important difference in chemical reactivity between Fe-
(TMM)L₃ complexes (L = PR₃ (3) and L = CO): a one-
electron transfer easily occurs from 3 on attempts of al-
kylation and the resulting 17-electron intermediate is
stabilized by the basic phosphines L, whereas Fe-
(TMM)(CO)₃ is the site of electrophilic substitutions,¹⁶ but
its one-electron abstraction is difficult to perform and
immediately undergoes decomposition, presumably via
initial cleavage of an Fe–CO bond. This illustrates the
deciding role of the nature of the ligands to modulate the
behavior of complexes within the same family.

Cyclc voltammetry of 6a–e showed a reversible one-
electron oxidation at negative potential correspondingly
to the formation of Fe(o-xylene)L⁺ (6'), and an irreversible
wave between 0.50 and 0.70 V was observed (Table VI).
The latter oxidation can be attributed to the 6⁺ /6²⁺ couple
which may decompose with phosphine dissociation. The
ease of the first oxidation of 6 confirms their electron-rich
character. The lowering of the corresponding potential
follows the expected increase in electron-donor ability of
phosphorus derivatives L according to the sequence P-
(PMe₂), < PMe₃Ph < PMe₃. Of special interest is the
comparison of the first oxidation potentials for 3 and 6 for
and the same ligand L. It shows a greater reducing power
for 6 vs 3 of more than 0.2 V. This is consistent with a
lower oxidation state of the iron center in 6 than in 3.
It confirms the nature of an (η⁴-diene)Fe⁰ complex of type
IV for derivatives 6.

Conclusion
Our results present an easy access to electron-rich
(η⁴-hydrocarbon)iron complexes via FeCl₃(PR₃), interme-

diates, magnesium, and halogenated substrates. This
method avoids the preliminary preparation of unstable
Grignard reagents and should offer a potential for or-
ganometallic syntheses. The nature and the number of
ancillary PR₃ groups play a determining role for the re-
activity of FeCl₃(PR₃) with o-BrMgCH₂C₆H₄CH₃, which
leads to either a C–H activation or a C–C coupling reaction.
The studied Fe(η⁴-TMM)L₃ and Fe(η⁴-o-xylene)L₃ com-
plexes show at the same time a high tendency to give an
electron transfer and an unexpected stability of the re-
sulting paramagnetic species. These two characteristics,
which are indicative of the electron-rich nature of the
complexes, appear to be modulated by the electron-donor
capability of the phosphorus groups and also by the type
of interaction between the hydrocarbon ligand and the
metal.

Acknowledgment. We are grateful to C. Mountassir
and Dr. S. Sinbandhit for helpful assistance and discus-
sions.

Registry No. 1a, 55853-16-2; 1b, 95911-92-5; 1d, 55059-43-3;
2, 1871-57-4; 3a, 95911-86-7; 3a(NO₃), 111409-19-9; 3b, 95911-83-4; 3b(NO₃), 95911-56-8; 3c,
111409-15-5; 3c(NO₃), 111409-17-7; 4a, 612-12-4; 5, 56812-
61-4; 6a, 111409-13-3; 6a⁺, 95911-14-4; 6b, 95911-91-4; 6d,
95911-89-0; 7, 952-80-7; 8b, 111409-18-8; 9b, 110074-88-9;
Me₂(C₃H₅)PMe₂, 77609-83-7; CH₂C(CH₂Cl)₃, 1067-09-0.

Supplementary Material Available: Tables of bond lengths,
bond angles, and anisotropic temperature factors and an ORTEP
of the structure including the anion (Figure 51) for 3a⁺(CF₃SO₃⁻)
(4 pages); a listing of calculated and observed structure factors
(10 pages). Ordering information is given on any current mshead page.

Metallacyclobutadiene versus Metallatetrahedrane Structures
for Cl₃MoC₃H₃ Complexes
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Received May 13, 1987

The bonding and energetics in the Cl₃MoC₃H₃ metallacyclobutadiene are contrasted with those in the
metallatetrahedrane. The complexes are found to be within 20 kcal/mol of one another, with the me-
metallatetrahedrane being the lower in energy. The Mo–C bonds are essentially covalent in both structures,
and the metallacyclobutadiene possesses resonance stabilization. Charge distribution in the carbon fragments
reveals no cyclopropenium character in the Cl₃H₃ ring of the metallatetrahedrane. Bonding, energetics,
and charge distributions are all discussed with an emphasis on structure and reactivity.

I. Introduction
Metallacyclobutadienes A are believed to play the same
role in acetylene metathesis¹ (eq 1) as metallacyclobutanes
B play in olefin metathesis² (eq 2). Strong evidence for

of the metallacylcobutadienes isolated are active catalysts.\(^6\)\(^-\)\(^7\) Species 2 and 3 are, however, isolated as end products from catalytically active systems.\(^4\)\(^5\) Despite the experimental observation of these species, there remain a number of questions concerning the bonding in these complexes. Are the complexes best viewed as metallacyclobutadienes, and, if so, is there resonance stabilization or destabilization in the \(\sigma\) system? The X-ray structures of 1, 2, and 3 reveal a particularly short \(W-\beta-C\) distance ranging from 2.093 to 2.159 \(\text{Å}\)—only about 0.2 \(\text{Å}\) longer than the \(W-\alpha-C\) bond.\(^5\)\(^-\)\(^6\) This naturally led to speculation that there might be a bonding interaction between \(W\) and the \(\beta\)-carbon and that 1, 2, and 3 might be structurally or electronically different from the metallacycles presumed to form in the catalytically active systems. Further studies of dialkylacetylene metathesis involved varying the steric bulk of the alkyl groups on the metallacyclic carbons and of the axial-equatorial ligands. The conclusion reached was that sterics are the overriding factor influencing the rate of metathesis and thus the stability of the metallacycle.\(^4\)\(^-\)\(^7\)

The fate of 1 in the presence of excess diethylacetylene is not that of metathesis but of cyclopentadienyl formation. Surprisingly, two cyclopentadienyl complexes (4 and 5) are formed.\(^7\) The simplest mechanism which would give rise to a cyclopentadienyl complex is one in which the alkyn coordinates and then inserts into a \(W-\alpha-C\) bond to yield a tungstabenzenes intermediate, which then collapses to a cyclopentadienyl complex. Such a route should yield only 4. A possible answer to this problem\(^7\) was suggested following the isolation of 6, \(\text{W}[\text{C}_2\text{H}_2(\text{p}-\text{Pr})_2]\text{O}_{1.5}\text{W}\) from the reaction of 1 and TMEDA. The complex is pseudoctahedral and contains a \(\text{symmetrically bound } \gamma^3\text{-cyclopentadienyl ligand. This interconversion between a metallacylcobutadiene and a metallatetrahedrane allows for the formation of the two cyclopentadienyl complexes by rotation of the \(\gamma^3\) ring and, thus, isomerization of the alkyl groups in the metallacyclobutadiene. The structure of 6 revealed that the ring substituents all bend up out of the plane of the carbon ring.\(^7\) The original investigators\(^7\) explain this in terms of steric interactions between the ring groups and the \(W\) ligands. The origin of this interaction is presumably due to the carbon ring being drawn close to the metal due to the latter's electron deficiency. These investigators\(^7\) proposed \(W(\text{IV})\) and \((\text{C}_3\text{R}_3)^+\) but did not rule out that the tungstetetrahedral is \(W(\text{VI})\) with a \((\text{C}_3\text{R}_3)^2\) ligand. They suggested that \((\text{C}_3\text{R}_3)^+\) with \(W(\text{II})\) seemed least satisfactory.

The interconversion between metallatetrahedrane and metallacylcobutadiene has been treated theoretically for the general case where an \(ML_n\) fragment cleaves a \(C-C\) bond of the metallatetrahedrane to give a metallacylcobutadiene.\(^8\) Equilibrium geometries for several \(ML_n\) and \(C_3R_3\) fragments were predicted, but detailed analyses of bonding and energetics were not presented. Therefore we undertook to study in detail a specific system of considerable current experimental interest.

The experimental studies suggest several questions which lend themselves to detailed theoretical analysis. What is the nature of the \(W-\beta\) bonding interaction in the metallacylcobutadiene and the metallatetrahedrane; is it mostly covalent or ionic? What are the charges on the carbon fragments and how large is the electron deficiency of each species? Is there any \(W-\beta-C\) bonding interaction in the metallacylcobutadiene? What are the relative energetics between the two structures and why do they interconvert so readily? Is there any barrier to rotation of the \(C_3R_3\) ring in the metallatetrahedrane species? What role do the electron-withdrawing ligands play in the energetics of the species?

### II. Calculational Details

#### A. Basis Sets and Effective Potentials

All electrons were considered explicitly for \(\text{C}\) and \(\text{H}\) but effective potentials were used to replace core electrons of \(\text{Cl}\) and \(\text{Mo}\). For \(\text{Cl}\), the \(\text{Ne}\) core was replaced with the \(\text{SHC}\) effective potential\(^9\) (treating neutral \(\text{Cl}\) with seven explicit electrons) and for \(\text{Mo}\), a relativistic effective potential\(^10\) was used for the \(\text{Zn}\) core (treating neutral \(\text{Mo}\) with 12 explicit electrons). All calculations used Cartesian Gaussian basis sets. For carbon,\(^9\) the 9s, 5p primitive Gaussian basis was contracted to valence double 6s, 3p, 2p. For hydrogen,\(^9\) the 3s primitive basis was scaled (\(\gamma = 1.2\)) and contracted to 2s.\(^11\) For \(\text{Mo}\),\(^9\) the basis set was contracted to 3s, 4p, 2d from the primitive 3s, 5p, 3d. For \(\text{Cl}\),\(^9\) the 3s, 2p primitive basis was contracted to 1s, 1p based on \(\text{TiCl}_4\).

#### B. Wave Functions

Wave functions were calculated at the Hartree–Fock (HF), generalized valence bond (GVB), and generalized valence bond configuration interaction (GVB-Cl) levels. For HF, the singlet state has all orbitals doubly occupied

\[
\phi_{\text{HF}} = \phi(1)\phi(2)(\alpha\beta - \beta\alpha)
\]

but optimized self-consistently. This leads to a good description of bonds constructed from highly overlapping orbitals but a poor description of bonds involving low overlap (e.g., \(M-C\) bonds). The GVB wave function introduces electron correlation effects by allowing each electron to have its own orbital, which is then optimized self-consistently. For a typical two-electron bond, the GVB wave function has the form

\[
\phi_{\text{GVB}} = \left[\phi_1(1)\phi_1(2) + \phi_2(1)\phi_2(2)\right](\alpha\beta - \beta\alpha)
\]

where \(\phi_1\) and \(\phi_2\) are, in general, fairly localized on opposite atoms in the bond.

Because some electron pairs are well treated in HF (as in 1), while others require correlation (as in 3), GVB cal-

---


calculations often correlate only a portion of the bond pairs.\(^{12}\)
Generally, we correlated the pairs that change significantly during a reaction sequence and those that have large correlation errors. To indicate the level of correlation, the GVB wave function is denoted as GVB \((n/m)\), where \(n\) is the number of electron pairs being correlated and \(m\) is the number of orbitals used for the correlated pairs (generally \(2n\)). In the various \(\text{C}_{2}\text{Mo}(\text{C}_{3}\text{H}_{3})\) complexes, the 12 electrons involved in Mo–C and C–C bonding are correlated since these electrons change dramatically as the structure is changed from 7 to 8. Thus we carry out GVB \((6/12)\) level calculations. The shape of the self-consistent GVB orbitals is used to interpret the bonding characteristics of the wave function. We find that these six bond pairs have the following character: the metallacyclobutadiene \(7\) has two metallacycle \(\pi\) bonds, two Mo–C \(\sigma\) bonds, and two C–C \(\sigma\) bonds (leaving the other orbitals as self-consistently double occupied orbitals). The metallatetrahedrane involves three Mo–C bonds and three C–C bonds.

The self-consistent GVB calculations were restricted (perfect pairing) such that each correlated pair is constrained to have the form in eq 4. This leads to an excellent description of systems that are well-described in terms of one bonding structure (e.g., 8) but not of systems involving strong resonance effects (as in 7). One approach to handling resonance in the GVB description is to optimize the orbitals for the resonating structures \((7a\) and \(7b)\) self-consistently, while allowing different orbitals for the two structures.\(^{13}\) This is termed GRVB (generalized resonating valence bond) and has been applied to similar systems (e.g., cyclobutadiene).\(^{14}\) An alternative approach is to start with the GVB orbitals for one bonding structure, say \(7a\), and then to carry out a configuration interaction calculation in which the occupation of the orbitals is allowed to change, permitting the orbitals to describe other resonance structures (e.g., \(7b\)). For the metallacyclobutadiene, we used the latter approach. Thus for \(7a\), the Mo–C and C–C \(\sigma\) bonds were allowed to have all spin pairings within each set of natural orbitals in a given bond pair (not just the one in eq 4); this is termed GVB-RCI. We allowed all excitations within the four natural orbitals representing the GVB \(\pi\) bonds (GVB-Cl). In addition, to allow readjustments in the shapes of the various orbitals in the presence of resonance, we allowed all excitations out of the dominant configurations into the entire valence space (GVB-RCI-S) for the Mo–C bonds in order to allow resonance. For the metallatetrahedrane, the C–C bonds were described with GVB-RCI and the Mo–C bonds with GVB-Cl.

**III. Results and Discussion**

First, we will examine the GVB description of the bonding in both the metallacyclobutadiene and the metallatetrahedrane with an emphasis on qualitative aspects. Next, we examine charge distribution, and finally we will compare relative energetics between the two complexes.

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Structures for Cl₃MoC₃H₃ Complexes

Figure 2. Contour plots for the GVB-PP orbital of metallacyclobutadiene. Spacing between contours is 0.05 au; solid lines are positive and dashed lines are negative.

B. Orbitals. The contour plots of the GVB orbitals for the metallacyclobutadiene are presented in Figure 2 and the GVB orbitals for a metallatetrahedrane in Figure 3. As discussed in section II.B, each GVB orbital has one electron, but the two orbitals of a bond pair are allowed to overlap, forming a wave function as presented in eq 2.

1. Bonding in the Metallacyclobutadiene. As indicated in Figure 2, the metallacyclobutadiene has covalent Mo-C₁, C₁-C₂, C₂-C₃, and Mo-C₃ σ bonds and covalent C₁-C₂ and Mo-C₃ π bonds. Thus, the name metallacyclobutadiene is appropriate. Considering both resonance structures (7a and 7b), the bond order is 1 1/2 for each Mo-C and C-C bond. The result is that three d orbitals on the Mo are involved in the bond to the C₃H₃ fragment. The other three valence electrons initially on Mo are involved in partially ionic bonds to the three Cl atoms.

The C₃H₃ framework has a central bond angle of 118.1°, and the C-C bond pairs are well directed along the bond axes (unstrained). However, parts a and b of Figure 2 show that the Mo-C single bonds are strained. The center of the C lobe of the GVB bond is at an angle of 120° with respect to the C₁-C₂ axis (as if it were going to point at the H of planar allyl), leading to an angle of 42° off the line joining Mo and C₁. Similarly, the d-like bonding orbital on the Mo points about 22° off of the Mo-C₁ vector, leading to a bent bond. The net result is that the two Mo d-like orbitals form Mo-C bonds lying at an angle of 129° with each other. This is quite consistent with the prediction of Rappé and Goddard who showed that in order for two pure d orbitals to each be symmetric about their respective bond axes, the angle must be 125.3°.

The C-C π bond (Figure 2h) is conventional, involving p orbitals on each carbon. The Mo-C π bond (Figure 2g)
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(a) Mo-C Bond Pair

(b) C-C Bond Pair

Figure 3. Contour plots for the metallatetrahedrane.

The results from ab initio calculations are quite different. The dπ bonds that were used to build molecular orbitals in the PMOT approach are instead used to make localized π bonds, which are then allowed to resonate (7a and 7b). This approach allows for a full 1.5 bond order between the carbons and also includes electron correlation in the bond pairs. This metalloaromatic system is thus stable because of the two dσ symmetry orbitals that allow the metal to form two π bonds simultaneously (by resonance) with bond orders each less than 1. In contrast, cyclobutadiene, uses a single carbon p orbital to form the π interaction with both adjacent carbon p orbitals.14

There is also an additional resonance structure which could arise from the PMOT approach by second-order mixing of the antibonding allyl orbital and the b2 orbital. This would yield the valence bond structure shown.

Figure 4. Mulliken populations for each atom in the metallatetrahedrane and metallacyclobutadiene.

There is no significant mixing of this state in our ab initio calculations, as indicated by the positive charge of 0.86 found on the Mo and the small charge (−0.03) on the Mo-C (see section C and Figure 4).

To form the two Mo-α-C bonds and two resonating π bonds, the Mo uses four of its five available orbitals. The empty orbital is of d4z character and leads to a coordination of a ligand along the x axis. An acetylene coordinated along this axis could then insert into an Mo-α-C bond to form a metallacyclohexatriene complex. With this insertion product formed, resonance is initially lost from the metallacyclobutadiene but is reintroduced in a hypothetical planar metallacyclohexatriene complex. While the C-Mo-C angle is preferred to be 125° to form the optimum directional dσ bonds, the axial Cl-Mo-Cl angle is expected to be large (166° in this case) due to high sd character (180° is optimal for two such bonds). Due to the high electronegativity of Cl, the Mo-Cl bonds are highly polarized toward Cl. These polar Mo-Cl bonds use Mo s orbitals since the ionization potential of the 5s electrons is less than for the 4d electrons. As a result of the charge transfer to the chlorines, the metal is positive,
leading to covalent bonds to the carbons that are 4d-like and highly directed. This is the case in both the metallocyclobutadiene and the metallatetrahedrane.

2. Bonding in the Metallatetrahedrane. For the metallatetrahedrane, all GVB wave functions led to three covalent Mo-C bonds. All attempts to force another description such as a π allyl complex only resulted in revision to the three covalent bond description. A typical Mo-C σ bond is in Figure 3a.

Each Mo-C bond involves a Mo d σ orbital pointing at an sp³-like orbital of the C₃H₃ fragment to form a σ bond. The metallatetrahedrane Mo-C σ bonds are less directional (more spatially diffuse) than in the metallocyclobutadiene because the metal now is forced to make three σ bonds rather than two. The bond is slightly bent (~12°) from the bond axis, leading to an angle of 63° between ends of the Mo d orbitals involved in each Mo-C bond. This small angle is created by the short C-C distance compared with the Mo-C distance. Rappé and Goddard² showed that three equivalent orthogonal d orbitals, each rotationally symmetric about its bond axis, must be at an angle of 54.7°, in reasonable agreement with the calculated results.

A contour map of the C-C bond is shown in Figure 3b. It involves typical sp³ orbitals localized on each carbon with no interaction with the metal.

C. Charges. Reactivity in transition-metal complexes is controlled by the charge distributions between the metal and its ligands and by the nature of the metal-ligand bonds. The proposed reactivity of the metallocyclobutadiene in coordinating an acetylene requires that the metal center have an empty orbital and be electrophilic. The metallocyclobutadiene is a 12-electron species and therefore has empty orbitals. It is also calculated to be electrophilic, with the metal having a high positive charge of 0.86. In this complex, each chlorine takes approximately 0.34 electrons (see Figure 4 for all Mulliken populations). The α carbons are slightly negative and the β carbon is neutral. This charge distribution of the organic ligand is basically covalent, with no large charge polarization between the Mo and the C's. This is also apparent by viewing the contour maps in Figure 2, parts a and b. Each one-electron orbital is essentially centered on the individual atoms.

The chlorine-to-molybdenum bonds behave the same in both complexes. The chlorines serve to remove s electron density from the metal. In each case, the chlorines take about 1.1 electrons and leave the positive charge to be dispersed among the Mo and the organic fragment.

The metallatetrahedrane has a similar charge distribution as the metallocyclobutadiene. Each chlorine takes about 0.38 of an electron and leaves the Mo with a positive 0.74 charge and the ring with a positive 0.34 charge (0.34 from the tetrahedrane, 0.19 for the butadiene). Again, the carbons have a slight negative charge, reflecting a higher electronegativity than Mo or H. This is as expected for a covalent Mo-C interaction.

In order to explore how charge transfer is affected by geometry of the C₃H₃ ring, we recalculated the wave function with a flat C₃H₃ ring (the original geometry had the H's bent back by 44°). Since C₃H₄⁺ is "antiaromatic," this might lead to a propensity for C₃H₄⁺ and hence charge transfer to the metal. Indeed, the carbons lose 0.18 electron to obtain a net charge of +0.62, while the metal gains 0.12 electron to obtain a net charge of 0.82. Basically, the strong tendency for the Mo and the C's to form covalent Mo-C bonds prevents charge flow from the ring in either the flat or bent up hydrogen geometry. Furthermore, metallatetrahedrane with the flat C₃H₃ geometry is 6 kcal/mol (GVB (6/12)) above the system with a nonplaner C₃H₃ geometry. Although this bent back geometry is favored by steric interactions with the metal ligands, we believe that the dominant factor is the covalent metal-carbon bonding. Hence, the "bend-back angle" is a guide to the true metallatetrahedrane character in the bonding. Every metallatetrahedrane may have different bend-back angles depending on the charge distribution between the metal and the carbon ring, the oxidation state of the metal, and the steric requirements of the ligands on the metal and the alkyl groups on the carbon ring. In summary, the charge transfer is dominated by the nature of the M-C bonds. Because of the greater polarity in its Mo-C σ bond, the Mo in the metallocycle is more electrophilic than in the metallatetrahedrane.

D. Geometries and Energies. A dominant factor in the interconversion of the metallocyclobutadiene and the metallatetrahedrane is the difference in ground-state energies. In Figures 5 and 6, the energy separations between the metallocycle and the metallatetrahedrane are shown for various levels of theoretical treatments.

The metallocycle has four σ bonds, two of which are Mo-C and two are C-C. It also has one Mo-C π bond and one C-C bond. The metallatetrahedrane also has six covalent bonds, three of which are Mo-C and three are C-C.

Taking GVB overlaps as a rough guide to relative bond overlap strengths, we used the following approach to estimate the potential energy required to interconvert the two forms. For each level of treatment, the energy difference was calculated for the interconversion of the metallacyclobutadiene and the metallatetrahedrane at the CI level (GVB overlap). Table I gives the orbital overlaps for the GVB pair orbitals.

![Figure 5](https://example.com/f5.png)

**Figure 5.** Ground-state energy difference of several metallatetrahedrane and the metallocyclobutadiene at the GVB (6/12) level.

![Figure 6](https://example.com/f6.png)

**Figure 6.** Ground-state energy differences of the metallatetrahedrane and the metallocyclobutadiene at the CI level.

<table>
<thead>
<tr>
<th>Orbital Overlaps for the GVB Pair Orbitals</th>
<th>Metallocyclobutadiene</th>
<th>Metallatetrahedrane</th>
</tr>
</thead>
<tbody>
<tr>
<td>bond overlap</td>
<td>bond overlap</td>
<td></td>
</tr>
<tr>
<td>Mo-Cσ</td>
<td>0.74</td>
<td>Mo-Cσ</td>
</tr>
<tr>
<td>Mo-Cσ</td>
<td>0.74</td>
<td>Mo-Cσ</td>
</tr>
<tr>
<td>Mo-Cτ</td>
<td>0.53</td>
<td>Mo-Cτ</td>
</tr>
<tr>
<td>C-Cσ</td>
<td>0.71</td>
<td>C-Cσ</td>
</tr>
<tr>
<td>C-Cσ</td>
<td>0.87</td>
<td>C-Cσ</td>
</tr>
<tr>
<td>C-Cσ</td>
<td>0.87</td>
<td>C-Cσ</td>
</tr>
<tr>
<td>total</td>
<td>4.46</td>
<td>total</td>
</tr>
</tbody>
</table>

The metallatetrahedrane has the higher energy, indicating that it is less stable than the metallocyclobutadiene. This is consistent with the observed preference for the metallocyclobutadiene in many coordination reactions.
the frontier orbitals of the cyclopropenium quartet state lead to the orbital interaction diagram of Figure 9 (no energy scale intended). The 1e and 1a1 orbitals on the metal are of the correct symmetry to mix with the equivalent symmetry orbitals of the cyclopropenium fragment only in an eclipsed geometry. Thus, covalent three-coordinate early-transition-metal systems (where the 1a1 and 1e set of orbitals is only singly filled) prefer to make covalent bonds to three other substituents in an eclipsed geometry. As these orbitals fill, the preference for a staggered (octahedral) geometry becomes stronger. Steric interactions could also lead to preference for staggered and are probably the dominant interactions in most experimental systems. Thus, in group VIII (group 8) metals, systems where these orbitals are filled (e.g., ClFeSMe2),
the geometry would be staggered due to the symmetry of the high-lying 2e and 2a1 orbitals.

Literature structures related to these conclusions are limited to the trigonal prism complexes of early transition metals with dithiolato bidentate ligands. It has been suggested that these complexes are eclipsed due to S-S bonding interactions. Since this suggestion, dithiolato complexes have been isolated with S-S distances greater than the sum of the covalent radii. We propose that the eclipsed preference is instead due to the electronic effects presented above.

The preference for eclipsed geometries derives from the high-spin MoCl3 fragment. This fragment has been calculated previously, and our calculations confirm the past results. The 1e orbitals are tilted from the axis due to mixing of dxy with d8y and d8z. The question of bonding then reduces to why the 1e orbitals mix in such a fashion so as to prefer eclipsing. Calculations on MoCl3 at several Cl-Mo-Cl angles reveal the preference for the high-spin orbitals to lie in the nodal planes of the Mo-Cl bonds. This allows for the least repulsion due to orthogonality between various occupied orbitals. It can be seen in Figure 8 that the dxy and d8 combination lies with the positive lobe directly between two Mo-Cl bonds. Consequently, the dxy and d8 combination has an angular node placed directly along an Mo-Cl bond. In this manner

---

**Figure 7.** Singly occupied orbitals localized over the Cl's of the quartet state of MoCl3.

**Figure 8.** Singly occupied orbitals of the quartet state of MoCl3.

---

the high-spin orbitals prefer to make bonds directly over the Mo–Cl bonds.

**E. Electron Correlation.** The above calculations used the simple GVB-PP wave function where the orbitals are optimized for a single bonding structure (for spin coupling). Thus these calculations are biased against this fully delocalized metallacycle, which requires two configurations.

In order to elucidate further the energetic differences between the metallatetrahedrane and the metallacyclobutadiene, we undertook several calculations designed to allow resonances in the metallacyclobutadiene.

An Mo atom with three unpaired d orbitals prefers the high-spin state (Hund’s rule) in which all three d electrons have the same spin, say \( \alpha \). However, bonding these electrons to ligands requires that the two electrons in the bond have opposite spins and that each electron has both \( \alpha \) and \( \beta \).

As a result, the d electrons on the metal can no longer all have the same spin. Thus the high-spin coupling of the metal tends to inhibit full bonding to the ligands and vice versa. A proper description of these spin-coupling effects requires bonding structures in which the two electrons of a given pair are allowed to be coupled either low spin (bonding) or high spin (antibonding). Such configurations are included in the GVB-RCI wave function, and we find that the metallacycle energy drops 6.28 kcal/mol more in the RCI than does the tetrahedrane. The contribution from such excitations within each GVB \( \pi \) bond pair resulted in a 19.8 kcal/mol lowering (half the total RCI lowering). Comparison to the analogous excitations from the C–C \( \sigma \) bonds of the metallatetrahedrane shows only a 7.63 kcal/mol lowering.

In order to allow resonance to build into the metallacycle, a full GVB-RCI in the \( \pi \) space was allowed, with simultaneous \( \sigma \) relaxations (single excitations out of the \( \sigma \) bonds). This gave a further lowering of 9.86 kcal/mol. The tetrahedrane was then subjected to a comparable calculation. It consisted of a full GVB-Cl in the Mo–C space while leaving the RCI in the C–C space. This gave a further lowering of 6.56 kcal/mol. The final separation between the two geometries is 8 kcal/mol, favoring the metallatetrahedrane. Due to the incorporation of resonance into the \( \pi \) system, these final CI’s introduced a greater lowering for the metallacyclobutadiene than for the metallatetrahedrane.

The long Mo–C bond distances (in comparison to C–C bond distances) places the \( \pi \) bonds much further from one another than in cyclobutadiene. This means that the repulsion from orthogonality that is so destabilizing in cyclobutadiene is much less evident in the metallacyclobutadiene and therefore the metallacycle takes advantage of some resonance stabilization. Analysis of the resonance configurations shows a resonance stabilization of 8.68 kcal/mol.

**F. Implications for Chemistry.** This study allows us to speculate about the mechanism of metathesis performed by metallacyclobutadienes. It is viewed as decomposition of the metallacycle to an acetylene–alkylidyne adduct which then loses the initial acetylene before or after coordinating a second acetylene. This reaction thus involves the slipping of an acetylene from a metallacycle to \( \pi \) coordination at the metal. The initial metallacyclobutadiene is a 12-electron species whereas the acetylene–alkylidyne complex is a 14-electron species. The movement of the acetylene thus increases the electron richness of the metal. A strong \( \pi \) electron donor ligand should slow this reaction. This is a direct consequence of filling the orbitals the acetylene is moving into and thus making them less accessible. This logic is supported by the calculation of Rappé and Upton for titanium metallacyclobutane olefin metathesis. In the titanium case, electron-donating substituents on the titanium slow the metathesis reaction. This in direct contrast to the tungsten–acetylene metathesis reaction where electron-withdrawing groups are indicated by Schrock et al. to slow the metathesis reaction. Thus, there is a contradiction and it would seem that the major effect operating in the tungsten metathesis systems is the steric environment imposed by the axial-equatorial ligands.

**IV. Summary**

The Mo–C bonding in both the metallacyclobutadiene and the metallatetrahedrane is covalent with no large polarization toward either Mo or C. The Cl’s remove 5s electron density from the Mo, leaving the d electrons to form hybrids that bond to the carbon fragments. The positive charge induced by the large electronegative chlorines is dispersed among the metal and the organic fragments. The metallacyclobutadiene is more electrophilic than the metallatetrahedrane. This supports the notion that the metallacycle can coordinate an acetylene. The electrophilicity of the Mo in the metallacycle argues in favor of accepting electron density from the axial-equatorial ligands. However, in contrast to the proposal by the original investigators, this acceptance would (on an electronic basis) slow the metallacycle decomposition to an acetylene–alkylidyne. The energy difference between the metallacycle and metallatetrahedrane shows that the two geometries are very close in energy (within 20 kcal/mol by all calculational procedures). This is due to covalent bonds in the metallatetrahedrane that make up for the increased strain energy. In contrast to cyclobutadiene, the metallacyclobutadiene shows resonance stabilization. There is no bonding interaction between the Mo and the \( \beta \) carbon in the metallacycle ring. The bending of the ring substituents out of the plane of the carbons in the metallatetrahedrane is due to bonding effects (hybridization at the carbons forming covalent bonds to the Mo) not steric interactions with the axial-equatorial ligands. There is a considerable barrier to rotation of the \( \text{C}_{2} \text{H}_{5} \) ring in the metallatetrahedrane due to strong electronic preference for the eclipsed geometry.

**Acknowledgment.** Partial support from the National Science Foundation (Grant No. CHE83-18041) is gratefully acknowledged.

**Registry No.** 7a, 110661-53-5; 8, 110661-54-6.