Reactivity of Transition-Metal Cluster Carbides Which Have an Exposed Carbon Atom

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In this paper we study the reactivity of butterfly and square-pyramidal transition-metal cluster carbides, compounds which are interesting because they have a sterically exposed carbon atom. First we discuss the stability of these compounds which have such an unusual coordination for carbon. Next we use perturbation theory arguments to discuss the relative reactivities of different sites on the butterfly cluster and of the carbon site on the square-pyramidal cluster. The results parallel the experimentally obtained products; in particular the carbon atom is not electronically the most favored site of attack. The orbital explanation of these trends gives us further insight into bonding in the cluster. We also discuss electron counting and π-bonding in the product of attack on the cluster and consider some relevant data on transition-metal and transition-metal-carbide surfaces.

I. Introduction

There exists a remarkable class of compounds, cluster carbides in which a carbon atom is completely exposed on one side of the molecule. The first known example was Fe₄(CO)₁₂C discovered by Braye and Dahl in 1962.¹ The iron framework is square pyramidal, so that carbon is bonded to nothing below the square base. Over the years a number of isostructural carbides were found, and their chemistry was determined.² In 1980 a new cluster, Fe₄-(CO)₁₂C²⁻, was isolated in which four iron atoms form a butterfly framework.³ Again the carbon atom which lies halfway in between the wing tips, 2, is asymmetrically bonded. In these structures, as drawn by us, each unlabeled vertex is an iron atom carrying three carbynols, Fe(CO)₃.

At first sight, it seems strange that these carbides should exist at all. Certainly, nothing in the realm of organic chemistry would indicate such an asymmetric environment for carbon. On the other hand, our experience with bulk carbides⁴ has been that it is not a good idea to make comparisons between the valence of carbon in carbides and the valence of carbon in organic compounds. Perhaps a better comparison is to surface chemistry. Much effort in recent years has been devoted to preparing clean metal surfaces. This is of course a very important endeavor if one is to understand details of how a metal surface catalyzes reactivity on an atomic scale; however, it shows that a clean metal surface is not a very stable entity. One of the most pervasive impurities found on surfaces is carbon. Anyone who has ever tried to clean an iron surface⁵ can convince him/herself that carbon binds very strongly to metal surfaces and stabilizes them, just as carbon stabilizes these surfacelike clusters.

We might also expect the carbon atom to be a site of attack by other ligands. In fact, a good deal is known about the chemistry of both of these classes of compounds. Two classes of reaction undergone by five-metal-atom clusters are (a) ligand substitution; for example, PR₃ substitutes CO,⁶ and (b) addition of a ligand to the metal framework, for example, I⁻ adds to Ru₄(CO)₁₂C. In this case there is a formal addition of electrons to the metal skeleton, causing it to change to the new geometry 3. There is, as yet, no example in which an attacking ligand attaches itself to carbon.

In contrast, there are three known examples of addition to the carbide carbon atom in Fe₄(CO)₁₂C²⁻ or a closely related molecule. Attack by an acid gives first a monohydride in which I is attached to the metal framework and then a dihydride in which the second H is attached to carbon (4).⁷ The dihydride has an extremely long C–H

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bond and is often mentioned in discussions of C-H activation. Attack by \( \text{CH}_3^+ \) presents a different twist. The product of attack has \( \text{CH}_3^+ \) attached to the carbide carbon atom but with the metal framework tetrahedral. What a difference the apparently minor perturbation (\( \text{CH}_3^+ \) replacing \( \text{H}^+ \)) makes! The second example is due to Bradley, who is responsible for a marvelous reaction in which \( \text{Fe}_6(\text{CO})_{16} \text{C}^- \) is used to make an ester from \( \text{CO} \) and \( \text{CH}_3\text{OH} \). Part of his reaction scheme is shown in 5 (all intermediates shown have been isolated). Carbon monoxide adds to the metal framework in the presence of \( \text{C}_2\text{H}_5^+ \), an oxidizing agent. In this case, addition of \( \text{CO} \) is accompanied by a loss of charge such that there is no formal addition of electrons to the cluster and no change in the geometry of the metal framework. Addition of methanol to \( \text{Fe}_6(\text{CO})_{16} \text{C}^- \) gives a product in which the carbide carbon atom is attached to a carbomethoxy ligand.

Hence, the carbon atom in a four-metal-atom cluster is more reactive than the carbon atom in a five-metal-atom cluster, but even in this case the initial product has the ligand attached to the metal framework. A relevant question is whether the structural preference is determined by the relative reactivities of the different sites on the cluster. It is perfectly plausible that ligand attack on carbide carbon is followed by migration around the cluster, provided that this migration is facile. Hydrogen migration is known to be facile, even on an NMR time scale. There are also examples of migration of \( \text{CO} \) from carbon to a metal framework, as well as many examples of facile CO migration around metal clusters. In this case the structural preference would reflect only thermodynamic considerations of the various products. The reactivity of the carbide cluster is the subject of our theoretical study.

II. Details of the Calculations

All calculations were performed by using the extended Hückel method, with parameters from previous work. A \( \text{C}_5 \) \( \text{Fe}(\text{CO})_3 \) cluster was chosen with the angle \( \text{OC}-\text{Fe}-\text{CO} = 90^\circ \) (experimental values are in the range 90-100). Four \( \text{Fe}(\text{CO})_3 \) clusters were arranged in a butterfly around carbon with the experimental values used for \( \text{Fe}(\text{C})_2 = 1.89 \) \( \text{Å} \), \( \text{Fe}(\text{C})_2 = 1.96 \) \( \text{Å} \), and \( \text{Fe}(\text{Fe})_2 = 2.53 \) \( \text{Å} \) (see 6). The other Fe-Fe bonding distances are forced to be 2.62 \( \text{Å} \). Concerning the four \( \text{Fe}(\text{CO})_3 \) fragments, we now (a) varied the angles \( \phi \) subtended by the threefold axes of the fragments and (b) rotated the fragment around its threefold axis. \( \phi_1 \) and \( \phi_4 \) were varied within the \( xz \) plane; \( \phi_2 \) and \( \phi_3 \) were varied within the \( xy \) plane. Certain \( \phi \) angles optimize metal-to-carbon bonding; others optimize metal-to-metal bonding. The angles we chose were a compromise: \( \phi_1 = \phi_4 = 155.3^\circ \), \( \phi_2 = \phi_3 = 165^\circ \). These values are close to the experimental angles for \( \text{Fe}_4(\text{CO})_{12} \text{C}^- \): \( \phi_1 = 154.5^\circ \), \( \phi_2 = 166.4^\circ \), and \( \phi_3 = 164.2^\circ \). Of course, steric reasons can cause the optimal values of \( \phi \) to be very different when additional ligands are bonded to the cluster. Finally, the fragments were rotated around their threefold axes until a minimum was obtained in the total energy corresponding to the \( C_{2v} \) geometry.

For \( \text{Fe}_6(\text{CO})_{16} \text{C}^- \), we used the same values for the metal-to-carbon distance; i.e., \( \text{Fe}(\text{C})_2 = 1.96 \) \( \text{Å} \), \( \text{Fe}(\text{Fe})_2 = 1.80 \) \( \text{Å} \). All angles \( \text{Fe}-\text{C}-\text{Fe} \) were ideal. The threefold axes of the four fragments in the \( xy \) plane were bent down (\( \phi_1 = \phi_4 = 170^\circ \), and \( \phi_2 = \phi_3 = 162^\circ \) (8). Rotation of carbonyls around the threefold axis was constrained to maintain an \( xz \) mirror plane, and a minimum energy geometry obtained was 9.

In both cases the position of the carbide carbon atom was idealized. For \( \text{Fe}_6(\text{CO})_{16} \text{C}^- \), carbon was chosen to bisect the edge of the wing (\( \theta = 180^\circ \)). Actually \( \theta = 176.3^\circ \), and the carbon atom moves slightly out of the wings of the butterfly. For \( \text{Fe}_6(\text{CO})_{16} \text{C}^- \), carbon was chosen to lie in the plane of the four basal Fe atoms, whereas it is actually displaced slightly (0.1-0.2 \( \text{Å} \)) below the plane.

III. \( \text{Fe}_6(\text{CO})_{12} \text{C}^- \)

Before considering the reactivity of the butterfly cluster, it helps to have some knowledge of its molecular orbitals. The bonding to carbon is highlighted by showing an interaction diagram, Figure 1, for the formation of \( \text{Fe}_6(\text{CO})_{12} \text{C}^- \) from \( \text{Fe}_6(\text{CO})_{12} \) and carbon. \( \text{Fe}_6(\text{CO})_{12} \) can be considered to be made up of four \( \text{Fe}(\text{CO})_3 \) fragments. The molecular orbitals of the fragment

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(15) We initially did the calculations with all Fe-Fe distances the same and all Fe-C distances the same. The differences in overlap population correctly predicted which bonds should shorten.
are well-known, they consist of a frontier set—the linear combinations of three orbitals directed toward the missing sites of an octahedron—above the remnants of the octahedral $t_{2g}$ set (three orbitals). The $t_{2g}$ set spreads into a narrow band due to metal-to-metal bonding, the frontier set into a wide band. The HOMO–LUMO gap occurs inside the frontier set: a cluster with charge 6− contains seven occupied orbitals in agreement with the calculations of Lauher, as well as with those for the isolobal nido boron hydride.

Bonding with carbon has been discussed in great detail for octahedral and trigonal-prismatic carbides. Interaction of carbon $s$ and $p$ orbitals in Figure 1 occurs primarily with four occupied frontier orbitals of symmetry $a_1$, $a_2$, $b_1$, and $b_2$ (the molecule has symmetry $C_{6v}$). The orbitals of the carbide have been numbered from the top down. Carbon $s$ lies very low in energy and is not shown. The other three bonding orbitals (73–75) lie below the $t_{2g}$ set. The three HOMO’s (58–60) are the metal-to-carbon nonbonding remnants of the occupied frontier set. Below them lies the $t_{2g}$ set (61–72), except that the top three and the bottom-most members of this set contain a substantial carbon content and are shown separately. The reason these orbitals have a high carbon content is that substantial overlap does exist between the $t_{2g}$ set and carbon, especially when threefold axes of the Fe(CO)$_6$ fragments do not point toward carbon. Among the unoccupied orbitals, the LUMO’s (56–57) are metal-to-carbon nonbonding, but there is a set of metal-to-carbon antibonding orbitals (53–55) above them. Actually, most metal-to-carbon antibonding is in orbitals that are off-scale.

The most impressive feature of the carbide is a HOMO–LUMO gap of 1.7 eV. This is remarkably high for a molecule this size in which no approximations (such as substituting H for CO) have been made. Note also how carbon stabilizes the molecule by opening up the HOMO–LUMO gap.

IV. Reactivity with a Model Nucleophile or Electrophile

In this section we use a model hydrogenic orbital $i$, whose energy $E_i$ is variable, as a probe of reactivity in butterfly and square-pyramidal clusters. Second-order perturbation theory and the extended Hückel approximation tell us that orbital $i$ will interact with each of the orbitals $j$ of the cluster, in a pairwise additive manner, according to the formula $S_{ij}^2/(E_i - E_j)$.

There are many reasons for separating the overlap term $S_{ij}$ from the energy term $E_i - E_j$. One reason is to make it easier to understand what would happen to a range of nucleophiles with different $E_i$. Another is to take account of the fact that our knowledge of these $E_i$’s is very imperfect; if the result we get is dependent on small changes in the energy of the nucleophile, we should be cautious.

The problem with separating out the overlap term is one of presentation: there may be a large number of orbitals of a molecule that interact with a nucleophile. This is true of large molecules such as the one we are considering; it is especially true of solids. Hence, we adopt a solution from solid-state physics, the same one we adopted when discussing bulk carbides. We draw the orbital energies in the way we normally construct an MO diagram, except that the line drawn for each orbital is weighted by the overlap of that orbital with the probe. For the bulk carbides, orbitals were weighted either by their percent carbon to give a projection of carbon or by their overlap population to give a COOP curve. In our case we have constructed an interacting orbital overlap (IOO) curve; six of these curves are plotted in Figures 2 and 3.

The geometries of attack corresponding to each profile are discussed in 10–15. Carbon-to-nucleophile distances in 10, 11, 12, and 15 are 2.0 Å. Carbon-to-nucleophile and
iron-to-nucleophile distances in 13 are 2.4 Å. In this case the threefold axis of the bridged Fe(CO)₃ unit is bent back ($\phi_r = 165.38^\circ$). The iron-to-nucleophile distance in 14 is 2.7 Å.

We start our discussion of these curves with some general comments about the reactivity of the butterfly cluster. As already mentioned, this cluster has a very large HOMO-LUMO gap of 1.7 eV, which tends to make it stable to attack. Figures 2 and 3 indicate that, in addition, most of its overlap with an incoming probe orbital is through occupied orbitals. This is in accord with the experimental facts described earlier: attack on the cluster is either with an electrophile (H⁺) or with a nucleophile and an oxidizing agent (CO and C₇H₇⁺).

The reason a nucleophile cannot act alone is that its major interaction is with occupied orbitals and is repulsive. Part of this repulsive energy is contained in a rising HOMO. When the energy of the HOMO rises above the energy of the LUMO on an oxidizing agent, electrons are transferred to the oxidizing agent, and there is a favorable bonding interaction between the HOMO and the nucleophile. If this orbital of the oxidizing agent lies inside the HOMO-LUMO gap of the carbide, the barrier to attack of a nucleophile is reduced, as shown schematically in 16.

The essence of this and the following arguments does not change if electron transfer to the oxidizing agent occurs before attack of CO on the butterfly carbide.

For a given nucleophile, which feature of a molecule makes this electron transfer efficient? To answer this question, we have constructed interaction diagrams in Figure 4 for the attack of H⁻ ($H_f = -13.6$ eV) either on the carbon atom, 12, or on the back iron-to-iron bond, 14. The major difference between the two interaction diagrams is that a much higher percentage of the repulsive energy of interaction is directed into the HOMO during attack on the iron-to-iron bond. IOO curves also show this difference. For attack on the back iron-to-iron bond the major interaction is with the HOMO (58). For attack on carbon the major interaction is with a deeper orbital (74) which is one of the iron-to-carbon bonds.

Hence, to understand the oxidizing-agent-assisted attack of a nucleophile, we need to consider the distribution of probe overlaps among occupied states of the carbide. A large overlap of the probe with states near the Fermi level is favorable. We now extend this concept to ask how we might favor attack on the carbon atom of Fe₄(CO)₁₂C₂⁻. If we compare the two asymmetric attacks 11 and 13 with direct attack 12, we see that the best attack is 13, across an iron-to-carbon axial bond. The reason for this is a good overlap with an orbital (59) which is near the HOMO.

We also have a means of comparing direct attack on the carbon atom of a butterfly cluster with a similar attack on a square-pyramidal cluster. The HOMO-LUMO gap is actually 1.3 eV in the square-pyramidal cluster, smaller than it is in the butterfly cluster. However, in the square-pyramidal cluster the HOMO does not have the correct $a_2$ symmetry to interact with a nucleophile, and hence the effective HOMO is an orbital $1/2$ eV below the real HOMO. More importantly, a larger percentage of the overlap of the square pyramid with a probe is with low-lying iron-carbon bonding orbitals. The net result is that attack on the carbide carbon atom of the square-pyramidal cluster is less favorable.

It is helpful to have a numerical criterion to summarize these results. Unfortunately, we are not able to draw a complete reaction profile since (i) the energy of the oxidizing agent is uncertain, and (ii) we are comparing apples with oranges: Fe–H bonds with C–H bonds. A criterion that gets around these objections, one that we shall use,
is the percentage of repulsive energy that is channeled into the HOMO. The higher the percentage of repulsive energy in the HOMO the faster it transfers its electrons to the oxidizing agent and the lower the barrier to attack. The components of this ratio are illustrated separately in Figure 5. As we go from left to right in this figure, 10–14, the repulsive change in the total energy of the system decreases, but the energy of the HOMO rises. Hence, the structures exhibit increasing reactivity from left to right in this figure. The reader will notice that this ordering is in accord with the qualitative picture implied by the interacting orbital overlap curves.

We can also change the energy of the probe to model a more electronegative ligand. The probe energy of $-13.6$ eV used in Figure 5 is close to the energy of the donor orbital on a CO or PH$_3$ ligand. In contrast, we use $-11.5$ eV in Figure 6; this energy is typical of the lone pair of I$^-$ or CH$_2^-$. This new probe is close in energy to the HOMO of the carbide and hence has a greater "reactivity" with the carbide; i.e., the change in total energy is approximately the same as in Figure 5, but the rise of the HOMO is greater. Hence, if we take two electrons out of this system, we obtain a large attractive energy. Secondly, the trend with different structures is the same as in Figure 5, but the differences are less marked. In other words, the "selectivity" of the probe is much less.

It is interesting to compare our results to previous theoretical discussions of these two carbides. Shriver et al. studied model carbides in which three CO ligands were replaced by one H. They correlated the greater reactivity of the carbon atom of a butterfly cluster when compared to that of the carbon atom of a square-pyramidal cluster to the fact that the HOMO of the butterfly carbide contains a significant charge on the carbon atom. We agree with this result; however, we do not agree with them that the HOMO–LUMO gap is less in the butterfly cluster. Housecroft and Fehlner did Fenske–Hall calculations on the product of interaction of a hydrogen atom with the butterfly cluster and showed that there is a greater overlap of hydrogen with the set of frontier orbitals of the cluster when hydrogen bridges an Fe–C bond as in 13 than when it is attached symmetrically to carbon. Our calculations on a reactant-like geometry are in agreement.

This trend in reactivity for different structures is also in excellent agreement with the experimentally determined thermodynamic products. In our introduction, we stated that (i) initial attack occurs on the back iron-to-iron bond as in 14; (ii) in the case of H$^+$ attack, the product of attack on carbon was asymmetric as in 13. There are also other examples of a product in which a ligand is bonded to the carbide in a manner analogous to 13. These are the attachment of OCH$_3^-$ to HF$_4$(CO)$_2$C$^{+}$ and the attachment of the carbene C(H)(t-Bu)$_2$ to HNiOS$_2$Cp(CO)$_2$C$^{+}$. A counterexample exists from the work of Bradley in which the attachment of CO(OCH$_3$)$_3$ to Fe$_4$(CO)$_2$C, which is in accord with 12 (this data is summarized in Table I). The reason for this exception may be steric. In any case it is not necessary to invoke mobility around the cluster to explain the preferred products as was implied in our discussion at the end of the Introduction.

V. Orbital Explanation

We now derive an orbital explanation of the above trends. The most surprising result of the previous section was the low reactivity of the carbide carbon atom. In order to explain this result, it is pedagogically useful to compare the carbide to its organic analogue, butterfly CH$_4$. Butterfly CH$_4$ does not exist, and a Walsh diagram for the transformation tetrahedral to butterfly depicted in 17

quickly explains why this is so. Of the three p orbitals on carbon, p$_z$ becomes more bonding on going from tetrahedral to butterfly, but p$_x$ rises in energy. These two orbitals cancel one another out leaving the largest effect, that due to the p$_y$ orbital which very rapidly loses bonding on one side of the molecule and becomes directed towards the
the probe ligand. Weakening the iron-to-carbon a bond decreases the angle and makes the (Fe-C) bond shorter than the average (Fe-C) bond. And carbon is approximately collinear with the other iron atoms; these are clearly signs of a thermodynamic phenomenon. However, our IO0 curves must consider the electrophile (such as H+, FeBr3 -) and the other views. The view in the figure is a cut off the carbon atom. In other words, the acidity of the carbide carbon atom in these clusters we can obtain both theoretical and experimental measures of the strength of this π-bond. The theoretical measure is that the overlap population of carbon px, with the rest of the cluster is not less than the overlap population of carbon px, with the rest of the cluster. Actual contributions of the different carbon orbitals to the total overlap population are s = 0.6282, px = 0.4944, py = 0.5014, and pz = 0.6324.

The experimental measure is structural; it is summarized in Table I. In those compounds in which there is no ligand attached to carbon, the (Fe-C) bond is shorter than the average (Fe-C) bond length, and carbon is not collinear with the axial iron atoms; these are clearly signs of π bonding. On the other hand, products of attack of a ligand on a carbide carbon atom have no need for this stabilizing π bond. In these products, structures 12 or 13, the average (Fe-C) bond length equals the average (Fe-C)xyz bond length, and carbon is not collinear with the axial iron atoms (θ < 180°). Deviation from collinearity increases with increasing steric bulk of the ligand.

It is also interesting to examine what happens to our IOO curves when carbon is no longer collinear with the axial iron atoms. The IOO curve for structure 15 in Figure 3 shows us what happens when θ is 150°. The HOMO moves up in energy and has a much bigger overlap with the probe ligand. Weakening the iron-to-carbon π bond also causes the antibonding orbital to come down in energy; we see an increasing overlap of the probe ligand with low-lying acceptor orbitals such as px. Our numerical criterion also shows that the reactivity goes up when we decrease the angle θ at the carbon atom.

Cluster Carbides with an Exposed Carbon

Table I. Structural Parameters of Fe6(CO)12Cl2, HFe6(CO)12Cl2, and Their Derivative Compounds in Which an Extra Ligand Is Attached to Carbon or to the Metal Framework

<table>
<thead>
<tr>
<th>framework</th>
<th>ligand</th>
<th>θ, deg</th>
<th>(Fe-C)x, Å</th>
<th>(Fe-C)xy, Å</th>
<th>structure</th>
<th>ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe6(CO)12Cl2</td>
<td>2e-</td>
<td>176.3</td>
<td>1.80</td>
<td>1.97</td>
<td>14</td>
<td>3</td>
</tr>
<tr>
<td>CO</td>
<td>176.0</td>
<td></td>
<td>1.80</td>
<td>1.99</td>
<td>12</td>
<td>11a</td>
</tr>
<tr>
<td>CO(OCH3)−</td>
<td>147.9</td>
<td></td>
<td>2.02</td>
<td>1.95</td>
<td>11</td>
<td>11b</td>
</tr>
<tr>
<td>FeBr3−</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HFe6(CO)12Cl2</td>
<td>2e−</td>
<td>174.0</td>
<td>1.80</td>
<td>1.99</td>
<td>13</td>
<td>7a</td>
</tr>
<tr>
<td>H−−</td>
<td>170.5</td>
<td></td>
<td>1.82, 1.92a</td>
<td>1.85</td>
<td>13</td>
<td>7a</td>
</tr>
<tr>
<td>OCH3−</td>
<td>162.4</td>
<td></td>
<td>1.92, 2.14a</td>
<td>1.86</td>
<td>13</td>
<td>7c</td>
</tr>
<tr>
<td>H2Ru3Cp(CO)6Cl2+</td>
<td>CH(t-Bu)3−</td>
<td>153.9</td>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

a The bridged axial bond is longer. b The isoelectronic BH4 complex exists in which the geometry of the B-H bond is similar to that of the C-H bond in CH4.

empty coordination site, 18. It is the px orbital which is responsible for CH4 not having a butterfly structure.

If such a "lone pair" existed in the butterfly carbide, we would expect our probe to feel a large interaction with an orbital near the HOMO. In fact, probe 12 finds the major px character to be in the much lower energy region of the iron-carbon bonds. The reason is that the carbon "lone pair" is lowered in energy through π bonding with iron orbitals (19). This π interaction is not possible when carbon is attached only to hydrogen with its s orbital.

![Figure 7. Contour plots corresponding to two views of the HOMO.](image)

π bonding can also be invoked to explain the greater reactivity of a butterfly carbide carbon atom when compared to a square-pyramidal carbide carbon atom. The reason is that carbon in the latter case π bonds with four in-plane Fe atoms 20, rather than two in-plane Fe atoms in the butterfly carbide 19. Hence the carbon px orbital is more strongly stabilized, and this result is reflected in the IOO curves.

At this point a word of caution is in order. The reactivity we have referred to above is explicitly related to attack by a nucleophile. In particular, if we discuss the acidity of the carbide carbon atom in these clusters we must consider the electrophile H+, which moves freely on and off the carbon atom. In other words, acidity is often a thermodynamic phenomenon. However, our IOO curves are still useful. This is because our probe also measures the thermodynamic stability of different sites on the reactant, as is obvious from the above discussion. The point is that the conjugate anion of this acid is stabilized by π bonding, and this stabilization is greater for the Fe5 cluster.

At this point we have shown why the carbide carbon atom is not reactive, but we have not shown why attack takes place at other sites on the cluster. To understand this it is helpful to examine some orbital plots. Clearly the nature of the HOMO is very important: it might even be able to predict the direction of nucleophilic attack. Figure 7 shows that this is so. The HOMO is shown in two views. The view in the xz plane passes through carbon, two axial iron atoms, and the center of the equatorial iron-to-iron bond. The position of the attacking probe in 12 is indicated by a H atom. This figure shows that the electron density is greater at −x, toward the back iron-to-iron bond, than it is at +x, in the direction of the attacking H atom. The other view of this orbital is a cut
through the equatorial iron-to-iron bond (yz plane), showing that this orbital is in fact strongly bonding between these iron atoms. Hence, we correctly predict the direction of preferred attack to be on this bond.

Figure 8 shows two other \(a_1\) orbitals, a low-lying unoccupied orbital (54) and one of the iron-to-carbon bonds (74). These were chosen because they have a relatively high component on the carbon atom. Compared to the HOMO, orbital 74 has a greater electron density at \(+z\), in the direction of the attacking nucleophile, and a smaller electron density at \(-x\). Hence, it is this orbital, shown schematically in 19, that interacts preferentially with a nucleophile, in agreement with everything we have said previously. The low-lying unoccupied orbital has too many nodes to interact well with the probe, and this explains our earlier observation that a probe does not interact well with unoccupied orbitals.

If we look at the orbital just below the HOMO, Figure 9, we can explain why 13 is the second most likely mode of attack. This orbital, schematized in 21, is essentially the minus combination of the Fe orbitals that stabilized the carbon \(p_z\) orbital in 19. This minus combination has \(b_2\) symmetry, but it cannot interact with the carbon \(z\) orbital since it has a node approximately along the Fe–C–Fe axial thread. Hence, it is high in energy when compared to the plus combination of the same iron orbitals. In order to take advantage of this orbital, the probe has to move in the \(z\) direction and bridge an iron-to-carbon bond. This is exactly the geometry described by 13.

**VI. Electron Counting in the Product**

The skeletal electron pair counting rule is a marvelous device for rationalizing the structures of clusters. In applying this formalism one occasionally runs across interesting ambiguities. One example is how to count the carbide carbon atom. Experimental evidence suggests that we maintain the same electron count in the carbide cluster as we do in the noncarbide cluster if we count carbon as neutral, a four-electron donor. This implies that no new occupied states are introduced on interacting carbon with the rest of the cluster. The calculations of both Lauher and ourselves support this statement, and we considered in detail the reasons why this was so for octahedral Ru-\((CO)_{18}C_2^+\).

In this section we shall consider the products of attack on the carbide. In Figure 10 we have constructed interaction diagrams for attack on the carbide carbon atom as well as on the back Fe–Fe bond. These diagrams are similar to those of Figure 4, except that the C–H (1.09 Å) and Fe–H (1.70 Å) distances have been shortened to form bonds. For attack on the back iron-to-iron bond, we also had to bend back the equatorial Fe(CO)\(_3\) fragments; \(\phi_2 = \phi_3 = 151.63^\circ\).

In Figure 10, as in Figure 4, \(\mathrm{H}^+\) interacts mainly with occupied orbitals. The result is that the HOMO is pushed up above the HOMO–LUMO gap and becomes empty. In other words the product has the same number of occupied orbitals as the reactant. We can think of this product either as attachment of a Lewis acid to Fe\(_4\)(CO)\(_{12}\)C\(_2^+\) or as...
attachment of a Lewis base to Fe₄(CO)₁₂C.

In Table I we took the latter point of view. Here we have listed the known derivatives of Fe₄(CO)₁₂C and HFe₂-(CO)₁₂C⁺ (protonated on the back iron-to-iron bond). The results justify our electron counting scheme.

We can also consider the product HFe₂(CO)₁₂CH as a CH group bridged across the wings of the Fe₄ butterfly. In this case CH is a five-electron donor, because there is interaction between the C-H bond and an iron framework orbital, analogous to 19. The point is emphasized by looking at the apparently isoelectronic compound 22.²³

Since the skeleton electron count for a tetrahedral cluster is two less than that for a butterfly cluster, the out-of-plane CCH₃ group in 22 is only a three-electron donor; i.e., there is no interaction between the C-C₃ bond and the metal framework. A similar analysis may applied to the claimed 64-electron butterfly clusters, made by Carty.²⁵

Another detail of Table I worth mentioning is the possible role of π interaction in determining the position of a ligand attached to carbon. We compared the two bridging geometries, 11 and 13, for CO bonded to the carbide. In both cases the distance C-CO was 1.5 Å, and the distance Fe-CO was 2.0 Å. We did not include terminal CO 12, since it was unreasonable to assume a constant C-CO distance in this case.

In this case of a π acceptor, the contribution of π orbitals to the overlap population was 0.6669 for geometry 11 and 0.5017 for geometry 13. Since σ interaction favors structure 13, it is possible, though not necessary, that π acceptors are found with structure 11. Unfortunately CO has only been isolated attached to the back iron-to-iron bond, and it is probably very reactive when attached to the carbon atom. The only "ligand" in our list with structure 11 is FeBr₂⁻. FeBr₂⁻ is d⁸ ML₂, a fragment derived from square-planar ML₄. Hence, it should have four low-lying nonbonding orbitals which are filled and two frontier orbitals which are empty. The best overlap is with the frontier orbitals; hence FeBr₂⁻ is a π acceptor. For the same reason, this fragment is a poor ρ donor, and this fact is consistent with the product being unstable.

π interaction of acceptors with the carbide is large and positive; i.e., interaction is mainly with occupied orbitals of the carbide. The interaction of occupied orbitals on the carbide with π donors such as OCH₃ is repulsive; hence donors have an additional reason to favor structure 13. We note that in our electron-counting scheme the carbene is a π donor since HNiRu₂Cp(CO)₂C⁺ is isolobal with HFe₂(CO)₁₂C⁺, and since interaction with the carbene gives a neutral product.

In concluding this section we note a recent study of bonding and reactivity in the butterfly clusters by Harris and Bradley.²⁴ Their theoretical work is in general in agreement with ours, with some differences as to the composition of the Fe₄(CO)₁₂⁻ HOMO and the reasons for the axial coordination of the organic group in Fe₄(CO)₁₂(CO₂C₂H₅)⁴°.

VII. Application to Surface Studies

In this section we broaden the scope of our discussion. First we consider what the appropriate geometry would have to be in order to obtain a stable carbide cluster containing three metal atoms. We then search for surfaces where the local environment of carbon is similar to that in the butterfly or square-pyramidal clusters. Finally, we consider some experimental data for these surfaces.

From our previous discussion, an essential structural feature that stabilizes the carbide is having carbon in the plane of, or collinear with, some neighboring iron atoms so that π bonding to carbon can take place. This means that a geometry in which carbon sits above the plane of a three-membered ring is not reasonable (this statement is supported by calculations of Shriver).²³ More reasonable is 23, in which an Fe-Fe bond opens up and allows carbon to bridge it. In fact, an analogous nitride does exist.²³

Similarly, we would expect that surface carbides are stabilized when carbon lies approximately in the plane of the surface layer of metal atoms. Consider the fourfold site on the (100) face of bcc, a site whose local environment resembles that of a square-pyramidal cluster. For a cube of side a, M-M = 3¹/₂a/2. If carbon lies in the surface layer, M-C = a/2, and the ratio (M-M)/(M-C) = 3¹/₂. For an Fe-Fe distance of 2.5 Å, Fe-C would be 1.44 Å. This means that carbon must lie ~0.5 Å outside the surface plane.

Carbon can get much closer to the surface of an fcc metal. In this case, M-M = a/(2¹/₂). If Fe-Fe = 2.5 Å, Fe-C = 1.77 Å, so carbon can be 0.1-0.2 Å outside the surface layer. Further, carbon in the fourfold site on a (100) face resembles carbon in a square-pyramidal cluster, and carbon in the long bridging site on a (110) face resembles carbon in a butterfly cluster. Both of these cases are shown in 24.

Unfortunately there is a paucity of good structural data for carbon attached to these surfaces. In Somorjai's²⁶ list of structures determined by LEED and ion-scattering techniques, we find a few examples where N or O adsorb on the fourfold sites of fcc (100). These are N on Cu, O on Co, and O on Ni. The metal-to-adsorbate distances suggest that these adsorbates are close to the surface plane of metal atoms (~0.2 Å outside this plane). It should be

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²⁴ Harris, S.; Bradley, J. S., submitted for publication in Organometallics.


noted that the analogous square-pyramidal nitride cluster also does exist.

Somorjai lists no examples of a small adsorbate on the long bridge site of a (110) face of a fcc crystal, although Weinberg has suggested such a site for O on Ir. Nevertheless there is indirect evidence that carbon may bind to a (110) face in the manner described above. The (100) and (110) surfaces of fcc Pt, Ir, and Au are not stable; they undergo reconstruction. Models for the (1 × 2) reconstruction of the (110) surface have been proposed, all of which destroy the butterfly sites for carbon. What is intriguing is that small amounts of carbon have been found in many cases to stabilize the unreconstructed surface. It is also interesting that in one of these cases carbon has been found to considerably enhance the ability of the surface to dissociate CO.

Another way to have carbon on the surface is to start with a carbide. The appropriate carbides are those with the rock-salt structure exemplified by NbC. Here the metal lattice is fcc, and carbon sits in octahedral holes. Since both a square pyramid and a butterfly are fragments of an octahedron, it is clear that if the rock-salt structure exists, carbon will have the environment we are looking for. Once again the appropriate cuts are (100) and (110), the main difference from the previous case being that the coverage of carbon is required to be 1 × 1. In the case of carbon on an fcc surface, C–C repulsion keeps the density of carbon atoms from getting as high as 1 × 1. The importance of high coverage is that in this case the butterfly or square-pyramidal "clusters" are not well isolated from one another, and this may affect their properties.

Even less is known about the surface of a carbide than about the carbonaceous surface of a metal. Rock-salt surfaces such as those of TaC and TiC have been shown to have high catalytic activity in hydrogenation as well as in Fisher–Tropsch chemistry, provided that the surfaces were activated by heating them in a vacuum. In the only single crystal study on surfaces of this structure, a (111) surface in which the surface is composed entirely of metal atoms was found to be better than a (100) surface at adsorbing O₂, H₂, and N₂.

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Transition-Metal Complexes Corresponding to the Insertion into a Group 4B Element–Carbon Bond. 3. Reactivity of Complexes with Unsaturated Carbon–Carbon Bonds. Crystal Structure of \((\eta^5\text{-Cyclopentadienyl})(\text{tri phenylgermyl})(\eta^3\text{-hexenyl})\text{nitrosylmolybdenum}\)

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Anions of the type \([\eta^5\text{-C}_5\text{H}_5\text{CH}_2\text{CO}]\text{(L)}\text{(M}_2\text{C}_2\text{Ph})\text{M}_2^+\text{]}\) (L = CO, NO; M = Si, Ge, Sn; M₂ = Mn, Mo, W) react with allyl halides, affording neutral \(\eta^5\)-bonded allyl derivatives. These new complexes can rearrange to \(\eta^1\)-allyl complexes and also lose \(\text{(allyl)}\text{M}_2\) to give \(\eta^1\)-allyl complexes. According to the nature of both the transition metal and the group 4B metal \(\eta^1\), \(\eta^5\), or \(\eta^3\) complexes are obtained. A mechanism for these successive reactions is proposed. Butenyl and hexenyl iodides also react with the anions, affording \(\eta^1\) complexes that can eliminate CO and rearrange to a \(\eta^5\) ligand as shown by X-ray structure determination.

Introduction

In previous papers, we reported the synthesis and reactivity of \((\eta^5\text{-cyclopentadienyl})\text{transition metal anions}\) that contain a transition metal–group 4B metal \(\eta^5\) bond. These anions were nucleophilic enough to undergo alkylation with alkyl iodides or benzyl bromides (Scheme I). The neutral complexes thus obtained were quite unreactive \((\text{Mo, W})^2\) or showed a peculiar reactivity \((\text{Mn})^2\).

We describe here the reactions of these anions with unsaturated halides and the reactivity of the neutral complexes thus obtained. The X-ray structure of a \(\eta^5\)-allylic