Seven-Coordination

Seven-Coordination and \textquoteleft even-coordinate\textquoteright complexes are isolable and well-defined.

Seven-Coordination. A Molecular Orbital Exploration of Structure, Stereochemistry, and Reaction Dynamics

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A systematic molecular orbital analysis of seven-coordinate molecules is presented, with an emphasis on the basic electronic structure, substituent site preferences, and pathways for interconversion of various geometries. Conformations studied in detail include the pentagonal bipyramid (PB), capped octahedron (CO), and capped trigonal prism (CTP). With respect to \( o \) or \( \sigma \) electronegativity effects the following conclusions are reached for the optimum site of substitution in a \( d^4 \) complex by a better \( \sigma \) donor: PB equatorial (eq), CO capping site (c), CTP edge (e). For \( \pi \) effects the following preferences are deduced for \( d^4 \) systems substituted by an acceptor and \( \sigma \) by a donor (\( > \) means preferred): PB ax, eq, eq, eq, eq, CO (cf = capping face, uf, uc, uf, uc, uf, u...
structure in seven-coordination although minor polyhedral variants of lower symmetry are useful for an accurate assessment of the more subtle structural details.

A constructive generation of the idealized seven-coordinate structures is the addition of a vertex (ligand) to the regular octahedron, a formal process analogous to the initial phase in an associative substitution reaction of a six-coordinate complex. For example, vertex addition along an octahedral edge, 1, accompanied by a minor motion of four vertices coplanar with the new vertex, generates the $D_{5h}$ pentagonal bipyramid 2.

![Figure 2](image_url)

Figure 2. The two most common polyhedra in eight-coordination. On the left is the $D_{4d}$ Archimedean square antiprism and on the right the $D_{2d}$ dodecahedron. All vertices are equivalent in the former polyhedron whereas there are two types of vertices (A and B), distinguished by symmetry and connectivity, in the dodecahedron.

which has been discussed as one possible way-point in an associative $ML_6$ substitution reaction—the other possible transition state in a concerted reaction is the pentagonal bipyramid.

Alternatively, the idealized seven-coordinate forms 2-4 may be conceptually generated by vertex removal from the favored or most common eight-coordinate geometries (see Figure 2). Although this approach is not readily envisaged nor readily illustrated by drawings (a careful inspection of models is required), there is a cogent reason for considering such a generation scheme. Seven-coordinate intermediates will be much more common in substitution reactions of eight-coordinate than of six-coordinate complexes. Starting from the $D_{2d}$ dodecahedral eight-coordinate form, there are two possible degradative processes because of the two different types of vertices (Figure 3). Vertex elision at an A site directly yields the $C_{3v}$ capped octahedron, 3, although with minor displacements of the other vertices either the $D_{5h}$ (2) or $C_{3v}$ (4) forms may be generated. Removal of a B vertex gives a form closely related to the pentagonal bipyramid (2). Starting with the $D_{4d}$ square antiprism, removal of a vertex generates a structure that is closely related to the capped trigonal prism 4 by a minor relaxation of the other vertices and again to the other forms (2 and 3) by slightly larger displacements of the other vertices.

There is an endless array of variants for 2, 3, and 4 of which possibly the most important are directly generated from the $D_{5h}$ pentagonal bipyramid by out-of-plane motions of the vertices in the relatively cramped pentagonal girdle as shown for 5 ($C_1$) and 6 ($C_2$) which combined with the symmetric and antisymmetric axial bending modes can not only generate the $C_1$ and $C_2$ polytopal forms but also describe the pathways for interconversion of the three high-symmetry seven-coordinate forms. Forms 5 and 6 were first identified by Bartell and Thompson in calculations based on the valence-shell electron-pair repulsion model (VSEPR). They examined seven points constrained to move on the surface of a sphere and subject to a repulsive force $\sum_i r_i^{n+}$, where $r_i$ is the distance between two of the points. For $n < 2.5$, that is soft potentials, the preferred geometry was $D_{4h}$. For $2.5 < n < 5$ the lowest energy structures were the $C_1$ and $C_2$ forms 5 and 6 which merged into the $C_{3v}$ capped trigonal prism at $n \approx 5$. For $n > 5.6$ the capped octahedron $C_{3v}$ was the preferred conformation. In the range of low $n$, there was little difference in energy between any of the structures. Other variants that may be relevant only to seven-coordinate complexes with stereo-
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A representation of the interconversion of the $D_{3h}$ pentagonal-bipyramidal and $C_{3v}$ capped trigonal-prismatic geometries. The dotted lines in the $D_{3h}$ form represent the edges that are "lost" in the diamond to square face transition from $D_{3h}$ to $C_{2v}$. All forms between the $D_{3h}$ and $C_{3v}$ limits have $C_2$ symmetry and this common axis is denoted in the limiting geometries.

Figure 3. A representation of the interconversion of the $D_{3h}$ pentagonal-bipyramidal and $C_{3v}$ capped trigonal-prismatic geometries.

A stylized representation of the interconversion of the $C_{2v}$ capped trigonal-prismatic and $C_{3v}$ capped octahedral geometries. All forms intermediate between the symmetry-defined limits have a mirror plane which in the representation is vertical and normal to the plane of the paper. In this interconversion, the square (or really rectangular) faces in $C_{2v}$ are converted into diamond faces in $C_{3v}$. The dotted lines in the figure to the right represent the new "diamond" edges.

Figure 4. A stylized representation of the interconversion of the $C_{2v}$ capped trigonal-prismatic and $C_{3v}$ capped octahedral geometries.

chemically constrained polydentate ligands or to special solid-state electronic or packing factors are the family of square base-trigonal cap polyhedral forms, of which one of the two $C_i$ forms is represented by 7, and the $C_{3v}$ capped trigonal prism is converted into diamond faces in $C_{3v}$. The dotted lines in the figure to the right represent the new "diamond" edges.

Figure 5. A possible mode for interconversion of the $D_{3h}$ pentagonal-bipyramidal and $C_{3v}$ capped octahedral geometries. All forms intermediate between the symmetry-defined limits have a mirror plane which is vertical in this perspective and normal to the plane of the paper. The dotted line edge in the $D_{3h}$ drawing is "lost" in the traverse, to be replaced by the orthogonal diamond edge shown in dotted outline for $C_{3v}$.

menting previous MO studies in the literature. The emphasis is on an understanding of substituent site preferences and the pattern of bond strengths in this important class of molecules. We proceed along the lines of our previous analysis of five-coordination, using extended Hückel calculations described in the Appendix.

The Basic Seven-Coordinate Geometries

There is an unavoidable complication in the discussion of the various seven-coordinate structures and their interconversions. A natural coordinate system exists for each structure—for instance, the $z$ axis as the vertical axis in 2 as drawn. However, the easy interconversion modes which connect these structures are such that they do not preserve this symmetry axis. We begin by discussing the orbitals of each of the primary structures in its natural coordinate system. In the spirit of attempting to dissect out $\sigma$ and $\pi$ effects, the initial discussion is of a hypothetical ML$_2$ system where L is a pseudoligand bearing a single $\sigma$ orbital and two electrons.

Pentagonal Bipyramid. Illustration 9 shows the familiar level pattern for this molecule. A low-lying $e_1''$ (xz, yz) does not mix in any ligand character at all. Relatively high above it lie the two metal-ligand $\sigma$-antibonding combinations $e_2$ (xy, $x^2-y^2$) and $a_1$ ($z^2$). The level pattern resembles that of five-coordination except that in the present case the second $e$ set, $e_1''$, is at much higher energy. The reason for this is that, unlike in the case of five-coordination, there is no metal $p$ orbital set which can interact in symmetry-allowed fashion with these $d$ orbitals and stabilize them. The population analysis for this structure is shown in 10. It applies to any species with 0–4 d electrons. The implication is that as far as $\sigma$ bonding is concerned the axial bonds should be marginally stronger. At first sight this conclusion appears to contrast with the trigonal-bipyramidal case in five-coordination. There the axial bonds were weaker for a phosphorane or a $d^0$ or $d^{10}$ transition.
metal complex. This was understandable from a Rundle model of normal equatorial and electron-rich three-center axial bonding. In the present case one could retain the three-center model for the axial bonding, but one is forced into a delocalized multicenter electron-deficient bonding scheme for the five equatorial bonds as well. This is because the three central-atom $s$ and $p$ orbitals in the equatorial plane transform as $a_1' + e_1'$, while the five equatorial ligand donor orbitals are $a_1' + e_1' + e_2'$. Only three fully bonding combinations result—the $e_1'$ ligand set is nonbonding until we allow mixing with the metal d orbitals. That mixing enhances equatorial bonding but apparently not to the extent of making those bonds stronger than the axial ones.

Known ML$_7$ structures are summarized in Table I. Prior to consideration of $\pi$-bonding effects it may be premature to point to any structural trends. The accuracy of some of the structures is limited. Among those which are Do$_{4h}$ or close to that limit, we have V(CN)$_4$ as, with nearly equal V–C distances, and IF$_7$, which has a distinctly shorter axial I–F bond. Shorter axial distances are typical of U(IV) PB complexes with oxgens in both axial and equatorial sites, although in these uranyl complexes, it is very likely that \( \pi \) bonding along the axis is responsible for this difference.

A consequence of the ML$_7$ charge distribution shown in 10, with apical ligands distinctly more negative, is that one would anticipate that more electronnegative ligands, better $\sigma$ acceptors, should preferentially enter the axial sites. Better $\sigma$ donors should favor the equatorial sites. This is for d$^0$-d$^2$ complexes. Were the number of d electrons to surpass 4, these conclusions would be modified. Cases where $\sigma$ effects alone are dominant are rare, but one such may be H$_2$Os(PMe$_2$Ph)$_3$. The better $\sigma$-donor hydrides are all equatorial.

Capped Octahedron. Within the C$_{3v}$ constraint and with equal M–L distances this 1:3:3 partition retains two degrees of freedom. These may be chosen as the spherical angles \( \theta_2 \) (= \( \theta_3 = \theta_4 \)) and \( \theta_3 \) (= \( \theta_6 = \theta_5 \)) characterizing the two triangles in 11. The capped regular octahedron, \( \theta_2 = 54.7^\circ \), \( \theta_5 = 125.3^\circ \), is clearly congested. The calculations of Thompson and Bartell$^{18}$ yield a relatively narrow range of optimal geometries progressing from \( \theta_2 = 73.3^\circ \), \( \theta_5 = 128.8^\circ \) for \( n = 1 \) to \( \theta_2 = 75.8^\circ \), \( \theta_5 = 131.4^\circ \) for \( n = 12 \) (\( n \) is the power in the repulsive potential). These values are not very different from \( \theta_2 = 75^\circ \), \( \theta_5 = 130^\circ \), which is what our molecular orbital calculation gives for L$_4$-; that is, seven ligands on a sphere without a central metal atom. If we insert the metal atom, we get for the d$^0$ configuration \( \theta_2 = 84^\circ \), \( \theta_5 = 138^\circ \); for low-spin d$^2$ \( \theta_2 = 80^\circ \), \( \theta_5 = 130^\circ \); for d$^4$ \( \theta_2 = 70^\circ \), \( \theta_5 = 130^\circ \). An earlier MO calculation by Gavrin$^{14}$ for the specific case of IF$_7$ gave \( \theta_2 = 75^\circ \), \( \theta_5 = 127^\circ \).

The MO results agree well with the VSEPR predictions. The slight trend that is noted is that $\theta_2$ and $\theta_5$ both decrease as the number of d electrons is increased from 0 to 4. This can be understood from the level scheme for the capped octahedron. Illustration 12 shows the d$^0$-block splitting pattern for a typical geometry, \( \theta_2 = 75^\circ \), \( \theta_5 = 130^\circ \). The $\alpha$ orbital metal contribution is nearly all $x^2$, while the $\epsilon$ orbitals are linear combinations of ($xy$, $x^2-y^2$) and ($xz$, $yz$), with some admixture of $x$ and $y$. Imagine the approach of a capping ligand to an octahedral face. The O$_2$ levels reduce in the resultant C$_{3v}$ symmetry as follows: $i_2g \rightarrow \epsilon_1 + \epsilon_4 \rightarrow \epsilon_2$. If the incoming seventh ligand bears only a $\sigma$ orbital, then it interacts exclusively with the $\alpha_1$ component of $i_2g$, destabilizing it.

The ligand–ligand repulsions, which make $\theta_2$ and $\theta_5$ greater than the regular octahedral values, can be found in the bonding molecular orbitals below the d block. The deformation which optimizes the total energy of the d$^0$ configuration actually has destabilized the 1e levels. That orbital set would be completely nonbonding in the regular octahedron. It does not interact with the incoming ligand directly, but indirectly the steric distortion pushes the six ligands into slight $\sigma$-antibonding positions relative to the metal d orbitals. Any deformation which would lead back toward the octahedron, i.e., lowering $\theta_2$ toward 54.7$^\circ$ and $\theta_5$ toward 125.3$^\circ$, would increase the energy of the lower lying $\sigma$ levels and would stabilize 1e. This is why d$^2$ and d$^4$ move to lower $\theta_2$ and $\theta_5$ relative to d$^0$. We should note that the slight geometrical trend we discuss for ML$_7$ is greatly attenuated if the ligand L is made a more realistic ligand like chloride. For MCl$_7$ we calculate an optimum $\theta_2 = 75^\circ$, $\theta_5 = 130^\circ$ for both d$^2$ and d$^4$ configurations.

A population analysis for the capped octahedron, d$^4$, is shown in 13. The d$^0$ case differs only by having less electron density in both triangles. We would conclude that as far as the $\sigma$ effect is concerned the weakest bond should be to the capping ligand for d$^4$. The better $\sigma$ donors should preferentially enter the capping site in d$^0$-d$^4$. There are no capped octahedra among the ML$_7$ structures, but there are several in the list of ML$_7$L$^-$, mixed-monodentate-ligand crystal structures that is given in Table II. The two d$^2$ structures all have $\theta_2$ near 74–75$^\circ$ and $\theta_5$ near 127$^\circ$. For some of the d$^4$ structures we do not have the required information; others are distorted toward the CTP geometry. If we ignore the asymmetries and simply average the angles, we get values such as 71.9, 128.6$^\circ$ for MoC$_2$(CO)$_3$P$_2$, 73.2, 125.6$^\circ$ for W(CO)$_2$(P$^+$)$_2$, and 72, 128$^\circ$ for WBr$_2$(CO)$_3$As$_2$. In the more symmetrical W(CO)$_3$Br$_3$ the angles are 74.1, 125.5$.\,$ No significant trend is visible. We noted above that the calculated ML$_7$ trend with L a model $\sigma$ ligand was very much diminished for L = Cl.
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<table>
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<th>Complex</th>
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$\alpha$ P = phosphate, $A$ = arsenic, $R$ = alkyl. $b$ Where two structures are indicated, the geometry is intermediate. $c$ Partitioning of ligands is indicated in an obvious way: 1:5:1 for PB, 1:3:3 for Co. 1:4:2 for CTP. $d$ U(VI). $e$ U(VI).

Table II. Structures of the $ML_4xL_3'xL_2y$ Type, with Seven Mixed Monodentate Ligands

The 1:3:3 partition of a capped octahedron points to another natural construction of level scheme 12. This is by interacting a $C_3v$ XMY$_3$ fragment with three additional ligands, a procedure that has already been implemented. $^{44}$

Capped Trigonal Prism. The capped ligand can enter a triangular or a tetragonal face of the trigonal prism, giving rise to a $C_3v$ or $D_3h$ geometry.8 or 4. The former has not been discussed in the literature, perhaps because there appear to be no known examples, perhaps because of a feeling that such a structure would be very crowded. We have not optimized this $C_3v$ geometry but have simply rotated the lower triangle of the capped octahedron so that it lies directly under the upper triangle. The resultant level scheme 14 once again shows an e orbital at low energy, $a_1$ + $e$ high up. These levels are most easily derived by bringing in a capping ligand onto a trigonal prism. The $D_3h$ structure has low-lying $a_1' + e'$ and high-energy $e''$. $^{46}$ A ligand approaching along the $z$ axis interacts strongly with $z^2$, destabilizing it.

The difference between the two $C_3v$ forms, capped octahedron vs. capped trigonal prism, is in the energy of the lower e orbital. This level is unaffected by the capping, differing little from $e'$ in $D_3h$ or a component of $t_2g$ in $O_h$. The lower e level is at higher energy in the capped trigonal prism, for there is some antibonding between $xy$, $x^2 - y^2$ and ligand $e$ functions that is absent in the octahedron. $^{46}$ Thus the $C_3v$ CTP is especially destabilized relative to the $C_3v$ CO for the $d^5$ configuration. For the $d^4$ configuration, e unoccupied, the situation changes. We in fact find the $C_3v$ CTP more stable than the $C_3v$ CO. A similar effect occurs in the uncapped $ML_4x$ structures, $^{46}$ and here as there we are not certain if the effect is real or if it is an artifact of the extended Huckel procedure.

The trigonal prism capped on a tetragonal face, 4, has received considerably more attention. The structural type is observed in NbBr$_3$- and Mo(CN-t-Bu)$_5^{2-}$ (see Table I) as well as in complexes with several more complicated ligand sets. $^5$ If equal ML bond lengths are assumed, the structure may be described by three independent polar angles: $\theta_1$ ($\theta_1 = \theta_2 = \theta_3$), $\varphi_1$ ($\varphi_1 = 180^\circ + \varphi_2 = 180^\circ - \varphi_3$), and $\theta_2$ ($\theta_2 = \theta_3$). Note that $\varphi_2$ is not constrained to 45$^\circ$; i.e., we do not require that the capped face of the trigonal prism be square.

Table III lists the Thompson and Bartell optimum geometries for $n = 1$, 6, and 12, our computed minima for several $d^5$ configurations and seven ligands alone, as well as the related parameters of the two observed $ML_7$ structures. Our $L_7$ and $ML_7$ (L is a pseudoligand bearing a single s-type orbital) $d^4$ cases are in good agreement with the observed structures and the VSEPR calculation. This is not so for the calculated $d^5$ geometry. However, when we optimized a more realistic MCl$_4$ structure, the $d^5$ case gave $\theta_1 = 86^\circ, \varphi_2 = 48^\circ$, and $\theta_2 = 145^\circ$.

The molecular orbitals of a model $C_3v$ CTP structure, corresponding to the points on a sphere optimal geometry with $n = 6$, are shown in 15. There are some minor discrepancies between this level ordering and that deduced by other workers $^{15,16,20,46,67}$ from a crystal field model, but that may be due to differing assumptions about geometries. The levels are once again easily derived from those of a trigonal prism. $^{46}$ The level ordering in $D_3h$ is $a_1' + e' > e'' > a_2' + e''$. The d levels are those appropriate to the geometrical situation at hand, namely, the $x$ axis being the threefold axis of the trigonal-prism fragment. The group theoretical reduction is $a_1' \rightarrow a_1 + b_1$, $e' \rightarrow a_2 + b_1$. The capping ligand is $a_1$ and interacts with the $y^2 - z^2$ component of $e''$ as well as with "$x^2$". These $a_1$ orbitals mix so that the lower $a_1$ becomes primarily $x^2 - y^2$ and the upper, destabilized one, $z^2$.

Note that for the first time we encounter a splitting of the two lowest levels. This is potentially an interesting phenomenon, for, if the splitting were large, a $d^5$ ML$_7$ complex might have a chance of having a low-spin ground state. In the geometry we calculated, however, the splitting of $a_1$ and $b_1$ is small, 0.16 eV. The CTP structure has been inferred for $d^4$ Mo(CN)$_7^{2-}$ $^{20d}$ and $d^4$ Ti(CN)$_7^{2-}$. $^{46}$ Electron spin resonance studies aimed at detecting the symmetry of the orbitals oc-
ocupied in these molecules would be of great interest.

Our calculations show an interesting trend on filling the lower d levels: an increase in \( \theta_2 \) and \( \theta_4 \), a slight decrease in \( \phi_2 \). To understand this behavior consider the relevant \( a_1 \) \((x^2 - y^2)\) and \( b_2 \) \((yz)\) levels, drawn in 16 and 17. The \( a_1 \), 16, level interacts with all ligands but most strongly with the unique edge 6-7. If \( \theta_6 = \theta_7 \) were to decrease, the metal–ligand, \( \sigma \)-antibonding character in \( a_1 \) would be enhanced. Therefore the larger \( \theta_6 \), the lower the energy of the \( a_1 \). Minimum antibonding with atoms 2, 3, 4, and 5 is also ensured by keeping \( \phi_2 \) near 45°. The \( b_2 \) orbital 17 has mixed into it only the tetragonal-face ligand orbitals. The greater \( \phi_2 \), the more \( \sigma \) antibonding. Similarly, maximum \( \sigma \) antibonding would occur at \( \theta_6 = 45^\circ \). Thus occupation of the \( b_2 \) orbital would favor a larger \( \theta_6 \) and a smaller \( \phi_2 \).

The electron distribution in the CTP geometry for \( d^{0} \) and \( d^{4} \) configurations is illustrated in 18. There is little relative change in the overlap populations and net charges as one goes from \( d^{3} \) to \( d^{4} \). In either case the weaker bonds are to the lower edge. Populating the \( a_1 \) and \( b_2 \) orbitals adds electron density to all ligands. In both \( d^{0} \) and \( d^{4} \) the better \( \sigma \) donors should enter the more positive sites on the unique edge.

**Trigonal Base–Tetragonal Base.** As we noted earlier, this is really a family of structures in which a set of four tetragonal or square vertices in a plane rotates against a trio of points in a parallel plane. Along the rotation itinerary 24 structures of \( C_{3v} \) symmetry are encountered: 12 of type 19, 12 of type 20.

We carried out a limited geometry optimization on 20, assuming square and equivalent triangle faces. This resulted in a geometry with \( \theta_1 = \theta_2 \) \((=0^\circ)\) = 50°, \( \theta_3 = \theta_4 = \theta_5 = \theta_6 = \phi_2 \) = 120° for \( d^{4} \) ML₇, MC₁₇, and M(CO)₇. This ML₇ geometry was then simply rotated into 19. Not unexpectedly the 12-fold barrier was minute. Type 20 was more stable by 0.002 kcal/mol. The level scheme in these structures is normal, two orbitals down and three up in energy, and the charge distributions only reveal the geometrical similarity to some of the other limiting structures.

**General Features.** The previous sections have examined in some detail the electronic structure of several of the higher symmetry ML₇ structures. Some lower symmetry structures, for instance 7 and 8, have been calculated but not described.

**Figure 6.** Correlation diagrams for the concerted elimination of two ligands from a capped trigonal prism leading to a square pyramid (top) and from a pentagonal bipyramid leading to a trigonal bipyramid (bottom). Not all levels are indicated in the diagram, but only the crucial metal–ligand bonding levels and the d block. Correlations are drawn only between the occupied levels. Note that in the bottom diagram the coordinate system, appropriate to the \( C_{3v} \) symmetry maintained, is an unconventional one for both reactant and product.

Their levels and electronic distributions resemble those of the higher symmetry structures just as their geometries do.

One general feature which emerges from our calculation is the ubiquitous level pattern of two low-energy orbitals and three high-energy or destabilized ones. Two low-energy orbitals of course implies a large gap between filled and unfilled levels, a low-spin ground state, relative kinetic stability, and a resistance to second-order Jahn–Teller distortions only to \( d^{4} \) complexes. Not surprisingly these are the ML₇ complexes that satisfy the 18-electron rule. They form by far the most numerous class of known seven-coordinate molecules, though there are several well-characterized series with more electrons. 49–51

Another feature of the calculations, not mentioned in the individual discussions, is the existence of small positive overlap populations among some of the ligands, especially in the \( d^{4} \) configuration. For example, the L–L population between the two ligands on the unique edge in the \( C_{3v} \) CTP is 0.024 and between two adjoining ligands in the pentagonal girdle of the PB is 0.020. These numbers are small. However, our past experience with overlap populations between nonbonded atoms attached to carbon in organic molecules leads us to suspect that in the context of these relatively crowded molecules such positive overlap populations, as small as they are, may be indicative of attractive interactions.

One way to look at these positive bond orders is to consider them as incipient bonding interactions. Extending that notion, we are led to analyze the reductive elimination or fragmentation reactions 21 and 22. Figure 6 shows correlation diagrams for these processes, assuming a least-motion \( C_{3v} \) reaction.
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path. Both are allowed reactions, though 22 leaves the ML₅ fragment in a low-spin d⁶ D₃h configuration, which almost certainly is an excited state for that geometry. A d⁶ D₃h molecule is likely to have a high-spin ground state. The low-spin configuration will distort to the square pyramid. In fact one can draw a correlation diagram going directly from the PB to a square-pyramidal ML₅ + L₃, and the result is a symmetry-allowed process. The reverse of reaction 21 is one mechanistic pathway invoked in some reactions of tungsten and molybdenum carbonyls. There are many pathways for mechanistic pathway invoked in some reactions of tungsten and molybdenum carbonyls. Just as there are many possible geometries for seven points on the surface of a sphere, so there are many pathways for interconverting the various limiting forms. The point was made in the introduction, and several of the possible interconversion modes were shown in 5, 6, and Figures 3–5. We have examined potential energy surfaces for these specific motions for ML₇, where L = pseudoligand, C₁, and CO. There was no geometry optimization along the reaction path but merely a progressions simultaneously in all of the spherical coordinates from one structure to the other. Because of the low C₅ or C₁ symmetry of the reaction paths, the lower d-block levels, the ones occupied in the d⁴ complexes, were split in energy. But they never split by much and always correlated smoothly from one structural limit to another. Consequently there were no significant barriers along any of the reaction paths. No level crossings were to be expected, at least for d⁰–d⁴ complexes—these are all symmetry-allowed reactions.

The only changes in energy along the rearrangement pathways were those caused by the calculated stability of the terminal structures. These we must discuss next. But before we do this we must remind the reader of two points. First, the limitations of the extended Hückel method for calculating isomer stabilities are nontrivial. Second, we have restricted ourselves to equal M–L distances. Each geometrical limit will adjust the M–L distances, and a calculation which assumes all such distances equal may not be a good guide to the actual stability.

With these cautionary points in mind we show in Table IV the calculated energies of three d⁴ geometries of ML₇, MC₁₁, and M(CO)₇, relative to the energy of the D₃h structure. There are some interesting variations in the relative energies of the various structures depending on the donor or acceptor properties of the ligands. We will discuss these in detail in the next section, but at this point note that the similarity in energy of the d⁶ MC₁₁, D₃h and C₁₁ structures that we calculate is supported by the structural data assembled in Table I. The known d⁶ fluorides are pentagonal bipyramids in the solid state, except for NbF₅, which is a capped trigonal prism. For a d⁴ M(CO)₇ we calculate a 4-kcal/mol preference for a PB. The observed Mo(CN-tert-Bu)₂⁺, for which M(CO)₇ might have been thought to be a model, is however a CTP. But another d⁴ complex, Mo(CN)₃⁺, is a PB.

As we mentioned above, there are no calculated barriers between these structures along pathways illustrated in Figures 3–5. Still further mobility in the system is introduced by the 3:4 structures 19 and 20, which are closely related to the CTP, CO, and PB geometries or way-points between these. The low barrier to rotation in the 3:4 structures was noted earlier. Such low barriers are analogous to those for intermediates in the turnstile rotation mechanism proposed for five-coordination.

Site Preferences of π Donors and Acceptors

Interactions of the π type may be probed in a number of ways. Either a test 2p orbital or a pair of them may be introduced on the pseudoligand L or actual models for π-acceptor or π-donor substituents, such as CO or C₁, can be used. Whichever probe is chosen one has a further alternative of focusing on the total energy, the energy of the specific substituent orbitals, or the overlap populations with the substituent as a measure of the extent of interaction. To some degree we have used all of these criteria in the subsequent discussion. We have also made a point of considering not only cylindrical π-interacting groups, bearing two π-donating orbitals as shown in 23, but also “single-faced” π systems, exemplified by −NR₃, −NO₂, and C₂H₄ and shown in 24. Some of the most interesting orientational effects are anticipated for this class of single-faced π systems. We begin with a discussion of the limiting structures.

Pentagonal Bipyramid. The four possible orientations of a substituent orbital are shown in 25. Reference to scheme 9 shows that the four d electrons enter the e" level which is pure xz, yz. The interactions are thus ordered ax, = ax, = eq, > eq, > eq, > eq, > eq, > eq. Before we draw the conclusions that follow from this ordering, we explore the alternative coordination geometries.

Capped Octahedron and Capped Trigonal Prism. The reason for considering these geometries together is that in both there are three distinct substitution sites: the apex, the capped face, and the uncapped face (CO) or unique edge (CTP). Let us denote these sites as c, cf, and uf in the CO and c, qf (quadrilateral face), and e (edge) in the CTP, following the notation introduced by Drew (see 28 and 29). The natural coordinate system that we defined previously for these geometries is suitable for a capping ligand but not...
Table V. Overlaps between a Substituent p Orbital and the Metal Orbital Interacting with It

<table>
<thead>
<tr>
<th></th>
<th>Capped octahedron</th>
<th>Capped trigonal prism</th>
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<tbody>
<tr>
<td>c_x</td>
<td>0.197</td>
<td>c_x</td>
</tr>
<tr>
<td>c_y</td>
<td>0.197</td>
<td>c_y</td>
</tr>
<tr>
<td>c_f</td>
<td>0.250</td>
<td>c_f</td>
</tr>
<tr>
<td>c_f</td>
<td></td>
<td></td>
</tr>
<tr>
<td>u_f</td>
<td></td>
<td></td>
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<tr>
<td>u_f</td>
<td></td>
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</table>

so for ligands occupying the other sites. For these we define a local coordinate system (x', y', z') as follows: (1) Place the ligand on the positive x' axis, i.e., the vector M–L is along x'. (2) Choose the z' axis orthogonal to x', and in the plane containing the apical M–L bond. (3) Let the y' axis complete a right-handed coordinate system. The procedure is illustrated in 30 for the e site of a CTP. We then define as the substituent p orbital along the y' axis and as the p orbital along z'. The parallel reference is to approximate parallel character to the principal z axis. The various possible substituent orientations are drawn in 31 and 32. In all cases except the CTP qf sites the ML bond lies in a mirror plane and this separation is "clean"; that is, the || and \( \perp \) orbitals cannot interact with each other and each p orbital interacts with different d orbitals.

The actual calculations were carried out with a single ligand bearing s and p orbitals and labeled as D (\( \pi \) donor) or A (\( \pi \) acceptor) dependent on the energy and occupation of the p orbitals. Details of the modeling are given in the Appendix. Also CH\(_2\)\(^+\) and NH\(_2\) were examined. The cases considered in detail were d\(^6\) ML\(_6\)D, d\(^5\) ML\(_6\)D, and d\(^4\) ML\(_6\)A. We did not analyze d\(^4\) ML\(_6\)A because the metal d orbitals that would overlap with the empty ligand p orbitals are empty; hence the only interaction that the p orbitals of A would have will be with the filled \( \sigma \)-type orbitals, and this interaction is small. Moreover, there appear to be few d\(^n\) ML\(_7\) structures containing \( \pi \)-acceptor ligands.

In both the CO and CTP polytopes there are two low-lying d orbitals that can interact with p orbitals on the ligands. In the CTP case one of the orbitals is largely \( x^2 - y^2 \) and one is largely yz. These are drawn in 33 and 34, and their primary interactions noted. In the CO case the two d orbitals are mixtures of xy and yz and of \( x^2 - y^2 \) with \( xz \), respectively. Their approximate shape is sketched in 35 and 36. These diagrams indicate the overlap possibilities. Just how good the interactions are depends on the relative values of the overlaps. Qualitative judgments are easy to make. Thus 35 and 36 imply that the cf\(_\perp\) and uf\(_\perp\) interactions should be greater than cf\(_\parallel\) and uf\(_\parallel\), respectively. Detailed comparisons are not so easy—for instance, the schematic drawing of orbital 35 does not reveal that the cf\(_\perp\) interaction is significantly greater than uf\(_\perp\). That is a consequence of the tilt of 35 and d–p hybridization which makes the orbital "point" better at a \( \pi \) orbital on a capped face substituent.

To facilitate the intercomparison of the various interactions we computed the overlaps between the metal orbitals (33–36) and a \( \pi \) orbital of a probe—a donor or acceptor at every distinct site.\(^5\) Table V summarizes this information. Note the general superiority of \( \perp \) interactions to \( \parallel \) ones. For a ligand bearing two \( \pi \)-type orbitals, that is, a cylindrically symmetrical \( \pi \) donor or acceptor, the capped octahedron provides maximum interaction in the capped face and minimal in the uncapped face. For the capped trigonal prism the quadrilateral face and edge are fairly well balanced and superior to the apical site, which by symmetry has a single interaction.

To proceed to the stage of actual predictions of substituent site preferences we must take into account the occupancy of the two d orbitals. The three cases we consider are shown schematically as
Seven-Coordination

<table>
<thead>
<tr>
<th>Substituent Site and Orientation Preferences$^a$</th>
<th>Pentagonal Bipyramid</th>
<th>Capped Octahedron</th>
<th>Capped Trigonal Prism</th>
</tr>
</thead>
<tbody>
<tr>
<td>$d^4$ A and $d^5$ B</td>
<td>$ax &gt; eq$</td>
<td>$eq &gt; ax$</td>
<td>$uf &gt; cf$, $cf &gt; uf$</td>
</tr>
<tr>
<td>$d^5$ D</td>
<td>$eq &gt; ax$</td>
<td>$eq &gt; ax$</td>
<td>$uf$</td>
</tr>
</tbody>
</table>

$a$ The symbols are defined in the text. $A = \pi$ acceptor; $D = \pi$ donor.

The first two are basically the same—if the $d$ orbitals are filled and the substituent is an acceptor or if the $d$ orbitals are empty and the substituent a donor, in either case one has two-electron stabilizing interactions. A substituent will seek out a site and an orientation (if it is "single-faced") that maximizes interaction. If the $d$ orbitals are occupied and the substituent a donor, then one encounters four electron repulsive interactions. The substituent will seek to minimize these. It may also seek interaction with unfilled $d$ levels, but our calculations seem to indicate that these are too high in energy to be significant.

We can thus proceed from the overlaps of Table V to a set of predictions for site preferences of cylindrical $\pi$ substituents and conformational preferences for single-faced donors or acceptors. Table VI summarizes this information, including for completeness the pentagonal-bipyramid case. The calculations that were done with model or specific donors or acceptors are in general accord with these predictions. In some cases, however, the difference in energy between various sites or conformations is not very large. Thus a model CO ML$_6$CH$_2^+$ prefers a cf, orientation to cfli by only 1.5 kcal/mol. In other cases the predicted rotational barriers are an order of magnitude larger.

Next we wish to compare these theoretical conclusions with the experimental evidence that is available. This is a hazardous task, for we recognize well that in these inevitably congested molecules the observed geometry may be set in large part or even entirely by the steric requirements of the ligands. Fully aware of this limitation, we nevertheless trace the full implications of the electronic effects analyzed in the previous sections.

In many $d^0$ PB complexes, the donor atoms, typically oxide or halide, preferentially occupy the axial sites. This is true not only for the ubiquitous uranyl systems but also for several Nb(V) and Ta(V) systems. There is an interesting class of $d^4$ molybdenum and tungsten(II) complexes which have ligand combinations of strong and weak $\pi$ acceptors with donor halides. All known complexes in this class have either CTP or CO structural form. Predicted substituent site preferences (Table VI) in this $d^4$ case give an ordering for the ligand extremes of strong $\pi$ acceptor (carbon monoxide or cyanide) and of strong donors (halide) of

acceptor $cf > e > uf$

<table>
<thead>
<tr>
<th>Ubuntu</th>
<th>$uf &gt; e &gt; cf + CO$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$eq &gt; ax &gt; cf$</td>
<td></td>
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</table>

The stereochemical guide would appear to be in accord with established structures in this class. For example, Mo(CN)$_3$(C$_2$H$_4$)$_3$ is a CTP with the halide at the capping position and W(CO)$_6$Br$_2$ is a CO with carbon monoxide ligands at the capping position and the capped face. Detailed discussion of the other structurally defined members in this class is complicated by the fact that few of these complexes have exact CTP or CO structural form and some seem to be intermediate between CTP or CO. In fact, the qualitative descriptions of structural form in the original crystallographic literature occasionally have been incorrect or misleading. Nevertheless, substituent site preferences maintain their validity in applications to alternative forms. The stereochemical features remain qualitatively intact irrespective of the reference (CO or CTP) form because the minor, one-step rearrangement path that interconverts CO and CTP forms (illustrated in Figure 4) permutes the vertex positions in CTP and CO forms in one-to-one correspondence with our substituent site preferences in the two forms as noted below for the unique vertices in the two forms

$c$ in CTP $\rightarrow$ $uf$ in CO
c in CO $\rightarrow$ $e$ in CP

This feature is nicely illustrated in the complex MoBr$_3$(C$_2$H$_4$)$_2$AsCH$_3$As(C$_2$H$_4$)$_2$ which first described as a CO with the carbon monoxide ligands at the capping position and at one of the capped face vertices, with bromine atoms at uncapped face positions, and with the unique diarsine ligand that is undentate at a capped face position. An alternative description as a CTP with bromines at the capping position and one of the qf positions and carbon monoxide at an e and a qf position. Our substituent site guide is in full accord with the stereochemical description of this complex in either CTP or CO form and the rearrangement mode shown in Figure 4 provides a simple geometrical correlation between the stereochemical descriptions in the two idealized limiting forms. Interestingly, the accurate description of this seven-coordinate complex places it at the midpoint of a path (see Figure 4) connecting the CTP and CO forms. Among the $d^0$ complexes in Table II is one CO and four PB. In several of the compounds we are faced with a difficult judgment, one that we would rather not make, between the donor capability of oxide and halide ligands. In the UO$_2$(NCS)$_2$ ion there is a clear partitioning between donors and acceptors, and as expected the former enter the axial sites. The above discussion has emphasized ML$_7$ structures. There are over 100 crystal structures with one or more bidentate and polydentate ligands. In these cases the constraint of the chelate ring(s) may become the determinant of both the overall geometry of the complex and the specific location of a substituent atom. Some of the consequences of such chelate constraints have been analyzed by Kept. In our discussion we mention but a handful of these complexes, those in which a clear distinction between the $\pi$-donating or -accepting characteristics of ligands may be drawn.

There is a series of $d^4$ complexes of the type M(L-L')(CO)$_3$X$_3$ where the bidentate ligand is a diarsine or a diposphine, X = Br, Cl, or I, and M = Mo or W. Most are capped octahedra. As expected, the carbonyls are in the capping or capped face sites and the halides in the uncapped face. In Mo(CO)$_3$(dppe)(SnCl)$_2$ the excellent $\pi$ acceptor (and $\delta$ donor) SnCl$_2$ occupies the capping site of a capped octahedron. W(CO)$_3$(bpy)GeBr$_3$Br has a structure intermediate between a CO and a CTP. Viewed as a CTP, it puts the donor Br in the capping site and the acceptors GeBr$_3$ and CO elsewhere. In the more clearly CTP structure of W(CO)$_3$(diars)$_2$ the capping ligand is in the capping site and the diars are located in the unique edge and quadrilateral face just as we would expect. A similar pattern is found in the CTP Mo(CO)$_3$(diars)$_2$Cl$^+$. In principle the arguments applied to site preferences should also allow us to predict bond strengths. In practice they do not seem to work well. For instance in a $d^4$ CTP system substituted by $\pi$ acceptors the combination of the $\sigma$ effect from 18 and the $\pi$ effect from Tables V and VI would lead one to predict that the bonds to the quadrilateral face ligands should.
be strongest. A group of crystal structures by Lippard and co-workers\textsuperscript{28,39,42} of MoL\textsubscript{6}X\textsuperscript{2} and MoL\textsubscript{7}\textsuperscript{2+}, \( L = \text{tert}-\text{butyl isocyanide}, \) definitely show shorter bonds to the two edge ligands.

**Prospects and Limitations**

The achievement of this investigation was the development of a molecular orbital view of seven-coordinate complexes wherein a hierarchy of substituent site preferences was established for each of the three major polytopal forms on seven-coordination. These preferences present a stereochemical view in accord with the established structural chemistry in this area. Hence it would appear that our guidelines could be utilized with fair accuracy for the prediction of stereochemistry in the three polytopal forms. Again we note that our predictions do not differ significantly from those that would be based solely on steric arguments—and steric factors are nontrivial considerations in complexes of high coordination number. We did not discern opportunities to differentiate between steric and electronic effects in seven-coordination; the directions of the two effects never seemed to diverge. If our substituent guidelines are to have predictive or rationalization value, it is essential to paint gray, at least with a broad brush, those areas where the guidelines should be applied only with great caution.

This molecular orbital view does not possess the resolution to differentiate among the three dominant polytopal forms in seven-coordination. All three forms are very similar in energy; hence molecular structure on this basis cannot be predicted. This is not a serious flaw—certainly not in a stereochemical sense—because stereochemistry can be fairly accurately assessed in each polytopal form. Most importantly, these preferred stereochemistries in each form are directly correlated by very minor intramolecular rearrangement modes such as the stylized cycle encompassed by the three paths traversing PB, CO, and CTP forms outlined in Figures 3–5. Thus, the guidelines are even applicable to complexes whose polytopal form is intermediate between two of the idealized forms. In this sense the molecular orbital view does have a dynamical range analogous to that developed for five-coordination. Application in this dynamic context would be contraindicated only if there were significant constraints due to chelate ligands and this class of ligands in itself presents complications as discussed below.

Our analysis has been limited to complexes in which the metal atom has d\textsuperscript{0} through d\textsuperscript{4} configurations, although there are seven-coordinate complexes based on metal ions of d\textsuperscript{5} configuration greater than four and also a large set based on lanthanides and actinides.\textsuperscript{5,6} The latter set may be characterized as largely electrostatic complexes in which case the \( \sigma \)-bonding considerations elucidated in the second section of this article may be sufficient for a substituent site preference guide. Actually, the majority of seven-coordinate complexes that lie outside the d\textsuperscript{0}–d\textsuperscript{4} range are primarily chelate structures.\textsuperscript{5,6} Chelate ligands can introduce substantial constraints in structure, stereochemistry, and dynamic stereochemistry. In the extreme case, the molecular structure simply may be a mirror extension beyond the preferred structure and formation of the chelate ligand. A classic example is the seven-coordinate manganese complex based on the sexadentate ethylenediaminetetraacetate ligand and one water molecule.\textsuperscript{49} The chelate ligand forms a basket about the manganese and the water molecule sits above this basket on an axis that is nearly a perfect twofold axis. This extreme example should not be structurally viewed in reference to the idealized seven-coordinate forms, nor should it be subjected to stereochemical analysis with the substituent site guide. In these extreme cases, the molecular mechanics approach, as developed by Kepert,\textsuperscript{59} should be used. The middle ground or gray area is difficult to assess. For example, there is a large class of seven-coordinate chelate structures of the form (chel),ML or (chel)\textsubscript{2}MX with the chelate ligand typically bidentate and uninegative and L a neutral donor and X a charged donor.\textsuperscript{5} In lanthanide chemistry, the common form is (chel)\textsubscript{2}ML with L most commonly a water molecule. Structural form is described either as CO or CTP with the unique donor ligand typically at the capping site in the CO form and a vertex of the unique edge in CTP form. These stereochemical forms are precisely correlated by the rearrangement mode shown in Figure 4. Stereochemical form in these complexes would seem nicely accommodated by substituent site preferences based on \( \sigma \) donors, a feature consistent with a rough characterization of the lanthanide complexes as largely electrostatic. Another common complex set within this chelate class consists of the d\textsuperscript{0} titanium group (chel)\textsubscript{2}MX species.\textsuperscript{5,6} Here a common structural form is the pentagonal bipyramid, and in this form the halide donor ligand lies at an axial site, a result that is consistent with \( \pi \)-donor site preference in the d\textsuperscript{0} case. The very same structure and stereochemistry is seen in (trop)\textsubscript{2}SnX\textsuperscript{2} and (trop)\textsubscript{2}SnOH\textsuperscript{6} which are d\textsuperscript{0} complexes and in Mo\textsubscript{2}(dtc)\textsubscript{2}, L = NO\textsubscript{6},\textsuperscript{59} NS\textsubscript{6},\textsuperscript{68} d\textsuperscript{4} cases with an acceptor substituent. Although the preference rules are consistent with molecular structure in this class, structure and stereochemistry may largely result from the dictates of the chelate ligands. These few examples should suffice to illustrate the possible pitfalls in the chelate area.

Molecular structure is sometimes inaccurately described in relation to idealized polytopal forms—even in crystallographic articles. There may be ambiguities in a description if it is referenced to idealized polyhedra. Analytical procedures have been developed for the accurate assessment of shape, and this can be done with precision if all seven ligands are identical. However, if the ligands are not identical, substantial tactical problems arise.\textsuperscript{5,6,49} Here, caution should be exercised in the assessment of structure in relation to idealized polyhedra.

An important potential application of the substituent site guide is in stereochemically outlining the reaction mechanism in substitution of six-coordinate complexes. The application is simple and will probably prove accurate for the elementary cases wherein a ligand L is added to an ML\textsubscript{6} complex with all L ligands identical. For this case, possible paths are neatly outlined in the 1 \( \rightarrow \) 2, 1 \( \rightarrow \) 3, and 1 \( \rightarrow \) 4 constructions and the resultant stereochemistries, using the new vertex as the L site, are those predicted on the basis of \( \sigma \)-bonding effects. Again these stereochemistries in each of the three polytopal forms are precisely related through minor rearrangements (Figures 3–5). With the introduction of \( \pi \)-type ligands the reaction path analysis becomes much more complicated but remains manageable. If, however, the octahedral complex has non-identical ligands of slightly varying electronic character, a realistic analysis would necessarily be quantitative in character—a requirement that could not be met with our qualitative scheme.
Seven-Coordination

Acknowledgment. We are grateful to Dr. Michael Drew for providing us with an advance copy of his comprehensive review of seven-coordination, to Professor S. J. Lippard for his interest and advice, to Professor S. Wrenford for information regarding recent calculations of shape in seven-coordinate complexes, and to E. Kronman and J. Jorgensen for technical assistance. Our research at Cornell was generously supported by the National Science Foundation and by the Advanced Research Projects Agency through the Materials Science Center at Cornell University.

Appendix

The method of calculation used is the extended Hückel procedure. The metal atom used was Mo, with a double-f 4d function. Coulomb integrals for the metal atom were obtained from a charge iterative calculation on Mo(CO)₆, with an assumed quadratic charge dependence of metal H atom.

The various parameters used in the calculations are listed in Table VII. L is a pseudogland bearing a single 1s orbital.

Bond distances used were M-L = 1.65 Å, M-Co = 1.97 Å, M-C = 2.00 Å, C-O = 1.14 Å, C-H = 1.10 Å, M-N = 2.00 Å, N-H = 1.04 Å, and M-Cl = 2.45 Å.

To probe π-donor and -acceptor character p orbitals with exponent 1.625 were added to the π ligand L. The p orbital H₂ and the total electron count were varied to make the p orbitals strong or weak donors or acceptors as desired.

References and Notes

(1) There are relatively simple guidelines that to a first approximation provide a perspective for structural inorganic chemistry. Formal systems are directly applicable to coordination compounds, boron hydrides, and metal clusters. A brief discussion of structural systems in inorganic chemistry may be found in "Boron Hydrides", E. L. Muetterties, Ed., Academic Press, New York, N.Y., 1975, Chapter 1.


(k) A summary of the relevant structural facts may be found in ref 5 and L. Cattalini, U. Croatto, S. Degetto, and E. Tondello, L. ibid., 38, 818 (1960).


(m) The general case refers to a coordination complex in which all ligands are directly applicable to coordination compounds, boron hydrides, and metal clusters. A brief discussion of structural systems in inorganic chemistry may be found in "Boron Hydrides", E. L. Muetterties, Ed., Academic Press, New York, N.Y., 1975, Chapter 1.


(r) In an MO calculation on V(CN)₄³⁻, Levenson obtained a greater population of the ligand's acceptor orbital and would call eq, than of eq. This was attributed to the large ligand overlap in the equatorial plane.
Pfeiffer Effect in Tetrahedral and Negatively Charged Octahedral Coordination Compounds

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The Pfeiffer effect can be utilized to study the optical rotatory and related properties of dissymmetric complexes which are difficult or impossible to resolve because of their optical lability. Most Pfeiffer-active systems are based upon optically active (racemic) six-coordinate complex cations. Examples of Pfeiffer-active systems based upon optically active (racemic) four-coordinate, tetrahedral complexes and negatively charged, octahedral complex anions are reported here, in which both optically active organic compounds and complex inorganic compounds are utilized as environment substances.

Introduction

The Pfeiffer effect is the change in rotation of a solution of an optically active substance (called the "environment" substance) upon the addition of a racemic mixture of dissymmetric, optically labile, complex inorganic compounds. Many Pfeiffer systems have been reported for tetrahedral complexes and negatively charged, octahedral complex anions. The only previously reported case of a tetrahedral complex exhibiting the Pfeiffer effect is for the bis(8-aminquinoline)zinc(I) ion. Another tetrahedral complex, bis(8-hydroxyquinoline-5-sulfonato)zinc(II) ion, is reported herein to exhibit the Pfeiffer effect with certain environment substances.

The only negatively charged racemic complexes to show a Pfeiffer effect have been complexes with the oxalate ion as a ligand, and this racemic complex was reported herein to exhibit the Pfeiffer effect with certain four-coordinate, tetrahedral complexes and negatively charged, octahedral complex anions.

Experimental Section

Optical rotations were determined on a Perkin-Elmer 141 photoelectric polarimeter at a wavelength of 589 nm and at a temperature of 23 °C. All optical rotatory dispersion and circular dichroism spectra were determined on a Cary 60 recording spectrophotometer at 23 °C. The nonpolarized absorption spectra were taken on a Cary 14 recording spectrophotometer at 23 °C. The analyses of the crystals thus obtained were in accord with the empirical formula. The analyses of the complex did not agree with the empirical formula.

The zinc complex H2[Zn(QS)2] (QS = 8-hydroxyquinoline-5-sulfonate) was prepared according to the method of Liu and Bailar. The analyses of the complex did not agree with the empirical formula, so a purification procedure was developed. This procedure was to dissolve 10 g of the complex in 200 ml of water and then filter the undissolved material. The filtrate was then made acidic by adding hydrochloric acid until the pH of the filtrate was approximately 3.5. The analyses of the crystals thus obtained were in accord with the empirical formula.

The synthesis of the Na[Ni(acac)3] complex was first reported by Dwyer and Sargeson. This complex was obtained from Mr. John Klein and Dr. Richard Lintvedt of Wayne State University and purified according to a method described by Klein. The complex (5.0 g) was dissolved in 250 ml of water and vigorously stirred for 1 h. It was then filtered and washed with methanol and then with ether. It was dried at room temperature over sulfuric acid. This purification separates the inner complex [Ni(acac)2] from Na[Ni(acac)3], and the analysis of the complex agreed with the empirical formula.

All solutions used in the Pfeiffer systems were aqueous, and these systems were prepared by mixing an aliquot of a stock solution of the environment substance with an aliquot of the stock solution of the metal complexes. They were then diluted to the appropriate volume.

Aqueous solutions of d-cinchonine hydrochloride and l-brucine hydrochloride were prepared from d-cinchonine and l-brucine by adding hydrochloric acid dropwise until the d-cinchonine or l-brucine dissolved (ca. 1.5 mol of HCI/mol of alkaloid). The pH of these solutions never went below 3.5.

Results and Discussion

The [Zn(QS)2]+ Systems. There is a significant difference between the absorption spectrum of the [Zn(QS)2]+ anion and that of the ligand, QS2- (Figure 1). The doublets of the QS2- anion at 340 and 250 nm converge to singlet peaks at 365 and 255 nm in the spectrum of the complex [Zn(QS)2]+.