nickel atom of $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhphy})]\text{Cl}_3$ was determined to be 2.74 B.M. at 40°C. Data for this calculation^{22,23} were obtained using a Varian A-60A Analytical NMR Spectrometer and aqueous solutions containing 2% by volume t-butanol as the indicator. This magnetic moment is in agreement with those of binuclear complexes of nickel reported by Ball and Blake.²⁴ Their complexes of the general formula $[\text{Ni}(\text{dhph})]_2\text{X}_4 \cdot n\text{H}_2\text{O}$ ($\text{X} = \text{Cl}$, Br, or I) had room temperature effective magnetic moments ranging from 2.79 to 2.89 B.M. As in the case of $[\text{Ni}(\text{dhph})]_2\text{X}_4 \cdot n\text{H}_2\text{O}$, where two Ni^{2+} ions are bridged by a conjugated system, spin-spin interaction is indicated in $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhphy})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$.

CHAPTER 3 X-RAY DIFFRACTION EXPERIMENTAL

Except where noted in the text, the experimental methods described in this section were used in preliminary crystallographic examination, collection and processing of data, and refinement of trial structures.

Data obtained using precession and Weissenberg X-ray photographic techniques²⁵⁻²⁷ were used in determining the preliminary space groups and cell constants. After centering fifteen intense reflections on a computer-controlled Syntax PI diffractometer and selecting an indexing consistent with preliminary photographs, accurate cell constants with estimated standard deviations were obtained from least-squares fittings of 2θ , Ω , x , and ϕ for those reflections. In each case the orientation matrix for data collection and the unit cell volume with its standard deviation were derived from these data. The calculated density was in agreement with the density measured by the flotation method²⁸ except in the cases of the metal-containing heterocycles. The specific gravity of the flotation liquid was measured to ± 0.01 with a precision hydrometer. Relevant crystallographic data for each of the compounds studied are given in Table 4.

The suitability of a crystal for data collection was determined by its physical shape and size, the ease with

Table 4
Crystallographic Data for A, $\text{C}\ell\text{Co}(\text{H}_2\text{dmg})(\text{dmg})(\text{cian}) \cdot 2\text{H}_2\text{O}$; B, $\text{C}\ell\text{Co}(\text{H}_2\text{dmg}) \cdot \text{C}_2\text{H}_5\text{OH}$;
C, $[\text{Co}(\text{Hdmg})_2(\text{cian})_2]\text{C}\ell$; D, $\text{H}_2\text{dhpipy}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$; E, $[\text{Ni}_2\text{C}\ell(\text{H}_2\text{O})_4(\text{dhpipy})]\text{C}\ell_3 \cdot 2\text{H}_2\text{O}$; F,
C₄(fph)₄Co(cp)(tpp); G, C₄(fph)4Rh(cp)(tpp); H, $\text{C}\ell\text{Co}(\text{H}_2\text{mpg})_2$ (cian); I, $\text{C}\ell\text{Co}(\text{H}_2\text{dmg})_2$ (4-nitroaniline) $\cdot 2\text{H}_2\text{O}$; J, $[\text{Co}(\text{H}_2\text{dmg})_2]\text{C}\ell$

Compound ^a	Formula	Crystal System	Systematic Absences	Space Group
A	$\text{C}_{14}\text{C}\ell_2\text{CoH}_{20}\text{N}_5\text{O}_4 \cdot 2\text{H}_2\text{O}$	triclinic	none	$\text{P}\bar{1}$
B	$\text{C}_{34}\text{C}\ell_2\text{CoH}_{28}\text{N}_5\text{O}_4 \cdot \text{C}_2\text{H}_5\text{OH}$	monoclinic	$h0\ell : h+\ell = 2n+1$	$\text{P}2_1/n$
C	$\text{C}_{20}\text{C}\ell_3\text{CoH}_{25}\text{N}_6\text{O}_4$	triclinic	none	$\text{P}\bar{1}$
D	$\text{C}_{20}\text{H}_{18}\text{N}_{10}\text{O}_6 \cdot 2\text{H}_2\text{O}$	monoclinic	$hk\ell : h+k=2n+1$ $h0\ell : \ell=2n+1$	$\text{C}2/\text{c}$
E	$\text{C}_{20}\text{C}\ell_4\text{H}_{24}\text{N}_9\text{Ni}_{12}\text{O}_4 \cdot 2\text{H}_2\text{O}$	monoclinic	$hk\ell : h+k=2n+1$ $h0\ell : \ell=2n+1$	$\text{C}2/\text{c}$
F	$\text{C}_{51}\text{CoF}_{20}\text{H}_{20}\text{L} \cdot ?$	triclinic	none	$\text{P}\bar{1}$
G	$\text{C}_{51}\text{F}_{20}\text{H}_{20}\text{PRh} \cdot ?$	triclinic	none	$\text{P}\bar{1}$
H*	$\text{C}_{24}\text{C}\ell_2\text{CoH}_{24}\text{N}_5\text{O}_4$	triclinic	none	$\text{P}1$ or $\text{P}\bar{1}$
I*	$\text{C}_{14}\text{C}\ell\text{CoH}_{20}\text{N}_6\text{O}_6 \cdot 2\text{H}_2\text{O}$	orthorhombic	$h\bar{k}0 : h+\ell=2n+1$	Pmn or $\text{Pm}2_1\text{n}(\text{Pmn}2_1)$
J*	$\text{C}_{22}\text{C}\ell\text{CoH}_{32}\text{N}_6\text{O}_4 \cdot ?$	monoclinic	$h0\ell : \ell=2n+1$	$\text{P}2/\text{c}$ or $\text{P}\bar{c}$

^a Data for compounds marked with an asterisk were obtained from photographic techniques.

Table 4 - extended

Compound	a \AA	b \AA	c \AA	α ($^\circ$)	β ($^\circ$)	γ ($^\circ$)	Volume \AA^3
A	7.494 (3)	11.838 (4)	13.758 (6)	106.31 (3)	91.25 (3)	112.79 (3)	1068.3 (7)
B	15.353 (13)	12.385 (3)	18.535 (13)	90	96.55 (7)	90	3503 (4)
C	6.336 (4)	2.710 (5)	12.719 (5)	90.55 (4)	105.16 (4)	98.83 (4)	673.9 (6)
D	20.480 (3)	11.166 (2)	10.704 (2)	90	102.99 (2)	90	2385.0 (8)
E	15.016 (6)	15.527 (7)	28.704 (17)	90	115.78 (3)	90	6027 (5)
F	11.680 (3)	14.008 (4)	20.455 (9)	114.08 (3)	107.41 (3)	106.72 (2)	2572.9 (1.7)
G	11.725 (4)	14.015 (6)	20.420 (6)	114.07 (3)	106.97 (3)	107.28 (3)	2574.3 (1.5)
H*	7.95	13.26	13.75	98.1	102.7	105.9	1330
I*	21.66	13.68	14.97	90	90	90	4436
J*	13.2	11.2	19.9	90	110.6	90	2750

Table 4 - extended

Compound	Molecular Weight	Z	ρ calc. (g/cm ³)	ρ meas. (g/cm ³)	Crystal Dimensions (mm ³)	Radiation Used	μ (cm ⁻¹)
A	488.22	2	1.512	1.52	0.24x0.18x0.07	MoK α	11.2
B	746.54	4	1.415	1.43	0.18x0.20x0.05	MoK α	7.1
C	578.75	1	1.426	1.44	0.19x0.31x0.35	MoK α	10.0
D	530.46	4	1.477	1.47	0.34x0.31x0.18	MoK α	1.3
E	735.73	8	1.622	1.63	0.29x0.30x0.14	MoK α	18.1
F	1102.79	2	1.423	1.59	0.27x0.31x0.50	MoK α	4.9
G	1146.57	2	1.479	1.60	0.14x0.24x0.43	MoK α	4.6
H*	576.3	2	1.439	1.47			
I*	498.8	8	1.494	1.50			
J*	538.9	4	1.300	1.44			

Table 4 - extended

Compound	μR	2θ Range	K	No. of Unique Reflections	No. of Observed Reflections
A	~0.2	0-45	2.0	2807	2000
B	~0.1	0-45	1.5	4364	2017
C	~0.2	0-45	2.0	1771	1662
D	~0.04	0-45	2.0	1573	1093
E	~0.5	0-45	2.0	3981	2959
F	~0.1	0-45	2.0	6772	5479
G	~0.1	0-45	2.0	6766	5235

which the reflections were centered on the diffractometer, and the values of the refined cell constants with their estimated standard deviations compared to the cell constants obtained by photographic methods. All intensity measurements were made with a Syntex PI diffractometer at ambient temperature. All unique reflections up to a limiting 2θ value were measured using a variable speed 0-20 scan technique. The scan rate was determined from a fast three-second counting scan of the reflection peak and varied linearly from $1^\circ/\text{minute}$ for counting rates of 150.0 c/sec. or less to $24^\circ/\text{minute}$ for 1500.0 c/sec. or more. The intensity, I, was defined:

$$I = (\text{scan rate}) \left[(\text{total scan counts}) - \frac{(\text{background counts})}{(\text{background to scan ratio})} \right].$$

Peaks were scanned from 1° below $K\alpha_1$ to 1° above $K\alpha_2$. Measurements of the background count were made at the limits of each scan. The estimated standard deviation, $\sigma(I)$, of each reflection was taken to be:

$$\sigma(I) = \left[(\text{total scan counts}) + \frac{(\text{background counts})}{(\text{background to scan ratio})^2} \right]^{1/2}.$$

For molybdenum radiation, the incident beam was monochromatized by a low order reflection of graphite. Any changes in the system were detected by measuring four standard reflections after each 96 intensity measurements.

A standardized data set was obtained by scaling the data to the initial value of the sum of the measured intensities of the standard reflections. The scaled in-

tensities of duplicate or equivalent reflections were averaged. Reflections with an intensity greater than $K\sigma(I)$, where K is given in Table 4, were considered reliable. The unreliable reflections with $I < K\sigma(I)$ were identified by a minus sign and not included in further steps of the structure solution. Corrections for Lorentz-polarization were of the form:

$$\frac{I}{I_p} = \frac{\sin 2\theta}{(1+\cos^2 2\theta)}.$$

To obtain a set of observed structure factors, F_{obs} 's, the monochromator was also assumed to be 50% perfect crystal and 50% mosaic crystal.

Scattering factors were obtained from Hanson, Herman, Lea, and Skillman;²⁹ Stewart, Davidson, and Simpson;³⁰ Doyle and Turner;³¹ and are uncorrected for anomalous dispersion. The natural log of the scale factor and the overall temperature factor were initially estimated from a Wilson pilot.³² The initial choice of a centric or acentric space group was made on the basis of calculated intensity statistics.³³

In the case where molecules contained at least one heavy atom (Atomic Number ≥ 16) the approximate positional coordinates were determined using a Patterson function³⁴ of the form:

$$P(UVW) = \frac{2}{v} \sum_{h=-\infty}^{\infty} \sum_{k=-\infty}^{\infty} \sum_{l=-\infty}^{\infty} |F(hkl)|^2 \cos 2\pi(hU+kV+lW).$$

Using the location of the heavy atom(s) in a structure

factor calculation allowed a sufficient number of reflection phases, $\alpha(hkl)$'s, to be assigned. The magnitude of the structure factor, $|F_{hkl}|$, and the phase may be defined by the following equations:²⁷

$$A_{hkl} = \sum_j f_j \cos 2\pi(hx_j + hy_j + lz_j)$$

$$B_{hkl} = \sum_j f_j \sin 2\pi(hx_j + hy_j + lz_j)$$

$$|F_{hkl}| = (A_{hkl}^2 + B_{hkl}^2)^{1/2}$$

$$\alpha_{hkl} = \tan^{-1}(B_{hkl} / A_{hkl}),$$

where f_j is the scattering factor for atom j.

Additional atomic positions could then be determined through the use of Fourier syntheses³⁴ of the form:

$$(XYZ) = \frac{2}{v} \sum_{h=0}^{\infty} \sum_{k=-\infty}^{\infty} \sum_{l=-\infty}^{\infty} |F_{hkl}| \cos 2\pi[(hx+ky+lz)-\alpha_{hkl}].$$

The positional coordinates of atoms in the trial structure were estimated from the Fourier generated electron density map using a FORTRAN computer program, BOOTHIT1, written in the course of this work. A description and listing of BOOTHIT1 is contained in Appendix A. Alternate structure factor calculations and Fourier syntheses were repeated until all nonhydrogen atoms were located.

In the case of a compound not containing a heavy atom but having a centrosymmetric space group, the direct method of symbolic addition was used. The FORTRAN computer programs, FAME-MAGIC-LINK-SYMPL, developed by E. B. Fleischer, R. B.

K. Dewar, and A.L. Stone^{35,36} were used to generate possible solutions to the phase problem. The programs first converted $|F_{obs}|$'s to normalized structure factors, E's, through the definitions:

$$(F_{absolute})^2 = \left(\frac{1}{K^2}\right) |F_{obs}|^2 e^{(T \sin \theta)/\lambda}$$

and

$$E^2 = (F_{absolute})^2 / \varepsilon \sum_i^N f_i^2$$

where the scale factor, K, and the overall temperature factor, T, were generated by a Wilson plot; where ε was a symmetry factor applied to reflections in special zones; and where f_i 's were the scattering factors for N atoms. The programs then assigned symbols representing the phases to six of the largest E's having the greatest number of interactions, i.e., for E_h and E_m there exists E_{h-m} . For such reflections the probability, p, that the phase of E_h is the same as E_{h-m} is given by:

$m=0$

$$p = 0.5 + 0.5 \tanh \left(\frac{\sigma_3}{\sigma_2^{1.5}} |E_h| \left| \sum_{m=0}^N E_m E_{h-m} \right| \right)$$

where

$$\sigma_n = \sum_{j=1}^N z_j^n$$

with N being the number of atoms in the unit cell and z_j being the atomic number of the jth atom. The programs, when given minimum acceptable probability criteria, iteratively assigned relative signs to the phase symbols. Combinations of these

signed phase symbols were finally used in conjunction with their structure factors to generate E-maps. The positional coordinates of most nonhydrogen atoms were determined from one of these E-maps. Structure factor calculations and Fourier syntheses were used to refine the atomic positions and, as in the heavy atom case, to locate any previously unfound nonhydrogen atoms of the trial structure.

The trial structure was refined by least-squares minimization³⁴ of the function:

$$\text{Residual} = \sum w(||F_{\text{obs}}| - |F_{\text{calc}}||)^2$$

where

$$\sqrt{w} = |F_{\text{obs}}|/|F_{\text{low}}| \quad \text{for } |F_{\text{obs}}| < |F_{\text{low}}|$$

$$\sqrt{w} = 1.0 \quad \text{for } |F_{\text{low}}| \leq |F_{\text{obs}}| \leq |F_{\text{high}}|$$

and

$$\sqrt{w} = |F_{\text{high}}|/|F_{\text{obs}}| \quad \text{for } |F_{\text{obs}}| > |F_{\text{high}}|$$

F_{low} and F_{high} are constants given in Table 4. Prior to refinement, an overall scale factor was chosen such that the sum of F_{obs} equaled the sum of F_{calc} . Isotropic temperature factors were used in the first three cycles of refinement and then anisotropic temperature factors of the form:

$$\exp[-(\beta_{11}h^2 + \beta_{22}k^2 + \beta_{33}\ell^2 + \beta_{12}hk + \beta_{13}h\ell + \beta_{23}k\ell)]$$

were used. The reliability index, R , was defined by:

$$R = \frac{\sum ||F_{\text{obs}}| - |F_{\text{calc}}||}{\sum |F_{\text{obs}}|}$$

Calculations were performed on an IBM 370/165 computer with programs written or modified by Dr. Gus J. Palenik, except where previously noted. The refinement of each structure is outlined in Table 5.

Schemes of Refinement

Table 5

Compound

R-index with all nonhydrogen atoms from Fourier synthesis

Refinement with isotropic thermal parameters

Refinement^a with anisotropic thermal parameters

	No. of cycles	R-index	No. of cycles	R-index
C ₁₂ Co(H ₂ dmg) (dmg) (clan) · 2H ₂ O	0.27	3	0.095	3
C ₁ ·Co(H ₂ dpg ₂) (clan) · C ₂ H ₅ OH	0.229	3	0.132	3
[Co(Hdmg) ₂ (clan) ₂]Cl	0.255	3	0.141	6
H ₂ dphphy(NO ₃) ₂ · 2H ₂ O	0.32	3	0.134	3*
[Ni ₂ Cl(H ₂ O) ₄ (dphphy)]Cl ₃ · 2H ₂ O	0.21	3	0.090	3
C ₄ (fph) ₄ Co(cp) (tpp)	0.26	3	0.137	9
C ₄ (fph) ₄ Rh(cp) (tpp)	0.168	3	0.105	9
				0.065

^aThe block-diagonal approximation to the full matrix was used except where marked with an asterisk.

Table 5 - extended

Compound	Refinement ^a with hydrogen atoms included isotropically but not refined			Refinement ^a with hydrogen atoms refined but not isotropically		
	No. of cycles	R-index	No. of R-index cycles		F _{low}	F _{high}
C ₂ Co(H ₂ dmg)(dmg)(clan)·2H ₂ O	3	0.052	6	0.047	18.0	49.0
C ₂ Co(H ₂ dpg ₂)(clan)·C ₂ H ₅ OH	3	0.087	9	0.075	55.0	145.0
[Co(Hdmg) ₂ (clan) ₂]Cl	3	0.038	9	0.033	4.5	12.0
H ₂ dphphy(NO ₃) ₂ ·2H ₂ O	-	-	6	0.050	8.0	22.0
[Ni ₂ Cl(H ₂ O) ₄ (dphphy)]Cl ₃ ·2H ₂ O	6	0.048	-	-	32.0	86.0
C ₄ (fph) ₄ Co(cp)(tpp)	-	-	-	-	17.5	35.0
C ₄ (fph) ₄ Rh(cp)(tpp)	-	-	-	-	17.5	35.0

CHAPTER 4

AN INVESTIGATION OF LIGAND-INDUCED PROTON SHIFT: THE CRYSTAL AND MOLECULAR STRUCTURES OF TRANS-CHLORO(DIMETHYLGlyOXIMATO)-(DIMETHYLGlyOXIME)(4-CHLOROANILINE)COBALT(III) DIHYDRATE, TRANS-CHLOROBIS(DIPHENYLGLYOXIMATO)(4-CHLOROANILINE)COBALT(III) ETHANOLATE, AND TRANS-BIS(DIMETHYLGlyOXIMATO)NIS (4-CHLOROANILINE)COBALT(III) CHLORIDE.

The stability of bis(dimethylglyoxime)metal complexes has long been known and their importance in both qualitative and quantitative analysis has been widely recognized.^{37,38} Metal complexes of Hdmg have been used to study the trans-effect³⁹ and the trans-influence^{40,41} of various ligands in octahedral complexes. Since the structural determination of the B₁₂ coenzyme the trans-bis(dimethylglyoxime)cobalt complexes have become of considerable interest.⁴²⁻⁴⁴ Schrauzer⁴² has stated that to be capable of mimicking B₁₂ a complex is required only to have a cobalt ion in the presence of a strong-binding planar ligand. Because Co(H₂dmg)₂ complexes successfully mimic the reactions of a cobalt ion in the corrin ring and because they are synthetically expedient, complexes of Co(H₂dmg)₂ have been investigated extensively in solution as models for B₁₂.⁴⁵

Until very recently there have been few structural data on Co(H₂dmg)₂ complexes.^{40,41,46-52} Except for the work of Palenik et al.⁴⁶ no structural investigation has been made of the interaction between the axial ligand and the equatorial Hdmg ligands. This interaction may be of considerable consequence.

Although sulfonamides are potent inhibitors of carbonic anhydrase they do not form strong coordination bonds with transition metal ions. Therefore, an interaction of the aromatic ring of the sulfonamide with the carbonic anhydrase protein has been proposed⁵³ to make a large contribution to the observed stability of the carbonic anhydrase-sulfonamide complex. Since a cobalt atom can replace the zinc atom in carbonic anhydrase with only a 50% decrease in activity, complexes of $\text{Co}(\text{H}_2\text{dmg}_2)$ may prove to be useful models for investigating the interaction of sulfonamides with carbonic anhydrase.

An apparent ligand-induced proton shift (LIPS) was observed⁴⁶ in $\text{ClCo}(\text{H}_2\text{dmg}_2)(\text{sulfa})$ which should be formulated $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{sulfa})$. To investigate further the LIPS phenomena and to examine interligand interactions within this type of complex the determination of the structures of $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{clan})$, $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}$, and $\text{ClCo}(\text{H}_2\text{dmg}_2)(\text{clan})$ was undertaken.

Structure Solution and Refinement
for $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{clan}) \cdot 2\text{H}_2\text{O}$

The heavy atom method was used with the positions of the cobalt atom and of the ionic chloride ligand estimated from a sharpened Patterson function. The magnitude of the Patterson function for the Co to Cl vectors was of the same order as that for the Co to Co vector. The positions of the heavy atoms, therefore, appeared ambiguous and several combinations were used in Fourier syntheses to determine their actual lo-

cations. Successive Fourier syntheses then revealed the locations of all nonhydrogen atoms in the compound. Three cycles of full-matrix least-squares refinement with individual isotropic thermal parameters and then three cycles of least-squares refinement using the block approximation with individual anisotropic thermal parameters reduced R to 0.066. A difference Fourier synthesis then indicated the absence of additional nonhydrogen atoms and revealed the positions of all hydrogen atoms. An outline of the refinement is given in Table 5. The refinement was terminated after the parameter shifts for the nonhydrogen atoms were less than one-tenth of their corresponding estimated standard deviations.

The scattering factors for cobalt, chlorine, oxygen, nitrogen, and carbon were from Hanson *et al.*²⁹ while those for hydrogen were from Stewart *et al.*³⁰ A list of the observed and calculated structure factors has been published and is available.⁴⁶ The final positional and thermal parameters are given in Tables 6 and 7.

Structure Solution and Refinement
for ClCo(H₂dpg₂) (clan) · C₂H₅OH

The nonstandard space group P2₁/n was chosen since the standard P2₁/c space group would require a very large value for β . The position of the cobalt atom was estimated from a sharpened Patterson function. The location of atoms and the refinement proceeded as in the case of ClCo(H₂dmg) (dmg) (clan) · 2H₂O. Two atoms, O(S1) and C(S1), of an apparent solvent mole-

Table 6
Final Atomic Parameters of Nonhydrogen Atoms for ClCo(H₂dmg)-
(dmg)(clan)^a

Atom	x	y	z	β_{11}	β_{22}
Co	19148(12)	36142(8)	21611(6)	1287(16)	454(7)
Cl(1)	-1353(2)	2445(1)	1882(1)	148(3)	65(2)
Cl(2)	8142(4)	10300(2)	3785(2)	512(9)	67(2)
O(11)	1633(7)	4573(4)	534(3)	305(13)	95(5)
O(12)	1742(6)	4621(4)	4260(3)	304(13)	89(5)
O(21)	2285(7)	2611(4)	3796(3)	314(13)	104(5)
O(22)	1944(7)	2468(4)	17(3)	298(13)	114(6)
N(1)	4821(7)	4589(5)	2375(3)	157(12)	93(6)
N(11)	1600(7)	4753(5)	1536(3)	179(12)	75(6)
N(12)	1693(7)	4792(4)	3347(3)	156(12)	58(5)
N(21)	2289(7)	2459(5)	2788(3)	191(13)	72(5)
N(22)	2144(7)	2398(5)	980(3)	168(12)	74(5)
C(11)	1403(8)	5758(6)	2145(5)	170(15)	67(7)
C(12)	1443(8)	5772(6)	3208(5)	168(15)	59(6)
C(13)	1247(10)	6794(6)	1785(5)	268(19)	85(8)
C(14)	1310(10)	6815(7)	4067(5)	281(20)	100(8)
C(21)	2584(9)	1516(6)	2201(17)	214(17)	66(7)
C(22)	2509(9)	1475(6)	1124(5)	164(15)	65(7)
C(23)	3010(14)	584(8)	2593(7)	530(32)	112(10)
C(24)	2779(12)	465(7)	277(6)	410(27)	121(10)
C(1)	5676(7)	5999(5)	2722(4)	96(13)	70(6)
C(2)	6105(9)	6655(6)	3753(5)	198(16)	71(7)
C(3)	6866(9)	7971(6)	4082(5)	221(17)	91(8)
C(4)	7201(10)	8629(6)	3382(6)	195(17)	73(7)
C(5)	6809(10)	7972(7)	2340(5)	237(19)	99(8)
C(6)	6047(9)	6644(6)	2005(5)	191(16)	82(7)
O(w1)	6682(7)	3785(5)	646(4)	264(13)	148(7)
O(w2)	6802(7)	3830(5)	3690(3)	276(13)	166(7)

^aAll values are $\times 10^4$ except for Co which are $\times 10^5$. The estimated standard deviations are given in parentheses. The temperature factors are of the form: $\exp[-(\beta_{11}h^2 + \beta_{22}k^2 + \beta_{33}\ell^2 + \beta_{12}hk + \beta_{13}h\ell + \beta_{23}k\ell)]$.

Table 6 - extended

β_{33}	β_{12}	β_{13}	β_{23}
276(4)	822(17)	219(13)	157(9)
53(1)	78(4)	25(3)	14(2)
138(2)	65(7)	46(7)	46(4)
36(3)	168(14)	7(9)	44(6)
32(3)	167(13)	70(9)	33(6)
41(3)	199(14)	74(10)	70(6)
30(3)	199(14)	31(9)	25(6)
37(3)	123(14)	29(10)	50(7)
35(3)	108(14)	24(10)	26(7)
35(3)	71(13)	30(9)	19(6)
41(3)	123(14)	41(10)	39(7)
35(3)	99(13)	21(10)	12(7)
53(4)	101(17)	25(13)	41(9)
51(4)	73(16)	30(13)	14(8)
76(5)	190(21)	-5(16)	55(10)
62(5)	214(22)	50(16)	-6(10)
56(4)	134(18)	47(14)	44(9)
60(5)	110(17)	52(13)	24(9)
102(7)	347(31)	148(24)	106(14)
73(6)	301(28)	87(20)	11(12)
44(4)	72(15)	23(11)	44(8)
47(4)	89(17)	4(13)	28(9)
42(4)	95(19)	4(14)	8(9)
82(6)	61(18)	23(16)	39(10)
69(5)	99(20)	83(16)	95(11)
52(4)	104(18)	56(13)	52(9)
80(4)	241(16)	102(11)	100(8)
53(3)	295(16)	20(10)	22(7)

Table 7

Final Parameters for the Hydrogen Atoms for ClCo(H₂dmg) (dmg) - (clan)^a

Atom [Bonded to]	Distance	x	y	z	B
H(B1) [O(22)]	1.16(8)	153(10)	335(7)	17(5)	6.3(1.8)
H(B2) [O(21)]	1.13(8)	184(10)	345(7)	402(5)	6.7(1.8)
H(2) [C(2)]	0.89(5)	591(7)	621(5)	420(4)	2.0(1.1)
H(3) [C(3)]	0.99(7)	726(10)	858(7)	478(5)	6.7(1.9)
H(5) [C(5)]	0.90(7)	709(10)	838(7)	187(5)	5.8(1.7)
H(6) [C(6)]	1.01(5)	568(7)	609(5)	127(4)	2.0(1.1)
H(7) [N(1)]	1.03(7)	522(10)	434(6)	166(5)	6.1(1.7)
H(8) [N(1)]	0.83(6)	518(9)	428(6)	278(5)	4.5(1.5)
H(11) [C(13)]	0.96(8)	243(11)	753(7)	212(6)	8.2(2.1)
H(12) [C(13)]	0.79(9)	37(11)	692(7)	203(6)	8.5(2.2)
H(13) [C(13)]	1.00(1)	102(10)	663(7)	103(5)	7.1(1.9)
H(14) [C(14)]	1.02(7)	46(9)	639(6)	453(5)	5.8(1.7)
H(15) [C(14)]	0.78(7)	62(9)	708(6)	388(5)	4.9(1.5)
H(16) [C(14)]	0.87(8)	228(11)	734(8)	456(6)	8.6(2.2)
H(21) [C(23)]	0.86(10)	414(12)	59(8)	248(6)	9.6(2.4)
H(22) [C(23)]	0.92(9)	239(12)	-21(8)	211(6)	9.3(2.4)
H(23) [C(23)]	0.97(9)	266(12)	52(8)	326(7)	9.8(2.5)
H(24) [C(24)]	1.03(9)	242(12)	44(8)	-46(7)	9.4(2.4)
H(25) [C(24)]	0.84(10)	396(12)	58(8)	29(6)	9.1(2.3)
H(26) [C(24)]	1.00(7)	210(10)	-38(7)	40(5)	5.9(1.7)
H(w1) [O(w1)]	0.70(8)	659(11)	319(7)	69(6)	7.2(2.0)
H(w1') [O(w1)]	0.80(13)	771(16)	438(11)	72(9)	15.2(3.7)
H(w2) [O(w2)]	0.79(7)	736(10)	420(7)	425(5)	6.6(1.8)
H(w2') [O(w2)]	0.71(7)	747(10)	371(6)	337(5)	6.0(1.8)

^aThe hydrogen atom is given followed by the atom to which it is bonded in brackets, the corresponding bond distance (in Å), the positional parameters with estimated standard deviations ($\times 10^{+3}$), and the isotropic thermal parameter (in Å²).

molecule were located before refinement. The scheme of the refinement is outlined in Table 5.

Although the compound was crystallized from ethanol, difference Fourier syntheses at various stages of refinement failed to indicate the position of an additional atom in the solvent molecule. Because a large region of relative high electron density existing near C(f1) could be indicative of an atom with high disorder and because ethanol was the solvent, a molecule of ethanol was assumed to be present for the purposes of determining the formula, molecular weight, and density.

The cobalt, chlorine, oxygen, nitrogen, and carbon scattering factors were taken from Hanson *et al.*²⁹ and those for hydrogen from Stewart *et al.*³⁰ Table B-1 is a list of observed and calculated structure factors for $\text{ClCo}(\text{H}_2\text{dmg})_2$ - (clan) . The final positional and thermal parameters are shown in Tables 8 and 9.

Structure Solution and Refinement
for $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}$

With one molecule per unit cell in the centric $P\bar{1}$ space group the cobalt atom and the chloride anion were required to lie on centers of symmetry. The sharpened Patterson function was in agreement with the chloride ion at $0\%0$ when the cobalt atom is placed at 000 . The remaining atoms were located in a similar manner as in $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{clan})$. An outline of the refinement is given in Table 5.

Table 8

The Final Atomic Parameters for the Nonhydrogen Atoms of ClCo
 $(H_2^{dpq})_2(\text{clan})^a$

Atom	x	y	z	β_{11}	β_{22}
Co	3339(1)	3017(2)	2961(1)	33(1)	58(1)
Cl(1)	3101(2)	4797(3)	3049(2)	45(2)	68(3)
Cl(2)	6313(4)	60(5)	1153(3)	96(4)	151(6)
O(11)	3953(5)	3496(7)	1583(4)	47(5)	82(9)
O(12)	4349(5)	2916(10)	4348(4)	41(5)	210(13)
O(21)	2763(6)	2628(8)	4340(4)	54(5)	130(11)
O(22)	2334(5)	3078(8)	1587(4)	42(5)	102(9)
N(1)	3495(6)	1461(9)	2903(6)	29(6)	97(12)
N(11)	4172(6)	3351(7)	2306(4)	41(6)	41(8)
N(12)	4368(7)	3168(9)	3660(5)	56(7)	78(11)
N(21)	2534(7)	2738(9)	3635(5)	50(6)	85(11)
N(22)	2312(6)	2882(9)	2294(4)	39(6)	62(9)
C(11)	4970(8)	3533(10)	2591(6)	24(7)	88(14)
C(12)	5080(8)	3361(11)	3373(6)	44(8)	101(15)
C(13)	5698(8)	3915(12)	2191(6)	38(8)	70(12)
C(14)	5956(9)	3431(11)	3862(7)	42(8)	82(14)
C(21)	1706(8)	2557(10)	3364(6)	46(8)	58(12)
C(22)	1575(7)	2672(10)	2562(5)	28(6)	66(12)
C(23)	1055(8)	2220(10)	3832(6)	54(8)	50(12)
C(24)	709(8)	2583(10)	2146(6)	52(8)	46(12)
C(1)	4167(8)	1048(11)	2436(6)	54(9)	46(11)
C(2)	5012(9)	858(11)	2798(7)	65(10)	65(14)
C(3)	5687(9)	544(11)	2387(8)	53(9)	62(13)
C(4)	5487(9)	448(11)	1654(8)	69(10)	53(13)
C(5)	4661(10)	592(13)	1341(7)	74(10)	123(18)
C(6)	3990(8)	883(11)	1733(7)	45(8)	67(13)
C(1A)	5975(8)	3387(11)	1614(7)	35(8)	92(15)
C(2A)	6642(9)	3760(13)	1251(7)	54(9)	111(16)
C(3A)	7077(8)	4683(12)	1485(7)	36(8)	97(15)
C(4A)	6831(9)	5248(13)	2052(8)	46(9)	108(15)
C(5A)	6157(9)	4877(12)	2413(7)	69(10)	93(15)

Table 8 - extended

β_{33}	β_{12}	β_{13}	β_{23}
19(0)	-7(3)	2(1)	-4(2)
40(1)	8(5)	11(3)	-13(4)
100(3)	71(9)	96(5)	-15(7)
21(3)	12(11)	6(6)	26(8)
16(3)	-17(16)	-8(6)	7(13)
20(3)	-35(13)	3(6)	8(9)
21(3)	-18(13)	-4(5)	12(10)
34(4)	-9(13)	7(8)	-15(12)
13(3)	23(11)	-6(6)	14(8)
26(4)	7(15)	-1(8)	7(12)
20(3)	-15(14)	6(7)	-21(11)
16(3)	-12(14)	13(6)	6(11)
23(4)	5(16)	16(9)	28(13)
18(4)	-5(18)	7(9)	0(13)
23(4)	20(17)	20(9)	23(14)
28(5)	-11(17)	-8(10)	5(13)
23(4)	4(15)	12(9)	-2(12)
15(4)	24(15)	-4(8)	7(11)
17(4)	-43(17)	3(3)	-9(12)
28(5)	26(16)	-1(10)	30(12)
30(5)	-5(18)	-18(10)	20(14)
36(6)	-17(19)	-7(11)	15(15)
45(6)	45(18)	4(12)	2(15)
46(6)	58(19)	40(12)	20(15)
37(6)	19(23)	44(12)	70(17)
33(5)	-69(17)	9(10)	-29(14)
39(5)	5(17)	31(10)	4(15)
29(5)	10(21)	22(11)	2(16)
45(6)	-12(19)	9(11)	46(16)
43(6)	-28(21)	30(11)	13(10)
32(5)	7(22)	11(11)	2(17)

Table 8 - continued

Atom	x	y	z	β_{11}	β_{22}
C(1B)	6675(9)	2815(14)	3687(7)	61(9)	113(16)
C(2B)	7444(9)	2801(13)	4142(7)	59(9)	89(15)
C(3B)	7498(9)	3363(15)	4781(7)	56(10)	189(23)
C(4B)	6828(9)	4051(16)	4937(7)	59(10)	207(22)
C(5B)	6047(10)	4094(14)	4476(7)	73(11)	148(19)
C(1C)	664(9)	1226(12)	3750(7)	66(10)	73(14)
C(2C)	20(9)	872(12)	4199(7)	66(10)	96(16)
C(3C)	-212(9)	1576(14)	4700(8)	32(8)	183(23)
C(4C)	184(9)	2563(12)	4813(7)	61(10)	112(17)
C(5C)	826(8)	2872(11)	4368(6)	60(8)	41(11)
C(1D)	593(9)	2121(12)	1437(6)	54(8)	83(14)
C(2D)	-224(9)	1992(14)	1046(7)	56(9)	116(16)
C(3D)	-951(9)	2410(12)	1345(7)	50(9)	105(17)
C(4D)	-888(8)	2847(12)	2044(7)	42(8)	77(14)
C(5D)	-69(8)	2975(12)	2447(6)	30(7)	63(12)
O(S1)	1418(9)	4904(10)	944(5)	190(13)	136(13)
C(S1)	1450(26)	4854(22)	182(12)	512(49)	196(30)

^aAll values are $\times 10^4$. The estimated standard deviations are given in parentheses. The temperature factors are of the form: $\exp[-(\beta_{11}h^2 + \beta_{22}k^2 + \beta_{33}\ell^2 + \beta_{12}hk + \beta_{13}h\ell + \beta_{23}k\ell)]$.

Table 8 - continued -- extended

β_{33}	β_{12}	β_{13}	β_{23}
39(6)	-11(22)	-24(11)	13(18)
39(5)	-24(21)	-4(11)	24(17)
32(5)	-59(24)	13(11)	-5(18)
18(5)	-47(26)	0(10)	-39(19)
28(5)	-90(25)	7(11)	0(18)
28(5)	-27(19)	-3(11)	-20(15)
28(5)	-24(21)	7(11)	13(16)
44(6)	-21(21)	-26(11)	-5(19)
26(5)	16(19)	26(10)	-10(14)
28(4)	-31(19)	-7(9)	8(14)
28(5)	-4(20)	7(10)	-16(16)
34(5)	13(23)	-34(10)	-39(19)
42(6)	11(19)	-31(11)	50(16)
51(6)	-13(20)	-8(11)	6(18)
40(5)	-30(18)	17(9)	7(16)
39(5)	67(23)	45(12)	-23(14)
59(11)	224(71)	194(38)	45(34)

Table 9
Final Parameters for Hydrogen Atoms for $\text{ClCo}(\text{H}_2\text{dpq}_2)(\text{clan})^{\text{a}}$

Atom [Bonded to]	Distance	x	y	z	B
H(B1)		303(9)	344(12)	153(8)	11.6(5.1)
H(B2)		352(7)	277(9)	439(5)	4.3(2.7)
H(2) [C(2)]	0.84(10)	514(7)	87(9)	325(5)	4.3(2.8)
H(3) [C(3)]	0.95(14)	627(9)	47(12)	260(7)	9.8(4.5)
H(5) [C(5)]	0.80(10)	445(6)	54(8)	93(5)	4.1(2.8)
H(6) [C(6)]	1.00(10)	336(7)	84(9)	152(5)	4.7(3.0)
H(7) [N(1)]	0.93(11)	305(7)	107(10)	264(6)	5.9(3.3)
H(8) [N(1)]	1.02(15)	382(9)	103(13)	332(8)	11.7(4.6)
H(1A) [C(1A)]	1.05(12)	558(7)	278(10)	135(6)	6.0(3.2)
H(2A) [C(2A)]	0.94(10)	680(6)	329(8)	89(5)	4.0(2.7)
H(3A) [C(3A)]	1.18(15)	759(9)	514(13)	117(7)	10.5(4.8)
H(4A) [C(4A)]	1.12(12)	711(7)	604(10)	227(6)	5.7(3.3)
H(5A) [C(5A)]	0.86(9)	601(6)	525(8)	227(5)	2.4(2.4)
H(1B) [C(1B)]	1.07(9)	654(6)	236(8)	320(5)	3.0(2.6)
H(2B) [C(2B)]	1.12(19)	796(11)	224(16)	398(9)	14.6(6.6)
H(3B) [C(3B)]	0.68(13)	792(9)	351(11)	487(7)	9.1(4.4)
H(4B) [C(4B)]	0.74(10)	690(7)	403(9)	533(6)	5.3(3.0)
H(5B) [C(5B)]	0.55(14)	586(9)	406(12)	471(7)	11.2(4.6)
H(1C) [C(1C)]	0.81(13)	76(8)	69(10)	352(7)	8.3(4.1)
H(2C) [C(2C)]	1.04(19)	3(12)	7(15)	403(9)	15.8(6.3)
H(3C) [C(3C)]	0.96(12)	-63(8)	140(10)	504(7)	8.2(4.0)
H(4C) [C(4C)]	0.92(12)	4(8)	305(10)	515(6)	6.0(3.0)
H(5C) [C(5C)]	1.00(8)	106(5)	362(6)	447(4)	0.5(1.9)
H(1D) [C(1D)]	1.05(9)	116(6)	175(7)	128(5)	3.0(2.4)
H(2D) [C(2D)]	1.10(9)	-41(6)	160(8)	52(5)	2.8(2.6)
H(3D) [C(3D)]	0.97(14)	-145(9)	219(12)	101(7)	10.1(4.4)
H(4D) [C(4D)]	1.04(10)	-136(6)	343(8)	212(5)	3.8(2.7)
H(5D) [C(5D)]	0.70(8)	25(5)	316(7)	224(4)	1.5(2.0)

^aThe hydrogen atom is given followed by the atom to which it is bonded in brackets, the corresponding bond distance (\AA), the positional parameters with estimated standard deviations ($\times 10^{+3}$), and the isotropic thermal parameters (\AA^2).

The scattering factors for cobalt, oxygen, nitrogen, and carbon were from Hanson *et al.*,²⁹ those for hydrogen were from Stewart *et al.*,³⁰ and those for chlorine were from Doyle and Turner.³¹ The observed and calculated structure factors are given in Table B-2. Lists of final positional and thermal parameters may be found in Tables 10 and 11.

Results and Discussion

The atomic numbering and thermal ellipsoids of $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{clan})$, $\text{ClCo}(\text{H}_2\text{dpg}_2)(\text{clan})$, and $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}$ are shown in ORTEP⁵⁴ drawings in Figures 1, 2, and 3, respectively. The individual bond distances for these three compounds together with those of two related compounds, $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{sulfa})$ ⁴⁶ and $[\text{Co}(\text{Hdmg})_2(\text{an})_2]\text{Cl}$,⁵² are tabulated in Table 12. The corresponding bond angles are given in Table 13.

In each case the two dmg or dpg groups are approximately planar as demonstrated by the deviations from least-squares planes in Tables 14-16. The dmg groups of each complex are linked by two intramolecular hydrogen bonds (see Table 17).

As in the case of $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{sulfa})$ ⁴⁶ the hydrogen bridges between the dmg groups in $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{clan})$ were found to be asymmetrical with both hydrogen atoms bonded to the same dmg ligand. The O(21)-H(B2) and O(22)-H(B1) distances of 1.13(8) and 1.16(8) Å, respectively, compared to the O(12)…H(B2) and O(11)…H(B1) distances of 1.36(8) and 1.37(8) Å indicate the formulation H_2dmg and dmg for the two

Table 10
 The Final Atomic Parameters for Nonhydrogen Atoms of $[\text{Co}(\text{Hdmg})_2(\text{clan})_2\text{Cl}]$.^a

Atom	x	y	z	β_{11}	β_{22}
Co	0(0)	0(0)	0(0)	817(11)	628(5)
Cl(1)	0(0)	50000(0)	0(0)	2653(30)	664(12)
Cl(2)	32052(23)	26440(24)	55881(8)	5254(48)	6743(52)
O(11)	4450(2)	1508(2)	573(1)	97(4)	126(3)
O(12)	-3514(3)	531(2)	-1814(1)	125(5)	144(3)
N(1)	-846(3)	1678(2)	816(2)	116(5)	77(3)
N(11)	2490(3)	1399(2)	-160(2)	105(5)	83(3)
N(12)	-1339(3)	923(2)	-1288(1)	118(5)	91(3)
C(1)	97(4)	1928(3)	1978(2)	148(7)	90(3)
C(2)	2160(4)	2812(3)	2368(2)	195(8)	136(4)
C(3)	3103(5)	3034(4)	3474(3)	224(9)	231(6)
C(4)	1982(6)	2386(5)	4181(3)	317(11)	311(8)
C(5)	-77(6)	1504(5)	3818(3)	320(11)	300(8)
C(6)	-1012(4)	1282(4)	2704(2)	193(8)	185(5)
C(11)	2176(4)	2274(3)	-990(2)	159(7)	76(3)
C(12)	-124(4)	1982(3)	-1665(2)	185(7)	87(3)
C(13)	3887(4)	3459(3)	-1239(2)	216(8)	116(4)
C(14)	-937(5)	2830(4)	-2658(3)	289(10)	169(5)

^aAll values are $\times 10^4$ except those for Co, Cl(1) and Cl(2) which are $\times 10^5$. The estimated standard deviations are given in parentheses. The temperature factors are of the form: $\exp[-(\beta_{11}h^2 + \beta_{22}k^2 + \beta_{33}\ell^2 + \beta_{12}hk + \beta_{13}h\ell + \beta_{23}k\ell)]$.

Table 10 - extended

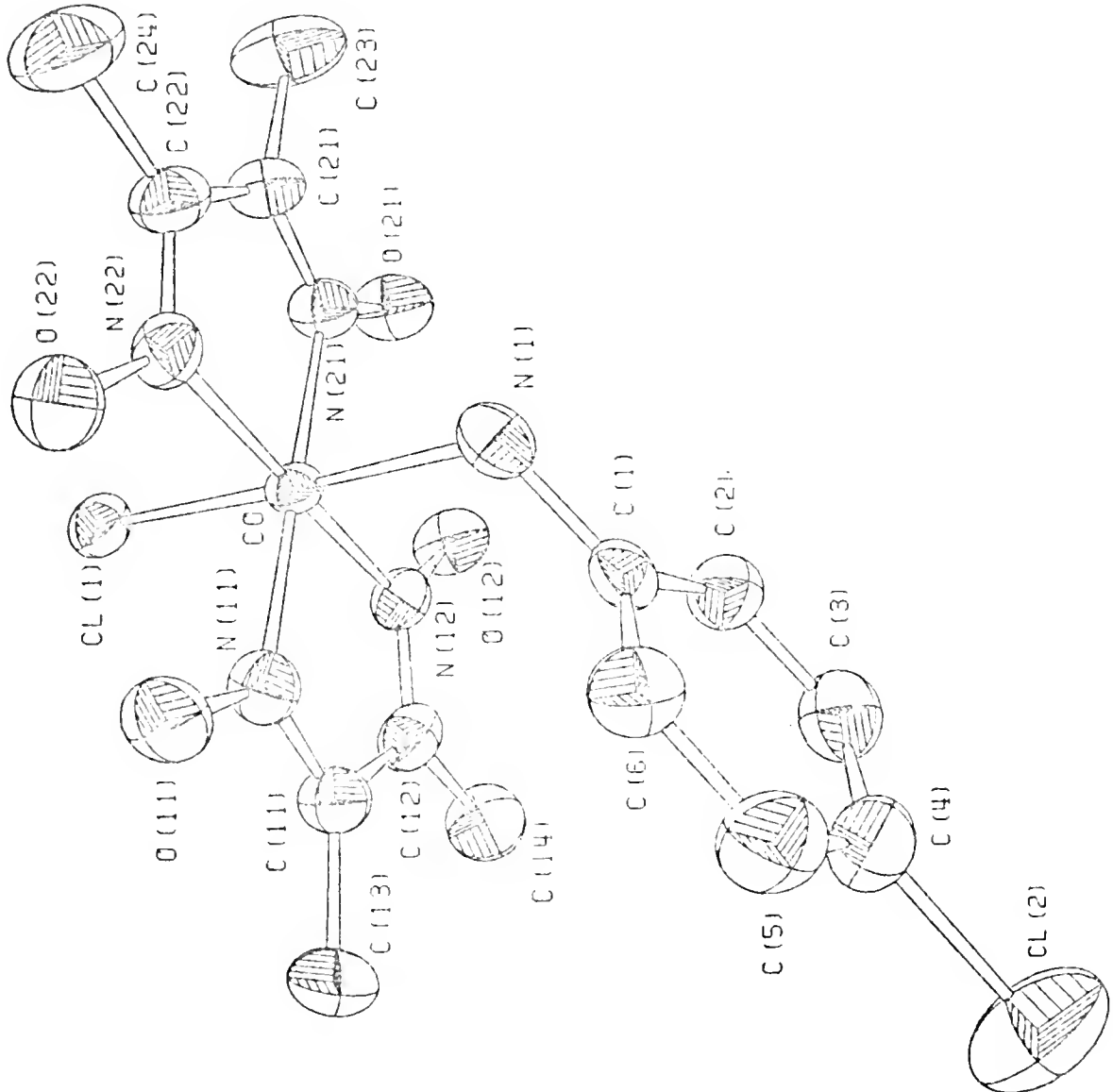
β_{33}	β_{12}	β_{13}	β_{23}
332(3)	226(12)	169(9)	-43(6)
873(8)	451(29)	1194(26)	131(15)
482(7)	-1987(79)	-431(28)	-982(29)
57(1)	9(5)	11(4)	-12(3)
57(1)	40(6)	-20(4)	15(3)
45(1)	28(6)	30(4)	-9(3)
47(1)	26(6)	30(4)	-21(3)
41(1)	53(6)	7(4)	-7(3)
47(2)	47(8)	27(5)	-24(4)
63(2)	-25(9)	59(6)	-42(5)
72(2)	-85(12)	-3(8)	-93(6)
49(2)	2(15)	-11(3)	-59(7)
51(2)	-22(15)	64(8)	1(7)
52(2)	-11(10)	36(6)	-12(5)
52(2)	42(7)	73(6)	-4(4)
46(2)	68(8)	56(6)	6(4)
77(2)	12(9)	109(7)	16(5)
68(2)	68(11)	47(8)	75(6)

Table 11
Final Parameters for Hydrogen Atoms for $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}^{\text{a}}$

Atom [Bonded to]	Distance	X	Y	Z	B
H(B1) [O(12)]	1.07(3)	-408(8)	-35(4)	-133(3)	5.5(0.8)
H(2) [C(2)]	0.85(3)	280(4)	321(3)	190(2)	3.5(0.6)
H(3) [C(3)]	0.91(4)	447(6)	361(4)	366(3)	6.0(0.8)
H(5) [C(5)]	0.98(4)	-92(6)	105(4)	431(3)	6.6(0.9)
H(6) [C(6)]	0.96(3)	-248(5)	73(3)	244(2)	3.9(0.6)
H(7) [N(1)]	0.88(2)	-299(4)	146(3)	64(2)	2.1(0.5)
H(8) [N(1)]	0.94(2)	-52(4)	262(3)	49(2)	2.7(0.5)
H(11) [C(13)]	0.90(4)	349(6)	440(4)	-131(3)	5.7(0.8)
H(12) [C(13)]	0.89(4)	417(7)	315(5)	-185(4)	9.0(1.2)
H(13) [C(13)]	0.91(4)	513(6)	353(5)	-67(3)	7.3(1.0)
H(14) [C(14)]	0.86(4)	-181(7)	217(5)	-314(3)	9.0(1.2)
H(15) [C(14)]	0.80(5)	-14(8)	300(6)	-274(4)	10.0(1.3)
H(16) [C(14)]	1.01(5)	-213(8)	337(6)	-252(4)	11.0(1.4)

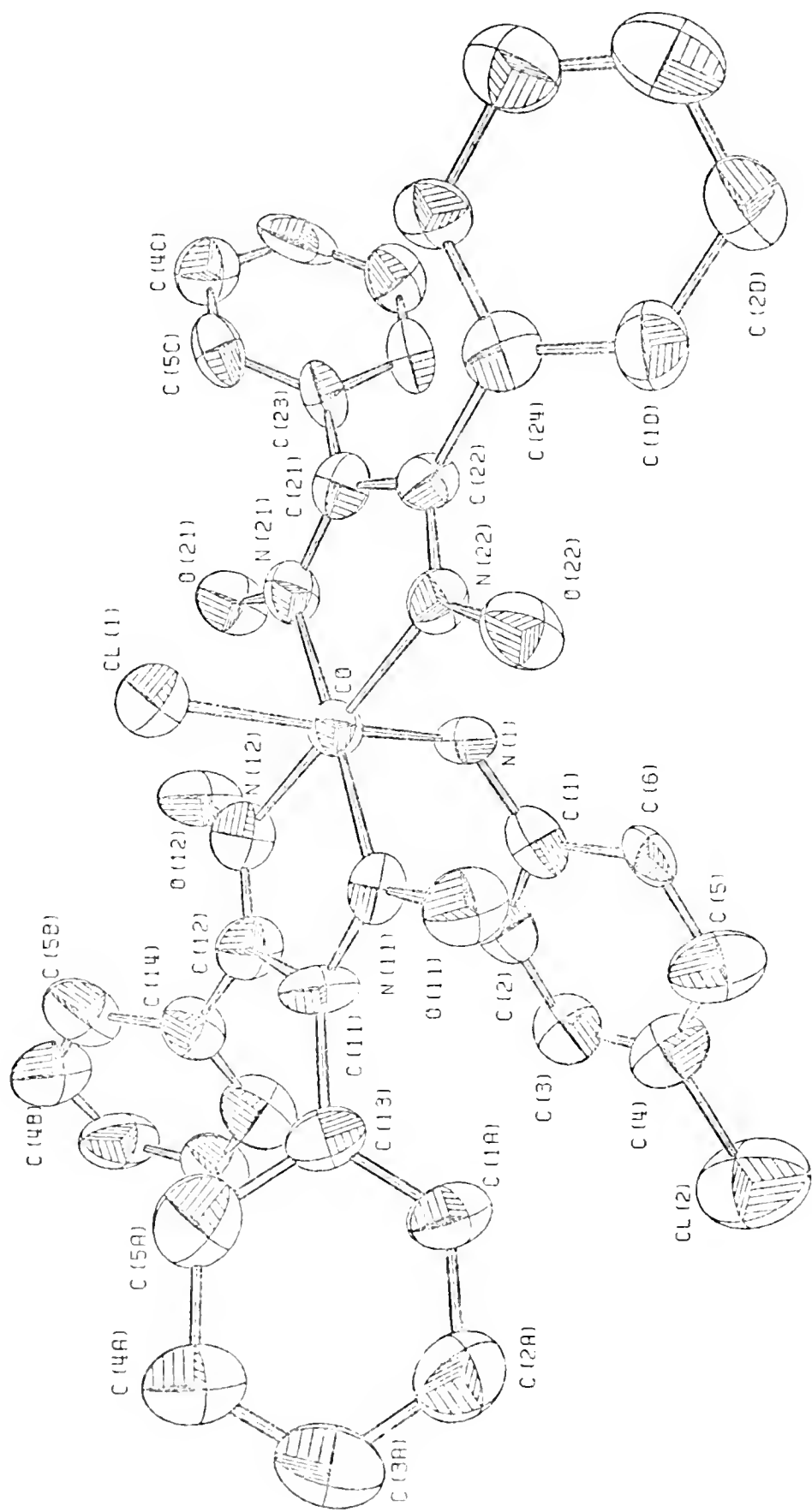
^aThe hydrogen atom is given followed by the atom to which it is bonded in brackets, the corresponding bond distance (\AA), the positional parameters with estimated standard deviations ($\times 10^{+3}$), and the isotropic thermal parameters (\AA^2).

Figure 1
An ORTEP drawing of $\text{C}_2\text{Co}(\text{H}_2\text{dmg})(\text{dmg})(\text{Clan}) \cdot 2\text{H}_2\text{O}$ showing the atomic numbering and thermal ellipsoids. The hydrogen atoms and water molecules have been omitted for clarity.



An Ortep drawing of $\text{C}_6\text{CO}(\text{CH}_2\text{Cl}_2)\cdot\text{C}_2\text{H}_5\text{OH}$ showing the atomic numbering and thermal ellipsoids. The hydrogen atoms and $\text{C}_2\text{H}_5\text{OH}$ molecule have been omitted.

Figure 2



An ORTEP drawing of $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}_2$ showing the atomic numbering and thermal ellipsoids. The hydrogen atoms have been omitted.

Figure 3

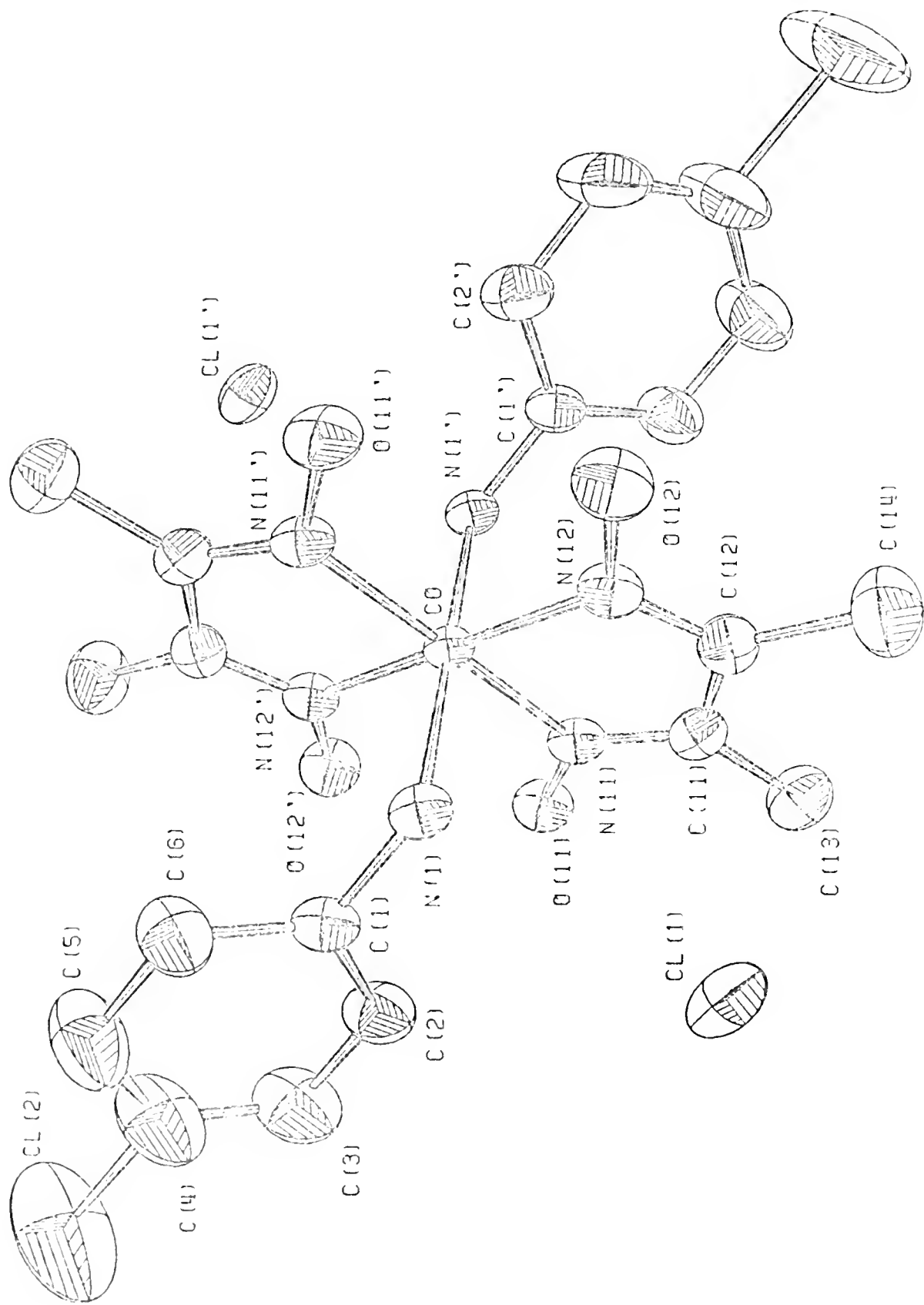


Table 12
Selected Interatomic Distances (\AA) in Some Cobaloxime
Complexes with Their Estimated Standard Deviations.^a

	$\text{C}_\ell\text{Co}(\text{H}_2\text{dmg})_2(\text{clan})$	$\text{C}_\ell\text{Co}(\text{H}_2\text{dpq})_2(\text{clan})$
Co-N(1)	1.999(6)	1.946(11)
Co-N(11)	1.872(5)	1.908(9)
Co-N(12)	1.884(5)	1.935(11)
N(11)-O(11)	1.337(6)	1.356(11)
N(12)-O(12)	1.329(6)	1.316(12)
N(11)-C(11)	1.311(8)	1.298(15)
N(12)-C(12)	1.308(8)	1.292(16)
C(11)-C(12)	1.457(9)	1.455(16)
C(11)-C(13)	1.488(10)	1.487(17)
C(12)-C(14)	1.487(10)	1.536(18)
O(11)...O(22)	2.497(7)	2.540(11)
O(11)-H(B)	1.37(8)	1.41(14)
O(12)-H(B)	1.36(8)	1.30(10)
Co-C ℓ (1)	2.257(2)	2.244(4)
Co-N(21)	1.908(5)	1.887(10)
Co-N(22)	1.906(5)	1.897(9)
N(21)-O(21)	1.348(6)	1.321(12)
N(22)-O(22)	1.359(6)	1.337(11)
N(21)-C(21)	1.280(8)	1.331(16)
N(22)-C(22)	1.288(8)	1.313(14)
C(21)-C(22)	1.468(9)	1.483(15)
C(21)-C(23)	1.486(11)	1.457(17)
C(22)-C(24)	1.498(11)	1.464(17)
O(12)...O(21)	2.479(7)	2.460(12)
O(21)-H(8)	1.13(8)	1.16(10)
O(22)-H(B)	1.16(8)	1.17(15)

*Distance given is for O(11)...O(12') \equiv O(12)...O(11')

^aValues for $[\text{Co}(\text{H}_2\text{dmg})_2(\text{an})_2\text{C}\ell]$ are listed with atomic numbering corresponding to the compounds of this work.

Table 12 - extended

$\text{ClCo}(\text{H}_2\text{dmg}_2)(\text{sulfa})^{46}$	$[\text{Co}(\text{H}_2\text{dmg}_2)(\text{clan})_2]\text{Cl}$	$[\text{Co}(\text{H}_2\text{dmg}_2)(\text{an})_2]\text{Cl}^{52}$
2.023(8)	2.003(2)	2.001(5)
1.870(8)	1.906(2)	1.885(6)
1.884(8)	1.889(2)	1.889(5)
1.323(11)	1.340(3)	1.353(6)
1.344(11)	1.362(3)	1.333(6)
1.289(14)	1.299(3)	1.286(10)
1.293(13)	1.290(3)	1.303(10)
1.494(16)	1.477(4)	1.463(7)
1.532(17)	1.483(4)	1.482(12)
1.488(16)	1.485(4)	1.476(11)
2.507(11)	2.495(3)*	2.491(8)*
1.50	1.44(3)	1.29
1.60	1.07(3)	1.21
2.235(3)		
1.905(8)		
1.896(8)		
1.326(10)		
1.338(11)		
1.292(12)		
1.290(14)		
1.447(17)		
1.494(17)		
1.488(16)		
2.479(11)		
0.90		
1.04		

Table 13
Selected Intramolecular Angles ($^{\circ}$) in Some Cobaloxime Complexes
with Their Estimated Standard Deviations.^a

	$\text{ClCo}(\text{H}_2\text{dmg}_2)$ (clan)	$\text{ClCo}(\text{H}_2\text{dpq}_2)$ (clan)
N(1)-Co-N(11)	90.5(2)	94.8(4)
N(1)-Co-N(12)	91.5(2)	92.1(4)
N(1)-Co-N(21)	88.4(2)	87.1(4)
N(1)-Co-N(22)	88.6(2)	88.6(4)
N(11)-Co-N(12)	82.6(2)	81.3(4)
N(11)-Co-N(22)	98.8(2)	100.0(4)
N(11)-Co-N(21)	178.8(2)	177.5(4)
N(12)-Co-N(21)	98.1(2)	97.0(4)
N(12)-Co-N(22)	178.6(2)	178.5(4)
N(21)-Co-N(22)	80.6(2)	81.7(4)
Cl(1)-Co-N(11)	90.6(2)	87.7(3)
Cl(1)-Co-N(12)	90.6(2)	89.1(3)
Cl(1)-Co-N(21)	90.5(2)	90.4(3)
Cl(1)-Co-N(22)	89.4(2)	90.2(3)
Cl(1)-Co-N(1)	177.8(2)	177.4(3)
Co-N(1)-C(1)	119.7(4)	118.6(8)
Co-N(11)-O(11)	121.9(4)	123.3(7)
Co-N(12)-O(12)	122.2(4)	121.2(8)
Co-N(21)-O(21)	123.2(4)	123.5(8)
Co-N(22)-O(22)	123.3(4)	120.7(7)
Co-N(11)-C(11)	116.0(4)	116.7(8)
Co-N(12)-C(12)	115.6(4)	114.1(9)
Co-N(21)-C(21)	116.6(4)	116.8(8)
Co-N(22)-C(22)	117.0(4)	117.4(8)
O(11)-N(11)-C(11)	122.1(5)	119.7(9)
O(12)-N(12)-C(12)	122.3(5)	123.8(11)
O(21)-N(21)-C(21)	120.3(5)	119.4(10)
O(22)-N(22)-C(22)	119.8(5)	121.7(10)
N(11)-O(11) ... O(22)	99.7(3)	95.9(6)
N(12)-O(12) ... O(21)	99.7(3)	99.2(7)
N(21)-O(21) ... O(12)	96.9(3)	98.2(7)
N(22)-O(22) ... O(11)	96.0(3)	100.1(6)

Table 13 - extended

$\text{ClCo}(\text{H}_2\text{dmg}_2)(\text{sulfa})^{46}$	$[\text{Co}(\text{H}_2\text{dmg}_2)(\text{clan})_2]\text{Cl}$	$[\text{Co}(\text{H}_2\text{dmg}_2)(\text{an})_2]\text{Cl}^{52}$
90.5(3)	89.8(1)	91.5(4)
91.7(3)	93.2(1)	93.0(5)
89.3(3)		
87.8(3)		
82.0(4)	80.8(1)	80.8(3)
98.7(4)		
179.3(4)		
98.7(3)		
179.2(4)		
80.6(3)		
89.6(3)		
88.5(3)		
90.5(3)		
91.9(3)		
179.7(2)		
119.1(6)	119.7(1)	119.5(7)
123.0(6)	121.3(1)	121.4(6)
122.6(6)	122.7(1)	122.9(7)
121.6(6)		
123.6(6)		
116.4(7)	116.9(2)	116.8(9)
117.4(7)	117.7(2)	117.8(9)
116.3(7)		
116.8(7)		
120.5(9)	121.8(2)	121.8(12)
120.0(8)	119.6(2)	119.2(10)
122.2(8)		
120.1(9)		
98.3(6)		
97.8(6)		
99.2(5)		
96.8(6)		

Table 13 - continued

	$\text{ClCo}(\text{H}_2\text{dmg}_2)$ (clan)	$\text{ClCo}(\text{H}_2\text{dpq}_2)$ (clan)
N(11)-C(11)-C(12)	112.8(6)	112.1(10)
N(11)-C(11)-C(13)	122.9(6)	125.6(11)
N(12)-C(12)-C(11)	113.1(6)	115.5(11)
N(12)-C(12)-C(14)	122.5(6)	119.5(11)
N(21)-C(21)-C(22)	113.5(6)	112.2(10)
N(21)-C(21)-C(23)	112.4(7)	120.9(11)
N(22)-C(22)-C(21)	112.3(6)	111.9(10)
N(22)-C(22)-C(24)	123.2(6)	126.0(11)
C(13)-C(11)-C(12)	124.2(6)	122.3(11)
C(14)-C(12)-C(11)	124.4(6)	125.0(11)
C(23)-C(21)-C(22)	124.1(6)	126.8(11)
C(24)-C(22)-C(21)	124.4(6)	122.2(10)

^aThe atomic numbering of $\text{Co}(\text{H}_2\text{dmg}_2)(\text{an})_2\text{Cl}^{52}$ has been changed to correspond to that of compounds of this work.

Table 13 - continued - extended

$\text{C}\ell\text{Co}(\text{H}_2\text{dmg}_2)(\text{sulfa})^{46}$	$[\text{Co}(\text{H}_2\text{dmg}_2)(\text{clan})_2]\text{C}\ell$	$[\text{Co}(\text{H}_2\text{dmg}_2)(\text{an})_2]\text{C}\ell^{52}$
113.3(9)	112.2(2)	112.4(10)
125.0(10)	125.0(2)	124.6(16)
110.7(9)	112.5(2)	112.2(9)
124.0(10)	124.1(2)	125.0(16)
113.1(9)		
120.7(10)		
113.1(9)		
122.9(10)		
121.7(10)	122.9(2)	123.0(12)
125.3(10)	123.4(2)	122.9(13)
126.1(10)		
123.6(10)		

ligands. This is in contrast to results reported for various $\text{Co}(\text{H}_2\text{dmg}_2)$ complexes^{40,47,48,50,52} as well as for $\text{Fe}(\text{H}_2\text{dmg}_2)\cdot(\text{imidazole})_2$,⁵⁵ $\text{Ni}(\text{H}_2\text{dmg}_2)$,⁵⁶ and $\text{Cu}(\text{H}_2\text{dmg}_2)$,⁵⁷ where either the hydrogen bridges were assumed to be equidistant from the two oxygen atoms or the ligand to be monoprotonated. The assumption of a symmetrical bridge may have in part been based on the earlier IR spectroscopic work on $\text{M}(\text{H}_2\text{dmg}_2)$ complexes where the weak band due to an O-H vibration near 1725 cm^{-1} was assumed to indicate a very short and symmetrical O-H-O bridge.^{19,20} McFadden and McPhail⁵¹ reported the structure of $\text{Co}(\text{H}_2\text{dmg}_2)(\text{CH}_3)(\text{H}_2\text{O})$ in which both bridging hydrogen atoms if ordered are required crystallographically to be on one dmg ligand. No comment was made concerning the bridging hydrogen atoms.

Although both hydrogen bridges in $\{\text{Co}(\text{H}_2\text{dmg}_2)(\text{clan})$ appear to be shifted toward one dmg where the O(21)-H(b2) and O(22)-H(B1) distances are $1.16(3)$ and $1.17(15) \text{ \AA}$ while the O(12)-H(B2) and O(11)-H(B1) distances are $1.30(10)$ and $1.41(14) \text{ \AA}$, the experimental uncertainty is too large to show that result to be significant.

The hydrogen bridges in $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}$ are not symmetrical and each dmg is singly protonated. The O(12)-H(B1) distance is $1.07(3) \text{ \AA}$ and the O(11)-H(B1) distance is $1.44(3) \text{ \AA}$. The gross structure is very similar to that of $[\text{Co}(\text{Hdmg})_2(\text{an})_2]\text{Cl}$.

Bowman *et al.*⁵⁵ suggested the N-O distance to be a sensitive indicator of the position of the bridging hydrogen.

Table 14
Deviations and Equations of Selected Least-Squares Planes
in $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{clan})^{\alpha}$

(a) Deviations ($\text{\AA} \times 10^{+3}$)

	Plane 1	Plane 2	Plane 3	Plane 4
Co	5	-1796	0*	72
O(11)	25		-2574	209
O(12)	-23		2533	80
N(11)	-2*		-1260	175
N(12)	2*		1218	140
C(11)	4*		-775	251
C(12)	-4*		682	220
C(13)	73		-1633	413
C(14)	41		1498	330
O(21)	34		2602	-22
O(22)	-79		-2548	-55
N(21)	48		1269	1*
N(22)	7		-1197	-1*
C(21)	113		795	-2*
C(22)	94		-673	2*
C(23)	230		1664	24
C(24)	143		-1482	-17
N(1)	2004	-41	0*	
C(1)	2752	-12*	0*	
C(2)	3092	5*	1197	
C(3)	3767	5*	1204	
C(4)	4105	-9*	18	
C(5)	3790	3*	-1193	
C(6)	3112	8*	-1204	
Cl(2)	4946	-28	5	
Cl(1)	-2252		-29	

(b) Coefficients of the Plane Equation⁵⁸
 $Ax + By + Cz = D$

Plane	A	B	C	D
1	0.8529	0.4975	0.1583	1.6954

Table 14 - continued

Plane	A	B	C	D
2	0.9995	0.0282	0.0142	1.6347
3	0.0208	-0.3098	0.9506	1.7440
4	0.8174	0.5536	0.1594	1.8108

^aThe deviations of atoms used to define the plane are marked with an asterisk.

Table 15
 Deviations and Equations of Selected Least-Squares Planes
 in $\text{ClCo}(\text{H}_2\text{dpg}_2)(\text{clan})^{\text{a}}$

(a) Deviations ($\text{\AA} \times 10^{+3}$)

	Plane 1	Plane 2	Plane 3	Plane 4
Co	19	-1932	0*	29
O(11)	-56		1350	-101
O(12)	122		-2807	192
N(11)	17*		160	2
N(12)	-18*		-1881	24
C(11)	-30*		-945	-30
C(12)	30*		-2118	62
C(13)	-179		-1048	-195
C(14)	94		-3571	149
O(21)	-5			60
O(22)	-37			-87
N(21)	-43			-8*
N(22)	28			8*
C(21)	-9			13*
C(22)	-1			-13*
C(23)	93			133
C(24)	-53			-89
N(1)	1958	-143	0*	
C(1)	2755	-184	0*	
C(2)	3166	4*	-1171	
C(3)	3843	16*	-1184	
C(4)	4069	-20*	10	
C(5)	3720	5*	1166	
C(6)	3076	14*	1205	
Cl(2)	4901	-1	-10	
Cl(1)	-2223		24	
C(1A)	676			
C(2A)	510			
C(3A)	-529			
C(4A)	-1117			
C(5A)	-1255			

Table 15 - continued

	Plane 1	Plane 2	Plane 3	Plane 4
C (1B)	1094			
C (2B)	1237			
C (3B)	427			
C (4B)	-641			
C (5B)	-821			
C (1C)			1232	
C (2C)			1330	
C (3C)			351	
C (4C)			-734	
C (5C)			-827	
C (1D)			571	
C (2D)			554	
C (3D)			-230	
C (4D)			-874	
C (5D)			-658	

(b) Coefficients of the Plane Equation⁵⁸
 $Ax + By + Cz = D$

Plane	A	B	C	D
1	0.1954	-0.9752	-0.1036	3.3476
2	-0.2301	-0.9634	0.1374	1.9549
3	0.5744	0.0297	0.8181	-7.1571
4	0.1976	-0.9769	-0.0813	3.2302

^aThe deviations of atoms used to define the plane are marked with an asterisk.

Table 16

Deviations and Equations of Selected Least Squares Planes
in $\text{Co}(\text{Hdmg})_2(\text{clan})_2\text{Cl}^a$

(a) Deviations ($\text{\AA} \times 10^{+3}$)

	Plane 1	Plane 2	Plane 3
Co	10	-1772	0*
O(11)	36		2519
O(12)	14		-523
N(11)	0*		1902
N(12)	0*		428
C(11)	1*		2567
C(12)	-1*		1677
C(13)	26		4046
C(14)	11		2173
N(1)	2009	-28	0*
C(1)	2799	-2*	0*
C(2)	3030	2*	1194
C(3)	3758	-1*	1208
C(4)	4257	0*	37
C(5)	4048	0*	-1163
C(6)	3312	1*	-1174
Cl(2)	5159	-33	53

(b) Coefficients of the Plane Equation⁵⁹
 $Ax + By + Cz = D$

Plane	A	B	C	D
1	-0.4938	0.6723	0.5515	-0.0101
2	-0.5672	0.8236	0.0096	1.7716
3	0.7336	0.6606	-0.1594	0.0000

^aThe deviations of atoms used to define the plane are marked with an asterisk.

Hydrogen Bonds with Estimated Standard Deviations Given in Parentheses.

Bond	Position σ_{\pm}^x			Distances (\AA)			Angles ($^\circ$)
	D-H...A ^a	A	D-H	H...A	D...A	D-H...A	
C(1)O(H ₂ dmg ₂) (c1an)							
O(21)-H(B2)...O(12)	x,y,z		1.13(8)		1.36(8)	2.479(7)	156(7)
O(22)-H(B1)...O(11)	x,y,z		1.16(8)		1.37(8)	2.497(7)	161(6)
N(1)-H(7)...O(w1)	x,y,z		1.03(7)		1.92(7)	2.900(7)	157(6)
N(1)-H(8)...O(w2)	x,y,z		0.93(6)		2.04(7)	2.849(7)	164(6)
O(w1)-H(w1)...C(1)	1+x,y,z		0.70(8)		2.76(8)	3.284(6)	134(3)
O(w1)-H(m1')...O(11)	1-x,1-y,-z		0.80(13)		2.34(12)	2.823(7)	120(11)
O(w2)-H(w2)...O(12)	1-x,1-y,1-z		0.79(7)		2.05(7)	2.813(6)	164(8)
O(w2)-H(w2')...C(1)	1+x,y,z		0.71(7)		2.56(7)	3.226(5)	157(7)
[Co(H ₂ dmg ₂) (c1an ₂)Cl]							
O(12)-H(B1)...O(11)	-x,-y,-z		1.07(3)		1.44(3)	2.495(3)	170(3)
N(1)-H(7)...O(11)	-1+x,y,z		0.93(2)		2.07(3)	2.918(3)	163(2)
N(1)-H(8)...C(1)	x,y,z		0.94(2)		2.17(2)	2.100(2)	168(2)
C(1)O(H ₂ dpg ₂) (c1an)							
O(21)-H(B1)...O(12)	x,y,z		1.16(10)		1.30(10)	2.460(12)	172(10)
O(22)-H(B2)...O(11)	x,y,z		1.17(15)		1.41(14)	2.540(11)	159(13)
O(S1)....O(22)	x,y,z					2.852(15)	

^a Mor-Hydrogen...Acceptor, D-H at x,y,z.

Dissimilar N-O bond lengths should indicate the hydrogen is not symmetrically located and is closer to the dm_g with the longer bond. This holds true in ClCo(H₂dm_g)(dm_g)(clan) where the N-O distances appear to be different. The N(21)-O(21) and N(22)-O(22) distances of 1.348(6) and 1.359(6) Å in the diprotonated dm_g are longer than the N(12)-O(12) and N(11)-O(11) distances of 1.329(6) and 1.337(6) Å in the dianionic dm_g. Using the significance test described by Cruickshank and Robertson⁶⁰ the N(21)-O(21) distance is possibly longer than the N(12)-O(12) with a t_o value of 2.24 and the N(22)-O(22) bond is significantly longer than the N(11)-O(11) bond with a t_o value of 2.59. Also, in [Co(Hdm_g)₂(clan)₂]Cl the N(12)-O(12) bond of 1.362(3) Å is significantly longer than the N(11)-O(11) bond of 1.340(3) Å, where the bridging hydrogen atom is bonded to O(12). Neither the N-O distances nor the bridging O-H distances in ClCo(H₂dpg₂)(clan) are significantly different. In [Co(Hdm_g)₂(an)₂]Cl where the hydrogen atoms are not significantly removed from a symmetrical position, the N(12)-O(12) distance is shorter than that of N(11)-O(11). The difference in these two bond lengths of 1.333(6) and 1.353(6) Å is of possible significance ($t_o = 2.36$). The sensitivity of the N-O bond as an indicator of the bridge position is questionable. The N-O bonds are not significantly different in ClCo(H₂dm_g)(dm_g)(sulfa) when both bridging hydrogen atoms are shifted to one dm_g. In the closely related dimethyl(3,3'-trimethylenedinitrilo)bis-(butan-2-one-oximato)cobalt(III) complex the two N-O distances are equal

even though an asymmetric hydrogen bridge is clearly indicated by the difference Fourier syntheses.⁶¹ Although a difference in the N-O bond lengths as a function of protonation is reasonable, there are very few structures so precisely determined that this comparison can be made. Hence, no general conclusion may be made. However, when a significant difference in the N-O distances has been found and the bridging hydrogen atom has been precisely located, the hydrogen atom is associated with the longer N-O bond.

Another point in support of the formulation $\text{ClCo}(\text{H}_2\text{dmg}) - (\text{dmg})(\text{clan})$ is the difference in the Co-N bond lengths. The Co-N distances on the H_2dmg side are 1.908(5) and 1.906(5) \AA compared to distances of 1.872(5) and 1.884(5) \AA on the dm^o side. The differences in the Co-N bond lengths are significant and the shorter distances involve the dianionic group. This holds true in the other cases where the presence of both H_2dmg and dm^o ligands has been indicated. In $\text{C}_6\text{Co}(\text{H}_2\text{dmg})(\text{dmg}) - (\text{sulfa})$ ⁴⁶ and in $\text{Co}(\text{H}_2\text{dmg}_2)(\text{CH}_3)(\text{H}_2\text{O})$ ⁵¹ the distances from the cobalt atom to the dianionic ligand are shorter than the distances to the neutral H_2dmg ligand. This is not the case in $\text{ClCo}(\text{H}_2\text{dpq}_2)(\text{clan})$ where the distances from the cobalt atom to what would be the dpq dianionic ligand, 1.935(11) and 1.908(9) \AA , appear to be longer than the corresponding distances to the H_2dpq ligand, 1.887(10) and 1.897(9) \AA . These differences together with the apparent positions of the bridging hydrogen atoms (vide supra) in $\text{ClCo}(\text{H}_2\text{dpq}_2)(\text{clan})$ are of questionable significance.

For the mononegative ligands in $[Co(Hdmg)_2(\text{clan})_2]Cl$ the Co-N distances are significantly different. However, N(12) which is bonded to the protonated oxygen atom is closer to the cobalt atom than is N(11) with distances of 1.889(2) and 1.906 (2) Å, respectively. The same relationship holds in $Fe(Hdmg)_2 - (\text{imidazole})_2$,⁵⁵ the only other $M(Hdmg)_2$ complex whose X-ray structure precisely places one bridging hydrogen on each dmfg and shows a significant difference in the metal to nitrogen distance.

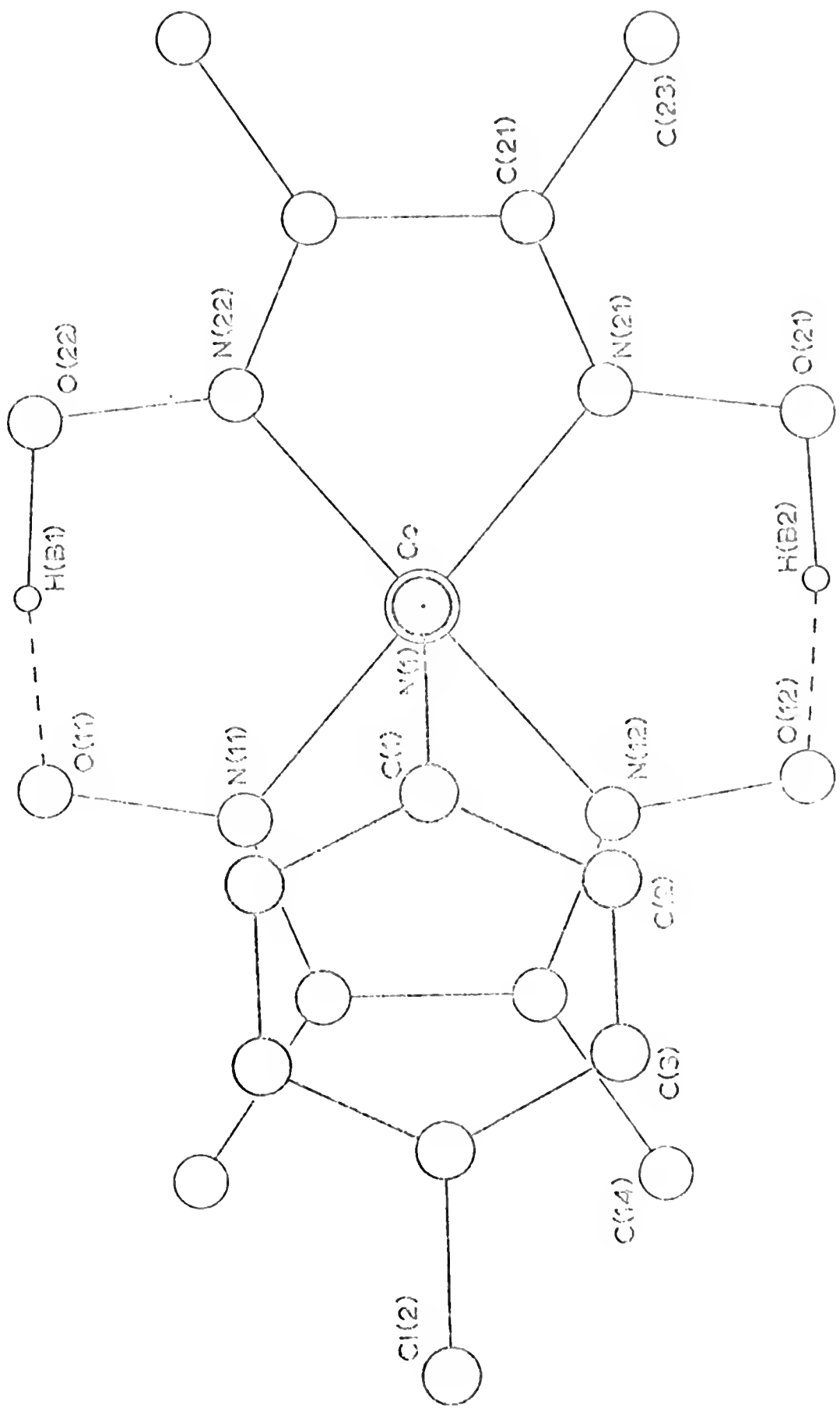
An unsymmetrical hydrogen-bonding system involving two similar atoms may be fluxional.⁶² In such a system two equilibrium positions, i.e. potential wells, exist for the hydrogen atom. Each of these positions may be considered as having the hydrogen atom covalently bonded to one atom and hydrogen bonded to the other. For the system to be truly fluxional the energy barrier between the two positions must be thermally accessible. Depending on the relative depths of the potential wells, the energy barrier between them, and the thermal energy of the system the position of the hydrogen atom as indicated by X-ray diffraction experiments would vary. Because of the diffuse appearance of the bridging hydrogen atoms of the $M(H_2dmg)_2$ complexes in difference Fourier syntheses, a fluxional system with two potential wells of unequal depth seems reasonable. The relative populations of the two positions will depend somewhat on the depths of the potential wells. The experimentally determined position (or positions) of the hydrogen atom will reflect these populations. As the depths of the

potential wells approach equivalence and as the energy barrier between them becomes smaller the position of the hydrogen atoms will become experimentally more uncertain. A fluxional system could, in part, account for the difficulty in precisely locating the bridging hydrogen atoms in $M(H_2\text{dmg}_2)$ complexes.

The orientation of the 4-chloroaniline ligand in the complexes of this study is quite intriguing. A projected view down the Co-N(1) bond for $C\ell\text{Co}(H_2\text{dmg})(\text{dmg})(\text{clan})$ is shown in Figure 4. A similar view for $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}$ is given in Figure 5(a) and one for $C\ell\text{Co}(H_2\text{dpq}_2)(\text{clan})$ is given in Figure 5(b). In $C\ell\text{Co}(H_2\text{dmg})(\text{dmg})(\text{clan})$, as in $C\ell\text{Co}(H_2\text{dmg})(\text{dmg})(\text{sulfa})$ ⁴⁶ the aromatic ring of the aniline is oriented over the dianionic dmg ligand. The orientation angle, i.e. the dihedral angle between the planes having Co-N(1) in common with one containing C(1) and the other containing the bisector of the angle N(11)-Co-N(12), for $C\ell\text{Co}(H_2\text{dmg})(\text{dmg})(\text{clan})$ is 0.9° and for $C\ell\text{Co}(H_2\text{dmg})(\text{dmg})(\text{sulfa})$ is 1.8° as given in Table 18. In $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}$ and in $[\text{Co}(\text{Hdmg})_2(\text{an})_2]\text{Cl}$ the benzene rings are skewed relative to the equatorial ligands with orientation angles of 53.9° and 58.3° , respectively. It seems significant that in the former pair of $\text{Co}(H_2\text{dmg})(\text{dmg})$ type complexes the rings align while in the latter pair of $\text{Co}(H-\text{dmg})_2$ type complexes the rings are skewed. Although the benzene ring of the aniline is tipped from being parallel to the dmg plane by ca. 30° as in other similar complexes (see Table 18) the alignment and the distances between the two planes in $C\ell\text{Co}(H_2\text{dmg})(\text{dmg})(\text{clan})$ suggest a π -type interaction.⁵¹

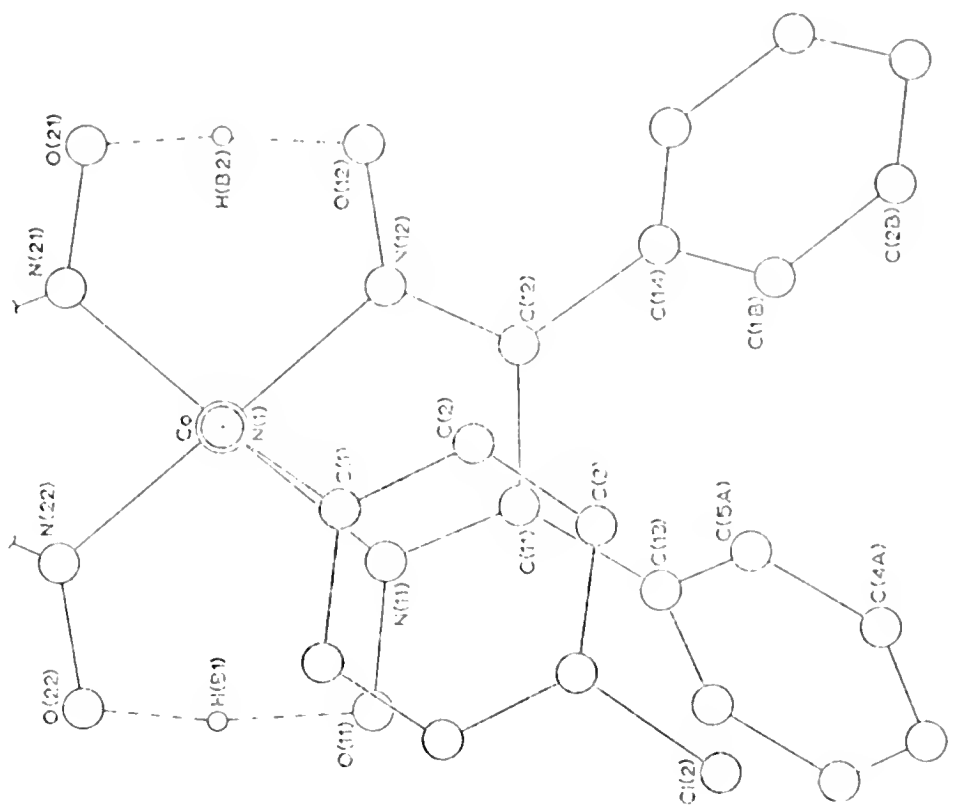
A projected view along Co-N(1) for C₂CO(₂^{dmrg}) (dmrg) (clan).

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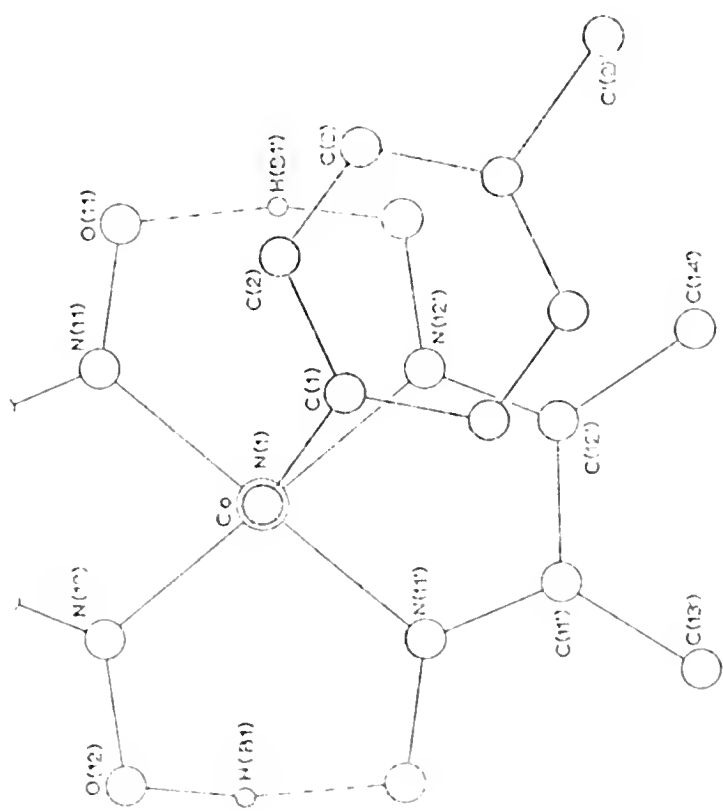


A projected view along Co-N(1) for (a) $[\text{Co}(\text{H}_2\text{dpg})_2(\text{clan})_2]\text{Cl}$ and (b) $\text{C}_{12}\text{Co}(\text{H}_2\text{dpg}_2)$ (clan).

Figure 5



(D)



(E)

Table 18
Dihedral Angles Formed by Selected Planes in Some Cobaloxime Complexes

Intersecting planes	Angles (°)		
	C ₆ Co(H ₂ dmg ₂) (clan)	C ₆ Co(H ₂ dpg ₂) (clan)	C ₆ Co(H ₂ dmg ₂) (sulfa) ⁴⁶
1-2	29.7	28.3	25.2
1-3	89.2	69.9	88.9
1-4	3.8	1.3	5.6
2-3	88.5	87.2	83.5
2-4	33.4	27.8	30.8
3-4	99.8	89.0	89.2
3-5	0.9	36.4	1.8

planes 1 to 4 are least-squares planes as defined in Tables 14-16. Plane 1 is defined by N(1¹), N(12), C(11), and C(12). Plane 2 is defined by the carbon atoms in the benzene ring of the aniline ligand. Plane 3 is defined by Co, N(1), and C(1). Plane 4 is defined by N(21), N(22), C(21), and C(22). Plane 5 contains Co, N(1), and the bisector of the angle N(11)-Co-N(12).

Intersecting
planes

Angles (°)

	$[\text{CO}(\text{H}_2\text{Cmg}_2)(\text{C}_2\text{H}_5)_2]\text{Cl}_2$	$[\text{CO}(\text{H}_2\text{Cmg}_2)(\text{an})_2]\text{Cl}_2$
1-2	33.0	32.9
1-3	89.7	86.9
1-4	(180)	(180)
2-3	82.7	86.9
2-4		
3-4		
3-5	53.8	59.3

distances from the dm_g plane to C(1), C(2), and C(6) given in Table 14 are substantially less than the 3.40 Å interplanar distance in graphite.⁶³ A proton transfer occurring from one Hd_{mg} ligand to the other would increase the electron density within the π-system of the formed dianion. An interaction by which the filled π orbitals of the dm_g overlap with the empty π* orbitals of the aniline would enhance the basicity of the aniline ligand. The complex formed would be stronger than might be expected based on the K_b value alone. This same argument applies to ClCo(H₂dm_g) (dm_g) (sulfa)⁴⁶ which was the first example of ligand-induced proton shift in a molecular complex. While the positions of the bridging protons in ClCo(H₂dm_g) (dm_g) (clan) and [Co(Hd_{mg})₂(clan)₂]Cl are well defined, the bridge in ClCo(H₂dpg₂) (clan) is ill defined and the orientation angle of 36.7° is an intermediate value (see Table 18). The O...O distances in this complex show more variation than those in other related complexes as shown in Table 12. The 0.08 Å difference in the O...O distances is the same as for the corresponding N...N distances. The N(12)...N(21) separation is 2.836(15) Å and the N(11)...N(22) distance is 2.914(13) Å. Concurring with these observed distances, the N(12)-Co-N(21) angle of 97.0(4)° is more acute than the N(11)-Co-N(22) angle of 100.0(4)°. None of the other compounds examined shows any significant differences in the corresponding distances and angles between the diglyoxime ligands.

A comparison of mean bonding distances for each of the reported $\text{Co}(\text{H}_2\text{dmg}_2)$ complexes may be made from Table 19. There appears to be little variation in the average Co-N distances or in the average dimensions within the equatorial dimethylglyoxime ligands as a function of the axial ligand.

Those complexes having chloride as an axial ligand show a definite variation with the nature of the trans ligand. The longest Co-Cl distance is found where tpp is the trans ligand. This is not surprising since phosphines are known to have a very large trans-influence⁶⁴ but the small influence the tpp ligand exerts on the trans-chlorine atom compared to that of an ammonia ligand is unexpected.⁴⁰ There is no significant difference in the Co-N(1) distance involving a clan ligand whether it is trans to a chlorine atom or trans to another clan ligand. The trans-influence appears to occur in $\text{Co}(\text{H}_2\text{dmg}_2)$ complexes but not to a large extent.

The Co-Y distances in the $\text{XCo}(\text{H}_2\text{dmg}_2)\text{Y}$ complexes where Y is a ligand with an sp^3 nitrogen, increase in the following order of Y: $\text{NH}_3 < \text{an} \sim \text{clan} < \text{sulfa}$ (see Table 19). This series can be rationalized in terms of the relative K_b 's for sulfa (2.3×10^{-12}),⁶⁵ clan (9.6×10^{-11}),⁶⁶ aniline (4.0×10^{-10}),⁶⁶ and ammonia (1.3×10^{-5}).⁶⁷ Brückner and Randaccio⁴⁰ did not consider the K_b 's of the different nitrogen donors in their argument of the trend in trans-influencing ligands, X, upon the Co-N bond. The same Co-N distances were used for NH_3 and aniline complexes in their argument for basing the extent of trans-influence on the σ -donor power of the H_2dmg_2

Table 19
Summary of the Average Bond Distances (Å) in XYCo(H₂dmg₂) Complexes. a,b,c,d

X	Y	Co-X	Co-Y	Co-N	N-O
clan	Cℓ	1.999(6)	2.257(2)	1.893(5)	1.343(6)
clan	clan	2.003(2)		1.898(2)	1.351(3)
clan	Cℓ	1.946(11)	2.244(4)	1.907(11)	1.333(12)
clan	Cℓ	2.023(8)	2.235(3)	1.839(8)	1.333(11)
sulFa	Cℓ	1.965(4)	2.251(1)	1.890(5)	1.346(10)
H ₃	Cℓ	2.327(4)	2.277(4)	1.89(1)	1.343(9)
tpp	Cℓ	2.54(2)	2.25(2)	1.95(2)	
Cl ₆ Pt ₄	an	2.001(5)		1.837(6)	1.343(6)
an	H ₂ C	1.990(5)	2.053(3)	1.890(3)	1.352(5)
CH ₂	P(n-Bu ₂) ₃	1.979(1)	2.339(11)	1.876(3)	1.339(7)
C-PY	DY	2.04	2.04	1.83	1.37
C ₁₄ H ₂ COOCH ₃	DY	1.97(1)	2.04(1)	1.88(1)	1.35(2)
DDT					

The ligands X and Y are approximately normal to the plane of the two ring groups.

The entry marked with an asterisk contains Hdmg rather than Hdmg and the distance under C-Cu₃ is C-C₆H₅.
The values given in parentheses are usually the mean of the estimated standard deviations. They are presented only to indicate the precision of the original values in a most general sense.

*C₆-Dy is a carbon-bonded pyridine and DDT is 1,1-bis(4-chlorophenyl)-2-chloroethylene.

Table 12 - extended

X	Y	C-N	C-C	C-CH ₃	O...O	Reference
clan	Cl	1.297(8)	1.463(9)	1.490(11)	2.488(7)	-
clan	clan	1.295(3)	1.477(4)	1.484(4)	2.495(3)	-
clan	clan	1.309(16)	1.459(26)	1.486(17)	2.500(12)	-
*clan	Cl	1.291(14)	1.472(17)	1.498(17)	2.493(11)	46
su12a	Cl	1.282(4)	1.483(6)	1.50(2)	2.486(7)	40
M ₃	Cl	1.300(14)	1.485(15)	1.501(8)	2.50(1)	40
t _{top}	Cl					49
short ₃	Cl					49
an	an	1.294(6)	1.463(7)	1.479(12)	2.491(8)	52
CH ₃	H ₂ O	1.302(5)	1.463(7)	1.494(7)	2.486(4)	51
C-OY	p(n-Bu ₂) ₃	1.295(7)	1.443(3)	1.499(3)	2.474(2)	48
CH ₂ COOC ₂ H ₃	py	1.28	1.46	1.50	2.50	47
DPT	py	1.30(2)	1.43(2)	1.50(11)	2.50(11)	50

ligand as are presented here.

In comparing $\text{ClCo}(\text{H}_2\text{dpg}_2)$ (clan) with $\text{ClCo}(\text{H}_2\text{dmg})$ (dmg) (clan) the distances from the cobalt atom to the equatorial nitrogens in the H_2dpg complex are longer and the distances to the axial ligands are shorter in the same complex. Because the phenyl substituents are inductively more electron withdrawing than methyl groups, Hdpg should be a weaker Lewis base than Hdmg. The equatorial distances to the Hdpg should, therefore, be longer. From an electronic standpoint the cobalt ion in the Hdpg complex would be more positively charged and a better Lewis acid toward the axial ligands than in the Hdmg complex. From a steric point of view the axial ligands are afforded a wider path of approach and will, therefore, be closer to the central cobalt ion when the equatorial ligands are farther away.

The benzene rings in the clan ligands of $\text{ClCo}(\text{H}_2\text{dmg}_2)$ (clan), $\text{ClCo}(\text{H}_2\text{dpg}_2)$ (clan), and $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}$ are planar (see Tables 14-16) having average C-C values of 1.376(3), 1.380(10), and 1.378(3) Å, respectively, with individual values reported in Table 20. The phenyl rings of the Hdpg ligands of $\text{ClCo}(\text{H}_2\text{dpg}_2)$ (clan) are also planar with pertinent values and equations of least-squares planes given in Table 21.

The crystals of $\text{ClCo}(\text{H}_2\text{dmg})$ (dmg) (clan) are held together by six hydrogen bonds where there are eight hydrogen atoms capable of hydrogen bonding. Relevant hydrogen-bonding data are presented in Table 17. Although the O-H...O bridges between the Hdmg and dmg groups are not symmetrical, the O-H

Table 2
Bond Distances and Bond Angles of Concentrated *t*-chloroaniline Molecules with their Estimated Standard Deviations.

Distances (Å)	$\text{C}=\text{CO}(\text{H}_2\text{Cmg}_2) (\text{C}^{\text{12a}})$	$\text{C}=\text{CO}(\text{H}_2\text{Cmg}_2) (\text{C}^{\text{12a}})$	$\text{C}=\text{CO}(\text{H}_2\text{Cmg}_2) (\text{C}^{\text{12a}})$	$\text{C}=\text{CO}(\text{H}_2\text{Cmg}_2) (\text{C}^{\text{12a}})$
$\text{C}(1)-\text{C}(1)$	1.461 (3)	1.451 (26)	1.442 (2)	1.451 (2)
$\text{C}(1)-\text{C}(2)$	1.374 (2)	1.380 (19)	1.383 (2)	1.383 (2)
$\text{C}(4)$	1.364 (10)	1.359 (20)	1.357 (1)	1.357 (1)
$\text{C}(4)-\text{C}(5)$	1.370 (10)	1.352 (20)	1.354 (1)	1.354 (1)
$\text{C}(5)-\text{C}(6)$	1.338 (5)	1.344 (21)	1.378 (2)	1.378 (2)
$\text{C}(6)-\text{C}(1)$	1.372 (10)	1.375 (20)	1.365 (1)	1.365 (1)
$\text{C}(2)-\text{C}(4)$	1.379 (9)	1.403 (17)	1.330 (4)	1.330 (4)
$\text{C}(2)-\text{C}(4)$	1.734 (3)	1.724 (15)	1.752 (3)	1.752 (3)
Angles (°)				
$\text{N}(1)-\text{C}(1)-\text{C}(2)$	119.4 (5)	122.0 (12)	129.4 (2)	129.4 (2)
$\text{N}(1)-\text{C}(1)-\text{C}(6)$	119.2 (5)	120.7 (11)	121.0 (2)	121.0 (2)
$\text{C}(1)-\text{C}(2)-\text{C}(3)$	119.7 (6)	122.4 (13)	120.2 (3)	120.2 (3)
$\text{C}(2)-\text{C}(3)-\text{C}(4)$	119.8 (7)	127.9 (23)	119.5 (3)	119.5 (3)
$\text{C}(3)-\text{C}(4)-\text{C}(5)$	120.7 (7)	120.8 (24)	121.8 (4)	121.8 (4)
$\text{C}(4)-\text{C}(5)-\text{C}(6)$	119.7 (7)	122.3 (14)	118.4 (3)	118.4 (3)
$\text{C}(5)-\text{C}(6)-\text{C}(1)$	118.7 (6)	119.4 (12)	120.7 (1)	120.7 (1)
$\text{C}(6)-\text{C}(1)-\text{C}(2)$	121.4 (6)	117.2 (12)	119.5 (2)	119.5 (2)
$\text{C}(2)-\text{C}(4)-\text{C}(3)$	120.5 (6)	117.7 (11)	119.4 (3)	119.4 (3)
$\text{C}(3)-\text{C}(4)-\text{C}(5)$	118.8 (6)	121.3 (11)	118.8 (3)	118.8 (3)

Table 21

Bond Distances, Bond Angles, and Least-Squares Planes of the Phenyl Rings in $\text{C}_6\text{Co}(\text{H}_2\text{dpq}_2)$ (clan) with Their Estimated Standard Deviations.

(a) Distances	n = 13	14	23	24
	$\ell = A$	B	C	D
C(n)-C(1 ℓ)	1.363(18)	1.411(20)	1.370(19)	1.426(17)
C(n)-C(5 ℓ)	1.421(20)	1.396(20)	1.356(17)	1.458(18)
C(1 ℓ)-C(2 ℓ)	1.368(19)	1.371(20)	1.432(19)	1.385(19)
C(2 ℓ)-C(3 ℓ)	1.370(21)	1.367(20)	1.351(21)	1.401(20)
C(3 ℓ)-C(4 ℓ)	1.352(20)	1.391(23)	1.371(23)	1.397(20)
C(4 ℓ)-C(5 ℓ)	1.374(20)	1.390(20)	1.409(18)	1.397(18)
(b) Angles ($^\circ$)				
C(n-2)-C(n)-C(1 ℓ)	123.9(11)	119.3(12)	120.9(11)	121.7(11)
C(n-2)-C(n)-C(5 ℓ)	119.9(11)	120.9(12)	121.2(11)	120.6(11)
C(n)-C(1 ℓ)-C(2 ℓ)	122.8(13)	120.6(13)	122.3(12)	122.6(12)
C(1 ℓ)-C(2 ℓ)-C(3 ℓ)	119.5(13)	119.3(14)	117.0(13)	117.7(13)
C(2 ℓ)-C(3 ℓ)-C(4 ℓ)	120.5(13)	121.0(14)	122.2(14)	122.5(13)
C(3 ℓ)-C(4 ℓ)-C(5 ℓ)	120.0(14)	120.4(14)	118.7(13)	120.2(13)
C(4 ℓ)-C(5 ℓ)-C(n)	121.0(13)	118.4(14)	121.7(12)	119.0(12)
C(5 ℓ)-C(n)-C(1 ℓ)	116.2(12)	119.8(13)	117.9(12)	117.7(11)
(c) Deviations ($\text{\AA} \times 10^{+3}$) from Least-Squares Planes of Phenyl Rings				
C(n)	2	41	-3	3
C(1 ℓ)	-7	-14	8	-12
C(2 ℓ)	10	-31	15	24
C(3 ℓ)	-10	48	-29	-28
C(4 ℓ)	5	-20	3	20
C(5 ℓ)	-2	-24	19	-7
C(n-2)	-3	172	-16	20
(d) Coefficients of the Plane Equation $PX + QY + RZ = S$				
	P	Q	R	S
Phenyl A	-0.5815	0.5296	-0.6176	4.7459
Phenyl B	-0.4144	-0.7611	0.4990	3.1793
Phenyl C	-0.6482	0.3950	-0.6509	4.0341
Phenyl D	-0.1592	-0.8986	0.4088	1.3642

distances are longer than might be expected. The two hydrogen atoms on N(1) of the clan ligand both hydrogen bond to different water molecules. The hydrogen atoms of one water molecule, O(w2), form reasonably strong hydrogen bonds to O(12) and Cl(1). The hydrogen atoms on O(w1), however, have only short contacts with angles indicating only weak hydrogen bonds.

While $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}$ and $\text{ClCo}(\text{H}_2\text{dpg}_2)(\text{clan})$ both exhibit the hydrogen bonding between the equatorial ligands, $\text{ClCo}(\text{H}_2\text{dpg}_2)(\text{clan})$ has no intermolecular hydrogen bonds. While the hydrogen atom on the solvent molecule was not located, a hydrogen bond may exist between O(81) and O(22). Each molecule of $[\text{Co}(\text{Hdmg})_2(\text{clan})_2]\text{Cl}$ possesses two intermolecular hydrogen bonds. Each clan molecule shows a hydrogen bond from N(1) to the O(11) of another molecule. The other hydrogen on each N(1) is hydrogen bonded to the ionic chloride. Relevant hydrogen-bonding data for these two compounds are also presented in Table 17.

All intermolecular distances less than 3.6 \AA were calculated and carefully examined. No unusually short intermolecular distances were found.

Ligand-induced proton shifts may be of biological significance. Since proton transfers in living systems are relatively common, the study presented here provides an important examination of orientation effects and enhanced stabilities which may be achieved by a small shift of one proton.

CHAPTER 5

A NOVEL BINUCLEATING LIGAND: THE CRYSTAL AND MOLECULAR STRUCTURES OF 1,4-DIHYDRAZINOPHTHALAZINEBIS(2-PYRIDINIUMCARBOXALDIMINE) NITRATE DIHYDRATE AND μ -CHLOROTETRAQUA[1,4-DIHYDRAZINOPHTHALAZINEBIS(2-PYRIDINECARBOXALDIMINE)] DINICKEL(II) CHLORIDE DIHYDRATE

Binuclear complexes of chelating ligands have been of interest recently for their potential activation of other ligands at an accessible bridging site⁶⁸⁻⁷³ and for their magnetic properties.^{24,74-80} The structure of $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhpipy})]\text{Cl}_3$ shows the planar chelating ligand, dhpipy, to be capable of binding two metal atoms simultaneously. In that complex a bridging site between the nickel ions is occupied by a chloride ion. Therefore, at least one bridging ligand in addition to dhpipy may be accommodated by $M_2\text{dhpipy}$ complexes.

While the study of magnetic interactions between metal ions through bridging atoms in such systems is convenient and theoretically significant, the catalytic possibilities of this type system are exceptional. The nitrogen-fixing enzyme nitrogenase has been considered to contain a polynuclear active site.^{6,7}

Although the mechanism of the reduction of N_2 to NH_3 by nitrogenase is not understood N_2 is believed to be coordinated to the metal ions of the enzyme.^{67,81,82} Nitrogenase has been shown to reduce a wide variety of small molecules which contain a triple bond.⁷ The distance between the metal

ions should be of importance in the activation of those molecules. In the complexes of Robson and coworkers⁶⁸⁻⁷³ and of Okawa *et al.*⁸³ the metal-metal distance is essentially controlled by a single bridging phenoxide ion. However, in diphpy complexes the metal ion separation is fixed at a greater distance by the geometry of the chelating ligand. Therefore, larger molecules which are reduced in the presence of nitrogenase, e.g., N₂, N₃, N₂O, C₂H₂, and HCN,⁷ should be suitable for incorporation as bridging molecules opposite the N-N bridge of diphpy. The syntheses and X-ray structures of H₂diphpy(NO₃)₂·H₂O and [Ni₂Cl(H₂O)₄(diphpy)]Cl₃·2H₂O were undertaken to examine the nature of the accessible bridging site in complexes of this type ligand.

Solution and Refinement of the Structure
of H₂diphpy(NO₃)₂·2H₂O

The direct method of symbolic addition was used in which the signs of two hundred large E's were assigned. All fourteen nonhydrogen atoms of the ligand within the asymmetric unit were located in an E-map computed from the signed E values. Two Fourier syntheses were used to validate the selected model, locate the remaining nonhydrogen atoms, and refine the atomic parameters. The refinement is outlined in Table 5. The observed and calculated structure factors are given in Table B-3. The final positional and thermal parameters are presented in Tables 22 and 23.

Table 22

Final Atomic Parameters^a for the Nonhydrogen Atoms for $\text{H}_2\text{dhpypy}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$

Atom	x	y	z	α_{11}	α_{22}	β_{33}	β_{12}	β_{13}	β_{23}
C(1)	378(2)	35336(3)	1603(3)	21(1)	80(4)	76(4)	0(3)	6(3)	-6(6)
C(2)	131(2)	2379(3)	2019(3)	21(1)	71(3)	83(4)	-1(3)	5(3)	3(6)
C(3)	354(2)	1293(3)	1534(3)	33(1)	74(4)	110(4)	3(3)	36(4)	-14(7)
C(4)	172(2)	240(3)	2015(4)	39(2)	73(4)	140(5)	-2(4)	43(4)	-13(7)
C(10)	1382(2)	4771(3)	-289(3)	24(1)	97(4)	86(4)	-1(3)	27(3)	-11(6)
C(11)	1579(2)	5947(3)	-658(3)	20(1)	86(4)	91(4)	-7(3)	9(3)	0(6)
C(12)	1963(2)	6072(4)	-1546(4)	27(1)	112(4)	103(4)	-8(4)	25(4)	2(7)
O(13)	2116(2)	7214(4)	-1912(4)	30(1)	142(5)	119(5)	-13(4)	43(4)	36(3)
O(14)	1883(2)	8189(4)	-1367(4)	39(1)	114(5)	143(5)	-15(4)	31(4)	44(8)
O(15)	1521(2)	8013(3)	-457(4)	23(1)	92(4)	149(5)	-9(4)	23(4)	8(8)
N(6)	290(5)	4549(2)	2054(2)	23(1)	77(3)	89(3)	4(3)	25(2)	-7(5)
N(7)	783(1)	3597(2)	753(3)	30(1)	77(3)	99(3)	-2(3)	35(3)	-6(5)
N(8)	972(1)	4712(2)	448(2)	25(1)	73(3)	95(3)	-4(3)	25(3)	1(5)
N(10)	1363(1)	6918(3)	-123(3)	27(1)	81(3)	111(3)	-5(3)	23(3)	7(5)
N(20)	1628(2)	1294(3)	-731(4)	68(2)	93(4)	223(6)	33(4)	149(5)	9(7)
O(1)	565(2)	6955(3)	1629(3)	53(1)	113(3)	162(4)	14(3)	62(3)	-50(6)
O(20)	1843(2)	1108(4)	-1701(5)	114(2)	159(5)	445(9)	15(5)	355(8)	-29(10)
O(21)	1086(2)	1953(3)	-974(3)	58(1)	123(3)	151(4)	34(3)	76(4)	-26(6)
O(22)	1861(2)	1062(4)	290(4)	89(2)	285(7)	127(6)	56(5)	109(11)	

^a All values are $\times 10^4$. The estimated standard deviations are given in parentheses. The temperature factors are of the form: $\exp[-(\beta_{11}h^2 + \beta_{22}k^2 + \beta_{33}l^2 + \beta_{12}hk + \beta_{13}hl + \beta_{23}kl)]$

Final Parameters for the Hydrogen Atoms in $\text{H}_2\text{Diphospho}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$ ^a

Atom bonded to	Distance	x	y	z	B
P(1) [O(1)]	0.78(5)	57(3)	629(4)	185(5)	12.5(1.5)
H(2) [O(1)]	0.88(4)	80(2)	737(4)	228(4)	10.1(1.3)
H(3) [C(3)]	1.00(3)	61(2)	129(3)	84(3)	6.2(0.9)
H(4) [C(4)]	1.01(4)	33(2)	-54(4)	171(3)	8.0(1.1)
H(10) [C(10)]	1.05(4)	157(2)	402(3)	-68(3)	6.5(1.0)
H(12) [C(12)]	1.02(3)	212(2)	533(3)	-196(3)	5.7(0.9)
H(13) [C(13)]	0.99(4)	235(2)	734(4)	-262(4)	9.4(1.3)
H(14) [C(14)]	0.91(4)	200(2)	395(4)	-152(4)	7.9(1.1)
H(15) [C(15)]	1.00(4)	136(2)	870(3)	-1(3)	6.5(1.0)
H(2) [N(2)]	0.95(4)	65(2)	293(4)	26(4)	9.4(1.2)
H(DY) [N(10)]	1.21(6)	105(3)	682(5)	69(5)	15.9(1.8)

^aOne hydrogen atom is given followed by the atom to which it is bonded in brackets, the corresponding bond distance (in Å), the positional parameters ($\times 10^{+3}$), and the isotropic thermal parameter (in \AA^2). The estimated standard deviations are given in parentheses.

Solution and Refinement of the Structure
of $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhphpy})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$

The position of Ni(1) was determined from a sharpened three-dimensional Patterson function. The positions of the remaining atoms were determined in a manner analogous to that used with $\text{C}_2\text{Co}(\text{H}_2\text{dmg})(\text{dmg})(\text{clan})$. After the hydrogen atoms were located they were included in further refinement with each having an isotropic thermal parameter one unit higher than the refined isotropic value for the atom to which the hydrogen atom was bonded. A summary of the refinement is given in Table 5. The scattering factors for the nonhydrogen atoms were from Hanson *et al.*²⁹ and the hydrogen scattering factors from Stewart *et al.*³⁰ Lists of observed and calculated structure factors are given in Table B-4. The final positional and thermal parameters are listed in Tables 24 and 25.

Results and Discussion

The atomic numbering and thermal ellipsoids of $\text{H}_2\text{dhphpy}\cdot(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$ are shown in Figure 6 and those of $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhphpy})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$ are shown in Figure 7. Selected interatomic distances of both compounds are listed in Table 26 and corresponding angles are given in Tables 27 and 28. Both compounds crystallize with the cationic complexes, their anions, and water molecules linked in a three-dimensional hydrogen-bonded network. The postulated hydrogen bonds in the structures are listed in Table 29. Diagrams illustrating the pack-

Table 24
The Final Atomic Parameters^a of the Nonhydrogen Atoms for
[Ni₂Cl(H₂O)₄(dhpypy)]Cl₃·2H₂O

Atom	X	Y	Z	β_{1J}
Ni (1)	11817 (7)	17289 (5)	9891 (3)	442 (5)
Ni (2)	11346 (6)	-4660 (5)	13854 (3)	407 (5)
Cl (1)	1046 (1)	315 (1)	650 (1)	63 (1)
Cl (2)	1196 (2)	4767 (1)	2315 (1)	86 (2)
Cl (3)	1297 (2)	-1548 (1)	3350 (1)	69 (1)
Cl (4)	1288 (2)	3725 (2)	4540 (1)	90 (2)
O (1)	-362 (4)	1864 (3)	639 (?)	55 (3)
O (2)	2743 (4)	1634 (3)	1329 (2)	55 (3)
O (3)	-423 (3)	-532 (3)	1026 (2)	48 (3)
O (4)	2664 (3)	-435 (4)	1708 (2)	48 (3)
O (5)	1626 (4)	-520 (4)	-194 (2)	54 (3)
O (6)	1072 (4)	5047 (5)	3678 (2)	74 (4)
N (1)	1178 (4)	1408 (3)	1690 (2)	50 (4)
N (2)	1153 (4)	576 (3)	1834 (2)	48 (4)
N (3)	1265 (4)	2854 (3)	1801 (2)	74 (5)
N (4)	1258 (4)	2873 (3)	1324 (2)	53 (4)
N (5)	1143 (4)	-469 (3)	2395 (2)	55 (4)
N (6)	1165 (4)	-1007 (3)	2025 (2)	45 (4)
N (10)	1234 (4)	2554 (4)	433 (2)	54 (4)
N (20)	1163 (4)	-1776 (4)	1227 (2)	55 (4)
C (1)	1233 (5)	2052 (4)	1992 (3)	42 (4)
C (2)	1251 (5)	1919 (4)	2495 (3)	36 (4)
C (3)	1280 (5)	2594 (5)	2826 (3)	50 (5)
C (4)	1300 (5)	2410 (5)	3302 (3)	56 (5)
C (5)	1288 (5)	1559 (5)	3450 (3)	51 (5)
C (6)	1260 (5)	987 (4)	3136 (2)	47 (5)
C (7)	1236 (4)	1057 (4)	2649 (2)	29 (4)
C (8)	1182 (4)	394 (4)	2288 (2)	29 (4)
C (10)	1281 (6)	3550 (4)	1076 (3)	71 (6)
C (11)	1291 (5)	3337 (5)	578 (3)	52 (5)

Table 24-- extended

β_{22}	β_{33}	β_{12}	β_{13}	β_{23}
273(4)	106(1)	-66(8)	224(5)	14(4)
244(3)	92(1)	2(8)	184(4)	13(4)
31(1)	10(0)	-10(2)	26(1)	-2(1)
35(1)	14(0)	8(2)	18(1)	0(1)
50(1)	17(0)	46(2)	39(1)	17(1)
72(1)	25(1)	-53(3)	26(2)	-1(1)
42(3)	21(1)	3(5)	26(3)	6(3)
40(2)	21(1)	-6(5)	29(3)	2(3)
58(3)	18(1)	-5(5)	23(3)	22(3)
69(3)	12(1)	17(5)	14(3)	-2(3)
62(3)	17(1)	1(5)	26(3)	-3(3)
111(5)	18(1)	1(7)	35(4)	-6(4)
23(2)	11(1)	-2(5)	25(3)	-1(2)
24(2)	11(1)	-1(5)	25(3)	1(2)
26(2)	15(1)	-14(6)	38(4)	-3(3)
31(3)	14(1)	-12(5)	29(3)	3(3)
25(2)	9(1)	1(5)	12(3)	1(2)
26(2)	11(1)	-1(5)	21(3)	3(2)
41(3)	14(1)	5(5)	33(3)	10(3)
32(3)	13(1)	13(5)	26(3)	0(3)
24(3)	14(1)	-9(6)	26(4)	0(3)
30(3)	13(1)	-11(6)	14(4)	-6(3)
37(4)	14(1)	-12(7)	27(4)	-10(3)
44(4)	14(1)	-8(7)	26(4)	-17(4)
55(4)	12(1)	-17(7)	23(4)	-11(4)
40(4)	9(1)	-2(6)	18(4)	1(3)
32(3)	10(1)	-8(6)	15(4)	-3(3)
28(3)	12(1)	-5(6)	20(3)	-2(3)
30(3)	20(2)	-7(7)	41(5)	7(4)
36(3)	17(1)	-3(7)	29(4)	7(4)

Table 24 - continued

Atom	x	y	z	β_{11}
C(12)	1343(6)	4055(5)	270(3)	83(7)
C(13)	1353(7)	3839(6)	-202(3)	86(7)
C(14)	1308(6)	2992(6)	-348(3)	73(6)
C(15)	1251(6)	2361(5)	-17(3)	59(6)
C(20)	1143(5)	-1829(4)	2053(3)	54(5)
C(21)	1147(5)	-2273(4)	1601(3)	48(5)
C(22)	1143(6)	-3175(5)	1577(3)	64(6)
C(23)	1165(6)	-3554(5)	1149(3)	77(6)
C(24)	1204(6)	-3048(5)	758(3)	76(6)
C(25)	1195(5)	-2157(5)	812(3)	55(5)

^aAll values are $\times 10^4$ except for those of Ni(1) and Ni(2) which are $\times 10^5$. The estimated standard deviations are given in parentheses. The temperature factors are of the form: $\exp\{-(\beta_{11}h^2 + \beta_{22}k^2 + \beta_{33}l^2 + \beta_{12}hk + \beta_{13}hl + \beta_{23}kl)\}$

Table 24 - extended - continued

β_{22}	β_{33}	β_{12}	β_{13}	β_{23}
49(4)	20(2)	-19(9)	36(6)	19(4)
68(5)	17(2)	-27(10)	34(6)	21(5)
68(5)	20(2)	7(9)	47(6)	15(5)
60(5)	14(1)	-6(8)	27(5)	5(4)
29(3)	14(1)	0(6)	29(4)	6(3)
31(3)	14(1)	10(6)	25(4)	5(3)
32(3)	21(2)	6(7)	31(5)	3(4)
39(4)	21(2)	15(8)	33(5)	-6(4)
43(4)	19(2)	8(8)	24(5)	-13(4)
39(4)	16(1)	2(7)	24(5)	-6(4)

Table 25
Final Parameters for the Hydrogen Atoms in $\text{Li}_2\text{C}_2(\text{H}_2\text{O})_6\text{(Chloro)}\text{LiCl}_2 \cdot 2\text{H}_2\text{O}^2$

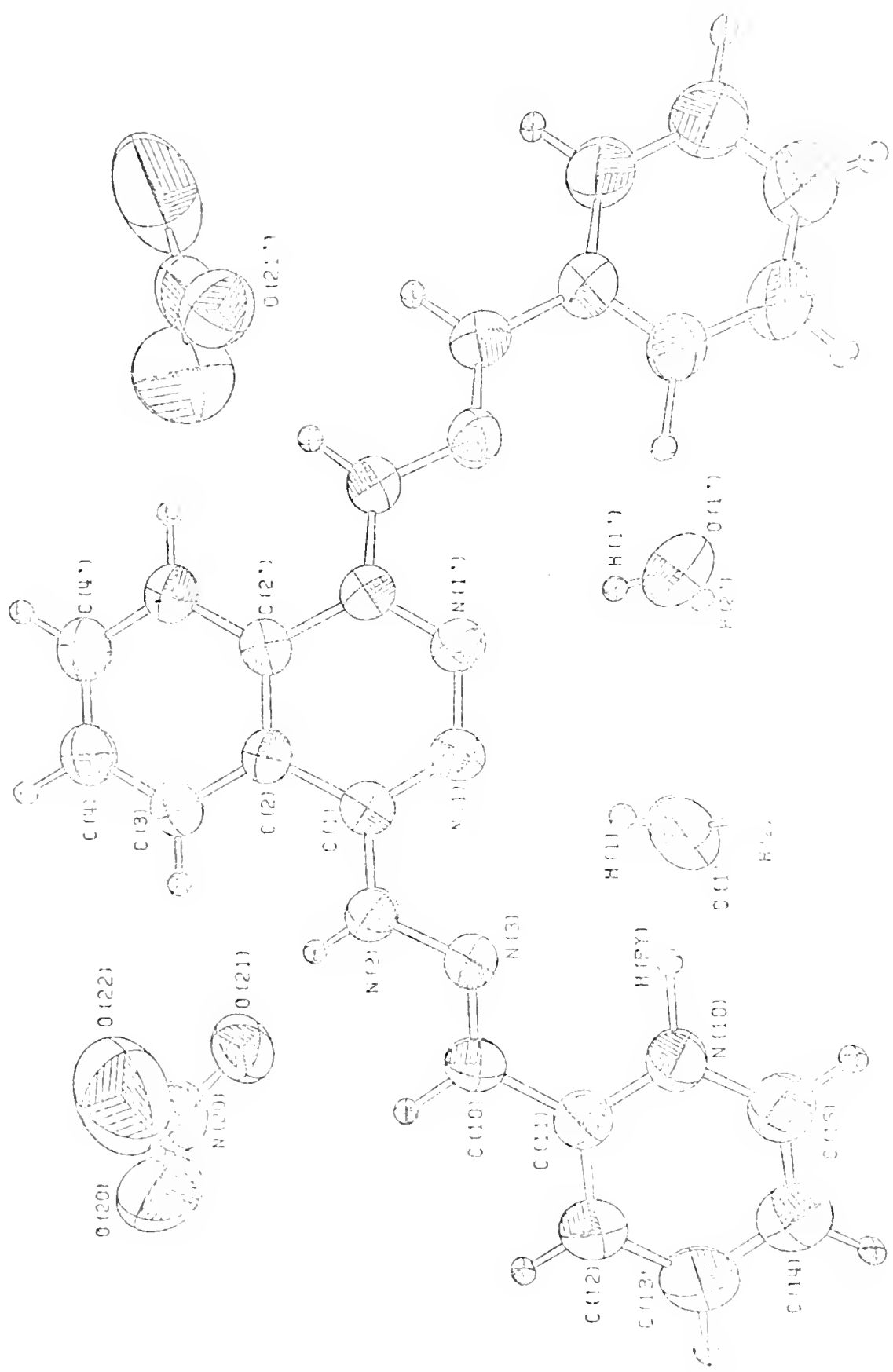
Atom Bonded to	Atom B	Distance x	Distance y	Distance z
H(N ²) [N(3)]		0.91	1397	3355
H(N ⁵) [N(5)]		0.97	1394	-703
H(C3) [C(3)]		1.09	1271	3289
H(C4) [C(4)]		1.05	1322	2949
H(C5) [C(5)]		0.99	1356	1515
H(C6) [C(6)]		1.02	1263	313
H(10) [C(10)]		1.06	1372	4129
H(12) [C(12)]		1.15	1514	4705
H(13) [C(13)]		1.10	1428	4964
H(14) [C(14)]		1.04	1445	2797
H(15) [C(15)]		1.04	1025	1751
H(20) [C(20)]		1.05	359	-2154
H(22) [C(22)]		1.04	1797	-2792
H(23) [C(23)]		1.02	1515	-1720
H(24) [C(24)]		1.17	274	-3299
H(25) [C(25)]		1.13	2426	-2547
H(O ¹) [O(1)]		1.12	798	1312
H(O ^{1'}) [O(1)]		0.95	-491	2465
H(C2) [C(2)]		0.64	2145	1240

Table 25 - continued

Atom bonded to	Distance	X	Y	Z	B
H(2') [O(2)]	1.00	3228	2098	1520	5.5
H(3) [O(3)]	0.91	-793	-51	891	5.6
H(3') [O(3)]	0.73	-693	-756	1149	5.6
H(4) [O(4)]	0.92	3049	-428	2050	5.4
H(4') [O(4)]	0.85	2003	-935	1646	5.4
H(5) [O(5)]	0.86	1353	-476	14	5.7
H(5') [O(5)]	0.90	2233	-622	-59	5.7
H(6) [O(6)]	1.15	853	4328	3630	7.4
H(6') [O(6)]	1.13	1345	4724	4067	7.4

aThe hydrogen atom is given followed by the atom to which it is bonded in brackets, the corresponding bond distance (in Å), the positional parameters ($\times 10^{+3}$), and the isotropic thermal parameter (in \AA^2).

Figure 6
An ORTEP drawing of H2diphenylno3)2·2H2O showing the atomic numbering and thermal ellipsoids. The hydrogen atoms are isotropic and small relative to the 50% probability thermal ellipsoids for nonhydrogen atoms.



An ORTEP drawing of $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhpipy})] \cdot 2\text{H}_2\text{O}$ showing atomic numbering and thermal ellipsoids. The hydrogen atoms and uncoordinated water molecules have been omitted.

Figure 7

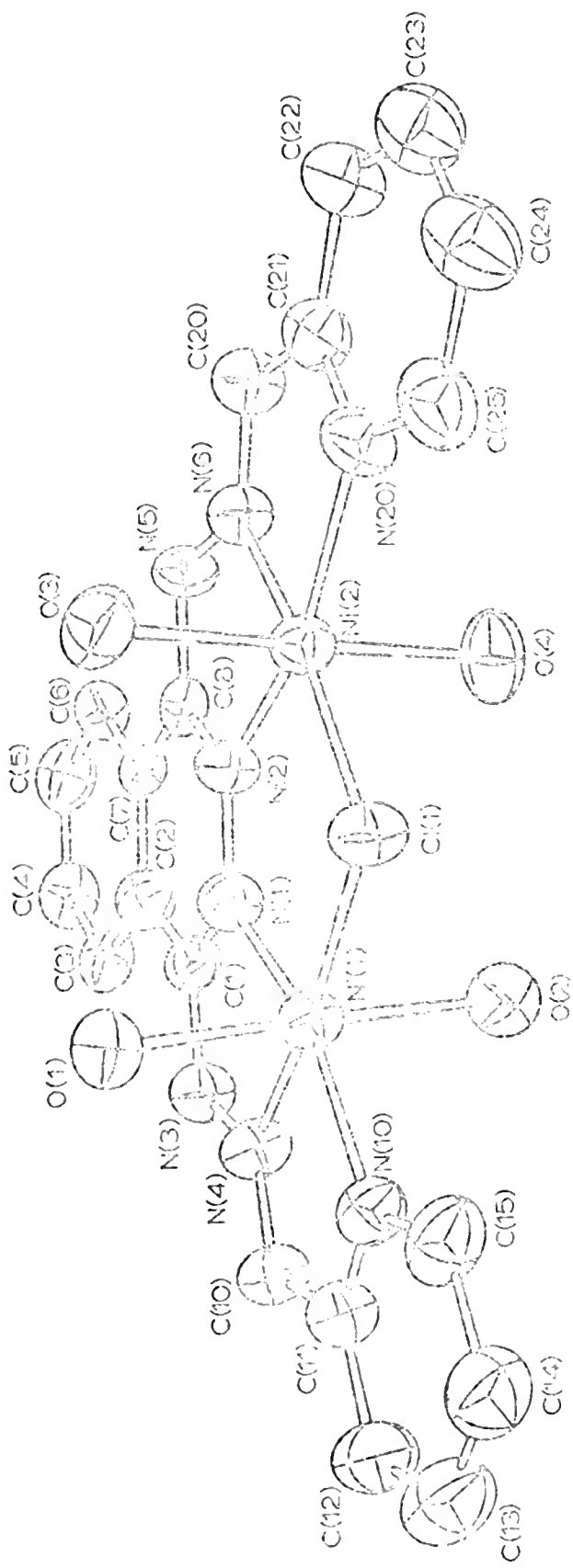
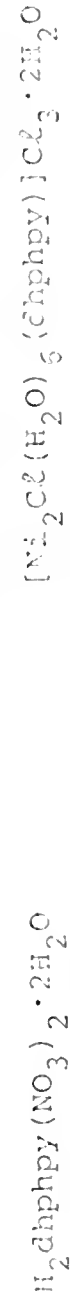


Table 25
Selected Intertatomic Distances for $\text{N}_2\text{C}_2\text{H}_2\text{O}_4$ (Chlorophyll) $\cdot \text{C}_2\text{H}_3 \cdot 2\text{H}_2\text{O}$

(a) Distance (\AA) in the coordination sphere in
 $[\text{Ni}_2\text{Cl}_2(\text{H}_2\text{O})_4\text{ (Chlorophyll)}]\text{C}_2\text{H}_3 \cdot 2\text{H}_2\text{O}$

$\text{Ni}(1)\text{-N}(1)$	2.074(5)
$\text{Ni}(1)\text{-N}(4)$	1.999(5)
$\text{Ni}(1)\text{-N}(10)$	2.074(5)
$\text{Ni}(1)\text{-C}(1)$	2.374(2)
$\text{Ni}(1)\text{-O}(1)$	2.093(5)
$\text{Ni}(1)\text{-O}(2)$	2.117(5)
$\text{Ni}(2)\text{-N}(1)$	2.061(5)
$\text{Ni}(2)\text{-N}(6)$	2.091(5)
$\text{Ni}(2)\text{-N}(2)$	2.029(6)
$\text{Ni}(2)\text{-C}(1)$	2.307(2)
$\text{Ni}(2)\text{-O}(3)$	2.203(6)
$\text{Ni}(2)\text{-O}(4)$	2.170(5)

(b) Distances (\AA) within the Ligand^a



$\text{N}(1)\text{-N}(2)$	2.363(7)
$\text{N}(1)\text{-C}(1)$	1.302(8)
$\text{C}(1)\text{-C}(2)$	1.447(10)
$\text{C}(2)\text{-C}(3)$	1.402(10)
$\text{C}(3)\text{-C}(4)$	1.383(10)
$\text{C}(2)\text{-C}(7)$	1.414(2)
$\text{C}(4)\text{-C}(5)$	1.390(11)
$\text{C}(1)\text{-N}(3)$	1.370(8)
$\text{N}(3)\text{-N}(4)$	1.366(8)
$\text{N}(4)\text{-C}(10)$	1.278(9)
$\text{C}(10)\text{-C}(11)$	1.457(11)
$\text{C}(2)\text{-N}(5)$	1.382(8)
$\text{N}(5)\text{-N}(6)$	1.364(7)
$\text{N}(6)\text{-C}(20)$	1.279(8)
$\text{C}(20)\text{-C}(21)$	1.464(10)

Table 26 - continued

$\text{H}_2\text{Naphpy}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$	$[\text{Ni}_2\text{Cl}_2(\text{H}_2\text{O})_6(\text{dmgpy})]\text{Cl}_2 \cdot 2\text{H}_2\text{O}$
C(11)-C(12)	1.397(5)
C(12)-C(13)	1.387(6)
C(13)-C(14)	1.370(6)
C(14)-C(15)	1.364(6)
C(15)-N(10)	1.331(5)
N(10)-C(11)	1.343(4)
O(1)...N(3)	3.007(4)
O(1)...O(1')	3.283(4)
Ni(1)...Ni(2)	3.603(1)
C(11)-C(12)	1.388(11)
C(12)-C(13)	1.400(13)
C(13)-C(14)	1.373(13)
C(14)-C(15)	1.392(12)
C(15)-N(10)	1.339(10)
N(10)-C(11)	1.351(9)
N(20)-C(21)	1.339(9)
C(21)-C(22)	1.402(10)
C(22)-C(23)	1.370(12)
C(23)-C(24)	1.392(12)
C(24)-C(25)	1.393(11)
C(25)-N(20)	1.349(10)

The estimated standard deviations are given in parentheses.

Selected Angles in $\text{H}_2\text{Naphthy}(\text{NO}_2)_2 \cdot 2\text{H}_2\text{O}$ ^a Table 27

Atoms	Angles (°)	Atoms	Angles (°)
N(1)-C(1)-N(2)	117.9(3)	C(1)-C(2)-C(3)	123.1(3)
N(1)-C(1)-C(2)	121.9(3)	C(2)-C(2')-C(2)-C(3)	119.7(3)
C(1)-C(2)-C(2')	117.2(3)	C(1)-N(1)-N(1')	120.8(3)
C(2)-C(3)-C(4)	119.4(3)	C(2)-C(1)-N(2)	120.1(3)
C(3)-C(4)-C(4')	120.3(4)	N(2)-N(3)-C(10)	117.2(3)
C(1)-N(2)-C(4')	116.9(3)	C(10)-C(11)-N(10)	118.5(3)
C(1)-N(2)-N(3)	118.4(3)	C(11)-C(12)-C(13)	119.0(4)
N(2)-C(10)-C(11)	118.4(3)	C(13)-C(14)-C(15)	121.5(4)
C(10)-C(11)-C(12)	121.2(3)	C(14)-C(15)-N(10)	120.1(4)
C(11)-C(13)-C(14)	119.5(4)	N(10)-C(11)-C(12)	120.3(3)
C(15)-N(10)-C(11)	120.6(3)	O(20)-N(20)-O(21)	118.1(4)
N(1)---O(1)---N(10)	108.3(1)	O(20)-N(22)-O(22)	123.2(5)
N(10)-O(21)---N(2)	123.0(3)		
C(11)-N(20)-O(22)	117.6(4)		

^aApproximate estimated standard deviations are given in parentheses.

Table 28
Selected Angles in $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_6(\text{dhphpy})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}^{\text{a}}$

Atom	Angle	Atom	Angle
N(1)-Ni(1)-Cl(1)	98.0(2)	N(2)-Ni(2)-Cl(1)	97.8(2)
N(1)-Ni(1)-N(4)	76.8(2)	N(2)-Ni(2)-N(6)	76.5(2)
N(1)-Ni(1)-N(10)	155.7(2)	N(2)-Ni(2)-N(20)	154.8(2)
N(1)-Ni(1)-O(1)	91.1(2)	N(2)-Ni(2)-O(3)	93.1(2)
N(1)-Ni(1)-O(2)	90.3(2)	N(2)-Ni(2)-O(4)	89.5(2)
N(4)-Ni(1)-Cl(1)	174.6(2)	N(6)-Ni(2)-Cl(1)	174.1(2)
N(4)-Ni(1)-N(10)	78.9(2)	N(6)-Ni(2)-N(20)	78.2(2)
N(4)-Ni(1)-O(1)	87.8(2)	N(6)-Ni(2)-O(3)	90.4(2)
N(4)-Ni(1)-O(2)	91.1(2)	N(6)-Ni(2)-O(4)	91.2(2)
N(10)-Ni(1)-Cl(1)	106.3(2)	N(20)-Ni(2)-Cl(1)	107.5(2)
N(10)-Ni(1)-O(1)	88.5(2)	N(20)-Ni(2)-O(3)	88.3(2)
N(10)-Ni(1)-O(2)	89.6(2)	N(20)-Ni(2)-O(4)	89.8(2)
O(1)-Ni(1)-Cl(1)	90.9(2)	O(3)-Ni(2)-Cl(1)	92.2(2)
O(1)-Ni(1)-O(2)	178.0(2)	O(3)-Ni(2)-O(1)	177.2(2)
O(2)-Ni(1)-Cl(1)	90.3(2)	O(4)-Ni(2)-Cl(1)	90.5(2)
N(10)-C(11)-C(12)	122.2(7)	N(20)-C(21)-C(22)	121.9(7)
C(11)-C(12)-C(13)	117.8(7)	C(21)-C(22)-C(23)	118.7(7)
C(12)-C(13)-C(14)	120.3(9)	C(22)-C(23)-C(24)	120.2(8)
C(13)-C(14)-C(15)	118.4(8)	C(23)-C(24)-C(25)	117.7(8)
C(14)-C(15)-N(10)	122.3(8)	C(24)-C(25)-N(20)	122.6(7)
C(15)-N(10)-C(11)	119.1(7)	C(25)-N(20)-C(21)	118.8(6)
N(10)-C(11)-C(10)	116.2(7)	N(20)-C(21)-C(20)	116.7(6)
C(12)-C(11)-C(10)	121.6(7)	C(22)-C(21)-C(20)	121.4(7)
C(11)-C(10)-N(4)	114.7(7)	C(21)-C(20)-N(6)	113.8(6)
C(10)-N(4)-N(3)	125.9(6)	C(20)-N(6)-N(5)	123.4(6)
N(4)-N(3)-C(1)	115.8(6)	N(6)-N(5)-C(8)	113.8(5)
N(1)-C(1)-N(3)	115.7(6)	N(2)-C(8)-N(5)	116.3(6)
C(2)-C(1)-N(3)	122.7(6)	C(7)-C(8)-N(5)	121.8(6)
N(1)-C(1)-C(2)	121.6(6)	N(2)-C(8)-C(7)	121.3(6)
C(1)-N(1)-N(2)	121.8(6)	C(8)-N(2)-N(1)	120.8(5)
C(1)-C(2)-C(7)	116.8(6)	C(2)-C(7)-C(3)	117.0(6)

Table 28 - continued

Atom	Angle	Atom	Angle
C(1)-C(2)-C(3)	123.5(6)	C(6)-C(7)-C(8)	123.5(6)
C(2)-C(3)-C(4)	119.7(7)	C(5)-C(6)-C(7)	119.4(6)
C(3)-C(4)-C(5)	120.1(7)	C(4)-C(5)-C(6)	121.6(7)
Ni(1)-N(1)-N(2)	122.4(4)	Ni(2)-N(2)-N(1)	123.3(4)
Ni(1)-N(1)-C(1)	115.8(5)	Ni(2)-N(2)-C(1)	115.9(4)
Ni(1)-N(4)-N(3)	115.9(4)	Ni(2)-N(6)-N(5)	117.3(4)
Ni(1)-N(4)-C(10)	118.2(5)	Ni(2)-N(5)-C(20)	119.1(5)
Ni(1)-N(10)-C(11)	111.9(5)	Ni(2)-N(20)-C(21)	112.1(5)
Ni(1)-N(10)-C(15)	128.9(5)	Ni(2)-N(20)-C(25)	129.1(5)
Ni(1)-C(1)-Ni(2)	98.4(1)		

^aThe estimated standard deviations are given in parentheses.

Hydrogen Bonds in $\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dihpipy})\text{Cl}_3 \cdot 2\text{H}_2\text{O}$

Position of ^a A	Distances (Å) ^b		Angles (°)	
	D-H...A	D-H	H...A	D-H...A
$\text{H}_2\text{Cl}(\text{dhpipy})(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$				
N(2)-H(N2)...O(21)	x,y,z	0.95(4)	1.85(4)	2.773(4)
N(1')-H(PX)...O(1)	x,y,z	1.21(6)	1.57(6)	2.758(4)
O(1)-H(1)...N(1)	x,y,z	0.78(5)	2.12(5)	2.855(4)
O(1)-H(2)...O(21)	x,1-y,1/2+z	0.88(4)	1.98(4)	2.824(4)

$\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dihpipy})2\text{Cl}_3 \cdot 2\text{H}_2\text{O}$

N(3)-H(N3)...Cl(2)	x,y,z	0.91	2.26	3.119(6)
N(5)-H(N5)...Cl(3)	x,y,z	0.97	2.24	3.135(6)
O(1)-H(1)...O(5)	-x,-y,-z	1.12	1.75	2.735(8)
O(1)-H(1')...Cl(4)	-x,y,1/2-z	0.95	2.24	3.150(6)
O(2)-H(2')...Cl(3)	1/2-x,1/2+y,1/2-z	1.00	2.20	3.121(6)
O(3)-H(3')...Cl(3)	-x,y,1/2-z	0.73	2.36	3.075(6)
O(4)-H(4)...Cl(2)	1/2-x,1/2+y,1/2-z	0.92	2.16	3.067(5)
O(5)-H(5)...Cl(1)	x,y,z	0.85	2.41	3.185(6)
O(5)-H(5')...Cl(4)	1/2-x,1/2+y,1/2-z	0.93	2.25	3.098(6)
O(6)-H(6')...Cl(4)	x,y,z	1.23	2.09	3.121(7)

Hydrogen-Hydrogen...Acceptor. D-H is at x,y,z.

The estimated standard deviations are given in parentheses.

ing and hydrogen bonding in $\text{H}_2\text{dhphy}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$ and in $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhphy})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$ are presented in Figures 8 and 9.

The most noticeable difference in the structures of the two dhphy ligands is that $\text{H}_2\text{dhphy}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$ contains a twofold rotation axis while the nickel complex does not. In both cases the ligand is approximately planar (see Table 30). The nickel atoms and the bridging chloride of $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4 \cdot (\text{dhphy})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$ lie slightly "below" the least-squares plane of the ligand (Plane 3) and both hydrazone portions are pivoted generally about an N(3) \cdots N(5) axis with both C(14) and C(24) "above" the plane. However, in the protonated ligand one hydrazone is pivoted "upward" and the other "downward" as required by the twofold axis. Also, the hydrazone "arms" in the nickel complex are drawn toward each other compared to the protonated form as indicated by the bond angles within the "arms." All of the pyridine rings are rotated about the C(n0)-C(n1) bond relative to the phthalazine plane with the pyridine nitrogen atoms tipped toward the coordinated species. In $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhphy})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$ the pyridine containing N(10) is rotated to a much greater extent than that containing N(20). This is shown by the deviations from plane 4 (Table 30) of N(10) and C(12), 0.121 and 0.222 \AA , compared to the deviations of N(20) and C(22), 0.148 and 0.161 \AA . The rings of the phthalazine fragment in each compound appear twisted relative to each other but by less than 2°.

Figure 8
A packing diagram of H₂C₆H₅N₃(NO₂)₂·2H₂O with atoms at X,Y,Z and X',Y',Z' labeled and those at X,1-Y,1/2-Z and X',Y',1/2-Z unlabeled. Proposed hydrogen bonds are indicated by broken lines.

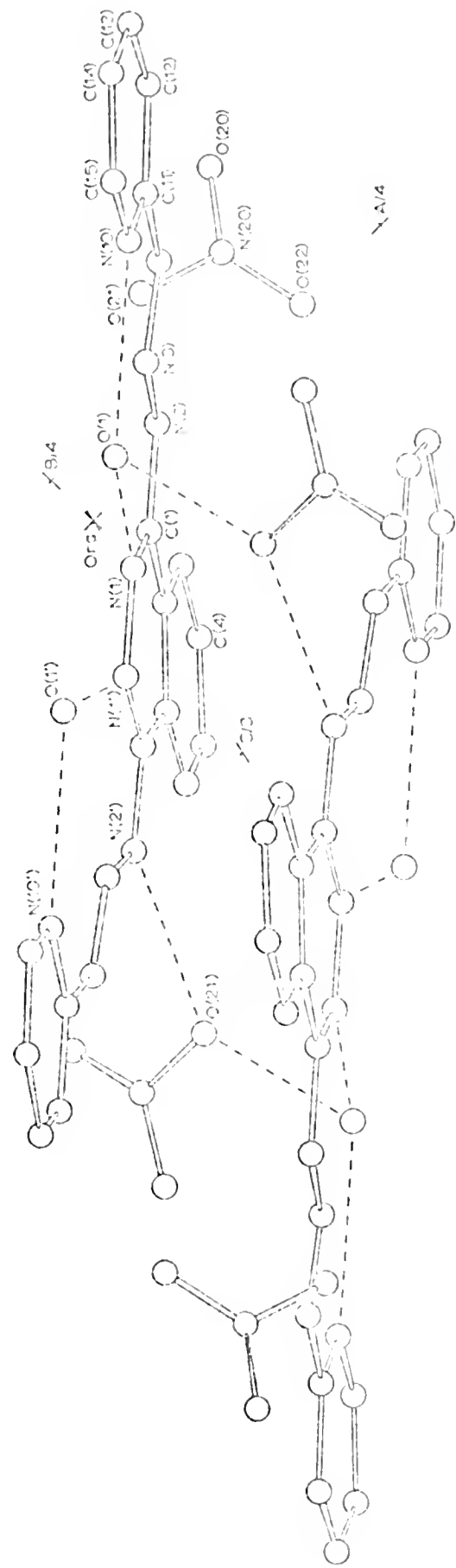


Figure 9
A packing diagram of $[\text{Ni}(\text{C}_2\text{O}_4)_4 \cdot 2\text{H}_2\text{O}]$ (diphopy)₁ where O(5) is at x, y, z ; O(6') is at $\bar{x}, \bar{y}, \bar{z}$; O(6'') is at $1/2-x, 1/2-y, 1/2-z$; and O(6''') is at $1/2+x, 1/2-y, z$. Proposed hydrogen bonds are indicated by broken lines.

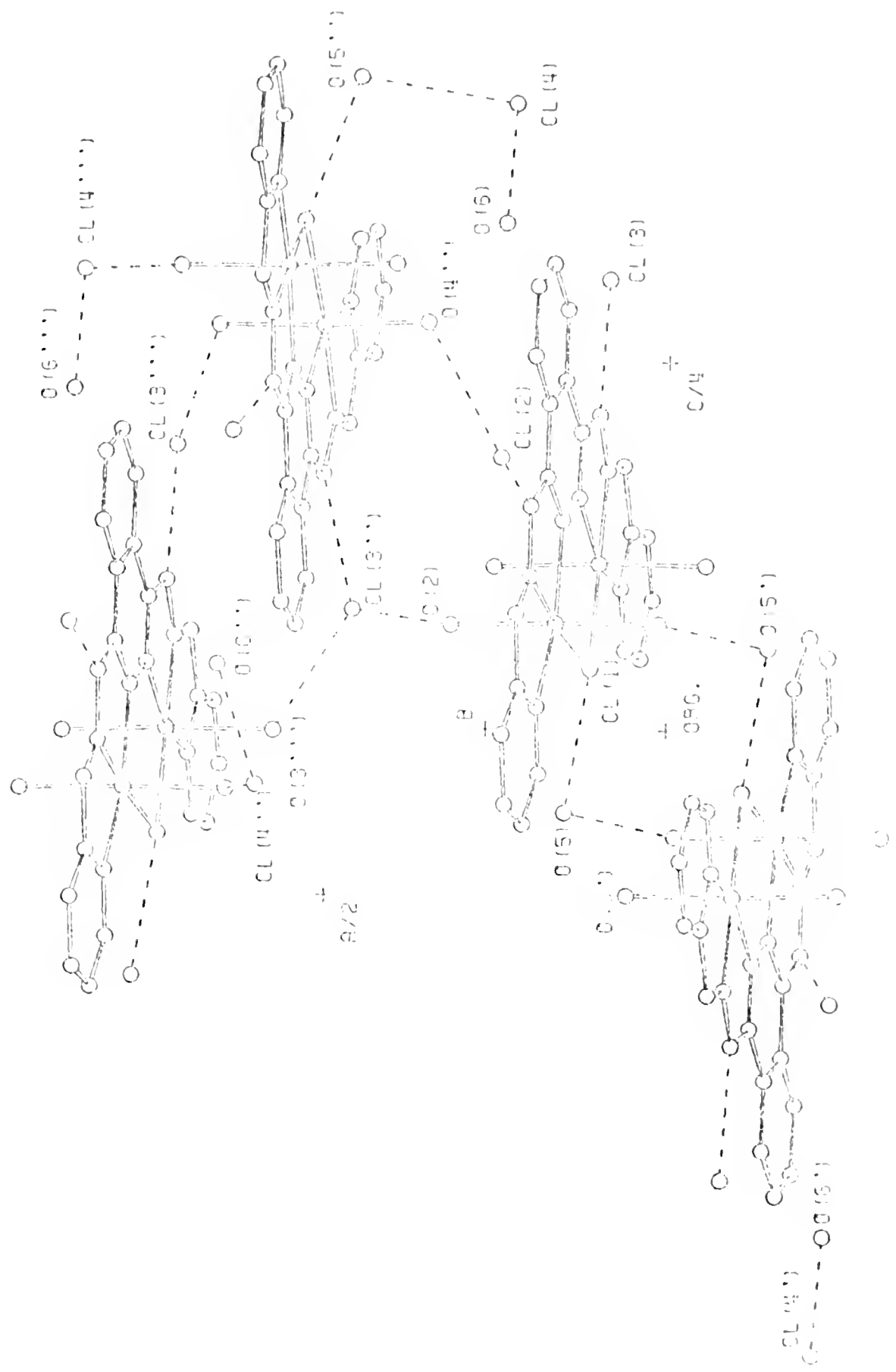


Table 30
Deviations and Equations of Selected Least-Squares Planes in $\text{H}_2\text{daphny}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$
and $[\text{Ni}_2\text{Cl}_6(\text{H}_2\text{O})_4(\text{daphny})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$
(a) Deviations ($\text{\AA} \times 10^{-3}$)
 $\text{H}_2\text{daphny}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$

Atom	Plane 1	Plane 2	Atom	Plane 3	Atom	Plane 4	Atom	Plane 3	Atom	Plane 4
N(1)	-18*	-2*	N(2)	-95*	-6*	N(2)	-92*	-15*		
C(1)	-41*	15*	C(1)	-23*	13*	C(8)	-44*	-5*		
C(2)	-2*	-12*	C(2)	5*	6*	C(7)	12*	17*		
C(3)	-15*	-14*	C(3)	27*	-8*	C(6)	56*	23*		
C(4)	5*	4*	C(4)	73*	-70*	C(5)	72*	-4*		
N(2)	-42*	-19	N(3)	-2*	2*	N(5)	-75*	-39		
N(3)	-26*	20	N(4)	-26*	7*	N(3)	-27*	53		
C(10)	62*	376	C(10)	-14*	34	C(20)	-22*	56		
C(11)	53*	429	C(11)	0*	155	C(21)	-16*	117		
C(12)	5*	448	C(12)	47*	222	C(22)	10*	161		
C(13)	-66*	472	C(13)	63*	283	C(23)	42*	233		
C(14)	-63*	424	C(14)	27*	270	C(24)	76*	296		
C(15)	36*	418	C(15)	-20*	292	C(25)	36*	242		
N(10)	79*	407	N(10)	-54*	124	N(20)	-13*	148		
O(1)	116	243	N(1)	-95	25	N(2)	-90	40		
			C(1)	-232	-55					

Table 30 - continued

(b) Coefficients of the plane equation $Ax + By + Cz = D^{59}$

Plane	A	B	C	D
<u>$\text{N}_2\text{O}_2\text{Cl}_2(\text{N}_2\text{O})_2\text{CH}_2\text{O}$</u>				
1	0.6518	-0.0289	0.7578	1.4465
2	0.7079	-0.0175	0.7061	1.3716
<u>$\text{N}_2\text{O}_2\text{Cl}(\text{N}_2\text{O})_4(\text{CH}_2\text{O})\text{Cl}\text{C}_3\text{H}_2\text{O}$</u>				
3	0.8991	-0.0193	0.4374	1.6470
4	0.9121	-0.0276	0.4091	1.4216

*The entries marked with an asterisk were used to define the plane.

All bonding distances involving nonhydrogen atoms are normal. The N-N distances in both compounds range from 1.363(7) to 1.374(4) Å and are comparable to the N-N distance in 4-FPYTSC of 1.365(3) Å.⁸⁴ Since this distance in both the phthalazine and hydrazone groups is significantly shorter than the accepted N-N single bond distance, 1.4414 Å,⁸⁵ and since the ligand is planar, a delocalized system is presumed to exist. In agreement with this assumption the C(n0)-N distances are longer than the pure C-N double bond distance and are all equivalent to the related C-N distance in 4-FPYTSC, 1.275(3) Å.⁸⁴ All other distances within the ligand are not significantly different from those in [Ni(dhph) (H₂O)₂Cl₄ · 2H₂O].⁸⁶

All Ni-N distances in [Ni₂Cl(H₂O)₄(dhphpy)]Cl₃ · 2H₂O are within the range of reported bonding distances of nickel(II) with aromatic nitrogen atoms (2.00 to 2.112 Å).⁸⁷

The bridging chloride is not symmetrically located between the two nickel atoms with Ni-Cl distances of 2.374(2) and 2.387(2) Å. The appearance of this bridge is remarkably similar to that in di- μ -chloro-sym-trans-dichlorobis-(2,9-dimethyl-1,10-phenanthroline)dinickel(II) · 2chloroform⁸⁸ where the Ni-C distances are 2.378(3) and 2.394(3) Å. Also, the Ni···Ni distance, 3.602(2) Å, and Ni-Cl-Ni angle, 98.0(1)°, in that compound are equivalent to the 3.603(1) Å separation and 98.36(7)° angle in [Ni₂Cl(H₂O)₆(dhphpy)]Cl₃ · 2H₂O. This distance between the nickel atoms is somewhat shorter than

the 3.791(4) Å distance found in the $[\text{Ni}(\text{dph})_2\text{O}_2]_2\text{Cl}_4 \cdot 2\text{H}_2\text{O}$ complex reported by Andrew and Blake⁸⁶ where both bridges are phthalazin nitrogen atoms. The separation between the nickel atoms in the dphpy complex, however, is substantially longer than the Ni-Ni distance of 2.879 Å in the doubly oxo-bridged complex of Hosking, Robson, and Schaap.⁷⁰ All these inter-nickel distances are much greater than twice the covalent radius of nickel and must be a function of the bridging atom.

The distorted octahedral coordination geometry about each nickel atom in $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dphpy})]\text{Cl}_3 \cdot 2\text{H}_2\text{O}$ is completed by two water molecules which lie on a line almost perpendicular to the ligand plane. The Ni-O bond distances are typical⁸⁷ for water coordinated to nickel(II) ranging from 2.070(6) to 2.117(6) Å.

A degree of uncertainty exists concerning the positions of hydrogen atoms about O(1) in $\text{H}_2\text{dphpy}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$. The O(1)-H(1) distance appears to be very short, 0.78 Å, while the N(10)-H(py) distance appears to be very long, 1.21 Å. Although the locations presented for the hydrogen atoms are the most reasonable interpretation of the difference map in terms of peak heights, distances, and H-O-H angles, other areas of positive density exist about the N(1), O(1), and N(10) positions. Disorder may exist with alternate forms having N(1) protonated or having a "coordinated hydronium ion."

Complexes of dphpy structurally provide a promising

uni-molecular system for the incorporation of a small molecule at a bridging position. Dinitrogen has been reported as a bridging ligand connecting two metal complexes in the μ -dinitrogen-bis{[1,2-bis(dimethylphosphino)ethane]hydrido- $[\eta$ -(1,3,5-trimethylbenzene)]molybdenum} cation and similar compounds.⁸⁹ No complex has been reported which could retain its structural integrity after the removal of a bridging dinitrogen. The structures presented here suggest complexes of ligands similar to dliphy may have such a capacity.

CHAPTER 6

MODELS OF PROPOSED INTERMEDIATES FOR THE CATALYZED CYCLIZATION OF ACETYLENES: THE CRYSTAL AND MOLECULAR STRUCTURES OF 1-(π -CYCLOPENTADIENYL)-1-TRIPHENYLPHOSPHINE-2,3,4,5-TETRAKIS(PENTAFLUOROPHENYL)COBALTOLLE AND 1-(π -CYCLOPENTADIENYL)-1-TRIPHENYLPHOSPHINE-2,3,4,5-TETRAKIS(PENTAFLUOROPHENYL)RHODOLLE

The catalysis of the oligomerization of acetylenes by transition metal complexes has been extensively studied.⁹⁰ A reaction mechanism involving a metallocyclopentadiene intermediate has been suggested⁸⁻¹³ for the trimerization of two molecules of acetylene with one of olefin in the presence of $\text{NiBr}_2(\text{tpf})_2$, $\text{Ni}(\text{CO})_2(\text{tpp})_2$, and other nickel catalysts. Metal-containing heterocycles, metallocycles, have been implicated^{14,91-93} as intermediates in the reactions of acetylenes with π -cyclopentadienyl dicarbonyl-metallocycles in which the metal was cobalt, rhodium, or iridium. Yamazaki et al.⁹⁴⁻⁹⁶ on the basis of chemical reactions assigned a metallocyclic structure to a phosphine-containing cobalt complex isolated from the reaction of diphenylacetylene with $\text{Co}(\text{cp})(\text{tpp})\text{I}_2$ and isopropylmagnesium bromide. They also isolated the same product from the reaction of excess diphenylacetylene with $\text{Co}(\text{cp})(\text{tpf})_2$. A preliminary report of the structure of a cobaltacycle formed by the reaction of $\text{Co}(\text{cp})(\text{tpp})(\text{PhC}\equiv\text{CCO}_2\text{Me})$ with dimethyl maleate has been reported.⁵⁷

Rausch and Gastinger¹⁵ prepared $C_4(fph)_4Co(cp)(tpp)$ by the reaction of bis(pentafluorophenyl)acetylene with π -cyclopentadienylcarbonyltriphenylphosphinecobalt. The analogous rhodium compound was prepared by the reaction of the corresponding rhodium compound.¹⁵

Except for one preliminary report⁹⁷ no structural data have been available for cobaltacyclopentadiene metallocycles. Therefore, the X-ray diffraction structural analysis of $C_4(fph)_4Co(cp)(tpp)$ was undertaken. The corresponding rhodacycle was studied for comparison with this cobaltacycle and related compounds.

Structure Solution and Refinement
for $C_4(fph)_4Co(cp)(tpp)$

The heavy atom method was used in which the positions of the cobalt and phosphorus atoms were estimated from a sharpened Patterson function. A Fourier synthesis based on these atoms was used to estimate the positions of eighteen additional atoms. Successive Fourier syntheses revealed the locations of all nonhydrogen atoms in the compound. A difference Fourier synthesis at that point revealed a region between the cobaltacycle: which was of relatively high electron density. Because this density was diffuse no additional atomic positions were estimated before starting refinement, $R = 0.27$. Three cycles of least-squares refinement with individual isotropic thermal parameters reduced R to 0.14. A difference Fourier synthesis again revealed relatively high elec-

tron density in the same location as before.

Because of the discrepancy of the calculated density (1.423 g/cm^3) from the measured density (1.59 g/cm^3), solvent molecules were presumed to be in the crystal. The deep red crystals of the compound were grown from Skelly C¹⁴ which is a saturated hydrocarbon fraction boiling between 83 and 98°C and consisting mainly of n-heptane, C_7H_{16} . If two solvent molecules were in the unit cell the calculated density would be much nearer the measured value at 1.55 g/cm^3 . Several maxima were observed in the difference Fourier synthesis within the region of high electron density. The distances between these points and the angles made by lines connecting them did not reasonably approximate a hydrocarbon chain.

The thermal parameters were converted to their anisotropic equivalent and nine least-squares cycles using a block approximation to the matrix reduced R to 0.077. The shifts of all parameters during the final cycle were less than one-tenth of their respective estimated standard deviations. A difference Fourier synthesis calculated at this stage again suggested the presence of an ill-defined solvent molecule. Although the distribution of the peaks, which were not well resolved, suggested a C₇ or C₈ chain, a closer examination of the distances and angles within the group showed them not to reasonably approximate a hydrocarbon chain.

Six peaks were selected which closely retained their positions in the final Fourier summation before refinement and in the difference Fourier synthesis just discussed.

which seemed the most reasonable in approximately a hydrocarbon chain. These locations were used isotropically as carbon atoms together with the seventy-three refined positions from the third full-matrix least-squares cycle used anisotropically in a structure factor calculation and in three cycles of block approximation least-squares refinement. Although almost all the poorly matched reflections ($|F_{\text{obs}} - F_{\text{calc}}| > 20$) improved, a Fourier synthesis revealed peaks at positions shifted to a less reasonable distribution from the linear hydrocarbon approximation used. The refinement was terminated at this point. An outline of the refinement is presented in Table 5.

Scattering factors for cobalt, phosphorus, fluorine, oxygen, and carbon were taken from Hanson *et al.*²⁹ A list of observed and calculated structure factors is available.¹⁴

Structure Solution and Refinement
for $C_4(\text{fph})_4\text{Rh}(\text{cp})(\text{tpp})$

The method of isomorphous replacement was used for the solution of the structure of $C_4(\text{fph})_4\text{Rh}(\text{cp})(\text{tpp})$. The cell constants of $C_4(\text{fph})_4\text{Co}(\text{cp})(\text{tpp})$ and $C_4(\text{fph})_4\text{Rh}(\text{cp})(\text{tpp})$ as reported in Table 4 are very similar with differences of less than one percent. The positional parameters from the third cycle of full-matrix least-squares refinement for the non-hydrogen atoms in the isomorphous compound $C_4(\text{fph})_4\text{Co}(\text{cp})(\text{tpp})$ were used in a structure factor calculation and a difference Fourier synthesis with the $C_4(\text{fph})_4\text{Ph}(\text{cp})(\text{tpp})$ data. The structure factor calculation resulted in an R of 0.17 and the

difference Fourier synthesis revealed no major structural differences in the two compounds. The same positional parameters were used in an isotropic least-squares refinement of the $C_4(fph)_4Rh(cp)(tpp)$ data. A summary of further refinement is given in Table 5.

A difference Fourier synthesis after refinement suggested the presence of an ill-defined solvent molecule. As in the case of the cobaltacycle the calculated density, 1.479 g/cm^3 , is significantly less than the density of 1.60 g/cm^3 obtained from flotation measurements of the yellow crystals. If two molecules of n-heptane are assumed within the unit cell the calculated density would be 1.60 g/cm^3 .

An attempt to fit a linear molecule to peaks in the difference Fourier synthesis was also unsuccessful and was not pursued.

The scattering factors used were taken from Hartman et al.²⁹ The observed and calculated structures are listed in Table B-5.

Results and Discussion for $C_4(fph)_4Co(cp)(tpp)$ and $C_4(fph)_4Rh(cp)(tpp)$

The final positional and thermal parameters for the nonhydrogen atoms of both $C_4(fph)_4Co(cp)(tpp)$ and $C_4(fph)_4Rh(cp)(tpp)$ are listed in Table 31. The atomic numbering and thermal ellipsoids of the cobaltacycle are shown in Figure 10. The atomic numbering of the rhodacycle is analogous. Selected bond distances and angles for the two compounds are listed in

Table 21
Final Atomic Parameters ($\times 10^4$) for the Nonhydrogen Atoms in $C_4(\text{fph})_4\text{Co}(\text{cp})(\text{tpp})$ and $C_4(\text{fph})_4\text{Ph}(\text{cp})(\text{tpp})$ with Estimated Standard Deviations Given in Parentheses.^a

Atom	x	y	z	β_{11}	β_{22}	β_{33}	β_{12}	β_{13}	β_{23}
Co	482(1) 453(2)	4958(1) 4917(1)	2130(1) 2125(0)	65(1) 62(1)	64(1) 63(1)	24(0) 25(0)	55(2) 54(1)	32(1) 29(1)	46(1) 44(1)
C(1)	1469(8) 1546(9)	4004(8) 4003(8)	2096(5) 2101(5)	78(10) 96(11)	71(8) 71(8)	27(3) 25(3)	70(15) 77(16)	32(10) 36(10)	52(9) 49(9)
C(2)	2439(8) 2532(9)	3318(8) 4207(8)	1903(5) 1901(5)	89(10) 87(10)	72(8) 61(8)	26(3) 26(3)	95(16) 63(15)	51(10) 41(10)	53(9) 43(9)
C(3)	2748(2) 2825(2)	5285(7) 5370(8)	2870(5) 1872(5)	72(9) 70(9)	71(8) 68(8)	27(3) 26(3)	84(15) 65(15)	46(10) 34(9)	51(9) 48(9)
C(4)	1955(8) 2016(3)	5881(7) 5854(8)	2001(5) 2903(5)	78(10) 72(9)	69(8) 75(8)	23(3) 25(3)	52(15) 57(15)	25(9) 27(9)	51(9) 56(9)
C(5)	1050(9) 1091(6)	2864(8) 2877(8)	2093(5) 2092(6)	82(16) 83(14)	72(9) 73(9)	38(4) 37(4)	80(16) 84(16)	60(11) 52(11)	63(10) 63(10)
C(12)	1267(9) 1293(10)	2877(8) 2836(9)	2792(6) 2751(6)	89(11) 94(11)	72(5) 92(10)	32(1) 42(4)	75(16) 84(17)	45(11) 55(12)	67(10) 83(11)
C(13)	889(10) 902(11)	1820(9) 1800(10)	2778(5) 2750(7)	195(14) 127(12)	116(12) 112(11)	51(5) 56(5)	134(21) 115(21)	62(14) 89(14)	117(13) 117(14)
C(14)	263(12) 273(13)	754(9) 735(10)	2043(7) 2027(8)	168(16) 156(15)	87(10) 83(10)	68(5) 73(7)	135(22) 131(22)	117(17) 122(18)	107(14) 125(15)
C(15)	19(12) 35(12)	726(9) 725(9)	1333(6) 1327(7)	151(15) 151(15)	80(10) 74(10)	47(5) 57(5)	115(20) 92(20)	97(14) 92(16)	60(11) 65(13)
C(16)	445(10) 438(11)	1799(8) 1777(9)	1279(6) 1263(6)	126(12) 122(13)	80(9) 82(9)	43(4) 44(5)	110(18) 109(19)	87(13) 84(13)	78(11) 75(11)

Table 32 - continued

Atom	x	y	z	δ_{11}	δ_{22}	δ_{33}	δ_{12}	δ_{13}	δ_{23}
F(12)	1875(5)	3866(5)	3516(3)	148(8)	97(5)	34(2)	21(11)	53(7)	67(6)
	1952(6)	3850(5)	3520(3)	153(6)	98(6)	38(2)	96(11)	54(7)	72(6)
F(13)	1138(7)	1855(6)	3474(4)	215(10)	147(7)	64(3)	164(14)	113(10)	157(9)
	1163(8)	1869(7)	3446(4)	210(11)	160(8)	68(4)	152(16)	101(10)	169(10)
F(14)	-138(8)	-275(6)	2022(5)	272(12)	103(7)	90(4)	162(15)	178(13)	145(9)
	-143(9)	-293(6)	1990(5)	263(13)	100(7)	101(5)	162(14)	158(14)	149(10)
F(15)	-612(7)	-301(5)	622(4)	239(12)	78(6)	65(3)	112(13)	132(10)	62(7)
	-610(6)	-305(6)	602(5)	249(12)	72(6)	70(4)	86(14)	139(12)	52(8)
F(16)	198(6)	1719(5)	670(3)	182(8)	37(5)	37(2)	106(11)	88(8)	54(6)
	188(7)	1725(5)	667(3)	188(9)	36(6)	39(3)	94(12)	79(8)	50(6)
C(21)	3171(9)	3601(8)	1668(5)	106(11)	71(8)	33(4)	87(16)	56(11)	58(10)
	3231(9)	3588(8)	1667(5)	87(10)	74(5)	35(4)	85(16)	51(11)	58(10)
C(22)	4034(10)	3453(9)	2197(6)	223(13)	95(10)	42(4)	135(19)	68(13)	67(11)
	4056(10)	3435(9)	2182(6)	218(12)	66(10)	41(5)	113(19)	60(13)	62(11)
C(23)	4718(12)	2839(11)	1981(7)	145(15)	140(13)	59(6)	190(25)	75(16)	103(15)
	4706(11)	2768(11)	1945(8)	123(14)	114(12)	63(6)	157(22)	67(15)	99(15)
C(24)	4545(12)	2345(11)	1201(8)	154(16)	126(13)	78(7)	198(25)	128(18)	29(16)
	4517(12)	2266(11)	1170(8)	127(15)	143(12)	65(6)	173(23)	92(16)	77(15)
C(25)	3682(11)	2660(9)	647(6)	252(15)	93(11)	48(5)	123(22)	107(15)	58(12)
	3684(12)	2412(10)	635(7)	256(25)	91(11)	45(5)	109(21)	100(15)	50(12)
C(26)	3029(10)	2061(8)	825(6)	17(12)	92(2)	42(2)	102(19)	73(23)	59(12)
	3043(10)	2059(9)	876(6)	100(11)	83(2)	42(4)	95(17)	65(12)	65(11)
N(27)	4256(7)	2923(6)	2957(4)	260(2)	145(7)	47(2)	226(14)	75(3)	102(5)
	4288(7)	3212(6)	2963(4)	172(2)	147(7)	46(2)	264(12)	68(3)	101(3)
N(28)	5532(3)	2681(8)	2501(5)	237(12)	215(10)	31(4)	345(20)	244(12)	252(12)
	5526(2)	2636(8)	2473(5)	226(22)	205(12)	30(4)	337(20)	296(12)	244(12)

Table 31 - continued

Λ_{atom}	α	β	γ	δ_{11}	β_{22}	β_{33}	β_{12}	β_{13}	β_{23}
F(24)	5233 (9) 5155 (9)	1775 (8) 1657 (8)	972 (5) 926 (6)	261 (13) 233 (13)	200 (10) 179 (10)	102 (5) 101 (5)	359 (20) 310 (20)	197 (14) 180 (14)	134 (12) 120 (12)
F(25)	3500 (3) 3478 (8)	1971 (6) 1910 (7)	-124 (4) -144 (4)	251 (12) 238 (12)	151 (8) 148 (8)	65 (4) 63 (4)	210 (16) 200 (17)	176 (11) 164 (11)	97 (9) 90 (9)
F(26)	2165 (6) 2223 (6)	3179 (5) 3177 (6)	329 (3) 332 (3)	177 (9) 167 (9)	111 (6) 118 (6)	37 (2) 38 (2)	143 (12) 150 (13)	79 (8) 77 (8)	71 (7) 75 (7)
C(31)	3869 (9) 3949 (2)	5823 (7) 5834 (8)	1701 (5) 1710 (5)	92 (10) 94 (11)	63 (8) 68 (8)	33 (4) 34 (4)	77 (15) 85 (16)	50 (11) 51 (11)	56 (9) 54 (10)
C(32)	3655 (9) 3746 (10)	5820 (8) 5826 (9)	992 (6) 1002 (6)	193 (12) 108 (12)	84 (9) 89 (10)	37 (4) 37 (4)	67 (17) 91 (18)	57 (12) 61 (12)	61 (11) 69 (11)
C(33)	4687 (11) 4764 (12)	6121 (10) 6138 (10)	782 (7) 805 (7)	161 (15) 153 (15)	119 (12) 106 (11)	55 (5) 51 (5)	151 (22) 125 (22)	135 (16) 114 (15)	110 (14) 96 (13)
C(34)	5991 (11) 6049 (11)	6489 (10) 6498 (10)	1326 (7) 1334 (7)	113 (13) 123 (13)	125 (12) 112 (12)	73 (6) 69 (6)	123 (21) 136 (21)	132 (16) 144 (16)	118 (15) 109 (14)
C(35)	6264 (9) 6328 (10)	6524 (9) 6544 (10)	2038 (7) 2048 (7)	80 (11) 81 (11)	109 (11) 100 (10)	63 (6) 60 (5)	105 (19) 104 (16)	89 (14) 73 (13)	92 (13) 87 (13)
C(36)	5234 (9) 5288 (10)	6200 (9) 6213 (9)	2213 (6) 2221 (6)	104 (11) 94 (11)	92 (10) 90 (10)	39 (4) 37 (4)	111 (18) 84 (18)	59 (12) 42 (12)	72 (11) 63 (11)
F(32)	2390 (6) 2468 (6)	5458 (5) 5439 (6)	454 (3) 459 (3)	127 (7) 122 (7)	147 (7) 147 (7)	43 (3) 40 (3)	129 (12) 125 (12)	63 (7) 56 (7)	107 (7) 104 (7)
F(33)	4392 (7) 4482 (8)	6050 (7) 6073 (7)	81 (4) 94 (4)	225 (11) 212 (11)	199 (9) 194 (10)	67 (4) 65 (4)	212 (17) 204 (17)	174 (11) 166 (11)	169 (10) 158 (10)
F(34)	6973 (7) 7059 (8)	6763 (7) 6827 (8)	1127 (5) 1164 (6)	178 (10) 164 (10)	213 (10) 208 (10)	111 (5) 111 (5)	200 (17) 198 (17)	224 (13) 207 (13)	206 (12) 203 (13)
F(35)	7531 (6) 7607 (5)	6879 (7) 6928 (7)	2557 (5) 2567 (5)	190 (7) 88 (7)	181 (9) 153 (8)	93 (4) 84 (4)	152 (13) 127 (13)	99 (9) 76 (9)	154 (10) 124 (10)

Table 31 - continued

Atom	α	β_{11}	β_{22}	β_{33}	β_{12}	β_{13}	β_{23}
F(35)	5552(5)	6243(6)	2923(3)	195(7)	157(7)	47(3)	125(12)
	5597(6)	6265(6)	2930(4)	197(7)	152(7)	46(3)	127(12)
C(41)	2261(9)	7041(8)	2072(5)	91(10)	75(2)	22(4)	95(16)
	2290(9)	7009(8)	2054(5)	80(10)	72(3)	32(4)	76(15)
C(42)	1472(9)	7178(8)	1482(5)	107(11)	99(9)	31(4)	104(18)
	1474(10)	7113(9)	1470(6)	98(11)	83(9)	34(4)	88(17)
C(43)	1764(10)	8273(9)	1579(6)	137(14)	102(11)	42(5)	159(21)
	1711(12)	8169(10)	1549(7)	150(15)	118(12)	50(5)	157(22)
C(44)	2816(11)	6458(9)	2269(7)	143(14)	79(9)	61(6)	110(20)
	2786(12)	9217(10)	2252(7)	144(14)	91(11)	68(6)	128(21)
C(45)	3644(10)	9168(8)	2245(6)	105(12)	79(9)	45(5)	54(18)
	3639(10)	9135(9)	2618(6)	103(12)	84(10)	45(5)	51(18)
C(46)	3377(8)	3666(8)	2729(5)	79(10)	68(8)	32(4)	54(15)
	3405(6)	8064(8)	2722(6)	94(7)	73(9)	37(4)	72(16)
F(42)	388(5)	6210(5)	778(3)	129(7)	110(6)	35(2)	99(11)
	402(6)	6129(5)	785(3)	127(7)	107(5)	38(2)	95(11)
F(43)	970(7)	8332(6)	986(4)	213(10)	152(8)	74(4)	214(15)
	887(2)	8225(7)	964(5)	18(11)	165(5)	73(4)	204(16)
F(44)	3063(8)	10324(6)	2362(5)	236(11)	96(5)	93(4)	155(14)
	3005(8)	10263(6)	2332(5)	226(12)	93(7)	109(5)	172(15)
F(45)	4746(6)	10156(5)	2532(4)	162(9)	21(7)	61(3)	35(11)
	4715(7)	10148(5)	3495(4)	164(8)	73(6)	64(3)	21(12)
F(46)	4247(5)	8032(5)	3215(3)	100(6)	94(5)	38(2)	68(10)
	4296(5)	8055(5)	3205(3)	100(6)	93(5)	37(2)	64(10)
C(+)	-1476(2)	1442(10)	1962(6)	63(10)	125(11)	49(4)	70(18)
	-1641(9)	4724(11)	1922(6)	65(10)	132(13)	43(5)	72(19)

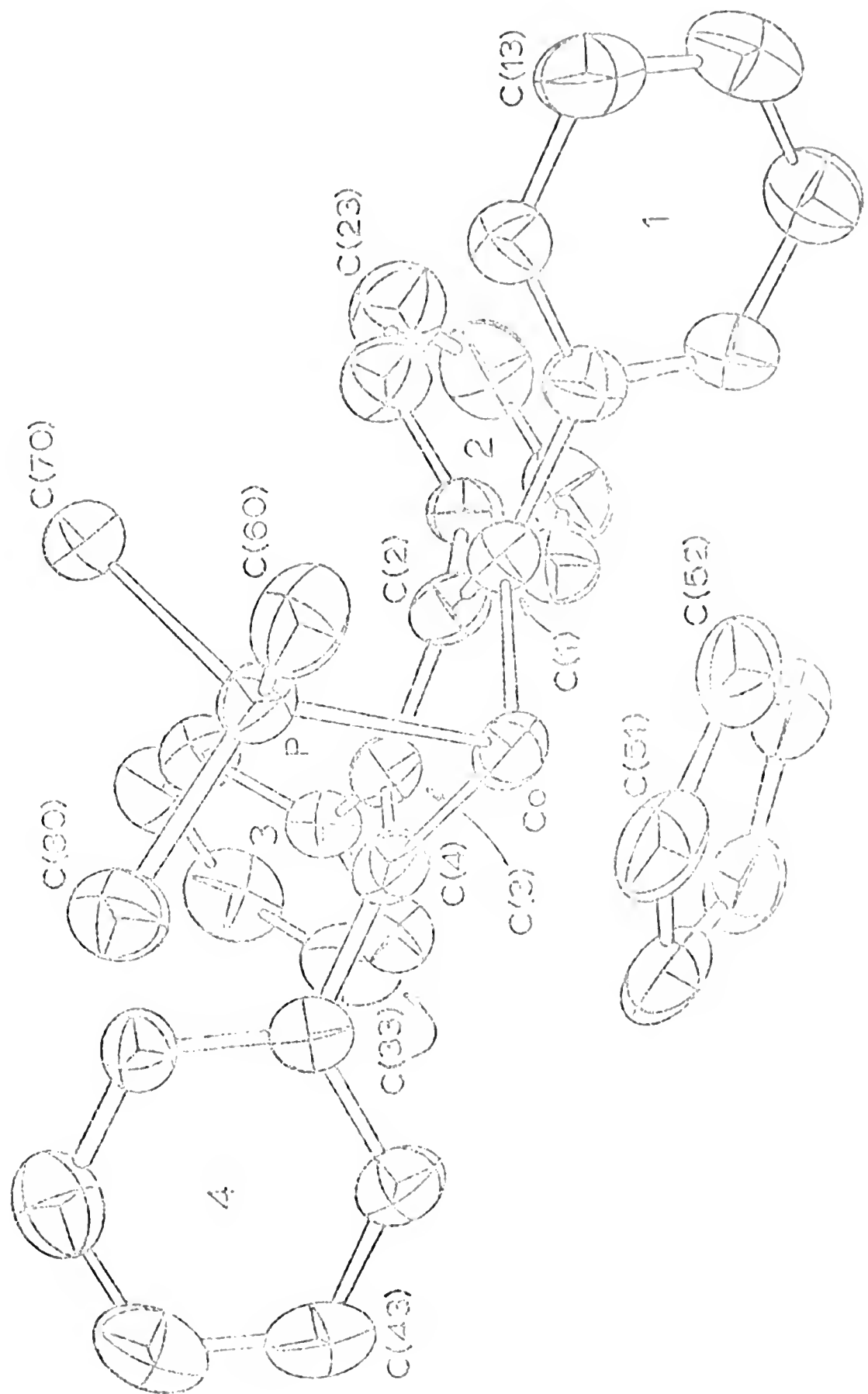
Atom	x	y	z	β_{11}	β_{22}	β_{33}	β_{12}	β_{13}	β_{23}
C(52)	-1486(9)	3709(9)	1756(6)	64(10)	112(11)	50(5)	25(18)	25(12)	90(13)
C(53)	-1672(10)	3638(11)	1750(7)	56(11)	125(13)	56(6)	6(19)	16(13)	103(14)
C(54)	-1196(10)	3326(9)	1216(6)	92(12)	84(10)	24(4)	61(18)	10(12)	41(11)
C(55)	-1405(10)	3198(10)	1083(7)	77(11)	99(10)	48(5)	29(18)	1(12)	54(12)
P	1381(2)	6265(2)	3462(1)	72(3)	71(2)	26(1)	61(4)	39(3)	52(2)
C(60)	174(9)	5972(8)	3855(5)	77(10)	97(10)	30(4)	67(16)	44(10)	61(10)
C(61)	-362(10)	4675(10)	3669(6)	118(13)	117(11)	53(5)	25(20)	67(14)	109(13)
C(62)	-1263(12)	4596(11)	4068(8)	143(15)	158(15)	66(6)	103(25)	119(17)	140(17)
C(63)	-1740(12)	5396(12)	4382(8)	160(17)	160(15)	65(6)	126(26)	134(13)	121(17)
C(64)	-1735(14)	5371(14)	4357(9)	159(18)	171(17)	74(7)	132(29)	136(20)	137(19)
C(65)	-1260(14)	6475(12)	4438(8)	206(20)	158(16)	80(8)	182(30)	202(22)	129(19)
C(70)	2932(9)	6458(8)	4204(5)	188(20)	184(18)	91(9)	213(33)	196(23)	156(22)
C(71)	3110(10)	6734(9)	4984(6)	130(13)	130(12)	59(6)	161(23)	140(16)	105(14)
	3141(11)	6760(10)	5001(6)	111(15)	118(12)	35(4)	147(24)	128(16)	99(15)

Table 31 - continued

Atcn	x	y	z	β_{11}	β_{22}	β_{33}	β_{12}	β_{13}	β_{23}
C(72)	4283(11)	6884(10)	5544(6)	122(14)	222(12)	42(5)	56(21)	24(13)	92(13)
	4315(12)	6902(12)	5555(7)	123(14)	145(24)	42(5)	77(23)	26(24)	102(14)
C(73)	5256(11)	6731(10)	5313(7)	113(14)	128(13)	54(5)	64(22)	22(14)	102(14)
	5271(12)	6770(11)	5344(7)	216(14)	130(13)	52(6)	64(23)	25(14)	101(15)
C(74)	5079(10)	6475(10)	4545(6)	102(13)	116(12)	46(5)	92(20)	27(13)	74(13)
	5119(11)	6506(11)	4572(7)	103(14)	118(13)	52(6)	84(22)	26(14)	78(14)
C(75)	3935(9)	6342(8)	3993(6)	76(10)	85(9)	36(4)	72(16)	32(11)	58(10)
	3933(9)	6358(9)	4925(6)	72(11)	92(10)	37(4)	72(17)	42(11)	59(12)
C(80)	1783(9)	7735(8)	3738(5)	107(11)	77(9)	31(4)	64(16)	65(11)	58(10)
	1836(9)	7761(8)	3744(5)	97(11)	95(9)	32(4)	60(16)	56(11)	66(10)
C(91)	857(9)	7053(8)	3225(6)	190(12)	93(12)	47(4)	215(18)	77(12)	76(12)
	938(10)	7245(9)	3425(6)	147(13)	24(12)	37(4)	2409(18)	65(12)	67(11)
C(92)	1415(11)	9122(10)	3249(7)	244(12)	112(12)	54(5)	154(21)	167(15)	100(15)
	1446(12)	9294(10)	3129(7)	140(15)	64(12)	52(5)	145(21)	67(15)	50(15)
C(93)	2295(11)	30114(10)	4154(7)	240(15)	94(12)	52(5)	109(21)	102(15)	91(12)
	2327(11)	30232(10)	4124(7)	169(16)	86(12)	53(5)	145(21)	66(15)	77(12)
C(94)	3215(11)	2048(9)	4655(7)	157(14)	64(12)	53(5)	50(12)	51(12)	51(12)
	3211(12)	2921(10)	4638(7)	153(15)	61(12)	55(6)	55(12)	67(15)	70(12)
C(95)	2962(10)	6774(9)	4465(6)	223(12)	63(12)	58(4)	55(12)	53(12)	52(11)
	2960(10)	6797(9)	4456(5)	125(12)	81(12)	42(5)	69(12)	54(12)	57(11)

The coordinates of the atoms in each structure are listed in order of the metal atom.

figure 10
An ORTEP drawing of $C_4(fph)_4Co(cp)(tpp)$ showing the atomic numbering and thermal ellipsoids. The fph rings are numbered 1-4 and the fluorines have been omitted for clarity. Similarly, the three phenyl rings of the tpp ligand have been omitted with only the first atoms C(60), C(70), and C(80) shown.



in Tables 32 and 33. Least-squares planes and deviations are given in Table 34.

The molecules are metallocycles with the metal atom also bonded to the cyclopentadienyl ring and to the triphenylphosphine ligand. The C(1) to C(4) fragment in both compounds is planar with the largest deviation from the best plane being 0.015 Å in the cobalt compound and 0.017 Å in the rhodium compound. The metal atoms, however, are significantly displaced from the plane in the direction of the cp ring by -0.203 and -0.239 Å. This perpendicular displacement is similar to that found in other similar metallocycles.⁹⁸

The metallocycles may be considered as a delocalized diene with the metal atom σ-bonded to the two carbon atoms of the ring, C(1) and C(4). The Co-C bond distances, 1.995 (11) and 1.993(11) Å, and the Rh-C bond distances, 2.060(12) and 2.007(11) Å, are similar to various values given by Churchill.⁹⁹ Values of 1.979(1) Å⁴⁸ and 1.990(5) Å⁵¹ have more recently been reported for Co-C bonds in cobaloxime complexes. Mague^{100,101} has reported structures of similar rhodacycles in which the Rh-C distances are 2.000(11), 1.964 (11), 2.047(16), and 1.998(16) Å. Also, Cotton and Norman¹⁰² report a single-bond covalent radius of 1.29 Å for Rh(III). When this value is added to half the 1.485 Å suggested length for a single-bond between sp^2 carbon atoms¹⁰³ the Rh-C distance is predicted to be 2.13 Å. The observed Rh-C distances where rhodium has a formal oxidation number of +1 are shorter than the above predicted single-bond distance. This differ-

Table 32
Selected Bond Distances (\AA) of $\text{C}_4(\text{fph})_4\text{M}(\text{Cl})(\text{tpp})$ ($\text{M}=\text{Co}, \text{Rh}$)
with Their Estimated Standard Deviations in Parentheses.

$\text{M} = \text{Co}$	Rh
$\text{M}-\text{C}(1)$	1.995(11)
$\text{M}-\text{C}(4)$	1.993(11)
$\text{M}-\text{P}$	2.234(3)
$\text{M}-\text{C}(51)$	2.157(12)
$\text{M}-\text{C}(52)$	2.121(13)
$\text{M}-\text{C}(53)$	2.119(11)
$\text{M}-\text{C}(54)$	2.104(9)
$\text{M}-\text{C}(55)$	2.133(12)
$\text{C}(1)-\text{C}(2)$	1.326(15)
$\text{C}(2)-\text{C}(3)$	1.467(16)
$\text{C}(3)-\text{C}(4)$	1.335(15)
$\text{C}(1)-\text{C}(11)$	1.487(16)
$\text{C}(2)-\text{C}(21)$	1.523(16)
$\text{C}(3)-\text{C}(31)$	1.481(15)
$\text{C}(4)-\text{C}(41)$	1.493(16)
$\text{P}-\text{C}(60)$	1.848(11)
$\text{P}-\text{C}(70)$	1.843(11)
$\text{P}-\text{C}(80)$	1.834(12)
$\text{C}(51)-\text{C}(52)$	1.463(20)
$\text{C}(52)-\text{C}(53)$	1.400(16)
$\text{C}(53)-\text{C}(54)$	1.426(18)
$\text{C}(54)-\text{C}(55)$	1.433(16)
$\text{C}(55)-\text{C}(51)$	1.457(17)
	1.431(18)

Table 33

Selected Bond Angles ($^{\circ}$) of $C_4(fph)_4M(cp)(tpp)$ with Their Estimated Standard Deviations Given in Parentheses. (M=Co, Rh)

M = Co	Rh
M-C(1)-C(2)	112.1(8)
C(1)-C(2)-C(3)	116.8(9)
C(2)-C(3)-C(4)	114.8(9)
M-C(4)-C(3)	113.1(7)
C(1)-M-C(4)	82.4(4)
P-M-C(1)	103.0(3)
P-M-C(4)	95.2(3)
C(11)-C(1)-M	127.0(7)
C(11)-C(1)-C(2)	119.6(9)
C(21)-C(2)-C(1)	123.9(9)
C(21)-C(2)-C(3)	119.2(9)
C(31)-C(3)-C(2)	119.7(9)
C(31)-C(3)-C(4)	125.5(9)
C(41)-C(4)-C(3)	119.8(9)
C(41)-C(4)-M	127.0(7)
C(51)-C(52)-C(53)	108.1(11)
C(52)-C(53)-C(54)	109.8(10)
C(53)-C(54)-C(55)	107.7(10)
C(54)-C(55)-C(51)	108.0(10)
C(55)-C(51)-C(52)	106.3(10)

Table 34

Deviations from and Equations of Some Least-Squares Planes of
 $C_4(fph)_4Co(cp)(tpp)$ and $C_4(fph)_4Rh(cp)(tpp)$.^a

(a) Deviations ($\text{\AA} \times 10^{+3}$)

Atom	Plane 1	Plane 2	Plane 3	Plane 4
Co	-203		1741	
Rh		-239		1908
C(1)	8*	9*		
C(2)	-15*	-17*		
C(3)	14*	17*		
C(4)	-8*	-9*		
C(51)	-931	-1058	-7*	2*
C(52)	-1211	-1307	15*	6*
C(53)	-2043	-2168	-16*	-12*
C(54)	-2265	-2437	11*	13*
C(55)	-1598	-1774	-2*	-9*
F	1884	1922	3025	3241

(b) Coefficients of the Plane⁵⁹ AX + BY + CZ = D

Plane	A	B	C	D
1	0.2201	0.0627	0.9735	3.2807
2	0.2193	0.0672	0.9733	3.3160
3	0.7356	0.1345	0.6639	-0.8420
4	0.7474	0.1591	0.6450	-1.0481

^aThe entries marked with an asterisk were used to define the plane.

ence could be indicative of multiple bonding between the terminal carbon atoms of the diene and the metal atom. The C-C distances in the metallocycle rings fall into two groups. The C(1)-C(2) and C(3)-C(4) distances are equal within experimental error to the accepted value of 1.337(6) Å for a simple C-C double bond.¹⁰⁴ The C(2)-C(3) distances are indicative of a C-C single bond between two double bonds.¹⁰⁴ The observations of Mague^{100,101} on two rhodacycles suggested a double-bond system similar to those in $C_4(fph)_4Co(cp)(tpp)$ and $C_4(fph)_4Rh(cp)(tpp)$.

The cp rings in the compounds are planar with the maximum deviations from the least-squares planes of -0.016 and -0.012 Å. The distances from the cp ring atoms to the metal atom show that the metal atom is slightly displaced from the center of the cp ring. The range of the Co-C(cp ring) distances is from 2.104(9) to 2.157(12) Å with a mean of 2.127(9) Å. These values are similar to those in other Co-cp complexes.^{105,106}

In both the cobalt and rhodium compounds the longest metal-C(cp ring) distance involves C(51), the carbon atom nearest the phosphine ligand. The mean Rh-C(cp ring) distance is 2.286(13) Å. This value is equivalent to the mean distance of 2.246(9) Å in $Rh(C_2F_5)(cp)I(CO)$ ¹⁰⁷ and falls within the 2.19 to 2.26 Å range reported for corresponding mean values for other cp-rhodium complexes.¹⁰⁸

The C-C bond distances within the cp rings range from 1.400(16) to 1.463(20) Å with a mean of 1.436(11) Å in the

cobalt compound and a range from 1.420(17) to 1.431(18) Å with a mean of 1.425 Å in the rhodium compound. These C-C distances are comparable to those found in other cp complexes.^{105,106,109} The cp rings are tipped relative to the C(1) to C(4) planes by 35.3° and 36.6°.

The Co-P distance of 2.234(3) Å is similar to the Co-P distance in five-coordinate complexes of cobalt where the range is reported¹¹⁰ to be from 2.192(6) to 2.27(1) Å. Also, in cobalt-carbonyl complexes such as $\text{Co}_4(\text{CO})_{10}(\text{Ph}_2\text{PC-CCF}_3)_2$ and $\text{Co}(\text{CO})_2(\text{NO})(\text{tpp})$ the Co-P distances are 2.236 and 2.229 Å¹¹¹ in the former and 2.224(3) and 2.230(3) Å¹¹² in the latter. The Rh-P distance of 2.293(?) Å is similar to those in phosphine complexes of rhodium(I).¹¹³ The metal to phosphine distance in metal-oxime complexes have been found to be somewhat longer.^{40,97} The Co-P distance in cobaloxime complexes has been reported as 2.327(4) Å⁴⁰ and 2.339(1) Å.⁴⁸ The Rh-P distance in $\text{RhCl}(\text{Hdmg})_2(\text{tpp})$ was reported to be 2.327(1) Å.¹⁰² Since the distances in oxime complexes in both cobalt and rhodium are equivalent, the phosphorus atom may be in the position of closest approach to the metal atom as limited by the steric constraints of the oxime ligands.

The distances in the fph rings have been summarized in Table 35. The individual values for the distances and angles in the fph rings on the metallocycles and the phenyl rings of the phosphines are given in Tables 36-38. The dimensions are not unusual and are in agreement with expected values.

Table 35

Average C-F and C-C Distances for the Pentafluorophenyl Groups in $C_4(\text{fph})_4\text{M}(\text{cp})(\text{tpp})$ with Estimated Standard Deviations Given in Parentheses (M=CO, Rh).

M =	(a) C-F Distances (Å)		(b) C-C Distances (Å)	
	CO	Rh	CO	Rh
All Rings	1.344(2)		1.344(2)	1.378(3)
Ring 1	1.342(6)		1.340(3)	1.385(1)
Ring 2	1.345(4)		1.347(4)	1.373(4)
Ring 3	1.340(3)		1.345(3)	1.384(3)
Ring 4	1.348(4)		1.346(3)	1.372(8)

a Standard deviations were estimated using the equation:

$$\sigma = \left[\sum_{i=1}^N (x_i - \bar{x})^2 / N(N-1) \right]^{1/2}$$

Table 36
Bond Distances and Bond Angles of Pentafluorophenyl Groups
in $C_4(fph)_4Rh(cp)(tpp)$.

(a) Distances (\AA)

n =	1	2	3	4
Cn1-Cn2	1.384(15)	1.342(16)	1.392(15)	1.385(15)
Cn2-Cn3	1.364(20)	1.400(20)	1.374(20)	1.351(20)
Cn3-Cn4	1.375(18)	1.358(18)	1.357(19)	1.389(18)
Cn4-Cn5	1.367(19)	1.365(20)	1.368(18)	1.355(19)
Cn5-Cn6	1.372(20)	1.373(19)	1.367(18)	1.362(19)
Cn6-Cn1	1.393(15)	1.389(14)	1.389(16)	1.386(14)
Cn2-Fn2	1.347(12)	1.354(12)	1.351(13)	1.344(12)
Cn3-Fn3	1.339(15)	1.341(18)	1.349(15)	1.348(16)
Cn4-Fn4	1.338(18)	1.337(19)	1.335(18)	1.340(18)
Cn5-Fn5	1.343(15)	1.358(14)	1.338(15)	1.357(14)
Cn6-Fn6	1.331(13)	1.343(14)	1.351(13)	1.342(13)

(b) Angles ($^\circ$)

Cn1-Cn2-Cn3	123.1(11)	122.4(12)	123.7(11)	122.9(11)
Cn2-Cn3-Cn4	119.6(13)	119.3(13)	118.8(13)	120.2(13)
Cn3-Cn4-Cn5	119.6(13)	119.3(14)	120.8(13)	117.9(13)
Cn4-Cn5-Cn6	119.8(13)	120.6(13)	119.0(12)	121.4(12)
Cn5-Cn6-Cn1	122.6(12)	121.1(11)	123.6(11)	122.0(11)
Cn6-Cn1-Cn2	115.3(11)	117.3(11)	114.1(10)	115.4(10)
Cn -Cn1-Cn2	124.2(10)	123.9(10)	123.1(10)	124.1(10)
Cn -Cn1-Cn6	120.5(10)	118.8(10)	122.5(10)	120.5(10)
Fn2-Cn2-Cn1	120.2(10)	121.3(11)	118.2(10)	119.4(10)
Fn2-Cn2-Cn3	116.7(11)	116.4(11)	118.0(11)	117.7(11)
Fn3-Cn3-Cn2	120.9(12)	120.7(13)	120.3(12)	120.8(12)
Fn3-Cn3-Cn4	119.5(12)	120.0(13)	120.9(12)	119.0(12)
Fn4-Cn4-Cn3	120.7(13)	121.3(14)	119.9(13)	119.9(12)
Fn4-Cn4-Cn5	119.7(13)	119.4(13)	119.3(13)	122.1(13)
Fn5-Cn5-Cn4	120.9(13)	120.1(13)	119.6(12)	118.8(12)

Table 36 - continued

n =	1	2	3	4
Fn6-Cn6-Cn5	115.7(10)	118.3(11)	117.6(10)	117.2(9)
Fn6-Cn6-Cn1	120.4(10)	118.8(10)	118.5(10)	119.7(9)

Table 37
Bond Distances and Bond Angles of Pentafluorophenyl Groups
in $C_4(fph)_4Co(cp)(tpp)$.

	(a) Distances (\AA)			
n =	1	2	3	4
Cn1-Cn2	1.387(14)	1.372(16)	1.394(14)	1.403(15)
Cn2-Cn3	1.388(19)	1.368(20)	1.398(19)	1.358(19)
Cn3-Cn4	1.387(17)	1.374(18)	1.370(18)	1.348(16)
Cn4-Cn5	1.382(17)	1.374(20)	1.372(18)	1.370(17)
Cn5-Cn6	1.382(18)	1.363(19)	1.362(18)	1.384(17)
Cn6-Cn1	1.385(14)	1.389(14)	1.408(15)	1.367(14)
Cn2-Fn2	1.322(11)	1.341(12)	1.339(13)	1.358(11)
Cn3-Fn3	1.350(14)	1.338(17)	1.339(14)	1.338(14)
Cn4-Fn4	1.360(17)	1.339(19)	1.334(17)	1.335(16)
Cn5-Fn5	1.336(13)	1.354(13)	1.330(14)	1.361(13)
Cn6-Fn6	1.341(12)	1.355(13)	1.356(12)	1.348(12)
	(b) Angles ($^\circ$)			
Cn1-Cn2-Cn3	122.4(11)	122.9(12)	123.4(11)	122.6(10)
Cn2-Cn3-Cn4	119.7(12)	119.2(13)	118.9(12)	120.5(12)
Cn3-Cn4-Cn5	119.6(12)	120.1(14)	120.2(13)	118.7(12)
Cn4-Cn5-Cn6	118.7(12)	119.0(12)	119.8(12)	120.4(11)
Cn5-Cn6-Cn1	123.9(11)	122.9(11)	123.8(11)	123.0(10)
Cn6-Cn1-Cn2	115.6(10)	116.0(10)	113.9(10)	114.5(10)
Cn -Cn1-Cn2	123.3(10)	123.8(10)	122.9(9)	124.2(9)
Cn -Cn1-Cn6	121.0(10)	120.2(10)	123.0(9)	121.3(9)
Fn2-Cn2-Cn1	121.4(10)	120.5(10)	119.1(10)	119.8(9)
Fn2-Cn2-Cn3	116.2(10)	116.6(11)	117.5(10)	117.7(10)
Fn3-Cn3-Cn3	120.4(11)	122.0(13)	119.9(11)	119.1(11)
Fn3-Cn3-Cn4	119.9(11)	118.8(13)	121.2(12)	120.4(11)
Fn4-Cn4-Cn3	120.0(12)	121.2(13)	118.9(12)	120.1(11)
Fn4-Cn4-Cn5	120.4(12)	118.7(13)	120.9(12)	121.2(11)
Fn5-Cn5-Cn4	119.7(11)	120.5(12)	119.9(12)	119.8(11)
Fn5-Cn5-Cn6	121.6(11)	120.5(12)	120.3(11)	119.8(10)

Table 37 - continued

n =	1	2	3	4
Fn5-Cn5-Cn6	119.4(12)	119.3(12)	121.4(12)	119.8(11)
Fn6-Cn6-Cn5	117.7(11)	119.0(11)	117.6(11)	117.9(10)
Fn6-Cn6-Cn1	119.7(11)	119.9(10)	118.9(10)	120.1(10)

Table 38
Bond Distances and Bond Angles of Triphenylphosphine in $C_4(\text{fPh})_4\text{M}(\text{cp})(\text{tPP})$.

(a) Distances (\AA)		n = 5				n = 6				n = 7				n = 8			
		Co	Rh														
P-C(n0)		1.348(11)		1.353(12)		1.843(11)		1.821(10)		1.834(12)		1.820(13)		1.820(12)		1.820(13)	
C(n0)-C(n1)		1.419(19)		1.400(20)		1.411(14)		1.387(17)		1.416(16)		1.395(16)		1.395(16)		1.395(16)	
C(n1)-C(n2)		1.414(20)		1.382(22)		1.394(17)		1.399(19)		1.413(19)		1.413(20)		1.413(20)		1.413(20)	
C(n2)-C(n3)		1.397(22)		1.372(25)		1.386(19)		1.347(21)		1.416(17)		1.413(19)		1.413(19)		1.413(19)	
C(n3)-C(n4)		1.394(25)		1.396(30)		1.390(17)		1.405(18)		1.422(19)		1.406(21)		1.406(21)		1.406(21)	
C(n4)-C(n5)		1.423(23)		1.367(25)		1.376(16)		1.403(18)		1.391(19)		1.380(21)		1.380(21)		1.380(21)	
C(n5)-C(n0)		1.387(18)		1.377(20)		1.394(16)		1.368(16)		1.413(14)		1.404(15)		1.404(15)		1.404(15)	
(b) Angles ($^\circ$)		n = 5				n = 6				n = 7				n = 8			
P-C(n0)-C(n1)		117.7(8)		117.4(9)		120.3(8)		121.4(9)		118.3(8)		118.5(9)		118.3(8)		118.5(9)	
P-C(n0)-C(n5)		122.2(9)		122.2(10)		119.6(8)		129.5(8)		121.8(8)		122.3(9)		122.3(9)		122.3(9)	
C(n0)-C(n1)-C(n2)		119.7(12)		120.1(13)		119.9(11)		120.3(12)		119.5(11)		121.0(11)		121.0(11)		121.0(11)	
C(n1)-C(n2)-C(n3)		120.0(13)		119.7(15)		119.0(12)		120.3(14)		120.7(12)		119.5(12)		119.5(12)		119.5(12)	
C(n2)-C(n3)-C(n4)		120.2(14)		119.2(15)		120.8(13)		120.8(14)		118.7(12)		118.2(13)		118.2(13)		118.2(13)	
C(n3)-C(n4)-C(n5)		120.2(14)		121.8(17)		120.8(12)		118.2(13)		120.8(12)		122.0(13)		122.0(13)		122.0(13)	
C(n4)-C(n5)-C(n0)		119.8(13)		118.8(14)		119.2(11)		121.3(11)		120.3(11)		120.1(12)		120.1(12)		120.1(12)	
C(n5)-C(n0)-C(n1)		120.0(11)		120.4(12)		120.1(10)		119.1(11)		119.9(10)		119.1(11)		119.1(11)		119.1(11)	

Table 38 - continued

$M = Co$	Rh
C(60)-P-C(70)	102.7 (5)
C(60)-P-C(80)	100.9 (5)
C(70)-P-C(80)	103.0 (5)
	103.5 (5)
	101.8 (5)
	103.8 (5)

The fluorinated metallocycles resist thermal decomposition better than the hydrocarbon analogs.^{14,15} Enhanced thermal stabilities have been observed in other highly fluorinated metallocycles relative to their hydrocarbon analogs.¹¹⁴ In the compounds of this study the triphenylphosphine ligand and the four fph rings provide an effective shield for the two double bonds in the metallocycles. Although the fluorine atoms of the fph rings and the phenyl rings of the tpp were omitted from Figure 10, the sterically hindered nature of the metallocycle may easily be seen. The lack of a convenient path for an attacking acetylene together with the enhanced thermal stability of the fluorinated derivatives may have allowed the isolation of these intermediate metallocycles. Metallocycles of cobalt and rhodium of the type presented are reasonable intermediates in the catalyzed oligomerization of acetylenes.

CHAPTER 7 CONCLUDING REMARKS

The structure of $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{clan})$ shows the same LIPS phenomenon as $\text{ClCo}(\text{H}_2\text{dmg})(\text{dmg})(\text{sulfa})$.⁴⁶ These two compounds exhibit the unusual feature of containing both neutral and dianionic dimethylglyoxime groups. Also, the orientation of the benzene ring of the sulfa and clan group in the respective compounds is over the dianionic dmg. The various distances and the relative orientation of the axial ligand in both compounds suggest a π -type interaction. LIPS supports the contention that "hydrophobic forces" are important in enzymic processes.³ The bis(diglyoximato)cobalt(III) complexes of aniline derivatives have here been shown to be useful models for the examination of this type interaction. An extension of X-ray structural determinations to similar compounds with other aniline derivatives and with other diglyoximes is suggested. Low-temperature X-ray studies could effect better resolution of the inter-dmg bridge structure and the N-O distances.

An investigation of the fluorescence spectra of these compounds could reveal additional information concerning the interaction between the equatorial and axial ligands. The fluorescence of 5-dimethylaminonaphthalene-1-sulfonamide was observed to be enhanced while the fluorescence of carbamic

anhydrase was diminished when a 1:1 complex of the two was formed.⁵¹ Although the major contribution to this observation is believed to be the ionization of the sulfonamide, a portion of the change is attributed to a hydrophobic interaction.^{51,115} The fluorescence spectra of cobaloxime complexes with aniline derivatives should help reveal the nature of the interligand interaction as a function of the orientation angle.

The novel ligand dhphy has been demonstrated as a bi-nucleating ligand. The bridging site occupied by a chlorine atom in $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4(\text{dhphy})]\text{Cl}_3$ clearly is accessible and of convenient dimensions to accommodate a molecule such as dinitrogen. Further development of this system as a possible model for nitrogenase should include use of molybdenum salts and work with the exclusion of oxygen. Synthesis of similar ligands with saturated "side arms" is also suggested.

The compounds $\text{C}_4(\text{fph})_4\text{Co}(\text{cp})(\text{tpp})$ and $\text{C}_4(\text{fph})_4\text{Rh}(\text{cp})(\text{tpp})$ contain a butadiene fragment with each end bound to a metal atom. The metal to carbon bonds are shorter than expected for the single-bonded distance. The metallocycles are, therefore, believed to contain a delocalized π -bonding system. While metallocycles should be highly susceptible to nucleophilic attack and thermal decomposition the two compounds studied here are very stable. The enhancement of thermal stability by the fluorinated substituents may be at least partially responsible. Also, the presence of the four fph rings

along with the tpp and cp ligands provides a shield from attack for the metallocycle.

The understanding of catalytic processes should improve the efficiency of our existence. Hopefully, enzymic processes occurring in nature can be duplicated in the laboratory by suitable models. These model enzyme systems may then be applied to cure the diseased and feed the hungry.

APPENDIX A BOOTHITL

A listing of the FORTRAN language computer program BOOTHITL follows. This program was designed to interpolate atomic positional parameters by Booth's method¹¹⁶ from the values of a Fourier synthesis calculation. The Fourier synthesis program written by Dr. Gus J. Palenik was modified to store the calculated values on a magnetic disk. After supplying BOOTHITL with input data of the approximate position of each atom, the stored values are retrieved. The program estimates the position of maximum electron density for each atom from these Fourier synthesis values. The positional parameters may be translated to equivalent positions and may be passed to a bond distance and angle program. The resulting fractional coordinates are punched into IBM cards in the format required for their input into the Fourier synthesis and least-squares refinement programs.

ROUTINE

THIS IS A SUBROUTINE WHICH TAKES ATOMIC POSITIONS DIRECTLY FROM HUMAN CALCULATIONS STORED ON DISK

SUMMARY OF THE ROUTINE FUNCTIONS

CARD 1. TITLE

CARD 2. NAME NAMES, IR, IS, IT, ISEL, ICEN, NORD, IDISP AS IN FIGURE. THESE ATTRIBUTES OF ATOMS IS CREATED THAN ZEROS. THE ATOMIC PARAS ARE THEN ADDED UP AND WRITTEN ONTO UNIT 04 TO BE PASSED ON TO THE SUBROUTINES PROGRAM. THEY WILL BE IN THE SAME ORDER AS THEY WERE READ.

CARD 3. EQUIVALENT POSITION CARD. MAY NEED MORE FOR TRANSLATIONS THAT WERE NEEDED FOR FCG16.

CARD 4. NV, NU, IRON FOR H, K, & IZ FOLLOWS BY NATOMS AND IDISK WHILE IZ IS THE UNIT ON WHICH THE FUNCTION IS STORED. THE NORMAL COORDINATE IS -1, -1, -1.

CARD 5. INFORMATION, OFFS, IR, IZ, A3Y IPICK(X), (Y), & (Z) ARE THE DIVISIONAL COORDINATES, NTRX, NTRY, AND

ENTER ANY DESIRED UNIT TRANSLATIONS. THE VALUES ENTERED ARE LESS THAN ONE.

IT IS, HOWEVER, TO OBTAIN THE PIGMENT WHERE IT IS THE LAST STEAMED INDULGE A LITTLE VAGUELY THAT A HIGHER VALUE SINCE THE PROGRAM IS STARTING AND TRY TO GET IT IN THOSE AT A TIME

NOTE 16. THE FESTIVAL POSITION CARD NO. 14 NAME IS NEGATIVE POSSITIVE CORRELATIVES ARE #1.

Douglas SULLIVAN UCT-1574

DIALECTON, PTC(100, 0.5), PTC(100, 1.20), PESAT(100, 1.20),
1 NT(100), NT(99), NT(98), NT(97), NT(96), NT(95), NT(94), NT(93), NT(92),
2 NT(91), NT(90), NT(89), NT(88), NT(87), NT(86), NT(85), NT(84), NT(83),
3 LUN(100), LUN(99), LUN(98), LUN(97), LUN(96), LUN(95), LUN(94), LUN(93),
4 FC(92, 5), FC(91, 5), FC(90, 5), FC(89, 5), FC(88, 5), FC(87, 5), FC(86, 5), FC(85, 5),
COPLEN, LPT(100, 0.5), LPT(100, 1.20), LPT(100, 1.20), LPT(100, 1.20),
1 PES, NFT(100), NFT(99), NFT(98), NFT(97), NFT(96), NFT(95), NFT(94), NFT(93),

01 FUSAL (MIA-65-12, J.S.A.)
 02 FUSAL (• 12-28, G.S. 614, 215)
 03 FUSAL (16-1)
 04 FUSAL (21-9 • M.G. = • 15, • NARL = • 15, • H = • 15, • I.C. = • 15, • T.B. = • 15, • I.C.NI = • 15, • NNUD = • 0
 2 • 15, • H(15) = • 15, • M.MAS = • 15, //)

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NVF = NVC(1)
NVT = NVC(2)
BLAD(1,1,SP) = IN,T,(CP,S1(1,NS,1,PF),IN,S=1,NVF),IN,r=1,NVF
BLAD(1,1,SP) = IN,T,(CP,S2(1,NS,1,PF),IN,S=1,NVF),IN,F=1,NVF
BLAD(1,1,SP) = IN,T,(CP,S3(1,NS,1,PF),IN,S=1,NVF),IN,F=1,NVF
BLAD(1,1,SP) = IN,T,(CP,S4(1,NS,1,PF),IN,S=1,NVF),IN,F=1,NVF
BLAD(1,1,SP) = IN,T,(CP,S5(1,NS,1,PF),IN,S=1,NVF),IN,F=1,NVF

NVT1 = NVT - ?
NVT1 = NVT + IN,T
IF(NVT1 .GT. 0.0) NVT1 = NVT1 - NVT
DO 200 JLT,VTL = 1,NVT1
BLVTL = IN,T - 1
IF(BLVTL .LT. 0.0) BLVTL = MIN(VTL - NVT
DO 150 IAT,VA7S = 1,NATAD
IF(CPIC(1,1,SP),IN).GT.NVT) CPIC(1,1,SP),IN = NVT
IF((1+IC(NAT,17)*EQUA(VTL)) .GE. VTL) GO TO 150
GO TO 160

150 CALL AGK
IF(CLK(NAT,17)*EQUA(VTL)) GO TO 160
CALL DMYTR
160 CALL CDMTR
IF(JLT,VTL .LT. NVT) GO TO 200
DO 170 IN,S = 1,NVS
DO 170 JLT,VA7P = 1,NVP
IF(S1(1,NS,1,PF) .EQ. S2(1,NS,1,PF))
IF(S2(1,NS,1,PF) .EQ. S3(1,NS,1,PF))
IF(S3(1,NS,1,PF) .EQ. S4(1,NS,1,PF))
IF(S4(1,NS,1,PF) .EQ. S5(1,NS,1,PF))
DO 170
CALL CDMTR
IF(CLK(NAT,17)*EQUA(VTL)) GO TO 160
200 CALL CDMTR
IF(S1(1,NS,1,PF) .EQ. S2(1,NS,1,PF)) IN,S=1,NVF,IN,r=1,NVF
IF(S2(1,NS,1,PF) .EQ. S3(1,NS,1,PF)) IN,S=1,NVF,IN,F=1,NVF
IF(S3(1,NS,1,PF) .EQ. S4(1,NS,1,PF)) IN,S=1,NVF,IN,F=1,NVF
IF(S4(1,NS,1,PF) .EQ. S5(1,NS,1,PF)) IN,S=1,NVF,IN,F=1,NVF
IF(S5(1,NS,1,PF) .EQ. S1(1,NS,1,PF)) IN,S=1,NVF,IN,r=1,NVF

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Y(1) = FC(1,2) / NV(2)
Z(1) = FC(1,3) / NV(3)
40 CONTINUE
WRITE(06,375)
WRITE(06,10) (I,CDOW(I),X(I),Y(I),Z(I)*B11(I)*B22(I)*B33(I),
1 B12(I)*B13(I)*B23(I)*LFT(I)*I = 1,NATOMS)
DO 60 I = 1,NATOMS
IF(NRLOC(I)*EQ.0) GO TO 554
N = LABS(NRFO(I))
IC1 = 1
IF(NRFO(I)) 45,550,550
545 IC1 = -1
550 CONTINUE
X(I) = IC1 * (X(I) * PTW(1,N) + PTW(10,N)) + NTWX(I)
Y(I) = IC1 * (Y(I) * PTW(5,N) + PTW(11,N)) + NTWY(I)
Z(I) = IC1 * (Z(I) * PTW(9,N) + PTW(12,N)) + NTWZ(I)
GO TO 60
554 CONTINUE
X(I) = X(I) + NTWX(I)
Y(I) = Y(I) + NTWY(I)
Z(I) = Z(I) + NTWZ(I)
60 CONTINUE
349 CONTINUE
WRITE(06,351)
WRITE(06,375)
WRITE(06,10) (I,CDOW(I),X(I),Y(I),Z(I)*B11(I)*B22(I)*B33(I),
1 B12(I)*B13(I)*B23(I)*LFT(I)*I = 1,NATOMS)
WRITE(07,C9) (X(I),Y(I),Z(I)*B11(I)*B22(I)*B33(I)*
1 B23(I)*LFT(I)*CDOW(I)*I = 1,NATOMS)
IF(AIPAS.EQ.0) GO TO 50
REWIND 04
WRITE(04,C9) (X(I),Y(I),Z(I)*B11(I)*B22(I)*B33(I)*B13(I),
1 B23(I)*LFT(I)*CDOW(I)*I = 1,NATOMS)
50 CONTINUE
WRITE(06,357)

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CONTINUE
STOP
END

C

SUMMING LOOPS

The purpose of this subroutine is to find the maximum F
IN Y₁₂ across six division volume.

DIMENSION IPIC(0,03), PFS1(60,120), PFS2(60,120),
1 MAX(60), IV(60), IV(60), FC(60,3)

COMMON IPIC,NAT,IR,L,IL,IV,IY,IVS,IV,MAX,IW,PFS1,PFS2,

1 PR,S,NS,IS,OUT,CC,MAXL,IVS,NVF

IPIC2 = IPIC(IR,IL) + IVS + 1
IPIC1 = IPIC(NAT,IL) + IVS + 1
IPIC2 = IPIC2 + CC*IPIC2 + NS
IPIC2 = IPIC2 + CC*IPIC2 + NS
MAX(NAT,3) = MAX(IPIC2,IPIC1)
IV(NAT) = IPIC2 + IVS - 1
IV(NAT) = IPIC1 + IVS - 1
IV(NAT) = OUT,NS
LG2 = IVS - 2
LG1 = IPIC2 + NS + 2
LG1 = LG1 + CC*IPIC1
LG1 = LG1 + CC*IPIC2 + 3
LG1 = LG1 + CC*IPIC1 + 3
DO 100 NS = 1,2
LG1 = LG1 + CC*IPIC2 + 3
IPIC2 = IPIC2 + NS - 3
DO 100 IPIC1 = 1,3
IPIC1 = IPIC1 + IPIC1 - 3
IPIC1 = IPIC1 + IPIC1 - 3
100 IY = 2*NS + 150
100 IY = 2*NS + 150
100 IY = 2*NS + 150

154 IVY = PFS2(IVF2,IVD1)

IVD3 = IVL*IVL

GO TO 160

165 IVY = PFS1(IVD2,IVD1)

IVD3 = IVL*IVL - 1

160 IF CMAX(CMAX) = IVY THEN 175,180,180

175 IMAX(CMAX) = IVY

IVNATS = IVD2 + IVL - 1

IUCNATS = IVD1 + IVL - 1

IF(IAT,0) = 1FD,

CONTINUE

174 IF(CMAX(CMAX+1) = IVL) IPICT(NATS,IR) = IV(NATS)

IF(CMAX(CMAX+1) = IVL) GO TO 180

GO TO 191

180 IV(NATS) = 0

IS(NATS) = 0

IUCNATS = 0

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350 TMAXT = TUTPT
TMINI = TINT
MINDT = IVCATS - 1
355 IF (MINDT.GT.0) MINDT = MINDT - NT
TMTF = (TMAXT - TMINF) / (TMID - TMINF)
TMDS = (TMAXS - TMINS) / (TMID - TMINS)
TMIF = (TMAYT - TMINT) / (TMID - TMINT)
BCTF = (BTAT - 4) / (ZPRTBF - 4)
BDS = (BTAS - 4) / (ZPKRS - 4)
EOT = (BTAT - 4) / (ZPRTBT - 4)
IF (BTOTF.LT.1.0 .OR. BTOTF.GT.1.5) TMAX(NATS) = 0.0
IF (BDS.GT.1.1 .OR. BDS.GT.1.5) TMAX(NATS) = 0.0
IF (BCTF.LT.1.1 .OR. BCTF.GT.1.5) TMAX(NATS) = 0.0
14 FORMAT(1H•14•,1D•.8,A3,I5,F10.5,• /•,I5,F10.5,•
12X,F10.1)
      WRITE (0,14) NATS,CGRH(NATS),LUNATS,BUOS,IVNATS,
1EOU1,FMAY(NATS)
      FC1 = R1NCF + F0CF
      FC2 = W1NFS + F0FS
      FC3 = M1NFT + F0FT
      IF (M1NCF.GT.IV(NATS)) FC1 = MINDF - BUUF
      IF (M1NFS.GT.IV(NATS)) FC2 = MINDS - ELES
      IF (M1NFT.GT.IV(NATS)) FC3 = MINDT - BCF
      FC(NATS•1F) = FC1
      FC(NATS,1S) = FCS
      FC(NATS,1Y) = FCY
      FC(TOTS)
END

```

APPENDIX B
OBSERVED AND CALCULATED STRUCTURE FACTORS

Table B-1
Observed and Calculated Structure Factors for $\text{ClCo}(\text{H}_2\text{dpg}_2) -$
 $(\text{clan}) \cdot \text{C}_2\text{H}_5\text{OH}$

L	T0	FC	L	T0	FC	L	T0	FC	L	T0	FC			
H=	0,	K=	0	19	-141	-100	11	-136	78	2	-142	-100		
			H=	0,	K=	4	12	292	270	3	-140	-138		
2	2949	-3127	0	1749	1724	13	-137	-48	4	-144	-10			
4	358	-673	1	2748	-2747	14	413	-407	5	-145	-64			
6	-108	-115	2	211	-203	15	280	-252	6	270	18			
8	461	405	3	694	706	16	271	270	7	268	139			
10	-112	46	4	659	-557	H=	0,	K=	8	8	-110			
12	427	-435	5	181	-195		0	-122	-22					
14	530	196	6	303	-279		1	-123	12	1	-146			
16	-133	65	7	210	-255		2	265	253	2	-146			
18	-131	-65	8	245	-234		3	-129	79	3	-146			
H=	0,	K=	1	9	-116	58	4	-128	-102	4	-153			
			10	287	371		5	439	-416	H=	0,	K=	13	
1	485	649	11	267	235		6	211	-107					
2	1334	1255	12	239	-703		7	-135	-66	H=	1,	K=	0	
3	440	-293	13	287	289		8	241	217	1	1980	-2098		
4	1079	1074	14	-136	152		9	-139	175	2	644	633		
5	-111	100	15	243	-747		10	-142	-61	5	419	442		
6	1139	1060	16	-134	139		11	724	-134	7	1764	-1753		
7	-107	127	17	-135	-39		12	-141	113	9	996	1025		
8	335	-347	18	267	-210		13	223	323	11	-113	0		
9	189	-175					14	223	-157	13	-124	179		
10	-112	-16	H=	0,	K=	5	15	-141	-103	15	282	350		
11	269	209					16			17	211	-303		
12	267	297	1	923	890	H=	0,	K=	9	19	310	339		
13	459	-459	2	228	151		2	-135	-66	-19	-140	60		
14	694	-701	3	1038	-1034		3	-130	168	-17	192	-156		
15	379	257	4	-112	96		4	-121	-120	-19	619	616		
16	542	530	5	-111	14		5	-133	-59	-13	690	-696		
17	-135	50	6	223	209		6	-134	40	-11	160	3		
18	-135	53	7	593	-601		7	196	41	-5	393	-456		
19	-142	30	8	-117	87		8	232	-132	-7	104	-177		
H=	0,	K=	2	9	337	326		9	-140	-59	-9	867	861	
			10	448	-407		10	247	198	-3	1436	-1533		
11	-120	-45					11	-143	-172	-1	2644	1697		
0	2055	-2207	12	-132	-52		12	-127	-81	H=	1,	K=	1	
1	1923	1405	13	401	-441		13	-129	24					
2	461	334	14	240	19		14	-140	47	H=	0,	K=	10	
3	195	-177	15	658	676		15	235	52	0	564	576		
4	258	76	16	-137	6		16	-147	147	1	1393	1364		
5	-111	48	17	346	-306		17	-147	-147	2	-98	50		
6	-108	55	18	-144	72	H=	0,	K=	10	3	1690	-1656		
7	464	448					8	428	433	4	1269	1259		
8	164	99	H=	0,	K=	6	9	279	225	5	-119	10		
9	-105	24					10	-140	-105	6	557	-563		
10	401	279	0	629	622		11	-127	-81					
11	-115	-20	1	1151	1149		12	-140	-105					
12	287	202	2	424	-423		13	-140	-105					
13	501	-255	3	538	-535		14	-141	133					
14	492	-512	4	-115	-177		15	-144	-152					
15	-137	172	5	457	445		16	-142	101					
16	229	15	6	275	226		17	-142	-150					
17	-133	-35	7	377	359		18	212	-145					
18	-135	-21	8	-123	-37		19	283	299					
19	-143	19	9	439	-450		10	-145	-54					
H=	0,	K=	3	10	297	299		11	222	-215				
			11	214	189		12	234	0	10	-138	68		
12	261	-173					13	-142	-91					
1	638	-561	13	350	-356	H=	0,	K=	11	-19	-133	103		
2	932	-584	14	-135	-67		15	-138	-67	-18	-135	-132		
3	654	-526	15	-135	150		16	-130	176	-18	-135	-132		
4	218	-100	16	-140	-134		17	-141	-141	-15	-71	512		
5	333	340	17	230	229		18	269	-313	-14	281	-252		
6	192	191					19	210	138	-12	217	-143		
7	312	272	H=	0,	K=	7	5	210	138	-12	-124	-173		
8	222	-229					6	294	322	-11	-118	-43		
9	-111	-27	1	572	-596		7	-141	-7	-10	175	-216		
10	189	169	2	-119	-170		8	217	-100	-9	347	342		
11	-119	-100	3	318	328		9	-147	111	-9	403	-403		
12	189	72	4	554	-561		10	-141	-26	-8	1108	-1110		
13	196	123	5	201	-187		11	-151	9	-7	643	651		
14	210	150	6	406	399		15	-129	-112	-5	427	432		
15	376	-299	7	-145	104	H=	0,	K=	12	-4	1275	-1245		
16	207	-130	8	235	-323		16	-145	-112	-2	324	323		
17	-135	217	9	-133	159		17	-129	-112	-2	-89	107		
18	212	-67	10	-135	70		18	-143	-170	-2	-89	107		

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
-1	585	-563	-7	356	330	-10	-125	-36	-7	369	-414
H= 1, K= 2			-6	-112	-72	-9	-119	-147	-6	-126	-108
0	-101	-17	-3	474	502	-8	404	-403	-5	722	740
1	1167	1077	-2	315	-280	-6	1334	1319	-3	251	-270
2	-107	-48	-1	2463	-2419	-5	163	-54	-2	840	-839
3	1256	1165				-4	821	-856	-1	518	-570
4	415	404	H= 1, K= 4			-3	426	-412			
5	436	435				-2	503	-594	H= 1, K= 8		
C	-110	10	0	981	-1005	-1	1052	1023	0	-126	0
7	-112	123	1	-115	-169	H= 1, K= 6			1	-126	61
8	452	-474	2	844	826	0	1499	1489	2	-124	38
9	-113	15	3	501	-490	1	178	-4	4	-132	57
10	-115	46	4	-112	51	2	301	-321	5	-138	-138
11	-118	78	5	206	-153	3	-117	150	6	-136	-54
12	-123	78	6	499	-490	4	-121	-239	7	-128	-30
13	489	-457	7	964	-571	5	-118	-5	8	-142	61
14	-135	-66	8	541	521	6	814	901	9	-144	-216
15	-137	-112	9	228	218	7	268	-276	10	-142	108
16	-134	12	10	-123	-112	8	620	-610	11	297	239
17	-134	79	11	-129	-194	9	251	253	12	241	154
18	-135	3	12	236	105						
19	-147	5	13	199	171	10	438	426	13	-139	-91
-19	246	-96	14	-136	-71	11	223	-60	14	-140	-118
-18	-139	-120	15	-139	147	12	251	-289	15	-146	12
-17	215	148	16	-134	-113	13	-140	49	-16	221	54
-16	-136	-205	17	310	-265	14	-128	48	-15	-144	-147
-15	451	-404	18	238	74	15	-136	-81	-14	-141	-141
-14	432	368	-19	-146	15	16	-145	204	-13	270	297
-13	491	490	-19	-136	-84	17	-149	3	-12	-128	17
-12	268	-255	-17	-134	-44	-17	-143	-42	-11	-138	-157
-11	512	518	-16	-139	194	-16	-142	-73	-10	-139	55
-10	-114	65	-15	-135	141	-15	-136	180	-9	-142	-33
-9	546	-547	-14	580	-582	-14	238	347	-8	-141	-68
-8	627	-594	-13	-134	-94	-13	285	-239	-7	-137	-60
-7	374	386	-12	-134	199	-12	411	-284	-6	219	217
-6	-109	-66	-11	213	159	-11	388	450	-5	-131	93
-5	204	96	-10	440	415	-10	212	273	-4	216	-205
-4	-114	-66	-9	333	-324	-9	258	-308	-3	269	320
-3	2196	2050	-8	-113	-119	-8	-125	-40	-2	-129	75
-2	1662	-1503	-7	221	146	-7	184	232	-1	366	-440
-1	2235	-2404	-6	305	276	-6	287	-410			
H= 1, K= 3			-5	810	614	-5	-116	14	H= 1, K= 9		
0	2381	-2325	-4	563	-534	-4	999	1028	0	-135	-111
1	1050	1050	-3	340	-370	-3	492	-510	1	-135	22
2	-118	-58	-2	322	328	-2	1315	-1314	2	-134	-41
3	806	-823	-1	521	591	-1	277	282	3	-124	46
H= 1, K= 5			H= 1, K= 7			H= 1, K= 7			4	-141	145
0	1758	1712	0	574	538	0	290	292	5	-145	-153
5	-112	-104	1	212	-190	1	534	552	6	-145	-215
6	228	-225	2	608	-880	2	-118	-150	7	-146	109
7	-111	-113	3	-112	-55	3	212	-225	8	-144	-59
8	754	-729	4	671	672	4	217	209	9	207	-150
9	707	702	5	-114	122	5	184	202	10	220	174
10	-117	29	6	683	-705	6	205	206	11	-145	-225
11	-123	-176	7	683	341	7	204	-286	12	-143	-106
12	-136	234	8	222	81	8	-135	47	13	230	301
13	-134	259	9	349	-366	9	-136	102	14	-140	90
14	349	-359	10	191	193	10	242	-331	14	-142	19
15	258	-517	11	-132	25	11	248	243	15	-139	-44
16	239	300	12	323	-314	12	298	256	16	267	-206
17	-139	-113	13	-139	35	13	212	-202	17	-149	201
18	197	-200	14	569	567	14	258	256	18	-140	95
19	-141	-32	15	257	-155	15	283	256	19	305	-257
-19	-145	-184	16	342	-374	16	211	161	-2	-142	-21
-18	228	42	17	342	-374	17	221	-220	-7	377	393
-17	-138	58	18	250	171	18	221	-220	-6	365	338
-16	-136	-157	19	-153	70	-16	-145	-169	-5	-137	-116
-15	229	-223	18	323	-258	-15	-142	132	-4	324	-334
-14	244	215	-17	-133	-6	-16	-124	0	-3	-138	-109
-13	-130	-107	-16	515	442	-13	-137	-113	-2	-130	55
-12	102	64	-15	-137	113	-14	273	260	-1	195	204
-11	-117	51	-14	362	-423	-13	-127	63			
-10	-116	-20	-13	-126	132	-10	401	-412	H= 1, K= 10		
-9	386	-517	-12	-121	-123	-9	133	67			
-8	-114	-125	-11	-123	-123	-8	231	239	0	261	176

L	E0	FC	L	E0	FC	L	E0	FC	L	E0	FC		
1	-143	-106	4	-149	-138	5	-133	113	0	1324	-1299		
2	-141	-99	4	-139	66	6	-102	112	1	462	-263		
3	-143	-73	3	-149	-12	7	-279	-212	2	113	-110		
4	-146	-1	2	-151	-53	8	-114	-95	3	216	-252		
5	242	731	1	-145	-79	9	-110	107	4	-117	266		
6	-143	87				10	-106	-101	5	-264	-200		
7	207	-145	H=	2,	K=	0	11	-102	-150	6	-113	-130	
8	294	-300				12	-194	-71	7	-350	561		
9	-143	174	0	592	560	13	-259	-243	8	413	474		
10	-143	129	2	927	-916	14	-136	-79	9	260	-261		
11	-146	-190	4	1131	1113	15	-119	101	12	-145	166		
12	-145	62	6	1693	-1823	16	-797	737	11	309	321		
-13	-145	-114	8	1767	1697	17	-100	-266	12	-123	-50		
-12	-147	-22	10	756	-777	18	-119	-175	13	-139	83		
-11	-145	173	12	746	72	19	-165	66	14	-173	-4		
-10	-140	-34	14	-137	125	+19	-157	-61	15	-128	-171		
-9	-140	-147	16	562	-500	+18	-141	156	16	-126	-54		
-8	-140	-70	18	460	461	+17	-125	-13	17	-130	65		
-7	-143	103	+20	-143	-131	+16	-625	-405	18	-140	127		
-6	-138	-3	+18	-140	-106	+15	-796	-594	+19	-140	-82		
-5	-147	224	+16	557	565	+14	-533	426	+18	-139	-15		
-4	-148	168	+14	556	-675	+13	-214	-210	+17	259	260		
-3	393	-369	+12	543	539	+12	-214	264	+16	-134	53		
-2	-145	-60	+10	444	-463	+11	-305	365	+15	372	-432		
-1	296	411	+8	372	-378	+10	-116	25	+14	215	-52		
			+6	456	446	+9	-112	105	+13	355	373		
H=	1,	K=	11	-4	956	-957	+8	200	-148	+12	-127		
			+2	1426	1073	+7	204	106	+11	436	-431		
0	-137	97				+6	1075	-1055	+10	171	26		
1	-140	125	H=	2,	K=	1	+5	-112	-251	+9	-115	18	
2	-140	-58				+4	-114	-73	+8	-113	0		
3	370	-370	0	314	307	+3	-113	-193	+7	269	224		
4	313	214	1	-100	-193	+2	-908	877	+6	407	397		
5	-149	291	2	1529	-1525	+1	631	671	+5	250	-260		
6	-143	-162	3	203	-231				+4	-110	84		
7	-143	-89	4	1255	1271	H=	2,	K=	3	+3	-112	-184	
8	-143	22	5	-114	-18				+2	1020	1537		
9	-148	-18	6	306	-215	0	252	-252	+1	-110	141		
10	-150	-23	7	307	300	1	400	400					
11	-153	183	8	359	-331	2	276	258	H=	2,	K=	5	
-11	-151	-132	9	-114	2	3	161	-171					
-10	-145	66	10	367	-373	4	-112	61	0	699	-620		
-9	-150	277	11	417	-414	5	132	663	1	881	-879		
-8	268	-227	12	626	-435	6	-105	-24	2	366	-344		
-7	306	-353	13	336	366	7	-113	56	3	972	850		
-6	-139	108	14	504	565	8	-114	-128	4	-115	-175		
-5	209	260	15	-135	-161	9	-117	110	5	393	-343		
-4	-135	-39	16	-139	-340	10	228	-267	6	216	277		
-3	-142	-139	17	-134	38	11	-125	51	7	-116	11		
-2	-138	-40	18	-126	-61	12	279	236	8	319	-317		
-1	-142	103	19	-145	69	13	62	-275	9	320	303		
			+19	-126	-65	14	399	-413	10	284	301		
H=	1,	K=	12	+18	350	-373	+16	245	323	11	509	-507	
			+17	220	170	+16	-134	114	12	-137	-115		
0	267	174	+16	241	235	+17	-142	-132	13	126	720		
1	-142	97	+15	265	-275	+18	-138	-72	14	-134	105		
2	-139	-38	+14	-132	171	+19	-141	121	15	161	-201		
3	-144	0	+13	-113	-52	+18	-129	102	16	-140	119		
4	-148	-94	+12	356	-292	+17	272	-224	17	-129	43		
5	-145	-61	+11	-113	29	+16	-122	-167	18	-152	-90		
6	-146	159	+10	623	619	+15	229	-103	+16	-154	-14		
7	-147	78	+9	561	-613	+14	-135	-63	+17	-143	200		
8	-151	-143	+8	1365	-1377	+13	212	106	+16	157	148		
-1	-148	-191	+7	602	591	+12	464	671	+15	-142	-217		
-7	219	20	+6	319	390	+11	-115	-105	+14	-124	-131		
-6	-143	19	+5	1273	-1232	+10	233	-328	+12	315	-244		
-5	-146	-126	+4	180	76	+9	400	629	+11	639	470		
-4	-143	55	+3	926	-494	+8	401	462	+10	-17	199		
-3	-146	87	+2	303	-272	+7	223	-205	+9	523	-560		
-2	304	-206	+1	-93	29	+6	356	-373	+8	249	-195		
-1	-143	-88				+5	-109	56	+7	693	670		
H=	1,	K=	13	H=	2,	K=	2	+4	272	-263	-6	-116	-66
						+3	603	790	+5	523	-521		
0	-150	-45	0	-106	1	+2	1252	-1250	+4	717	-703		
1	-153	66	1	609	-620	+1	-117	+28	+3	170	-169		
2	-151	157	2	1317	-1307	H=	2,	K=	4	+2	1140	103	
3	242	-16	3	1243	-1223	H=	2,	K=	4	+1	1397	1397	

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC	
H= 2, K= 6	0	195	-153	11	-146	-15	9	494	-493			
0	269	-293	3	285	290	12	-146	55	11	273	-247	
1	646	-625	4	-134	-124	-13	-147	-88	13	403	-429	
2	584	-576	5	-145	-142	-12	-144	111	18	714	-729	
3	-119	-168	6	-141	198	-11	-144	-59	17	254	304	
4	203	79	7	-142	-89	-10	-142	5	19	254	-155	
5	748	786	8	-141	56	-9	-140	206	-19	-136	-64	
C	197	-204	9	217	-162	-8	-140	-98	-17	613	634	
7	541	-642	10	213	127	-7	-141	-74	-18	531	-542	
8	258	275	11	-141	7	-6	-139	-25	-13	497	493	
9	337	334	12	-139	-131	-5	-148	-148	-11	400	-436	
10	319	-365	13	-144	6	-4	370	-355	-9	356	-310	
11	-133	13	14	-141	-9	-3	-143	19	-7	1011	1026	
12	-145	-5	15	-144	121	-2	488	487	-5	1442	-1403	
13	-141	-126	16	281	-187	-1	201	226	-3	203	221	
14	-135	56	-15	-148	-45				-1	1290	-1240	
15	-137	81	-14	-143	128							
16	-139	-58	-13	-137	60	H= 2, K= 11						
17	-145	-117	-12	-136	-88	0	-144	198	H= 3, K= 1			
-17	-145	-104	-11	-141	152	1	-143	-73	0	551	-573	
-16	-137	208	-10	-145	181	2	-143	-352	1	269	-216	
-15	347	344	-9	-147	-284	3	-143	62	2	1756	1736	
-14	-141	-293	-8	-140	-161	4	-143	324	3	261	-271	
-13	510	-308	-7	380	415	5	-143	-166	4	583	-569	
-12	421	416	-6	-134	-34	6	-144	-73	5	241	-271	
-11	204	210	-5	-134	-85	7	-145	177	6	754	-771	
-10	360	-358	-4	481	499	8	-146	-112	7	510	-492	
-9	-128	58	-3	260	304	9	-149	-29	8	-114	124	
-8	199	256	-2	-129	-129	10	327	240	9	716	708	
-7	262	-294	-1	425	-431	11	322	163	10	-121	-169	
-6	-122	-92				12	-152	230	11	796	-805	
-5	-120	246	H= 2, K= 9			13	-144	-146	12	343	359	
-4	532	-517				14	262	-355	13	440	432	
-3	609	-601	0	291	-309	15	331	270	14	307	-268	
-2	1065	1096	1	-137	-52	16	-142	-102	15	199	-127	
-1	1220	1224	2	-141	145	17	-143	13	17	-136	-102	
			3	-137	79	18	-144	-220	19	-146	150	
H= 2, K= 7	4	-137	0	-1	-142	105	-19	-144	-179	-18	213	130
C	601	579	5	231	241	H= 2, K= 12	-17	-137	-222	-16	-135	-103
1	206	210	6	-143	-42	0	-144	126	-15	-138	198	
2	324	-435	7	-146	-126	1	-145	-6	-14	-129	-88	
3	376	-341	8	-144	-38	2	-147	-35	-13	412	-424	
4	449	405	9	-142	183	3	214	-107	-12	-122	194	
5	-130	151	10	-135	-31	4	-149	-163	-11	886	870	
6	-132	-277	11	-143	-271	5	-149	224	-10	270	-288	
7	-130	-152	12	236	223	6	294	144	-9	453	-438	
8	-133	60	13	-151	136	7	226	-249	-8	624	615	
9	-136	-139	14	-149	-72	8	222	-42	-7	282	241	
10	222	226	15	-139	-8	9	-151	-17	-6	333	-399	
11	335	347	16	-143	-60	10	-151	-29	-5	-110	179	
12	574	-601	17	-141	34	11	-149	-122	-4	505	-520	
13	256	-188	18	313	257	12	-153	-122	-3	-112	68	
14	300	308	19	224	-310	13	-151	251	-2	393	323	
15	-147	152	20	-140	25	14	226	277	-1	1687	1663	
16	215	-71	21	-140	117	15	335	-359				
-17	-146	-97	22	-140	71	16	-143	-177				
-16	-143	89	23	285	-275	H= 3, K= 2						
-15	200	16	24	-138	-68	H= 2, K= 13	0	450	-473	1	413	382
-14	-136	85	25	539	507	1	-180	109	2	822	849	
-13	219	126	26	263	377	2	-148	15	3	222	-194	
-12	-138	-170	27	301	-252	3	-148	-71	4	852	831	
-11	219	-257	H= 2, K= 10	0	-149	53	5	1072	1074			
-10	457	419	1	-142	-149	1	-180	109	6	635	-636	
-9	403	441	2	-140	-77	2	-151	52	7	426	-421	
-8	476	-503	3	276	-243	3	-151	-8	8	426	448	
-7	240	-215	4	-142	-149	4	278	-188	9	190	233	
-6	437	469	5	-140	-77	5	-148	-71	10	-117	-94	
-5	189	-150	6	-143	-99	6	-149	-59	11	-122	-173	
-4	-124	72	7	-147	163	7	-151	-59	12	124	-116	
-3	-122	143	8	-145	153	8	586	-564	13	-154	-317	
-2	382	-610	9	300	-262	9	795	-758	14	327	309	
-1	293	-221	10	-141	-3	10	1501	-1510	15	-141	261	
H= 2, K= 6	9	-143	81	10	-149	-216	11	1092	1081	16	266	-217

L	F0	FC									
17	-130	-185	13	248	179	11	-140	51	-15	-146	82
16	285	127	14	261	-300	12	256	-230	-14	-128	2
-19	-140	95	15	-130	-169	13	373	370	-13	-137	101
-18	242	-216	16	-145	160	14	374	318	-12	236	142
-17	225	-273	17	-147	192	15	-144	-177	-11	-141	-175
-16	-138	160	18	-145	-165	16	-150	-210	-10	385	-390
-15	421	420	-19	-143	75	-17	-147	129	-9	-134	110
-14	302	-279	-18	278	244	-18	274	307	-8	-140	207
-13	-124	-197	-17	-134	-25	-15	-137	49	-7	-136	-110
-12	416	449	-16	421	-406	-14	378	-337	-6	-135	-22
-11	402	-409	-15	-135	-34	-13	-138	19	-5	-138	258
-10	-114	49	-14	250	267	-12	271	314	-4	260	280
-9	516	522	-13	-130	94	-11	-136	-196	-3	590	-574
-8	554	-529	-12	220	24	-10	236	192	-2	447	-490
-7	172	-156	-11	-125	-8	-9	-124	-7	-1	-129	84
-6	553	547	-10	-121	-121	-8	526	-500	H=	3,	K= 9
-5	262	-220	-9	-119	-224	-7	538	523	0	-137	-79
-4	-112	-16	-8	548	608	-6	-120	191	1	228	102
-3	210	-161	-7	245	283	-5	407	-407	2	-141	131
-2	654	610	-6	248	233	-4	573	-562	3	-139	-82
-1	369	-230	-5	232	-186	-3	316	314	4	-137	-36
H=	3,	K= 3	-4	-110	-36	-2	693	920	5	-146	93
0	-115	-9	-3	579	603	-1	-114	44	6	286	-323
1	331	-281	-2	738	-778	H=	3,	K= 7	7	241	75
2	1146	1182	-1	858	-858	0	633	626	8	221	213
3	298	-231	H=	3,	K= 5	1	466	-456	9	232	-157
4	485	-444	0	587	-574	2	288	253	10	-143	-67
5	428	428	1	219	47	3	442	472	11	-145	67
6	-112	55	2	940	567	4	-123	-11	12	217	273
7	162	72	3	-113	-47	5	-126	-38	13	-151	-70
8	217	244	4	710	-718	6	-130	29	-14	222	147
9	-117	-29	5	-118	-152	7	182	-102	-12	-144	6
10	328	260	6	280	-277	8	-135	-83	-12	-140	-41
11	257	256	7	181	-153	9	-141	22	-11	-140	-66
12	335	-209	8	386	398	10	203	156	-10	-141	-128
13	221	-222	9	187	228	11	242	-225	-9	-141	133
14	-138	193	10	380	-379	12	353	-377	-8	-144	-9
15	-137	199	11	-134	34	13	-143	194	-7	-142	101
16	-132	159	12	636	638	14	-140	199	-6	263	198
17	-139	-83	13	-157	64	15	-143	-16	-5	-139	-170
18	-143	-74	14	218	-210	16	-144	-2	-4	294	-245
-19	-140	89	15	-140	-214	-16	-143	-33	-3	285	329
-18	-138	13	16	-137	-97	-15	-139	-9	-2	-133	43
-17	-135	-80	17	-140	-8	-14	-136	39	-1	352	-366
-16	-135	-193	-18	273	247	-13	-136	-57	H=	3,	K= 10
-15	-133	-97	-17	254	220	-12	243	-204	0	266	-172
-14	-132	99	-16	-136	-48	-11	318	248	1	-147	-10
-13	315	324	-15	223	-246	-10	314	287	2	-147	60
-12	338	-282	-14	-138	-162	-9	561	-558	3	-145	91
-11	496	-505	-13	-140	181	-8	-134	163	4	-147	-29
-10	636	649	-12	-135	162	-7	-133	110	5	315	-292
-9	352	367	-11	-132	-27	-6	274	-148	6	-148	10
-8	-112	-34	-10	620	-629	-5	-126	86	7	229	253
-7	-109	-78	-9	-119	39	-4	294	250	8	-147	-136
-6	516	-555	-8	969	954	-3	447	-470	9	-149	-48
-5	-110	-103	-7	-115	7	-2	646	-672	10	-148	35
-4	275	-270	-6	200	53	-1	524	546	11	-152	50
-3	557	566	-5	-115	-7	H=	3,	K= 8	12	-152	52
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H=	3,	K= 4	-2	1008	1015	2	-134	4	-11	-151	-140
0	199	-205	H=	3,	K= 6	3	-133	-170	-10	-145	-37
1	549	535	0	-118	96	4	257	-131	-9	278	-159
2	-109	-23	1	302	280	5	247	208	-7	-152	284
3	-112	-7	2	531	-538	6	-137	104	-6	-146	64
4	504	-510	3	268	264	7	-140	-178	-5	357	-379
5	529	-560	4	659	651	8	-142	65	-4	-151	-88
6	359	389	5	678	-694	10	-142	-79	-3	349	380
7	-117	85	6	847	-851	11	-137	1	-2	262	327
8	208	-160	7	372	398	12	205	-52	-1	-145	-134
9	246	314	8	413	451	13	-141	-196			
10	209	93	9	290	-268	14	-144	18			
11	288	-231	10	-135	-160	15	200	97			

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC		
H=	3+, K= 11		0	237	-227	-1	-113	105	-3	-110	-53		
1	-147	-69	1	-114	-135	1	-111	-61	-2	-109	-58		
2	-146	86	2	629	599	H=	4+, K= 3	-1	,241	224			
3	-151	151	3	270	-273								
4	-147	-126	4	-109	83	0	-111	-61	H=	4+, K= 5			
5	266	25	5	401	389	1	-111	-114	0	245	-328		
6	-150	146	6	252	-205	2	643	633	1	255	348		
7	-153	-226	7	536	570	3	430	395	2	-114	89		
8	-149	-32	8	1196	1187	4	458	-449	3	-117	94		
9	-161	248	9	505	-487	5	-112	6	4	-118	-92		
10	-155	-109	10	844	-824	6	240	363	5	284	-238		
11	-153	3	11	333	334	7	588	-618	6	-121	130		
12	-161	248	8	432	403	8	264	-375	7	389	264		
13	-155	-109	9	-132	23	9	374	354	8	-129	-168		
14	-153	3	10	152	46	10	286	289	9	724	-735		
15	-150	-43	11	-137	-201	11	288	-243	10	-134	-14		
16	-149	-41	12	-138	127	12	-132	-89	11	690	729		
17	-148	186	13	-140	54	13	-133	-118	12	-137	84		
18	-150	-52	14	294	191	14	265	292	13	-139	-103		
19	-149	-47	15	-143	74	15	-136	23	14	-139	-170		
20	379	-295	16	-142	-85	16	200	-127	15	-140	-131		
21	298	-279	17	-140	-74	17	-140	39	16	-142	112		
22	299	333	18	201	23	18	-145	-68	17	222	168		
23	521	524	19	-131	26	19	-145	-63	18	-140	24		
24	-150	-13	20	562	-602	20	-139	-56	19	-137	-66		
25	-12	-13	21	287	298	21	-137	-211	20	-138	-199		
26	H= 3+, K= 12		22	680	709	22	-16	197	19	-135	-92		
27	-11	269	23	-282	-282	23	306	286	15	-136	166		
28	0	-123	24	615	-623	24	226	8	14	-136	166		
29	1	-147	25	455	453	25	-135	-111	13	331	352		
30	2	-153	26	221	184	26	230	-204	12	-135	-141		
31	3	-149	27	257	-179	27	271	301	11	480	-451		
32	4	-155	115	-107	105	28	-119	218	10	183	-26		
33	5	238	100	-5	-109	57	29	381	-288	9	400	411	
34	6	291	-195	-4	741	-773	30	380	390	8	537	545	
35	7	-151	-119	-3	391	393	31	172	-160	7	313	-346	
36	8	-158	-93	-2	271	226	32	-110	-124	6	-118	3	
37	9	-158	-70	-1	343	-367	33	878	694	5	249	-202	
38	10	312	281	-3	-164	171	34	655	640	4	651	-648	
39	11	288	249	H= 4+, K= 2		35	164	171	3	661	678		
40	12	422	-424	12	726	-734	36	-113	135	2	-114	134	
41	13	-3	316	327	1	243	-172	37	-113	-112	1	-112	-86
42	14	-152	117	2	-112	-227	H= 4+, K= 4						
43	15	3	-112	-113	H= 4+, K= 6								
H=	3+, K= 13		4	201	-147	0	-111	-175	0	376	-431		
1	5	489	-517	1	197	145	1	260	-213				
2	0	-161	229	6	-110	-23	2	-113	196	2	367	338	
3	1	223	-56	7	226	225	3	440	-460	3	861	898	
4	2	-159	-113	8	-113	7	4	191	-260	4	414	-446	
5	3	-159	95	9	285	311	5	559	532	5	1018	-1014	
6	6	-161	-234	10	-124	101	6	795	730	6	727	699	
7	7	-161	-109	11	240	-257	7	202	-209	7	383	379	
8	8	12	414	-412	8	312	-319	8	219	-346			
9	9	-123	-113	9	-123	4	9	-138	190				
10	H= 4+, K= 0		10	353	359	10	-222	10	-142	-136			
11	11	14	-138	153	10	234	-222	11	241	-272			
12	12	15	-133	-51	11	302	304	12	-139	179			
13	13	6	-137	6	12	218	292	13	-140	174			
14	14	7	-137	61	13	268	-158	14	-128	-163			
15	15	8	-137	62	14	-133	-55	15	296	-226			
16	16	9	-139	62	15	245	69	16	-143	108			
17	17	10	-137	89	16	225	156	17	295	217			
18	18	11	-137	89	17	215	-184	18	-140	14			
19	19	12	-137	61	18	-140	156	19	-140	174			
20	20	13	-137	56	19	-144	-200	15	-140	-218			
21	21	14	-137	210	20	234	-204	14	-134	-218			
22	22	15	-137	210	21	-137	176	13	-130	20			
23	23	16	-137	378	22	314	330	12	-126	56			
24	24	17	-137	378	23	276	130	11	291	197			
25	25	18	-137	359	24	234	-224	10	377	-377			
26	26	19	-139	186	25	-111	373	9	349	-354			
27	27	20	-139	186	26	-126	-193	8	347	321			
28	28	21	-139	924	27	786	790	7	617	648			
29	29	22	-139	907	28	-116	73	6	281	-206			
30	30	23	-139	846	29	-116	691	5	738	-750			
31	31	24	-139	413	30	-116	681	4	878	892			
32	32	25	-139	-110	31	121	-292	3	134	-204			
33	33	26	-139	-110	32	830	643	2	308	-315			
H=	4+, K= 1		34	757	752	33	189	150					
35	35	2	1159	-1135	34	-1135							

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC		
-1	172	-145	6	-140	92	3	-152	178	1	-109	-160		
E=	4+	K=	7	7	-141	69	4	203	131	2	501	564	
			8	-142	-77	5	223	-240	3	511	499		
			9	237	-73	6	-149	-75	4	650	-605		
0	314	-242	10	-140	59	7	-146	94	5	20	189		
1	-122	-67	11	-143	34	8	-150	123	6	571	599		
2	229	231	12	-145	-53	9	223	-159	7	250	230		
3	201	-240	13	-150	-151	10	-149	-44	8	266	235		
4	-132	-65	14	-148	97	11	-145	156	9	420	-61		
5	455	464	15	-149	-36	12	-140	173	10	347	-344		
6	345	-310	16	-150	-65	13	-142	13	11	357	-304		
7	397	-208	14	-138	79	14	-132	17	12	-132	17		
8	329	413	15	-137	-98	15	-140	-69	13	-140	-69		
9	-143	192	16	-139	136	16	569	581	14	-140	190		
10	325	-259	17	222	6	1	569	581	15	-138	69		
11	313	-238	18	-132	-43	2	236	-2345	16	-135	146		
12	-140	-15	19	-141	93	3	1216	1526	17	-122	70		
13	330	226	20	314	-34	4	-105	60	18	-147	67		
14	-145	146	21	-143	111	5	658	-661	19	269	-693		
15	-147	-116	22	404	420	6	672	674	18	-144	220		
-16	-140	75	23	313	-261	7	304	311	17	245	37		
-15	-136	-14	24	-142	-245	8	-125	35	16	-123	-63		
-14	-140	-64	25	-137	-11	9	-137	-11	15	-137	-118		
-13	-138	52	26	6	K=	10	-19	16	14	269	-291		
-12	-140	-20	27	-145	-133	11	299	264	13	262	-10		
-11	-141	115	28	-148	54	12	-137	-261	12	335	384		
-10	-142	-207	29	-141	148	13	-132	-100	11	762	735		
-9	226	-232	30	-141	-148	14	537	-535	10	315	-295		
-8	206	212	31	257	141	15	1210	1201	9	211	-157		
-7	202	-234	32	267	-396	16	1254	-1369	8	-113	140		
-6	-137	-177	33	-143	-69	17	387	246	7	-113	51		
-5	687	715	34	332	335	18	-109	-67	6	555	-545		
-4	414	-414	35	-142	-129	19	208	172	5	415	316		
-3	360	-321	36	-142	-69	20	-169	-29	4	456	471		
-2	673	711	37	-142	171	21	855	880	3	353	-367		
-1	-122	119	38	-148	-158	22	-111	90	2	157	-44		
			39	-149	-29	23	293	-375	1	-109	53		
E=	4+	K=	8	-12	206	24	1	-111	90	E=	5+	K=	3
			40	-147	-160	2	293	-375					
0	-135	-257	41	-142	-95	3	737	705	0	781	758		
1	-135	-104	42	-141	145	4	482	-495	1	502	-442		
2	-134	59	43	-140	222	5	1230	-1223	2	728	-709		
3	-135	-61	44	-143	76	6	601	591	3	627	-623		
4	475	481	45	354	-365	7	662	-664	4	249	-310		
5	-141	171	46	-143	-106	8	372	-386	5	500	-402		
6	-144	-210	47	275	290	9	612	-624	6	201	163		
7	-143	-9	48	-146	23	10	272	305	7	326	-353		
8	-143	11	49	272	-183	11	292	366	8	520	508		
9	334	-248	50	-143	-162	12	-132	64	9	383	-434		
10	-142	161	51	-143	-162	13	404	-409	10	258	-250		
11	-144	94	52	-144	-162	14	-124	-65	11	-120	63		
12	-140	-164	53	-147	-160	15	-137	115	12	-196	-174		
13	-143	-185	54	281	-304	16	-138	56	13	-145	274		
14	-140	101	55	-141	99	17	407	461	14	276	-54		
15	-145	111	56	-144	93	18	-146	-155	15	188	-65		
-14	-144	111	57	-144	17	19	-132	-67	16	-137	-66		
-13	-140	50	58	-142	71	20	-138	-160	17	-142	-141		
-12	-137	-185	59	-144	-57	21	-137	122	18	205	165		
-11	-141	-176	60	257	-296	22	708	139	19	-139	-76		
-10	-143	219	61	-148	55	23	-137	107	20	-159	-76		
-9	-142	243	62	214	232	24	-132	111	21	-177	-123		
-8	546	-498	63	-153	-32	25	609	590	16	-133	12		
-7	-139	-276	64	-151	-131	26	-127	-61	17	226	-190		
-6	346	443	65	-146	-13	27	518	-522	18	-126	-10		
-5	-137	-160	66	-146	214	28	643	662	19	224	-126		
-4	-135	-63	67	-144	-1	29	112	17	20	324	336		
-3	242	275	68	-135	6	30	249	-344	21	-172	36		
-2	-131	6	69	-144	-6	31	521	524	22	164	237		
-1	-132	-67	70	363	-273	32	446	-403	23	-119	-25		
E=	4+	K=	9	-2	360	347	33	606	617	24	297	-256	
			-1	455	-404	34	606	617	25	907	948		
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1	-142	175		0	-145	-33	H=	5+	K=	2	272	-175	
2	-144	102		1	-146	-65		-1	-110	55	280	179	
3	-141	-10		2	-147	-61				1	199	277	

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
H= 5, K= 4			H= 5, K= 6			H= 5, K= 6			H= 5, K= 11		
0 -116	-147		0 -121	-256		4 -138	-108		-5 -140	44	
1 -113	94		1 301	301		5 292	-227		-4 -143	98	
2 724	-724		2 521	548		6 -140	126		-3 -139	-8	
3 -113	-37		3 378	-298		7 -143	87		-2 -145	24	
4 901	910		4 505	-484		8 -140	0		-1 -144	37	
5 472	-451		5 373	392		9 -145	83				
6 -118	-134		6 240	173		10 -138	66				
7 474	-416		7 -130	152		11 238	-254		H= 5, K= 11		
8 450	-470		8 -124	151		12 -138	17		0 -144	49	
9 -128	-39		9 -140	-248		13 -151	107		1 249	177	
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11 -137	104		11 257	247		15 -141	12		3 -143	146	
12 -138	186		12 291	287		16 -141	-33		4 -144	-165	
13 235	-153		13 -138	-76		17 -141	-190		5 244	-240	
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-17 235	-17		19 -138	-108		23 -137	-180		-7 -151	-84	
-16 -139	-105		20 -142	190		24 -138	609		-6 -145	-24	
-15 232	353		21 228	202		25 -138	4		-5 -143	-3	
-14 237	168		22 263	202		26 -138	-531		-4 -137	-34	
-13 587	-610		23 280	-262		27 -138	261		-3 -143	103	
-12 342	-382		24 559	-504		28 -140	-1		-2 -142	-10	
-11 215	119		25 190	149		29 -140	-146		-1 -146	-237	
-10 438	430		26 472	492		30 -142	-88				
-9 322	300		27 273	-339		H= 5, K= 9			H= 5, K= 12		
-8 208	-179		28 438	-421		0 223	228		0 -151	-190	
-7 222	-371		29 -121	131		1 -145	166		1 217	-180	
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-4 -114	56		32 -120	102		4 -147	-190		4 385	-297	
-3 403	284		33 -119	114		5 -150	221		5 -154	-19	
-2 -111	-23		34 -119	114		6 -145	89		6 -153	100	
-1 -114	168		H= 5, K= 7			7 -141	-176		7 -152	-160	
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0 299	372		9 -142	-12		9 -142	88		9 -146	94	
1 689	662		10 -129	-53		10 -144	60		-3 -153	-157	
2 281	-301		11 195	-88		11 213	-124		-2 -151	93	
3 319	352		12 -132	-24		12 -144	65		-1 -149	79	
4 890	-895		13 231	333		H= 6, K= 0					
5 209	-132		14 281	-291		0 -148	123				
6 1020	1068		15 381	-393		1 -148	-82				
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8 1006	-1040		17 514	507		3 -148	123				
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12 384	-297		21 -136	-165		7 -141	-20		0 -107	123	
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14 205	212		23 -140	-152		9 -141	-55		2 1232	1246	
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16 -149	35		25 -134	58		11 -144	243		8 603	-619	
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-17 -138	-10		27 -135	-85		13 -141	37		12 -132	-21	
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-13 219	218		31 -133	-90		17 -141	-82		-16 -130	42	
-12 573	-598		32 -133	-90		H= 6, K= 10			-14 -136	-102	
-11 -133	-97		33 -212	-105		0 -142	-86		-12 718	-744	
-10 570	563		34 -212	-105		1 -146	232		-10 832	850	
-9 -128	2		35 -213	-90		2 -143	101		-8 -112	-158	
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			47 -142	805		14 -142	805		7 181	-113	
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L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC	
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6	-170	89	87	553	-576	88	-140	-65	-1	-143	-47	
7	-117	74	88	-136	-14	89	-140	203				
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-9	-142	-152	13	318	-308	13	-137	-63	14	-141	47	
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-11	250	124	8	-122	82	-2	-112	110				
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-9	228	-235	4	691	-680	H=	8,	K=	3			
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-16	-138	-84	-2	-138	59				5	-127	93	
-18	-138	-13	-1	-138	-114	0	-155	201	6	-131	-61	
-14	-129	66	H=	8,	K=	8	1	-153	-184	7	-130	-127
-13	237	-163					2	-216	-276	8	-135	228
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-10	-139	-113	1	-143	1	-7	-152	106	11	-131	29	
-9	229	-150	2	-141	50	-6	-157	235	12	-137	-27	
-8	-136	-7	3	-138	65	-5	-214	-59	13	275	-244	
-7	429	439	4	212	-197	-4	-147	-79	14	-141	83	
-6	-130	700	5	-139	-154	-3	-147	-52	15	-151	207	
-5	579	-544	6	209	325	-2	-151	12	16	377	215	
-4	-123	-59	7	-141	178	-1	-149	135	15	397	423	
-3	-125	168	8	-141	-174	H=	9,	K=	0	-14	236	
-2	192	-94	9	-143	42				-13	265	-268	
-1	-129	250	10	-148	-21				-12	-135	132	
			11	-141	-101				-11	268	-267	
H=	8,	K=	6	-13	-147	77	1	-118	-92	-10	-135	-10
0	-132	36	-12	-143	-97	2	472	454	-9	190	194	
1	-137	-186	-11	-135	-88	3	222	-243	-8	-128	117	
2	-134	-76	-9	-136	94	4	225	294	-7	289	-306	
3	-132	-95	-8	-143	210	5	267	-256	-6	-124	158	
4	-137	72	-7	217	147	6	395	373	-5	723	720	
5	342	344	-6	419	-453	7	500	-493	-4	447	-425	
6	318	-218	-5	-140	-231	8	243	322	-3	163	-23	
7	514	-552	-4	-145	145	9	224	-211	-2	229	226	
8	426	419	-3	-142	107	10	198	208	-1	311	-223	
9	393	328	-2	-141	13	H=	9,	K=	3			
10	313	-344	-1	-147	-100							
11	-142	-86										
12	-142	212	H=	8,	K=	9						
13	-146	-36										
-16	-146	37	0	387	-393							
-15	-143	135	1	-138	-44							
-14	-138	-103	2	-141	202	H=	9,	K=	1			
-13	301	-205	3	-140	119							
-12	267	152	4	-138	-45							
-11	387	331	5	-146	-224							
-10	214	-1	6	-126	38							
-9	-141	-122	7	-140	31							
-8	-136	78	8	-167	62							
-7	342	-292	9	-147	-6							
-6	196	188	10	-151	33							
-5	365	334	-12	-154	150							
-4	-136	-146	-11	-145	-35							
-3	-133	-17	-10	-146	-160							
-2	278	212	-9	-143	0							
-1	-129	76	-8	-141	102							
			-7	-144	89							
H=	8,	K=	7	-6	-137	-77						
0	410	454	-5	-136	20							
1	486	467	-4	-136	-12							
2	453	-460	-3	259	-223							
3	330	-373	-2	-141	153							
4	-143	229	-1	-143	160							
5	-139	111	H=	8,	K=	10						
6	-141	-214										
7	-159	177	0	-144	-22							
8	276	223	1	-138	-65							
9	-157	-109	2	-141	-35							
10	-139	-61	3	-149	127							
11	-141	-39	4	-145	89							
12	-143	-42	5	-146	56							
-16	265	-128	6	307	-263							
-14	233	-132	7	226	-18							
-13	-143	193	-10	-152	-121							
-12	-135	20	-9	265	-232							
-11	-140	-250	-8	-142	27							
-10	-137	99	-7	236	102	H=	9,	K=	2			
-9	300	401	-6	-144	107							
-8	304	-305	-5	-142	-61							
-7	249	-252	-4	-148	-209							
-6	369	375	-3	-138	52							

L	FU	FC	L	FU	FC	L	FU	FC	L	FU	FC	
6	579	-191	-8	-136	-174	-8	-142	149	-12	-137	190	
10	210	-144	-7	-139	153	-7	-143	-74	-11	-124	161	
11	214	704	-6	451	473	-6	-144	-160	-10	-133	-300	
12	269	190	-5	206	-705	-5	242	-245	-9	-132	99	
13	-164	107	-4	466	-496	-4	-141	74	-8	216	191	
14	-143	-113	-3	210	283	-3	299	215	-7	368	-397	
-16	591	-203	-2	275	302	-2	-141	96	-6	-144	142	
-15	-136	-56	-1	361	-366	-1	302	-206	-5	-126	100	
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-13	212	302	H=	9+	K=	7	H=	S+	K=	10	-3	
-12	-132	-192	0	423	398	0	-140	11	-2	291	271	
-11	171	-133	1	450	-452	1	-151	-48	-1	678	700	
-10	-136	-111	2	325	-227	2	251	-6	H=	10+	0-	
-9	-133	2	3	204	206	3	222	257	-10	-144	513	
-8	166	226							-2	453	-419	
-7	-127	62	4	-140	65	4	-151	-21	0	494	513	
-6	475	-462	5	241	-232	5	-147	-104	1	-144	9	
-5	-130	155	6	-141	161	6	-150	-35	2	453	-419	
-4	232	223	7	-138	-131	-8	-147	256	3	-131	112	
-3	205	78	8	-142	-95	-7	-148	190	4	156	171	
-2	-122	43	9	281	227	-6	-142	-76	5	793	-281	
-1	-130	-208	10	-146	115	-5	-147	-142	6	-137	59	
			11	249	-155	-4	-148	?	7	-126	150	
H=	9+	K=	9	-14	247	-220	-3	254	164	8	-135	-63
			-13	-142	-55	-2	-140	-15	9	-136	100	
0	567	-537	-12	-144	-125	-1	257	-167	10	-136	59	
1	193	-50	-11	-143	32				11	-139	-160	
2	683	725	-10	-135	-55				12	218	-244	
3	-131	137	-9	-120	-46				13	279	234	
4	-135	-203	-8	-135	-104	0	324	-235	14	-149	227	
5	-137	133	-7	-137	48	1	340	-255	-15	-129	170	
6	-136	-20	-6	-144	121	2	284	231	-16	-136	-102	
7	-137	178	-5	-141	-145	-4	-157	20	-14	-132	83	
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9	227	-182	-3	-140	-141	-2	-147	cc	-12	-132	32	
10	261	-205	-2	473	-494	-1	256	250	-11	-136	206	
11	-140	50	-1	216	290				-10	354	319	
12	360	302							-9	276	-169	
13	-143	-102	H=	9+	K=	8			-8	454	-502	
-16	-146	-38							-7	203	165	
-15	-129	-81	0	290	-282	2	-126	100	-6	-130	28	
-14	-137	-132	1	-143	103	4	364	-218	-5	-127	-68	
-13	-135	179	2	-163	179	6	472	448	-4	-147	190	
-12	-141	187	3	-141	-199	8	378	-213	-3	254	201	
-11	-135	-41	4	239	-200	10	-136	-140	-2	274	-219	
-10	-138	-180	5	-136	83	12	368	416	-1	211	-223	
-9	-135	79	6	-135	73	14	486	-463				
-8	290	237	7	-140	-106	-16	340	-337	H=	10+	K=	
-7	-134	118	8	-141	-126	-15	-121	100	-	-	-	
-6	-130	-17	9	-144	64	-12	-136	130	0	-129	17	
-5	-133	-174	10	-141	-2	-10	212	-198	1	240	-210	
-4	156	-66	-12	-143	-34	-8	212	252	2	255	183	
-3	777	217	-11	-150	-85	-6	506	-517	3	-130	-108	
-2	-126	-105	-10	225	-123	-4	678	601	4	-126	32	
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H=	9+	K=	6	-7	-135	-118			7	273	-212	
			-6	-140	-205			8	212	-179		
0	-131	56	-5	-142	214	0	194	-255	9	-134	119	
1	-134	76	-6	-141	-190	1	221	185	10	264	71	
2	383	-214	-3	246	-193	2	228	185	11	-132	-11	
3	703	190	-2	462	446	3	123	-167	12	-129	-193	
4	-139	160	-1	207	-55	4	-121	73	13	-141	51	
5	-141	-58				5	-121	100	-14	-143	-115	
6	248	-242	H=	9+	K=	9	6	227	-219	-15	254	216
7	-141	178				7	-125	73	-16	253	-216	
8	268	286	0	252	-261	8	252	74	-12	271	-240	
9	-135	107	1	340	352	9	-134	-62	-12	250	-169	
10	-163	-134	2	-146	-236	10	245	-171	-11	261	350	
11	-146	175	3	-140	-123	11	-132	108	-10	-137	130	
12	-140	-79	4	-139	14	12	-111	165	-9	-116	-197	
-13	-140	-62	5	-146	0	13	-142	-135	-8	299	294	
-14	304	-227	6	-143	23	14	-143	-44	-7	-135	-111	
-15	-129	55	7	-140	2	15	-141	-12	-6	-129	-79	
-12	-134	116	8	-145	-98	-16	272	226	-5	234	266	
-11	-133	-7	-11	-148	-123	-15	-127	0	-4	223	271	
-10	-138	63	-10	-141	-54	-16	-136	-187	-3	-120	-167	
-9	-137	-173	-9	-142	160	-15	-121	-89	-2	369	-430	

L	FC	FC	L	FO	FC	L	FO	FC	L	FO	FC	
-1	527	505	10	213	-179	-9	-147	62	1	315	-331	
			11	-142	-235	-8	-142	70	2	286	227	
H=	10,	K=	4	-14	-141	79	-7	-140	-56	3	207	235
			-13	-138	66	-6	-144	-150	4	-152	-104	
0	272	-235	-12	-137	94	-5	-144	71	5	-157	-112	
1	279	252	-11	-134	-113	-4	-144	162	6	-134	165	
2	-130	-77	-10	-135	36	-3	-143	53	7	-135	-102	
3	-132	19	-9	217	-99	-2	-145	-208	8	-133	23	
4	-135	-84	-8	192	29	-1	-137	59	9	-133	38	
5	-137	224	-7	206	222				10	-136	-1	
6	-139	272	-6	-146	-90	H=	10,	K=	10	11	-137	
7	-136	-54	-5	281	-254				12	-141	160	
8	-133	-157	-4	-143	155	0	-146	-46	13	310	293	
9	-131	-29	-3	-134	71	1	-150	63	14	-148	-147	
10	253	297	-2	268	-286	2	-144	112	15	-139	-6	
11	251	223	-1	-139	188	3	-146	-46	16	-128	70	
12	-138	-75				4	-147	-56	17	-137	-168	
13	256	-269	H=	10,	K=	7	-6	-158	-242	18	-122	-132
-16	229	-56				-5	-145	-72	19	-137	64	
-15	-143	245	0	461	-478	-4	-220	240	20	-136	-17	
-14	-139	120	1	-129	-39	-3	-145	46	21	340	312	
-13	-134	48	2	336	360	-2	-234	-103	22	373	205	
-12	336	-407	3	-135	-172	-1	-150	-51	23	232	282	
-11	-134	-233	4	-131	-41				24	-133	-196	
-10	-139	201	5	211	174	H=	11,	K=	0	25	226	
-9	-137	195	6	-138	-212				26	-150	78	
-8	-137	-10	7	245	-220	1	609	633	27	433	-476	
-7	-136	-250	8	-141	246	2	258	-354	28	230	199	
-6	300	-220	9	234	239	5	278	395	31	904	890	
-5	296	234	10	313	-238	7	-132	-54	H=	11,	K=	
-4	218	236	-13	217	-190	9	246	243	3	129	-92	
-3	204	-227	-12	-137	43	11	193	152	4	-132	44	
-2	-130	184	-11	-134	141	13	534	-470	5	206	-202	
-1	-129	-67	-10	255	-67	-15	-141	124	6	199	1	
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3	273	-178	-3	-137	-265	-1	652	-601	13	-127	48	
4	-135	-124	-2	281	263	H=	11,	K=	1	14	-138	
5	-135	-22	-1	380	345				15	-138	-48	
6	-136	65				H=	10,	K=	8	16	-138	
7	-135	178	H=	10,	K=	8	0	619	637	17	236	234
8	-133	-8				1	-130	140	18	-135	-113	
9	332	-310	0	-139	140	2	-131	-183	19	-137	-132	
10	-133	-120	1	379	435	3	-129	11	20	-124	13	
11	271	306	2	272	-248	4	-120	20	21	-124	13	
12	-144	86	3	-128	-70	5	409	-420	22	-125	103	
-15	301	-568	4	-136	88	6	-134	64	23	-127	-132	
-14	-137	71	5	-135	-22	7	297	283	24	-130	-188	
-13	-139	169	6	-134	104	8	-137	-161	25	-129	-189	
-12	-133	-12	7	-141	-146	9	-133	62	26	-129	84	
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-10	211	-157	9	-145	81	11	-128	131	28	-129	470	
-9	-130	178	-11	-142	-58	12	-135	-82	29	-121	-101	
-8	-136	23	-10	-135	114	13	-144	145	30	516	550	
-7	243	-190	-9	213	180	14	-142	76	31	-136	-216	
-6	-136	-213	-8	210	-281	15	272	-267				
-5	246	-139	-7	-135	-164	H=	11,	K=	4			
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-3	608	641	-5	-135	200	-12	-135	52	1	293	260	
-2	224	-212	-4	-136	-48	-11	-135	210	2	340	318	
-1	746	-734	-3	-135	57	-10	214	-210	3	356	340	
			-2	-140	122	-9	-135	74	4	336	268	
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1	219	-30				-6	-133	-211	7	-130	-43	
2	239	-76	0	273	169	-5	-131	-229	8	-136	-69	
3	-140	34	1	-143	-67	-4	210	750	9	199	57	
4	-138	-151	2	-145	-63	-3	220	-210	10	-137	46	
5	217	-274	3	-135	14	-2	504	-531	11	-135	-83	
6	-130	75	4	-143	-130	-1			12	266	-212	
7	-137	232	5	207	-10	H=	11,	K=	2	-140	-1	
8	-133	43	6	-152	182				3	-143	7	
9	-135	24	7	-144	154	0	-132	-165				

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
-13	203	-105	6	370	-248	6	240	242	-2	-135	-148
-12	-131	-79	7	227	227	7	-133	-67	-1	-137	-214
-11	-132	128	8	-149	112	8	184	10			
-10	547	365	9	243	-167	9	-143	145	H=	12,	K= -4
-9	-131	-11	-12	-143	78	10	-137	-36			
-8	335	-275	-11	-144	-25	11	-141	-34	0	-138	-58
-7	-139	66	-10	-143	23	-15	-139	62	1	546	-503
-6	-134	-17	-9	-140	22	-14	-141	65	2	-139	13
-5	159	179	-8	-137	-138	-13	-133	22	3	-135	77
-4	-133	50	-7	-137	-47	-12	-136	-26	4	-134	215
-3	-137	-197	-6	238	228	-11	-132	-111	5	-131	-76
-2	365	-294	-5	-137	5	-10	-132	-273	6	-132	-50
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			-2	-140	3	-7	-122	71	9	-135	55
			-1	-160	-207	-6	280	-212	10	-142	142
0	421	372				-5	-140	176	-14	-145	17
1	-130	-110	H=	11,	K= 8	-4	341	258	-13	267	-49
2	-138	-116				-3	-134	-151	-12	-139	60
3	265	-250	0	239	210	-2	622	-650	-11	-134	100
4	-139	-125	1	-141	-154	-1	-131	5	-10	-127	-75
5	-136	127	2	-144	-116				-5	-136	-193
6	-136	178	3	-144	24				-8	305	-285
7	-132	68	4	-145	113				-7	-134	47
8	-138	-184	5	-143	-39	0	835	-824	-6	311	170
9	-142	-53	6	-149	120	1	-135	227	-5	-139	219
10	-147	198	7	-143	-58	2	-132	40	-4	-140	-268
11	274	107	-10	-145	97	3	-134	-173	-3	408	-409
-14	234	170	-9	270	-120	4	-139	169	-2	-136	-41
-13	-143	-5	-8	-140	-124	5	-121	34	-1	794	810
-12	253	-219	-7	266	194	6	191	-159			
-11	-137	-137	-6	-138	-29	7	-130	-44	H=	12,	K= 5
-10	-138	192	-5	-138	-64	8	-136	107			
-9	-137	172	-4	-142	91	9	-134	0	0	225	161
-8	-140	63	-3	-140	-75	10	-145	-100	1	-124	72
-7	-134	-104	-2	-141	-89	11	270	208	2	-137	-100
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-5	-132	-5				-14	-142	-144	4	-131	45
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-3	-137	-3				-12	-137	116	6	-142	-95
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0	548	-523	-7	-149	-154	-6	-134	153	-11	-136	-31
1	262	249	-6	-140	-22	-5	-130	2	-10	204	167
2	363	-264	-5	242	226	-4	218	-167	-9	-136	-1
3	-139	-86	-4	-141	-15	-3	-124	49	-8	219	-65
4	314	-336	-3	214	-151	-2	608	817	-7	-126	-160
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6	-138	195	-1	-144	190				-5	-136	222
7	225	-96							-4	-128	-10
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9	-143	-91				0	-138	139	-2	369	-311
10	-149	6	0	947	991	1	212	-179	-1	356	-297
-13	-142	-103	2	-133	-136	2	-141	-79	H=	12,	K= 6
-12	-143	63	4	-138	353	3	-136	-30			
-11	-135	-3	6	250	-218	4	-139	57			
-10	-136	-58	8	-147	3	5	-137	-101	0	-135	-72
-9	-138	80	10	-143	186	6	-133	-41	1	346	37
-8	-124	25	12	294	-357	7	150	-60	2	-125	-160
-7	-135	-67	13	-137	-43	8	-135	-61	3	207	-172
-6	211	-131	12	195	127	9	-135	-9	4	-126	141
-5	-131	-51	13	-128	-1	10	-139	10	5	-141	30
-4	-135	179	8	-135	-156	11	-146	-162	6	287	-159
-3	-138	-128	6	-131	11	-10	-144	-119	7	-147	118
-2	326	291	4	380	269	-12	-124	-3	8	-143	-68
-1	-137	-53	-2	1112	-1090	-14	-128	-6	-12	-143	-10
H=	11,	K= 7	H=	12,	K= 1	-13	-129	73	-10	-140	11
						-9	209	-273	-9	214	-203
0	253	256	0	-130	104	-8	-139	-153	-8	206	-113
1	-137	171	1	-131	-52	-7	217	327	-7	-135	30
2	290	-200	2	-131	49	-6	-137	174	-6	-145	262
3	-139	5	3	279	207	-5	-136	-707	-5	-134	-154
4	-137	146	4	-130	-161	-4	250	-241	-4	-130	52
5	-140	-124	5	-130	112	-3	421	523	-5	-137	32

L	FC	FC	L	FO	FC	L	FO	FC	L	FO	FC	
-2	192	73	H=	13,	K=	0	5	-134	-177	2	-135	-41
-1	304	-263		1	-136	-86	6	-139	-16	3	-140	-20
H=	12,	K=	7	3	202	98	7	299	292	4	195	124
				5	-135	167	8	-138	-51	5	230	-13
0	193	128		7	207	-190	9	233	-158	6	-141	-206
1	200	-196		9	212	160	10	-143	46	7	-141	-77
2	-143	127	-13	-140	32	-12	-143	-120	8	-146	128	
3	205	135	-11	-136	38	-11	198	-166	9	-143	46	
4	-142	-67	-9	-134	-100	-10	-133	187	-12	-145	100	
5	-143	-48	-7	-134	0	-9	-138	99	-11	-140	44	
6	-146	141	-5	-138	30	-8	-136	-150	-10	-140	-153	
7	-140	-16	-3	683	-674	-7	-134	99	-9	264	203	
-10	-144	7	-1	430	442	-6	-130	39	-8	-136	70	
-9	-147	-184				-5	390	-270	-7	194	-20	
-8	-140	17	H=	13,	K=	1	-4	233	143	-6	-133	154
-7	202	270				-3	505	552	-5	-140	-245	
-6	-140	-100	0	-132	-139	-2	344	-286	-4	400	-404	
-5	-143	-109	1	-133	117	-1	-138	-173	-3	230	100	
-4	-130	144	2	-130	65				-2	406	428	
-3	201	164	3	285	-284	H=	13,	K=	-1	268	195	
-2	-134	-102	4	-135	19							
-1	-137	-72	5	298	294	0	202	-223	H=	13,	K=	
			6	-138	-179	1	-133	-117	5	-135	-99	
H=	12,	K=	8	7	-135	-128	2	-137	150	1	-139	-143
			8	209	111	3	-136	29	2	-135	54	
0	298	-195	9	-138	24	4	298	-249	3	-138	193	
1	232	-135	10	-146	-37	5	-134	-74	4	240	228	
2	-147	63	-14	-146	-83	6	213	246	5	-138	-80	
3	-141	28	-13	-139	55	7	-140	-160	6	290	-298	
4	-144	-42	-12	-139	106	8	-141	65	7	-143	111	
5	-150	171	-11	-136	-195	9	-144	64	8	257	99	
-8	254	143	-10	-133	-18	-12	-138	-33	-11	-142	-12	
-7	-144	-104	-9	266	298	-12	-139	123	-10	-141	149	
-6	-143	28	-8	-130	-8	-11	-141	109	+9	-140	66	
-5	-140	144	-7	-130	-116	-10	-139	-137	-8	-143	-264	
-4	-144	-194	-6	-132	-22	-9	553	-285	-7	-142	60	
-3	275	-102	-5	271	169	-8	-135	210	-6	318	293	
-2	-148	195	-4	218	-310	-7	-136	182	-5	-134	121	
-1	293	184	-3	258	-257	-6	240	-274	-4	241	-221	
			-2	220	270	-5	410	-417	-3	-138	-207	
H=	12,	K=	9	-1	277	-196	-4	-135	69	-2	-135	140
						-3	547	525	-1	315	287	
0	-191	55	H=	13,	K=	2	-2	-143	273	H=	13,	K=
1	-145	-74				-1	-136	-262	6			
-4	-151	-90	0	210	78				-10	-142	16	
-3	-146	-90	1	226	-193	H=	13,	K=	-9	-141	-123	
-2	-150	28	2	-131	-64				-8	-143	39	
-1	-146	81	3	-136	29	0	210	-248				
			4	-132	89	1	-136	-15				

Table B-2
Observed and Calculated Structure Factors for $[\text{Co}(\text{Hdmg})_2(\text{Clan})_2]\text{Cl}$

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC	
H= 0, K= 0	0	155	-142	H= 0, K= 0	6	15	-6					
1 765	730	3	134	-126	1	134	-126	7	238	230		
2 469	420	4	40	35	2	95	-80	8	241	254		
3 137	121	5	28	30	3	116	-113	9	64	62		
4 202	177	6	295	270	4	137	-113	10	229	234		
5 437	405	7	140	137	5	135	131	11	126	131		
6 48	35	8	135	131	6	172	172	12	122	126		
7 357	342	9	184	184	7	43	43	13	42	41		
8 75	73	10	60	61	8	70	74	14	147	155		
9 229	219	11	95	95	9	68	63	15	193	194		
10 55	58	12	-12	-10	-10	131	130	9	111	110		
11 230	229	-12	60	69	-9	92	91	-7	231	223		
12 103	108	-11	-12	-15	-8	101	99	-6	414	384		
13 43	39	-10	42	-44	-7	75	70	-5	24	25		
H= 0, K= 1	-9	135	133	-6	140	141	-4	311	300			
0 651	638	-6	15	23	-5	200	204	-3	30	-21		
1 426	400	-5	234	238	-2	190	188	-2	43	45		
2 349	309	-4	173	163	-1	185	189	H= 1, K= 1	653	627		
3 177	164	-3	41	29								
4 235	-217	-2	304	-292	H= 0, K= 7	0	238	-229				
5 172	161	-1	239	-225	0	76	78	1	99	94		
6 315	-276	H= 0, K= 4			1	27	28	2	315	-301		
7 226	191		211	199	2	59	58	3	110	-115		
8 101	105	0	194	191	3	124	123	4	343	330		
9 60	54	1	333	332	4	97	100	5	72	-61		
10 190	193	2	333	329	5	54	48	6	223	217		
11 81	83	3	235	224	6	-11	0	7	34	32		
12 174	177	4	202	201	7	32	37	8	287	281		
13 27	21	5	328	329	8	110	112	9	-11	11		
-13 53	57	6	132	131	9	-12	2	10	127	127		
-12 44	-48	7	204	199	-8	-11	10	11	17	32		
-11 101	103	8	68	67	-7	-11	-2	12	-12	-4		
-10 -14	15	9	138	136	-6	-11	-7	-13	-12	14		
-9 67	-62	10	60	58	-5	69	68	-12	60	60		
-8 -12	-8	11	132	137	-4	214	211	-11	100	99		
-7 31	1	12	109	107	-3	242	247	-10	144	148		
-6 565	519	-12	104	105	-2	126	128	-9	175	-173		
-5 231	216	-11	104	105	-1	61	64	-8	166	177		
-4 349	337	-10	146	147				-7	55	53		
-3 103	-103	-9	224	230				-6	88	69		
-2 64	-73	-8	75	73	H= 0, K= 8	-5	164	-164				
-1 443	431	-7	194	190	0	104	111	-4	399	-370		
H= 0, K= 2	-6	160	155	1	67	65	-3	285	292			
	-5	210	214	2	93	99	-2	176	-174			
0 475	-471	-3	31	30	3	111	112	-1	426	421		
1 394	367	-2	21	19	4	103	106	H= 1, K= 2				
2 299	279	-1	112	-102	5	54	57					
3 204	198	H= 0, K= 5			6	90	90	0	413	394		
4 146	131		-10	8	-7	83	83	1	59	53		
5 40	-36		-5	51	-6	51	52	2	411	392		
6 487	464	0	271	266	-5	79	78	3	81	85		
7 269	256	1	193	190	-4	193	194	4	284	260		
8 320	303	2	89	94	-3	280	282	5	37	38		
9 115	114	3	89	94	-4	273	276	6	241	225		
10 297	291	4	58	54	-1	192	193	7	194	188		
11 74	78	5	138	140				8	213	204		
12 91	87	6	93	91	H= 0, K= 9	9	9	9	322	323		
-13 34	38	7	21	20	0	-11	5	10	81	79		
-12 52	47	8	59	-61	1	-11	17	11	70	72		
-11 36	38	9	51	-49	2	26	30	12	73	-78		
-10 102	103	10	28	26	3	26	24	-13	74	74		
-9 -14	21	-11	92	94	-2	45	45	-12	145	151		
-8 238	241	-12	195	196	-1	25	29	-11	110	113		
-7 67	73	-9	90	93				-9	146	144		
-6 486	472	-8	16	-9				-8	126	132		
-5 547	512	-7	20	-16	H= 1, K= 0	0	699	676	-7	269	266	
-4 476	450	-6	61	60	1	250	242	-6	140	143		
-3 458	450	-5	60	53	2	45	40	-5	159	160		
-2 255	228	-4	34	-30	3	671	651	-4	44	-37		
-1 673	668	-3	73	-72	4	337	324	-3	306	323		
H= 0, K= 3	-2	36	42	3	5	320	491	-2	300	296		
	-1	25	-19	4	5	20		-1	302	233		

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC	
H=	1+	K=	-3	H=	1+	K=	6	H=	1+	K=	-7	
0	156	159	0	221	225	-5	127	125	0	244	242	
1	237	-131	1	154	182	-4	175	175	1	163	161	
2	146	140	2	166	166	-2	152	157	2	595	394	
3	247	219	3	176	172	-1	104	107	3	627	617	
4	33	37	4	188	197		63	65	4	164	163	
5	130	127	5	244	233		0	27	7	144	141	
6	145	-142	6	14	151		1	30	8	126	113	
7	157	156	7	37	34		2	33	9	126	117	
8	-11	-12	8	102	104		3	63	10	75	63	
9	129	123	-10	105	104		3	107	107	-1	12	26
10	0	68	-9	72	69		4	33	-12	105	109	
11	-11	9	-8	30	28		5	16	-11	114	115	
-13	96	99	-7	113	110		6	38	-10	71	75	
-12	101	102	-6	200	252		7	100	-8	170	168	
-11	-11	10	-5	309	314		8	99	-7	133	134	
-10	108	-109	-4	246	240		-5	93	-7	-11	-72	
-9	133	137	-3	176	179		-7	41	-6	232	251	
-8	50	47	-2	160	158		-6	40	-5	765	269	
-7	96	96	-1	152	153		-5	36	-4	300	356	
-6	-10	15					-4	52	-3	214	207	
-5	29	28	H=	1+	K=	7	-3	45	-2	194	193	
-4	257	249					-2	46	-1	381	380	
-3	131	-132	0	53	59		-1	22	H=	1+	K=	
-2	186	178	1	16	-14				1	71	66	
-1	183	177	2	-11	-5	H=	1+	K=	0	629	-617	
H=	1+	K=	4	4	52	52	0	134	141	1	162	152
0	298	295	6	-12	0	1	229	228	2	375	361	
1	405	400	7	93	101	2	242	243	3	314	297	
2	233	230	-2	-12	19	3	165	165	5	59	-53	
3	294	284	-3	-12	-17	4	124	127	6	70	67	
4	246	244	-7	30	-28	5	120	125	7	30	36	
5	208	203	-6	22	-20	6	101	101	8	159	162	
6	142	136	-5	36	35	7	113	114	9	-11	3	
7	113	110	-4	79	80	8	112	111	10	-11	-9	
8	28	26	-3	68	64	10	134	126	11	-11	-19	
9	25	22	-2	62	62	-10	113	112	12	26	25	
10	148	151	-1	77	75	-9	83	85	-12	62	62	
-12	62	86				-8	97	94	-11	16	+21	
-11	65	64	H=	1+	K=	8	-7	121	123	-10	107	105
-10	128	127	0	95	98	-6	78	72	-9	-11	-12	
-9	127	127				-5	115	117	-8	79	82	
-8	223	232	1	114	116	-4	141	141	-7	40	37	
-7	266	267	2	114	116	-3	129	125	-6	231	219	
-6	263	291	3	104	107	-2	73	70	-5	165	163	
-5	254	251	4	100	98	-1	79	76	-4	632	521	
-4	231	240	-7	82	80	H=	1+	K=	-3	542	507	
-3	114	117	-6	63	63	0	99	98	-2	266	268	
-2	320	319	-5	54	52				-1	265	261	
-1	495	489	-4	84	87							
H=	1+	K=	5	-2	119	122	0	145	143	H=	1+	K=
0	152	140	-1	82	83	2	145	143	0	105	-102	
1	314	313	H=	1+	K=	-9	5	55	1	411	392	
2	136	138				6	62	61	2	310	328	
3	39	44	0	66	68	7	40	38	3	143	184	
4	139	137	1	113	119	8	28	32	4	503	482	
5	95	94	2	64	83	9	63	66	6	405	396	
6	183	179	2	21	19	10	77	79	7	143	119	
7	99	-105	-3	36	33	11	37	40	8	191	149	
8	46	-44	-2	34	37	-11	119	124	9	264	221	
9	56	54	-1	25	24	-10	46	41	10	31	29	
-11	38	37				-9	71	65	11	76	70	
-10	71	76	H=	1+	K=	-8	-6	-11	-7	12	61	
-9	-11	0				-7	41	-41	-12	33	35	
-8	55	52	0	109	110	-6	75	-63	-12	106	103	
-7	179	175	1	107	104	-5	72	69	-11	57	57	
-6	298	302	2	174	177	-4	-10	-2	-10	216	201	
-5	244	251	3	180	186	-3	17	-15	-9	764	745	
-4	20	-11	4	136	139	-2	16	11	-8	167	193	
-3	-9	14	5	102	101	-1	48	-51	-7	436	404	
-2	13	4	6	122	123				-6	23	26	
-1	67	76	7	131	131	H=	1+	K=	-5	312	291	

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC	
-4	421	384	-11	177	178	-11	111	2	-5	62	64	
-3	1149	1097	-10	-12	8	-10	60	61	-4	137	132	
-2	1301	1250	-9	42	44	-9	170	173	-3	209	215	
-1	189	167	-8	48	-62	-8	342	349	-2	140	143	
H=	1,	K=	-1	-7	97	102	-7	344	349	-1	83	83
0	874	852	-4	50	52	-4	179	180	H=	2,	K=	
1	67	-60	-3	395	384	-3	216	224	0	22	16	
2	120	110	-2	90	88	-2	90	87	1	-11	4	
3	112	112	-1	94	-84	-1	171	175	2	26	-26	
4	85	-71							-3	33	31	
5	406	383	H=	2,	K=	2	H=	2,	K=	5	-2	
6	19	-23							-1	17	16	
7	130	135	0	441	445	0	35	35	H=	2,	K=	
8	-10	20	1	172	181	1	64	62	0	79	81	
9	225	222	2	282	275	2	98	102	1	79	83	
10	46	45	3	244	238	3	112	108	2	98	100	
11	105	106	4	107	-100	4	149	156	3	114	113	
12	87	92	5	147	153	5	212	210	4	101	103	
13	43	40	6	180	-177	6	150	150	5	115	113	
12	18	-6	7	271	266	7	43	38	6	130	132	
11	62	61	8	187	189	8	73	69	7	102	105	
10	19	24	9	164	159	-11	23	23	8	102	105	
9	359	359	10	80	85	-10	-11	6	-7	120	120	
8	186	176	-13	98	103	-9	27	32	-6	82	83	
7	316	302	-12	76	82	-8	32	29	-5	82	83	
6	86	-88	-11	163	167	-7	117	121	-4	69	72	
5	45	-38	-10	77	80	-6	104	102	-3	65	85	
4	47	-52	-9	268	259	-5	63	65	-2	93	102	
3	18	3	-8	141	147	-4	49	46	-1	100	108	
2	734	686	-7	184	178	-3	-10	-6	H=	2,	K=	
-1	166	-177	-6	170	182	-2	24	-26	0	37	35	
			-5	88	95	-1	79	-80	-1	46	51	
H=	2,	K=	0	-4	679	670	H=	2,	K=	6	2	
			-3	581	564		H=	2,	K=	3	79	
0	264	232	-2	672	646				4	51	52	
1	495	467	-1	319	301	0	164	169	5	26	26	
2	301	285				1	176	178	6	32	32	
3	190	184	H=	2,	K=	3	2	177	182	7	44	
4	423	413				3	195	203	8	45	45	
5	119	120	0	108	-108	4	139	141	-9	26	28	
6	438	454	1	71	72	5	160	167	-8	47	45	
7	93	97	2	103	100	6	49	47	-7	61	66	
8	121	121	3	156	152	7	85	89	-6	-11	-5	
9	108	-110	4	43	-38	-10	113	110	-5	-16	-20	
10	-11	-6	5	120	-123	-9	93	96	-4	-11	-15	
11	85	92	6	32	36	-8	44	47	-3	28	-25	
-13	48	48	7	221	217	-7	64	64	-2	-11	-1	
-12	79	84	8	185	186	-6	99	99	-1	19	7	
-11	119	122	9	47	47	-5	95	94	H=	2,	K=	
-10	329	329	10	152	156	-4	43	46	0	407	415	
-9	35	33	-12	-12	0	-3	35	28	1	316	313	
-8	193	193	-11	47	-49	-2	179	169	2	165	162	
-7	18	-4	-10	82	78	-1	161	182	3	119	122	
-6	22	13	-9	199	197	H=	2,	K=	7	119	122	
-5	272	256	-8	304	310				8	152	155	
-4	94	-35	-7	108	109	0	38	34	5	158	165	
-3	368	350	-6	50	64	1	59	63	6	73	71	
-2	66	58	-5	63	65	2	116	110	7	48	42	
-1	534	524	-4	203	201	3	124	129	8	-11	7	
H=	2,	K=	1	-2	209	205	4	38	39	9	22	29
			-1	385	374	5	57	-54	9	22	29	
0	648	627				-8	-11	6	-10	92	96	
1	188	176	H=	2,	K=	4	-7	-11	-9	103	106	
2	201	194				-6	-11	-4	-8	155	157	
3	170	-150	0	92	95	-5	-11	-8	-7	167	165	
4	24	-18	1	208	206	-4	55	-60	-6	88	90	
5	101	-98	2	308	306	-3	21	15	-5	117	113	
6	22	-21	3	231	237	-2	66	68	-4	100	97	
7	82	83	4	283	282	-1	43	45	-3	117	115	
8	48	46	5	40	47				-2	126	124	
9	78	82	6	340	332	H=	2,	K=	8	-1	176	
10	116	-117	7	168	169	0	74	76	H=	2,	K=	
11	56	56	8	142	140	1	75	77	0	364	370	
-13	-12	-9	9	169	173	2	109	109				
-12	40	43	-12	62	62							

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC
1	225	230	1	383	363	-1	126	119	H=	3+	K= -4
2	126	124	2	326	318	H=	3+	K= -1	0	295	297
3	20	14	3	256	263				1	152	155
4	25	11	4	171	115				2	152	156
5	69	69	5	454	439	0	293	376	3	111	111
6	20	-14	6	243	250	1	67	69	4	120	118
7	42	-42	7	280	282	2	143	-134	5	198	204
8	36	-79	8	174	161	3	89	96	6	233	233
9	27	21	9	263	249	4	-10	9	7	164	179
10	66	69	10	103	98	5	372	374	8	36	41
-11	69	69	11	100	105	6	48	-47	-11	42	42
-10	35	24	-13	66	73	7	121	122	-10	-11	-6
-9	39	31	-12	58	65	8	95	93	-9	103	105
-8	60	64	-11	122	120	9	-11	12	-8	80	83
-7	45	45	-10	123	123	-13	66	67	-7	100	99
-6	239	236	-9	360	362	-12	115	117	-6	74	74
-5	199	199	-8	152	143	-11	290	290	-5	161	165
-4	85	84	-7	241	239	-10	135	147	-4	310	314
-3	43	-29	-6	115	149	-9	114	106	-3	172	172
-2	56	-100	-5	176	163	-8	22	22	-2	152	150
-1	220	220	-4	643	406	-7	64	63	-1	171	173
			-3	257	269	-6	173	170			
H=	2+, K= -4		-2	402	381	-5	17	12	H=	3+, K= 5	
			-1	349	-341	-4	324	319			
0	128	121				-3	97	92	0	143	147
1	234	229	H=	2+, K= -1		-2	137	126	1	132	133
2	329	323				-1	97	99	2	39	40
3	233	230	0	319	-310	H=	3+, K= 2		3	26	-24
4	190	193	1	246	252				4	45	-47
5	65	63	2	173	161				5	72	73
6	203	199	3	318	314	0	296	297	6	45	41
7	146	148	4	116	113	1	373	379	-10	24	27
8	104	103	5	417	409	2	181	175	-9	31	34
9	91	87	6	202	194	3	181	176	-8	11	17
10	153	154	7	51	51	4	222	237	-7	27	27
-12	140	162	8	59	59	5	100	176	-6	21	21
-11	108	109	9	62	-70	6	100	166	-5	64	58
-10	120	129	10	6	64	7	222	222	-4	91	90
-9	40	39	11	38	33	8	241	248	-3	77	-74
-8	68	63	-13	102	101	9	34	33	-2	22	22
-7	119	119	-12	-11	-11	-13	49	50	-1	93	99
-6	473	475	-11	21	-21	-12	92	90			
-5	579	574	-10	228	225	-11	192	189	H=	3+, K= 6	
-4	512	498	-9	104	107	-10	301	303			
-3	280	267	-8	92	92	-9	177	179	0	139	136
-2	167	176	-7	162	-145	-8	222	225	1	190	199
-1	548	549	-6	79	79	-7	121	121	2	171	169
			-5	29	-10	-6	191	180	3	108	112
H=	2+, K= -3		-4	17	15	-5	75	80	4	24	18
			-3	234	-230	-4	147	141	5	-12	-5
0	138	-133	-2	222	-200	-3	281	283	-9	107	102
1	-9	4	-1	204	-203	-2	225	217	-8	90	89
2	73	70				-1	495	481	-7	121	115
3	36	31	H=	3+, K= 0		H=	3+, K= 3		-6	157	166
4	85	87							-5	211	208
5	30	-24	0	479	465				-4	58	56
6	241	241	1	97	99	0	73	77	-3	60	44
7	233	221	2	64	105	1	140	146	-2	106	102
8	119	122	3	23	23	2	292	260	-1	99	101
9	138	136	4	144	151	3	55	59			
10	77	75	5	303	303	4	175	172	H=	3+, K= 7	
11	126	122	6	246	245	5	38	-33			
-12	104	100	7	32	77	6	224	200	0	-11	-7
-11	106	110	8	34	-39	7	147	153	1	44	42
-10	61	57	9	114	114	8	52	50	2	127	120
-9	80	78	10	46	43	-12	12	12	-7	57	56
-8	33	-26	-13	68	59	-11	-11	-12	-5	52	59
-7	97	67	-12	122	121	-10	23	22	-6	57	56
-6	41	41	-11	242	244	-9	58	62	-5	52	59
-5	349	330	-10	14	138	-8	64	66	-4	30	26
-4	408	391	-9	196	194	-7	42	33	-3	66	69
-3	319	338	-8	107	114	-6	-10	15	-2	28	19
-2	333	312	-7	324	324	-5	88	87	-1	46	-27
-1	20	-2	-6	44	422	-4	33	-31			
			-5	502	493	-3	120	130	H=	3+, K= -3	
H=	2+, K= -2		-4	206	210	-2	-9	2			
			-3	192	189	-1	147	115	0	38	35
0	316	306	-2	330	328				1	77	81

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC
2	110	113	4	216	220	4	211	215	7	46	47
3	87	88	5	129	126	5	86	89	-12	56	56
4	46	42	6	178	182	6	233	239	-11	92	99
-6	87	91	7	176	174	7	100	-105	-10	82	79
-5	93	94	8	121	126	8	66	-63	-9	28	35
-4	134	138	9	170	173	9	-11	-8	-8	83	81
-3	191	197	-12	155	155	10	23	25	-7	169	172
-2	219	227	-11	178	186	-13	39	40	-6	136	143
-1	118	117	-10	61	55	-12	-12	15	-5	134	133
H=	3, K=	-7	-9	57	57	-11	47	-48	-4	63	60
			-8	49	39	-10	169	161	-3	365	378
			-7	129	127	-9	-11	2	-2	247	251
0	35	32	-6	152	148	-8	66	65	-1	208	204
1	27	26	-5	157	154	-7	104	102			
2	33	37	-4	63	80	-6	195	188	H=	4, K=	3
3	46	45	-3	109	105	-5	376	365			
4	-11	1	-2	343	338	-4	112	-114	0	-10	-6
5	-11	8	-1	378	383	-3	156	155	1	107	-100
6	26	27				-2	168	165	2	27	-26
-8	157	166	H=	3, K=	-3	-1	150	153	3	17	-14
-7	115	125							4	25	19
-6	17	14	0	197	202	H=	4, K=	0	5	91	90
-5	-13	0	1	97	94				6	-11	14
-4	41	40	2	142	148	0	219	212	-11	32	32
-3	125	128	3	263	254	1	221	234	-10	36	-41
-2	132	138	4	28	29	2	200	205	-9	33	30
-1	29	22	5	25	21	3	402	398	-8	19	-9
H=	3, K=	-6	6	36	-38	4	205	205	-7	39	37
			7	62	62	5	411	425	-6	-11	24
			8	62	62	6	124	126	-5	63	-62
0	283	285	9	136	143	7	49	60	-4	153	165
1	189	189	10	124	119	8	107	108	-3	228	231
2	144	145	-12	103	99	-12	160	155	-2	212	214
3	173	171	-11	46	49	-11	180	178	-1	61	61
4	199	196	-10	133	-125	-10	136	144			
5	156	157	-9	23	14	-9	165	159	H=	4, K=	4
6	70	68	-8	19	-15	-8	122	129			
7	90	87	-7	-10	19	-7	110	105	0	122	123
-10	97	104	-6	32	-24	-6	247	238	1	45	48
-9	187	189	-5	71	-72	-5	143	141	2	109	-115
-8	300	304	-4	108	114	-4	242	262	3	17	13
-7	277	281	-3	32	-29	-3	159	158	4	43	38
-6	215	217	-2	149	132	-2	322	317	5	113	117
-5	138	139	-1	103	102	-1	204	199	-10	90	56
-4	133	128							-9	137	139
-3	143	146	H=	3, K=	-2	H=	4, K=	1	-8	177	176
-2	64	57							-7	147	146
-1	165	161	0	36	39	0	181	178	-6	102	94
H=	3, K=	-5	1	206	202	1	185	186	-5	102	96
0	96	98	4	276	273	2	142	144	-4	223	226
1	74	75	5	196	201	3	220	221	-3	85	82
2	86	86	6	55	44	4	214	205	-2	190	189
3	104	104	7	30	-23	7	-11	0	H=	4, K=	5
4	217	220	8	82	76	8	110	114			
5	130	131	9	62	63	-12	63	68	0	85	89
6	60	58	10	120	122	-11	79	74	1	-11	17
7	53	50	-13	117	117	-10	126	130	2	57	-62
8	84	81	-12	93	98	-9	-11	3	3	22	19
-11	64	65	-11	31	-23	-8	90	90	4	62	61
-10	68	67	-10	86	91	-7	-10	4	-9	47	46
-9	-11	10	-9	46	44	-6	-10	-2	-6	87	86
-8	77	73	-8	143	138	-5	-10	4	-7	109	113
-7	131	132	-7	136	131	-4	70	-72	-9	91	91
-6	190	189	-6	212	221	-3	91	94	-5	118	115
-5	104	104	-5	242	233	-2	107	105	-4	-11	-17
-4	-10	0	-4	362	354	-1	312	315	-3	78	-61
-3	32	-23	-3	270	263				-2	-11	-7
-2	42	-26	-2	218	214	H=	4, K=	2	-1	84	85
-1	175	171	-1	290	279	0	196	201	H=	4, K=	6
H=	3, K=	-4	H=	3, K=	-1	1	137	139			
0	266	266	0	192	-186	2	250	253	0	145	146
1	214	210	1	47	-33	3	155	157	1	111	109
2	264	271	2	47	-50	4	293	296	2	146	143
3	253	260	3	123	-118	5	-11	10	-7	149	151
						6	47	46	-6	175	177

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC	
-5	127	133	5	-11	-14	-8	92	91	-7	-11	-6	
-4	-12	15	6	-11	3	-7	191	184	-6	48	-58	
-3	51	47	7	62	86	-6	195	196	-5	100	-99	
-2	82	84	-11	119	119	-5	166	163	-4	133	129	
-1	116	111	-10	37	39	-4	104	102	-3	131	134	
H=	4,	K=	-8	-9	103	101	-3	121	119	-2	85	84
			-8	97	98	-2	185	182	-1	39	35	
			-7	129	127	-1	186	-189				
0	82	80	-6	149	148				H=	5,	K=	
1	121	121	-5	68	71	H=	5,	K=	0	0	125	
-5	89	91	-4	149	149					1	120	
-4	119	121	-3	140	140	0	174	173		2	94	
-3	175	179	-2	171	170	1	223	227		-8	108	
-2	178	182	-1	169	181	2	175	174		-7	62	
-1	86	68				3	83	84		-6	34	
			H=	4,	K=	-3	4	98			29	
H=	4,	K=	-7			5	94	102		-5	123	
			0	70	-65	6	125	120		-4	152	
0	49	49	1	17	12	-11	-12	14		-3	165	
1	88	89	2	122	123	-10	140	147		-2	165	
2	112	114	3	117	122	-9	111	118		-1	163	
3	80	78	4	-11	-20	-8	144	144	H=	5,	K=	
4	56	53	5	151	-146	-7	119	124	0	5,	5.	
-7	59	66	6	68	-76	-6	200	191				
-6	30	29	7	49	53	-5	319	321	-6	23	30	
-5	42	40	8	99	100	-4	123	120	-5	54	56	
-4	59	61	-12	85	87	-3	87	87	-4	61	62	
-3	76	77	-11	25	-25	-2	95	92	-3	89	89	
-2	26	20	-10	17	19	-1	183	184	-2	95	96	
-1	-11	-3	-9	69	66				-1	107	105	
			-8	125	127	H=	5,	K=	1			
H=	4,	K=	-6	-7	57	59			H=	5,	K=	
			-6	-11	5	0	22	11	-7			
0	157	188	-5	101	102	1	-11	-3	0	24	-20	
1	170	171	-4	220	224	2	180	185	1	28	-22	
2	152	154	-3	106	103	3	81	79	-5	82	83	
3	153	154	-2	20	-20	4	90	94	-4	96	97	
4	200	193	-1	71	71	5	-11	-8	-3	106	105	
5	173	175				-11	36	48	-2	89	90	
6	169	152	H=	4,	K=	-2	-10	55	54	-1	44	
-9	103	109				-9	32	34				
-8	125	125	0	78	-71	-8	27	27	H=	5,	K=	
-7	101	104	1	49	53	-7	59	60	-6			
-6	104	105	2	203	202	-6	133	135	0	111	112	
-5	144	147	3	172	174	-5	221	215	1	33	33	
-4	171	168	4	210	208	-4	29	-17	2	25	-20	
-3	147	145	5	38	-40	-3	29	23	3	-12	15	
-2	84	87	6	124	127	-2	73	-68	-7	93	92	
-1	155	156	7	144	142	-1	100	-106	-6	115	117	
			8	135	142				-5	115	115	
H=	4,	K=	-5	-12	84	83	H=	5,	K=	2	-4	
			-11	-11	11				-3	107	108	
0	185	186	-10	141	140	0	-11	2	-2	146	151	
1	163	165	-9	179	188	1	60	62	-1	147	150	
2	57	59	-8	285	284	2	151	154	H=	5,	K=	
3	-11	4	-7	202	267	3	122	128	0	136	138	
4	35	33	-6	143	146	4	72	77	1	70	67	
5	48	44	-5	335	326	5	131	128	2	46	-48	
6	66	64	-4	282	270	-10	116	117	3	45	-46	
7	93	88	-3	332	324	-9	117	116	-7	52	53	
-10	44	43	-2	199	190	-8	70	72	-8	59	57	
-9	-12	6	-1	89	93	-7	90	82	-7	107	105	
-8	-11	-4				-6	113	117	-6	106	106	
-7	20	18	H=	4,	K=	-1	-5	25	22	-5	106	
-6	20	21				-4	203	211	-6	-11	-10	
-5	20	-23	0	73	-65	-3	245	247	-5	43	-50	
-4	21	-19	1	23	22	-2	107	106	-4	45	-44	
-3	43	43	2	195	163	-1	51	-47	-3	33	-33	
-2	87	87	3	183	174				-2	41	39	
-1	116	113	4	99	110	H=	5,	K=	3			
			5	176	170				-1			
H=	4,	K=	-4	6	140	149	0	31	-35	H=	5,	K=
			7	63	63	1	110	111	0	210	214	
0	267	268	8	37	42	2	26	20	1	189	188	
1	271	278	-12	46	45	3	51	45	2	129	126	
2	227	234	-11	47	52	4	36	34	3	57	62	
3	132	127	-10	21	19	-5	95	94				
4	45	-46	-9	121	125	-8	64	61				

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
4	115	112	-5	218	221	-7	23	25	-1	210	215
5	119	122	-4	307	308	-6	146	153			
-10	167	169	-3	223	225	-5	70	67	H=	6, K=	-3
-9	129	134	-2	218	219	-4	17	1			
-8	127	129	-1	165	165	-3	23	-27	0	80	83
-7	163	164				-2	-11	6	1	67	65
-6	148	149	H=	5, K=	-1	-1	35	35	2	74	78
-5	95	97							3	97	95
-4	64	62	0	126	132	H=	6, K=	2	-9	32	29
-3	36	31	1	215	213				-8	62	47
-2	27	27	2	73	71	0	127	127	-7	70	71
-1	172	169	3	75	77	1	216	219	-6	88	91
			4	21	-30	-7	80	0	-5	40	43
H=	5, K=	-2	5	140	140	-6	120	117	-6	82	83
			6	104	102	-5	73	70	-3	130	130
0	141	139	-11	-12	5	-4	149	145	-2	176	176
1	136	138	-10	20	17	-3	127	123	-1	147	147
2	158	160	-9	25	25	-2	104	102			
3	83	81	-8	-11	15	-1	148	149	H=	6, K=	-2
4	71	72	-7	25	18						
5	29	-31	-6	91	92	H=	6, K=	3	0	126	130
6	82	83	-5	114	116				1	93	92
-11	45	48	-4	257	254	-5	84	85	2	31	29
-10	79	75	-3	139	140	-4	100	100	3	78	79
-9	83	84	-2	127	127	-3	142	137	-9	98	97
-8	68	87	-1	84	85	-2	94	95	-8	135	139
-7	58	61							-7	116	122
-6	30	23	H=	6, K=	0	H=	6, K=	-5	-6	61	61
-5	110	110							-5	49	55
-4	44	45	0	111	116	0	98	100	-4	200	203
-3	16	3	1	35	36	-6	44	46	-3	183	184
-2	43	38	2	105	106	-5	-11	8	-2	180	194
-1	155	157	3	95	94	-4	26	20	-1	201	204
			-9	64	63	-3	54	52			
H=	5, K=	-2	-8	80	82	-2	57	57	H=	6, K=	-1
			-7	151	147	-1	95	94			
0	246	246	-6	216	222	H=	6, K=	-4	0	94	92
1	270	265	-5	233	235				1	59	59
2	224	233	-4	53	-33				2	28	-32
3	202	197	-3	52	-55	0	185	186	3	17	-16
4	60	66	-2	20	19	1	139	139	-9	50	51
5	54	52	-1	44	43	2	115	117	-8	119	113
6	136	144				-8	120	125	-7	111	114
-11	71	75	H=	6, K=	1	-7	160	155	-6	80	17
-10	-12	25				-6	140	197	-5	65	66
-9	98	98	0	40	37	-5	101	104	-4	-11	-11
-8	111	110	1	29	30	-4	69	72	-3	45	-49
-7	134	134	2	86	92	-3	133	134	-2	15	2
-6	157	160	-8	-11	-9	-2	182	185			

Table B-3
Observed and Calculated Structure Factors for $\text{H}_2\text{dhpipy}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC
H= 0, K= 0			H= 20, K= 0			H= -2, K= 0			0	418	115
2 321 -223	0	32	25	4 343 -334	2 1071 -1111	1 126 -130					
4 105 -108	2	32	85	6 58 99	2 -18 -4						
6 292 -272			8 97 -98	3 -18 22							
8 247 -242	H= -22, K= 0		10 -21 -5	4 62 84							
10 37 -20		2 40 22	H= 1, K= 1	5 37 -50							
H= 2, K= 0			H= -18, K= 0			6 100 97					
0 334 342	H= -20, K= 0		0 401 -389	7 40 56							
2 247 -261	2 36 47		1 354 -325	8 -22 11							
4 210 220	4 38 -83		2 77 -84								
6 26 22	6 -23 42		3 125 -133	H= 13, K= 1							
8 96 -95			4 159 -157	0 88 -97							
10 73 -77	H= -18, K= 0		5 120 123	1 128 -137							
H= 4, K= 0		2 35 40	6 68 -76	2 -19 -10							
0 361 -397	4 45 -54		7 47 -49	3 -19 15							
2 7484 -2899	6 -22 1		8 29 17	4 39 41							
4 135 128	8 148 -147		9 -20 -18	5 239 -253							
6 26 35			10 -20 18	6 117 114							
8 169 -160			11 -21 -4	7 32 30							
10 89 100	2 30 -21			H= 15, K= 1							
	4 30 -40		H= 3, K= 1								
H= 6, K= 0	0 134 138		0 113 -105	0 35 -27							
	8 110 -104		1 419 -425	1 -20 37							
0 650 640			2 453 441	2 -20 -16							
2 531 -525	H= -14, K= 0		3 469 455	3 53 59							
4 131 120			4 165 170	4 -21 7							
6 -18 -37	2 265 266		5 206 207	5 -22 -18							
8 58 -55	4 -20 -3		6 113 115	H= 17, K= 1							
10 -22 14	6 65 66		7 163 163	0 113 114							
	8 -21 -23		8 -19 -14	1 133 131							
H= 8, K= 0	10 101 108		9 90 89	2 -21 23							
			10 61 -60	3 -22 21							
0 296 -799	H= -12, K= 0		H= 5, K= 1	4 -22 11							
2 62 -69			0 549 551	5 -22 -19							
4 903 907	2 234 221		1 969 -986								
6 60 52	4 194 192		2 499 493	H= 19, K= 1							
8 40 43	6 88 85		3 107 -106	0 -21 2							
	8 142 -121		4 233 228	1 -21 12							
H= 10, K= 0	10 78 84		5 97 90	2 -22 1							
			6 110 117	3 -22 -9							
0 54 41	H= -10, K= 0		7 47 -52	H= 21, K= 1							
2 324 -329	2 30 -32		8 -20 23	0 -22 -20							
4 239 229	4 585 -543		9 40 42								
6 46 -45	6 -16 -5		10 44 -27	H= -21, K= 1							
8 -21 20	8 49 -46										
H= 12, K= 0	10 42 48		H= 7, K= 1								
			0 72 75								
0 89 -89	H= -8, K= 0		1 65 55	1 -23 -17							
2 67 76			2 226 -229	2 -22 -22							
4 -26 23	2 404 382		3 54 102	3 -22 -4							
6 390 -412	4 523 -523		4 309 -300	4 -23 -28							
8 50 -56	6 317 -355		5 213 -217	5 -23 22							
	8 -26 -12		6 54 -55								
H= 14, K= 0	10 118 118		7 52 -60	H= -19, K= 1							
			8 46 -51	1 87 83							
0 -18 5	H= -6, K= 0		9 -21 -17	2 -21 -3							
2 -19 1				3 47 43							
4 222 223	2 421 439		H= 9, K= 1	4 42 52							
6 79 -87	4 267 273		0 111 -111	5 -22 -11							
	6 415 395		1 23 30	6 -22 -12							
H= 16, K= 0	8 42 -35		2 183 -174	7 68 -70							
	10 64 41		3 62 61								
0 156 -152			4 244 -250	H= -17, K= 1							
2 -21 -7	H= -4, K= 0		5 50 -24	1 111 111							
4 -21 5	2 196 184		6 76 -76	2 64 59							
H= 18, K= 0	4 587 -581		7 72 -72	3 47 50							
	6 523 524		8 32 -40								
0 167 -163	8 103 97		9 40 35	4 65 62							
2 51 -45	10 35 -24		H= 11, K= 1	5 38 28							
4 -22 -6			6 31 31	6 31 39							

L	F0	FC	I	F0	FC	L	F0	FC	L	F0	FC
7	-22	-34	6	133	-123	H=	C, K=	2	0	-22	3
8	-23	29	7	31	-26				1	-37	-31
9	39	-34	8	99	-93	0	181	183	2	-23	16
			9	32	9	1	-15	-1			
H= -15, K=	1		10	49	-47	2	69	-66	H= -20, K=	2	
			11	52	-45	3	61	-56			
1	-19	-17				4	65	-59	1	49	45
2	-19	20	H=	-34, K=	1	5	152	-144	2	46	45
3	149	-140				6	64	-59	3	-22	-18
4	40	-29	1	125	141	7	-20	-18	4	37	-11
5	52	-58	2	202	-206	8	-70	24	5	73	-70
6	56	-60	3	181	-161	9	-21	21	6	-23	20
7	33	32	4	45	-51						
8	-21	12	5	100	-119	H=	E, K=	2	H= -16, K=	2	
9	36	52	6	80	-71						
			7	-18	-16	0	163	164	1	34	-42
H= -13, K=	1		8	-19	-5	1	167	-181	2	-21	8
			9	87	80	2	-16	2	3	62	52
1	295	284	10	-20	0	3	104	-117	4	31	-36
2	216	-214	11	-22	2	4	-17	3	9	93	87
3	232	-236				5	58	63	6	46	-55
4	55	-58	H=	-14, K=	1	6	53	55	7	-22	7
5	-20	-4				7	-21	37	8	-23	-3
6	23	-90	1	397	-422	8	-20	9			
7	34	-12	2	413	-414	9	-22	30	H= -16, K=	2	
8	-21	-22	3	115	111	H=	10, K=	2	1	119	-108
9	58	71	4	152	-149				2	-20	-24
10	42	-41	5	348	-345	0	114	112	3	69	-74
			6	51	60	1	79	86	4	110	-107
H= -11, K=	1		7	284	-283	2	-17	-9	5	37	19
			8	145	145	3	29	-20	6	40	26
1	126	133	9	-21	32	4	100	108	7	112	110
2	27	27	10	46	47	5	20	14	8	-74	-14
3	91	-87	11	-22	16	6	-21	19	9	39	38
4	43	-35				7	-21	13			
5	71	68	H=	0, K=	2	8	21	30	H= -14, K=	2	
6	79	88									
7	42	40	0	431	-401	H=	12, K=	2	1	53	-61
8	50	51	1	510	498				2	52	52
9	-21	3	2	93	94	0	140	133	3	69	66
10	-21	0	3	322	322	1	-18	8	4	101	-102
11	-22	-10	4	282	291	2	128	-129	5	70	-72
H= -9, K=	1		5	149	147	3	81	80	6	-21	26
			6	63	-59	4	-20	-24	7	80	-80
1	334	-330	7	-18	-22	5	96	100	8	91	93
2	129	122	8	51	-43	6	29	2	9	32	19
3	306	-301	9	104	-105	7	-21	-26	10	-23	9
4	119	107	10	38	-38						
5	167	176	11	43	-43	H=	14, K=	2	H= -12, K=	2	
6	38	37	H=	2, K=	2	0	65	-62	1	66	58
7	71	-71				1	55	56	2	175	176
8	73	74	0	178	153	2	58	-56	3	154	155
9	-21	23	1	720	-716	3	-20	10	4	-18	12
10	-21	26	2	313	-311	4	66	-56	5	103	100
11	-23	-43	3	226	231	5	49	46	6	108	109
H= -7, K=	1		4	33	-49	6	47	-39	7	-20	-2
			5	117	133	7	-21	-26	8	-21	-26
1	81	-54	6	80	-81	0	65	-62	1	66	58
2	86	-82	7	50	-55	H=	16, K=	2	2	175	176
3	96	101	8	-20	-6				3	154	155
4	162	154	9	-20	-14	0	40	-51	4	-18	12
5	275	268	10	-20	2	1	-20	-24	H= -10, K=	2	
6	-17	12	H=	4, K=	2	2	30	-23	1	160	151
7	88	-89				3	30	-16	2	159	162
8	-20	-34	0	127	126	4	84	46	3	73	70
9	51	-40	1	404	394	5	52	-47	4	39	-43
10	30	-31	2	128	131	H=	18, K=	2	5	224	-229
11	-22	-24	3	157	-162				6	120	114
H= -5, K=	1		4	186	-181	0	-21	-24	7	71	75
			5	45	-56	1	-21	-3	8	91	-95
1	332	-332	6	85	-86	2	31	31	9	50	56
2	315	314	7	64	-58	3	-22	-5	10	83	-84
3	264	250	8	74	76	4	-23	16	11	-23	8
4	64	-40	9	34	-27						
5	323	314	10	-21	24	H=	20, K=	2			

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC
			5	37	39				9	-21	-17
H = -6, K = 2	-6	2	6	-18	-5	H = 17, K = 3	18	15	10	-22	19
1	173	174	8	-20	-4	0	-21	15	H = -9, K = 3	-9	3
2	313	-313	9	-21	22	1	-21	-10	43	33	
3	141	138	10	-21	1	2	-21	3	77	-76	
4	121	-106				3	46	-43	62	-68	
5	77	-77	H = 5, K = 3			4	-22	-18	136	131	
6	69	67							-18	-1	
7	117	-122	0	107	-102	H = 19, K = 3	19	5	96	96	
8	103	-105	1	150	131	0	-22	-9	141	144	
9	-21	-5	2	25	25	1	37	41	98	101	
10	-21	-21	3	327	325	2	53	-44	-21	8	
11	38	46	4	135	145	5	122	114	36	29	
H = -6, K = 2	-6	2	6	47	40	H = -21, K = 3			H = -7, K = 3		
1	243	237	7	-20	14						
2	408	-394	8	31	-36	1	53	-51	235	230	
3	360	-342	9	-21	6	2	57	-59	175	180	
4	207	-204	10	35	25	3	-23	28	41	-47	
5	27	36	H = 7, K = 3			4	-22	1	-16	1	
6	90	90							94	96	
7	111	114	0	279	-275	H = -19, K = 3			102	105	
8	62	-59	1	88	-93	1	35	-25	84	91	
9	102	-98	2	54	-56	2	61	-76	221	225	
10	66	68	3	246	248	3	50	-62	77	78	
11	46	-42	4	97	92	4	-22	8	-21	9	
			5	-18	-1	5	-20	-14	11	-22	3
H = -4, K = 2	-4	2	6	-19	5	6	67	61			
1	710	-709	7	-20	-2	7	-23	-16	H = -5, K = 3		
2	229	-222	8	-20	-16						
3	467	-464	9	-21	27	H = -17, K = 3			133	127	
4	122	127	H = 9, K = 3			1	42	41	2	-15	-22
5	163	-166				2	-21	29	220	-212	
6	-17	-9	0	117	-122	3	69	64	153	-156	
7	186	187	1	55	57	4	74	75	92	-87	
8	-20	2	2	-17	26	5	32	-38	50	-49	
9	108	99	3	39	38	6	36	40	-20	5	
10	95	95	4	-18	11	7	-22	-17	125	-121	
11	-22	2	5	74	-73	8	-23	21	75	-66	
			6	57	-61				22	18	
H = -2, K = 2	-2	2	7	51	-52	H = -15, K = 3			H = -3, K = 3		
1	216	209	8	45	-47						
2	78	68	H = 11, K = 3			1	45	58	67	-84	
3	93	-87				2	-20	-7	557	559	
4	370	370	0	193	-199	3	43	50	94	93	
5	251	239	1	28	24	4	121	129	157	-153	
6	35	36	2	103	98	5	36	39	130	135	
7	-18	-10	3	59	-57	6	-21	-3	59	-66	
8	65	-68	4	32	37	7	-22	-10	-19	-26	
9	-21	8	5	107	-114	8	38	-41	84	-82	
10	54	52	6	-20	15	9	62	47	81	-78	
11	44	47	7	-21	-3	H = -13, K = 3			119	-116	
			8	-22	-2				54	-48	
H = 1, K = 3	1	3	H = 13, K = 3			1	46	41			
0	150	-134				2	-18	16	H = -1, K = 3		
1	774	-775	0	76	75	3	-19	9			
2	640	-641	1	89	83	4	52	-58	114	123	
3	264	-278	2	43	41	5	57	-57	187	188	
4	130	-131	3	-20	-8	6	110	-114	111	-113	
5	140	-138	4	-21	-14	7	-21	-2	58	72	
6	182	187	5	33	-27	8	64	-64	242	235	
7	-18	-26	6	-21	4	9	-22	25	66	65	
8	-19	7	7	-22	20	10	-22	-1	26	-9	
9	-20	1							43	-35	
10	-20	7	H = 15, K = 3						21	3	
H = 3, K = 3	3	3	0	114	113	1	21	-24	10	-21	-6
1	-20	9	1	-20	9	2	112	-111	11	61	54
0	678	675	2	74	77	3	80	-83	H = 0, K = 4		
1	463	-475	3	-21	23	4	-18	1	343	-351	
2	100	-169	4	55	-53	5	89	-90	277	279	
3	112	114	5	-22	19	6	233	-237	253	-251	
4	133	-123	6	-22	-19	7	53	-43	70	-62	

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC				
4	407	404	4	-21	10	2	-19	10	1	112	-106				
5	87	-94	5	-20	15	2	30	-40	2	105	157				
6	35	36	6	-21	-8	4	132	-129	3	118	110				
7	95	91	7	62	-53	5	130	142	4	52	51				
8	-19	-20	H=	14	K=	4	6	282	294	5	208	-206			
9	-20	21				7	225	236	6	212	-209				
10	97	95	0	102	91	8	71	69	7	101	-94				
						9	-21	-2	8	-20	-4				
						10	43	-30	9	66	60				
									10	-21	-1				
H=	2,	K=	4	1	61	-61									
	2	111	-107				H=	-10,	K=	4	H=	3,	K=	5	
0	361	355	3	63	-61										
1	332	320	4	-21	-7	1	266	-263	0	50	58				
2	156	-146	5	-21	-2	2	153	-154	1	79	-86				
3	121	-117	6	-22	-10	3	75	-68	2	243	243				
4	105	102	H=	16,	K=	4	4	27	10	3	100	98			
5	112	112				5	80	-84	4	260	257				
6	38	-30	0	-21	-3	6	-20	20	5	225	-226				
7	79	72	1	-21	37	7	62	65	6	65	68				
8	-20	-5	2	-21	-21	8	221	-219	7	-19	14				
9	-20	21	3	-21	-17	9	-21	28	8	-20	16				
10	46	49	4	68	71	10	-22	20	9	-21	1				
H=	4,	K=	4	H=	18,	K=	4	H=	-8,	K=	4	H=	5,	K=	5
0	119	112	0	-21	11	1	37	-36	0	56	47				
1	429	428	1	-21	-4	2	107	109	1	66	-63				
2	58	67	2	40	-43	3	63	-66	2	-16	6				
3	140	-140	3	-23	20	4	-17	-26	3	167	167				
4	76	76	H=	20,	K=	4	5	62	59	4	72	-74			
5	34	-16				6	165	171	5	89	-89				
6	233	-240	H=	20,	K=	4	7	65	-77	6	39	29			
7	31	-23	0	-23	-11	8	275	-276	7	98	97				
8	51	-48	1	-23	-11	9	117	-126	8	58	57				
9	38	-32	H=	-20,	K=	4.	10	48	-49	9	33	44			
H=	6,	K=	4	1	46	23	H=	-6,	K=	4	H=	7,	K=	5	
0	165	-170	2	111	110	1	421	412	0	123	125				
1	79	-81	3	162	161	2	173	-109	1	40	-38				
2	102	-103	4	-23	0	3	253	245	2	46	-40				
3	81	-88	5	42	36	4	-16	10	3	53	56				
4	117	121	H=	-16,	K=	4	5	82	-90	4	41	-42			
5	57	56				6	-18	-3	5	56	-61				
6	53	-43	1	33	-40	7	40	45	6	71	-64				
7	124	-129	2	-22	18	8	56	-59	7	97	98				
8	-21	20	3	-22	17	9	-22	-28	8	-21	-21				
9	48	-51	4	240	-240	10	141	134							
H=	8,	K=	4	5	-22	11	H=	-4,	K=	4	H=	9,	K=	5	
0	540	536	6	46	43	H=	-4,	K=	4	H=	93	44			
1	254	-253	7	46	54	1	202	-201	0	53	-235				
2	-16	-5	H=	-16,	K=	4	2	661	-665	1	233	-169			
3	122	-128	3	-21	22	3	275	277	2	169	-165				
4	-18	10	4	33	39	4	36	29	3	-19	0				
5	134	141	5	68	-68	5	31	-27	4	47	-47				
6	58	-53	6	203	-209	6	55	58	5	71	-72				
7	30	25	7	266	-261	7	93	-92	6	-21	15				
8	94	99	8	256	-256	8	125	-126	7	60	62				
			9	63	-58	9	-21	25	8	-22	-11				
H=	10,	K=	4	7	39	-38	H=	-2,	K=	4	H=	11,	K=	5	
0	145	146	8	-23	12	H=	-2,	K=	4	0	160	165			
1	181	174	H=	-14,	K=	4	1	112	-113	1	157	-155			
2	-15	2	1	32	37	2	-15	38	2	31	-27				
3	-19	-1	2	-19	7	3	254	-241	3	-20	-13				
4	-19	1	3	110	111	4	62	63	4	-20	-36				
5	48	33	5	-20	-2	5	44	-42	5	32	-6				
6	64	-69	6	-21	-17	6	-18	0	6	31	20				
7	-21	-3	7	294	304	7	42	47	7	-21	20				
8	36	30	8	42	-49	8	62	-59							
H=	12,	K=	4	9	52	-57	H=	1,	K=	5	H=	13,	K=	5	
0	27	-16	8	44	-47	9	36	41	0	-20	28				
1	34	31	H=	-12,	K=	4	H=	1,	K=	5	1	87	-86		
2	171	-173	1	36	27	0	149	-138	2	79	78				
3	54	55							3	87	91				

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC			
4	114	115	H=	-9,	K=	5	H=	2,	K=	6	H=	16,	K=	6
5	-21	1		1	81	-71	0	206	195					
6	-21	0		2	65	-66	1	126	144	0	-21	8		
			H=	15,	K=	5		2	73	-68	1	44	-49	
				3	67	-66	3	152	148	2	-22	12		
				4	93	-102	4	26	-10	3	-22	-9		
0	37	-44		5	168	172				H=	18,	K=	6	
1	-21	-5		6	65	76	5	32	29					
2	114	-107		7	219	-227	6	-19	32					
3	64	83		8	86	90	7	33	-34	0	39	-43		
4	-22	19		9	66	71	8	39	-29	1	34	-40		
5	-22	1		10	90	85	9	-21	-15					
			H=	17,	K=	5	H=	-7,	K=	H=	-18,	K=	6	
				1	306	-296	0	197	-191	1	73	76		
0	-21	0		2	179	-169	1	217	-221	2	111	-110		
1	-22	-21		3	109	-107	2	199	-192	3	83	81		
2	-21	-25		4	41	-34	3	82	76	4	-22	0		
3	43	52		5	-19	32	4	122	-115	5	47	-32		
			H=	19,	K=	5		6	115	111				
				7	-21	-37	6	-19	-4	H=	-16,	K=	6	
0	-23	30		8	223	-220	7	-20	8		1	44	-46	
1	-22	-7		9	179	180	8	54	-42	2	-21	-3		
				10	-22	0	9	-22	30	3	58	58		
H=	-19,	K=	5				H=	-5,	K=	H=	6,	K=	6	
1	37	-21					H=	6,	K=	6	4	45	27	
2	131	119		1	365	-354	0	165	-151	5	70	73		
3	183	-179		2	31	22	1	62	-56	6	47	-49		
4	-22	9		3	122	119	2	138	-139	7	80	-78		
5	-23	28		4	245	242	3	104	-102	H=	-14,	K=	6	
				5	34	-37	4	47	46					
H=	-17,	K=	5					5	85	-82	1	85	-89	
				6	143	138	6	52	45	2	56	55		
				7	171	-169	7	-20	7	3	146	-150		
1	55	56		8	75	-62	8	-21	0	4	179	176		
2	57	59		9	173	171				5	75	-85		
3	149	-145		10	84	-83	H=	8,	K=	6	6	-21	22	
4	101	105								7	59	51		
5	89	92	H=	-3,	K=	5		0	31	24	8	-22	12	
6	72	68						1	237	231				
7	-23	-9		2	220	-215	2	93	93	H=	-12,	K=	6	
				3	321	320	3	32	31					
H=	-15,	K=	5					4	94	94	1	118	-116	
				4	317	318	5	85	-82	2	55	-67		
1	-21	11		5	150	147	5	86	-87	3	-20	18		
2	-21	6		6	86	87	6	81	78	4	34	-14		
3	40	39		7	62	-63	7	68	-64	5	154	-147		
4	135	-135		8	-21	24				6	-21	-27		
5	263	263		9	54	50	H=	10,	K=	6	7	45	-44	
6	-22	-16		10	107	104		0	70	-59	8	61	55	
7	-22	-18						1	-19	16	9	99	95	
8	38	-27	H=	-1,	K=	5		2	54	61				
								3	91	84	H=	-10,	K=	6
H=	-13,	K=	5					4	-20	23				
1	51	-56		2	310	-293	4	-21	23					
2	30	30		3	345	327	5	61	62	1	138	133		
3	107	-111		4	261	-259	6	-21	-25	2	-19	42		
4	89	-86		5	52	-47	7	-22	26	3	130	132		
5	212	217		6	-20	33	H=	12,	K=	6	4	48	-60	
6	62	-64		7	76	-74				5	176	174		
7	87	-83		8	111	114	0	50	-42	6	196	-205		
8	82	-80		9	63	49	1	-20	5	7	57	59		
9	-22	-14						2	-21	22	8	-21	6	
			H=	0,	K=	6		3	66	-72	9	-22	-8	
H=	-11,	K=	5					4	-21	-34	H=	-8,	K=	6
1	-18	7		0	263	260	5	-21	-2	1	50	56		
2	-19	-25		1	153	-151	6	41	-28	2	95	66		
3	49	-53		2	197	196				3	152	159		
4	76	-81		3	110	-105	H=	14,	K=	6	4	35	27	
5	-20	-11		4	54	-45				5	-20	-18		
6	187	191		5	60	62	0	-21	-18	6	44	-35		
7	253	-252		6	-19	17	1	42	25	7	106	108		
8	57	49		7	42	-36	2	54	50	8	-21	7		
9	-22	-4		8	35	28	3	37	-35	9	-22	-11		
				9	-21	-2	4	-22	-24					
							5	34	-35					

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC
H= -6, K= -6			1 63	58	5 55	64	0 35	28			
			2 79	77	6 -21	-20	1 366	369			
			3 45	45	7 83	-63	2 466	-62			
1 129	-136		4 -21	6	8 67	-68	3 -19	18			
2 119	117		5 -20	14			4 -81	-89			
3 104	-100		6 61	-56	H= -9, K= 7		5 -20	-1			
4 40	-43		7 -22	7			6 -20	0			
5 33	-27				1 49	-64	7 41	36			
6 39	50		H= 9, K= 7		2 -19	21	8 -21	-3			
7 118	-120				3 36	-54					
8 144	143		0 -19	-36	4 111	124	H= 6, K= 8				
9 73	-76		1 -19	6	5 -21	-25	0 125	129			
			2 72	-71	6 113	104	1 52	62			
H= -4, K= -6			3 -20	9	7 51	-86	2 53	-50			
			4 -20	-4	8 33	-23	3 -20	-16			
1 309	320		5 33	-25	H= -7, K= 7		4 -20	-7			
2 -16	-31		6 -21	-7			5 45	-34			
3 146	-142		H= 11, K= 7		1 227	-275	6 46	52			
4 -18	-8				2 42	39	7 -21	-19			
5 42	-36		0 50	-51	3 59	58	H= 6, K= 8				
6 -19	5		1 97	-91	4 54	-40	0 39	31			
7 34	29		2 57	-55	5 73	82	1 48	-51			
8 46	48		3 -20	16	6 63	-64	2 70	-72			
9 71	-67		4 60	-56	7 52	-52	3 21	-22			
10 -22	-18		5 40	-29	8 -21	24	4 -21	-12			
H= -2, K= -6					9 45	40	5 -20	-8			
H= 13, K= 7			H= -5, K= 7				6 -21	29			
1 94	-96						7 -22	-7			
2 71	51		0 -20	0	H= 8, K= 8						
3 -16	-7		1 72	-71	1 65	-63					
4 135	-133		2 -21	-10	2 40	57					
5 -18	-21		3 -21	-14	3 82	82					
6 63	58		4 32	20	4 48	-58					
7 -21	3				5 102	107					
8 -21	-20		H= 15, K= 7		6 110	-112	0 67	-69			
9 105	101		0 -21	12	7 61	66	1 -20	-27			
			1 -22	-9	8 65	-60	2 59	-73			
H= 1, K= 7			2 -21	24	9 84	78	3 71	-69			
0 24	0		3 -22	19	H= -3, K= 7		4 45	42			
1 30	13						5 30	4			
2 39	-37		H= 17, K= 7		1 80	85	H= 10, K= 8				
3 53	51				2 -17	4					
4 43	38		0 -22	17	3 101	95	0 -21	26			
5 42	-35				4 90	-100	1 -21	-29			
6 83	80		H= -17, K= 7		5 100	-100	2 -21	-20			
7 35	23				6 -19	-6	3 -21	12			
8 -21	1		1 53	-44	7 61	-60	4 -97	45			
9 -21	24		2 -23	37	8 -21	31	5 134	134			
H= 3, K= 7			3 -23	-34	9 49	45	H= 12, K= 8				
0 -17	-28		H= -15, K= 7		H= -1, K= 7						
1 65	-61				1 -17	-12	0 -20	11			
2 37	-48		1 48	51	2 145	-148	1 38	41			
3 52	-52		2 57	-61	3 87	85	2 -21	-16			
4 93	92		3 50	-51	4 -18	-8	3 -21	0			
5 -19	-17		4 -21	-5	5 118	-113	4 33	50			
6 37	28		5 63	53	6 -20	14	H= 14, K= 8				
7 46	41		6 77	66	7 67	-68	0 -21	-19			
8 -21	16				8 41	55	1 -21	-17			
H= 5, K= 7			4 -23	-22	9 44	-39	2 -22	-22			
			H= -13, K= 7								
0 186	196		1 -21	54	H= 0, K= 8						
1 71	-66		2 58	-56							
2 -18	17		3 -21	25	0 40	48	H= 16, K= 8				
3 -19	-6		4 100	-94	1 -18	20	0 34	-33			
4 -19	0		5 -21	29	2 60	-62	H= -16, K= 8				
5 -20	-4		6 -21	2	3 47	41					
6 -21	-37		7 40	-48	4 43	-44					
7 55	43		H= -11, K= 7		5 53	50	1 55	-50			
8 -22	-21		1 -20	-3	6 29	-11	2 -22	13			
H= 7, K= 7			2 38	33	7 -21	25	3 40	-19			
0 41	24		3 49	-59	H= 2, K= 8		H= -14, K= 8				
			4 101	108			1 -22	-33			

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC				
	2	45		-38		H=	-5,	K=	11	4	47	56			
H=	-5,	K=	11	H=	-9,	K=	11	1	-21	-22	H=	-1,	K=	11	
0	-21	-27				2	65	-67	1	-20	2				
1	-21	-32	1	37	-29	3	-21	30	2	-21	17				
2	-21	-13				4	-21	4	3	-21	11				
3	-36	-35	H=	-7,	K=	11	H=	-3,	K=	11	4	61	54		
H=	7,	K=	11	1	63	-52	1	-21	31	H=	0,	K=	12		
0	70	-72	2	51	-52	2	58	61	0	107	-106				
1	54	40	3	-22	-10	3	35	30							

Table B-4
Observed and Calculated Structure Factors for $[\text{Ni}_2\text{Cl}(\text{H}_2\text{O})_4 \cdot (\text{dhpipy})\text{Cl}_3 \cdot 2\text{H}_2\text{O}]^{4-}$

L	FD	FC	L	FD	FC	L	FD	FC	L	FD	FC
H=	0, K=	0	H=	6, K=	0	-4	412	-419	-23	194	-212
2	963	-490	2	3015	-2957	-2	607	-632	-27	-93	-106
4	1938	-2036	4	911	-905	0	1671	-1722	-26	264	-325
6	563	-593	6	363	-391	2	494	-506	-22	363	-384
8	135	-253	8	965	-1004	4	427	-434	-21	-96	-160
10	659	-607	10	317	-322	6	-67	-152	-20	227	-270
12	314	-323	12	-79	6	8	-84	-75	-19	326	-352
14	-64	103	14	445	-434	8	-84	-75	-18	-97	-29
16	130	-183	16	229	-230	-20	501	-541	-17	-17	-422
18	148	-113	18	133	-93	-20	241	-200	-16	-90	412
20	242	-261	20	-64	-106	-20	194	187	-15	-96	-59
22	177	-186	-30	602	-576	-22	132	-120	-12	-98	103
24	115	-142	-33	550	-567	-20	362	-399	-14	-96	-297
26	700	-691	-26	747	754	-18	177	-177	-13	563	595
			-24	349	-383	-16	357	357	-12	-98	103
H=	2, K=	0	-22	765	-760	-14	249	256	-11	493	-522
			-20	163	165	-12	161	-180	-10	190	-247
0	483	373	-18	276	315	-10	-79	38	-9	847	-507
2	2936	-2814	-16	929	-921	-8	-79	61	-8	445	512
4	137	-124	-14	157	197	-6	-80	-14	-7	201	-232
6	345	-303	-12	197	119	-4	374	362	-6	793	-703
8	1693	1682	-10	147	-99	-2	1073	1111	-5	1418	1462
10	124	-119	-8	1553	1483				-4	827	-696
12	115	107	-6	973	930	H=	14, K=	0	-3	393	249
14	645	649	-4	816	741				-2	867	723
16	555	-547	-2	2496	-2505	0	586	-594	-1	124	51
18	255	251				2	433	436	H=	3, K=	1
20	348	362	H=	8, K=	0	-24	-83	-16			
22	342	-354				-22	346	-328			
24	413	-402	0	3957	4100	-20	-81	65	0	758	-639
26	311	292	2	411	421	-18	284	277	1	348	317
-28	519	-541	4	429	-429	-16	306	-343	2	-70	-130
-26	448	-459	6	252	278	-14	171	-167	3	381	-331
-24	206	181	8	-70	-83	-12	-31	-111	4	390	-403
-22	426	443	10	-81	-123	-10	197	-229	5	559	559
-20	144	-109	12	199	-199	-8	141	136	6	-67	19
-18	269	-274	14	-79	57	-6	394	399	7	-67	-43
-16	860	866	16	-81	-11	-4	371	360	8	472	-485
-14	568	-551	30	472	474	-2	378	-411	9	447	-461
-12	283	-322	-28	541	520				10	-66	-92
-10	273	332	-26	341	-352	H=	16, K=	0	11	500	-513
-8	2573	-2519	-24	-78	18				12	-69	59
-6	1388	-1666	-22	-78	-24	-16	322	-311	13	621	605
-4	329	-390	-20	-77	92	-14	293	-207	14	195	233
-2	3358	3505	-18	139	148	-12	192	202	15	301	279
			-16	446	-446	-10	-84	5	16	475	-473
H=	4, K=	0	-14	315	-276	H=	1, K=	1	17	388	-382
			-12	126	109	H=	1, K=	1	18	-81	51
0	7329	-8496	-10	277	-252				19	426	-456
2	242	-331	-8	138	100	0	1935	2008	20	411	393
4	1140	1162	-6	-55	48	1	1546	-1410	21	-82	-59
6	305	-285	-4	1229	-1237	2	1354	-1915	22	450	-430
8	351	-306	-2	1643	-1663	3	617	-569	23	273	243
10	99	120				4	1209	-1109	24	-84	44
12	425	433	H=	10, K=	0	5	712	-730	-29	-103	-191
14	199	-227				6	347	323	-28	-100	-233
16	-80	36	0	655	660	7	338	-273	-27	764	291
18	-80	27	2	942	-955	8	187	-142	-26	136	-171
20	271	-285	4	424	-433	9	172	193	-25	200	247
22	195	-153	6	-30	95	10	256	-241	-24	569	577
-30	469	-434	8	529	527	11	700	-693	-23	400	-403
-28	396	-387	10	257	-260	12	1135	1242	-22	316	327
-26	574	573	12	140	-155	13	-70	46	-21	233	-240
-24	218	209	-30	514	502	14	430	457	-20	663	-674
-22	216	-217	-23	226	-223	15	733	747	-19	-74	-161
-20	467	-417	-25	559	-552	16	-76	145	-18	571	573
-18	192	162	-24	113	120	17	310	335	-17	392	392
-16	674	657	-22	668	675	18	353	-364	-16	352	353
-14	134	126	-20	131	-133	19	-82	-57	-15	1334	1345
-12	312	246	-18	642	-619	20	501	505	-14	663	-631
-10	565	512	-16	759	758	21	251	-253	-13	105	-103
-8	659	588	-14	343	349	22	-82	-47	-12	1674	-1664
-6	219	-147	-12	-72	57	23	336	-356	-11	1773	-1757
-4	427	380	-10	319	325	24	670	-656	-10	733	640
-2	1524	1519	-8	576	-550	25	-91	134	-9	-67	33
			-6	519	-554	26	-95	-23	-3	143	-144

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
-7	281	230	12	117	38	-9	735	-721	-11	-81	-21
-6	753	-765	13	123	-155	-8	144	157	-10	-80	-32
-5	697	-574	14	175	-151	-7	517	-504	-9	201	245
-4	1044	1036	15	-80	-67	-6	206	-191	-8	-92	-103
-3	459	-483	16	149	153	-5	428	431	-7	299	332
-2	1737	1695	17	-34	150	-4	385	-385	-6	-80	-10
-1	1005	-945	18	126	70	-3	203	210	-5	257	-240
H=	5, K=	1	19	201	254	-2	443	-455	-4	319	322
			-30	-84	33	-1	-73	-39	-3	225	-215
			-29	200	223				-2	357	369
0	1566	-1596	-28	480	473	H=	11, K=	1	-1	-84	17
1	942	945	-27	312	-317						
2	959	923	-26	295	263	0	-82	34	H=	15, K=	1
3	465	439	-25	333	-339	1	-82	58			
4	1415	1442	-24	477	-464	2	391	-378	-22	158	-118
5	350	372	-23	393	407	3	-80	-44	-21	126	162
6	340	-347	-22	514	-331	4	-81	-4	-20	277	273
7	390	-379	-21	246	208	5	273	270	-19	-85	55
8	335	-338	-20	431	477	6	221	226	-18	-64	55
9	356	-360	-19	-73	22	7	-82	77	-17	-34	9
10	570	553	-18	450	-455	8	234	-199	-16	297	-277
11	645	640	-17	-70	-15	9	171	-137	-15	415	-415
12	452	-450	-16	337	-328	10	236	-214	-14	-35	-113
13	176	218	-15	632	-629	11	228	-237	-13	208	-219
14	365	-342	-14	212	209	-29	242	-240	-12	300	302
15	466	-452	-13	292	-278	-28	320	-307	-11	416	414
16	230	-205	-12	1029	1043	-27	205	215	-10	-65	-95
17	204	-229	-11	1070	1055	-26	357	-349	-9	168	174
18	-81	124	-10	395	-403	-25	307	300	-8	-85	-31
19	-81	-11	-9	-70	70	-24	300	305	-7	181	-173
20	276	-275	-8	-70	17	-23	297	-295	-6	252	253
21	-64	79	-7	485	-480	-22	298	283	-5	157	-123
22	-84	3	-6	997	903	-21	316	-302	-4	-86	-14
-30	501	-545	-5	-70	-35	-20	632	-630	-3	-87	47
-29	246	240	-4	886	-880	-19	155	-143	H=	0, K=	2
-28	-81	-14	-3	574	587	-18	117	120			
-27	-82	192	-2	2150	-2151	-17	-80	89			
-26	421	419	-1	1041	1025	-16	524	541	0	321	-349
-25	219	-236				-15	753	770	1	90	250
-24	319	293	H=	9, K=	1	-14	-78	13	2	761	-703
-23	582	-593				-13	249	275	3	238	-327
-22	667	-681	0	700	712	-12	579	-554	4	402	425
-21	-77	4	1	348	-347	-11	594	-591	5	614	-490
-20	363	369	2	187	-153	-10	143	151	0	143	57
-19	551	583	3	-77	-53	-9	-77	-105	7	875	821
-18	519	483	4	443	-447	-8	-76	-23	8	550	537
-17	690	694	5	213	-245	-7	123	89	9	650	652
-16	506	-477	6	-80	-27	-6	297	-287	10	376	-376
-15	123	55	7	-80	43	-5	-73	-39	11	-71	-8
-14	-64	55	8	119	83	-4	290	292	12	715	-742
-13	702	-622	9	129	150	-3	141	-144	13	1073	-1095
-12	566	-550	10	272	-286	-2	895	904	14	975	981
-11	328	285	11	243	-255	-1	407	-427	15	-73	-38
-10	443	393	12	-81	41				16	409	426
-9	140	148	13	221	-192	H=	10, K=	1	17	-26	-31
-8	425	408	14	162	146	-12	611	-641	16	432	-457
-7	319	275	15	295	232	0	228	254	19	-94	-50
-6	345	297	-30	497	471	1	-53	-16	20	264	-311
-5	475	-440	-29	190	-110	2	-64	71	21	235	-230
-4	109	110	-28	154	177	3	-64	71	22	-35	66
-3	-68	-70	-27	101	-110	4	379	364	23	240	-274
-2	505	448	-25	156	-129	5	149	141	24	276	-359
-1	-68	96	-25	-80	89	-27	124	106	25	323	312
H=	7, K=	1	-24	311	-305	-25	187	175	26	267	-265
			-23	441	428	-64	-35	27	-85	65	
			-22	355	349	-24	296	259	H=	2, K=	2
0	473	456	-21	-78	11	-23	320	-340			
1	-67	-62	-20	127	-109	-22	303	-293			
2	812	843	-19	373	-347	-21	-79	-47	0	94	59
3	-68	65	-18	474	-480	-20	-79	-5	1	183	-132
4	-69	-6	-17	545	-555	-19	213	179	2	195	123
5	428	-443	-16	254	255	-16	303	318	3	94	12
6	-72	-81	-15	-72	-55	-17	267	248	4	494	510
7	-73	-55	-14	352	321	-16	155	-112	5	547	589
8	360	371	-13	930	887	-15	115	110	6	1059	-1043
9	-77	105	-12	415	427	-14	140	-124	7	500	513
10	-78	67	-11	-59	-97	-13	303	-320	8	-67	-23
11	219	222	-10	270	-265	-12	359	-355	9	375	860

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC
10	366	-339	-24	541	520	117	6+	K= 2	-20	118	-136
11	756	-760	-23	143	-193				-19	470	-463
12	1086	1099	-22	-79	-56				-18	-76	14
13	-71	-62	-21	202	-269	0	337	-342	-17	549	-553
14	360	401	-20	429	433	1	131	253	-16	936	932
15	1101	1109	-19	-75	-71	2	214	-243	-15	744	703
16	410	-423	-18	1108	1065	3	242	262	-14	676	581
17	146	-143	-17	362	375	4	152	163	-13	465	450
18	263	-313	-16	473	-477	5	145	-140	-12	1029	-1004
19	749	-756	-15	-276	-283	6	113	116	-11	924	-913
20	-69	-28	-14	1643	-1630	7	260	282	-10	151	-135
21	-80	-43	-13	1572	-1672	8	150	113	-9	179	-192
22	173	158	-12	737	721	9	260	275	-8	284	255
23	-61	-31	-11	397	-375	10	191	-142	-7	178	166
24	159	-154	-10	614	610	11	170	155	-6	206	214
25	-66	35	-9	604	794	12	497	-495	-5	290	279
-29	-89	-37	-8	293	-263	13	455	-477	-4	433	-432
-28	158	-147	-7	550	602	14	218	293	-3	-75	137
-27	-54	32	-6	886	893	15	-84	-95	-2	-76	-131
-26	305	-321	-5	-69	-37	16	355	343	-1	-77	-94
-25	-64	-3	-4	418	-397	17	170	179	H= 12+, K= 2		
-24	-61	28	-3	-70	-82	-20	127	-126	0	231	237
-23	-62	-142	-2	371	364	-29	153	-155	-1	-81	-60
-22	-81	-59	-1	-69	-164	-28	333	322	-2	-82	67
-21	-82	31				-27	-79	-50	-3	140	-142
-20	-115	200	H=	6+, K= 2		-26	-79	128	-2	-82	67
-19	717	-717				-25	472	-473	-3	140	-142
-18	-82	143	0	542	-523	-24	387	-410	-4	-82	23
-17	499	-541	1	352	355	-23	127	95	5	155	174
-16	655	660	2	238	245	-22	-79	-65	6	131	-71
-15	1196	1184	3	374	390	-21	-78	6	7	-85	-31
-14	-66	-120	4	496	-482	-20	251	-229	8	186	-186
-13	-72	111	5	360	-365	-19	-76	-19	-28	247	-260
-12	2047	-1794	6	253	259	-18	738	-732	-27	-69	5
-11	1667	-1751	7	401	-435	-17	-73	-27	-26	258	-270
-10	408	369	8	357	349	-16	-71	75	-25	355	351
-9	451	342	9	507	-501	-15	362	-343	-24	330	315
-8	797	730	10	594	593	-14	1003	1074	-23	151	-71
-7	1069	1154	11	372	402	-13	1003	1027	-22	-79	-27
-6	754	703	12	461	-460	-12	230	-222	-21	-50	-52
-5	1507	1321	13	121	68	-11	637	601	-20	-61	41
-4	1370	-1344	14	253	-251	-10	853	-343	-19	-81	-4
-3	120	1	15	573	-542	-9	831	-853	-18	603	604
-2	352	-380	16	-82	38	-8	132	140	-17	-83	52
-1	152	-211	17	-80	17	-7	1146	-1108	-16	149	97
			18	143	92	-6	144	105	-15	235	207
H=	4+, K= 2		19	565	340	-5	194	182	-14	320	-315
			20	127	101	-4	137	157	-13	390	-310
0	554	534	-30	404	-408	-3	437	436	-12	-31	-54
1	526	-547	-29	-55	78	-2	572	-549	-11	23	-284
2	266	299	-28	-30	12	-1	121	135	-10	122	115
3	129	-163	-27	-32	-32	H=	10+, K= 2		-9	245	227
4	601	-395	-26	420	433				-8	-80	-77
5	438	414	-25	-51	13				-7	457	466
6	290	-283	-24	-60	-30	0	343	319	-6	-80	66
7	605	-673	-23	-79	74	1	180	-189	-5	-80	46
8	555	-544	-22	131	57	2	-70	-52	-4	142	-148
9	933	-905	-21	-78	-70	3	232	-243	-3	171	-140
10	636	641	-20	-75	-14	4	213	212	-2	378	372
11	113	-107	-19	602	653	5	-7	74	-1	-83	-114
12	1036	1023	-18	120	-124	6	115	-120	H= 14+, K= 2		
13	1170	1154	-17	653	606	7	135	174			
14	828	-887	-16	746	-737	8	150	-181	0	241	-233
15	-60	100	-15	605	-600	9	246	252	1	151	65
16	635	-564	-14	561	-540	10	403	-501	2	-86	24
17	163	-229	-13	-56	3	11	291	-305	-23	-84	5
18	234	-227	-12	1039	907	12	105	179	-25	-84	93
19	-81	22	-11	936	952	13	152	-137	-24	-84	93
20	164	152	-10	115	84	-30	303	211	-23	-53	42
21	197	168	-9	315	-331	-29	151	-113	-22	-52	19
22	-85	-105	-8	-68	-63	-28	121	120	-21	121	-103
23	231	217	-7	341	-327	-27	-69	71	-20	-61	92
-30	-86	44	-6	422	-444	-26	202	-340	-19	-63	103
-29	-67	140	-5	703	-739	-25	-78	-18	-18	-63	64
-28	543	-318	-4	1155	1103	-24	-79	-89	-17	252	265
-27	-82	90	-3	107	-100	-23	-60	-42	-16	341	-339
-26	-81	92	-2	226	247	-22	-80	-3	-15	231	-231
-25	465	465	-1	377	365	-21	153	142	-14	417	-412

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC		
-13	303	-317	-2	1489	1483	16	483	-471	-10	436	-410		
-12	402	415	-1	-64	181	17	136	-157	-9	316	-282		
-11	406	410				18	436	418	-8	1335	-1311		
-10	132	121	H=	3,	K=	3	19	350	359	-7	928	-519	
-9	273	272				20	154	141	-6	523	557		
-8	325	-320	0	-68	-30	21	240	-235	-5	780	757		
-7	-81	-66	1	-68	8	-30	-84	69	-4	807	911		
-6	150	-171	2	1777	-1730	-29	137	-100	-3	531	515		
-5	303	-304	3	335	418	-28	278	-297	-2	222	219		
-4	378	347	4	264	-292	-27	209	-233	-1	583	-621		
-3	-85	-96	5	685	674	-26	232	-226					
-2	170	120	6	597	583	-25	319	333	H=	9,	K=	3	
-1	141	106	7	954	904	-24	560	550					
			8	1112	-1116	-23	279	272	0	245	-239		
H=	16,	K=	2	9	-66	-55	-22	177	-161	1	203	191	
			10	956	928	-21	810	-799	2	204	201		
-16	143	-150	11	538	-598	-20	595	572	3	-78	78		
-15	271	-282	12	1624	1034	-19	345	-349	4	275	297		
-14	307	358	13	-77	-5	-18	263	-247	5	278	-276		
-13	220	241	14	-76	-34	-17	345	352	6	212	219		
-12	197	199	15	256	253	-16	943	-946	7	125	-138		
-11	349	324	16	831	-651	-15	139	-160	8	225	-230		
			17	335	-339	-14	602	-558	9	280	310		
H=	1,	K=	3	18	494	-473	-13	165	163	10	441	-453	
			19	-82	-39	-12	913	900	11	-81	60		
0	243	-269	20	395	401	-11	747	599	12	-83	-27		
1	989	947	21	630	621	-10	1502	1509	13	233	-220		
2	620	684	22	189	-204	-9	491	523	14	233	257		
3	-69	25	23	-86	24	-8	115	-154	15	190	166		
4	940	853	24	317	289	-7	492	-473	-30	133	-117		
5	677	-686	25	-35	142	-6	1037	1060	-29	-83	54		
6	513	565	26	-86	-35	-5	644	-597	-28	295	293		
7	-67	-7	27	-98	-119	-4	342	315	-27	249	250		
8	918	-902	28	269	-303	-3	363	-375	-26	332	336		
9	557	521	29	-91	170	-2	874	-879	-25	183	-162		
10	331	-354	30	-38	227	-1	295	-280	-24	444	-461		
11	123	-189	31	253	239				-23	208	-193		
12	495	527	32	-87	-176	H=	7,	K=	3	-22	156	165	
13	356	-507	33	727	742				-21	468	478		
14	-91	351	34	-31	143	0	262	-287	-20	277	-289		
15	-95	249	35	555	-674	1	214	-182	-19	370	393		
16	547	590	36	-18	1435	1503	2	634	646	-18	-77	-47	
17	-115	87	37	-74	-119	3	236	238	-17	273	-275		
18	906	-986	38	312	-307	4	-70	57	-16	737	735		
19	-116	-593	39	-15	540	510	5	-73	-113	-15	-75	1	
20	-100	-238	40	-14	937	-923	6	207	-203	-14	910	903	
21	415	460	41	-13	903	-826	7	397	-403	-13	161	158	
22	-94	220	42	-12	1017	-1003	8	684	711	-12	775	-763	
23	210	274	43	-11	859	-871	9	-79	-57	-11	481	-481	
24	-90	-187	44	-10	1038	1072	10	287	-295	-10	1555	-1529	
25	-87	109	45	-9	802	808	11	221	197	-9	827	-817	
26	206	211	46	-8	1650	1549	12	760	-772	-8	439	439	
-28	140	143	47	-7	478	391	13	-83	68	-7	303	303	
-27	127	139	48	-6	853	-808	14	-80	-36	-6	-71	-74	
-26	-82	81	49	-5	1071	-1007	15	130	-87	-5	495	503	
-25	-87	-200	50	-4	506	-606	16	281	277	-4	337	-339	
-24	491	-488	51	-3	356	-366	17	133	110	-3	-73	31	
-23	-83	-73	52	-2	608	-526	18	361	333	-2	266	275	
-22	172	204	53	-1	-69	119	-30	285	-277	-1	-73	40	
-21	651	651					-29	269	-262				
-20	325	-355	H=	5,	K=	3	-28	-51	45	H=	11,	K=	3
-19	-78	143					-27	163	183				
-18	402	415	0	365	361	-26	303	337	0	205	206		
-17	214	-223	1	756	-770	-25	-80	-97	1	-81	49		
-16	557	602	2	378	-429	-24	144	-100	2	197	-215		
-15	174	160	3	137	-152	-23	221	-197	3	206	-214		
-14	271	296	4	737	-793	-22	240	226	4	-79	19		
-13	-71	-80	5	892	944	-21	810	-840	5	-80	-57		
-12	1523	-1529	6	779	-754	-20	-77	-76	6	-81	81		
-11	617	-621	7	113	131	-19	201	211	7	239	253		
-10	1373	-1803	8	710	693	-18	1237	-1192	8	510	-527		
-9	416	-402	9	743	-717	-17	-74	107	9	-35	106		
-8	1357	1283	10	915	899	-16	162	133	10	-35	-3		
-7	1544	1493	11	-77	65	-15	237	-240	-29	196	133		
-6	596	-503	12	258	-240	-14	-68	105	-28	-80	50		
-5	1175	1208	13	376	330	-13	515	435	-27	141	-161		
-4	339	-356	14	356	-490	-12	531	553	-26	261	-274		
-3	126	-119	15	298	-289	-11	716	630	-25	-80	29		

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
-24	-79	-8	-5	-85	31	-12	1394	-1362	8	501	463
-23	-80	133	-4	164	192	-11	-70	-33	9	612	-623
-22	296	-293	H= 0, K= 4			-10	146	-180	10	152	220
-21	766	781				-9	1076	1051	11	158	-191
-20	153	-80				-8	935	1008	12	437	-497
-19	-82	-100	0	350	316	-7	1053	1054	13	208	-169
-18	1002	1005	1	1374	1346	-6	1111	1073	14	169	-147
-17	202	-197	2	516	-539	-5	239	-216	15	-80	53
-16	117	150	3	1039	981	-4	410	-370	16	-81	3
-15	299	301	4	279	333	-3	1273	-1252	17	360	321
-14	171	-193	5	597	-531	-2	892	805	18	-33	24
-13	224	-226	6	-57	-82	-1	790	803	19	-80	47
-12	350	-359	7	1431	-1415				20	-35	-25
-11	445	-449	8	-69	-24	H= 4, K= 4			-30	142	-176
-10	-78	90	9	406	435				-29	518	-510
-9	-78	96	10	153	-120	0	235	-170	-73	-34	-29
-8	515	502	11	315	322	1	561	-516	-27	223	199
-7	299	269	12	226	-221	2	415	432	-26	-32	-2
-6	-79	-80	13	-66	10	3	700	-732	-25	607	643
-5	150	-152	14	1093	1098	4	212	-216	-24	-84	121
-4	190	-172	15	486	484	5	178	163	-23	182	160
-3	126	-111	16	343	365	6	190	-193	-22	-30	63
-2	-81	-113	17	249	-262	7	1319	1332	-21	630	-567
-1	216	203	18	191	-207	8	155	-190	-20	207	156
			19	1006	-1005	9	270	-260	-19	545	-551
H= 13, K= 3			20	260	-269	10	223	208	-18	-75	-77
			21	143	120	11	517	-519	-17	596	511
0	181	188	22	-81	112	12	452	447	-16	204	-168
1	198	-160	23	297	303	13	-78	29	-15	503	573
2	-85	-73	24	-65	44	14	833	-831	-14	214	-215
3	157	-172	25	160	202	15	451	-453	-13	-67	-62
4	152	-153	26	-90	-145	16	312	-325	-12	843	841
5	154	168	27	411	-439	17	-82	-23	-11	-66	70
-26	357	-353				18	163	153	-10	482	442
-25	-85	83	H= 2, K= 4			19	790	705	-9	213	-203
-24	286	293				20	162	133	-8	567	-515
-23	197	214	0	238	324	21	-82	21	-7	474	-461
-22	-82	-100	1	705	708	22	145	-99	-6	820	-651
-21	226	-213	2	563	-620	-29	-99	100	-6	-70	-17
-20	179	153	3	474	-463	-28	233	-377	-4	755	204
-19	290	-294	4	116	-51	-27	469	-457	-3	1244	1241
-18	143	116	5	434	428	-26	-24	48	-2	519	-491
-17	-80	44	6	1033	-1032	-25	-74	122	-1	657	-650
-16	226	-223	7	423	431	-24	-82	-73	H= 8, K= 4		
-15	-80	60	8	532	-569	-23	522	529	0	-72	-8
-14	478	-494	9	315	602	-22	222	-220	1	296	291
-13	-80	-68	10	124	-142	-21	398	403	2	169	-154
-12	118	118	11	210	216	-20	331	340	3	570	562
-11	202	185	12	677	858	-19	1317	-1400	4	221	232
-10	679	645	13	230	230	-18	521	511	5	157	131
-9	367	381	14	238	256	-17	503	-495	6	-80	121
-8	173	-178	15	257	-245	-16	396	-303	7	493	-490
-7	-81	-7	16	-31	-30	-15	704	679	8	-81	24
-6	133	-139	17	713	-713	-14	1312	-1319	9	-81	21
-5	203	-230	18	-85	-101	-13	-67	90	10	-79	-11
-4	134	162	19	-80	-19	-12	227	203	11	235	220
-3	-81	71	20	-64	99	-11	373	393	12	307	-298
-2	-81	63	21	514	550	-10	194	145			
-1	131	114	22	-67	6	-9	658	650	13	-61	10
			23	213	180	-8	277	-247			
H= 15, K= 3			24	-64	-7	-7	844	-846	14	323	317
			25	502	-284	-6	454	370	15	230	227
-21	377	-386	-26	-88	42	-5	169	-154	16	211	172
-20	-68	118	-27	226	-216	-4	209	-103	-22	126	-140
-19	-86	-42	-16	-34	2	-3	456	-433	-23	368	372
-18	454	-433	-25	338	-309	-2	366	370	-27	502	492
-17	151	127	-24	-30	-79	-1	120	-33	-26	-79	115
-16	-86	-131	-23	-77	-9				-25	-77	17
-15	168	-190	-22	-33	-3				-24	126	133
-14	-85	69	-21	405	462				-23	301	-224
-13	-83	9	-20	-81	-138	0	432	-420	-22	201	161
-12	329	322	-19	200	280	1	146	-1413	-21	528	-601
-11	290	282	-18	-47	19	2	400	405	-20	236	-264
-10	-65	-60	-17	844	-849	3	712	724	-19	734	736
-9	-63	9	-16	197	269	4	-63	-58	-18	452	-440
-8	457	-445	-15	517	-534	5	726	725	-17	414	-425
-7	261	-263	-14	-37	15	6	512	543	-16	-75	71
-6	-66	14	-13	204	230	7	130	-170	-15	452	-433

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC	
-14	961	943	-19	580	-583	-25	372	-361	-2	251	-218	
-15	203	-108	-18	307	300	-24	204	-207	-1	502	-473	
-12	-70	99	-17	350	-353	-23	457	-437				
-11	274	-270	-16	-82	-13	-22	-82	110	H=	K=	5	
-10	251	-239	-15	176	194	-21	164	165				
-9	716	-709	-14	450	-430	-20	-80	47	0	488	-470	
-8	131	122	-13	-81	102	-19	621	615	1	702	-676	
-7	560	562	-12	120	-76	-18	126	172	2	371	-374	
-6	-69	6	-11	-60	117	-17	215	-195	3	309	-306	
-5	654	655	-10	-81	-14	-16	-73	-73	4	614	-554	
-4	-59	19	-9	296	321	-15	522	-516	5	1032	1040	
-3	-69	-52	-8	143	-88	-14	107	-91	6	228	233	
-2	379	-380	-7	-61	-53	-13	284	282	7	359	361	
-1	549	-574	-6	-82	82	-12	514	526	8	411	-411	
			-5	183	-162	-11	423	425	9	770	-785	
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			-3	-81	0	-9	491	-480	11	259	-275	
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1	375	380	-1	413	410	-7	2513	2452	13	305	-294	
2	160	-161				-6	1324	-1274	14	324	319	
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-26	232	-240	6	-56	-157	5	815	-823	-13	-78	111
-25	-83	87	7	-84	58	6	537	-533	-12	-76	-78
-24	308	283	8	-87	-135	7	-73	50	-11	135	-142
-23	-82	-79	9	306	293						
-22	499	505	-25	-85	-101	8	687	-695	-10	437	-432
-21	265	-277	-24	140	-150	9	467	440	-9	537	603
-20	282	-279	-23	-83	-13	10	294	322	-8	750	770
-19	156	-136	-22	406	-415	11	-78	-102	-7	300	333
-18	-82	-125	-21	322	325	12	149	93	-6	651	603
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-14	-76	47	-17	403	-471	16	174	171	-2	355	-411
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-6	344	-343	-9	179	-160	-23	144	-149	3	149	-134
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7	-81	-65	-23	-86	92	-7	892	906	-25	103	171
8	142	-144	-22	-84	-8	-6	649	651	-24	-84	52
9	201	165	-21	151	132	-5	390	415	-23	330	337
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11	158	151	-19	-84	-83	-3	-60	3	-21	402	-383
12	140	-76	-18	270	-258	-2	759	-785	-20	225	-235
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-23	-80	-32	-13	257	-271	0	944	975	-15	122	-113
-22	-77	-29	-12	-32	94	1	112	-42	-14	-79	-56
-21	-80	-98	-11	439	-292	2	465	474	-13	653	542
-20	339	-357	-10	-82	-64	3	105	109	-12	-77	-110
-19	105	185	-9	193	200	4	280	-297	-11	290	319
-18	465	478	-8	209	227	5	269	235	-10	-75	34
-17	-80	9	-7	429	424	6	296	-272	-9	407	401
-16	137	109	-6	-54	31	7	832	878	-8	-74	-57
-15	154	-171	-5	-63	84	8	-74	-83	-7	504	-501

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC	
-6	331	-348	-11	236	-244	15	171	171	-22	243	237	
-5	423	-408	-10	-81	-57	10	-61	102	-21	200	168	
-4	722	-733	-9	373	-373	17	369	-373	-20	-82	108	
-3	248	-263	-8	132	-120	18	182	-190	-19	570	-576	
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2	701	-682	H+	11,	K= 9	1	220	214	-10	653	-672	
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12	-63	5	8	-21	401	405	11	456	442			
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14	-86	33	10	-19	-82	-51	13	-82	47	0	211	138
-26	129	-81	11	-18	-81	27	14	204	-213	1	308	-332
-25	218	235	12	-17	-31	17	15	470	-453	2	452	-455
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-23	156	-143	14	159	-155	17	454	-437	4	351	-328	
-22	231	215	15	-14	209	-225	18	218	233	5	378	373
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-19	219	227	18	-11	102	-150	-23	155	136	8	-50	19
-18	128	153	19	-10	257	-200	-22	341	-349	9	313	-293
-17	118	-52	20	-9	147	113	-21	226	202	10	434	-343
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-9	225	-191	28	-1	-34	-7	-14	256	262	-22	453	450
-8	527	-544	H+	13,	K= 9	-13	-78	-7	-13	214	-232	
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-5	617	614	H-	15	-67	122	-10	438	-439	-18	-79	80
-4	377	383	H+	16	-30	-72	-9	610	634	-17	203	214
-3	184	190	H-	15	190	-173	-8	193	-191	-16	127	64
-2	777	730	H+	14	-65	-30	-7	145	146	-16	526	530
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H+	9,	K= 9	-12	-65	20	-5	295	-291	-13	147	111	
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3	115	129	-7	-57	-45	H+	4,	K= 10	-7	231	-215	
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6	-83	-66	-4	420	-405	2	344	343	-4	-76	-13	
7	219	-216	H+	0,	K= 10	3	425	-397	-3	583	580	
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9	235	191	H+	113	-1196	5	-79	52	-1	261	-253	
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-24	-84	35	H-	3	415	395	8	345	401	1	312	334
-23	328	-325	H+	4	820	823	9	301	273	2	233	-237
-22	477	-493	H-	5	153	-151	10	400	-402	3	352	324
-21	233	230	H+	6	442	-443	11	277	-283	4	324	305
-20	-81	58	H-	7	355	-355	12	-61	-109	5	-80	-13
-19	328	351	H+	8	384	-384	13	257	-250	6	151	-140
-18	130	105	H-	9	141	-153	14	139	153	7	201	-219
-17	221	-253	H+	10	528	536	15	195	-275	8	241	-254
-16	-79	101	H-	11	305	375	16	-84	11	9	159	-160
-15	-80	81	H+	12	159	-100	17	218	220	10	198	190
-14	-80	6	H-	13	352	423	18	233	-233	1	159	-160
-13	362	-367	H+	14	117	-93	19	422	419	10	198	190
-12	-81	26	H-	14	117	-93	H+					

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC		
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-22	179	-172		2	259	248	-6	311	311	-8	538	544	
-21	240	-209		3	163	169	-5	-79	-137	-7	-82	-46	
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3	241	-225		29	403	-376	-20	522	-527	-11	-81	-22	
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6	-87	-105		32	638	710	-17	255	-247	-8	194	-216	
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-21	242	202		34	687	-727	-15	340	305	-6	531	-521	
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				55	447	-440							
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-16	172	-184		60	432	-393	8	467	-456	-2	187	-192	
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-14	-87	15		62	-86	85	10	-87	122				
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-5	-87	-29		71	532	609	18	295	293	0	169	-204	
-4	250	-226		72	262	241	19	431	413	7	291	298	
-3	129	60		73	1035	1033	14	162	-174	8	-80	-62	
-2	320	-332		74	564	563	15	117	-1	9	125	-100	
				75	521	-500	16	832	-339	10	651	600	

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC		
11	569	559	-11	504	519	H=	10	K=	12	-18	233	-302	
12	293	224	-10	714	-720				-17	249	342		
13	168	-176	-9	158	-160				-16	126	92		
14	510	-492	-8	-91	15		0	175	-140	-15	181	-165	
15	-80	43	-7	273	239		-17	-83	135	-14	322	244	
16	399	-386	-6	-31	91		-16	534	-526	-13	-32	-143	
17	-82	5	-5	117	65		-15	192	121	-12	-60	-13	
18	345	333	-4	-51	79		-14	253	-210	-11	-60	112	
19	-65	43	-3	464	-472		-13	-84	-64	-10	163	-224	
H=	2,	K=	12	-2	232	-222		-12	477	469	-9	176	-159
							-11	-87	-95	-8	-81	37	
							-10	238	231	-7	303	324	
0	-80	-99	H=	5,	K=	12	-9	126	123	-6	123	-137	
1	208	-283		0	133	142	-8	271	-270	-5	302	317	
2	387	393		1	237	228	-7	227	-230	-4	120	-39	
3	322	-316		2	337	-338	-6	-36	64	-3	253	-279	
4	712	-695		3	254	230	-5	304	301	-2	500	-500	
5	653	600		4	590	593	-4	506	496	-1	601	-587	
6	-82	-63		5	542	-540	-3	134	103				
7	122	-105		6	-31	55	-2	-85	-73	H=	5,	K=	13
8	280	266		7	-82	-29	-1	-86	-82	0	303	310	
9	195	171		8	288	-281				1	504	559	
10	-81	11		9	-85	-68	H=	1,	K=	13	2	402	-397
11	-62	42		10	134	-111	0	375	-384	3	526	536	
12	502	-563		11	143	-45	1	706	-769	4	-81	-41	
13	165	-132		12	257	234	2	424	436	5	-82	-117	
14	273	-263		13	194	-161	3	464	-510	6	220	-153	
15	-83	-33		14	-19	-33	4	-83	40	7	435	-433	
16	316	280		15	-19	-29	5	150	175	8	-35	37	
17	-86	-37		16	401	475	6	298	303	9	159	141	
-20	-86	153		17	-63	-68	7	452	463	10	180	-173	
-19	-65	115		18	506	490	8	-60	12	11	-66	-2	
-18	498	-508		19	138	-53	9	116	-129	-13	354	343	
-17	-83	32		20	-81	90	10	164	161	-18	167	-177	
-16	527	-510		21	-62	133	11	-82	103	-17	-83	-77	
-15	-79	16		22	555	-565	12	-16	103	-15	301	293	
-14	124	105		23	175	129	13	-32	-40	-15	296	-327	
-13	205	-197		24	239	-302	14	210	-249	-14	135	132	
-12	340	856		25	272	-221	15	193	-172	-13	-30	32	
-11	-81	-66		26	213	254	16	340	-277	-12	347	-343	
-10	150	140		27	-32	135	17	-55	115	-11	-32	-73	
-9	211	185		28	-81	-64	18	-13	115	-10	147	-112	
-8	409	-466		29	692	-723	19	-83	32	-9	156	103	
-7	242	-238		30	737	-742	20	237	-315	-8	400	403	
-6	183	180		31	-31	-42	21	205	-224	-7	268	269	
-5	810	830		32	194	195	22	-19	151	-6	307	-290	
-4	946	935		33	230	247	23	13	-112	-5	416	-442	
-3	116	25		34	-	-	24	410	422	-4	161	-138	
-2	367	-380	H=	8,	K=	12	25	-11	195	-9	-3	146	-163
-1	165	-164		26	-81	-92	26	162	165	-2	236	243	
H=	4,	K=	12	27	-81	46	27	471	-429	-1	-81	5	
				28	131	-90	28	406	-354				
0	275	277		29	220	-220	29	300	339	H=	7,	K=	13
1	218	194		30	-83	-75	30	551	552				
2	-80	14		31	103	-132	31	147	157	0	-83	22	
3	410	402		32	-85	-74	32	277	282	1	270	245	
4	-81	61		33	-50	113	33	303	-403	2	347	-318	
5	-80	62		34	153	153	34	153	-112	3	119	-129	
6	194	207		35	-84	-3	35	-112	4	150	-141		
7	252	-261		36	517	553	H=	3,	K=	13	5	346	-350
8	254	274		37	-83	15	37	406	-354	6	137	140	
9	343	276		38	146	-143	38	149	121	7	-85	73	
10	507	-552		39	-81	-13	39	221	-173	-18	250	238	
11	329	-312		40	524	-551	40	301	372	-17	219	-214	
12	426	-412		41	364	345	41	-81	57	-16	-83	-33	
13	-80	65		42	126	-60	42	-50	35	-15	-84	94	
14	436	414		43	350	-371	43	501	477	-14	237	-221	
-21	-83	-51		44	-81	643	44	275	-265	-13	163	193	
-20	206	-203		45	-80	51	45	175	-143	-12	-82	-20	
-19	-84	73		46	-30	103	46	419	405	-11	-82	-62	
-18	534	-594		47	172	-193	47	136	-181	-10	-82	73	
-17	-82	-3		48	210	-206	48	-43	57	-9	140	157	
-16	351	365		49	182	-188	49	127	75	-8	-79	53	
-15	-80	42		50	-81	-73	50	408	-422	-7	227	-238	
-14	654	666		51	322	297	51	100	-111	-6	166	160	
-13	252	-271		52	323	303	52	-87	-53	-5	428	-450	
-12	208	-164		53	395	404	-19	120	-2	-4	-81	4	

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
-3	-80	49	-6	165	183	-12	-63	-36	0	501	496
-2	512	516	-5	411	410	-11	-85	-115	1	353	345
-1	562	562	-4	201	-203	-10	-84	-29	2	-86	33
H=	9, K=	13	-3	130	104	-9	202	202	3	242	-250
0	264	-235	-2	542	-575	-8	313	-325	4	361	-362
-15	269	270	-1	173	-189	-7	140	-157	-13	466	474
-14	315	-295	H=	4, K=	14	-6	204	216	-12	191	147
-13	-64	0	0	565	555	-5	263	-270	-11	152	-167
-12	170	183	1	-83	-112	-4	130	145	-10	-84	11
-11	-65	64	2	133	133	-2	206	210	-8	161	-140
-10	149	139	3	475	438	-1	-84	62	-7	-84	36
-9	-83	-20	4	-33	4	H=	1, K=	15	-6	379	-381
-8	434	-425	5	117	-59	-5	422	-428	-5	372	377
-7	278	-280	6	357	-309	0	455	-429	-3	263	-245
-6	193	187	7	313	-293	1	290	-242	-2	230	-209
-5	270	254	8	-65	83	2	135	115	-1	178	164
-4	240	256	9	225	-14	3	421	404	H=	7, K=	15
-3	189	154	10	142	113	4	491	460	-8	274	-287
-2	-83	-23	-17	213	-162	5	-82	97	-7	230	229
-1	-83	73	-16	211	-209	6	147	-119	-6	170	-171
H=	0, K=	14	-14	166	205	7	198	-165	-5	-86	28
0	746	-779	-13	-83	85	8	280	-242	-4	390	386
1	-81	-59	-12	126	-73	9	-85	-114	-3	328	-334
2	-80	-41	-11	210	220	10	-84	55	H=	0, K=	16
3	502	-479	-10	142	60	11	245	221	-8	237	-274
4	-80	62	-9	148	-160	-12	130	-47	1	-35	-93
5	313	315	-8	270	281	-11	375	377	2	153	113
6	411	409	-7	251	270	-10	-82	-6	3	140	91
7	407	399	-6	314	-345	-9	265	273	4	-84	79
8	117	-125	-5	208	240	-8	191	153	5	273	253
9	281	-271	-4	-81	-47	-7	199	-158	6	199	-193
10	-81	-54	-3	328	-320	-6	433	436	7	132	140
11	-82	65	-2	-31	"116	-5	402	-438	8	-84	-38
12	146	120	-1	-79	-16	-4	-83	93	H=	2, K=	16
13	157	112	H=	5, K=	14	-3	300	294	-8	2	16
14	191	-200	-2	-2	371	-402	-2	-82	15	-85	46
15	-84	46	0	339	332	H=	3, K=	15	0	-85	46
H=	2, K=	14	1	275	233	-83	302	310	1	-84	-77
0	128	-140	2	361	-378	-6	-83	-33	2	-85	123
1	123	-164	3	283	-271	1	360	346	3	-85	133
2	465	463	5	365	-355	2	268	288	4	185	132
3	-81	41	6	-86	30	3	-85	86	5	-86	-72
4	277	252	-16	128	114	4	249	-257	-9	351	354
5	344	353	-15	-85	-67	5	249	-207	-8	157	155
6	-80	-32	-14	-84	92	6	326	-307	-7	404	-410
7	355	-335	-13	254	229	7	127	-100	-6	289	-262
8	132	-110	-12	154	-179	8	200	-185	-5	174	-191
9	161	-134	-11	-82	-41	-13	207	202	-4	241	-256
10	199	182	-10	-84	51	-12	190	149	-3	167	196
11	-85	25	-9	-83	34	-11	427	398	-2	146	-125
12	-87	-144	-8	168	-163	-10	-65	30	-1	-85	17
13	238	-205	-7	447	418	-9	-32	-18	H=	4, K=	16
-16	194	-182	-6	131	-151	-8	298	310	-82	264	249
-15	-82	4	-5	300	-324	-7	229	-219	1	-86	103
-14	-84	-64	-4	-81	55	-6	-34	57	-7	253	-249
-13	204	-182	-3	320	-294	-5	-85	-23	-6	-85	-53
-12	170	220	-2	422	442	-4	537	-525	-5	230	217
-11	-81	106	-1	-83	50	-3	346	347	-4	-86	-103
-10	-81	-122	H=	8, K=	14	-2	173	-120	-3	129	134
-9	-80	-104	-1	339	-377	H=	5, K=	15	-2	140	-122
-8	224	201	0	332	-377	-13	-34	-53			
-7	462	-451	-13	-34	-53						

Table B-5
Observed and Calculated Structure Factors for $C_4(fph)_4Rh(cp)(tpp)$

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC		
H= 0, K= 0	-18	177	173	-14	223	336	-7	461	-489				
	-17	-65	22	-13	374	348	-6	526	-544				
	-16	190	-191	-12	410	-402	-5	-56	49				
1	-21	1092	-15	187	183	-11	496	-515	-4	-56	-6		
2	1897	-1630	-14	172	157	-10	176	180	-3	967	-560		
3	871	-822	-13	-61	-60	-9	409	431	-2	90	0		
4	213	-104	-12	617	-621	-8	217	206	-1	470	-415		
5	908	856	-11	541	-504	-7	238	-202					
6	334	-211	-10	463	463	-6	180	-225	H= 0, K= 7				
7	1247	-1240	-9	313	289	-5	1102	1068	0	690	-603		
8	631	-596	-8	522	-537	-4	838	897	1	105	-107		
9	719	723	-7	1485	-1483	-3	550	-509	2	406	415		
10	361	244	-6	-54	-53	-2	682	-613	3	-66	-8		
11	-50	39	-5	1687	1629	-1	-57	25	4	-67	1		
12	342	-393	-4	353	429	H= 0, K= 5	0	353	347	5	252	-241	
13	144	-172	-3	-45	189		1	337	-371	6	173	-164	
14	539	536	-2	920	-548				7	244	249		
15	214	234	-1	105	156				8	113	132		
16	325	-722					0	353	347	9	205	-188	
17	138	-173	H= 0, K= 3	2	648	640					242	237	
				3	593	586	-21						
				4	301	-269	-20	-67	81				
H= 0, K= 1	0	1047	699	5	460	-475	-19	93	-86				
	1	307	-760	6	167	165	-18	-64	-71				
0	1644	-2456	2	559	525	7	235	223	-17	-65	-41		
1	155	-81	3	464	442	8	165	151	-16	652	449		
2	301	-154	4	149	186	9	143	-156	-15	217	216		
3	1179	1084	5	274	218	10	305	-305	-14	395	-309		
4	332	-259	6	-55	-12	11	135	164	-13	180	-186		
5	1289	-1263	7	257	279	12	238	282	-12	300	309		
6	128	-108	8	-62	104	13	-20	-68	-11	441	415		
7	537	519	9	211	-197	14	-19	21	-10	145	-149		
8	389	423	10	310	-338	15	-19	-307		438	-458		
9	254	-248	11	-69	85	16	-15	-52	-9	378	-381		
10	410	-423	12	293	296	17	423	428	-7	176	199		
11	-63	-6	13	-69	-84	18	218	237	-6	102	41		
12	427	403	14	130	-128	19	184	175	-5	320	-337		
13	120	134	15	244	-243	20	222	-225	-4	-54	-20		
14	282	-283	16	311	-310	21	108	96	-3	134	128		
15	-53	1	17	357	368	22	862	852	-2	326	367		
16	225	237	18	-66	94	23	720	741	-1	275	292		
-18	-67	-9	19	239	-232	24	94	122					
-17	158	174	20	158	-146	25	487	-477	H= 0, K= 8				
-16	192	187	21	110	-89	26	341	-323					
-15	-69	-90	22	140	145	27	470	-464	0	230	235		
-14	383	-375	23	382	406	28	417	341	1	242	-241		
-13	-53	-52	24	-54	-154	29	662	-722	2	-67	57		
-12	432	425	25	196	-201	30	1307	-1306	3	-70	-94		
-11	123	134	26	220	201	31	1195	1196	4	140	-118		
-10	820	-842	27	359	322	32	130	165	5	262	269		
-9	331	-304	28	163	-210	33	418	475	6	-70	73		
-8	257	250	29	964	-1036	H= 0, K= 6	0	1116	1108	7	214	-245	
-7	631	607	30	254	-223		1	385	379	8	108	-72	
-6	308	235	31	949	-776		2	188	-178	-21	326	-325	
-5	599	-915	32	116	253		1	-18	-57	-20	-68	-103	
-4	102	-36	33	1145	-997		-1		60				
-3	503	553											
-2	1102	1449	H= 0, K= 4	0	-57	-64	3	215	-210	-18	173	167	
-1	552	-650		1	286	307	4	169	159	-17	-68	146	
			2	591	-610	5	355	357	-16	401	-380		
H= 0, K= 2	0		1	286	307	6	145	129	-15	-65	-7		
			2	591	-610	7	213	-212	-14	445	455		
0	797	195	3	624	-619	8	192	-205	-13	247	232		
1	444	-363	4	324	328	9	132	147	-12	148	-176		
2	611	-772	5	628	605	10	-71	98	-11	625	-414		
3	1243	-1260	6	122	-103	11	-72	-88	-10	194	206		
4	157	223	7	450	-488	12	-67	-86	-9	464	474		
5	1147	1107	8	141	-112	13	177	177	-8	116	-86		
6	77	14	9	-56	60	14	139	147	-7	541	-540		
7	348	-375	10	165	153	15	163	-191	-6	300	-295		
8	368	-264	11	260	-261	16	627	-638	-5	573	557		
9	243	252	12	358	-352	17	-65	-144	-4	390	392		
10	329	334	13	194	-125	18	279	283	-2	-60	-22		
11	-66	-30	14	160	-136	19	128	-148	-2	799	-323		
12	323	-317	15	292	384	20	330	-344	-1	145	-142		
13	-67	36	16	-66	2	21	693	-680					
14	301	301	17	414	-407	22	-52	61	H= 0, K= 9				
15	-70	38	18	-64	-18	23	596	589	0	444	-449		
-19	104	86	19	-63	71	24	209	211					

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC	
1	360	-393	-2	-67	17	-15	521	492	-14	129	111	
2	310	302	-1	-67	-80	-14	267	297	-13	-58	-10	
3	323	311				-13	300	-287	-12	806	-761	
4	-68	12	H=	0,	K=	12	-12	236	-266	-11	423	-445
5	112	-95				-11	324	-303	-10	483	511	
6	-70	-40	0	117	108	-10	149	172	-9	962	939	
-20	108	118	-19	265	250	-9	-52	64	-8	464	-436	
-19	-66	79	-18	260	270	-8	803	-736	-7	430	-413	
-18	193	-760	-17	-56	-35	-7	545	-567	-6	575	-433	
-17	-65	63	-16	254	-233	-6	721	732	-5	1231	123	
-16	444	649	-15	-67	-63	-5	1005	914	-4	603	367	
-15	181	-170	-14	177	174	-4	704	-717	-3	1367	-1613	
-14	522	-534	-13	233	252	-3	314	-593	-2	1400	-1647	
-13	-63	11	-12	166	-220	-2	682	-950	-1	-51	317	
-12	-62	25	-11	345	-351	-1	1574	2416	H=	1,	K=	
-11	-62	27	-10	100	-49				1		3	
-10	108	-182	-9	282	270	H=	1,	K=	0	904	-878	
-9	228	-250	-8	133	172				1	303	303	
-8	-61	-24	-7	243	-255	0	514	-454	2	771	757	
-7	283	308	-6	169	-150	1	139	-216	3	463	421	
-6	203	221	-5	-69	64	2	792	929	4	119	107	
-5	271	-265	-4	129	159	3	188	134	5	505	-471	
-4	190	-230	-3	218	-229	4	267	-356	6	98	-80	
-3	287	-277	-2	145	-139	5	875	-755	7	315	329	
-2	529	527	-1	-71	-48	6	854	856	8	-64	-12	
-1	-68	-65	H=	0,	K=	13	7	789	788	9	313	-325
H=	0,	K=	10			8	272	-293	10	276	-268	
0	430	426	-17	125	103	9	501	-492	11	223	198	
1	110	104	-16	132	134	10	210	-214	12	320	317	
2	-68	-99	-15	100	92	11	153	132	13	120	-176	
3	112	-125	-14	147	-138	12	403	401	14	245	-266	
4	-71	-49	-13	225	-231	13	-66	-30	-10	-69	65	
-20	174	-173	-12	-66	34	14	264	-258	-19	354	-345	
-19	-67	76	-11	-70	122	15	769	-63	-18	-60	-69	
-18	290	286	-10	-69	-56	16	-67	-30	-17	329	328	
-17	360	-277	-9	109	-107	17	192	217	-16	132	106	
-16	488	-493	-8	-69	51	18	380	388	-15	120	132	
-15	241	251	-7	107	136	19	161	-174	-13	247	-230	
-14	575	566	-6	166	-146	20	210	-307	-12	696	613	
-13	397	397	-5	-73	-128	21	103	109	-11	595	567	
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-8	-66	82	-15	137	-122	-8	115	190	-6	191	206	
-7	378	-370	-14	160	186	-7	424	419	-5	269	-242	
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-3	-70	110	-10	-70	47	-3	73	-414	-1	715	741	
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-19	244	-254	3	284	-369	4	775	804	4	626	620	
-18	174	-190	4	710	-636	5	214	269	5	341	319	
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-16	160	-172	6	1064	-1021	7	463	-481	7	117	103	
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-11	340	296	11	240	-267	12	405	-403	-19	261	259	
-10	172	-188	12	408	-404	13	107	147	-18	-64	6	
-9	226	-236	13	-68	102	14	203	225	-17	142	-155	
-8	161	-121	14	482	496	15	133	95	-16	235	-216	
-7	152	143	15	-69	-6	-19	188	180	-15	135	-151	
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-5	156	-177	-18	-69	-39	-17	213	-312	-13	119	223	
-4	285	-273	-17	393	-280	-18	181	-179	-12	975	-938	
-3	-69	90	-16	150	-173	-15	134	122	-11	242	203	

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC	
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-9	275	261	-2	704	-730	-19	267	-279	-15	-65	-74	
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-7	505	-454	H=	1.	K=	7	-17	416	-13	177	196	
-6	255	-214				-16	307	299	-12	258	-265	
-5	952	1021	0	552	-550	-15	496	-501	-11	210	-207	
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3	136	130	-22	-66	-67	-4	355	-333	13	.	.	
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7	224	241	-16	432	441	H=	1.	K=	-14	258	-269	
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9	341	-334	-14	507	-516	1	-70	102	-12	202	207	
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11	180	169	-12	-57	67	3	179	-200	-10	-69	-13	
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-19	320	-328	-9	797	-809	-18	104	102	-6	179	175	
-18	122	-159	-8	89	57	-19	135	141	-5	183	-187	
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-13	183	164	-3	-58	87	-14	502	529	-11	-71	-71	
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-11	185	171	-1	-59	0	-12	239	-265	11	188	-169	
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-9	666	-691				-10	198	216	H=	1.	K=	
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4	134	151	-15	194	201	-18	-66	-96	2	137	154	
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6	-68	-5	-13	254	239	-16	122	144	4	282	-274	
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9	-71	87	-10	311	328	-13	-66	-97	7	-63	40	
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-15	274	269	-2	433	-435	-5	271	-280	15	102	124	
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-7	218	-208	4	-71	79	-19	252	256	3	-66	21	
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-5	423	415	-21	98	87	-17	191	-173			.	

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
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6	368	-287	4	221	-260	18	114	-64	-7	-62	15
7	179	-195	5	375	-349	19	125	-104	-6	377	-356
8	218	228	6	-58	-7	20	-66	79	-5	652	-655
9	347	225	7	346	343	-10	174	-157	-4	80	-70
10	-66	81	8	219	-229	-9	203	-171	-3	348	375
11	406	-290	9	449	-455	-8	190	163	-2	364	363
12	155	-163	10	224	207	-7	208	326	-1	145	146
13	202	217	11	305	297	-6	210	-276	H=	1+	K= -4
14	105	149	12	200	209	-5	117	-100	0	1574	1465
15	92	-61	13	-62	-4	-4	125	-110	1	490	425
16	-64	-42	14	197	-202	-3	421	298	2	743	-921
17	-68	-40	15	-66	18	-2	598	590	3	244	-312
18	304	296	16	357	402	-1	186	-182	-1	1010	1064
19	-67	0	17	-63	-37	H=	1+	K= -6	5	-54	92
-1	177	192	18	119	-109	-6	-66	-90	6	1126	-1002
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3	110	-95	-3	294	285	5	429	471	13	321	329
4	358	-364	-2	-67	83	6	674	-680	14	191	185
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6	210	209				8	-50	36	16	407	-422
7	190	159	H=	1+	K= -8	9	305	295	17	182	-203
8	-64	25				10	210	211	18	206	233
9	256	-263	0	320	340	11	385	-390	19	170	175
10	99	-37	1	568	-572	12	280	-296	20	-68	62
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13	187	-152	4	619	577	15	256	-235	23	133	-141
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16	147	146	7	479	-487	18	293	287	26	-62	-31
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-3	274	271	11	328	-330	-11	127	-117	30	821	-795
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1	362	-352	18	130	118	-4	298	272	0	2217	-1992
2	357	-362	19	148	160	-3	289	-383	1	654	563
3	-64	12	20	190	-190	-2	426	-414	2	1744	1647
4	303	286	-9	138	106	-1	592	617			
5	180	197	-8	165	-182	H=	1+	K= -5	3	283	163
6	571	-578	-7	278	-275	H=	1+	K= -5	4	823	-778
7	593	-601	-6	175	180				5	267	-253
8	320	312	-5	253	275	0	845	-887	6	196	-123
9	431	424	-4	-60	-40	1	1124	-1087	7	265	250
10	140	-105	-3	-63	0	2	1322	1200	8	-53	-30
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13	519	527				5	377	-320			
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15	207	-194				7	-54	119	12	386	370
16	589	-613	0	146	-166	8	146	-161	13	170	-129
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19	-66	16	3	140	130	11	851	870	16	325	324
20	177	-164	4	626	542	12	562	574	17	-60	53
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L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC		
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2	689	-822	4	132	-145	7	254	-250	10	-68	-48		
3	292	-234	5	261	247	8	344	341	11	160	-173		
4	793	823	6	862	-632	9	348	359	12	243	-231		
5	858	751	7	599	-973	10	131	234	13	156	164		
6	599	-540	8	266	259	11	254	-264	14	416	414		
7	745	-193	9	195	184	12	128	-107	15	218	-206		
8	190	151	10	-63	89	13	-69	123	16	118	-153		
9	586	68	11	286	-273	14	-73	124	15	211	181		
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11	699	-701	13	551	581	16	412	405	13	512	-318		
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-16	165	-135	-14	294	280	11	92	-66	5	691	666		
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-11	-62	92	-9	893	847	6	705	601	H=	2+	K=	5	
-10	581	564	-8	938	-943	5	1305	1291					
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-8	475	-479	-6	310	319	3	102	132	1	-57	2		
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-4	201	-256	-2	212	-558				4	257	-257		
-3	1120	-1111	-1	1120	909				5	257	-257		
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-1	142	729	H=	2+	K=	1	1	-55	7	280	269		
H=	1+	K=	-1	0	82	126	2	445	496	8	118	15	
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5	970	-904	7	203	236	8	105	-134	18	-63	2		
6	1006	674	8	307	-512	9	303	-368	17	151	111		
7	1558	1555	9	276	-267	10	217	-245	16	243	270		
8	76	85	10	153	-134	11	292	301	15	-60	-15		
9	540	-520	11	262	281	12	-60	42	14	293	-301		
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12	200	228	14	129	-169	15	107	-99	11	-53	19		
13	118	-162	15	150	-150	16	305	311	10	315	-308		
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15	-69	59	17	437	440	18	635	-635	7	467	431		
16	248	230	18	102	176	19	224	-210	6	108	-143		
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-13	221	214	-10	523	-528	25	-54	-102					
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-11	162	-170	-8	452	431	27	268	248	1	164	177		
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L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC	
9	193	198	-6	361	-348	-7	201	216	2	207	-203	
-21	195	-198	-5	108	127	-6	-68	-42	3	205	209	
-20	103	85	-4	582	590	-5	432	-442	4	200	202	
-19	150	180	-3	334	-362	-4	192	-103	5	210	-101	
-18	156	141	-2	463	-462	-3	220	208	6	302	-305	
-17	162	-164	-1	135	130	-2	104	149	7	93	-71	
-16	352	-353				-1	-71	-6	8	250	240	
-15	266	284	H=	2+	K= 9	H=	2+	K= 12	10	247	-243	
-14	594	617	0	278	-272	H=	2+	K= 12	11	275	-270	
-13	129	114	1	-69	37	-18	123	67	12	-65	-20	
-12	435	-637	2	131	108	-17	-68	-93	13	115	100	
-11	192	218	3	104	113	-16	129	-104	14	113	118	
-10	223	245	4	188	-167	-15	-60	42	15	194	150	
-9	595	564	5	-21	-66	-14	229	228	16	194	-193	
-8	231	-243	6	-38	-38	-12	-69	103	17	-66	73	
-7	987	-984	7	106	-94	-12	204	-191	18	233	234	
-6	261	-239	8	189	-184	-11	161	-163	-2	112	104	
-5	136	87	9	-64	-56	-10	-65	15	-1	131	138	
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2	-68	-50	17	132	-152	H=	2+	K= 13	6	195	177	
3	105	182	18	444	445	-16	143	156	7	164	160	
4	121	107	19	332	322	-15	122	-66	8	-64	73	
5	152	-142	20	312	308	-14	151	-168	9	153	-172	
6	189	-189	21	416	-421	-13	-69	44	10	111	67	
7	-71	69	22	104	-31	-12	163	166	11	434	473	
8	265	259	23	390	365	-11	-71	22	12	59	-67	
9	-21	140	24	144	-152	-10	-70	-64	13	346	-343	
10	-66	-31	25	453	-459	-9	-72	-94	14	-62	-112	
11	-64	93	26	-85	H=	2+	K= 10	-6	15	-63	-33	
12	-66	-85	27			H=	2+	K= -14	16	229	242	
13	293	301	28	362	0	202	191	H=	2+	K= -14	17	
14	349	-366	29	-71	53	4	259	237	18	301	-294	
15	445	-470	30	120	-123	5	-70	-50	19	-68	25	
16	-60	31	31	-68	-30	6	220	-203	20	302	245	
17	440	439	32	-67	-59	7	-67	5	249	256		
18	230	236	33	267	267	8	198	194	2	105	93	
19	406	-421	34	192	177	9	225	220	-1	-67	0	
20	222	-202	35	292	-295	10	-69	-45	H=	2+	K= -10	
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22	305	336	37	228	252	12	-67	-103	1	414	-456	
23	-138	-146	38	362	377	13	298	251	2	307	-299	
24	-57	-31	39	99	58	14	-67	-2	3	229	240	
25	346	-323	40	272	-261	15	193	-166	4	373	363	
26	297	286	41	245	-261	16			5	113	-114	
27	441	473	42	257	262	17	265	261	6	403	-409	
28	-64	37	43	-8	183	159	H=	2+	K= -13	7	310	-309
H=	2+	K= 8	44	260	-270	0	-68	-25	8	94	-110	
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1	123	96	46	225	226	2	162	165	10	260	-262	
2	128	-139	47	106	96	3	143	-125	11	241	-113	
3	99	-34	48	374	-368	4	195	-126	12	215	212	
4	164	139	49	-69	1	5	-66	-29	13	424	477	
5	250	239	50	-69		6	139	146	14	304	249	
6	-67	-63	H=	2+	K= 11	7	147	162	15	373	-355	
7	-20	-20	8	-69	-5	10	-64	-34	16	232	-233	
8	-65	27	9	393	-393	11	211	305	17	130	119	
9	-64	48	10	130	-164	12	-65	-2	18	307	306	
10	477	-453	11	-65	40	13	160	-168	19	-67	-30	
11	237	-242	12	-65	40	14	-65	64	20	394	340	
12	-65	41	13	155	117	15	-70	124	21	201	213	
13	154	160	14	99	-103	16	147	197	-4	293	-299	
14	100	120	15	237	-232	17	-67	16	-3	137	-104	
15	206	-215	16	-66	-23	17	-67	16	-2	218	217	
16	341	-348	17	449	472	H=	2+	K= -12	-1	404	403	
17	453	447	18	-66	30	0	131	152				
18	131	174	19	-10	173	-214	1	156	-171			
19	222	-222	20	-66	26							
20	343	-320	21	134	144							

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC	
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0 737	-732	12	410	411	-14	-69	-102	-11	-62	36		
1 160	144	13	526	-548	-13	-69	107	-10	425	-403		
2 265	277	14	-63	4	-12	231	236	-9	-57	-78		
3 124	112	15	206	207	-11	-65	-61	-8	100	115		
4 295	-294	16	274	268	-10	286	-270	-7	404	403		
5 110	-139	17	-65	100	-9	-63	-64	-6	-56	-13		
6 513	506	18	96	-63	-8	123	116	-5	283	-249		
7 427	445	19	212	-220	-7	-58	-2	-4	-52	15		
8 366	-341	-11	-59	19	-6	247	-231	-3	120	13		
9 -59	-74	-8	359	363	-3	703	735	-1	344	78		
10 -60	14	-7	-65	9	-2	91	-154	H= 2, K= -2				
11 91	60	-6	251	-236	-1	332	322	0	966	1066		
12 394	387	-5	271	-276	H= 2, K= -4	1	1474	1446				
13 -64	-77	-4	253	-244	0	96	127	2	1016	-976		
14 401	-398	-3	458	447	1	348	-216	3	1064	1036		
15 294	319	-2	161	163	2	947	-838	4	322	315		
16 184	166	-1	709	-700	3	225	-215	5	118	-96		
17 194	-210	H= 2, K= -6	0	96	4	753	726	6	99	-101		
18 328	-325	-24	104	24	5	136	114	7	677	-654		
19 -65	267	0	1029	-1015	6	298	-294	8	394	404		
-7 136	114	1	253	-770	7	536	-536	9	428	405		
-6 141	-178	2	881	-84	8	689	-646	10	112	-150		
-5 99	-120	3	100	-124	9	505	590	11	200	-198		
-4 -65	15	4	122	-124	10	125	103	12	285	-298		
-3 120	161	5	95	-131	11	545	-548	13	191	173		
-2 314	323	6	553	-570	12	-59	-47	14	618	624		
-1 496	-495	7	886	-874	13	-61	-16	15	-67	10		
H= 2, K= -8	8	135	177	H= 2, K= -8	14	-63	16	16	254	-261		
9 761	720	15	-63	14	15	-63	-99	-17	208	-218		
10 -54	11	447	-438	16	-66	-87	-16	230	-231			
0 649	644	12	231	-279	17	-66	-40	-15	130	124		
1 -56	-26	13	250	258	18	276	284	-14	201	198		
2 431	-429	14	118	104	-15	-69	92	-13	94	-113		
3 315	312	15	151	154	-14	259	276	-12	448	-458		
4 662	637	16	304	-295	-13	160	-171	-11	158	126		
5 191	125	17	-64	39	-12	308	-319	-10	528	521		
6 703	-710	18	198	204	-11	147	118	-9	240	-212		
7 277	-288	19	206	178	-10	195	297	-8	453	-434		
8 698	701	20	109	-105	-9	49	-96	-7	614	-624		
9 305	312	-13	200	-200	-8	197	176	-6	404	443		
10 111	88	-12	258	-200	-7	317	-349	-5	-53	94		
11 329	-331	-11	-58	10	-6	260	287	-4	518	-511		
12 344	-338	-10	171	176	-5	1327	1306	-3	1459	-1084		
13 278	293	-9	-66	-65	-4	135	128	-2	1356	-1697		
14 260	264	-8	399	-386	-3	229	-232	-1	195	-540		
15 237	-232	-7	255	-242	H= 2, K= -1	112	-215					
16 135	-122	-6	166	170	0	846	-733	0	773	-489		
17 -63	40	-5	523	524	1	201	-59	1	618	625		
18 220	238	-4	172	-178	2	1288	1338	2	525	528		
19 115	83	-3	333	-363	3	-54	3	3	665	-622		
-10 165	216	-2	403	442	4	172	-181	4	428	-459		
-9 -68	5	-1	811	786	5	492	480	5	649	597		
-8 273	-283	H= 2, K= -5	6	649	610	6	495	495	6	107	145	
-7 354	-337	7	1472	1453	5	496	495	7	646	-658		
-6 325	319	0	649	610	6	940	929	8	202	-196		
-5 263	269	1	1472	1453	7	940	929	9	306	-289		
-4 -64	300	2	364	441	8	449	-438	10	237	236		
-3 345	-322	3	250	-219	9	603	-621	11	237	236		
-2 272	-266	4	828	-587	10	202	188	12	271	259		
-1 748	773	5	761	-754	11	532	529	13	893	-940		
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0 393	-266	7	510	473	13	95	88	15	183	177		
1 201	164	8	175	-129	14	239	-235	16	292	295		
2 376	423	9	251	255	15	-66	32	-18	-67	28		
3 860	830	10	664	674	16	231	236	-17	179	161		
4 159	-176	11	127	117	17	-66	-9	-16	-67	113		
5 583	-622	12	126	-67	18	209	-220	-15	143	-154		
6 717	746	13	-65	-117	19	163	175	-14	96	-56		
7 227	213	14	107	-127	20	140	-160	-13	-62	38		
8 893	-7	15	146	147	21	247	-256	-12	370	350		
9 539	-513	16	-64	-61	22	158	174	-11	-56	-10		
10 617	-592	17	257	-252	23	158	174					

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
-10	748	-736	-6	384	-305				6	118	-157
-9	683	-702	-5	1429	-1534	H=	3, K= 4		7	332	-310
-8	519	-482	-4	937	559				-21	180	-173
-7	367	235	-3	903	970	6	270	309	-20	100	96
-6	108	-64	-2	133	126	1	-56	-43	-19	288	285
-5	978	-693	-1	172	-262	2	584	-363	-18	-64	81
-4	257	316				3	308	370	-17	452	-452
-3	1128	1530	H=	3, K= 2		4	216	119	-16	154	-131
-2	652	-25				5	168	703	-15	408	470
-1	880	-1233	6	476	509	6	101	-69	-14	268	276
			7	247	-235	7	683	-295	-13	173	-166
H=	3, K= 0		8	1216	-1233	8	159	175	-12	142	-153
			9	450	-401	9	264	246	-11	198	-198
0	131	-6	4	849	662	10	122	-69	-10	668	691
1	344	-294	5	202	203	-21	110	-178	-9	526	502
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3	100	-71	7	126	-97	-19	-66	63	-7	807	-611
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6	546	-554	10	105	113	-16	162	-174	-4	194	179
7	-56	-90	11	369	-366	-15	273	260	-3	571	-571
8	380	366	12	209	-226	-14	514	522	-2	288	-219
9	-63	60	-20	-71	129	-13	261	-354	-1	370	357
10	195	202	-19	322	303	-12	423	-415			
11	293	-310	-18	177	-106	-11	226	223	H=	3, K= 7	
12	-66	-18	-17	292	-273	-10	-82	30			
13	239	239	-16	141	-138	-9	261	271	0	397	-380
14	133	148	-15	159	154	-8	370	-397	1	-69	22
-19	252	254	-14	249	274	-7	481	-151	2	225	211
-18	-67	56	-13	94	-86	-6	-56	145	3	-71	116
-17	357	-260	-12	1065	-1055	-5	73	125	4	196	-191
-16	262	263	-11	-53	11	-4	264	254	5	213	-215
-15	507	493	+10	801	771	-2	323	-372	6	203	194
-14	-63	51	-9	92	106	-2	241	-230	-21	-68	110
-13	-60	104	-8	644	-654	-1	329	341	-20	110	-114
-12	286	-295	-7	841	-711				-19	192	-194
-11	168	-201	-6	479	463	H=	3, K= 5		-18	194	-116
-10	670	720	-5	138	210				-17	357	344
-9	402	-367	-4	158	6	0	515	-455	-16	129	116
-8	962	-926	-3	961	-854	1	136	117	-15	590	-607
-7	385	-336	-2	942	-935	2	379	362	-14	224	-212
-6	776	-715	-1	218	-245	3	402	-422	-13	100	84
-5	171	-179				4	159	-158	-12	-59	19
-4	1654	1190	H=	3, K= 3		5	143	-144	-11	-60	-75
-3	1724	-1491				6	115	136	-10	565	-559
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-1	1633	1429	1	554	527	8	-70	-84	-8	511	513
			2	634	629	9	226	-228	-7	156	173
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			4	579	-567	-20	307	-314	-5	575	-560
0	496	484	5	107	-70	-19	177	-169	-4	205	-195
1	728	635	6	136	154	-18	103	-178	-3	327	309
2	622	627	7	256	243	-17	240	246	-2	320	325
3	-56	-2	8	-68	-67	-16	318	306	-1	457	-442
4	620	-570	9	451	-461	-15	219	-220			
5	-56	92	10	-69	96	-16	210	-219	H=	3, K= 6	
6	313	303	11	194	180	-13	327	309			
7	-59	7	-21	206	184	-12	407	407	0	100	119
8	132	-123	-20	117	-162	-11	145	-157	1	-70	64
9	221	-215	-19	233	-227	-10	135	-133	2	222	-210
10	157	-172	-18	134	176	-9	261	-304	3	179	-169
11	658	459	-17	299	289	-8	326	323	4	248	216
12	213	206	-16	-62	30	-7	1011	1021	-21	-67	-67
13	131	-125	-15	267	-251	-6	162	-171	-20	-65	64
-20	158	-155	-14	319	-204	-5	567	-515	-19	230	232
-19	351	-276	-13	245	267	-4	610	634	-18	110	113
-18	-65	-19	-12	1075	1027	-3	527	530	-17	242	-234
-17	314	327	-11	195	183	-2	47	130	-16	99	-104
-16	-65	-11	-10	-51	54	-1	205	-298	-15	119	106
-15	294	-274	-9	242	235				-14	-65	-1
-14	421	-425	-8	714	757	H=	3, K= 6		-13	228	223
-13	-56	-79	-7	95	133				-12	413	-392
-12	455	461	-6	477	-412	0	341	351	-11	-63	11
-11	190	-211	-5	577	-479	1	190	-204	-10	425	427
-10	891	-892	-4	765	-716	2	191	-215	-9	55	92
-9	539	531	-3	412	-409	3	134	58	-8	-64	-63
-8	559	562	-2	309	377	4	290	272	-7	159	-163
-7	892	889	-1	570	-602	5	260	290	-6	-66	15

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
-5	562	573	-18	137	97	17	159	140	15	230	249
-4	-66	52	-17	180	-198	-3	-69	-94	16	-64	59
-3	503	-511	-16	265	-283	-2	108	127	17	287	-262
-2	128	126	-15	-69	43	-1	208	222	18	115	-135
-1	-70	100	-14	287	284	H=	3,	K= -11	-9	-68	69
H=	3,	K= 9	-13	-70	-46				-8	-67	37
0	264	-235	-12	234	-235	0	112	-139	-7	-66	1
1	-70	0	-11	137	-146	1	334	283	-6	164	-161
2	143	178	-9	165	176	2	176	199	-5	334	-334
-21	-67	63	-8	-72	-98	3	409	-410	-4	119	108
-20	111	132	-7	213	-210	4	171	-153	-3	191	177
-19	-66	-73	-6	104	65	5	160	165	-2	-64	-46
-18	-65	-25				6	166	182	-1	354	-330
-17	160	157	H=	3,	K= -15	7	119	71	H=	3,	K= -8
-16	245	-258				8	253	-232			
-15	150	-150	8	-70	-74	9	315	-300	0	287	-289
-14	247	235	9	108	116	10	607	501	1	511	-542
-13	-65	23	10	106	106	11	299	303	2	216	-226
-12	343	239	11	130	152	12	528	-535	3	-54	87
-11	-67	2				13	115	-134	4	368	394
-10	194	-208	H=	3,	K= -14	14	-64	4	5	91	-87
-9	175	162				15	122	123	6	840	-824
-8	232	233	3	122	119	16	95	50	7	272	-295
-7	-68	86	4	192	160	17	-57	-97	8	169	173
-6	-68	-69	5	123	-92	18	281	-264	9	228	246
-5	-70	-138	6	180	-188	-5	-78	-54	10	-59	1
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-3	-69	74	8	106	118	-3	101	111	12	205	-240
-2	-70	-66	9	128	121	-2	-67	-85	13	515	520
-1	283	-276	10	113	-132	-1	-64	-17	14	-65	-82
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0	113	125	13	220	216	0	-65	-67	17	-66	-2
-20	101	-73	14	-66	-52	1	605	-709	18	249	258
-19	204	210	15	143	-129	2	-73	64	-11	-69	79
-18	-68	91	H=	3,	K= -13	3	177	165	-10	132	119
-17	166	-177				4	-61	64	-9	-69	-141
-16	116	-107	0	117	-83	5	115	-110	-8	430	-442
-15	-65	-15	1	-69	99	6	385	-382	-7	-68	68
-14	-66	36	2	123	124	7	305	-305	-6	461	467
-13	108	80	3	271	-241	8	517	492	-5	137	134
-12	348	-352	4	194	-196	9	217	183	-3	138	-111
-11	-69	1	5	-64	66	10	333	-347	-2	-61	49
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-7	115	-93	9	263	-272	14	-65	51	0	112	-118
-6	-68	-40	10	136	140	15	243	-235	1	306	314
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2	513	-592	9	914	898	12	260	-713	17	291	-218
3	271	235	10	-57	72	13	550	359	16	273	262
4	768	741	11	260	-252	14	-69	-56	15	173	185
5	145	170	12	-62	35	15	518	-235	16	-61	39
6	950	-373	13	497	521	16	194	-177	13	246	-247
7	325	262	14	359	737	17	159	-259	12	597	-510
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9	192	213	16	331	-322	19	741	256	10	279	273
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11	352	-371	18	-70	-55	21	155	-163	8	744	-767
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3	568	-518	8	609	-630	12	489	-494	17	286	258
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8	346	-746	13	210	-209	17	313	296	12	243	292
9	647	-637	14	362	-356	18	-64	38	11	284	-270
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11	547	537	16	312	294	20	156	-124	9	784	705
12	-62	-83	17	162	-147	21	-66	80	8	207	234
13	393	-415	18	131	104	22	312	304	7	422	287
14	440	-457	19	-67	70	23	119	107	6	776	-800
15	267	253	20	340	-341	24	135	90	5	1192	-1016
16	227	247	21	138	-147	25	278	-261	4	430	324
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-9	155	-142	29	457	-438	33	779	-475	2	765	-761
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L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
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-17	232	-210	-8	404	-389	3	-70	-40	-19	151	163
-16	191	189	-7	191	-140	4	269	-271	-18	-68	-78
-15	428	429	-6	1356	1284	-21	-67	-69	-17	316	-347
-14	159	154	-5	625	635	-20	113	-87	-16	-68	-93
-13	327	-420	-4	101	-102	-19	150	-142	-15	234	241
-12	441	-406	-3	353	-352	-18	-65	-25	-14	239	252
-11	865	849	-2	519	-537	-17	367	389	-13	-66	-59
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-7	524	-499				-13	-63	-39	-9	-67	63
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-3	629	-607	3	-68	-7	-9	-62	-7	-5	370	393
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9	179	-185	-11	-57	25	-20	127	134	-10	154	-178
10	-69	48	-10	472	-476	-19	236	239	-9	-69	-52
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12	261	-254	-8	533	550	-17	233	-333	-7	160	184
13	151	-133	-7	-50	91	-16	450	-439	-6	-73	-125
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22	610	-612				-7	-66	-43	-12	167	-200
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24	647	636	1	366	-364	-5	268	277	-10	158	174
25	383	372	2	223	-203	-4	235	-257	H=	4+	K= -19
26	673	-936	3	-67	68	-3	-69	-74			
27	905	-868	4	253	267	-2	-68	27			
28	140	129	5	-69	34	-1	-68	36	7	115	-100
29	590	557	6	273	-253	H=	4+	K= 9	8	152	-145
30	370	-335	-22	225	-223	-22			9	123	-126
31	809	-851	-21	-68	-83	-20	188	190	10	139	125
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9	126	141	-8	497	-463	-11	98	-80	8	103	113
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17	-62	76				-3	227	219	0	141	131
18	627	-656	H=	4+	K= 7	-2	-70	-79	1	275	307
19	243	-246				-1	257	-226	2	-66	-43
20	155	158	0	176	-177	H=	4+	K= 10	3	253	-244
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13	161	-179	18	231	-222	4	645	-657	14	144	146	
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11	257	-254	2	150	-158	15	249	-252	-11	359	352	
12	192	-200	3	599	576	16	202	-194	-10	365	370	
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L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC	
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13	432	-455	-16	-64	-80	-9	716	716				
14	448	-468	-15	134	-130	-8	102	-59	0	177	-164	
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-17	168	163	-13	221	230	-6	769	-736	2	-66	-20	
-16	-65	-15	-12	420	441	-5	301	-289	3	-68	-15	
-15	256	-258	-11	-56	-33	-4	759	689	4	299	216	
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-13	317	316	-9	409	418	-2	187	-163	6	330	-323	
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11	474	-461	-18	209	-205	-10	579	549	1	263	238	
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2	165	176	-13	146	-131	13	405	361	14	212	225
-21	127	162	-12	229	-243	14	124	118	15	269	278
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-19	159	-164	-10	204	217	-4	194	-160	-10	-71	47
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-16	-65	-91	2	-69	-26	-3	90	-73	15	299	-301
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-13	239	-230	5	253	-245	0	-65	-3	-10	130	110
-12	293	-307	6	183	-168	1	404	-467	-9	378	-381
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-7	304	-314	11	-67	-63	6	165	-165	-3	225	-238
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-14	210	-219	9	-65	-68	21	234	-243	10	155	-210
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-3	139	-115	3	242	-234	11	243	230	-11	305	-264
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4	259	261		-2	250	218		-9	-72	63	-5	-69	116
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-21	183	159	H=	6,	K=	7	H=	6,	K=	-14	-3	109	143
-20	201	189		0	164	-181		1	110	-120	-2	-66	-73
-19	-63	22		-21	-68	65		2	179	152	-1	231	-233
-18	152	-151		-20	202	-200		2	277	271	H=	6,	K= -10
-17	-64	-37		-19	120	-107		4	-68	-45			
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-15	510	512		-17	270	251		6	-67	-13	1	308	-326
-14	90	-73		-16	295	297		7	-67	54	2	-65	-94
-13	150	-131		-15	271	-268		8	221	220	3	173	150
-12	132	-128		-14	315	-300		9	-69	-73	4	486	460
-11	286	302		-13	-67	-52		10	225	-223	5	172	-129
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-9	-62	147		-11	176	172				7	259	264	
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-7	112	143		-9	346	-337				9	93	-53	
-6	229	239		-8	299	305		0	221	205	10	174	-140
-5	-62	-40		-7	211	224		1	215	-14	11	336	-344
-4	335	-277		-6	102	-62		2	-66	10	12	-64	-30
-3	320	-245		-5	104	-115		3	217	-220	13	233	210
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-1	298	273		-3	307	277							

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
15	153	-131	5	305	326	-11	240	-251	-1	880	-857
-9	318	-306	6	90	94	-10	481	-501	H=	6	K= -2
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-7	208	199	8	603	-604	-8	295	283			
-6	136	129	9	174	-162	-7	712	-642	0	161	161
-5	-66	-43	10	541	528	-6	685	-689	1	555	-541
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-3	-65	15	12	382	-386	-4	569	520	3	342	342
-2	503	530	13	174	-178	-3	261	346	4	550	546
-1	227	240	14	-66	69	-2	309	-287	5	212	-214
			15	170	176	-1	321	-329	6	343	-336
									7	-65	58
H=	6,	K= -9	-13	287	280				8	-67	-60
			-12	-68	-13	H=	6,	K= -4	9	229	239
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1	205	191	-10	250	245	0	176	-192	11	253	-244
2	-61	60	-9	235	237	1	362	-336	12	-69	18
3	277	-310	-8	163	150	2	-54	-88	-18	177	-175
4	166	-138	-7	-66	-11	3	221	235	-17	258	-274
5	249	257	-6	354	-397	4	286	305	-16	220	239
6	130	115	-5	212	-226	5	210	-212	-15	380	371
7	129	-151	-4	133	131	6	408	-457	-14	291	-269
8	104	-104	-3	-59	22	7	92	46	-13	-64	-8
9	-64	-14	-2	330	-312	8	506	507	-12	-63	-127
10	305	291	-1	333	-300	9	108	-91	-11	432	447
11	309	321	H=	6,	K= -6	10	406	-407	-10	355	355
12	307	-311				11	-64	-86	-9	-55	-32
13	315	-306				12	-67	78			
14	190	162	0	-54	-48	13	232	235	-8	-54	-112
15	138	154	1	96	-155	-17	-70	-98	-7	-53	106
-10	-69	57	2	-55	-34	-16	118	126	-6	-92	-10
-9	155	149	3	403	414	-15	105	86	-5	257	-268
-8	-66	-34	4	95	75	-14	215	-211	-4	738	-695
-7	135	-127	5	424	-412	-13	246	-237	-3	1030	-966
-6	209	-208	6	-59	0	-12	159	-171	-2	-51	-11
-5	-65	-25	7	-60	-2	-11	-64	39	-1	385	374
-4	337	330	8	195	200	-10	399	412	H=	6,	K= -1
-3	-66	-70	9	-63	-36	-9	398	-375			
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-1	388	-384	11	163	-174	-7	507	546	0	288	300
			12	252	248	-6	527	586	1	1014	976
			13	261	264	-5	212	153	2	553	535
H=	6,	K= -8	14	-67	39	-4	238	-277	3	-59	-95
0	247	-232	-14	-71	-41	-3	-52	-53	4	-60	19
1	443	-451	-13	272	-272	-2	-62	81	5	231	206
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3	731	726	-11	483	510	H=	6,	K= -3	7	-65	25
4	-60	22	-10	106	119				8	408	-349
5	429	-402	-9	100	-103	0	244	270	9	177	-184
6	194	-182	-8	259	-258				10	122	117
7	-60	21	-7	95	-67	1	449	436	11	-71	108
8	272	285	-6	541	550	2	482	460	-19	97	-78
9	132	130	-5	-58	26	3	456	-454	-18	102	126
10	545	-571	-4	357	-387	4	776	-757	-17	245	242
11	-64	28	-3	141	-153	5	216	212	-16	105	-107
12	461	460	-2	640	642	6	779	747	-15	-65	-51
13	144	136	-1	617	440	7	349	-354	-14	148	169
14	159	-177				8	227	-224	-13	198	196
15	167	-196	H=	6,	K= -5	9	287	273	-12	308	301
-12	-70	-52				10	109	108	-11	404	-301
-11	230	231	0	155	119	11	218	212	-10	540	-530
-10	-69	-94	1	369	400	12	134	-95	-9	403	470
-9	269	-273	2	177	-183	13	424	-403	-3	913	864
-8	-65	-48	3	603	-659	-17	111	112	-7	-53	18
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-6	267	264	5	549	531	-15	251	-255	-5	-82	-40
-5	113	-164	6	282	301	-14	172	204	-4	467	521
-4	449	-460	7	-61	8	-13	130	160	-3	331	356
-3	141	-146	8	325	-377	-12	221	203	-2	402	-411
-2	354	352	9	149	149	-11	88	-43	-1	498	-483
-1	290	295	10	490	436	-10	215	-238	H=	7,	K= 0
			11	173	175	-9	464	425			
H=	6,	K= -7	12	117	-153	-8	193	173			
			13	170	-179	-7	014	-830	0	262	228
0	192	203	14	-68	-77	-6	173	-127	1	490	-487
1	657	696	-15	124	-133	-5	114	18	2	201	-205
2	155	146	-14	172	185	-4	310	238	3	201	169
3	481	-512	-13	355	367	-3	220	235	4	-65	-60
4	126	93	-12	121	-123	-2	705	-663	5	196	-195

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC	
6	-66	-59	-5	-61	36	-12	118	106	-17	-67	13	
7	190	170	-4	-62	-69	-11	278	-260	-16	-70	-75	
8	273	260	-3	100	-168	-10	140	-109	-15	319	-322	
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-18	164	-150	-1	230	219	-8	216	219	-13	255	248	
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-16	604	606	H=	7+	K=	3	-6	152	-141	-11	-69	-79
-15	247	247				-5	-67	-71	-10	-71	-44	
-14	-65	47	0	-67	53	-4	424	419	-9	232	249	
-13	425	-407	1	273	269	-3	428	433				
-12	-62	-21	2	-65	-40	-2	344	-348				
-11	145	145	3	246	-237	-1	204	-211	H=	7+	K= -14	
-10	-59	-66	4	224	-218							
-9	159	150	5	-70	69	H=	7+	K=	2	-71	110	
-8	421	-387	-21	-69	-64				3	216	198	
-7	197	191	-22	227	-240	0	-71	-11	4	-70	-71	
-6	390	416	-19	110	-83	-20	205	177	5	-70	-131	
-5	-56	-112	-18	115	118	-19	-67	74	6	114	-120	
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			-13	414	410	-14	95	62				
H=	7+	K=	1	-12	-63	-13	223	-230	0	215	214	
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1	286	272	-9	440	435	-10	106	133	2	177	-171	
2	-69	-152	-8	330	328	-9	-68	-103	3	279	-239	
3	235	-224	-7	164	146	-8	242	-246	4	-69	-22	
4	120	140	-6	439	-439	-7	103	120	5	127	134	
5	-64	81	-5	150	-148	-6	115	88	6	-67	68	
6	161	172	-4	364	354	-5	-68	84	7	142	-158	
7	-70	-95	-3	157	217	-4	402	-394	8	170	-159	
-20	183	-173	-2	-66	-7	-3	193	-196	9	191	187	
-19	104	84	-1	226	-215	-2	385	356	10	349	355	
-18	164	135				-1	211	180	11	-69	50	
-17	144	-157	H=	7+	K=	4			-2	194	-169	
-16	171	-155				H=	7+	K=	-1	-69	-19	
-15	370	-370	0	153	-142				H=	7+	K= -12	
-14	185	169	1	176	-159	-20	218	-202				
-13	309	307	2	-69	10	-19	-68	-10	0	276	-297	
-12	-63	-62	3	130	147	-18	426	421	1	127	-134	
-11	140	-146	-21	-63	45	-17	-68	-22	2	201	224	
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-9	168	166	-19	-65	40	-15	194	-199	4	-65	75	
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-3	166	-136	-13	247	-226	-9	107	122	10	275	-295	
-2	198	-191	-12	105	-73	-8	190	183	11	174	-170	
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H=	7+	K=	2	-9	415	-212	-5	-70	-70	-5	117	-116
0	114	-93	-8	245	-252	-4	375	368	-4	203	-180	
1	328	-328	-6	173	109	-2	235	-241	-2	201	247	
2	130	121	-5	-66	-90				-1	132	136	
3	183	190	-4	034	-810	H=	7+	K=	8	H=	7+	K= -11
4	-65	47	-3	213	-134							
5	-66	-2	-2	115	150	-19	-67	24	0	320	309	
6	207	-209	-1	395	612	-18	158	-148	1	166	167	
-20	200	274	H=	7+	K=	5	-17	105	2	319	-269	
-19	-64	25				-15	-68	-69	3	169	-105	
-18	-65	-42				-14	-68	-58	4	152	174	
-17	-65	-93	0	-68	-30	-13	301	-293	5	611	610	
-16	-65	8	1	-68	49	-12	-67	34	6	265	269	
-15	263	259	2	133	118	-11	197	174	7	520	-537	
-14	340	-347	-21	-59	-62	-10	190	176	8	266	-268	
-13	301	-272	-20	176	-187	-9	185	-182	9	-67	114	
-12	-62	47	-19	116	-27				10	-65	112	
-11	356	376	-18	172	207	-8	216	-202	11	-64	2	
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-9	114	-97	-16	237	-236	-6	232	261	13	267	-243	
-8	393	-375	-15	-64	-60	-5	119	113	-7	218	-170	
-7	231	212	-14	-65	-5				-6	-70	-50	
-6	423	411	-13	257	223	H=	7+	K=	9			

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC
-5	-66	0	-9	100	-144	5	323	323	-11	430	-408
-4	109	88	-8	-65	-63	6	234	246	-10	186	-207
-3	-68	75	-7	232	239	7	280	-298	-9	593	627
-2	297	-309	-6	238	273	8	319	-331	-8	207	204
-1	146	-142	-5	217	-241	9	-66	100	-7	-57	-110
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H=	7,	K= -10	-2	233	237	12	-69	-110	-4	719	646
0	210	-237	-1	162	150	-16	127	-125	-3	361	361
1	110	-86	H=	7,	K= -7	-15	121	-120	-2	1056	-1004
2	-54	47	H=	7,	K= -7	-14	194	194	-1	198	-186
3	219	219				-13	-65	27			
4	142	-138	0	171	186	-12	229	-235	H=	7,	K= -2
5	765	-782	1	554	556	-11	261	-275			
6	151	-120	2	257	256	-10	168	-160	0	119	-91
7	320	344	3	259	-295	-9	299	318	1	143	-160
8	127	131	4	-62	74	-8	-62	-20	2	142	-147
9	-62	-58	5	263	277	-7	294	-323	3	440	469
10	201	-193	6	-63	30	-6	99	73	4	427	422
11	-64	-27	7	200	-209	-5	173	170	5	292	-292
12	325	331	8	404	-401	-4	488	503	6	138	-156
13	222	227	9	-63	64	-3	347	340	7	-66	85
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-7	137	138	12	143	-137				10	207	-217
-6	130	89	13	96	-28	H=	7,	K= -4	-18	340	-341
-5	-66	-87	-13	309	311				-17	164	-190
-4	267	-268	-12	117	-137	0	434	-407	-16	486	495
-3	-65	51	-11	94	-116	1	294	-273	-15	365	338
-2	343	349	-10	-65	46	2	-57	-34	-14	-65	-40
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1	-64	92	-4	518	520	6	269	266	-8	641	-594
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5	400	399				12	145	163	-4	226	-254
6	204	216	H=	7,	K= -6	-17	133	-117	-3	565	-567
7	141	-139				-16	116	85	-2	357	374
8	116	-105	0	285	-286	-15	201	190	-1	593	595
9	-63	92	1	607	-626	-14	241	-251	H=	7,	K= -1
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12	385	-402	4	-60	-46	-11	591	600	0	116	-119
13	132	-145	5	251	-248	-10	347	352	1	455	459
-10	-67	10	6	-63	-55	-9	293	-247	2	334	315
-9	145	167	7	144	122	-8	113	-119	3	520	-533
-8	-64	13	8	313	315	-7	172	125	4	129	-104
-7	228	-223	9	140	-153	-6	318	317	5	390	397
-6	290	-308	10	260	-277	-5	152	-122	6	152	145
-5	108	123	11	-64	67	-4	518	-535	7	316	-327
-4	272	272	12	194	180	-3	377	-394	8	358	-377
-3	99	-96	13	-68	92	-2	88	128	9	-67	-21
-2	408	-402	-14	204	-207	-1	261	266	-19	-67	42
-1	-63	-27	-13	212	-200	H=	7,	K= -3	-18	208	219
H=	7,	K= -8	-12	188	170				-17	-66	-15
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1	216	-235	-8	245	-251	2	-57	-10	-13	259	274
2	106	-73	-7	240	217	3	34	-337	-12	-63	-58
3	193	187	-6	273	272	4	271	-264	-11	661	-641
4	116	116	-5	356	-336	5	192	179	-10	-60	6
5	123	-84	-4	443	-423	6	133	116	-9	802	808
6	138	-141	-3	303	-314	7	139	-113	-8	247	234
7	94	87	-2	159	-121	8	-64	-5	-7	174	-198
8	347	379	-1	246	250	9	110	-113	-6	315	-326
9	-65	-138				10	270	264	-5	171	-170
10	236	-243	H=	7,	K= -5	11	222	213	-4	345	285
11	-63	42				-17	134	101	-3	107	103
12	-66	45	0	314	343	-16	362	-350	-2	454	-468
13	-66	69	1	433	424	-15	264	-276	-1	221	-216
-12	-70	90	2	322	-333	-14	374	374			
-11	273	298	3	238	-231	-13	307	337			
-10	-66	-23	4	195	136	-12	263	262			

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
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1	229	-237	-3	-67	-61	-2	118	-100	9	-172	-148
2	-66	90	-2	359	349	-1	-67	-10	10	150	-166
3	291	299	-1	383	370	H=	8, K= 6		11	-68	78
4	-64	5				-19	-68	-29	-2	205	-212
5	178	-186	H=	8, K= 3		-18	253	-266	-1	172	60
6	260	-251				-17	156	165	H=	8, K= -11	
-19	119	-115	0	203	211	-16	156	165	0	188	186
-18	170	-147	1	137	139	-15	123	152	1	-64	-33
-17	-64	-11	2	-69	-65	-14	-64	16	2	156	-195
-16	113	117	3	-70	-119	-13	270	-279	3	152	-192
-15	367	371	-20	203	-205	-12	-65	-58	4	139	125
-14	-66	-46	-19	-67	6	-11	190	167	5	306	309
-13	432	-415	-18	208	187	-10	-66	0	6	-64	-48
-12	-65	-57	-17	-66	58	-9	130	-138	7	272	-263
-11	151	138	-16	196	-185	-8	99	-89	8	132	-120
-10	108	-64	-15	-66	-103	-7	-65	60	9	113	92
-9	329	-352	-14	192	185	-6	220	226	10	124	118
-8	286	-302	-13	174	162	-5	120	138	-7	-66	-49
-7	-62	63	-12	-66	-82	-4	177	-177	-7	151	-166
-6	363	355	-11	295	-271	-3	-69	1	-5	-68	61
-5	-60	29	-10	217	-201	H=	8, K= 7		-6	-69	-29
-4	461	-482	-9	284	287				-5	-68	61
-3	111	-127	-8	270	249	H=	8, K= 2		-4	235	232
-2	261	256	-7	398	-409	-18	199	204	-3	-67	59
-1	336	342	-6	322	-328	-17	-69	-43	-2	-68	-147
H=	8, K= 1		-5	284	282	-16	241	-212	-1	-66	7
			-4	157	144	-15	271	-272			
			-3	-66	38	-14	-66	12	H=	8, K= -10	
0	265	256	-2	371	-370	-13	295	293	0	216	-228
1	290	286	-1	327	-323	-12	-67	10	1	137	-111
2	199	-196	H=	8, K= 4		-11	241	-222	2	232	-229
3	251	-262				-10	-67	-69	3	170	192
4	-68	-76	H=	8, K= 4		-9	-67	99	4	106	-116
5	209	203	0	-63	-78	-8	210	222	5	113	-113
-19	133	132	1	-69	-1	-7	-70	-64	6	151	142
-18	168	166	-20	212	197	-6	259	-253	7	210	228
-17	219	205	-19	-67	48	H=	8, K= 8		8	133	116
-16	163	-199	-18	271	-262	H=	8, K= 8		9	108	-95
-15	345	-242	-17	-65	-55				10	166	-165
-14	364	344	-16	113	76	-15	217	240	11	-68	43
-13	197	187	-15	-65	-4	-14	131	-140	12	374	380
-12	124	-123	-14	-64	-72	-13	202	-264	-3	-69	73
-11	-63	-16	-13	124	-139	-12	108	141	-6	-65	41
-10	88	24	-12	-65	9	-11	142	122	-5	-66	-3
-9	218	222	-11	342	346	-10	-70	21	-4	169	-166
-8	268	256	-10	-66	90	H=	8, K= -14		-3	-63	27
-7	128	-141	-9	269	-259	H=	8, K= -14		-2	220	227
-6	96	-98	-8	-64	-46				-1	129	-121
-5	311	281	-7	234	228	4	121	-101			
-4	436	423	-6	-65	64	5	159	-157	H=	8, K= -9	
-3	267	241	-5	-66	-59	H=	8, K= -13		0	280	264
-2	189	-197	-4	132	-120	H=	8, K= -13		1	-67	130
-1	423	-427	-3	129	-134				2	263	272
H=	8, K= 2		-2	343	364	0			3	-67	-218
			-1	-67	11	1	-69	23	4	-67	-60
0	306	-297	H=	8, K= 5		2	144	-177	5	484	471
1	317	-328				3	-65	8	3	374	363
2	-66	80	-20	174	-184	4	-67	72	6	114	-127
3	201	188	-19	-65	-19	5	237	230	7	202	-192
4	101	93	-18	273	272	6	-68	-48	8	-62	-51
-20	-71	126	-17	-66	74	7	226	-216	9	-66	59
-19	-66	-63	-16	-65	-44	8	-70	-80	10	174	168
-18	228	-231	-15	185	-179	9	296	272	11	145	-134
-17	255	-252	-14	-64	30	-1	-69	59	12	321	-325
-16	225	243	-13	151	153	H=	8, K= -12		-10	-63	52
-15	-63	17	-12	-66	22				-9	-64	143
-14	294	-271	-11	213	-232	0	134	-138	-8	-66	-5
-13	271	-267	-10	-64	18	1	-66	-10	-7	-66	-20
-12	-65	89	-9	266	255	2	-65	31	-6	-64	-2
-11	109	53	-8	116	140	3	271	251	-5	172	132
-10	218	223	-7	155	-170	4	-67	-13	-4	257	263
-9	157	-185	-6	172	-187	5	345	-307	-3	213	-187
-8	262	-251	-5	-67	-72	6			-4		

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
-3	209	-211	-10	-63	52	5	444	439	-5	-59	-40
-2	278	-268	-9	256	-253	6	-66	77	-4	244	223
-1	-66	67	-8	225	-193	7	131	-124	-3	-60	0
H=	8,	K= -8	-6	134	134	8	141	-144	-2	305	-294
			-5	215	-243	9	-68	104	-1	192	-186
0	309	-314	-4	299	-229	-17	312	-303	H=	9,	K= 0
1	277	-278	-3	-63	-71	-15	-66	79			
2	570	578	-2	239	209	-14	242	236	0	172	-185
3	274	264	-1	89	133	-13	-64	28	1	199	-198
4	216	-244				-12	312	-202	2	102	105
5	357	-346	H=	8,	K= -5	-11	428	-409	3	299	315
6	-64	-26				-10	-64	-73	4	-68	-18
7	277	280	0	140	172	-9	563	549	-18	-67	-33
8	238	247	1	-61	25	-8	-61	43	-17	-66	-6
9	-65	-77	2	96	-95	-7	253	-268	-16	169	155
10	277	-276	3	403	-407	-6	114	-121	-15	-66	133
11	-67	15	4	-67	-139	-5	178	196	-14	413	-412
12	220	209	5	323	336	-4	727	719	-13	368	-343
-12	155	167	6	-64	28	-3	255	-262	-12	254	221
-11	155	168	7	333	-338	-2	875	-887	-11	227	220
-10	-65	0	8	247	-238	-1	-60	20	-10	96	-113
-9	-68	-122	9	177	190				-9	214	-201
-8	-64	-62	10	-67	89	H=	3,	K= -2	-8	-65	-53
-7	-63	30	11	99	17				-7	458	463
-6	-65	101	-15	119	-91	0	-63	73	-6	303	295
-5	151	-161	-14	212	201	1	170	-186	-5	373	-364
-4	313	-321	-13	140	128	2	95	-34	-4	-66	57
-3	-64	-33	-12	-64	-19	3	510	507	-3	349	334
-2	184	180	-11	-65	-19	4	180	-178	-2	296	293
-1	146	139	-10	-66	59	5	413	-413	-1	-66	29
H=	8,	K= -7	-8	-64	11	6	106	-51			
			-7	487	-464	7	184	180	H=	9,	K= 1
0	434	449	-6	191	-203	-18	224	-203	0	246	247
1	478	485	-5	376	392	-17	-69	79	1	225	189
2	514	-511	-4	476	494	-16	158	156	2	179	-176
3	294	-282	-3	118	99	-15	-54	-29	3	257	-254
4	137	-125	-2	-60	7	-14	402	-412	-19	-68	34
5	-65	102	-1	279	-271	-13	268	-302	-18	132	134
6	126	130				-12	274	249	-17	-67	34
7	271	-292	H=	8,	K= -4	-11	642	658	-16	418	-442
8	444	-440				-10	-63	9	-15	-64	-24
9	-66	53	0	398	-402	-9	692	-704	-14	172	167
10	423	429	1	222	-217	-8	-60	-34	-13	-64	-43
11	-67	-12	2	-62	6	-7	116	123	-12	201	-194
-13	111	111	3	505	505	-6	343	323	-11	154	-165
-12	154	-151	4	-65	68	-5	-59	-54	-10	103	37
-11	171	-167	5	261	-267	-4	826	-814	-9	367	364
-10	-63	36	6	-65	24	-3	191	-232	-8	269	266
-9	170	182	7	314	315	-2	569	594	-7	412	-438
-8	192	185	8	150	134	-1	-61	0	-6	186	-175
-7	347	-363	9	215	-216				-5	-67	32
-6	467	-467	10	-68	-97	H=	8,	K= -1	-4	-68	-81
-5	212	234	-16	288	292				-3	-66	61
-4	-64	59	-15	-56	31	0	-63	-18	-2	275	-280
-3	204	-195	-14	-64	-81	1	342	397	-1	-65	6
-2	145	-149	-13	-64	-25	2	809	-279			
-1	160	-165	-12	117	100	3	263	-276	H=	9,	K= 2
H=	8,	K= -6	-11	144	114	4	158	180			
			-10	-64	44	5	140	157	0	360	-361
0	360	-360	-9	461	-447	6	-68	121	1	232	-226
1	-62	-67	-7	545	553	7	140	-114	-19	104	-4
2	173	163	-6	411	390	-19	-71	77	-18	274	-294
3	297	301	-5	-59	-44	-18	126	143	-16	264	241
4	266	268	-4	359	-357	-17	-64	2	-15	104	91
5	132	-139	-3	161	-148	-16	136	-126	-14	166	-164
6	198	-186	-2	317	313	-15	167	-134	-13	150	-131
7	289	276	-1	174	150	-14	-67	439	-12	114	124
8	284	281				-13	253	264	-11	168	159
9	-67	-61	H=	8,	K= -3	-12	117	-115	-10	-66	67
10	242	-238				-11	323	-319	-9	459	-455
11	-66	2	0	240	238	-10	434	447	-8	-66	-65
-14	195	-208	1	156	136	-9	604	607	-7	452	459
-13	-69	-105	2	-63	39	-8	238	230	-6	226	227
-12	109	101	3	325	-325	-7	174	-180	-5	-65	25
-11	192	163	4	172	-165	-6	494	-484	-6	-66	-89

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
-3	244	-257	-7	-71	84	8	145	-146	-13	-67	-34
-2	253	246	H=	9,	K= -13	9	136	131	-12	134	159
-1	-66	33				10	205	214	-11	-64	61
						-10	112	111	-10	-64	0
H=	9,	K= 3	0	156	201	-9	126	151	-9	126	-146
			1	-69	-51	-8	-77	-20	-8	-69	-20
0	238	235	2	-66	-91	-7	175	-173	-7	362	612
-19	-69	52	3	203	-176	-6	119	-131	-6	117	76
-18	248	259	4	150	159	-5	160	127	-5	-66	-151
-17	-66	-42	5	191	153	-4	145	162	-4	165	-173
-16	169	-187	6	-68	-18	-3	222	-227	-3	-64	-37
-15	150	-146				-2	-64	50	-2	315	320
-14	250	213	H=	9,	K= -12	-1	-64	-31	-1	-65	-68
-13	248	266	0	156	-159	H=	9,	K= -8	H=	9,	K= -5
-12	-64	-83	1	-65	-14						
-11	233	-209	2	317	301	0	370	-384	0	372	381
-10	102	-69	3	183	169	1	-65	-76	1	251	249
-9	162	206	4	314	-294	2	165	158	2	324	-317
-8	-63	52	5	205	-231	3	-65	7	3	345	-326
-7	203	-226	6	103	81	4	160	-142	4	-64	-34
-6	254	-258	7	164	161	5	184	-175	5	221	216
-5	109	-89	8	-67	13	6	-62	-17	6	91	87
-4	352	353	-3	140	155	7	376	400	7	200	-201
-3	219	224	-2	245	245	8	202	203	8	144	-122
-2	277	-285	-1	-68	-106	9	215	-219	9	122	116
-1	100	-76				10	158	-186	-15	-67	73
H=	9,	K= 4	H=	9,	K= -11	-11	145	126	-14	121	133
						-10	-67	-78	-13	-65	60
-19	-68	-63	0	126	126	-9	185	-217	-12	190	-168
-18	264	-273	1	159	-131	-8	-65	-24	-11	136	-141
-17	-63	7	2	214	-214	-7	226	232	-10	136	169
-16	177	165	3	-63	-15	-6	165	177	-9	392	373
-15	152	140	4	220	207	-5	281	-269	-8	-66	-9
-14	249	-245	5	242	222	-4	-65	-111	-7	334	-350
-13	242	-262	6	-68	-109	-3	126	128	-6	161	150
-12	-64	65	7	220	-210	-2	139	123	-5	117	65
-11	229	266	8	-66	-54	-1	113	79	-1	195	196
-10	192	175	9	115	120	H=	9,	K= -7	-3	237	256
-9	-68	-121	-6	-67	-12	H=	9,	K= -7	-2	524	-532
-8	-65	-8	-5	-65	75				-1	122	-119
-7	173	178	-4	197	189	0	520	516	H=	9,	K= -4
-6	196	206	-3	114	-148	1	202	-201			
-5	-66	16	-2	239	-201	2	312	-300	0	221	-209
-4	383	-395	-1	144	124	3	124	-120	1	305	-303
-3	102	-88				4	170	174	1	214	225
-2	171	172	H=	9,	K= -10	5	285	319	2	267	265
H=	9,	K= 5	0	130	66	6	-64	12	4	-66	117
			1	274	-273	7	319	-308	5	308	-295
-18	276	280	2	214	203	8	113	-120	6	161	-171
-17	-68	27	3	424	420	-13	-68	52	7	270	234
-16	165	-155	4	-64	18	-12	153	-146	8	101	153
-15	118	-91	5	274	-267	-11	-65	-62	-16	-63	-8
-14	162	157	6	-68	66	-10	162	118	-15	-65	-35
-13	278	288	7	252	251	-9	184	174	-14	160	-186
-12	-67	-60	8	-64	45	-8	-64	-19	-13	214	-210
-11	375	-373	9	127	-136	-7	407	-405	-12	168	209
-10	-66	-4	10	209	-191	-6	-66	-81	-11	-65	67
-9	215	214	-8	-68	-3	-5	338	334	-10	185	-205
-8	-68	62	-7	-67	90	-4	132	100	-9	458	-476
-7	137	-123	-6	145	143	-3	-65	-69	-8	-66	43
-6	251	-249	-5	159	-138	-2	292	-200	-7	375	375
-5	-60	26	-4	323	-379	-1	-64	-40	-6	101	-122
-4	315	301	-3	181	170	H=	9,	K= -6	-5	-63	-73
H=	9,	K= 6	-2	165	194				-4	350	-358
			-1	-60	-133	0	376	-380	-3	-64	-56
-17	-70	84	H=	9,	K= -9	1	-65	64	-2	420	411
-16	-68	18				2	477	409	-1	201	-199
-15	-67	49	0	101	92	3	344	304	H=	9,	K= -3
-14	160	-155	1	233	235	4	-66	-50			
-13	232	-247	2	160	-201	5	213	-230	C	100	135
-12	113	111	3	263	-278	6	-64	-10	1	219	204
-11	248	252	4	-64	-20	7	144	119	2	158	-143
-10	107	-61	5	199	181	8	-68	66	3	500	-500
-9	178	-149	6	-62	-22	9	194	-195	4	162	153
-8	185	-165	7	205	-195	-14	227	-231	5	563	560

L	FO	FC	L	FO	FC	L	FO	FC	L	FO	FC
6	-66	67	2	209	224	-16	233	224	-4	-66	43
7	179	-195	-17	-66	-29	-15	-65	28	-3	184	-211
-17	174	-187	-16	231	249	-14	-69	-125	-2	182	-172
-16	-67	-77	-15	-66	-34	-13	-68	-112	-1	-64	-58
-15	-63	-52	-14	393	-380	-12	100	119	H=	10+	K= -8
-14	152	159	-13	-66	-69	-11	267	310	0	362	-360
-13	127	107	-12	196	189	-10	-67	-47	1	-64	-70
-12	190	-196	-11	160	195	-9	223	-237	2	328	341
-11	-65	-64	-10	-63	22	-8	-67	-42	3	256	254
-10	254	250	-9	159	-182	-7	-68	51	4	-65	-30
-9	451	487	-8	-65	-77	-6	173	218	5	298	-291
-8	188	-172	-7	397	378	H=	10,	K= 5	6	-66	-18
-7	206	-196	-6	144	128	H=	10,	K= -12	7	310	311
-6	155	-164	-5	236	-241	0	138	-105	-10	159	-159
-5	244	226	-4	293	-265	1	-70	100	-9	145	-151
-4	662	676	-3	-65	-55	2	226	228	-8	160	138
-3	454	-472	-2	331	336	3	186	197	-7	355	362
-2	535	-526	-1	-67	5	-12	209	-201	-6	-66	-30
-1	-64	16	H=	10+	K= 1	-11	241	-231	-5	272	-266
H=	9,	K= -2	0	269	274	-10	-68	66	-4	154	-181
0	245	-240	-17	-67	-72	H=	10,	K= -12	-3	-64	15
1	218	-227	-16	176	-131	0	138	-105	-2	253	249
2	391	303	-15	262	286	1	-70	100	-1	-66	-103
3	269	230	-14	245	252	2	226	228	H=	10,	K= -7
4	475	-457	-13	59	104	3	-69	-10	0	335	360
5	558	-378	-12	212	-225	4	207	-199	1	262	261
6	-69	-38	-11	312	-297	-1	-70	-108	2	194	-190
-17	-69	120	-10	-62	-33	H=	10,	K= -11	3	241	-259
-16	136	106	-9	403	414	H=	10,	K= -11	-1	-66	23
-15	-64	67	-8	-65	-6	0	-68	79	5	319	319
-14	131	-135	-7	445	-431	1	-69	9	6	113	67
-13	209	199	-6	-65	-21	2	123	-150	7	323	-310
-12	391	343	-5	230	226	3	184	-174	-12	193	-130
-11	109	130	-4	357	349	4	238	219	-11	-66	41
-10	193	-163	-3	-67	-24	5	165	167	-10	166	145
-9	264	-309	-2	271	-273	6	-70	-107	-9	216	211
-8	-64	65	-1	-67	-28	-5	122	124	-8	-67	-83
-7	233	226	H=	10,	K= 2	-4	-69	89	-7	426	-441
-6	339	323	H=	10,	K= -2	-3	136	-148	-6	-67	-103
-5	411	-430	-17	181	184	-2	142	-137	-5	198	203
-4	592	-617	-16	172	178	-1	121	80	-4	94	118
-3	502	497	-15	217	-206	H=	10,	K= -10	-3	-64	-8
-2	500	493	-14	266	-306	H=	10,	K= -10	-2	308	-310
-1	210	217	-13	97	-89	H=	9,	K= -1	-1	-67	136
H=	9,	K= -1	-12	218	247	0	279	-282	H=	10,	K= -6
0	405	384	-10	158	-176	1	169	-160	0	238	-229
1	125	112	-9	420	-404	2	296	323	1	-63	-23
2	158	-147	-8	-68	115	3	170	187	2	115	129
3	202	-230	-7	359	333	4	262	-277	3	-63	50
4	123	104	-6	-66	58	5	-65	-59	4	-67	-78
5	363	350	-5	99	-95	6	-67	88	5	302	-311
-18	134	110	-4	318	-308	-7	113	138	6	-63	-70
-17	-67	-66	-3	-68	80	-6	-68	28	7	274	274
-16	-64	-13	-2	190	104	-5	170	-199	-12	-96	55
-15	117	134	H=	10,	K= 3	-4	-67	-72	-12	-66	51
-14	121	140	H=	10,	K= -3	-3	306	318	-11	125	125
-13	-65	32	-17	176	-174	-2	-67	82	-10	-65	-34
-12	179	-130	-16	244	-243	-1	-65	-48	-9	267	-265
-11	243	-241	-15	-66	61	H=	10,	K= -9	-8	-65	38
-10	-66	102	-14	203	209	0	268	288	-7	255	294
-9	264	264	-13	-65	75	1	-64	6	-6	-64	-45
-8	-63	-33	-12	265	-250	2	300	-303	-5	179	-201
-7	318	-313	-11	202	-246	3	156	-167	-4	163	-164
-6	164	-144	-10	-65	53	4	260	250	-3	180	174
-5	427	417	-9	-65	53	5	103	122	-2	504	522
-4	310	307	-8	343	356	6	-67	83	-1	-66	-70
-3	498	-490	-7	147	145	7	127	-100	H=	10,	K= -5
-2	402	-296	-6	232	-226	-8	159	160	0	220	236
-1	227	-224	-5	134	-149	-9	-68	4	1	138	-159
H=	10,	K= 0	-4	251	248	-7	146	-108	2	187	-200
0	295	-205	-3	-70	-15	-6	-68	6	0	220	236
1	214	-202	H=	10,	K= 4	-5	342	342	1	138	-159

L	F0	FC	L	F0	FC	L	F0	FC	L	F0	FC	
3	110	-139	-14	-67	-67	-10	189	-116	-2	103	-97	
4	170	149	-13	-65	-50	-9	140	-119	-1	-35	256	
5	204	313	-12	172	102	-8	104	105	H=	114	-12	
6	-67	-70	-11	-63	17	-7	202	198	-6			
-14	101	79	-10	161	-205	-6	-68	61	0	396	-408	
-13	-66	55	-9	-66	-64	H=	114	K= -3	1	-60	-19	
-12	208	-201	-3	95	35				2	142	121	
-11	144	-160	-7	227	230	-13	175	169	3	227	236	
-10	-65	69	-6	134	141	-12	192	-119	4	-70	-100	
-9	138	159	-5	120	-131	-11	233	-216	-12	131	140	
-8	-65	-104	-4	106	-123	-10	152	153	-11	154	173	
-7	320	-323	-3	111	108	-9	177	158	-10	202	-149	
-6	-64	41	-2	97	106				-9	104	-146	
-5	249	262	-1	-64	-48	H=	114	K= -11	-8	165	160	
-4	273	262					0	168	184	-7	198	212
-3	371	-290	H=	104	K= -1				-6	127	138	
-2	556	-586				0	245	242	-5	-65	-91	
-1	186	172	0	325	337	H=	114	K= -10	-4	135	-122	
H=	104	K= -4	1	245	242				-3	-67	90	
0	336	-324	-17	-68	-16	0	267	-269	-2	136	146	
1	-64	53	-16	221	-208	1	-67	-31	-1	245	-244	
2	195	196	-15	267	-282	2	320	232	H=	114	K= -5	
3	293	298	-14	187	176	3	-69	70				
4	171	-165	-13	148	164	-5	293	-263	0	284	276	
5	358	-374	-12	295	-206	-4	-71	-2	1	134	135	
6	152	146	-11	-63	-64	-3	340	311	2	-69	-84	
-15	-68	-30	-10	146	140	-2	148	147	3	195	-183	
-14	-67	-123	-9	128	111	-1	124	-141	-13	-67	22	
-13	-66	-12	-8	-64	-26	H=	114	K= -9	-12	193	-202	
-12	281	286	-7	325	-326				-11	125	-116	
-11	133	124	-6	297	-301	0	263	260	-10	249	250	
-10	94	-92	-5	233	220	1	-66	20	-9	141	143	
-9	216	-235	-4	105	110	2	268	-304	-8	-62	-20	
-8	156	130	-3	102	121	3	-66	25	-7	225	-221	
-7	260	249	-2	281	-267	4	220	218	-6	-64	-28	
-6	147	-114	-1	91	-4	5	255	-284	-5	93	116	
-5	109	20				-7	255	-284	-4	193	212	
-4	-66	65	H=	114	K= 0	-6	-69	106	-3	-64	-23	
-3	909	517	-5	104	-86	-5	199	192	-2	326	-345	
-2	-66	69	-14	203	-185	-4	284	-255	-1	123	97	
-1	320	-310	-13	-66	54	-1	140	143	H=	114	K= -4	
H=	104	K= -3	-12	169	175							
0	274	273	-10	117	-127	H=	114	K= -8	0	-65	-49	
1	116	114	-9	255	-276	0	287	-287	1	-67	9	
2	309	-315	-8	-65	17	1	-68	102	2	159	162	
3	279	-273	-7	277	275	2	203	213	-14	154	-171	
4	360	374	-6	-66	9+	3	-65	-35	-13	-69	-60	
5	253	274	-5	127	-112	4	101	-152	-12	243	241	
-16	-66	-35	-4	100	-140	-9	144	-133	-11	186	177	
-15	-66	35	-3	130	129	-8	171	190	-10	130	-119	
-14	178	187	-2	131	150	-7	123	116	-9	-63	-59	
-13	-65	44				-6	155	-194	-8	-64	3	
-12	345	-230	H=	114	K= 1	-5	152	-147	-7	111	161	
-11	-65	-84	-4	119	147	-4	119	117	-6	115	94	
-10	257	246	-5	-66	28	-3	307	302	-5	140	-161	
-9	198	194	-4	-67	74	-2	101	117	-4	205	-200	
-8	-64	14	-3	-66	-29	-1	214	-204	-3	-65	-61	
-7	-65	-70	-2	115	-152	H=	114	K= -7	-2	270	264	
-6	-64	41	-1	263	-254				-1	111	-52	
-5	238	213	-10	220	220	H=	114	K= -3				
-4	-66	-44	-9	257	239	0	353	350	H=	114	K= -3	
-3	502	-435	-8	-67	-55	1	143	-147	0	-68	-81	
-2	110	-125	-7	151	-162	2	222	-233	1	-67	-31	
-1	212	192	-6	-67	-55	3	159	-127	2	185	-173	
H=	104	K= -2	-5	132	146	4	-70	81	-14	176	167	
0	186	-201	H=	114	K= 2	-11	-69	-91	-13	-68	61	
1	133	-101	-4	112	144	-10	143	138	-12	307	-307	
2	262	295	-15	134	-136	-9	216	203	-11	105	-100	
3	-69	83	-14	100	-114	-8	216	-216	-10	-66	115	
4	187	-205	-13	110	-113	-6	121	90	-9	-66	61	
-16	142	112	-12	169	143	-4	-65	-11	-7	269	-305	
-15	-66	94	-11	243	222	-3	119	-143	-6	100	78	

L	FO	FC									
-5	430	435	-6	-66	66	0	118	-88	H=	12,	K= -3
-4	124	155	-5	157	170	-7	-69	-80			
-3	252	-246	-4	-67	99	-5	107	103	-12	255	-249
-2	111	-121	-3	-68	-119	-7	-68	88	-11	-67	42
-1	210	181	-2	222	-204	-6	203	-205	-10	141	120
			-1	-68	53	-5	346	-371	-9	138	150
H=	11,	K= -2				-4	-67	65	-8	-67	-34
0	160	-174	H=	12,	K= 0	-3	305	309	-7	222	-249
1	-69	-65	-11	114	114	-2	180	145	-6	110	84
-15	-68	-40	-10	207	-213	-1	197	-216	-5	223	225
-14	233	-248	-9	236	-208	H=	12,	K= -5	-4	-67	23
-13	-67	-22	-8	101	102				-3	123	-151
-12	285	303				-11	-67	-38	-2	167	-148
-11	-66	-9	H=	12,	K= -8	-10	211	218	H=	12,	K= -2
-10	-64	-32				-9	-66	15			
-9	259	-254	0	172	-199	-8	142	-137			
-8	-65	-78	-6	-68	-25	-7	106	-66	-12	-70	132
-7	271	274	-5	149	-155	-6	142	120	-11	-65	60
-6	-67	-55	-4	120	-138	-5	293	267	-10	114	-82
-5	402	-409	-3	-68	98	-4	-69	60	-9	183	-201
-4	-66	-62	-2	160	143	-3	208	-233	-8	103	116
-3	199	194	-1	194	-181	-2	117	-157	-7	270	302
-2	-66	35				-1	217	229	-6	126	-81
-1	-63	7	H=	12,	K= -7	H=	12,	K= -4	-5	210	-219
H=	11,	K= -1	0	146	122				-4	116	-101
0	355	352	-8	155	-143	-11	-69	-25	H=	12,	K= -1
-15	152	136	-7	130	-134	-10	226	-235			
-14	377	375	-6	-69	60	-9	-66	-75	-12	165	-152
-13	-70	-115	-5	264	270	-8	-65	26	-11	97	-103
-12	267	-288	-4	119	101	-7	150	176	-10	193	185
-11	-66	-68	-3	210	-236	-6	-67	42	-9	201	220
-10	-66	94	-2	132	-119	-5	153	-158	-8	127	-129
-9	298	301	-1	177	191	-4	103	-65	-7	213	-214
-8	-64	-14	H=	12,	K= -6	-3	177	147	-6	-67	46
-7	301	-288				-2	111	141			
						-1	150	-144			

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BIOGRAPHICAL SKETCH

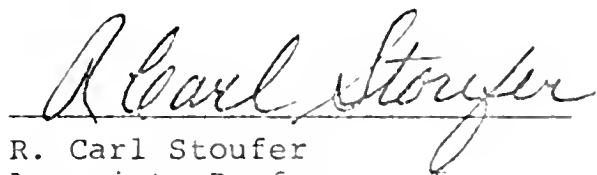
Douglas Allen Sullivan was born November 9, 1945, in Huntington, West Virginia. In May, 1963, he was graduated from Vinson High School, Huntington, West Virginia. He received the degree of Bachelor of Science in Chemistry from Marshall University in May, 1967. After studying at the University of Florida from September, 1967, to August, 1968, Mr. Sullivan taught chemistry, physics, physical science, and mathematics for the Wayne County (West Virginia) Board of Education. He then returned to the University of Florida in September, 1972, and received a Master of Science in Teaching degree majoring in chemistry in December, 1974. He is a member of the American Chemical Society. Mr. Sullivan is married to the former Jeanie Delaine Puckett of Titusville, Florida. They have a three-year-old son, David O'Donald Sullivan.

I certify that I have read this study and that in my opinion it conforms to acceptable standards of scholarly presentation and is fully adequate, in scope and quality, as a dissertation for the degree of Doctor of Philosophy.



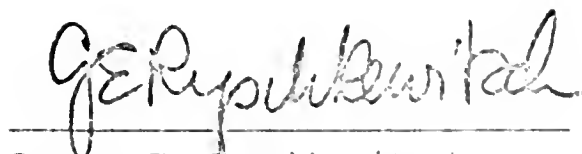
Gus J. Palenik, Chairman
Professor of Chemistry

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R. Carl Stoufer
Associate Professor of
Chemistry

I certify that I have read this study and that in my opinion it conforms to acceptable standards of scholarly presentation and is fully adequate, in scope and quality, as a dissertation for the degree of Doctor of Philosophy.



George E. Ryschkewitsch
Professor of Chemistry

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Associate Professor of
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I certify that I have read this study and that in my opinion it conforms to acceptable standards of scholarly presentation and is fully adequate, in scope and quality, as a dissertation for the degree of Doctor of Philosophy.

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Professor of Education

This dissertation was submitted to the Graduate Faculty of the Department of Chemistry in the College of Arts and Sciences and to the Graduate Council, and was accepted as partial fulfillment of the requirements for the degree of Doctor of Philosophy.

December, 1975

Dean, Graduate School

