

#### 4. A Class of Trinuclear Clusters with Carbonyl Bridging

by Allan R. Pinhas<sup>1a)</sup>, Thomas A. Albright<sup>1b)</sup>, Peter Hofmann<sup>1c)</sup> and Roald Hoffmann<sup>1a)</sup>

Department of Chemistry, Cornell University, Ithaca, New York 14853

University of Houston, Houston, Texas 77004

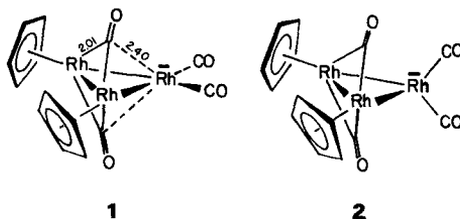
Institut für Organische Chemie, Universität Erlangen, Henkestrasse 42, D-8520 Erlangen,  
Federal Republic of Germany

(24.IX.79)

##### Summary

The impetus for this work was the structure of a trinuclear complex with two carbonyl groups showing incipient triple bridging -  $\text{Cp}_2\text{Rh}_3(\text{CO})_4^-$ . Its structure, barrier to rotation of one  $\text{Rh}(\text{CO})_2^-$  piece vs. the rest of the molecule, and the nature of the bridging carbonyl interaction are analyzed. Isolobal analogies form an interesting connection between this complex and a bridged isomer of the recently synthesized carbene complexes,  $\text{Cp}_2\text{Rh}_2(\text{CO})_2\text{CR}_2$ , one isomer of  $\text{Cp}_2\text{Rh}_3(\text{CO})_3$ , and hypothetical carbyne complexes  $\text{Cp}_2\text{Rh}_2(\text{CO})_2\text{CH}^{+,-}$ . A general bonding model for  $\text{Cp}_2\text{Rh}_2(\mu\text{-CO})_2\text{X}$  complexes is constructed. The model, rich in geometrical detail, allows minima for the bridging carbonyl groups bending toward and away from the bonded ligand X.

**Introduction.** - Interest and expediency, two of the motive forces of science, lay behind the initiation of this work. The interest was provided by the preparation and structure determination of  $\text{Cp}_2\text{Rh}_3(\text{CO})_4^-$  (**1**) by Bergman *et al.* [1]<sup>2)</sup>. We were intrigued by the semi-triply bridging disposition of two carbonyl groups in **1**. In transition metal clusters there exists a spectrum of terminally bonded, doubly bridging, semi-doubly bridging, and triply bridging carbonyl groups [2] [3]. The complex **1** is a so far unique example of a molecule caught between the doubly

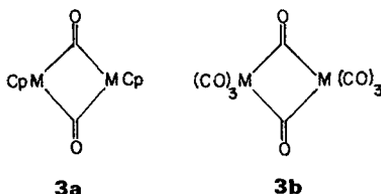


<sup>1a)</sup> Cornell University.

<sup>1b)</sup> University of Houston.

<sup>1c)</sup> Universität Erlangen.

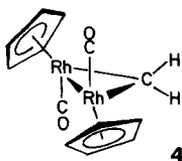
<sup>2)</sup> Throughout this paper we use the abbreviation Cp for an  $\eta^5$ -cyclopentadienyl ring.



bridging and triply bridging extremes. Moreover there is an interesting orientational choice made in **1** - the  $\text{Rh}(\text{CO})_2$  unit is coplanar with the two other Rh-atoms, rather than perpendicular, as in **2**, a choice that might have seemed better on steric grounds.

The expediency was grounded in a study we had just completed of  $\text{CpM}(\mu\text{-CO})_2\text{MCp}$  and  $(\text{CO})_3\text{M}(\mu\text{-CO})_2\text{M}(\text{CO})_3$  complexes (**3a** and **3b**) focusing on the planarity or puckering of the inner rhomboid and the metal-metal interaction [4]. The orbitals of a  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  were thus fresh in mind. That structural unit is obviously a piece of the trirhodium cluster **1**. It was a simple matter to think about constructing the orbitals of **1** or **2** from  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  and a  $\text{Rh}(\text{CO})_2^-$  fragment.

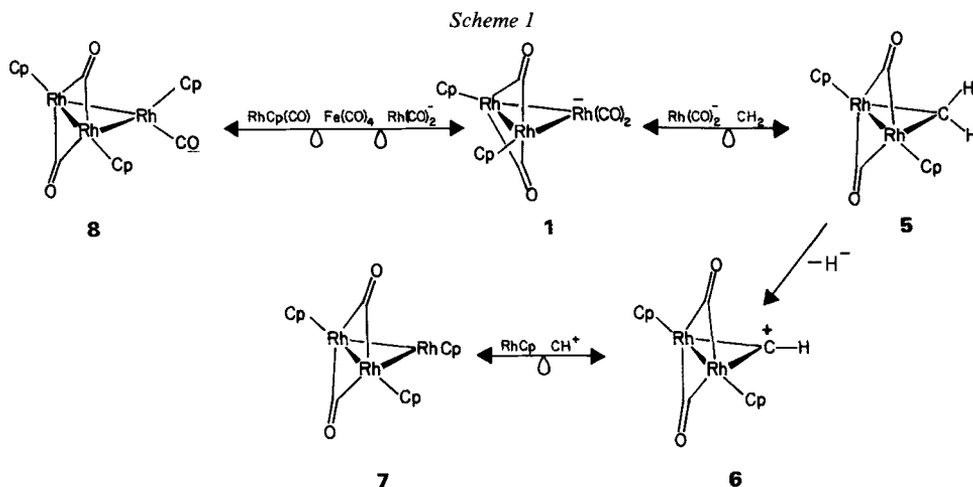
This was done, and the details will be given below. At this point another incentive appeared in the perception of a connection between  $\text{Cp}_2\text{Rh}_3(\text{CO})_4^-$  and a beautiful series of compounds prepared by *Herrmann's* group and structurally characterized by *Bernal et al.*, the carbene complexes of type **4** [5]. A second step took us to a hypothetical symmetrical  $\text{Cp}_3\text{Rh}_3(\text{CO})_2$  with two triply bridging carbonyl groups.



The chain of reasoning that we followed may be worth presenting at this point. We [6] and others [7] have established the utility of an isolobal analogy between the orbitals of transition metal carbonyl (or cyclopentadienyl) fragments and pieces of organic molecules. The analogy pairs  $\text{Mn}(\text{CO})_5$  with  $\text{CH}_3$ ,  $\text{Fe}(\text{CO})_4$  with  $\text{CH}_2$ ,  $\text{Co}(\text{CO})_3$  with  $\text{CH}$ . A cyclopentadienyl anion,  $\text{Cp}^-$ , obviously takes the place of three carbonyl groups, so further identifications of  $\text{Co}(\text{CO})_3$  with  $\text{NiCp}$ ,  $\text{Fe}(\text{CO})_4$  with  $\text{CoCp}(\text{CO})$  are possible. An extension of the analogy [6d] shows that the orbitals of a  $d^8\text{ML}_4$  are similar to those of a  $d^{10}\text{ML}_2$ . Let us apply several of these isolobal transformations to **1** (*Scheme 1*).

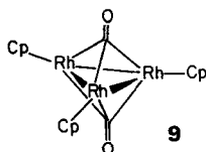
A  $d^{10}\text{ML}_2$  fragment is isolobal with methylene. This replacement leads us back from **1** to **5**. Compound **5** is not the observed structure of *Herrmann's* complexes, but could represent a bridged carbonyl group waypoint in a hypothetical carbonyl groups pathway in these molecules. This is apparently a high energy process [5c, d].

Let us next remove a hydride ion from the bridging methylene to give a hypothetical bridging carbyne complex,  $\text{Cp}_2\text{Rh}_2(\text{CO})_2\text{CH}^+$  (**6**). Isolobal replacement of

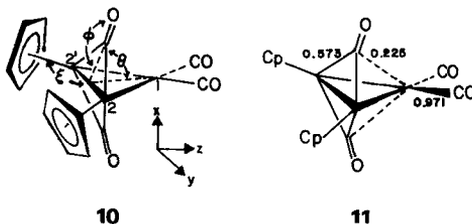


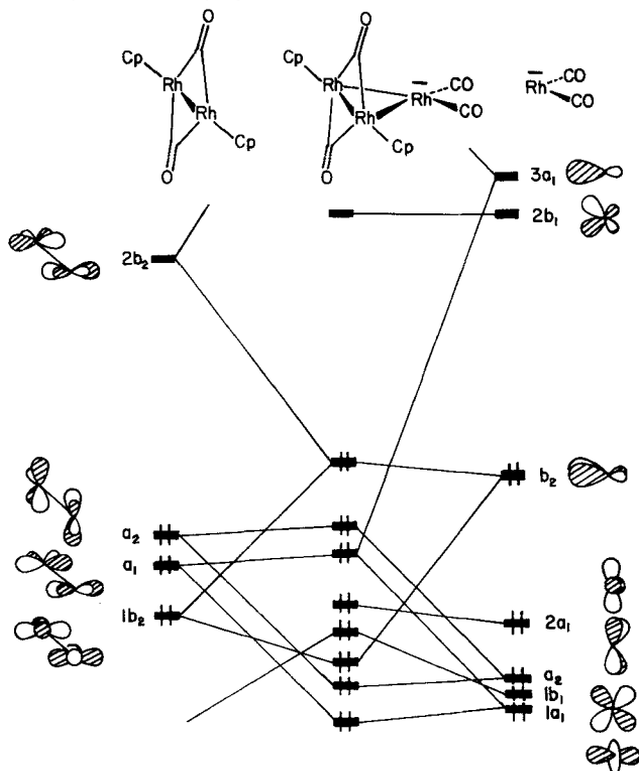
$\text{CH}^+$  by  $\text{RhCp}$  produces  $\text{Cp}_3\text{Rh}_3(\text{CO})_2$  (7). Another isolobal replacement, that of the  $d^{10}\text{Rh}(\text{CO})_2^-$  by  $\text{RhCp}(\text{CO})$  (both similar to a  $d^8\text{ML}_4$ ), leads one to the real structure of one isomer of  $\text{Cp}_3\text{Rh}_3(\text{CO})_3$  (8).

The  $\text{Cp}_3\text{Rh}_3(\text{CO})_2$  structure is drawn unsymmetrically in the scheme, to bring out its relationship to 1. However, one can inquire whether the structure can be symmetrized by having the carbonyl groups move into fully symmetrical triply bridging positions, as in 9. This compound is not known. Our calculations indicate a triply bridging structure for it, with some interesting possibilities for deformation.



$\text{Cp}_2\text{Rh}_3(\text{CO})_4^-$ . - A series of extended *Hückel* calculations, parameters specified in the Appendix, were carried out. In these we optimized independently the three variables shown in 10. The dihedral angle between the  $\text{Rh}_1\text{-Rh}_2\text{-Rh}_2'$  and the  $\text{Rh}_2\text{-C-Rh}_2$  planes is designated by  $\theta$ ; the bending of the Cp groups away from the y axis is given by  $\xi$ , the  $\text{Rh}_2\text{-Rh}_2'\text{-Cp}$  midpoint angle; and  $\varphi$  is the tilting of the bridging CO away from  $\text{Rh}_1$ . The optimized values were  $\theta = 74^\circ$ ,  $\xi = 157^\circ$ ,  $\varphi = 176^\circ$ . These agree quite well with the experimental values of  $\theta = 77.5^\circ$ ,  $\xi = 159.4^\circ$ ,  $\varphi = 170^\circ$  [1]. The Rh-C(O) overlap populations shown in 11 indicate significant



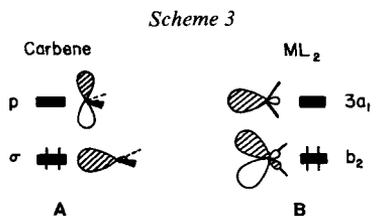
Scheme 2. Construction of the orbitals of  $Cp_2Rh_3(CO)_4^-$  from  $Cp_2Rh_2(CO)_2$  at left and  $Rh(CO)_2^-$  at right

interaction between the bridging carbonyl groups and  $Rh_1$ , although the other two types of  $Rh-C(O)$  bonding are substantially larger. When the two carbonyl groups are placed in a more idealized doubly bridging geometry,  $\theta = 90^\circ$ ,  $\xi = \varphi = 180^\circ$ , the  $Rh_1-C(O)$  overlap population is diminished to 0.083, while that for  $Rh_2-C(O)$  increases to 0.617.

Let us construct the orbitals of  $Cp_2Rh_2(CO)_4^-$  from those of  $Cp_2Rh_2(CO)_2$  and  $Rh(CO)_2^-$ , for simplicity taking a doubly-bridging reference geometry,  $\theta = 90^\circ$ ,  $\xi = \varphi = 180^\circ$ . This is done in *Scheme 2*. The valence orbitals of a  $d^{10}ML_2$  fragment analogous to  $Rh(CO)_2^-$  have been fully described elsewhere [6b, e] [8] [9]. Basically there is a nest of four low-lying orbitals,  $1a_1$ ,  $1b_1$ ,  $a_2$ , and  $2a_1$ , which are largely nonbonding. Their resemblance to the four occupied orbitals of a square-planar,  $d^8ML_4$  is obvious. At higher energy is  $b_2$ . This is comprised of metal  $yz$  and  $y^3$  in such a way that the orbital is hybridized out away from the carbonyl ligands. At still higher energy is  $2b_1$ , an orbital largely of carbonyl  $\pi^*$  character with some metal  $xz$  and  $x$  mixed in an antibonding and bonding manner, respectively.

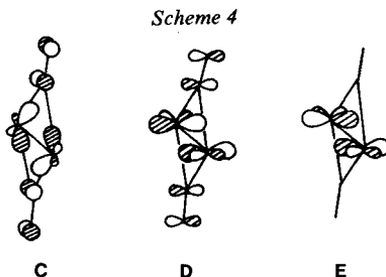
Because  $2b_1$  does not carry a heavy metal contribution, it will be much less active in  $\pi$  interactions with a ligand than  $b_2$ . Highest in the scheme is a well-

<sup>3)</sup> Throughout this paper we use the notation  $z^2$ ,  $x^2-y^2$ ,  $xy$ ,  $xz$ ,  $yz$  for the  $nd$  orbitals,  $s$ ,  $x$ ,  $y$ ,  $z$  for  $(n+1)s$  and  $p$ .



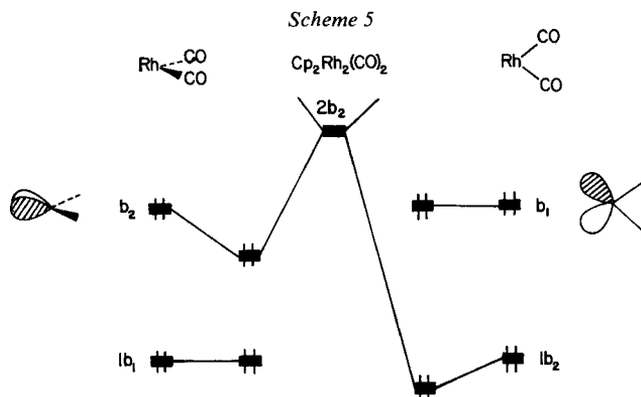
directed hybrid of metal  $s$ ,  $z$  and  $z^2$ . The orbitals of  $\text{Rh}(\text{CO})_2^-$  which figure in the isolobal analogy to methylene, *i.e.* those which are analogous to a  $\text{CH}_2$   $\sigma$  and  $p$  orbital [10] are  $3a_1$  and  $b_2$  respectively. This is shown below in **A** and **B** (Scheme 3). Note the energy ordering in the metal fragment differs from that in the carbene.

The orbitals of  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  are a bit more complicated. The reader is referred to our detailed analysis [4], while here we sketch the outlines of the story. Two  $\text{MCp}$  fragments, each with three low-lying orbitals and three high-lying directed hybrids, combine to give a nest of six low-lying levels and six higher valence orbitals. One of the latter six, the  $\text{MM}\sigma^*$ , is very high in energy. The bridging carbonyl lone pairs destabilize two of the remaining five valence orbitals. There remain behind the six low-lying metal centered orbitals (the highest of these,  $1b_2$ , appears in Scheme 2) and three valence orbitals,  $a_1$ ,  $a_2$  and  $2b_2$ . In the  $d^8$ - $d^8\text{Cp}_2\text{Rh}_2(\text{CO})_2$  fragment the six lower orbitals plus  $a_1$  and  $a_2$  are occupied. There is a great differentiation in energy between  $a_2$  and  $a_1$ , both stabilized by bridging carbonyl  $\pi^*$  mixing, as shown in **C** and **D**, and  $2b_2$ , **E** (Scheme 4), which does not have the right symmetry for that stabilization.



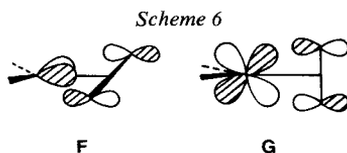
Application of the eighteen electron rule leads one to a Rh-Rh double bond in this fragment. We don't find much metal-metal bonding, a typical result for carbonyl group bridged systems where the bonding in the bridging region is highly delocalized [6c] [11]. But note that the  $2b_2$  orbital of  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  is clearly a  $\pi^*$  orbital, available for acceptance of electrons from another fragment.

This  $\pi$  interaction is in fact the major bonding factor in the trinuclear complex (see Scheme 2). The interaction is slightly complicated by the further involvement of the  $1b_2$  orbital of the  $\text{Rh}_2$  fragment. These three orbitals ( $1b_2$ ,  $2b_2$  of  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$ ,  $b_2$  of  $\text{Rh}(\text{CO})_2^-$ ) interact, producing three molecular orbitals - one at low energy, one high (not shown in the Scheme 2), one nonbonding. The two lower orbitals are filled, producing a net stabilization. Further bonding is provided by donor  $a_1$  of  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  and acceptor  $3a_1$  of  $\text{Rh}(\text{CO})_2^-$ .

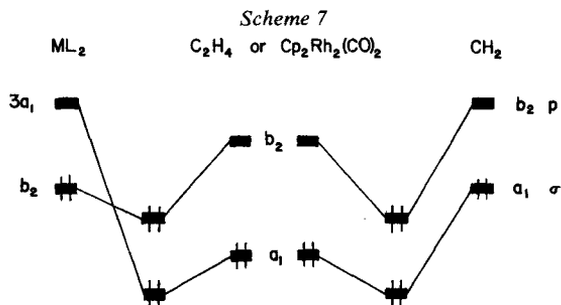


The way is now clear to an understanding of the conformational preference for the observed geometry **1** over **2**. An interaction diagram similar to that of *Scheme 2* but with the  $\text{Rh}(\text{CO})_2^-$  fragment rotated by  $90^\circ$  could be constructed. The orbitals remain, of course, the same, but in that  $\text{ML}_2$  fragment the symmetry labels  $b_1$  and  $b_2$  switch. The net result, sketched schematically in *Scheme 5*, is dramatic. The  $\text{ML}_2$  fragment donor orbital,  $b_2$  in orientation **1**, becomes uncoupled from the bonding in conformation **2**, since it is now  $b_1$ . Its place in the interaction with  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  is taken by a more distant and less effectively overlapping lower energy orbital. In the observed trimer geometry we calculate a rigid rotation barrier of 21 kcal/mol.

What we have described here for the conformational preference in  $\text{Cp}_2\text{Rh}_3(\text{CO})_4^-$  is analogous to what occurs in the rotational barrier of  $d^{10}$  ethylene- $\text{ML}_2$  complexes [6d]. The ethylene  $\pi^*$  orbital has a choice of interacting with either  $b_2$  in a  $d^{10}\text{ML}_2$  fragment, isoelectronic with  $\text{Rh}(\text{CO})_2^-$ , as in **F** or  $b_1$  in **G** (*Scheme 6*). The geometry **F** is greatly favored for the same reasons as those that were described in the previous paragraph: larger overlap and smaller energy gap. The relationship between  $\text{Cp}_2\text{Rh}_3(\text{CO})_4^-$  and ethylene- $\text{ML}_2$  does not stop here. The bonding in both molecules could be described in terms of a classical *Dewar-Chat-Duncanson* scheme (*Scheme 7*) in which  $3a_1$  and  $\sigma$  on the one hand and  $b_2$  and  $p$ , on the other, play analogous roles. The bonding in both complexes is entirely analogous to that in cyclopropane. At this point the reader might also compare our previous analysis of the  $\text{L}_4\text{Pt}_2\text{CR}_2$  system [6e].

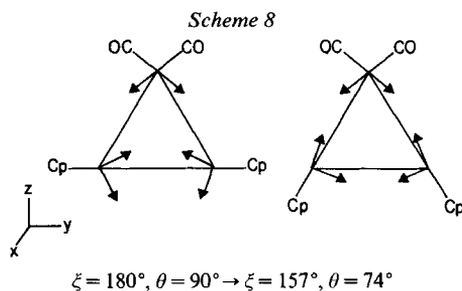


The actual geometry of the  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  fragment in the  $\text{Rh}_3$  complex differs substantially from the idealized structure with  $\theta = 90^\circ$ ,  $\xi = \varphi = 180^\circ$  in which the  $\text{CpRhRhCp}$  axis is linear and the bridging carbonyl groups do not lean toward the third rhodium. Bending the carbonyl groups over (reducing  $\theta$  and  $\varphi$ ) does not decrease the total energy. Stabilization ensues upon simultaneous bending back

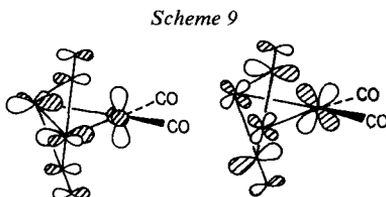


of the Cp rings (reducing  $\xi$ ). Motion of the Cp units is inextricably coupled to the puckering of the bridging carbonyl rhombus. We have examined this motion previously for the  $\text{Cp}_2\text{M}_2(\text{CO})_2$  complexes themselves [4], and will have something further to say about it below.

For the trimer the consequences of puckering, which involves simultaneous reduction of  $\xi$  and  $\theta$ , has important stabilizing consequences. These enter through an energy lowering of the  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  fragment  $2b_2$  orbital (through mixing of carbonyl  $\pi^*$ 's, absent in the planar rhomboid, turned on with puckering [4]) and a reorientation of that orbital for better overlap with the  $\text{Rh}(\text{CO})_2^-$  fragment. The latter factor is shown schematically in Scheme 8 where the main lobes of the interacting  $b_2$  orbitals are drawn as arrows. The important  $b_2$  coupling overlap increases from 0.120 to 0.205.



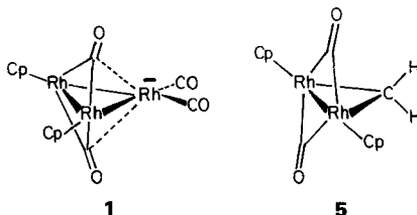
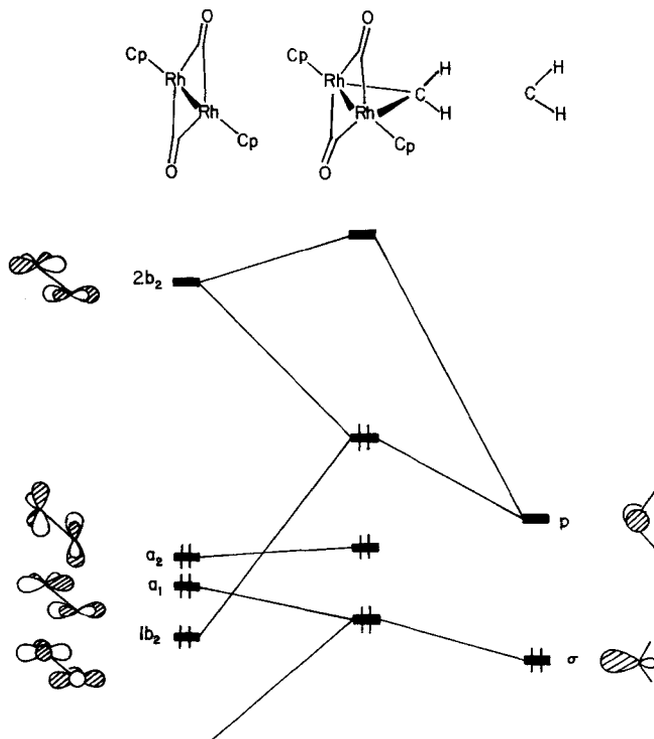
We return to the question of incipient triple bridging of the carbonyl groups. As  $\theta$  is decreased there are several stabilizing interactions turned on between the bridging carbonyl group and  $\text{Rh}_1$ . Energetically this effect is smaller than that described previously. It is also partially masked by intermixing between the many lower-lying orbitals of the same symmetry which have been left out of Scheme 2 and slight destabilization of the dimer levels themselves with the distortion. How-



ever, a fragment analysis allows a partial tracing of the interaction. Significant portions of it are to be found in the  $a_1$  and  $b_2$  orbitals shown in *Scheme 9*.  $\text{Rh}(\text{CO})_2^-$  lone pair interactions with bridging  $\text{CO } \pi^*$ , not the reverse charge transfer from carbonyl  $\pi$  to the 16-electron Rh-center, are implicated. We will return to the general problem of semi-bridging carbonyl groups in a separate paper.

**Carbene complexes.** - The introduction indicated succinctly how the isolobal analogy connects a hypothetical carbonyl bridged isomer of *Herrmann's* carbene complexes **5** to *Bergman's* compound **1**. The correspondence carries over in substantial detail, as *Scheme 11*, an interaction diagram for  $\text{Cp}_2\text{Rh}_2(\text{CO})_2\text{CH}_2$  constructed from  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  and  $\text{CH}_2$ , shows.

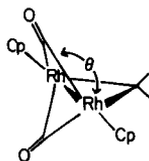
Scheme 10

Scheme 11. Construction of orbitals of  $\text{Cp}_2\text{Rh}_2(\text{CO})_2(\text{CH}_2)$  from  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  at left and  $\text{CH}_2$  at right

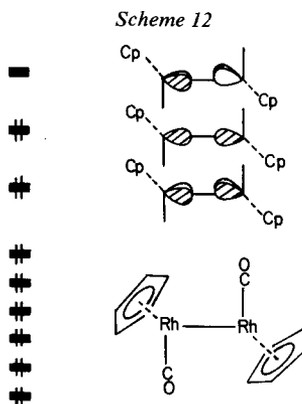
Even though the carbene begins with its  $\sigma$  orbital below the p, it is the bonding combination of carbene p with  $\text{Cp}_2\text{Rh}(\text{CO})_2$   $2b_2$  that is occupied. The three highest occupied levels in *Scheme 11*,  $b_2$ ,  $a_2$ ,  $a_1$ , are very similar to the three highest levels in *Scheme 2*, the  $\text{Rh}(\text{CO})_2^-$  case. The isolobal analogy is a very strong one.

Some differences emerge. When the flap angle  $\theta$  (see **10**) is allowed to vary, the carbonyl groups bend away from the  $\text{CH}_2$  group (see **12**) quite opposite to what happened in the *Bergman's* compound, where the carbonyl groups bent toward the  $\text{Rh}(\text{CO})_2^-$ . In the methylene complex  $\theta$  optimizes at  $115^\circ$ . This value is quite insensitive to the Cp orientation, as measured by  $\xi$ .

We now have a curious feature. The bridging carbonyl groups bend one way in the  $\text{Rh}(\text{CO})_2^-$  complex, the other way in the methylene compound. Why that is so will be discussed in the next section.

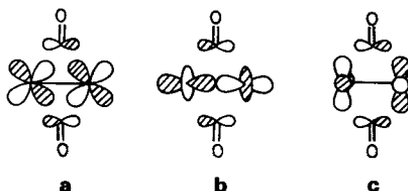

**12**

Let us not forget that the *Herrmann's* compounds in fact do not have bridging carbonyl groups, but terminal ones (**4**). What is their electronic structure, and how is it related to the hypothetical bridging isomer discussed by us up to this point? The electronic structure of unbridged  $\text{Cp}_2\text{Rh}_2(\text{CO})_2(\text{CH}_2)$  has been treated in detail by one of us [12]. It is appropriate to write for the  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  fragment of it a double bonded structure, for above a nest of six metal centered orbitals the unbridged dimer has occupied levels of  $\pi$  and  $\sigma$  symmetry (as well as a low-lying  $\pi^*$  level), as shown in *Scheme 12*.



The unbridged  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  may have a nice double bond, but it is unstable with respect to bridging. There are a couple of ways to see this. First, second-order *Jahn-Teller* considerations [13] turn our attention to the low-lying  $\pi^*$  orbital in the unbridged form. Second a detailed examination of the levels on each side shows a

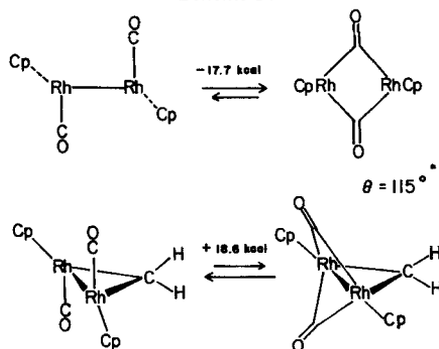
Scheme 13



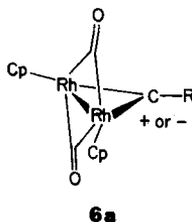
general stabilization on the bridged side through carbonyl stabilization of formally metal-metal antibonding orbitals such as **a**, **b**, **c** in *Scheme 13*. A detailed discussion will be given elsewhere [12b]. Here it will suffice to say that the bridged form is calculated to be more stable by some 21.7 kcal/mol when  $\theta = 90^\circ$ , 17.7 kcal/mol for  $\theta = 115^\circ$ .

While the unbridged  $\text{Cp}_2\text{Rh}_2$  fragment is the less stable one, it is far better set up to bind a methylene than the bridged isomer. To a great extent this is the consequence of the low-lying  $\pi^*$  orbital (see *Scheme 12*) so beautifully set up to interact with a carbene p. The unbridged  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  interaction with  $\text{CH}_2$  is not shown in detail here [12], but it is a lovely case of metallocyclopropane formation. The way the energetics of bridging-unbridging is coupled with carbene addition is summarized in *Scheme 14*.

Scheme 14

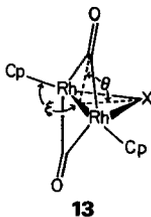


**Carbyne complexes.** - While **6** served just as a waypoint in an isolobal analogy in *Scheme 1*, perhaps it should be given more serious consideration, for other carbyne complexes are known. We have computed a potential energy surface for carbonyl bending in both the cation and anion of  $\text{Cp}_2\text{Rh}_2(\text{CO})_2\text{CH}$  (**6a**).

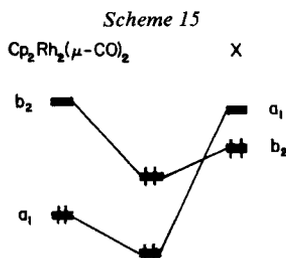


The cation is relatively undistorted,  $\theta = 175^\circ$  (see **10** for the definition of the angle). The anion shows a rather different picture. In a curious mixture of the behaviour of the  $\text{Rh}(\text{CO})_2^-$  and  $\text{CH}_2$  complexes, the hypothetical  $\text{Cp}_2\text{Rh}_2(\text{CO})_2(\text{CH})^-$  has two well defined minima. In one the carbonyl bridges bend toward the  $\text{CH}^-$ ,  $\theta = 70^\circ$ ,  $\zeta = 155^\circ$ ; in the other they bend away,  $\theta = 120^\circ$ ,  $\zeta = 175^\circ$ . These geometries are 4.5 kcal/mol apart in energy, with the one bent away from the carbonyl groups being more stable. Secondary interactions between the carbanion lone pair and bridging carbonyl  $\pi^*$  orbitals are clearly identifiable in the low  $\theta$  minimum. There is high  $\text{C}(\text{O})-\text{C}(\text{H})$  overlap population of +0.62 there. The barrier between the minima is approximately 13 kcal/mol. It would be most interesting to obtain such a carbyne complex and explore its structure and dynamics.

**A general bonding pattern for  $\text{Cp}_2\text{Rh}_2(\mu\text{-CO})_2\text{X}$  complexes.** - So far we have discussed several compounds, one real one with  $\text{X} = \text{Rh}(\text{CO})_2^-$ , and several hypothetical ones with  $\text{X} = \text{CH}_2$  or  $\text{CH}^+$ ,  $\text{CH}^-$ . Still another known species remains,  $\text{X} = \text{RhCp}(\text{CO})$ , but even before we discuss its structure we must face up to the curious geometrical conclusion of the previous sections, namely that for  $\text{X} = \text{Rh}(\text{CO})_2^-$  the bridging carbonyl groups bend toward  $\text{X}$ ,  $\theta < 90^\circ$ , while for  $\text{X} = \text{CH}_2$  the bridge is calculated to bend away from  $\text{X}$ ,  $\theta > 90^\circ$  (see **13** for a reminder of the definition of the angles).



The explanation of this phenomenon is to be found in a general analysis of the bonding of a  $\text{Cp}_2\text{Rh}_2(\mu\text{-CO})_2$  fragment to a fragment  $\text{X}$ , where  $\text{X} = \text{Rh}(\text{CO})_2^-$ ,  $\text{CH}_2$ ,  $\text{Fe}(\text{CO})_4$ ,  $\text{RhCp}(\text{CO})$  or an isolobal group. As we have already said, the bonding pattern carries the usual *Dewar-Chatt-Duncanson* features of metal-ligand interaction, namely forward and back donation, involving orbitals of  $a_1$  and  $b_2$  symmetry (*Scheme 15*).



When we analyze the carbonyl bending in our two  $\text{Cp}_2\text{Rh}_2(\text{CO})_2\text{X}$  complexes, much of the driving force for bending is to be found in the HOMO,  $b_2$ . It is stabilized as  $\theta$  increases or decreases from  $90^\circ$ . Why? The reasons are discussed in detail in our earlier paper [4]. At  $\theta = 90^\circ$ , a planar  $\text{M}_2(\text{CO})_2$  rhomboid, this orbital

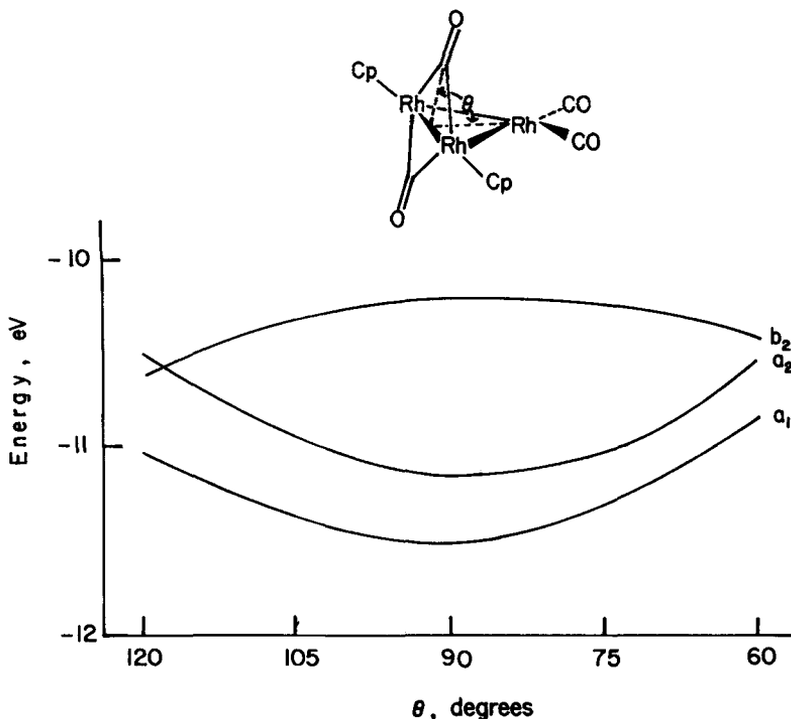


Fig. 1. Frontier orbitals of  $\text{Cp}_2\text{Rh}_3(\text{CO})_4^-$  as a function of  $\theta$ . Both bridging carbonyls are bending.

cannot interact with the carbonyl  $\pi^*$ 's by symmetry. As the rhomboid puckers this favorable, stabilizing interaction becomes increasingly important.

In the  $d^8-d^8\text{Cp}_2\text{Rh}_2(\text{CO})_2$  this  $b_2$  orbital, which favors bending, is *not* occupied. But, as *Scheme 2* and *11* clearly show, interaction with  $\text{Rh}(\text{CO})_2^-$  or  $\text{CH}_2$  populates this orbital. *Figure 1* indicates the evolution of the higher occupied orbitals in  $\text{Cp}_2\text{Rh}_3(\text{CO})_4^-$  at  $\xi = 180^\circ$  as  $\theta$  is varied. Note how the  $b_2$ , which favors puckering, is stabilized approximately equally as  $\theta$  departs to greater or smaller angles than  $90^\circ$ . The lower orbitals counteract this trend.

The case at hand,  $\text{X} = \text{Rh}(\text{CO})_2^-$ , is perturbed by further steric (Cp-carbonyls on unique Rh) and electronic (bridging CO-unique Rh) interactions. But quite generally if these secondary interactions are minimized we expect the primary potential energy curve for puckering in  $\text{Cp}_2\text{Rh}_2(\text{CO})_2\text{X}$  to resemble that for a  $\text{Cp}_2\text{M}_2(\text{CO})_2$  fragment with the  $b_2$  orbital occupied, *i.e.* a broad flat minimum in a deformation coordinate in which  $\theta$  and  $\xi$  are coupled. This is the dashed line in *Figure 2*.

In a hypothetical partitioning of interactions, the next factor to consider is the electronic role of the X group. As it interacts with  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$  it will seek out maximum overlap with the  $a_1$  and  $b_2$  orbitals of the binuclear fragment. The directionality or hybridization of  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$   $a_1$  and  $b_2$  is a function of  $\xi$ , as shown below in *Scheme 16*, and as we discussed earlier. This will shift the minimum of *Figure 2* to smaller  $\theta$  and  $\xi$ , the solid line.

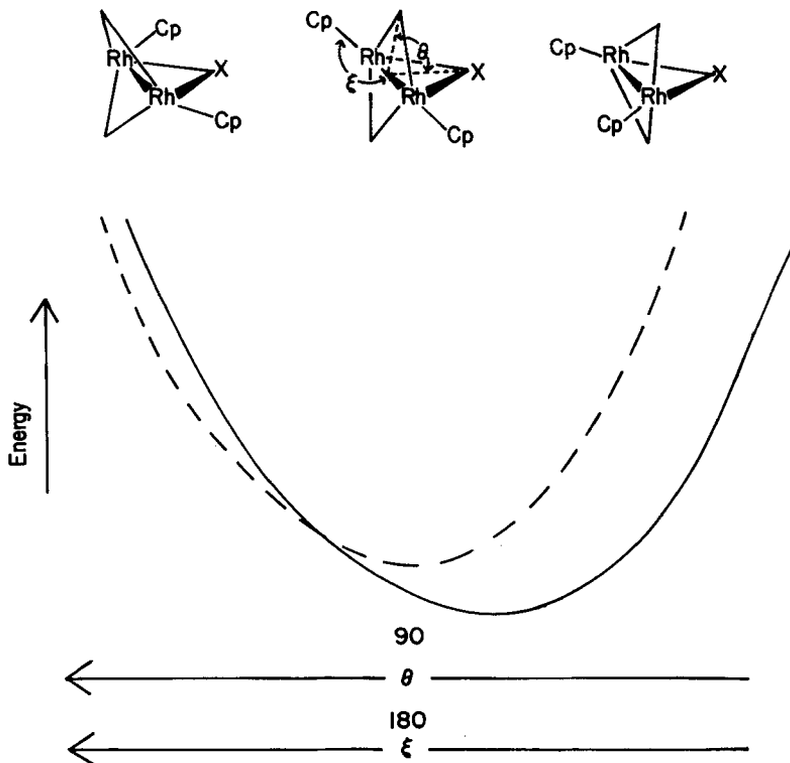
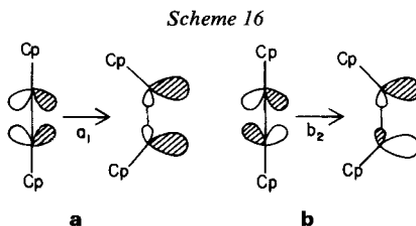


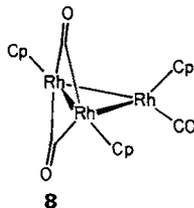
Fig. 2. A schematic potential energy surface for carbonyl and cyclopentadienyl bending in  $\text{Cp}_2\text{Rh}_2(\text{CO})_2\text{X}$  complexes. The dashed line is the hypothetical curve resulting from simple transfer of electrons to the  $b_2$  orbital, while the solid line represents schematically the result of including rehybridization of  $a_1$  and  $b_2$  for better bonding to X.  $\theta$  and  $\xi$  increase to the left.



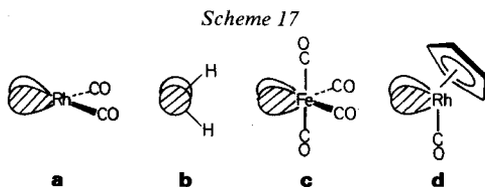
Superimposed upon this basic picture are weak attractive (secondary orbital interaction) and variable, often strong repulsive (steric) factors. These are highly encumbered geometries, and relatively small variations of  $\theta$  and  $\xi$  bring on unacceptable steric contacts. In the case of  $\text{X}=\text{CH}_2$  a decrease of  $\theta$  brings the bridging carbonyl groups too close to the methylene. The minimum shifts to  $\theta > 90^\circ$ . The hypothetical  $\text{Cp}_2\text{Rh}_2(\text{CO})_2\text{CH}^-$  has transformed the deformable surface into a double minimum curve with shallow wells, the one for small  $\theta$  created by secondary interactions between  $\text{CH}^-$  and the bridging carbonyls. For  $\text{X}=\text{Rh}(\text{CO})_2^-$  the region

of small  $\theta$  and  $\xi$  is doubly deepened, by the secondary interactions between the bridging carbonyls and the unique Rh, and by the severe steric difficulties for  $\xi > 180^\circ$ . We will soon see another example of the steric effects at work.

**Cp<sub>3</sub>Rh<sub>3</sub>(CO)<sub>3</sub>.** - Another compound obtained by isolobal replacement of Rh(CO)<sub>2</sub> in **1** by RhCp(CO) is Cp<sub>3</sub>Rh<sub>3</sub>(CO)<sub>3</sub> (**8**). There are in fact two compounds of this stoichiometry obtained from the photolysis of CpRh(CO)<sub>2</sub>. One has a symmetric structure with three bridging carbonyl groups [14], the other geometry **8** [15].



At first sight it seems that the RhCp(CO) fragment orientation is different from that of Rh(CO)<sub>2</sub>. So it is, but in fact the orientation is entirely consistent with the isolobal analogy. Recall that the crucial bonding feature in these Cp<sub>2</sub>Rh<sub>2</sub>(CO)<sub>2</sub>X molecules is the interaction with a<sub>1</sub> and b<sub>2</sub> orbitals of the X group. The b<sub>2</sub> interaction sets the orientational preference. The low-lying b<sub>2</sub> orbitals of Rh(CO)<sub>2</sub> and CH<sub>2</sub> are shown in *Scheme 17* (a and b). RhCp(CO) is isolobal with Fe(CO)<sub>4</sub>. The latter has a well-known valence active b<sub>2</sub> orbital (*Scheme 17*, c [5d]), and, not surprisingly, so does the d<sup>8</sup>MCp(CO) fragment **d** (*Scheme 17*, [12]). We have deliberately oriented all of these orbitals in the same way, the mode in which they should enter bonding with a Rh<sub>2</sub>Cp<sub>2</sub>(CO)<sub>2</sub> fragment coming from the left.



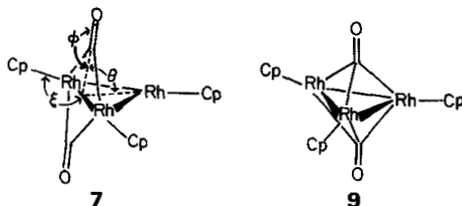
So the observed geometry of isomer **8** of Cp<sub>3</sub>Rh<sub>3</sub>(CO)<sub>3</sub> is, in fact, a consequence of an electronic fit. Calculations on **8** confirm this. An estimate of the barrier to rotation of the RhCp(CO) fragment is difficult to obtain because of steric crowding of Cp groups in the Rh<sub>3</sub> plane in the rotated form. We calculate a barrier of 36 kcal/mol even when the positions of the Rh<sub>2</sub> Cp rings are adjusted.

In the observed structure of **8** the Cp<sub>2</sub>Rh<sub>2</sub>(CO)<sub>2</sub> carbonyl groups are bent away from the third rhodium ( $\theta = 95^\circ$ ), in contrast to the Cp<sub>2</sub>Rh<sub>3</sub>(CO)<sub>4</sub><sup>-</sup> structure. We are reasonably certain that this is a steric effect, that decreasing  $\theta$  below  $90^\circ$  is precluded by crowding between the bridging carbonyl groups and the CO and Cp on the third rhodium.

Our knowledge of the bridging tendencies in these molecules is far from complete. Cp<sub>3</sub>Rh<sub>3</sub>(CO)<sub>3</sub> opts for the two structures mentioned above. Cp<sub>3</sub>Co<sub>3</sub>(CO)<sub>3</sub> finds its

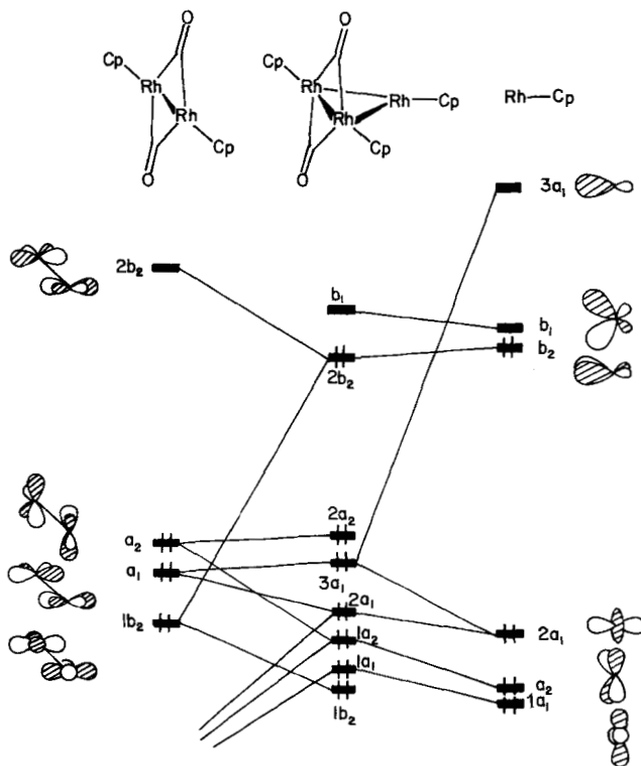
equilibrium geometry in the solid state in a third isomer, which has two doubly bridging carbonyl groups (on distinct edges), and one triply bridging CO [16]. Yet in solution its infrared spectrum is indicative of structure **8a** [16].

**Cp<sub>3</sub>Rh<sub>3</sub>(CO)<sub>2</sub> and other hypothetical molecules.** - A second application of the isolobal analogy led us, in the Introduction, to an initially unsymmetrical Cp<sub>3</sub>Rh<sub>3</sub>(CO)<sub>2</sub> cluster, which has the appealing possibility of achieving three-fold symmetry through a small motion of the carbonyls into a fully triply bridging geometry, **7** → **9**. A discussion of this pathway follows.



The interaction diagram for an unsymmetrical trimer,  $\theta = 90^\circ$ ,  $\xi = 180^\circ$ ,  $\varphi = 180^\circ$  is given in *Figure 5*. The orbitals of an MCp fragment, isolobal to M(CO)<sub>3</sub>, have been discussed previously [6a, c]. They consist of three low-lying levels,  $1a_1$ ,  $a_2$ ,  $2a_1$ ,

Scheme 18. Construction of the orbitals of Cp<sub>3</sub>Rh<sub>3</sub>(CO)<sub>2</sub> from Cp<sub>2</sub>Rh<sub>2</sub>(CO)<sub>2</sub> at left and CpRh at right



remnants of the octahedral  $t_{2g}$  set, and three high-lying levels,  $b_2$ ,  $b_1$  and  $3a_1$ , which are hybridized away from the Cp unit. Upon interaction the lower set of three are destabilized slightly by lower  $Cp_2Rh_2(CO)_2$  orbitals.  $a_2$ ,  $a_1$ ,  $1b_2$  and  $2b_2$  in the dimer fragment interact in a familiar way, as we have seen previously for  $Cp_2Rh_3(CO)_4$  and  $Cp_2Rh_2(CO)_2CH_2$ . The nonbonding molecular orbital labelled  $2b_2$  is the HOMO of the complex. The  $b_1$  level of the CpRh fragment is now high-lying. It forms a three orbital interaction with  $b_1$ , a primarily carbonyl  $\pi^*$  level, and its bonding counterpart, not shown in the diagram. Note that  $b_1$  remains a low-lying LUMO.

Next we studied the approach to triply bridging. A *Walsh* diagram for a symmetrical cut of the potential surface varying  $\theta$ ,  $\xi$  and  $\phi$  is presented in *Figure 3*. As  $\xi$  decrease the interaction between  $2b_2$  of the dimer and  $b_2$  of CpRh increases and the HOMO  $2b_2$  drops. Also as the bridging carbonyl groups move to the triply bridging positions, a strong stabilization in the CpRh  $b_1$  orbital is turned on. Thus

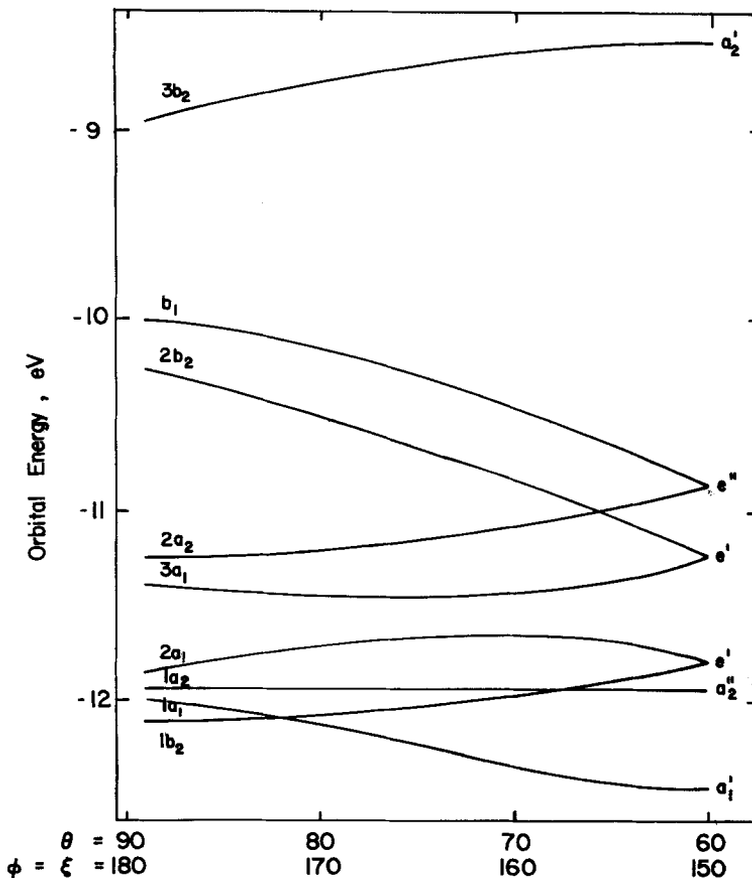


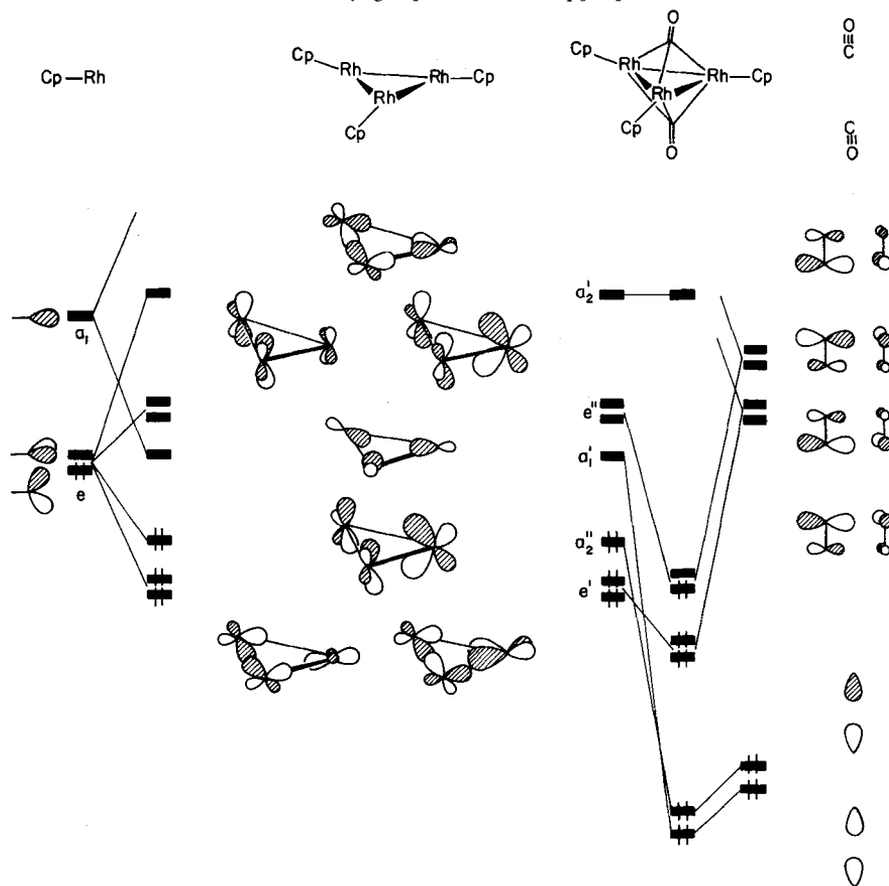
Fig. 3. A *Walsh* diagram for an approach to the symmetrical triply bridging structure of  $Cp_3Rh_3(CO)_2$ . The starting point, at left, is the doubly bridging structure with  $\theta = 90^\circ$ .

the molecular  $b_1$  level also falls.  $2a_2$  rises in energy because some of the carbonyl  $\pi^*$  bonding is lost. Finally  $3a_1$  remains relatively constant in energy until the later stages of distortion where it undergoes an avoided crossing with  $2a_1$ .  $1a_1$  falls in energy because its interaction with lower orbitals of the dimer decrease as the Cp rings are canted back. The other valence levels of the molecule do not change much along this path.

The effective symmetry of the triply bridging  $\text{Cp}_3\text{Rh}_3(\text{CO})_2$  is  $D_{3h}$  even though the maximal real symmetry is  $C_{3v}$ . The levels at the right side of *Figure 3* are labelled according to  $D_{3h}$ .

The constitution of the trimer orbitals is better seen from an alternative construction, which begins from a three-fold symmetric  $(\text{CpRh})_3$  triangle, and adds two carbonyl groups. This is done in *Scheme 19*. At left are the seven valence orbitals of  $(\text{CpM})_3$  lying above a nest of nine metal orbitals, which are not shown in the scheme, but which have room for 18 electrons. The  $\text{Cp}_3\text{M}_3$  system is isolobal to

Scheme 19. At left the orbitals of a symmetrical  $\text{Cp}_3\text{Rh}_3$  are built up from three CpRh units, at right two carbonyl groups are added to  $\text{Cp}_3\text{Rh}_3$

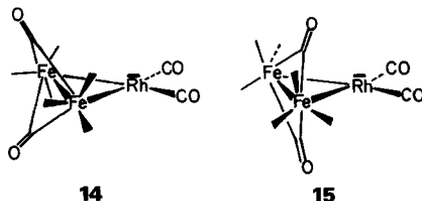


$M_3(CO)_9$  and the reader is referred to our paper on that system [17] for a detailed discussion of the  $(M(CO)_3)_3$  or  $(MCp)_3$  orbitals as they are built up from three  $M(CO)_3$  or  $MCp$  units. Two capping carbonyl groups are brought in from right. Of the seven starting orbitals of the  $Rh_3$  fragment two,  $a_2'$  and  $a_1'$ , are destabilized by the symmetric and antisymmetric carbonyl lone pair combinations. Four orbitals,  $e'$  and  $e''$ , are stabilized by carbonyl  $\pi^*$ . This leaves us with  $a_2'$ , which does not find a partner among the carbonyl orbitals. There are two electrons in each  $CpRh$  e set (over and above a  $d^6$  count) giving a total of six electrons for the valence orbitals in  $Cp_3Rh_3(CO)_2$ . Four go into  $e'$ , leaving two for  $e''$ .

Thus the  $D_{3h}$  triply bridged  $Cp_3Rh_3(CO)_2$  is an open shell complex. The molecule itself is not known, but there are some related species. With three electrons more, one reaches the known paramagnetic  $Cp_3Ni_3(CO)_2$  [18]. A diamagnetic molecule with two electrons more is  $(C_6H_6)_3Co_3(CO)_2^-$  [19]. Related species with triply bridging chalogens are  $Cp_3Co_3S_2$  [20], its cation [21],  $Fe_3(CO)_9As_2$  [22],  $Fe_3(CO)_9Se_2$  [23],  $Fe_3(CO)_9S_2$  [24] and  $Cp_3Co_3(CO)(O)$  [25].

As a consequence of its electronic structure, two electrons in a degenerate orbital,  $Cp_3Rh_3(CO)_2$  has several options open to it. It could remain a symmetrical high-spin species, as does the  $Cp_3Co_3S_2$  system with four electrons less [20]. A second possibility is deformation of the Cp rings, such as that observed in a number of *Jahn-Teller* active metallocenes [26]. The degeneracy of  $e''$  could be lifted in a number of ways. One metal-metal bond may stretch or contract, as in the  $Fe_3(CO)_9X_2$  structures [22-24]. Alternatively the molecule may pause somewhere between the doubly and triply bridging structures, while keeping the metal-metal separations approximately constant, an excursion along the reaction coordinate of *Figure 3*. We have explored this last possibility, optimizing  $\theta$ ,  $\varphi$  and  $\zeta$  independently. The energy is minimized at  $\theta = 65^\circ$ ,  $\varphi = 158^\circ$ ,  $\zeta = 150^\circ$  (the  $C_{3v}$  structure is defined by  $\theta = 60^\circ$ ,  $\varphi = 150^\circ$ ,  $\zeta = 150^\circ$ ). This geometry is only 2.4 kcal/mol below the symmetrical triply bridging structure and has a gap between filled and unfilled levels of 0.4 eV. We have not optimized the metal separations, and it could well be that the molecule, when it is made, will combine in its structure several of these distortions.

Isolobal replacements can be applied in the dirhodium fragment as well. For instance, let us consider the substitution of two  $RhCp$  units by  $Fe(CO)_3$ . Two possible conformations of  $Fe_2Rh(CO)_{10}^-$ , **14** and **15**, were considered.



Based on our previous work [4], we realized that the tendency for the bridging carbonyl groups to pucker should be much larger in the  $Fe_2(CO)_8$  fragment itself compared with  $Cp_2Rh_2(CO)_2$ . Furthermore, the sense of the puckering is tied to the orientation of the terminal carbonyl groups. The bridging carbonyl groups distort so as to stagger the terminal carbonyl groups. Optimization of  $\theta$ , defined

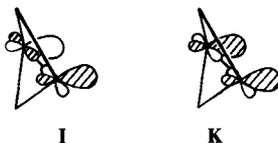
in an analogous manner to that in **10**, was consistent with this idea. For **14**  $\theta$  was found to be  $109^\circ$  while for **15** it was  $74^\circ$ , similar to that in the triply bridging  $\text{Cp}_2\text{Rh}_3(\text{CO})_4^-$ . The Rh-C overlap population for the bridging carbonyl group was again quite large - 0.266 in **15**.

While the  $\text{Rh}_3$  system bent the bridging carbonyl groups toward the unique Rh, this does not happen here. The stability of **15** is calculated to be 59 kcal/mol less than that of **14**. The reason behind this is fairly simple. There is a relatively low-lying  $b_2$  orbital **I** (Scheme 21), in the  $\text{Fe}_2(\text{CO})_6$  fragment, analogous to  $2b_2$  in  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$ . It is hybridized away from the bridging carbonyl groups [4]. Therefore, its overlap is much larger when  $b_2$  of  $\text{Rh}(\text{CO})_2^-$  is brought in from the right side of **I** (Scheme 20) rather than the left. There is also a high-lying filled orbital in  $\text{Fe}_2(\text{CO})_8$  of  $a_1$  symmetry (**K** in Scheme 21), which is analogous to  $a_1$  of  $\text{Cp}_2\text{Rh}_2(\text{CO})_2$ . It is hybridized like the  $b_2$  and interacts with the empty  $3a_1$  of  $\text{Rh}(\text{CO})_2^-$  much more strongly in conformation **14** than in **15**.

Scheme 20



Scheme 21



A simple way to view the hybridization in these two orbitals is the following: Consider the  $\text{Fe}_2(\text{CO})_8$  fragment as two coupled  $\text{ML}_5$  units and disregard any potential Fe-Fe bond [27]. Each  $\text{ML}_5$  unit will have an orbital basically derived from one member of the  $e_g$  set in octahedral  $\text{ML}_6$  [6]. A top view of the linear combination of these two orbitals is shown in Scheme 20. The interaction of  $\text{Rh}(\text{CO})_2^-$  in conformation **14** restores the octahedron with the formation of one Rh-Fe bond at each  $\text{ML}_5$  unit. It is a brief step from here to the orbitals of  $\text{Fe}_3(\text{CO})_{12}$ . All one has to do is to bring in an  $\text{Fe}(\text{CO})_4$ , isolobal to  $\text{Rh}(\text{CO})_2^-$ .

We have also examined a series of complexes based on  $\text{Cp}_2\text{Rh}_3(\text{CO})_4^-$  in which the bridging carbonyl groups are replaced by SR,  $\text{PR}_2$  and halide bridges. There are some comprehensible changes in the bonding pattern and conformational preferences in these molecules, but we will constrain ourselves from presenting them in detail here.

This paper presents a rational analysis of a number of the formula  $\text{Cp}_2\text{Rh}_2(\mu\text{-CO})_2\text{X}$ . If the reader now feels comfortable with the electronic order now apparent in what seemed to be a confused picture of different conformations and diverse  $\text{X}=\text{Rh}(\text{CO})_2^-$ ,  $\text{CR}_2$ ,  $\text{RhCp}$ ,  $\text{RhCp}(\text{CO})$ , then we have accomplished our task.

We are grateful to *R. Bergman* and *W.A. Herrmann* for communication of experimental data prior to publication and to *J.R. Shapley* for discussions of great value. The work at *Cornell* was supported by the *National Science Foundation* through Research Grant CHE 7828048 and DMR-7681083 to the *Materials Science Center* (R.H. and A.R.P.), the *Robert A. Welch Foundation* (T.A.A.), *Deutsche Forschungsgemeinschaft* and the *Fond der Chemischen Industrie* (P.H.).

*Appendix.* - The molecular orbital calculations were of the extended *Hückel* type [28], with a weighted  $H_{ij}$  formula [29]. *Table* lists the parameters used. The  $H_{ii}$ 's and orbital exponents for Fe and Rh were taken from previous work [11, 30]. The bond lengths and angles in **1** and other molecules containing the  $Cp_2Rh_2(CO)_2$  unit were based on the experimental crystal structure of  $Cp_2Rh_3(CO)_4^-$  [1]. The geometry of **14** was based on that of  $Fe_2(CO)_9$  [31], with the Fe-Rh distance set at the sum of the single-bonded radii, 2.42 Å.

Table. *Extended Hückel Parameters*

| Orbital | $H_{ii}$ eV | Exponents <sup>a)</sup> |               |
|---------|-------------|-------------------------|---------------|
|         |             | $\zeta_1$               | $\zeta_2$     |
| Rh 5s   | - 8.09      | 2.135                   |               |
| 5p      | - 4.57      | 2.10                    |               |
| 4d      | - 12.50     | 4.29 (0.5807)           | 1.97 (0.5685) |
| Fe 4s   | - 9.17      | 1.90                    |               |
| 4p      | - 5.37      | 1.90                    |               |
| 3d      | - 12.70     | 5.35 (0.5366)           | 1.80 (0.6678) |
| C 2s    | - 21.40     | 1.625                   |               |
| 2p      | - 11.40     | 1.625                   |               |
| O 2s    | - 32.30     | 2.275                   |               |
| 2p      | - 14.80     | 2.275                   |               |
| H 1s    | - 13.60     | 1.30                    |               |

a) Two Slater exponents are listed for the d functions. Each is followed in parenthesis by the coefficients in the double zeta expansion.

## REFERENCES

- [1] *W.D. Jones, M.A. White & R.G. Bergman*, *J. Amer. chem. Soc.* **100**, 6770 (1978).
- [2] See *R.B. King*, *Prog. inorg. Chemistry* **15**, 287 (1972); *F.A. Cotton & J.M. Troup*, *J. Amer. chem. Soc.* **96**, 1233, 5070 (1974); *F.A. Cotton*, *Prog. inorg. Chemistry* **21**, 1 (1976); *E. Band & E.L. Muetterties*, *Chem. Rev.* **78**, 639 (1978).
- [3] *J. Evans*, *Adv. organomet. Chemistry* **16**, 1 (1977); *S. Aime & L. Milone*, *Prog. NMR. Spectr.* **11**, 183 (1977); *R.D. Adams & F.A. Cotton* in 'Dynamic Nuclear Magnetic Resonance Spectroscopy', *L.M. Jackman & F.A. Cotton*, eds., Academic Press, New York, N.Y. 1975, p. 489-520.
- [4] *A.R. Pinhas & R. Hoffmann*, *Inorg. Chemistry* **18**, 654 (1979).
- [5] a) *W.A. Herrmann*, *Angew. Chemie* **90**, 855 (1978); *Angew. Chemie Int. Ed.* **17**, 800 (1978); b) *W.A. Herrmann*, *Chem. Ber.* **111**, 1077 (1978); c) *W.A. Herrmann & I. Schweizer*, *Z. Naturforsch.* **33b**, 1128 (1978); d) *W.A. Herrmann*, personal communications; e) *W.A. Herrmann, C. Krüger, R. Goddard & I. Bernal*, *J. organomet. Chemistry* **140**, 73 (1977); *W.A. Herrmann, I. Schweizer, M. Creswick & I. Bernal*, *ibid.*, **165**, C17 (1979); *W.A. Herrmann, C. Krüger, R. Goddard & I. Bernal*, *Angew. Chemie* **89**, 342 (1977); *Angew. Chemie Int. Ed.* **16**, 334 (1977); *M. Creswick, I. Bernal, W.A. Herrmann & I. Steffl*, *Chem. Ber.*, in press.
- [6] a) *M. Elian, M.M.L. Chen, D.M.P. Mingos & R. Hoffmann*, *Inorg. Chemistry* **15**, 1148 (1976); b) *M. Elian & R. Hoffmann*, *ibid.* **14**, 1058 (1975); c) *J.W. Lauher, M. Elian, R.H. Summerville & R. Hoffmann*, *J. Amer. chem. Soc.* **98**, 3219 (1976); d) *T.A. Albright, R. Hoffmann, J.C. Thibeault & D.L. Thorn*, *J. Amer. chem. Soc.* **101**, 3801 (1979); e) *A. Dedieu & R. Hoffmann*, *ibid.* **100**, 2074 (1978).

- [7] a) *K. Wade*, 'Electron Deficient Compounds', Nelson, London 1971; *Adv. inorg. Chemistry Radiochemistry* 18, 1 (1976); b) *D. M. P. Mingos*, *Nature* 236, 99 (1972); *Adv. organomet. Chemistry* 15, 1 (1977); c) *A. S. Foust, M. S. Foster & L. F. Dahl*, *J. Amer. chem. Soc.* 91, 5633 (1969) and references therein; d) *J. Halpern*, *Adv. chem. Series*, No 70, 1 (1968).
- [8] *P. Hoffmann*, *Angew. Chemie* 89, 551 (1977).
- [9] *D. M. P. Mingos*, *J. chem. Soc. Dalton Trans.* 1977, 602.
- [10] a) *R. Hoffmann, G. D. Zeiss & G. W. VanDine*, *J. Amer. chem. Soc.* 90, 1485 (1968); b) *W. Kirmse*, 'Carbene Chemistry', 2nd Edition, Academic Press, New York 1971; especially the Chapter by *J. F. Harrison*.
- [11] *R. H. Summerville & R. Hoffmann*, *J. Amer. chem. Soc.* 98, 7240 (1976); 101, 3821 (1979).
- [12] a) *P. Hoffmann*, *Angew. Chemie* 91, 591 (1979); b) *P. Hoffmann*, to be published.
- [13] *R. G. Pearson*, 'Symmetry Rules for Chemical Reactions', Wiley-Interscience, New York 1976, and references therein.
- [14] *O. S. Mills & E. F. Paulus*, *J. organomet. Chemistry* 10, 331 (1967).
- [15] a) *E. F. Paulus, E. O. Fischer, H. P. Fritz & H. Schuster-Woldan*, *ibid.* 10, P3 (1967); b) *E. F. Paulus*, *Acta crystallogr. Sect. B.* 25, 2206 (1969); c) *R. J. Lawson & J. R. Shapley*, *Inorg. Chemistry* 17, 772 (1978); *J. Amer. chem. Soc.* 98, 7433 (1976).
- [16] *F. A. Cotton & J. D. Jamerson*, *J. Amer. chem. Soc.* 98, 1273 (1976).
- [17] *B. E. R. Schilling & R. Hoffmann*, *J. Amer. chem. Soc.* 101, 3456 (1979).
- [18] a) *E. O. Fischer & C. Palm*, *Chem. Ber.* 91, 1725 (1958); b) *A. A. Hock & O. S. Mills*, *Proceedings, 6th International Conference on Coordination Chemistry, Detroit 1961*, p. 640.
- [19] a) *E. O. Fischer & D. Beckert*, *Angew. Chemie* 70, 744 (1958); b) *P. Chini & R. Ercoli*, *Gazz. chim. Ital.* 88, 1171 (1959).
- [20] a) *S. Otsuka, A. Nakamura & T. Yoshida*, *Liebigs Ann. Chem.* 719, 54 (1968); *Inorg. Chemistry* 7, 261 (1968); b) *H. Vahrenkamp, V. A. Uchtmann & L. F. Dahl*, *J. Amer. chem. Soc.* 90, 3272 (1968); c) *M. Sorai, A. Kosaki, H. Suga, S. Seki, T. Yoshida & S. Otsuka*, *Bull. chem. Soc. Japan* 44, 2364 (1971).
- [21] *P. D. Frisch & L. F. Dahl*, *J. Amer. chem. Soc.* 94, 5082 (1972).
- [22] *L. T. J. Delbaere, L. J. Kruczynski & D. W. McBride*, *J. chem. Soc. Dalton Trans.* 1973, 307.
- [23] *L. F. Dahl & P. W. Sutton*, *Inorg. Chemistry* 2, 1067 (1963); *W. Hieber & J. Gruber*, *Z. anorg. allgem. Chem.* 296, 91 (1958).
- [24] *C. H. Wei & L. F. Dahl*, *Inorg. Chemistry* 4, 493 (1965).
- [25] *V. A. Uchtmann & L. F. Dahl*, *J. Amer. chem. Soc.* 91, 3763 (1969).
- [26] *A. Almennigen, A. Haaland & S. Samdal*, *J. organomet. Chemistry* 149, 219 (1978); *J. H. Ammeter*, *J. magn. Res.* 30, 299 (1978); *D. P. Freyberg, J. L. Robbins, K. N. Raymond & J. C. Smart*, *J. Amer. chem. Soc.* 101, 892 (1979); and references therein.
- [27] *M. Benard*, *J. Amer. chem. Soc.* 100, 7740 (1978).
- [28] *R. Hoffmann*, *J. chem. Physics* 39, 1397 (1963); *R. Hoffmann & W. N. Lipscomb*, *ibid.* 36, 2179 (1962); 37, 2872 (1962).
- [29] *J. H. Ammeter, H.-B. Bürgi, J. C. Thibeault & R. Hoffmann*, *J. Amer. chem. Soc.* 100, 3686 (1978).
- [30] *T. A. Albright, P. Hofmann & R. Hoffmann*, *J. Amer. chem. Soc.* 99, 7546 (1977).
- [31] *F. A. Cotton & J. M. Troup*, *J. chem. Soc. Dalton Trans.* 1974, 800.