brane feature would be a specified arrangement of different substrates in different binding sites depending on their properties. We can also envisage a wider scope of other applications, in both mechanistic and practical senses, to barrier-dependent phenomena such as encapsulation and transport of various substrates.

Quadruply Bonded

Tetramethyltetrakis(trialkylphosphine)dimolybdenum Compounds: Phosphine Exchange Kinetics, Acetone Formation with Carbon Monoxide, and Crystal Structure of Mo₂Me₄(PMe₃)₄

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Creation of coordinative unsaturation about a metal center is crucial for the generation of a catalytic cycle. One of the most general methods for achieving such unsaturation is reversible dissociation of neutral ligands, such as phosphines. A reasonable amount is known concerning the energetics of phosphine exchange in mononuclear transition-metal systems,² but relatively little is known about polynuclear ones.3 The quadruply bonded Mo(II) dimers are interesting candidates for such a study, since generation of vacant coordination sites about two metal atoms ca. 2 Å apart may allow them to cooperate in promoting reactions which are not otherwise possible. Unfortunately, little is known about the intimate reaction mechanisms which govern this class of molecules.⁴ As part of our efforts to explore the reactivity of quadruply bonded dimers,⁵⁻⁷ we have examined phosphine exchange kinetics and reductive cleavage by carbon monoxide in the alkyl compounds $Mo_2Me_4(PR_3)_4$

Blue Mo₂Me₄(PEt₃)₄⁸ was prepared from Mo₂(O₂CCMe₃)₄, MgMeCl, and PEt₃ in diethyl ether, followed by crystallization from diethyl ether at -10 °C. This complex undergoes ligand exchange with excess PMe₂Ph or PMe₃ in toluene solution within minutes at room temperature to give the known Mo₂Me₄-

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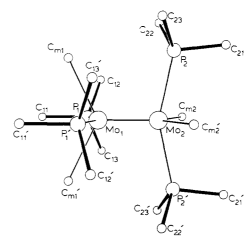


Figure 1. Perspective ORTEP drawing of the nonhydrogen atoms of Mo₂Me₄(PMe₃)₄. All atoms are represented by arbitrary-sized open circles for purposes of clarity. Atoms labeled with a prime are related to those labeled without a prime by the crystallographic twofild axis which passes through Mo1 and Mo2.

(PMe₂Ph)₄ or Mo₂Me₄(PMe₃)₄, respectively.⁹ The NMR parameters of all three complexes are similar, and they are doubtless isostructural. An X-ray crystal structure analysis of Mo₂Me₄-(PMe₃)₄ has been performed, ¹⁰ and the molecular geometry (Figure 1) is in accord with that previously deduced spectroscopically. The structure analysis reveals that the single crystals are composed of discrete binuclear molecules (Figure 1) which presumably contain a Mo-Mo quadruple bond. 10a These crystals are essentially isomorphous with those of W₂Cl₄(PMe₃)₄¹¹ and have a solid-state packing arrangement which is quite similar to Mo₂Cl₄(SEt₂)₄. ¹² Although the rigorous solid-state crystallographic symmetry of $Mo_2Me_4(PMe_3)_4$ is only C_2 , it approximates rather closely to full D_{2d} symmetry with the two MoL₂L'₂ units having a nearly eclipsed (to within 0.8°) conformation. Relevant bond lengths and angles include Mo₁-Mo₂, 2.153 (1)Å; Mo-P, 2.513 (2,0,0,2) Å; Mo-C, 2.439 (5,18,18,2) Å; Mo-Mo-P, 102.5 (5,9,9,2)°; Mo-Mo-C, 115.3 (1,4,4,2)°. 10b

It is of interest to determine the kinetics and mechanisms of phosphine exchange in these binuclear clusters. But before an examination of the kinetic parameters is possible, the stoichiometry of the reaction and stereochemistry of the products must be established. This is accomplished conveniently by monitoring the ³¹P{¹H} NMR spectra as a function of temperature during the progress of the reactions.

A toluene- d_8 solution of $Mo_2Me_4(PEt_3)_4$ (0.036 M) and PMe₂Ph (0.75 M) at -80 °C gives a spectrum that consists of two singlets due to these two species. Warming the sample to -50 °C gives free PEt₃, which eventually accounts for $25 \pm 3\%$

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⁽⁹⁾ Andersen, R. A.; Jones, R. A.; Wilkinson, G. J. Chem. Soc., Dalton Trans. 1978, 446-453.

^{(10) (}a) Single crystals of $Mo_2Me_4(PMe_3)_4$ are monoclinic, space group $C_{2/c}-C_{2h}^6$ (No. 15) with a=18.507 (5), b=9.462 (3), c=17.513 (4) Å; $\beta=116.18$ (2)°; Z=4 (dimeric units). Three-dimensional X-ray diffraction data were collected for 3172 independent reflections having 20 data were collected for 3172 independent reflections having $2\theta_{\rm MoK} \alpha < 55^{\circ}$ on a Nicolet P1 autodiffractometer by using graphite-monochromated Mo K α radiation and full (1° wide) ω scans. The solid-state structure was solved by using the "heavy-atom" technique, and the resulting structural parameters for all nonhydrogen atoms have been refined anisotropically to convergence [R-(unweighted, based on F) = 0.044 for 2372 independent reflections having $2\theta_{\text{MoK}\alpha} < 55^{\circ}$ and $I > 3\sigma(I)$ by empirically weighted full-matrix least-squares techniques on a Data General Eclipse S-200 computer using locally modified versions of the Nicolet E-XTL interactive crystallographic software system. (b) The first number in parenthesis following an averaged value of a bond length or angle is the root mean square estimated standard deviation of an individual datum. The second and third numbers, when given, are the average and maximum deviations from the averaged value, respectively. The fourth number represents the number of individual measurements which are included in the average value.

⁽¹¹⁾ Cotton, F. A.; Felthouse, T. R.; Lay, D. G. J. Am. Chem. Soc. 1980, 102, 1431-1433.

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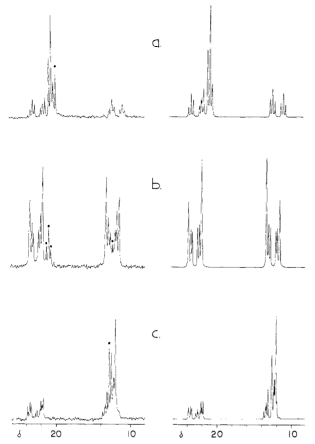


Figure 2. Observed (left-hand side) and calculated (right-hand side) spectra; A = I, B = IIb, and C = III. Solid circles indicate previous or subsequent substitution products.

of the total PEt₃ that is present initially.¹³ Simultaneously, the resonance due to Mo₂Me₄(PEt₃)₄ gradually disappears and is replaced by new multiplets due to coordinated PEt₃ and PMe₂Ph. These new resonances comprise an A₂BC spin system¹⁴ and correspond to the monosubstitution product Mo₂Me₄(PEt₃)₃-(PMe₂Ph) (I) (see Figure 2a). At -50 °C, no further changes

in the spectrum are evident, but warming to -30 °C causes liberation of a further molar equivalent of PEt₃. Concurrently, the A₂BC resonances diminish and new multiplets appear which correspond to the disubstitution product Mo₂Me₄(PEt₃)₂-(PMe₂Ph)₂. Two isomers of this mixed phosphine complex are possible, IIa and IIb, and these possibilities are distinguishable, since the former is an A₂B₂ spin system, while the latter is AA'BB'. The observed spectrum¹⁵ (Figure 2b) is consistent with isomer IIb only, and thus, the second phosphine substitution does *not* occur on the same molybdenum atom as did the first substitution. Replacement of a third equivalent of phosphine occurs at a temperature of ca. -10 °C, and the product formed, Mo₂Me₄-(PEt₃)(PMe₂Ph)₃ (III), is another example of an A₂BC spin system¹⁶ (figure 2c). Finally, at 0 °C, the last PEt₃ group is replaced by PMe₂Ph, giving Mo₂Me₄(PMe₂Ph)₄. The only other resonances remaining are those due to free PEt₃ and free PMe₂Ph.

It is apparent that the phosphine exchange process in these molecules is stepwise. Having established the stoichiometry of the individual steps and the stereochemistries of the respective products, it becomes possible to measure kinetic parameters, viz., the rate of substitution of the first PEt₃ group in Mo₂Me₄ (PEt₃)₄ by PMe₂Ph. At -50 °C, the rate of disappearance of Mo₂Me₄-(PEt₃)₄ is first order in [Mo₂Me₄(PEt₃)₄] and zero order in [PMe₂Ph], when the latter is in excess. Further, the rate is invariant on [PMe₂Ph], when the latter is varied over a 10- to 30-fold excess. Under these conditions, the first-order rate constant is $k_{\rm obs} = (5.5 \pm 0.1) \times 10^{-4} \, \rm s^{-1}$ over three half-lives. An Arrhenius plot of $\ln (k_{\rm obs}h/k_{\rm B}T)$ vs. 1/T yields the activation parameters, $\Delta H^{\dagger} = 20.6$ kcal mol⁻¹ and $\Delta S^{\dagger} = 19.7$ eu. Further, replacement of PEt₃ in Mo₂Me₄(PEt₃)₄ by PMe₃ proceeds via the same stepwise reaction path as did the PMe₂Ph experiment and follows identical kinetics within experimental error: $k_{\text{obs}} = (5.3 \pm 0.3) \times 10^{-4} \,\text{s}^{-1}$ at -50 °C.

The invariance of the reaction rates on the nature of the incoming group and the positive ΔS^* both are indicative of a dissociative mechanism. Surprisingly, associative attack by phosphine does not occur, even though these complexes possess vacant axial sites. However, previous studies suggest that axial ligation is weak at best and does not involve covalent interaction. The driving force for these dissociative reactions is most reasonably ascribed to relief of steric congestion, as PEt₃ is replaced by the smaller PMe₂Ph or PMe₃ ligands. Consistent with this, Mo₂Me₄-(PMe₂Ph)₄ does not react at all with excess PMe₃ until a temperature of 0 °C is reached. Further, neither Mo₂Me₄(PMe₃)₄ nor Mo₂Me₄(PMe₂Ph)₄ react with excess PEt₃ at 25 °C, as shown by $^{31}P^{\{1}H\}$ NMR spectroscopy.

The $Mo_2Me_4(PR_3)_4$ compounds $(PR_3 = PEt_3, PMe_2Ph, PMe_3)$ also react with carbon monoxide under 18 atm at room temperature in benzene. NMR spectroscopy shows the formation of acetone and Mo[O] compounds of the type $Mo(CO)_{6-x}(PR_3)_x$ where x = 0-3. Qualitatively, the rates of reaction at room temperature of CO with $Mo_2Me_4(PR_3)_4$ depend on the coordinated phosphine: $PEt_3 \gg PMe_2Ph > PMe_3$. This suggests that the rate-determining step involves phosphine dissociation by analogy with the results described above. On the basis of studies of the reaction of quadruply bonded dimers with isoelectronic *tert*-butyl isocyanide, it is possible that reductive elimination of acetone may not occur from the intact binuclear alkyl, but from a mononuclear one.⁷ Further studies are under way in an attempt to resolve this question. 19,20

⁽¹³⁾ The values are corrected for differing NOE's. Spectra were acquired at 72.9 MHz.

⁽¹⁴⁾ $\delta_A = 19.7$, $\delta_B = 21.4$, $\delta_C = 10.9$; $J_{AB} = 21.5$, $J_{AC} = 21.2$, $J_{BC} = 102.9$ Hz. Simulations were performed by using the program ITRCAL, written by Nicolet Technology Corp. Note that this is the first time P-P coupling across a M-M quadruple bond has been directly observed.

⁽¹⁵⁾ $\delta_A = 22.5$, $\delta_B = 12.3$; $J_{AA'} = 20.0$, $J_{AB} = 104.1$, $J_{AB'} = 23.0$, $J_{BB'} = 19.2$ Hz.

⁽¹⁶⁾ $\delta_{A} = 12.3$, $\delta_{B} = 12.7$, $\delta_{C} = 22.7$; $J_{AB} = 23.0$, $J_{AC} = 23.0$, $J_{BC} = 106.8$

⁽¹⁷⁾ Cotton, F. A.; Norman, J. G. J. Am. Chem. Soc. 1972, 94, 5697-5702. Garner, C. D.; Senior, R. G. J. Chem. Soc., Dalton Trans. 1975, 1171-1174.

⁽¹⁸⁾ The 18-atm reactions do not represent the minimum pressure for reaction but is a convenient pressure that gives satisfactory rates of reaction at room temperature. Some of the phosphine carbonyls may arise from scrambling after acetone has been eliminated.

⁽¹⁹⁾ For kinetic studies of acetone formation in other systems, see: Bergman, R. G. Acc. Chem. Res. 1980, 13, 113-120.

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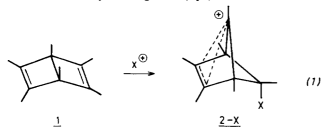
(20) trans-Mo(CO)₄(PMe₃)₂ and other products have been observed in the reaction of Mo₂Me₄(PMe₃)₄ with CO in refluxing toluene. Cotton, F. A.; Darensbourg, D. J.; Kolthammer, B. W. S. Southwest-Southeast American Chemical Society Regional Meeting, New Orleans, L.A., Dec 10-13, 1980; INOR 235.

Sulfenium Ion Addition to Hexamethyl(Dewar benzene). A Case for a Hypervalent Sulfonium Ion Intermediate

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The addition of electrophiles to hexamethyl(Dewar benzene) (1) induces its transformation to reactive intermediates that provide a rich array of rearrangement products.¹ In the addition of hetero atom derived electrophiles, i.e., halonium ions, the pivotal intermediate is the bicyclo[2.2.1.]hexenyl cation (2), the formation of which is initiated by addition to a double bond from the endo face of 1 followed by rearrangement (eq 1).^{1,2} We find now that



reaction of 1 with sulfenium ion (RS⁺) equivalents involves a significant deviation from this course of reaction.

Phenylsulfenyl chloride (3) and 1 were mixed in equimolar amounts at -120 °C in CH_2Cl_2 under an N_2 atmosphere. The solution became homogeneous at -70 °C and the 1H NMR spectrum taken immediately revealed in the methyl region three new singlets of equal intensity at δ 1.49, 1.57, and 1.67 (the absorptions for 1 are found at δ 1.55 and 1.05). Within 10 min the 1H NMR spectrum becomes complex, but after warming to room temperature a clean spectrum is obtained which arises from 1 (48% yield) and dichloride 5, formed in 31% yield. The latter is known to be the end product of the chlorination of 1.2b Diphenyl disulfide is also formed.

An abbreviated scheme consistent with these observations is given in eq 2. The intermediate immediately formed at low temperature is assigned structure 4-Cl (endo attack, exo orientation of phenyl substituent) consistent with the normal reaction course of sulfenyl halides.⁴ The complicated spectrum that soon appears is probably the consequence of subsequent formation of covalent adducts. However, the formation of 4, as well as any adducts, must be reversible. This is indicated by the failure to obtain sulfur

containing stable adducts as well as by the recovery of 48% of the starting material 1. The other half of the starting material (1) is ultimately irreversibly consumed by chlorination via a chloronium ion from 3.⁵ Reversible thiiranium ion formation has been previously demonstrated by us.⁶

The chloride counterion in 4 is clearly too nucleophilic. Phenylbis(thiophenyl)sulfonium hexachloroantimonate (6) has the less nucleophilic diphenyl disulfide as the leaving group.⁷ treatment of 1 with 6 in CH₂Cl₂ (this was the only suitable solvent found) a solid formed at low temperatures, but at -60 °C the solution became homogeneous. In the methyl region four new singlets were observed in the ¹H NMR spectrum (60 MHz) at δ 1.10 (3 H), 1.37 (6 H), 1.43 (6 H), and 1.51 (3 H). A small and variable amount of 1 remained; the phenyl absorptions were obscured by diphenyl disulfide. On warming the three absorptions at δ 1.37, 1.43, and 1.51 broaden and coalesce at -43 °C. At -30 °C the spectrum consists of two sharp singlets at δ 1.43 (15 H) and 1.10 (3 H). These spectra are shown in Figure 1. Owing to the limited solubility and stability we have been unable to obtain ¹³C NMR spectra. Above -20 °C the spectrum broadens irreversibly, but after standing for some hours the sharp absorption for hexamethylbenzene appears. A rationalization for these observations is given in eq 3.

At low temperatures 4-SbCl₆ is formed as a solid that at -60 °C rearranges to 7, which is a single geometrical isomer with the phenyl group oriented toward the five-membered ring.⁸ That the structure of the ion is 7 and not 2-SC₆H₅, which has the same symmetry, is shown by the high field positions of the methyl groups in the ¹H NMR spectrum.⁹ Moreover, this structural assignment is in agreement with the observation that quenching with $(C_2H_5)_3N$ gives in 45% yield tricyclic alkene 9^{10} as shown in eq 3. This is

630.
(7) Gybin, A. S.; Smit, W. A.; Bogdanov, V. S.; Krimer, M. Z.; Kakyarov,

(7) Gybin, A. S.; Smit, W. A.; Bogdanov, V. S.; Krimer, M. Z.; Kakyarov, J. B. Tetrahedron Lett. 1980, 383.

(8) No indication for a second isomer could be found from the 360-MHz ¹H NMR spectrum. Examination of models indicates that the isomer drawn has the phenyl group in the least hindered position. The formation of only one of two possible isomers of a thiiranium ion has been observed previously. See, for example: Raynolds, P.; Zonnebelt, S.; Bakker, S.; Kellogg, R. M. J. Am. Chem. Soc. 1974, 96, 3146. Capozzi, G.; De Lucchi, O.; Lucchini, V.; Modena, G. Tetrahedron Lett. 1975, 2603.

(9) The carbon bound methyl groups in the methyl sulfonium salt of cis-2,4-dimethylthietane absorb at δ 1.64 (Trost, B. M.; Schinski, W. L.; Chen, F.; Mantz, I. B. J. Am. Chem. Soc. 1971, 93, 676), which value should be compared with the chemical shifts for 2-Cl cited in ref 3.

(10) As has been shown (Hogeveen, H.; Zwart, L. Israel. J. Chem., in press) ¹³C NMR spectroscopy provides a powerful tool for structural assignments of these types of compounds. The characteristic chemical shifts are sp² carbon atoms 135–165 ppm; cyclopropane quaternary atoms 29–36 ppm; cyclobutane quaternary atoms 48–60 ppm; bicyclobutane quaternary atoms 19–32 (nonbridgehead) and 37–47 ppm (bridgehead atoms) and the methyl groups 1.5–4 ppm (nonbridgehead connected methyl groups) and 4–9 ppm (bridgehead connected methyl groups). Spectra for 9: ¹H NMR (CDCl₃) δ 0.91 (s, 3 H), 1.05 (s, 3 H), 1.18 (s, 3 H), 1.22 (s, 3 H), 1.34 (s, 3 H), 4.32 (s, 1 H), 4.35 (s, 1 H), 7.1–7.5 (m, 5 H); ¹³C NMR (CDCl₃) δ 4.2 (CH₃), 6.1 (CH₃), 6.4 (CH₃), 7.4 (CH₃), 20.4 (CH₃), 40.0 (C), 44.6 (C), 48.7 (C), 60.2 (C), 62.1 (C), 85.2 (sp² CH₂), 126.2 (aromatic CH, 2×) 127.6 (aromatic C), 128.0 (aromatic CH, 2×), 136.1 (aromatic CH), 158.1 (sp² C).

⁽¹⁾ Review: Hogeveen, H.; Kwant, P. W. Acc. Chem. Res. 1975, 8, 413 and references cited therein.

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⁽³⁾ For comparison purposes in 2-Cl the methyl absorptions are found at δ 1.64 (3 H), 1.71 (6 H), 1.97 (3 H), and 2.30 (6 H). See: Schaefer, W.; Hellmann, H. Angew. Chem. 1967, 79, 566

Hellmann, H. Angew. Chem. 1967, 79, 566.

(4) See, for example: Smit, W. A.; Zefirov, N. S.; Bodrikov, I. V.; Krimer, M. Z. Acc. Chem. Res. 1979, 12, 282. These authors also discuss the problem whether thiiranium salts have a covalent sulfurane structure or an ionic structure as drawn for 4-Cl.

⁽⁵⁾ With 2 equiv of 3, addition to both double bonds of 1 occurs to give a product: ¹H NMR δ 1.46 (s, 12 H), 1.60 (s, 6 H), and 7.25-7.50 (br abs, 10 H). This is apparently a dithiranium salt derived from addition of phenylsulfenium ion to both double bonds of 1. The C₂₀ symmetry apparent from the ¹H NMR spectrum is consistent with endo-endo or exo-exo addition.

(6) Bolster, J. M.; Kellogg, R. M. J. Chem. Soc., Chem. Commun. 1978,