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Mechanism of the photochemical disproportionation reactions of (.eta.5-C5H5)2M2(CO)6 (M = Cr, Mo, W)

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Katz has suggested that the general lack of stereospecificity in Lewis acid cocatalyzed metathesis systems is due to a reversible Lewis acid induced cleavage of transient metallacyclobutanes to a 3-metallapropyl cation, which can then undergo C-C bond rotation, ultimately resulting in scrambling of stereochemistry. 16,17 The similarity of Katz's suggestion to our proposal is apparent. The tendency of a given metathesis catalyst system to show high stereospecificity in the presence of Lewis acids may depend on steric and electronic factors that can determine the susceptibility of a transient metallacyclobutane to transmetalation by the Lewis acid cocatalyst.

Acknowledgment. We gratefully acknowledge the support of the National Science Foundation (Grant CHE 8016528), the Department of Energy, and the use of the Southern California Regional NMR Facility (NSF Grant 7916324A1).

(16) Katz, T. J.; Hersh, W. H. Tetrahedron Lett. 1977, 585.

(17) Not all aluminum alkyl cocatalyzed metathesis systems show such a dependence of stereospecificity on the Lewis acid concentration. For example, the stereochemistry of ring-opening metathesis polymerization of cyclopentene with MoCl₃/AlEt₃ is unaffected by raising the AlEt₃ concentration: Dall'Asta, G.; Motroni, G. Angew. Makromol. Chem. 1971, 16/17, 51.
(18) All reactions described herein were followed by NMR spectroscopy

with either a Bruker WM-500 (500.13 MHz 1H, 76.76 MHz 2H) or a JEOL

FX-90Q (89.60 MHz ¹H) spectrometer.
(19) Reaction of *trans*·I-d₁ with 1 equiv of AlMe₃ resulted in less than 5% isomerization in 20 min at room temperature. Reaction of *trans*·I-d₁ with AlCl₃ resulted in decomposition of the metallacycle, but in the initial stages of reaction, the metallacycle was observed to be isomerized. Reaction of trans-I-d₁ with diphenylacetylene resulted in no isomerization of the metallacycle, and trans-3,3-dimethyl-1-butene-,- d_1 is observed as the only deuterated olefin. The half-life for thermal isomerization of trans-I- d_1 is 2 h at 50 °C: Lee, J. B.; Ott, K. C.; Grubbs, R. H. to be submitted for publication.

Mechanism of the Photochemical Disproportionation Reactions of $(\eta^5 - C_5 H_5)_2 M_2(CO)_6$ (M = Cr, Mo, W)

Albert E. Stiegman and David R. Tyler*

Department of Chemistry, Columbia University New York, New York 10027 Received October 30, 1981

Very little is known about the mechanism of the photochemically induced disproportionation reactions of metal-metal bonded dimers.1 For this reason we have been investigating the photochemical disproportionation reactions of the $Cp_2M_2(CO)_6$ (M = Cr, Mo, W; Cp = η^5 -C₅H₅) complexes, an example of which is shown in eq 1. The photochemical reactions of metal-metal

$$Cp_2Mo_2(CO)_6 \xrightarrow[NEt_3]{h\nu} CpMo(CO)_3^- + CpMo(CO)_2(NEt_3)_2^+$$
(1)

bonded dimers are generally thought to proceed via homolytic cleavage of the metal-metal bond,2 but the disproportionation reactions might be an exception to this generalization; it has been suggested that these reactions proceed via photoinduced heterolytic cleavage of the metal-metal bond. 1a We communicate here the results of experiments that strongly suggest that the disproportionation reactions of the Cp₂M₂(CO)₆ complexes result from neither homolytic nor heterolytic cleavage of the metal-metal bond but via a third pathway instead.

Scheme I

$$\begin{split} & \operatorname{Cp_2M_2(CO)_5L} \xrightarrow{h\nu} \operatorname{CpM(CO)_3} + \operatorname{CpM(CO)_2L} \\ & \operatorname{CpM(CO)_3} + \operatorname{CpM(CO)_2L} \xrightarrow{\operatorname{electron\ transfer}} \operatorname{CpM(CO)_3^-} + \\ & \operatorname{CpM(CO)_2L^+} \\ & \operatorname{CpM(CO)_2L^+} + \operatorname{L} \to \operatorname{CpM(CO)_2L_2^+} \end{split}$$

Scheme II

$$\begin{array}{c} \operatorname{Cp_2M_2(CO)_5L} \xrightarrow{h\nu} \operatorname{CpM(CO)_3} + \operatorname{CpM(CO)_2L} \\ \operatorname{CpM(CO)_2L} + \operatorname{Cp_2M_2(CO)_5L} \to \operatorname{CpM(CO)_2L^+} + \operatorname{Cp_2M_2(CO)_5L^-} \\ \operatorname{Cp_2M_2(CO)_5L^-} \to \operatorname{CpM(CO)_2L} + \operatorname{CpM(CO)_3^-} \\ \operatorname{CpM(CO)_2L^+} + \operatorname{L} \to \operatorname{CpM(CO)_2L_2^+} \end{array}$$

Irradiation ($\lambda > 290 \text{ nm}$) of $Cp_2Mo_2(CO)_6$ (1.0 × 10⁻² M) in degassed cyclohexane solution containing PPh₃ (5.0 \times 10⁻² M) proceeds as in eq 2.3 The substitution product, Cp₂Mo₂(CO)₅-

$$Cp_{2}Mo_{2}(CO)_{6} \xrightarrow{h\nu} Cp_{2}Mo_{2}(CO)_{5}(PPh_{3}) \xrightarrow{h\nu} CpMo(CO)_{2}(PPh_{3})_{2}^{+} + CpMo(CO)_{3}^{-} (2)$$

(PPh₃), was identified as an intermediate by its infrared absorption bands at 1968, 1900, and 1827 cm⁻¹. The products CpMo-(CO)₂(PPh₃)₂⁺ and CpMo(CO)₃⁻ were identified by their infrared absorption bands at 1974 and 1895 cm⁻¹ and at 1771 cm⁻¹, respectively.4 Note the small mole ratio of PPh3 to metal complex (5:1) required to give the disproportionation products. This small ratio shows that the disproportionation is not attributable to a solvent-induced dipolar excited state as originally proposed.⁵

Disproportionation reactions similar to the one in eq 2 occur for the $Cp_2M_2(CO)_6$ (M = Cr, Mo, W) dimers in hexane solution with ligands other than PPh₃. A list of ligands that lead to disproportionation includes NEt3, CH3CN, pyridine, aniline, AsPh₃, P(O-i-C₃H₇)₃, and P(OCH₃)₃. In each case, irradiation initially produces the monosubstituted dimer, Cp₂M₂(CO)₅L (identified by IR spectroscopy), and then this complex photolyzes to give the ionic products $CpM(CO)_2L_2^+$ and $CpM(CO)_3^-$. In no case were ionic products observed to form without the initial formation of the monosubstituted dimer. In order to check the idea that the monosubstituted dimer is an intermediate in the formation of the ionic products, the complex Cp₂Mo₂(CO)₅(PPh₃) was synthesized⁶ and its photochemistry studied. Irradiation (290 nm) of this complex $(1.0 \times 10^{-2} \text{ M})$ in hexane solution containing 0.1 M PPh₃ gave CpMo(CO)₂(PPh₃)₂⁺ and CpMo(CO)₃⁻. The products were identified by infrared spectroscopy.4 No infrared bands other than the ones attributable to these products were

The disproportionation reactions show a marked wavelength dependence. Irradiation of (MeCp)₂Mo₂(CO)₆⁷ in cyclohexane containing excess PPh₃ at 505, 435, 405, and 366 nm⁸ does not give the ionic disproportionation products; only the monosubstituted product (MeCp)₂Mo₂(CO)₅(PPh₃) is formed.⁹ quantum yield for substitution of (MeCp)Mo₂(CO)₆ by PPh₃ ([PPh₃] = 0.018 M, cyclohexane solution) at 405 nm is 0.04 \pm 0.01. Disproportionation does not occur until the excitation wavelength is 290 nm. At 290 nm the quantum yield for disproportionation of $(MeCp)_2Mo_2(CO)_5(PPh_3)$ is 0.40 ± 0.04 in

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⁽³⁾ All irradiations were carried out with a 200-W high-pressure mercury arc. Solutions were degassed with a stream of argon or nitrogen. The 290-nm wavelength band was isolated with a Corion interference filter

⁽⁴⁾ Haines, R. J.; Nyholm, R. S.; Stiddard, M. H. B. J. Chem. Soc. A

⁽⁵⁾ The disproportionation reactions were initially observed to occur in neat solvents such as pyridine, Me_2SO , and DMF. It was suggested that the polar solvent molecules induced a dipole in the excited state, ${}^{5}M-M^{\delta^{+}}$, which sub-

sequently relaxed to give ionic products.

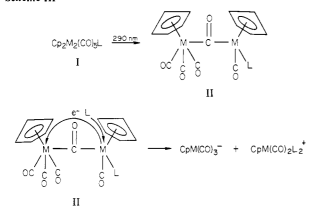
(6) Barnett, K. W.; Treichel, P. M. *Inorg. Chem.* 1967, 6, 294-299.

(7) The methylcyclopentadienyl compounds were used in some of these experiments as they were more soluble in alkane solvents than their cyclopentadienyl counterparts.

⁽⁸⁾ These wavelengths were isolated with interference filters from Edmund Scientific Co.

⁽⁹⁾ The CO stretching region of the infrared spectrum of (MeCp)₂Mo₂-(CO)₅(PPh₃) is identical with that of Cp₂Mo₂(CO)₅(PPh₃).

Scheme III



benzene; substitution of (MeCp)₂Mo₂(CO)₆ by PPh₃ ([PPh₃] = 0.018 M, cyclohexane solution) at 290 nm has a quantum yield of 0.35 ± 0.04 . The two lowest energy electronic absorption bands at approximately 500 and 380 nm in the Cp2M2(CO)5L complexes have been assigned to the $d\pi \rightarrow \sigma^*$ and $\sigma \rightarrow \sigma^*$ transitions, respectively.^{2g} Because electronic excitation at 505, 435, 405, and 366 nm does not lead to disproportionation, we must conclude that these excited states are inactive toward disproportionation. The dependence of the disproportionation reaction on wavelength is independent of the ligand. Wavelength results similar to those obtained with PPh, were also found for the other ligands used in our study.

Homolytic cleavage of the metal-metal bond occurs upon $\sigma \rightarrow \sigma^*$ or $d\pi \rightarrow \sigma^*$ excitation of the $Cp_2M_2(CO)_5L$ complexes.¹⁰ Therefore, the wavelength dependence of the disproportionation reaction has an important mechanistic implication: homolytic cleavage of the metal-metal bond is not sufficient to induce disproportionation. Consequently, the outer-sphere electrontransfer pathway in Scheme I and the radical-chain pathway¹¹ of Scheme II are not responsible for disproportionation of the Cp₂M₂(CO)₆ complexes. In addition, the previously proposed substitution-induced outer-sphere electron-transfer mechanism can also be eliminated from consideration.¹²

The results above suggest that disproportionation results from excitation to an excited state that is higher in energy than the $d\pi \rightarrow \sigma^*$ or $\sigma \rightarrow \sigma^*$ states. A possible pathway is outlined in Scheme III. In this scheme, the effect of 290-nm excitation is to produce intermediate II, a species with no metal-metal bond but a CO bridge. One of the metal atoms in II is coordinatively unsaturated and it undergoes nucleophilic attack by ligand L. This addition of another ligand to the metal puts sufficient electron density¹³ on the metal so as to induce an inner-sphere electron transfer. Note that reaction intermediates similar to II have been proposed before in the reactions of binuclear metal carbonyl complexes. 14-16

The quantum yield data support our suggestion that a coordinatively unsaturated intermediate such as II forms upon 290-nm excitation of the Cp₂M₂(CO)₅L complexes. Note that the quantum yields for substitution of Cp2Mo2(CO)6 and disproportionation of Cp₂Mo₂(CO)₅L at 290 nm are identical within experimental error (0.35 \pm 0.04 and 0.40 \pm 0.04, respectively). This constant value suggests that structurally related intermediates form with constant quantum efficiency when the Cp₂M₂(CO)₆ and Cp₂M₂(CO)₅L complexes are irradiated at 290 nm; we suggest that the M-CO-M bridged intermediate is common to both the substitution and disproportionation reactions at 290 nm. When Cp₂M₂(CO)₆ is irradiated, attack of L on the intermediate simply leads to substitution. When Cp₂M₂(CO)₅L is irradiated, the bridged intermediate forms with the same quantum efficiency as when Cp₂M₂(CO)₆ is irradiated. This time, however, coordination of L (two L's are now coordinated to the same metal) polarizes the M-CO-M unit enough so as to induce electron transfer.¹⁷ Attempts to stabilize II by irradiating Cp₂M₂(CO)₅L in lowtemperature glasses are in our laboratory.

Acknowledgment. We thank Professors H. B. Gray, T. L. Brown, and M. S. Wrighton for helpful discussions. S. Brawner McCullen is thanked for sending us a preprint of ref 11. Acknowledgment is made to the donors of the Petroleum Research Fund, administered by the American Chemical Society, and to the Research Corp. for the support of this research.

Registry No. $Cp_2Mo_2(CO)_6$, 12091-64-4; $Cp_2Cr_2(CO)_6$, 12194-12-6; $Cp_2W_2(CO)_6$, 12566-66-4; $Cp_2Mo_2(CO)_5(PPh_3)$, 12119-01-6; (MeCp)₂Mo₂(CO)₆, 33056-03-0; PPh₃, 603-35-0; NEt₃, 121-44-8; CH_3CN , 75-05-8; AsPh₃, 603-32-7; $P(O-i-C_3H_7)_3$, 116-17-6; $P(OCH_3)_3$, 121-45-9; pyridine, 110-86-1; aniline, 62-53-3.

(17) Although the quantum yield data are consistent with the formation of intermediate II in Scheme III, our results cannot rule out direct heterolysis of the M-M bond at 290 nm. The $\sigma\sigma^*$ singlet excited state of a metal-metal bonded complex is a bound ionic state. Irradiation at 290 nm may excite the molecule to a vibrational energy level of the $\sigma\sigma^*$ singlet state that is above the dissociation limit and ions may result.

Stereoselective Total Synthesis of $1\alpha,25$ -Dihydroxycholecalciferol

Enrico G. Baggiolini,* Jerome A. Iacobelli, Bernard M. Hennessy, and Milan R. Uskoković

Chemical Research Department, Hoffmann-La Roche Inc. Nutley, New Jersey 07110

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The isolation and structure determination of the physiologically active vitamin D_3 metabolite $1\alpha,25$ -dihydroxycholecalciferol (1)¹ and its use as a lifesaving drug for osteodystrophy due to renal failure have stimulated significant efforts toward synthesis of this natural product.² We report here the first³ total and chiral synthesis of 1α ,25-dihydroxycholecalciferol, which can also be used efficiently in the preparation of other 1α -hydroxy vitamin D

Lythgoe and co-workers have shown^{4,5} that the lithium phos-

⁽¹⁰⁾ Our results indicate that homolytic cleavage of the metal-metal bond occurs with low-energy excitation of Cp2Mo2(CO)5(PPh3). Irradiation (405 nm) of this complex in CCl4 solution yields CpMo(CO)3Cl and CpMo-(CO)₂(PPh₃)Cl. These products were identified by infrared spectroscopy. (See: Burkett, A. R.; Meyer, T. J.; Whitten, D. G. J. Organomet. Chem. 1974, 67, 67-73.) In addition, irradiation (405 nm) of Cp₂Mo₂(CO)₅(PPh₃) in benzene solution gives Cp2Mo2(CO)6. No CO stretching bands attributable to other products were observed in the infrared spectrum. The products of the Cl atom abstraction reaction and the cross-coupling reaction are consistent

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