driving force for reaction 2, this result is in reasonable agreement with the findings of Guarr and McLendon which indicate that the thermal reactions between $Ru(bpy)_3^{3+}$ and homologues of methyl viologen, having driving forces of 0.1 to 0.6 V, require a mean separation between the reactants ranging from a contact distance to 10 Å in glycerol at 0 °C.93

Conclusion

These data indicate that disproportionation on PVG occurs via a reaction sequence very similar to that which occurs in aqueous solution.⁴⁰ Biphotonic excitation photoionizes the complex and the photodetached electron reduces a second $Ru(bpy)_3^{2+}$ ion. In aqueous solution, product lifetime is limited by the thermal back reaction which occurs because the products are free to diffuse about the reaction medium. On this support, however, the reactants are fixed, whereas the photodetached electron, after thermal excitation from initially populated surface sites, is a mobile intermediate. Consequently, disproportionation only occurs when the mean separation between the reacting adsorbate ions is within the electron migration distance. On the other hand, the electron migration distance, which these and other experiments indicate is on the order of 50 Å^{46,81,92} exceeds the mean separation, ≤ 13 Å, necessary for the thermal back reaction, and the reaction

(93) Guarr, T.; McLendon, C., Abstracts, 182nd National Meeting of the American Chemical Society, New York, 1981: INORG 20.

products are stable. Increasing the amount adsorbed increases the probability of the thermal back reaction by decreasing the mean product separation, and the quantum efficiency of the reaction declines.

This apparent "photochemical diode" effect³² appears to arise from two factors on this support. First is the availability of electron acceptors sites on the surface. These are thought to be shallow energy wells from which the electron can be thermally activated but, nevertheless, present an energy barrier, albeit slight, which prevents immediate recombination. Energetically, they perform a function similar to band bending which promotes charge separation on the more conventional metal oxide semiconductors.94 Second is that the electron migration distance exceeds that necessary for the thermal back reaction.

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CO Activation by Biscyclopentadienyl Complexes of Group 4 Metals and Actinides: η^2 -Acyl Complexes^{\perp}

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Abstract: The electronic and geometrical structure and reactivity of η^2 -acyl complexes of group 4 transition metals and of actinides are the subject of this paper. The directionality of the LUMO of d⁰ Cp₂MI₂ systems favors a lateral or outside approach of a CO over a central one. Next the electronic structures of the η^2 acyls themselves are analyzed, and overlap reasons given for the preference for η^2 coordination in the Ti, Zr, and actinide complexes that are the main subject of this study, in contrast to the η^1 coordination in 18-electron Mn acyls. Our potential energy surfaces and qualitative analysis point to a preference for the O-inside η^2 -acyl conformer for Ti and Zr, and approximately equal energies for O-inside and O-outside conformers for U and Th. We also find an unexpected η^1 minimum in the O-outside surfaces, with an orbital symmetry imposed barrier to slipping to the η^2 minimum. This suggests a pathway for the interconversion of the O-outside and O-inside η^2 isomers through an as yet unobserved η^1 structure. Separate oxycarbene isomers are also located, but these are at high energy. We believe the electrophilic reactivity of the η^2 acyls does not depend on reaching the oxycarbene isomers, but may be traced to a low-lying carbenium-ion-like acceptor orbital in the undistorted η^2 -acyl structure.

Introduction

A particularly fascinating aspect of the activation of carbon monoxide by transition metal compounds is the carbonylation chemistry of biscyclopentadienyl dialkyls (diaryls), haloalkyls, and related derivatives of group 4 d metals¹ and of actinides,^{2,3} Cp₂MR₂ and Cp₂MRX. CO insertion processes into the M-R bonds of these complexes are extremely facile and while not catalytic have

served as models for the CO activation and reductive CO coupling steps of some potentially industrially important catalytic processes.⁴

¹ In this paper the periodic group notation is in accord with recent actions by IUPAC and ACS nomenclature committees. A and B notation is elimi-nated because of wide confusion. Groups IA and IIA become groups 1 and 2. The d-transition elements comprise groups 3 through 12, and the p-block elements comprise groups 13 and 18. (Note that the former Roman number designation is preserved in the last digit of the new numbering: e.g., III \rightarrow 3 and 13.) [†]Osaka University.

¹Anorganisch-chemisches Institut der Technischen Universität München. [§]Cornell University.

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the other one with the opposite C–O orientation (2). Acyl complexes of group 4 metals, such as $Cp_2Ti(COCH_3)Cl^5$ and $Cp_2Zr(COCH_3)(CH_3)$,^{la,b} show structures of type 2. In contrast, the actinide analogues favor structure 1, or at least 1 can be as stable as 2. Thus the acyl ligand of Cp_2 *Th[COCH₂C(CH₃)₃]Cl⁷

(3) (a) Intriguing is a CO insertion into the U=C double bond in Cp₃UCHP(CH₃)(C₆H₃)₂, which leads to the $\eta^2 \beta$ -keto ylide structure: Cramer, R. E.; Maynard, R. B.; Paw, J. C.; Gilje, J. W. Organometallics **1982**, 1, 869-871. (b) A coordinated carbonyl in CpMn(CÕ)₃ or [CpFe-(CO)₂]₂ can also be inserted into the U=C bond in Cp₃U CHP-(CH₃)(C₆H₃)(R) (R = CH₃, C₆H₃): Cramer, R. E.; Higa, K. T.; Gilje, J. W. J. Am. Chem. Soc. **1984**, 106, 7245-7247. (c) Remarkably facile CO insertion into M-C and M-N bonds (M = Th, U) were also found for Cp₃ThR, Cp₃UR, and Cp₃UNR₂: Sonnenberger, D. C.; Mintz, E. A.; Marks, T. J. *Ibid.* **1984**, 106, 3484-3491. Paolucci, G.; Rossetto, G.; Zanella, P.; Yünlü, K.; Fischer, R. D., J. Organomet. Chem. **1984**, 272, 363-383. (d) Insertions of isocyanides into the U-C bonds of Cp₃UR, Cp^{*}₂URCl, and Cp^{*}₂UR₂ have recently appeared: Dormond, A.; Elbouadili, A. A.; Moise, C. J. Chem. Soc., Chem. Commun. **1984**, 749-751. Zanella, P.; Paolucci, G.; Rossetto, G.; Benetollo, F.; Polo, A.; Fischer, R. D.; Bombieri, G. *Ibid.* **1985**, 96-98.

(4) For general aspects of CO insertion reactions, see, for example: Collman, J. P.; Hegedus, L. S. "Principles and Applications of Organotransition Metal Chemistry"; University Science Books: Mill Valley, CA, 1980. See also ref 1f.

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(5) Cp₂Ti(COCH₃)Cl was formed by the reaction between Cp₂Ti(CO)₂ and CH₃COCl: Fachinetti, G.; Floriani, C.; Stöckli-Evans, H. J. Chem. Soc., Dalton Trans. 1977, 2297-2302.

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orients as 1, while in the related carbamoyl compounds Cp_2 *Th(CONR₂)Cl and Cp_2 *U(CONR₂)Cl,^{8a} both conformers are in equilibrium. Cp_2 *Th(η^2 -COC₆H₅)Cl has the η^2 O-inside structure, 2.^{8b} Furthermore, the biscyclopentadienyl actinides (Cp_2 *An) accomodate as many as two η^2 carbamoyl ligands,⁸ 3.



These are formally 20-electron compounds and a group 4 metal complex of this type is unlikely to be stable.

The recent mechanistic study of the carbonylation of Cp_2ZrR_2 enables us to view the geometrical choice between 1 and 2 in a more profound way.⁹ The O-inside conformer 2, as found in the X-ray analysis of the acyl compound, is not the initial kinetic product. Instead, CO insertion into the Zr-R bond seems to yield an intermediate, very possibly of structure 1, which subsequently isomerizes irreversibly to its thermodynamically more stable form 2.

Another interesting point to be mentioned here is the reactivity of such η^2 -acyl biscyclopentadienyl complexes, e.g., reactions with nucleophiles such as metal halides,¹⁰⁻¹² acyl to ketene or ketenimine transformations,¹³⁻¹⁵ ketone ligand formation,^{16,17} deprotonation of the acyl ligands,¹⁴ and reductive CO coupling.¹⁸ These features have often been attributed to the "oxycarbene" character of η^2 -acyl ligands, invoking a resonance structure **4b** which is thought to contribute to the bonding in addition to the η^2 -acyl structure **4a**.

The observed C–O stretching frequencies (ν_{CO}) of η^2 acyls may provide an argument in favor of the contribution of **4b**. Typical ν_{CO} 's for transition metal η^1 acyls range from 1630 to 1680 cm⁻¹, while those for η^2 acyls are in the range ~1530–1620 cm⁻¹. The decrease in ν_{CO} on forming η^2 acyl is significant in the case of Cp₂*Th(COR)X and Cp₂*Th(COR)R', (ν_{CO} ~1450–1480 cm⁻¹).^{2a,h} However, in the X-ray structures of Cp₂Ti(COCH₃)Cl,⁵ Cp₂Zr(COCH₃)(CH₃),^{1b} and Cp₂*Th(COCH₂'Bu)Cl,⁷ the C–O distances are not elongated as much as one might expect from the resonance form **4b**. There remain some doubts about significant participation of resonance form **4b** in the ground-state

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structures of these acyl complexes. Or it may be that there is



yet another metastable structure of an oxycarbene isomer 5, from which the carbene-type reactions could proceed.

CO Insertion Process

Lauher and Hoffmann have predicted that the initial attack of CO on d^0 Cp₂MR₂ (M = Ti and Zr) occurs along the line perpendicular to the Cp centroid-metal-Cp centroid plane, as marked by A in 6.¹⁹ This favorable approach was anticipated



6

from the spatial extention of the LUMO of $d^0 Cp_2ML_2$, approximately of a "y²" type, **7a**. The shape of the LUMO should allow for better overlap with the carbon lone pair (HOMO), **7b**, through a "lateral" approach A compared with the alternative "central" pathway B. Note that the coordinate system chosen for Cp_2ML_2 complexes in 6 and throughout this paper is different from the one defined in the paper by Lauher and Hoffmann.¹⁹ In ref 19, the choice is tied to C_{2v} symmetry, with z along the twofold axis instead of x.



Let us begin our theoretical analysis with a study of the CO insertion processes of the model molecules $Cp_2Zr(CH_3)_2$ and $Cp_2U(CH_3)_2^{2+}$. Since the extended Hückel method is not very well suited for describing energetics of bond-breaking and bond-formation steps, we examine the initial stages of a CO attack. The calculations were performed by varying the carbon (CO)to-metal distance, L, from 4.5 to 3.0 Å for $Cp_2Zr(CH_3)_2 + CO$ and from 5.0 to 3.5 Å for $Cp_2U(CH_3)_2^{2+} + CO$. The uranium complex with 2+ charge has a 5f⁰ 6d⁰ electronic configuration and corresponds to $Cp_2Th(CH_3)_2$.²⁰ At each point along the central attack pathway B, the CH_3 -M-CH₃ angle was optimized, while the geometry of the Cp_2M fragment was kept fixed. C_{2v} molecular symmetry was preserved throughout the association path. For the lateral approach A, we independently optimized



Figure 1. Total energy profiles for the two CO approaches toward $Cp_2Zr(CH_3)_2$ (top) and $Cp_2U(CH_3)_2^{2+}$ (bottom). The curves connecting filled circles are for a "lateral" approach A and those connecting open circles are for a "central" approach B.

the angular positions of all three ligands, two CH_3 groups and CO, thereby allowing for the best ligand arrangement within the xy plane as CO approaches.

Figure 1, top, gives the total energy curves for the CO attack on $Cp_2Zr(CH_3)_2$ as a function of the Zr-CO distance L. It is found that the lateral pathway A, accompanied by an appropriate relaxation of the methyl positions, is energetically favored. The result confirms that a CO molecule will indeed keep out of the central route B as Lauher and Hoffmann had suggested. The two curves for the Zr system go through a small barrier at about L = 3.5 Å. This is the region where closed-shell-closed-shell repulsions still outweigh the attractive interaction between the HOMO 7b and the LUMO 7a. Pathway A is characterized by stronger HOMO-LUMO bonding interaction, due to the better overlap. Therefore, the calculated activation energy of pathway A is smaller by about 0.2 eV (\sim 5 kcal/mol) than that of pathway **B.** The difference is not that much, but we think it will easily suffice to induce regiospecificity for the primary CO attack.²¹ On the left side of Figure 1, top the energy drops. In this connection we note that reaction enthalpies of -12.7 and -11.7 kcal/mol have been found for CO insertion into the Hf-R bond of Cp₂HfR₂ (R = CH_3 and $CH_2C_6H_5$).^{1b}

The potential energy curves for the two pathways of carbonylation of $Cp_2U(CH_3)_2^{2+}$ are shown in Figure 1, bottom. Again the lateral pathway A is slightly preferred. Our calculations give a barrier of 0.1 eV (~2 kcal/mol) for B, while attack along A does not encounter an appreciable barrier. The energy difference between the two approaches is even smaller than in the Zr case, e.g., 0.1 eV (~2 kcal/mol) at L = 4.25 Å. The regiospecificity

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Figure 2. Potential energy curves of Cp₂Ti(COCH₃)Cl, Cp₂U-(COCH₃)Cl²⁺, and Mn(CO)₅(COCH₃) as a function of M-C-C(CH₃) angle α . The experimentally observed angle α for Cp₂Ti(COCH₃)Cl, $Cp*_{2}Th[COCH_{2}C(CH_{3})_{3}]Cl, or [Mn(CO)_{4}(COCH_{3})(COPh)]^{-} is lo$ cated by a mark "Obs." in each corresponding curve.

for A, albeit weak, can be attributed again to the better overlap between the vacant orbital of type 7a of $Cp_2U(CH_3)_2^{2+}$ and the CO lone pair 7b. We have also computed the analogous potential energy curves for the U system with two more electrons, Cp₂U- $(CH_3)_2 + CO$, in which the U atom now has a $5f^2 6d^0$ configuration. The energy profile obtained is very similar to the one in Figure 1, bottom. The presence of the two additional electrons in f orbitals does not affect the choice of preferred stereocourse for CO attack.

We should note here that the calculated small energy barriers are in harmony with what we know from experiment. That is, CO insertion into $d^0 Cp_2 MR_2$ complexes is an extremely facile process. Although an activation energy for such a process has not been experimentally determined, CO insertion into Cp₂Zr- $(CH_3)_2$ is known to occur at temperatures as low as -130 °C in solution. Carbonylation of the actinide complexes might well be still more facile. In toluene solution at -80 °C, Cp_2 *Th(CH_3)₂ and $Cp_2^*U(CH_3)_2$ take up 2 equiv of CO (<1 atm) within 1 h.² Even a compound with a bulkier alkyl group, Cp2*Th(CH2'Bu)Cl, reacts rapidly with 1 equiv of CO at room temperature,⁷ and the reaction of $Cp_2*M[CH_2Si(CH_3)_3]Cl$ (M = Th, U) takes place at -78 °C.2a

CO attack via the favorable route A should lead to an immediate acyl product of "O-outside" structure 1. In contrast, the isolated acyl complexes of group 4 metals reveal only the alternative "O-inside" structure 2. This apparent contradiction was resolved by Erker and Rosenfeldt,⁹ who detected spectroscopically an intermediate of structure 1 in the carbonylation of $Cp_2Zr(p CH_3C_6H_4)_2$, $Cp_2Zr(p-CH_3OC_6H_4)$, and $Cp_2Zr(CH_3)_2$. The unstable intermediate isomerizes to structure 2 irreversibly, by a mechanism which will be analyzed later.

Acyl-Ligand Coordination

To investigate more closely the coordination modes of acyl ligands, we have calculated potential energy curves for acetyl pivoting in $Cp_2Ti(COCH_3)Cl$ and $Cp_2U(COCH_3)Cl^{2+}$. The model uranium complex could be a model for the known Cp_2 *Th(COCH₂'Bu)Cl. The first variable we vary is the M-C-CH₃ angle α , while all other geometrical parameters are still kept fixed for each molecule. Figure 2 superimposes the three energy curves, including for comparison also the energy of an acyl ligand in $Mn(CO)_5(COCH_3)$ so rotated.²² As Figure 2 shows, the



Figure 3. Contour plots of the two frontier orbitals, d_{π} and d_{σ} of a Cp_2TiCl^+ fragment. The orbitals are shown in the xy plane. The contour levels of each diagram are ± 0.01 , ± 0.02 , ± 0.055 , ± 0.1 , ± 0.2 , and ± 0.4 . The solid lines trace contours of one sign of ψ , the dashed lines of the opposite sign.

optimum mode of acyl coordination in $Mn(CO)_5(COCH_3)$ is the η^1 type ($\alpha = 128^\circ$). On the other hand, the acyl ligands of Cp₂Ti(COCH₃)Cl and Cp₂U(COCH₃)Cl²⁺ tend to bend toward an η^2 geometry, pronouncedly so in the latter molecule. The calculated trend is in good agreement with the experimentally observed structures of Cp₂Ti(COCH₃)Cl and Cp₂*Th-(COCH₂'Bu)Cl. Note that in calculating the curves in Figure 2 we assumed the orientation of the η^2 acyl (O-inside or O-outside) to be the same as observed for each. We will turn to the interesting choice between the two conformations of the acyl in a moment.

The diversity of optimal angles α in Figure 2 may be easily understood in terms of optimal interaction between frontier orbitals of each metal fragment and acetyl. Knowing that a bent d⁰ Cp₂M



unit has three low-lying vacant orbitals, $1a_1$, b_2 , and $2a_1$ (8),²⁰ one may derive the frontier orbitals of a d^0 Cp₂MX⁺ fragment. The addition of a ligand X ($X = R^{-}$, Cl⁻, etc.) destabilizes one combination and rehybridizes the two acceptor orbitals remaining at low energy.²³ One is a somewhat perturbed version of the $1a_1$ which is the LUMO of $d^0 Cp_2 MX^+$. Toward a fourth ligand L, an acyl group in this case, this LUMO exhibits π symmetry with respect to the M-L bond. For simplicity let us abbreviate this MO as d_{π} . The other acceptor orbital is locally of σ type, thus named d_{σ} here. The two low-lying vacant orbitals are plotted for Cp_2TiCl^+ in Figure 3.

An acyl ligand, CH₃CO⁻, brings two donor orbitals. The high-lying HOMO, n_, consists mostly of a carbon lone pair with an out-of-phase admixture of the oxygen p orbital. The lower donor orbital, n_+ , is an in-phase combination of the same orbitals, with a larger oxygen p component. The n_{-} and n_{+} orbitals, which are sketched in 9, find a symmetry match with the $Cp_2TiCl^+ d_{\pi}$ and d_{α} orbitals, respectively. The dominant interaction occurs between n_{-} and d_{r} ; this is, in fact, the major factor governing the η^2 coordination mode of acyl in Cp₂Ti(COCH₃)Cl and also in $Cp_2Zr(COCH_3)(CH_3).$

⁽²²⁾ Studies of the carbonylation of RMn(CO)₅ to give RCOMn(CO)₅ are numerous. For example, Closson, R. D.; Kozikowski, J.; Coffield, T. H. J.

^{2153-2174. (}b) Silver, M. E.; Eisenstein, O.; Fay, R. C. Inorg. Chem. 1983, 22, 759-770.



The in-phase combination of the n_{-} and d_{π} orbitals yields the HOMO of Cp₂Ti(COCH₃)Cl. The primary bonding is set by an overlap between the large lobe of d_{π} and the carbon lone-pair portion of n_{-} , as shown in 10. This represents the metal-to-carbon



 σ bonding for the O-inside conformer. There is a further interaction available between the O p in n₋ and the smaller lobe of d_{π}. This additional interaction will be increased as acyl pivots toward an n^2 structure, thus stabilizing the HOMO.

Figure 4 shows how the frontier orbitals evolve as a function of α . The HOMO level (2a') first goes down in energy as the acyl ligand bends from an η^1 geometry, falling to a minimum at $\alpha \approx 157^\circ$. This energy profile can be anticipated from the phase relationship of 10. The total energy curve for the acyl pivoting of Cp₂Ti(COCH₃)Cl in Figure 2, having a minimum at $\alpha \approx 165^\circ$, follows well the HOMO energy change. The empty 3a' orbital, which is the antibonding counterpart of 10, moves up steeply on going from η^1 to η^2 . In the course of its ascent 3a' crosses another empty orbital 2a". The 2a" stays low in energy over a wide range of bending angle α , and plays a critical role as the low-lying LUMO of the η^2 -acyl complexes governing their unique reactivity. It consists mostly of the acyl π^* orbital lying perpendicular to the xy plane, thus designated π^*_{CO} , stabilized through an interaction with an empty d orbital of Cp₂Ti(COCH₃)Cl, based on the character of its HOMO, parallels that of the isostructural η^2 -acyl complex, Cp₂Zr(COCH₃)(CH₃).

For the actinide molecule Cp₂U(COCH₃)Cl²⁺, we observed a similar stabilization of the HOMO as the acyl bends. Note that the acyl orientation is now O-outside. As shown in the Walsh diagram of Figure 5, the HOMO has a minimum at $\alpha \approx 173^\circ$; this is the primary reason the acyl ligand tends to bend toward an η^2 geometry (see the total energy curve in Figure 2). Again the HOMO comprises mostly the acyl n_orbital with U d orbitals as well as f orbitals mixing in. As a consequence of the additional f orbital contribution, the detailed shape of the HOMO is somewhat complicated and its contour map is drawn in 11, where





(e<

Energy

110

Figure 4. Walsh diagram for acyl pivoting (α) in Cp₂Ti(COCH₃)Cl. The orientation of acyl is O-inside.

150

a°

170

190

130



Figure 5. Walsh diagram for acyl pivoting (α) in Cp₂U(COCH₃)Cl²⁺ with a 5f⁰ electronic configuration. The acyl orients as O-outside.

the composition of the U orbitals in the HOMO is also shown. From the contour map of 11, it is clear how the η^2 -acyl structure stabilizes the HOMO. The carbon and oxygen orbital lobes in the acyl n₋ orbital overlap nicely with this U hybrid orbital. To



Figure 6. Walsh diagram for acyl pivoting (α) in Mn(CO)₅(COCH₃). The LUMO is at around -9 eV in energy and is not shown in the figure.

investigate how much U f orbitals participate in the net U-acyl bonding, we performed a population analysis for Cp₂U (COCH₃)Cl²⁺ at $\alpha = 170^{\circ}$. The calculated overlap populations associated with U f orbitals are 0.062 (U(f)-C) and 0.046 (U-(f)-O), while the total U-C and U-O overlap populations are 0.372 and 0.0723, respectively.

The LUMO in the Walsh diagram is very much similar to the 2a'' of $Cp_2Ti(COCH_3)Cl$ in Figure 4. It contains the $acyl \pi^*_{CO}$ orbital with a strong admixture of U f orbitals. Higher in energy there are seven U f orbitals. There is an interesting question as to where the electrons go into the system in the f^2 complexes, e.g., $Cp_2^*U(COR)Cl$. If they reside in the 2a'', then the CO bond should be much weakened. The observed CO stretching frequencies are 1546 and 1559 cm⁻¹ in $Cp_2^*Th[CO(N(CH_3)_2)]Cl$ and the analogous U derivative, respectively.^{8a} The trend is opposite to expectations based on 2a'' occupation. It may be that the CO vibration is strongly coupled with M–C and M–O modes, for these acyls are far from normal, or, alternatively, our 5f parameters put the mainly f orbitals too high, and really they should be below the 2a''.

The Walsh diagram for the acyl pivoting in Mn(CO)₅(COCH₃) is shown in Figure 6. In contrast to the group 4 metal and the actinide systems, the HOMO of Mn(CO)₅(COCH₃) moves to higher energy as the acyl ligand bends from the η^1 minimum at $\alpha \approx 131^\circ$. The reason behind this trend is rather straightforward. The Mn(CO)₅⁺ fragment, having a d⁶ electronic configuration, carries an acceptor orbital of a₁ symmetry, **12**,²⁴ while two d_π



orbitals are filled. The d_{π} orbitals are mutually orthogonal, and one of them is depicted in 13. The interaction of the occupied d_{π} 13 with the occupied n₋ of acyl turns out to be repulsive in the HOMO of Mn(CO)₅(COCH₃), when the acyl ligand coordinates to Mn in an η^2 mode. This reasoning carries over to an analysis of the geometrical preference of many other η^1 -acyl complexes, e.g., Fe(HBpz₃)(CO)₂(COCH₃),²⁵ Ni(PMe₃)₂Cl(COR),²⁶ and Rh(OEP)(COH).²⁷

An interesting conclusion follows from Figure 6. For an acyl complex with two electrons less than $Mn(CO)_5(COR)$, η^2 -acyl bonding should become feasible. The HOMO (3a') in Figure 6 is empty for the d⁴ electron system. Then the lower 2a' orbital, consisting of the in-phase combination of a metal d_π and the acyl n₋, governs the geometry of the coordinated acyl ligand. The 2a' molecular orbital is analogous to the HOMO (2a') of Cp₂Ti-(COCH₃)Cl in Figure 4, which makes the η^2 -acyl structure possible.

Indeed acyl compounds of this type, $V(\eta^2-COC_3Ph_3H_2)$ -(CO)₃(Ph₂AsCH₂CH₂PPh₂),^{28a} $V(\eta^2-COPh)(CO)_3(dppe)$ (14a),^{28b} and Mo(η^2 -COCH₂SiMe₃)Cl(CO)(PMe₃)₃ (14b),^{28c} have been synthesized. Interestingly, in Mo(COCH₃)(S₂CNMe₂)(CO)-(PMe₃)₂ (14c) the acyl ligand does not pivot toward an η^2 form,



but moves in the opposite direction by virtue of a strong interaction with a β -C-H bond of acyl.^{28c} Not unrelated to the η^2 -acyl complexes of the d⁴ M(COR)L₅ is d⁶ Ru(η^2 -COR)I(CO)(PPh₃)₂ (14d),^{28d} where a d_{\pi} orbital in the equatorial plane of the RuI-(CO)(PPh₃)₂⁺ fragment is vacant and behaves like the d_π of Cp₂TiCl⁺ in Figure 3. The electronic structures and bonding of the vanadium complexes will be discussed elsewhere.²⁹

O-Inside vs. O-Outside Conformations

Let us now turn to the interesting geometrical choice between the O-outside 1 and O-inside 2 isomers. In an attempt to compare energies of the two conformers of Cp₂Ti(COCH₃)Cl, we have calculated potential energy surfaces varying the Ti-C-C angle α and the Cl-Ti-C angle β independently, as shown in 15a and 15b. The parameter α is the one used for plotting the energy curve of Figure 2. The geometry of the Cp₂TiCl portion was common and kept fixed for both isomers. The calculated surfaces

⁽²⁴⁾ The orbitals of M(CO), fragments have been extensively described. See, for instance: (a) Elian, M.; Hoffmann, R. *Inorg. Chem.* 1975, 14, 1058–1076. (b) Hoffmann, R. *Science* 1981, 211, 995–1002. (c) Hoffmann, R. *Angew. Chem.*, *Int. Ed. Engl.* 1982, 21, 711–724.

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^{(26) (}a) Huttner, O.; Orama, O.; Bejenke, V. Chem. Ber. 1976, 109, 2533-2536. (b) Carmona, E.; Gonzalez, F.; Poveda, M. L.; Atwood, J. L.; Rogers, R. D. J. Chem. Soc., Dalton Trans. 1980, 2108-2116.

⁽²⁷⁾ Wayland, B. B.; Woods, B. A.; Pierce, R. J. Am. Chem. Soc. 1982, 104, 302-303.

^{(28) (}a) Franke, U.; Weiss, E. J. Organomet. Chem. 1979, 165, 329-340.
(b) Schiemann, J.; Weiss, E. Ibid. 1983, 255, 179-191. (c) Carmona, E.; Sanchez, L.; Marin, J. M.; Poveda, M. L.; Atwood, J. L.; Priester, R. D.; Rogers, R. D. J. Am. Chem. Soc. 1984, 106, 3214-3222. (d) Roper, W. R.; Taylor, G. E.; Waters, J. M.; Wright, L. J. J. Organomet. Chem. 1979, 182, C46-C48.

⁽²⁹⁾ Schiemann, J.; Weiss, E.; Hofmann, P., in preparation.



are presented in Figure 7a and Figure 7b for the O-outside and O-inside conformations, respectively. The potential surface of Figure 7b has a minimum at $\alpha = 165^{\circ}$ and $\beta = 112^{\circ}$, corresponding to the O-inside η^2 structure 2. This minimum is 0.39 eV (9 kcal) more stable than that of the O-outside η^2 structure 3 in Figure 7a, which appears at $\alpha = 165^{\circ}$ and $\beta = 85^{\circ}$. The result agrees well with the experimental finding that only O-inside structures were found by the X-ray structure analyses for Cp₂Ti(COCH₃)Cl and for its Zr congener, CpZr(COCH₃)(CH₃).

However, the most interesting feature of the energy surface of Figure 7a is the presence of two potential minima for the O-outside conformation. One is for the aforementioned η^2 type 1, and the other corresponds to an η^1 -acyl coordination, appearing at $\alpha = 100^\circ$ and $\beta = 120^\circ$. This finding will be important when we consider the full reaction pathway of Cp₂Zr(CH₃)₂ + CO later.

Before examining the electronic reasons for this unexpected existence of an η^1 minimum, we take a look at the origin of the different stability of the two η^2 structures, **1** and **2**. The orientational preference is again a consequence of the better overlap between the occupied acyl n₋ (**9**) and the vacant d_{π} (Figure 3) of the d⁰ Cp₂TiCl⁺ fragment. For the O-inside orientation, the larger lobe of d_{π} can overlap with the larger lobe on carbon of n₋. The smaller lobes of both frontier orbitals also retain a good overlap with each other. On the other hand, the O-outside orientation has a smaller d_{π}-n₋ interaction, because the larger d_{π} lobe now faces the smaller lobe on oxygen of n₋ and vice versa. This situation is drawn schematically in **16a** and **16b**, where the



calculated overlaps, S, between the d_{π} and n₋ orbitals for the two acyl orientations are also compared. The different magnitudes of the overlap integrals lead to a different stability of the HOMO of Cp₂Ti(COCH₃)Cl as shown in the interaction diagrams for both orientations in Figure 8. The HOMO of the O-inside isomer is more stabilized, owing to the better d_{π} -n₋ interaction, which in turn gives a lower total energy.

Next we consider an actinide complex, $Cp_2U(COCH_3)Cl^{2+}$. Figure 9 shows the computed potential energy surfaces as a function of the U–C–C angle α and the C–U–Cl angle β for the two acyl conformations. The two surfaces are similar in shape to those for Cp₂Ti(COCH₃)Cl in Figure 7. However, the Ooutside minimum at $\alpha = 170^{\circ}$ and $\beta = 96^{\circ}$ is now lower in energy than the O-inside minimum at $\alpha = 170^{\circ}$ and $\beta = 112^{\circ}$. The energy difference is merely 0.03 eV, 0.7 kcal/mol, and the stabilities of the two minima are essentially equal. Experimentally Cp₂*Th(COCH₂'Bu)Cl is known to assume the O-outside configuration, while for the related carbamoyl complexes, Cp*2M- $(CONR_2)Cl (M = Th, U)$, the two conformations coexist either in solution or in crystal. $Cp_{2}^{*}Th(COC_{6}H_{5})Cl$ has an O-inside structure.^{8b} The computed potential surface for the O-outside conformation contains an additional minimum at $\alpha = 105^{\circ}$ and $\beta = 120^{\circ}$, though this minimum is a shallow one. It corresponds to an η^1 acyl structure, just as in the Ti case.



Figure 7. Potential energy surfaces as a function of α and β for the O-inside (top) and O-outside (bottom) orientations of acyl in Cp₂Ti-(COCH₃)Cl. The point marked by X in b indicates the experimentally observed structure of Cp₂Ti(COCH₃)Cl.



Figure 8. Interaction diagrams for the frontier orbitals of Cp₂TiCl⁺ and COCH₃⁻ in the η^2 O-outside (left) and η^2 O-inside (right) structures of Cp₂Ti(COCH₃)Cl.

The Cp₂UCl³⁺ fragment carries a vacant d_{π} orbital, 17, like the one plotted for Cp₂TiCl⁺ in Figure 3. Although the shape of 17 is essentially analogous to the Cp₂TiCl⁺ d_{π} orbital, its two lobes pointing toward the vacant site are nearly the same in size. Thus the two acyl orientations should not differ in their d_{π}-n₋ interactions as much as they do in the Ti complex. This is probably one reason why O-outside and O-inside conformers of Cp₂U-(COCH₃)Cl²⁺ are energetically well balanced. There appears to be another factor coming in. The U atom has vacant f orbitals



(in addition to d_{π}) which can interact with the occupied acyl norbital. The outcome is a complicated hybridization of d and f orbitals in the HOMO of Cp₂U(COCH₃)Cl²⁺, the contour map of which has been shown in **11** for the O-outside orientation. The f orbitals mix into the HOMO in such a way that the hybrid-noverlap becomes nearly equal for the two conformers. One may see in **11** that reversing the acyl orientation from O-outside to O-inside will not weaken the interaction between U and acyl.

We now come back to the unexpected double minimum found in the potential energy surfaces of the O-outside conformers in Figures 7a and 9a. Our eventual aim in this portion of the study is to gain insight into Erker's experiment on the CO insertion process of Cp_2ZrR_2 , so that we shift our theoretical analysis from the Ti system to $Cp_2Zr(COCH_3)CH_3$ at this point. As will be shown soon, our conclusions based on the calculations for $Cp_2Ti(COCH_3)Cl$ carry over well to the Zr case.

To check if the energetics obtained for Cp₂Ti(COCH₃)Cl are not particular to the Ti system, we computed also the potential surfaces for O-outside and O-inside isomers of Cp₂Zr-(COCH₃)(CH₃). The geometrical optimization was upgraded by allowing both CH₃ and COCH₃ ligands to move, thus varying independently three parameters, α , β' , and γ , as defined in 18.



The sum of the new variables, β' and γ , corresponds to β in **15a** and **15b**. Again one η^2 minimum was found for **18b** at $\alpha = 170^\circ$, $\beta' = 55^\circ$, and $\gamma = 60^\circ$, while for **18a** two minima showed up at $\alpha = 110^\circ$, $\beta' = 55^\circ$, and $\gamma = 50^\circ$, and at $\alpha = 170^\circ$, $\beta' = 25^\circ$, and $\gamma = 60^\circ$, being the η^1 and η^2 structures, respectively. The η^2 O-inside geometry is most stable of all. The η^2 O-outside is about 5 kcal/mol higher in energy than that, and the η^1 O-outside is still less stable by 5 kcal/mol.

The presence of two minima for the O-outside conformation means that there is an energy barrier on going from η^1 to η^2 , and vice versa. The η^1 geometry carries 16 electrons at the metal center. Then the attachment of the oxygen lone pair to the electron-deficient metal center, in the η^2 coordination mode, makes the molecule an 18-electron compound. It is rather surprising that, depending on the acyl orientation, the acyl pivoting, an interconversion between 16- and 18-electron systems, should pass through a barrier. Let us try to understand the reason behind this. In the η^1 structures, the phase relationship for the important d_{π} -n_ interaction is set so as to form an overlap between the large lobe of d_{π} and the carbon lobe of acyl n_. The HOMO of each



Figure 9. Potential energy surfaces as a function of α and β for the O-inside (top) and O-outside (bottom) orientation of acyl in Cp₂U-(COCH₃)Cl²⁺. The point marked \times is the experimentally observed structure of Cp*₂Th[COCH₂C(CH₃)₃]Cl.

 η_1 complex, O-outside or O-inside, is made up of this bonding combination of d_{π} and n_{-} ; see **19a** and **20a**.



The notable difference between 19a and 20a lies in their nodal properties. It is clear from 20a and 20b that the η^1 O-inside structure relaxes to the η^2 structure without destroying the phase-match between d_{π} and n_{-} . The $\eta^1 \rightarrow \eta^2$ transformation occurs smoothly, simply increasing the interaction between the small lobe of d_{π} and the oxygen lobe of n_{-} . Therefore, the O-inside conformer ends up with a single minimum of the η^2 type. On the contrary, the $\eta^1 \rightleftharpoons \eta^2$ interconversion for the O-outside case involves a phase discontinuity. The carbon lobe has to switch its phase during the acyl pivoting, and the same for the oxygen lobe. at the place where the phase discontinuity takes place, the molecule loses its d_{π} - n_{-} bonding interaction, thus causing an energy barrier.

The origin of the barrier described above is evident in the Walsh diagram of Figure 10 for the O-outside $\eta^1 \rightleftharpoons \eta^2$ process. The HOMO on each side, which is **19a** or **19b**, tends to correlate to the LUMO on the other side. Each HOMO ascends in energy until they meet each other at a maximum inbetween the η^1 and η^2 energy wells. What actually happens is that there is a strongly avoided crossing between the LUMO and the HOMO, because of the low C_s molecular symmetry. The HOMO energy profile is responsible for the double minimum found in the potential energy surfaces for the Cp₂M(COCH₃)X complexes.

Noteworthy here is the metal-oxygen antibonding interaction seen in **19a**. A consequence is the large M-C-O angle of \sim 130-135° at the η^1 O-outside minima found by our calculations



Figure 10. Orbital correlation diagram for the interconversion between η^1 and η^2 structures of the O-outside acyl orientation in Cp₂Zr-(COCH₃)(CH₃).

on the Ti and Zr complexes. Incidentally, a related Zr complex, $Cp_2Zr(CH_2PPh_2)Cl$, has been recently synthesized.³⁰ Its X-ray diffraction study shows that the CH_2PPh_2 ligand is bound to Zr in an η^1 manner, as indicated in **21**. This complex is analogous



to the η^1 O-outside structure **19a** of Cp₂Zr(COCH₃)CH₃. The CH₂PPh₂⁻ ligand has a HOMO made out of an out-of-phase combination of carbon and phosphorus lone-pair orbitals, which is very similar to the n_{-} orbital of COCH₃⁻. Because of the antibonding interaction between the d_{π} lobe and the phosphorus lone pair in 22, the P atom tends to stay away from the electron-deficient Zr center with its 16-electron count, a result which would seem puzzling if it were not for these orbital symmetry arguments. The opening up of the Zr-C-P angle from its ideal tetrahedral angle of 109° is a result. The observed Zr-C-P angle is as large as 130°.³⁰ For details of the theoretical analysis of $Cp_2Zr(CH_2PPh_2)Cl$, the reader should refer to ref 23a where alternative geometries and steric effects for 21 have been discussed in detail. A double-minimum situation similar to the one described above has also been found for Cp₂Zr(S₂CH)Cl by Silver, Eisenstein, and Fay.23b

The η^2 O-Outside to η^2 O-Inside Rearrangement

As discussed in an earlier section of this paper, we believe a CO molecule attacks the central metal at a lateral site of Cp₂MRX or Cp₂MR₂. For M = Zr, the isolable product assumes the η^2 O-inside structure, **2**, so that the formation of an immediate kinetic product η^2 O-outside, **1**, should be followed by an isomerization





Figure 11. Computed energy profile for $Cp_2Zr(COCH_3)(CH_3)$ along the proposed isomerization pathway from the η^2 O-outside structure, which is the kinetic product of CO insertion into $Cp_2Zr(CH_3)_2$, to the thermodynamically more stable form, η^2 O-inside. Energies are shown in kcal/mol units.

Scheme I



step leading to the thermodynamically more stable structure 2. We want to understand here how the isomerization takes place.

The simplest isomerization process, i.e., the direct rotation of the η^2 acyl about the axis *m* in Scheme I, is not likely to occur. A bent Cp₂M fragment is known to form strong bonds with ligands only when the coordination occurs in the *xy* plane.¹⁹ It is clear why this is so: the three frontier orbitals of Cp₂M, 8, all lie in the plane. Thus the rotation of acyl about *m* would result in disruption of much of the bonding interaction between acyl and Cp₂MR, importantly the n_-d_{\pi} interaction. As a result, the calculated barrier to that rotation is fairly large, as large as 2.5 eV (58 kcal/mol) for Cp₂Ti(COCH₃)Cl and only slightly smaller than that in the case of Cp₂Zr(COCH₃)(CH₃).

We now propose an alternative route via the η^1 O-outside intermediate, 23, which was unexpectedly found in our potential energy surfaces for acyl O-outside conformations of Cp₂Ti-(COCH₃)Cl and of Cp₂Zr(COCH₃)(CH₃). The potential energy surfaces reveal that the transformation of 1 to 23 is an easier process than the direct rotation, requiring 7.5 kcal/mol for Cp₂Zr(COCH₃)(CH₃),³¹ and about 14 kcal/mol for Cp₂Ti-(COCH₃)Cl.

The subsequent step from 23 to 2 consists of a rotation around the M-C single bond, passing through the "perpendicular" η^1 structure, 24, as a transition state. To obtain a theoretical estimate of the rotation barrier, we optimized the transition state 24 for Cp₂Zr(COCH₃)(CH₃), by varying the angles β' and γ as we did in 18b, as well as the Zr-C-CH₃ angle α and the Zr-C-O angle α' . The computed barrier is 8.5 kcal/mol, much smaller than the rotation about the axis *m*, as one might easily anticipate from the fact that the former is a rotation about the Zr-C single bond. At the transition state, 24, the Zr-C-CH₃ angle α opens up slightly to 130°, in order to avoid a contact between the Cp ring and the

⁽³¹⁾ In the calculations on the $Cp_2Zr(COCH_3)(CH_3)$ system, the position of the metal-bound methyl group was also optimized independently.



methyl group of acyl. The other optimized numbers include 120° for α' and 95° for $\gamma + \beta'$. The overall energy profile is now in our hands for the $1 \rightarrow 2$ isomerization of Cp₂Zr(COCH₃)(CH₃). Figure 11 summarizes it in kcal/mol units. The net activation energy on going from the elusive η^2 O-outside structure to the thermodynamically stable η^2 O-inside form amounts to 13.5 kcal/mol. The experimentally observed activation free energy for the isomerization of Cp₂Zr(COR)(R) ranges from 11.4 kcal/mol (R = CH₃) to 15.6 kcal/mol (R = *p*-tolyl or *p*-anisyl).⁹ The agreement between the theoretical estimates and the experimental observations is surprisingly good, considering the approximate nature of our calculations.

For Cp₂Ti(COCH₃)Cl, there appeared to be steric problems at the transition state analogous to 24. A rigid rotation of COCH₃ from η^1 O-outside meets a prohibitively high barrier of ~ 2.5 eV, due to the steric congestion associated with the smaller ionic radius of Ti compared with Zr. A better geometrical optimization could result in a reduction of the barrier,³² but we think that the calculated large barrier provides us with an interesting hypothesis. If an acyl complex of the type $Cp_2M(COR)X$ (M = group 4 metals) is not allowed to undergo the rotation for steric reasons, then the CO insertion product of Cp2MRX may be unable to reach the stable O-inside η^2 -acyl product. Two possibilities ensue. The metastable η^2 O-outside complexes of group 4 metals could be isolated. Alternatively, if the steps are reversible, the CO insertion process itself might not be seen, and other reaction channels followed. As a matter of fact, we know of no experimental evidence that acyl complexes of Ti, Cp₂Ti(acyl)X, are formed by CO insertion of Cp₂TiRX. Remember that Cp₂Ti(COCH₃)Cl was synthesized from Cp₂Ti(CO)₂ by a reaction with CH₃COCl.⁵

The isolated product of Cp2*Th[COCH2'Bu]Cl possesses an η^2 O-outside geometry.⁷ This fact and the potential energy surfaces in Figure 9 for the model complex, $Cp_2U(COCH_3)Cl^{2+}$, indicates that the kinetic η^2 O-outside product of CO insertion is thermodynamically stable too. The trimethylsilylmethyl derivatives, $Cp*_2M[COCH_2Si(CH_3)_3]Cl (M = Th, U)$, which were formed by the reaction of $Cp*_2M[CH_2Si(CH_3)_3]Cl$ with CO, were de-tected by ¹H NMR at low temperature.³³ Although their detailed structures are yet unknown, the molecules have been assigned bihaptoacyl structures. Not to be ruled out is the possibility that the two isomeric bihaptoacyl structures, η^2 O-outside and η^2 O-inside, of these actinide complexes are in equilibrium in solution. If the equilibrium exists, it is likely to occur via an η^1 O-outside intermediate, just as we think occurs for the zirconium congeners. Figure 9 shows that the η^1 O-outside minimum is 0.55 eV (12.6 kcal/mol) less stable than the η^2 O-outside structure. The activation energy for the $\eta^2 \rightarrow \eta^1$ transformation is about 0.55 eV as well, since the η^1 potential well is shallow. The barrier to the subsequent rotation leading to an η^2 O-inside structure would be sensitive to the bulk of acyl substituents and that of Cp*. Unfortunately, the synthesized actinide complexes are too large to treat by our calculations, but the potential surfaces of Figure 9 should aid us in considering the isomerization process. For the related carbamoyl complexes, $Cp_{2}M[CON(CH_{3})_{2}]Cl$ (M = Th,

U), the two η^2 conformations, **25a** and **25b**, are essentially equal



in free energy in solution. The observed activation free energy ΔG^* for their interconversion is ca. 10 kcal/mol.⁸ The transformation from η^2 O-outside to η^1 O-outside of the carbamoyl complexes would require less energy than the one of the model Cp₂U(COCH₃)Cl²⁺, because of the π conjugation present between C-O and C-N bonds of the carbamoyl ligand.

Reactivity of the η^2 -Coordinated Acyls

We finally analyze the origin of the unique reactivity of acyl ligands in Cp₂M(COR)X. As mentioned in the Introduction, such η^2 -acyl biscyclopentadienyl complexes exhibit a rich and promising chemistry. The dominant features of their reactivity have often been rationalized in terms of a contribution of the oxycarbene-type resonance hybrid **4b** to their ground-state properties, lending carbenoid character to the η^2 -coordinated acyl ligands. In view of strong oxophilicity of group 4 metal and actinides,³⁴ one might anticipate that acyl ligands could adopt still another mode of coordination, as shown in **5**, where only an oxygen atom of the acyl is attached to the d⁰ central metal, leaving a carbon site free.

In order to test whether, in fact, the alternative structure can be an intermediate in the many reactions associated with coordinated acyl ligands, we carried out a geometrical optimization for the two conformations, **26** and **27**, while the geometry of the acyl portion is unchanged, with a Zr–O distance of 2.15 Å. The electronic configuration of the carbene part is assumed to be singlet.³⁵ The structures **26** and **27** formally derive from carbon-metal bond disruption in the η^2 O-outside and η^2 O-inside structures, respectively. They are interrelated by a simple rotation



about the Zr-O bond, and rotation about the C-O bond from them gives two more isomers, **28** and **29**. From our geometrical search,

⁽³²⁾ In order to attenuate the steric problems at the transition state, we intentionally elongated the Ti-C(Cp) and Ti-C(acyl) distances from 2.388 to 2.5 Å, and from 2.07 to 2.197 Å, respectively. The longer distances are those used for calculations on Cp₂Zr(COCH₃)CH₃. The rotational barrier was decreased to 0.9 eV (21 kcal/mol), which is still higher in energy than the Zr case.

⁽³³⁾ See ref 2a, and ref 6 and 33 cited in that paper.

^{(34) (}a) McDade, C.; Green, J. C.; Bercaw, J. E. Organometallics 1982, 1, 971–973. (b) Dias, A. R.; Salema, M. S.; Simoes, J. A. M. Ibid. 1982, 1, 971–973. (c) See ref 2, 3a, and Bruno, J. W.; Marks, T. J.; Morss, L. R. J. Am. Chem. Soc. 1983, 105, 6824–6832.

⁽³⁵⁾ The EH-HOMO/LUMO gap is about 1 eV for the zirconoxycarbene structures, as found for alkyl-alkoxycarbenes, which are singlet ground states. See, for instance: Davidson, E. R. "Singlet Triplet Energy Separations in Carbenes and Related Diradicals"; In "Diradicals"; Borden, W. T., Ed.; Wiley: New York, 1982; p 73.

we obtained the following results: (i) Each zirconoxy carbene structure, 26 or 27, is a local minimum of the potential energy surface. (ii) The optimal Zr–O–C angles θ are both close to 180° so that there is no practical difference in structure between 26 and 29, and between 27 and 28.36 (iii) In our model calculations, the energies of 26 and 27 are both very high, ca. 2 eV higher than those of corresponding η^2 acyl geometries.

Although Cp₂Zr(COCH₃)CH₃ has local minima corresponding to zirconoxy carbene structures, it is a high-energy process for the molecule to move from its ground-state η^2 -acyl structure to these metastable states. Therefore, the so-called carbene-type reactivity of such η^2 -coordinated acyls is probably not derived from a preequilibrium with the oxycarbene isomers, 26-29, which could subsequently undergo the observed reactions. Instead, the electronic properties of the ground-state η^2 -acyl structures themselves should be responsible for the unique reactivity of acyl groups attached to $d^{\bar{0}} Cp_2 MX$ fragments.

Recall here the molecular orbital diagrams of Cp2Ti(COCH3)Cl in Figures 4 and 8, of $Cp_2U(COCH_3)Cl^{2+}$ in Figure 5, and of $Cp_2Zr(COCH_3)CH_3$ in Figure 10. A common feature of these diagrams is that they all possess a low-lying LUMO, 30, made



up of the acyl C–O π^*_{CO} which is perpendicular to the acyl plane. The LUMO may appear even lower in energy than the π^*_{CO} orbital of acyl itself, because metal xz and/or yz orbitals can mix in π^*_{CO} , thus pushing it *down*. The presence of a low-lying vacant π^*_{CO} makes the acyl carbon strongly electrophilic, which accounts for the reactivity of these d⁰ η^2 -acyl compounds. This is in contrast to acyl complexes of the later transition metals where d orbitals, being capable of interacting with acyl π^*_{CO} , are usually low in energy and are occupied by electrons, thereby pushing $\pi^*_{CO} up$. The higher positioning of the π^*_{CO} level weakens the electrophilic nature of the acyl carbon. Therefore, the observed reactivity of η^2 -acyl complexes is attained only when acyl ligands are bound to electron-deficient early-transition metal (and actinide) fragments, namely d⁰ Cp₂MX.

Then can we say that the η^2 -coordinate-acyl ligands in Cp₂M-(acyl)X complexes have "carbene" character, as has often been invoked? We must first clarify what "carbene" character means. Obviously there is a similarity between an η^2 -coordinated acyl and a carbene in that both contain a low-lying LUMO, 30 vs. 31, the large component of which is a carbon p_{π} . However, resonance form 4b is somewhat misleading, because it might lead to an expectation that "carbene-type" behavior would also manifest itself in an elongated C=O bond. The experimental observations available so far argue against this expectation. No visible C=O elongation was observed in the X-ray structures of $Cp_2Ti-(COCH_3)Cl$ (1.18 Å), $Cp_2Zr(COCH_3)(CH_3)$ (1.211 Å), and $Cp*_{2}Th[COCH_{2}C(CH_{3})_{3}]Cl$ (1.18 Å). Indeed the calculated C=O overlap populations, P(C=O), tell us that the C=O bond of an η^2 acyl is as strong as that of an η^1 acyl. This bond even gets slightly stronger on going from η^1 to η^2 , e.g., for Cp₂Ti-(COCH₃)Cl, $P(C=O, \eta^1 O$ -outside) = 1.005, $P(C=O, \eta^2 O$ -outside) = 1.029, and $P(C=O, \eta^2 O$ -inside) = 1.028. The trend is what one would expect from our bonding picture presented in an earlier section. Coordination of acyl in an η^2 manner takes electrons out of a C-O antibonding orbital through the n_-d_{π} interaction, 10, (and n_-f interactions as well as in the actinide case, 11).³⁷ On the other hand, there is no filled metal orbital

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Table I. Extended Hückel Parameters

		exponents ^a		
orbital	H_{ii} , eV	ζ1	\$2	
Ti 4s	-8.97	1.075		
Ti 4p	-5.44	0.675		
Ti 3d	-10.81	4.550 (0.4206)	1.40 (0.7839)	
Mn 4s	-8.63	1.8		
Mn 4p	-5.06	1.8		
Mn 3d	-11.59	5.15 (0.5311)	1.90 (0.6479)	
Zr 5s	-9.87	1.817		
Zr 5p	-6.76	1.776		
Zr 4d	-11.18	3.835 (0.6211)	1.505 (0.5796)	
U 7s	-5.50	1.914		
U 7p	-5.50	1.914		
U 6d	-5.09	2.581 (0.7608)	1.207 (0.4126)	
U 5f	-9.01	4.943 (0.7844)	2.106 (0.3908)	
U 6p	-30.03	4.033		
C 2s	-21.4	1.625		
C 2p	-11.4	1.625		
O 2s	-32.3	2.275		
O 2p	-14.8	2.275		
Cl 3s	-30.0	2.033		
Cl 3p	-15.0	2.033		
H 1s	-13.6	1.30		

^aSlater-type orbital exponents. The numbers of parentheses are coefficients used in the double- ζ expansion.

available to transfer electrons back to the empty π^*_{CO} orbital of acyl.

Since the so-called "carbene-type" reactivity may be accounted for by the low-energy position of π^*_{CO} , the ensuing electrophilicity and the positive charge of the acyl carbon, without the C=Oelongation implied by 4b, perhaps it would be more appropriate to describe the reactivity of the η^2 acyls as "carbenium-type".

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Appendix

The calculations were carried out using the extended Hückel method,³⁸ with parameters as listed in Table I. Exponents for the Ti and Mn orbitals are from the works of Richardson et al.³⁹ and of Summerville and Hoffmann, 40 respectively. For Zr the exponents are those given by Basch and Gray.⁴¹ Metal H_{ii} 's for Ti⁴² and Mn⁴³ were taken from other work, while those for Zr were determined by charge iterative calculations on $Cp_2Zr(\eta^1$ -CH₂PPh₂)Cl.²³ The U parameters are taken from previous work,⁴⁴ while those for C, O, Cl, and H are standard ones. A modified Wolfsberg-Helmholz formula⁴⁵ for H_{ij} was used throughout the calculations.

Assumed geometries not listed in the text are as follows: $Cp_2Ti(COCH_3)CI$: Ti-C(Cp), 2.388 Å; Ti-C(acyl), 2.07 Å; Ti-Cl, 2.49 Å; C-O, 1.18 Å; C-C(acyl), 1.47 Å, C-H(acyl), 1.09 Å; C-C(Cp), 1.42 Å; C-H, 1.09 Å; Cp-Ti-Cp, 132°; O-C-C-

(37) The $n_+ - d_+$ interaction weakens the acyl C-O bond, but this effect is

cancelled out by the stronger n.-d. interaction.
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⁽³⁶⁾ Note in this connection the M-O-M angles in [Cp₂Zr(CH₃)]₂O and related compounds, which fail in the range of 169-180°: Hunter, W. E.; Hrncir, D. C.; Vann Bynum, R.; Penttila; R. A.; Atwood, J. L. Organometallics 1983, 2, 750-755, and references therein.

(acyl), 126°. Mn(CO)₅(COCH₃): MnC(CO), 1.82 Å; C–O(CO), 1.14 Å; Mn–C(acyl), 2.07 Å; C–O(acyl), 1.21 Å; C–C(acyl), 1.51 Å; C–H, 1.09 Å; O–C–C(acyl), 115°. Cp₂Zr(COCH₃)(CH₃): Zr–C(Cp), 2.5 Å; Zr–C(CH₃), 2.336 Å; Zr–C(acyl), 2.197 Å; C–O, 1.211 Å; C–C(acyl), 1.492 Å; C–H(CH₃), 1.1 Å; C–H(Cp), 1.08 Å; C–C(Cp), 1.40 Å; Cp–Zr–Cp, 130°; O–C–C(acyl), 120°. Cp₂U(COCH₃)Cl: U–C(Cp), 2.804 Å; U–C(acyl), 2.44 Å; U–Cl, 2.67 Å; C–O, 1.18 Å C–C(acyl), 1.47 Å; C–C(Cp), 1.42 Å; C–H, 1.09 Å; Cp–U–Cp, 138°; O–C–C(acyl), 118°.

Registry No. $Cp_2Zr(CH_3)_2$, 12636-72-5; $Cp_2U(CH_3)_2^{2+}$, 87136-09-2; $Cp_2Ti(COCH_3)Cl$, 66320-88-5; $Cp_2U(COCH_3)Cl^{2+}$, 96729-30-5; Mn-(CO)₅(COCH₃), 13963-91-2; $Cp_2^*Th[COCH_2C(CH_3)_3]Cl$, 74587-36-3; [Mn(CO)₄(COCH₃)(COPh)]⁻, 55318-15-5; Cp_2TiCl^+ , 96729-31-6; $Cp_2Zr(COCH_3)(CH_3)$, 60970-97-0; CO, 630-08-0.

Why the Energetic Minimum Aluminum Vinylidene Is Not Observed in Low-Temperature Aluminum + Acetylene Reactions

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Abstract: Ab initio molecular electronic structure theory has been used to investigate the Al + C_2H_2 potential energy surface. Particular emphasis was placed on resolving the apparent conflict between theory and experiment by examining the barrier to rearrangement between *cis*-AlHCCH (observed by ESR at 4 K) and AlCCH₂ (theoretically predicted global energy minimum). Analytic SCF gradients were employed with a double- ζ basis set to locate and characterize stationary points on the energy surface. Single point CI calculations using a double- ζ + polarization basis set have been carried out at the DZ-SCF stationary points. A DZP-CI barrier of 39.1 kcal for the *cis*-AlHCCH \rightarrow AlCCH₂ isomerization has been found. Such a barrier is clearly prohibitive to this isomerization under the experimental conditions of 4 K. By comparing this isomerization to the bare HCCH \rightarrow CCH₂ isomerization, the effect of the aluminum atom has been found to be significant.

The synthesis of the aluminum atom-acetylene adduct which was first reported by Kasai, McLeod, and Watanabe¹ and the more recent ESR analysis of this adduct by Kasai² suggest strongly the formation of a σ -bonded vinyl structure, **3**. Kasai's second paper² also discusses a trans \rightarrow cis ($3 \rightarrow 2$) conversion of this σ -bonded complex brought about by photoirradiation. A perhaps surprising feature is that no evidence was found to indicate the formation of the vinylidene complex, **1**, which has been predicted theoretically to be the global minimum on the Al-C₂H₂ energy surface.³



The purpose of the present research was twofold: first, to re-examine the vinylidene complex and the cis and trans (to the aluminum atom) σ -bonded vinyl radicals (structures 1, 2, and 3, respectively) with analytic gradient techniques;⁴ and second, to investigate the barrier to $2 \rightarrow 1$ rearrangement. As speculated in the previous theoretical study,³ the presence of a large barrier to rearrangement would prohibit the formation of the low-energy vinylidene structure at the liquid helium temperatures of the ESR experiments.^{1,2}

Table I. Calculated DZ-SCF Harmonic Vibrational Frequencies (in cm^{-1}) for AlCCH₂ (1), *cis*-AlHCCH (2), and *trans*-AlHCCH (3) and the $2 \rightarrow 1$ Transition State^{*a*}

AICCH ₂		cis-AlHCCH		trans-AlHCCH	
3360 3272 1796 1543 1109 1048 520 239 183	CH ₂ a stretch CH ₂ s stretch ν_{C-C} CH ₂ scissors CH ₂ wag CH ₂ rock ν_{AI-C} $\tau_{AI-C_{a}-C_{\beta}}$	3369 3238 1671 1286 1019 973 568 565 212	$\begin{array}{c} \nu_{C_{\beta}-H_{\beta}} \\ \nu_{C_{\alpha}-H_{\alpha}} \\ \nu_{C-C} \\ \delta_{AI-C_{\alpha}-H_{\alpha}} \\ \tau_{H-C-C-H} \\ \delta_{H-C-C-H} \\ \nu_{AI-C} \\ \tau_{AI-C_{\alpha}-C_{\beta}-H_{\beta}} \\ \delta_{U-C} \\ c \end{array}$	3388 3210 1664 1270 993 928 571 522 192	$\begin{array}{l} {}^{\nu}C_{\sigma}\text{-}H_{\beta} \\ {}^{\nu}C_{\alpha}\text{-}H_{\alpha} \\ {}^{\nu}C_{-C} \\ \delta_{A -C_{\sigma}\text{-}H_{\alpha}} \\ \tau_{H-C-C-H} \\ \delta_{H-C-C-H} \\ {}^{\nu}A_{I-C} \\ \tau_{A -C_{\alpha}-C_{\sigma}\text{-}H_{\beta}} \\ \delta_{D-C} \\ \epsilon_{-C} \end{array}$
$\frac{cis-AlHCCH}{AlCCH_2} \rightarrow \frac{cis-AlHCCH}{AlCCH_2}$ transition state			$\frac{cis-AIHCCH}{AICCH_2} \rightarrow \frac{AIC_2 - 2g}{AICCH_2}$ transition state		
$\begin{array}{c c} 3346 & \nu_{C_{g} \neg H_{\tilde{g}}} \\ 3058i & \nu_{C_{g} \neg H_{\alpha}} \\ 2383 & \nu_{C_{g} \neg H_{\alpha}} \\ 1805 & \nu_{C-C} \\ 927 & \delta_{C_{\alpha} \neg C_{g} \neg} \end{array}$		$ \begin{array}{c} - \nu_{C_{\alpha} - H_{\alpha}} \\ + \nu_{C_{\alpha} - H_{\alpha}} \end{array} $	513 276 258 237	$ \begin{array}{c} \nu_{A \mid -C} \\ \tau_{A \mid -C_{\alpha} - C_{\beta} - H_{\alpha}} \\ \delta_{A \mid -C_{\alpha} - C_{\beta} - H_{\beta}} \\ \tau_{A \mid -C_{\alpha} - C_{\beta} - H_{\beta}} \end{array} $	

^a The carbon center bonded to aluminum is designated C_{α} and the hydrogen bonded to this carbon is designated H_{α} . The remaining carbon and hydrogen are designated C_{β} and H_{β} respectively.

Method

All geometries were precisely optimized by using self-consistent-field (SCF) analytic gradients.⁴ The contracted double- ζ (DZ) basis set of Dunning and Hay,⁵ designated Al(11s 7p/6s 4p), C(9s 5p/4s 2p), and H(4s/2s), was employed for these optimizations. A series of calculations based on finite nuclear displacements from the optimized DZ-SCF geometries were carried out to obtain harmonic vibrational frequencies for all stationary points. With use of the DZ-SCF structures, single point calculations with a DZ + polarization (DZ+P) basis set and including configuration interaction (CI) were also carried out. The polarization

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