a-Alkyl and -Aryl Complexes of the Group 4-7 Transition Metals

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I. lnfroducfion

Transition metal complexes containing alkyl or aryl ligands have proliferated in the past decade. One of the most striking developments is the synthesis of "binary" complexes containing only σ -alkyl or -aryl ligands which violate previously accepted stability rules, such as the principle that stable metal alkyls must be diamagnetic or that they must be coordinatively saturated (the 18 electron rule).'

We intend to summarize recent advances in transition metal alkyl and aryl chemistry and will emphasize the early transition metals (groups 4-7) since the most exciting fundamental developments involve these elements. Generally group 8 metal alkyl chemistry is still well represented by earlier comprehensive reviews.^{2,3} More recent discussions of the metal-carbon bond are also available $4-7$ (see also section VI), while other sources cover earlier or more specialized material.⁸⁻¹⁶

II. Binary Alkyl Complexes+

Transition metal compounds bearing only σ -alkyl ligands have previously been so scarce and so little studied that the impression arose that binary metal alkyls are inherently unstable. The key to dispelling this fallacy came with the understanding that transition metal alkyls can decompose via β hydrogen ''elimination" (or "abstraction"), which is not common among main group alkyls under similar mild experimental conditions. Subsequently people recognized that alkyls which have no β -hydrogen should form stable complexes. (Stability normally refers to thermal stability, a property which has not yet been measured in a standard way.) Several research groups reported examples almost simultaneously. Particularly striking are WMe₆ and Ti(CH₂SiMe₃)₄, which grossly violate the 18 electron stability rule, and paramagnetic $V(CH_2Ph)_4$.

Others prepared stable compounds containing tert-butyl or norbornyl groups. In the former the β hydrogen appears to be sterically inaccessible to the metal atom. In the latter β -elimination is unfavorable because unsaturation at a bridgehead carbon would result (Bredt's rule).

Though suppression of intramolecular decomposition (primarily β -elimination) has received the greatest emphasis, the fact that binary complexes with large alkyl ligands lacking a β hydrogen are prepared more easily and are more stable than the corresponding methyl complexes almost certainly can be attributed also in part to the concomitant suppression of intermolecular decomposition reactions. The interesting consequence in some instances is that intramolecular α -hydrogen abstraction becomes a favored decomposition mode.

For brevity, "alkyl" will usually be a general term including aryl and alkaryl, and ligands containing Si in the β position such as $-CH_2$ SiMe₃.

Quite generally, the new binary metal alkyls react with air and moisture and are only modestly stable thermally. The usual synthesis involves reaction of a Grignard or organolithium reagent with a transition metal halide as in

4PhCH₂MgCI + $ZrCl_4 \xrightarrow{E t_2O} Zr(CH_2Ph)_4$

$$
4PhCH2MgCl + ZrCl4 \xrightarrow{Et2O} Zr(CH2Ph)4
$$

Exact experimental details are often critical. We classify binary alkyls by ligand type and discuss their preparation and chemistry below.

A. Neutral

1. Methyl

In general, binary methyl complexes are extraordinarily stable *relative to* those containing alkyls bearing a β hydrogen. Complexes are known for Ti, Zr, Ta, Nb, Cr, W, and Mn.

Clauss and Beermann¹⁷ first prepared bright yellow TiMe₄ in 1959 from methyllithium or a methyl Grignard reagent and TiCI4 in diethyl ether at -78° . Inverse addition (TiCI4 to the alkylating agent) was later found to give a smoother reaction.¹⁸ Tetramethyltitanium and ether codistill from the reaction mixture at ca. -30° . In ether it decomposes near room temperature to give methane, traces of ethylene and ethane, and a black residue containing carbon, hydrogen, and Ti.^{19,20} Crystalline TiMe4, obtained by removing ether in vacuo from a hexane-ether solution at -78° (ref 18), begins to decompose a few degrees above -78° .

Tetramethylzirconium is prepared in ether-toluene at -45° from methyllithium and ZrCl₄. Codistillation gives a red-brown ether solution of ZrMe₄ which evolves 4.0 mol methane per zirconium on hydrolysis.^{21,} No further characterization or reactions have been reported.

Methyllithium (2 mol) reacts with TaMe₃Cl₂ at -78 to 0° in ether to give TaMe₅ which can be isolated as volatile, pale yellow needles which melt at ca. 10° to a yellow oil.²² It decomposes smoothly at 25° in several hours to give methane (ca. 3.2 mol) as the only detectable gaseous product. Its ¹H NMR spectrum in toluene- d_8 at -10° shows a single sharp singlet at τ 9.18 and its mass spectrum a peak corresponding to TaMe₄⁺. Like TiMe₄, its solutions in ether appear somewhat more stable than the pure complex; it may be stabilized by coordination of ether. The analogous reaction of $NbMe₂Cl₃$ with 3 mol of LiMe gives a yellow-orange ether solution of a product which decomposes smoothly between -20 and -10° . Though a stable, seven-coordinate adduct, NbMe₅L₂ (L₂ = e.g., a chelating phosphine), was isolated after adding L_2 to the ether solution, firm conclusions concerning the existence of NbMe₅ must await further studies.

Hexamethyltungsten can be prepared in ca. 40% yield from WCI₆ and 3 mol of LiMe in ether at -20° (ref 23, 24). The yield is reportedly extremely sensitive to this reactant ratio, and evidence suggests WMe₆ does not form directly. Traces of oxygen may also be required.25 It has also been prepared from WCI_6 and AIMe₃ in aliphatic hydrocarbons.²⁵ WMe₆ can be extracted into pentane or sublimed onto a -10° probe and melts and decomposes near room temperature to give ca. 3 mol of methane containing **<5%** ethane. In toluene- d_8 it gives a single ¹H NMR resonance at τ 8.38. The $WMe₅$ ⁺ peak was the highest found in its mass spectrum. It can be characterized further by reaction with O₂, R₂NH, and halogens (X₂) to give W(OMe)₆, W(NR₂)₆, and WX₆, respectively, and with NO to give $WMe₄{ON}(Me)N=O₂$ (see section VI1.G). A photoelectron spectrum is consistent with octahedral W(VI) bound to six carbon ligands.²⁶

Isolation of WMe $_6$ and TaMe₅ suggests that other third-row binary methyl complexes, $HfMe₄$ and $ReMe₇$, should be isolable though a direct method of preparing the latter is not obvious. ReMe₆ has recently been mentioned briefly.²⁵

Only binary methyl complexes with the metal in its highest common formal oxidation state $(Ti^{\mid V}, Ta^{\mid V}, W^{\mid V})$ are well characterized. A few lower valent complexes have been reported, but only dimethylmanganese has been isolated. Reaction of TiCI₃(THF)₃ with LiMe in ether¹⁷ or MeMgCl in THF or ether²⁷ gives a dark-green solution believed to contain trimethyltitanium. Brown-red CrMe₃ and black CrMe₂ have been postulated in analogous reaction schemes,¹⁷ while CrMe₄ is probably formed from $Cr(OCMe₃)₄$ and 4LiMe in pentane at -78 ° (ref 28, 29). All are exceedingly unstable and no adducts have been isolated. It was noted²⁹ that codistillation of CrMe4 and petroleum in vacuo gave a considerably less stable solution (it decomposed slowly at -60°) than when lithium reagent was present. Dimethylmanganese (from MnI₂ and 2 LiMe in ether $30,31$), unlike the complexes discussed so far, is insoluble in ether and assumed to be polymeric. It has also been prepared in THF,³² coordination of which apparently prevents polymerization.

Binary methyl complexes have low formal valence electron counts (e.g., 8 for TiMe₄) and consequently readily form adducts with donor ligands when sterically possible. Adducts of tetramethyltitanium have been studied to the greatest extent.³³⁻³⁵ Both five-coordinate (TiMe₄L) and six-coordinate complexes (TiMe4L2) have been isolated. Those which can be characterized are monomeric in aromatic solvents. In general, TiMe4L complexes are less stable. A qualitative thermal stability series is $L = \text{dios} < N \text{Me}_3 < \text{PMe}_3 < \text{py}$. For TiMe₄L₂ it is L₂ = TMEDA \leq 2PMe₃ \leq phen \leq bpy \leq dmpe. For example, TiMe₄(diox) explodes at 0° while TiMe₄(dmpe) sublimes in vacuo at 30-40° without decomposition. Greatest stability occurs when two coordination sites are strongly occupied, the implication being that vacant metal coordination sites allow at least initiation of decomposition. This phenomenon has been observed somewhat more directly for remarkably stable TaMe₅(dmpe) and NbMe₅(dmpe).²² The ¹H NMR spectrum of TaMe_s(dmpe) at 105° exhibits a triplet resonance for five methyl groups coupled to two phosphorus nuclei. At slightly higher temperatures the triplet becomes a singlet, coordinated dmpe exchanges with added dmpe, and the complex rapidly decomposes. Weak 1:l adducts of WMe₆ with PPh₂Me, PPhMe₂, and PPh₂Et are isolable at low temperature, but the ligands dissociate readily in solution at room temperature.²⁴ Finally, apparently even weak solvent adducts are more stable than the parent compound judging by enhanced stability in solvents such as diethyl ether.

2. Phenyl

Contrary to popular belief, phenyl complexes are not inherently more stable than alkyl complexes, unless the alkyl contains a β hydrogen. Neutral binary phenyl complexes are scarce and difficult to prepare. Metal reduction and biphenyl formation are common. A low energy decomposition route involving ortho hydrogen abstraction, strictly analogous to *p*hydrogen abstraction, may be responsible for these difficulties (see section V1.A).

The MPh₄ complexes where M = Ti,³⁶ Zr,³⁷ or V^{38} have been prepared at low temperature in ether and characterized by reaction with HgCl₂ to give PhHgCl. All decompose readily
at 0°:
MPh₄ → MPh₂ + biphenyl (M = Ti, V) at **0':**

$$
MPh_4 \rightarrow MPh_2 + biphenyI (M = Ti, V)
$$

$$
ZrPh_4 \rightarrow [ZrPh_2(\text{ether})]_2 + 0.5 \text{bipheny} + C_6H_6
$$

No details of the latter, which apparently requires a source of H, were proposed. Black, crystalline $[ZrPh_2(\text{ether})]_2$ can be isolated in 95% yield and has been characterized well. It is very soluble in ether, THF, dioxane, and aromatic hydrocarbons and poorly soluble in saturated hydrocarbons. It decomposes at high temperatures according to the stoichiometry

$$
[ZrPh2(ether)]2 \xrightarrow{50-180^{\circ}}
$$
 2Zr(Ph)(OEt) +

$$
C_6H_6 + 0.5C_6H_5 - C_6H_5 + 0.5C_2H_6 + 1.5C_2H_4
$$
.

The exact nature of black, pyrophoric TiPh₂ and VPh₂ has not been fully elucidated.

Reduction can be minimized by appropriate choice of reactants, order of addition, rate of addition, and temperature. For example, good yields of $TiPh_4$ are obtained by reaction of TiCl₄L₂ (L = py, pip, or quin) and PhMgBr in ether at -16° (ref 39). The authors attribute this favorable result to precipitation of $MgBr_2L_x$ during the reaction. They also suggest that the crystalline "TiPh₄" which was isolated at -80° in ether³⁶ was actually sparingly soluble TiPh₄. $nMgX_2$ (X = halide). Further studies showed that substitution of $Ti(OCMe₃)₄$ for $TiCl₄$ and use of inverse addition gives $Ti(OCMe₃)₄ \cdot nPhMgBr$ which disproportionates on addition of dioxane to give TiPh₄, $Mg(OCMe₃)₂$, and sparingly soluble $MgBr₂(diox)₂$. The best system appears to be that employing $Ti(OCMe₃)₄$ and MgPh₂ which gives a smooth reaction to form TiPh₄ with essentially no reduction of Ti'" (ref 40). These authors conclude that, at least in the case of titanium, magnesium halides and other Lewis acids contribute to instabilities of reaction mixtures containing TiR4. Accordingly, in contrast to previous reports, $36,41$ they found pure TiPh₄ does not decompose in refluxing ether.⁴² In the crystalline state, however, $TIPh₄$ turns black in a few minutes at room temperature. It is monomeric in benzene.

A few adducts of TiPh4 were prepared for spectroscopic purposes.³⁵ Apparently only orange TiPh₄(bpy), stable at O'C, was isolated, though they observed adducts with TMPDA, py, TMEDA, and TMMDA in solution.

The major class of binary phenyl complexes are those of chromium. The early work of Hein was shown by Zeiss and Tsutsui in 1954 to involve η^6 -arene complexes of Cr(0) and $Cr(1).43$ At about the same time, the first σ -organochromium compound, blood-red CrPh₃(THF)₃, was prepared from $CrCl₃(THF)₃$ and PhMgBr in THF at -20° (ref 44, 45). It decomposed on heating to a black solid which gave $Cr(\eta^6 C_6H_6$)₂ and $Cr(\eta^6-C_6H_6)(\eta^6-biphenyl)$ on hydrolysis. More recent studies^{46,47} have not altered the overall interpretation though details are still not fully understood. This aspect of chromium chemistry as well as the preparation and properties of σ -bonded organochromium compounds in general have been discussed in detail.¹³ Only adducts, CrPh3L3 and, in one case, CrPh₃L₂, have been isolated. Decomposition to Cr(0) appears determined to a large extent by the lability of L.

Addition of L-L to $CrPh₃(THF)₃$ gives sparingly soluble CrPh₃(L-L)(THF) (L-L = bpy or phen).⁴⁸ These adducts are considerably more stable than $CrPh₃(THF)₃$ and decompose at 135° (bpy) and 160° (phen).

When CrPh₃(THF)₃ is dissolved in neat PPhR₂ (R = Et, Bu) and THF removed in vacuo, five-coordinate, red-violet $CrPh₃(PPhR₂)₂$ complexes can be isolated.⁴⁹ They may be recrystallized from hexane and are monomeric in benzene. Only traces of aromatic complexes are found after refluxing for several hours in ether, benzene, or hexane even though they dissolve in THF to give $CrPh₃(THF)₃$ instantly.

A Cr(II) species, $CrPh₂(bpy)₂$, can be prepared by a direct or indirect method⁵⁰ (eq 1). It does not decompose at 80[°].

CrCl₂
$$
\frac{1.2 \text{Liph.}-78^{\circ} \text{(THF)}}{2.2 \text{bpy,}-20^{\circ} \text{(THF)}} \text{CrPh}_2(\text{bpy})_2
$$

\n($\mu = 3.22$)
\nCrPh₂(CHPh₂)(THF)_x $\frac{2 \text{bpy}}{-1/2 \text{Ph}_2 \text{CHCHPh}_2}$ (1)

 $CrPh₂(CHPh₂)(THF)₂$ was prepared in THF from $Cr(CHPh₂)Cl₂(THF)₂$ and 2LiPh.

The mesityl derivatives of Cr(lll) and Cr(ll), in contrast, apparently do not give Cr(0) on decomposition. Violet Cr(Mes)₃THF (μ = 3.74) was prepared from CrCl₃(THF)₃ and MesMgX in THF.51 THF is lost in noncoordinating solvents and refluxing causes decomposition to mesitylene and Cr(lll); subsequent hydrolysis did not give Cr(0) species. The authors suggest that absence of an ortho hydrogen and/or steric hindrance stabilizes Cr(Mes)₃ toward reduction. Intermolecular decomposition was not ruled out. Violet $Cr(Mes)_{2}(THF)_{3}$, black-violet $Cr(Mes)_{2}(bpy)(THF)$, and blue $Cr(Mes)Cl(THF)_{2}$ were reported a year later.⁵² Decomposition in ether or THF also gave no Cr(0) species. Similar compounds containing the 1-naphthyl and 2-methyl- 1-naphthyl ligands were reported but not well characterized.

Violet-blue V(Mes)₃ \cdot 1.25THF (μ = 2.74) was prepared⁵³ as for the Cr analog. It is stable to 120° in vacuo and loses THF in dioxane or benzene. The solvent-free complex could not be isolated.

The existence of green diphenylmanganese³⁰ has been disputed. 31 If MnPh₂ was indeed prepared, its insolubility in ether would suggest a polymeric structure analogous to $MnMe₂$. However, as with $MnMe₂$, soluble $MnPh₂$ can be prepared in THF.54

Finally, several rhenium complexes containing σ -aryl groups are known.⁵⁵ Blue RePh₃(PPhEt₂)₂, Re(p -tolyl)₃(P-PhEt₂)₂, and $[RePh₂(PPh₃)₂]$ _n are prepared from $ReCl₃L₃$ or Re(O)Cl₃L₂, and orange Re(N)Ar₂(PPh₃)₂ (Ar = Ph, p-tolyl, σ tolyl) from Re(N)Cl₂(PPh₃)₂. The former are stable in refluxing benzene; the latter decompose.

3. *Benzyl*

Preparation of binary benzyl complexes marked the beginning of a rapid development in the area of "stable" binary alkyl complexes. Alkyl ligands such as neopentyl or trimethylsilylmethyl, stable complexes of which were soon discovered, resemble the benzyl ligand in that they too have no β hydrogen and are rather bulky. In addition to their scientific import the $M(CH_2Ph)_4$ compounds, especially the zirconium derivative, have practical potential as olefin polymerization catalysts.56

Tetrabenzyltitanium was prepared in solution in 196740,57 and isolated a year later.⁵⁸ It is most commonly prepared from TiCl₄ and PhCH₂MgX ($X = Cl$, Br) in diethyl ether but variations have been studied carefully.⁵⁹⁻⁶² An interesting indirect method employing TiMe₄ and B(CH₂Ph)₃ has also been reported.⁶³ In general, however, $Ti(CH_2Ph)_4$ is much easier to prepare than TiPh₄. One important finding⁶⁰ is that some ether is necessary apparently in order to form an adduct with TiCI4.

The preparations of $Zr(CH_2Ph)_4^{64,65}$ and $Hf(CH_2Ph)_4^{66-69}$ are entirely analogous to that of $Ti(CH_2Ph)_4$, while green $V(CH_2Ph)_4$ is best prepared from VCl_4 and $Mg(CH_2Ph)_2$ in pentane in the presence of a small amount of ether,⁷⁰ or in neat ether.⁷¹ Apparently only black, crystalline $V(CH_2Ph)_4$ (ether) has been isolated.⁷¹ Use of PhCH₂MgCI gave large quantities of VCI₃ and little V(CH₂Ph)₄. Tetrabenzyltungsten^{72a} and tetrabenzylmolybdenum^{72b} have also been mentioned, but details are not yet available.

The synthesis, properties, and some reactions of $Ti(CH_2Ph)_4$ and $Zr(CH_2Ph)_4$ have been described fully^{65,73,74} (Figure 1). Red Ti(CH₂Ph)₄ (mp 70°) and yellow Zr (CH₂Ph)₄ (mp 133') are slightly soluble in aliphatic hydrocarbons, more so in aromatic hydrocarbons and ethers, and are monomeric in benzene.

Both Ti(CH₂Ph)₄ and Zr(CH₂Ph)₄ decompose in refluxing heptane in several hours (Ti \gg Zr). The titanium compound

Figure 1. Some reactions of $M(CH_2Ph)_4$ (M = Ti, Zr).

yields the maximum theoretical amount of toluene which can arise from methylenic hydrogen atom transfer (2.66 mol). No deuterated toluene is produced in deuterated solvents. In the solid state both decompose to give benzene, bibenzyl, diphenylmethane, and ethylene in addition to toluene.73 The black Zr-containing residue on deuterolysis gave a mixture of 50% H2, 25% HD, and **25%** D2 (suggesting catalyzed H/D exchange), d_0 - to d_4 -toluene, and deuterated mixtures of methane and ethane, while reaction with $HgCl₂$ gave PhCH₂HgCI, MeHgCI, and EtHgCI, all in trace amounts.

Interestingly, $Zr(CH_2Ph)_4$ is photosensitive, more so than Ti(CH₂Ph)₄. Irradiation in toluene at -60° gave a brown product believed to be $Zr(H)(CH_2Ph)_2(CH_2C_6H_4CH_2Ph)$. Hydrolysis gave hydrogen, toluene, and a 1:1 mixture of o - and p -benzyltoluene. Though methanolysis with CH3OD gave HD, infrared or 'H NMR spectroscopy did not confirm a Zr-H bond. Photochemical activation is apparently necessary to observe rapid olefin polymerization by Zr(CH₂Ph)₄.⁵⁶

Single-crystal x-ray structural determinations have been completed for the three $M(CH_2Ph)_4$ species (M = Ti, Zr, or Hf). $66,75,76$ in contrast to Sn(CH₂Ph)₄, an essentially perfect tetrahedron with C-Sn-C angles of 108-111° and a mean Sn-C-C angle of 111^o, the M(CH₂Ph)₄ structures are distorted with rather large variations in the C-M-C angles (e.g., **99-1** 18' for Hf) and small mean M-C-C angles (Hf, **94';** Zr, **92';** Ti, 103'). Several authors have suggested that metal interaction with the phenyl ring contributes to their stability though no abnormal 'H NMR spectral features support this proposal. The structure of $V(CH_2Ph)_4$ is also a distorted tetrahedron as suggested by the well-resolved ESR spectrum at room temperature.⁷⁰

Several adducts of tetrabenzyl species are known. Ti(CH₂Ph)₄(bpy) has been isolated at room temperature^{35,63} and adducts with TMEDA, TMPDA, TMMDA, and py are formed in solution.³⁵ At low temperature $Ti(CH_2Ph)_4(e^{t}her)_2$ crystallizes from ether.⁶³ Adducts of $Zr(CH_2Ph)_4$ with bipyri $dy1^{67,73,77}$ and py, THF, thioxane, and other donors⁷⁷ have also been isolated. An 'H NMR study67-69 of Lewis base interactions with $Zr(CH_2Ph)_4$ and $H(CH_2Ph)_4$ has shown that (i) only monoadducts form for $L = py$, quin, PMe₃, or THF; (ii) the equilibrium constant for pyridine adduct formation in chlorobenzene is ca. 13 for Zr(CH2Ph)4 and ca. **460** for Hf(CH₂Ph)₄; (iii) no adducts form with NR₃ (R = Et, Pr, Bu, or Ph), TEEDA, diethyl ether, dioxane, PPh₃, P(OMe)₃, P(OEt)₃, cyclooctene, cyclohexene, or 1,1-diphenylethylene; and (iv) bipyridyl forms insoluble M(CH₂Ph)₄(bpy). Clearly steric factors play an important role.

Reaction of $TiCl_3(THF)_3$ with PhCH₂MgCI in ether at -45° gives a brown solution which may contain $Ti(CH_2Ph)_3$ as suggested by reaction with $3.5l_2$ to give Til₄ and 3PhCH₂1.78 When the temperature of the initial mixture rose to **20'** a red color developed, ethane, methane, and smaller amounts of ethylene, propane, and pentane were evolved, and $Ti(OEt)_{2}(CH_{2}Ph)_{2}$ (ether)₂ was subsequently isolated. The solvent-free form, $\left[\text{Ti}(\text{OEt})_2(\text{CH}_2\text{Ph})_2\right]_2$, was prepared from Ti(CH₂Ph)₄ and 2 mol of ethanol.^{58,63}

The reaction between $TiCp_2Ph_2$ and 2LiCH₂Ph reportedly gives Ti(CH₂Ph)₂.⁷⁹ The authors suggested a polymeric solidstate structure. It dissolved in tetrahydrofuran in which it reacted with HCI to give **2** mol of toluene.

Neither $Cr(CH_2Ph)_4$ nor $Cr(CH_2Ph)_3$ has been isolated. ESR evidence for the former was obtained^{29,80} while the latter probably exists at low temperature.⁸¹ On hydrolysis at 20 $^{\circ}$ the mixture from the -78° reaction of 3PhCH₂MgCI and $CrCl₃(THF)₃$ gave H₂, toluene, bibenzyl, 2-benzyltoluene, and $[Cr(\eta^6-2-benzyltoluene)(\eta^6-toluene)]^+$, isolated as its tetraphenylborate salt.

The dimeric complexes, $M_2(CH_2Ph)_6(M = Mo, W)$, are probably structurally similar to $Mo_{2}(CH_{2}SiMe_{3})_{6}^{82}$ (section 4, below). Preparative details have not yet been published.

Finally, $Mn(CH_2Ph)_2$ was prepared in THF but not isolated.32

4. Neopentyl, Trimethylsilylmethyl, and Related Alkyls

Neopentyl and trimethylsilylmethyl ligands have been used to prepare what is now the most extensive class of binary metal alkyls.

The group 4 M(CH₂CMe₃)₄ complexes are usually prepared from neopentyllithium and MCI4 in an ethereal solvent and red-purple $Cr(CH_2CMe_3)_4$ by air oxidation or disproportionation of $[Cr(CH_2CMe_3)_4]^-$ in hydrocarbons or ether.^{29,80,83-85} (In THF Cr(CH₂CMe₃)₃(THF)₃ does not readily disproportionate.²⁸) The pale yellow to colorless group 4 complexes are all volatile, low-melting solids which sublime in vacuo without decomposition. Attempted preparation of $V(CH_2CMe₃)₄$ from either VCI₄ or VOCI₃ and neopentyllithium in saturated hydrocarbons gave brown solutions whose **ESR** spectra were not consistent with its formation. 84

Corresponding M(CH₂SiMe₃)₄ complexes have been prepared for $M = Ti$, Zr, Hf, and Cr, and also for $M = V$ by analogous The group **4** complexes are liquids at room temperature. The green vanadium complex is somewhat unstable at room temperature in contrast to the group **4** complexes. $Cr(CH_2SiMe_3)_4$ is remarkably inert toward water, dilute mineral acids, amines, phosphines, alkenes, phenylacetylene, and CO at 100° and 200 psi, but reacts with oxygen. In contrast, V(CH₂SiMe₃)₄ decomposes in aqueous solutions, alcohols, and chlorinated solvents and is oxidized to V(O)(CH₂SiMe₃)₃ on passage through a cellulose column, yet it does not react under mild conditions with CO, alkenes, acetylene, or phosphines.

Related MR_4 species such as $Cr(CH_2CMe_2Ph)_4$ and $Cr(CH_2CPh_3)_4, ^{29,80}$ $Zr(CH_2SiMe_2Ph)_4, ^{87}$ and Ti(CH₂Si-Me₂Ph)₄⁹⁰ have been mentioned. A crystal structure⁹¹ of Cr(CH₂CMe₂Ph)₄ clearly shows the dense packing around the metal which is assumed to be at least partly responsible for its chemical stability. It is a slightly distorted tetrahedron with Cr-C distances from **2.01** to 2.07 **A.**

Because the Cr(lV) oxidation state is rather unusual, the ESR and electronic spectra of CrR4 species have been thoroughly studied.^{28,29,80,92} Several ESR signals are generally observed, possibly because the alkyls cannot rotate about the metal-carbon bond. Both electronic and ESR spectra suggest a distorted tetrahedral symmetry.

Attempts to prepare binary -CH₂CMe₃ or -CH₂SiMe₃ complexes from NbCl₅, TaCl₅, MoCl₅, and WCl₆ gave unanticipated results, probably in part because more than four of these bulky ligands on a single metal is sterically unfavorable. The niobium and tantalum reactions gave both bridging and terminal carbene complexes (see section **ll.A.5),** probably by α -hydrogen abstraction.

Attempts to prepare $-CH₂SiMe₃$ complexes of Mo(V) or W(VI) employing Me₃SiCH₂MgCI gave the dimers, $M_2(CH_2SiMe_3)_6$, which are stable in air for a short time and sublime in vacuo at 120° (ref 83, 88). A crystal structure⁸² of $Mo₂(CH₂SiMe₃)₆$ shows the following essential features: (i) a short (2.167 Å) Mo-Mo distance indicates a multiple metalmetal bond; (ii) each molybdenum has a distorted tetrahedral coordination and the three alkyl groups on one atom are staggered relative to the three on the second: (iii) the Mo-CH₂-Si angle (121^o) and the Mo-Mo-CH₂ angle (101^o) are probably a consequence of the mutual repulsion of the bulky trimethylsilyl groups; and (iv) the mean $Mo-CH₂$ distance is 2.131 A. The tungsten analog is isostructural.

To avoid low yields the analogous $Mo₂(CH₂CMe₃)₆$ compound must be prepared with neopentyllithium instead of the Grignard reagent.⁸⁴ Though it is sufficiently volatile to sublime (130 $^{\circ}$), it is slightly more unstable than the $-CH₂SiMe₃$ analog and decomposes at this temperature.

All complexes in this section have been fully examined by infrared, 'H NMR, or ESR (for V and Cr) spectroscopy, but only qualitative reactions reported. Unfortunately, the large ligands hinder a reagent's approach to the metal and thereby limit to some extent reaction under conditions where the complex remains intact. Ironically, this is probably one reason why the complexes can be prepared in the first place.

5. Alkylidenes

Neopentyl and CH₂SiMe₃ alkylating agents react with NbCI₅ and TaCI₅ to give compounds containing carbene ligands probably because more than four bulky alkyl groups cannot fit around a single metal atom.

Addition of MCl₅ to 5.5 mol of Me₃SiCH₂MgX in ether did not give $M(CH_2SiMe_3)_5$ (M = Nb or Ta) but $[M(CH_2Si Me₃$ ₂(CSiMe₃)]₂.^{83,84} These red (Nb) or orange (Ta) complexes do not sublime in vacuo though they are soluble in pentane and decompose only at their melting points, 152 and 170°, respectively. Formulation of the Nb complex has been confirmed by a crystal structure⁹³ shown schematically in Figure 2. The two crystallographically independent dimers do not differ significantly. The metals and bridging carbon atoms lie in a plane but neither the Nb-C(bridging)-Si angles nor the Nb-C(bridging) distances are identical in a given molecule. Nevertheless, the Nb-C-Nb-C ring can be described as quasi-aromatic, a fact which explains the diamagnetism of these complexes and possibly also why they do not readily react with NO, CO, tertiary phosphines, amines, or alkenes. Their mechanism of formation is unknown and neopentyl analogs were not obtained.

The attempted preparation of $Ta(CH_2CMe_3)_{5}$ from $Ta(CH_2CMe_3)_3Cl_2$ and 2 mol of LiCH₂CMe₃ in pentane gave 2 mol of LiCI, 1 mol of neopentane, and very soluble, orange, crystalline Ta(CH₂CMe₃)₃(CHCMe₃).⁹⁴ Its ¹H NMR, mass spectrum, and molecular weight are consistent with formulation as

Hydrolysis with **5** mol of DCI produced 3.7 mol of a mixture of three parts neopentane- d_1 and one part neopentane- d_2 while **13C** NMR confirmed the presence of a carbene-carbon with a 250-ppm chemical shift. Labeling experiments suggest formation via Ta(CH₃CMe₃)₅ by intramolecular α -hydrogen abstraction. An exactly analogous reaction gives wine-red $Nb(CH_2CMe_3)_3$ (CHCMe₃).⁹⁵ The former is extraordinarily stable thermally, the latter considerably less so. Their isolation demonstrates clearly that, under some conditions, (i) alkylidene complexes are stable and (ii) intramolecular α -hydrogen abstraction is a viable mode of "decomposition" of binary alkyl complexes.

Interestingly, elimination of a second mole of neopentane

Figure 2. The structure of $[Nb(CH_2SiMe_3)_2(CSiMe_3)]_2$ (ref 93).

Figure 3. The structure of $Ta(CH_2CMe_3)_3[C(CMe_3)(Li·N,N'-dimethyl$ piperazine)].

to give the carbon analog of $[M(CH_2SiMe_3)_2(CSiMe_3)]_2$ (vide supra), i.e., perazine)].
9 give the carbon analog of $[M(CH_2SiMe_3)_2(CSiMe_3)]_2$ (vide
upra), i.e., \longrightarrow 0.5 $[M(CH_2CMe_3)_2(CCMe_3)]_2$
 $M(CH_2CMe_3)_3(CHCMe_3)$

$$
M(CH_2CMe_3)_3(CHCMe_3) \stackrel{-C_5H_{12}}{\longrightarrow} 0.5[M(CH_2CMe_3)_2(CCMe_3)]_2
$$

does not occur readily despite the fact that lithium reagents react with $M(CH_2CMe_3)$ ₃(CHCMe₃) to give $M(CH_2CMe_3)$ ₃[C(C- Me_3 (Li-L_x)] (L_x = N,N'-dimethylpiperazine, for example), ⁹⁴ i.e., the neopentylidene α proton is more acidic than a neopentyl α proton. Failure to isolate M(CH₂SiMe₃)₃(CHSiMe₃) may possibly be ascribed to the greater acidity of protons α to silicon vs. α to carbon.

A crystal structure of the lithiated Ta species has been completed⁹⁶ (see Figure 3). The extraordinarily short tantalum to "carbene"-carbon distance (1.76 **A)** and large Ta-C-CMe3 angle suggest that this compound should be viewed as a "carbyne" complex analogous to group 6 carbyne complexes.⁹⁷

6. Other Alkyls

a. Bulky Alkyls

Complexes of the type $M(1-norborn)/₄$ where $M = Ti$, Zr, Hf, V, and Cr (also Mn, Fe, and Co) were prepared from the metal halides and 1-norbornyllithium in pentane.⁹⁸ Disproportionation to yield M(IV) species was the rule when lower valent halides were used. Chromium (2,3,3-trimethylbicy- $\text{clo}[2.2.1]$ hept-1-yl)₄ could also be prepared but with 2,2,3trimethylbicyclo^[2.2.1]hept-1-yl only CrR₃ (61% yield) resulted. Ligand interrepulsions must be too large in CrR₄ in

this case. All are pentane-soluble, monomeric, and pass through an alumina column unchanged. The authors attribute their thermal stability $(t_{1/2} = 29 \text{ h at } 100^{\circ} \text{ for TiR}_4)$ to the fact that β -hydrogen abstraction is unfavorable (unsaturation at a bridgehead carbon would result) and their chemical stability $(M = Ti, V, or Cr)$ to the ligands' close packing around these relatively small metals. It is also interesting to note 1-norbornyl has no α proton which would allow decomposition by an α -abstraction process. ESR⁹⁸⁻¹⁰⁰ and preliminary x-ray data (for Cr(1-norbornyl)4) suggest tetrahedral structures.

Apparently only three $CH(SiMe₃)₂$ ligands can fit conveniently about a first-row transition metal. Paramagnetic M[CH(SiMe3)2]3 species (M = Ti, **V,** Cr) have been isolated^{101a} and at least the Cr complex is quite stable thermally; it decomposes at ca. 80° . Excess LiCH(SiMe3)₂ reacts with MCI₄ (M = Zr, Hf) in ether to give only M[CH(SiMe₃)₂]₃CI while 1 mol reduces TiCp₂Cl₂ to [TiCp₂Cl]₂. CH(SiMe₃)₂ must be about the size of $N(SiMe₃)₂$ for which many three-coordinate complexes are known.^{101b}

Though the tert-butyl ligand is very susceptible in most circumstances to β -hydrogen abstraction, Cr(CMe₃), can be prepared in 60% yield from $Cr(OCMe₃)₄$ and LiCMe₃ in pentane.²⁸ It can be chromatographed on alumina and sublimes at 10⁻⁴ mm. The activation energy for decomposition $(t_{1/2} =$ **4.5** min at 70' in heptane) measured over the range *55* to 80° was 29 \pm 3 kcal/mol. At 80° the decomposition products were isobutane and isobutylene (6.5: 1) containing small amounts of methane, ethane, and propane. Its stability is attributed to the fact that the methyl groups are tightly packed and cannot orient properly for β -hydrogen abstraction to occur.

Colorless Ti(1-adamantyl)₄ was prepared recently in 18% yield from TiC14, Na, and 1-chloroadamantane in refluxing cyclohexane. **102a** Considerable biadamantane was also obtained, consistent with a radical reaction. The extraordinary thermal stability of Ti(1-adamantyl) $_4$ (mp 233-235°; "the compound proved difficult to decompose with $HNO₃-HF H₂O₂$ mixtures at 170° when analyzing for the metal") combined with the fact that the adamantyl radical is fairly longlived (ref 13 in ref 102a) suggests that homolytic metal-carbon bond cleavage is not a common *primary* decomposition step (see section VI.E). β -Hydrogen abstraction is unfavorable for reasons analogous to those stated for 1-norbornyl; 1-adamantyl also has no *a* hydrogen.

b. "Unstable" **Alkyls**

Except for special cases (e.g., $Cr(CMe₃)₄$) and in monoalkyl complexes with an 18 valence-electron count (e.g., $Mn(CO)_{5}Et$), complexes of alkyl groups containing a β -hydrogen atom are accessible only at low temperature or in coordinating solvents. For example, TiR₄ species where $R = Cy$, Bu, CMe₃, Prⁱ, Pr, and Et can be prepared without accompanying reduction to Ti(III) by adding $Ti(OCMe₃)₄$ to RMgX at low temperatures.⁵⁹ The CrR₄ species where R = Prⁱ, s-Bu, Bu, and cyclohexylmethyl, prepared as for $Cr(CMe₃)₄$, are "more or less stable between -78° and room temperature" in pentane.²⁸ For R = Bu and cyclohexylmethyl, CrR₄ appeared stable for several hours at **0'** in pentane. Decomposition rates of $MnR₂$ in THF³² are almost certainly moderated by formation of MnR₂(THF)₂. They vary in the order R = Pr' > CMe₃ > Et > Pr ~ Bu >> Me ~ CH₂CMe₃ ~ CH₂Ph. ("MnPr₂" decomposes 10% in 2 h at **25'.)** Though decomposition of longchained alkyl complexes produced in situ may be accelerated by metal salts, reduced metal complexes, etc., the basic contention concerning their instability appears sound. Exceptions, where the valence-electron count is less than 18, will probably have a crowded coordination sphere similar to that in $Cr(CMe₃)₄$.

c. **"Mixed" Alkyls**

A few complexes containing more than one type of alkyl are known. Not unexpectedly, properties of the mixed binary complexes are roughly intermediate between those of the parent species.

Exchange between TiMe₄ and BPh₃ or B(CH₂Ph)₃ at -78° in diethyl ether yields TiMe₂Ph₂ or TiMe(CH₂Ph)₃, respectively, and $BMe₃$.⁶³ (TiMe(CH₂Ph)₃ has also been prepared from $Ti(CH_2Ph)_3CI$ and MeMgl. 65) in the latter case raising the temperature to -30° caused complete exchange to give $Ti(CH_2Ph)_4$. The reaction with BPh₃ was not altered by chang-

ing the reactant ratio. One suggested mechanism
\nTiMe₄ + BPh₃
$$
\rightleftharpoons
$$
 [TiMe₃]⁺ [BMePh₃]⁻ \rightleftharpoons

TiMe₃Ph + BMePh₂ \rightleftharpoons etc.

should be compared with the proposed mechanism of methyl exchange between $Time₄$ and $AlMe₃$ in the presence of ether: **102b**

$$
\text{Time}_4 + \text{AlMe}_3 \rightarrow [\text{Time}_3]^+ [\text{AlMe}_4]^+
$$

The reaction of $Cr(CHPh_2)Cl_2(THF)_2$ with phenyllithium to give Cr(CHPh₂)Ph₂(THF)₂ was briefly mentioned⁵⁰ as was a bipyridyl adduct.

B. Anionic

Most "anionic" metal alkyls contain one or more alkali metal counterions. Since lithium alkyls contain bridging alkyl groups, it is not surprising that the "anionic" complexes contain the Li-R-M arrangement. The truly anionic complex, $[Li(THF)_4]$ ⁺[TaPh₆]⁻, probably exists because a structure containing Li-Ph-Ta bonds is sterically unfavorable.

1. Methyl

Addition of methyllithium to TiMe₄ in ether followed by 2 mol of dioxane gives a precipitate, $Li[TiMe₅]\cdot2diox.^{103,104}$ Conductivity measurements in THF favor an ionic formulation but the manner in which dioxane is bound is uncertain. Similarly, TiMe₄ and LiPh or LiCH₂Ph (LiR) gave Li $[TiRMe₄]-2diox$ ane, and $Time_2Ph_2$ and LiPh, Li $[Time_2Ph_3]$.2dioxane. All decompose at -20° to 0°

Reaction of $ZrCl₄$ with >6 mol of methyllithium in ethertoluene gives $Li_2[ZrMe_6]$.¹⁰⁵ The bright yellow crystals are stable under nitrogen at **0'** for several hours and dissolve in ether, tetrahydrofuran, and toluene, but not in aliphatic hydrocarbons. In toluene, alcoholysis produced only methane **(6** mol per Zr). **A** structure was not proposed but may well contain Li-Me-Zr bridges as found in $Li_3[CrMe_6]$.3(solvent) (vide infra). Some bridging structure is probably retained even in coordinating solvents, though $[ZrMe_6]^{2-}$ does not seem unreasonable.

Similar anionic niobium and tantalum complexes, $[MMe_n]^{5-n}$ ($n = 6$ or 7), are probably formed in ether from appropriate quantities of $\text{Table}_3\text{Cl}_2$ or NbMe_2Cl_3 and methyllithium.²² However, the ¹H NMR spectrum of $Li_2[TaMe_7]$ in THF suggests equilibria between at least two unidentified species and methyllithium. Ether solutions are remarkably stable compared with those of TaMe₅ or NbMe₅ yet methyllithium is readily lost on addition of dmpe and $MMe₅(dmpe)$ isolated.

The chromium-methyl system has been studied most extensively so far. Reaction between CrCl₃ and 6LiMe in ether at -18° gives Li₃[CrMe₆], isolable as blood-red Li₃CrMe₆-3dioxane on addition of dioxane.'06 It is insoluble in hexane and benzene (it slowly decomposes in these solvents) but is soluble and stable in ether and dioxane in which it is monomeric and apparently undissociated. Its reactions with H_2O and I₂ give 6 mol of methane and 6 of methyl iodide, respec-

Figure 4. A schematic drawing of $Li_3CrMe₆(diox)₃$.

Figure 5. Partial schematic structure of $[Li_2CrMe_4(THF)_2]_2$.

tively. Its structure¹⁰⁷ is shown schematically in Figure 4. Six methyl groups are disposed in a slightly distorted octahedral arrangement around Cr, and each tetrahedral lithium atom is surrounded by two methyls and two dioxanes. The dioxanes bridge to lithiums in other molecules. The Ti and **Zr** complexes discussed above are most likely structurally related.

In ethers Li₃CrMe₆ slowly decomposes to give a dimeric $Cr(II)$ complex: 106

$$
2 \text{Li}_3 \text{CrMe}_6 \stackrel{\text{R}_2 \text{O}}{\longrightarrow} [\text{Li}_2 \text{CrMe}_4(\text{OR}_2)_2]_2 + 2 \text{LiMe} + 2 \text{``Me·''}
$$

The dimer may also be prepared directly from $CrCl₂$ and methyllithium or from $Cr_2(\eta^3-C_3H_5)_4$ and 8LiMe in THF.¹⁰⁸ Its reduced paramagnetism $(\mu = 0.57)$ suggests a Cr-Cr bond. The structure¹⁰⁹ of $[L_2CrMe_4(THF)_2]_2$ is partially shown in Figure 5. Each of the four equivalent faces of the tetragonal prismatic unit composed of eight methyl groups and two joined chromium atoms is capped with a lithium atom to which a THF molecule is also bound. The authors postulate a quadruple Cr-Cr bond.

 $MoCl₃(THF)₃$ is reduced by 5LiMe to give Li₂MoMe₄(THF)₂ and 0.5 mol of ethane, according to the authors.^{110a} (Li₃-MoMe₆ is apparently not isolable.) Li₂MoMe₄(THF)₂ is a diamagnetic dimer analogous to $[Li_2CrMe_4(ether)_2]_2$ (vide supra). This is confirmed by an x-ray structure^{110d} where Mo-Mo = 2.148 **A** and Mo-C(mean) = 2.29 **A.**

Also in this class is $[Li_2Cr(C_4H_8)_2(\text{ether})_2]_2$ (see section 11.6.3 and Figure 6).11' The metallocyclic rings are staggered with Cr-Cr = 1.975 Å. The Cr-CH₂ distances and angles are variable.

The only other "anionic" methyl complex is $Li[MnMe₃]^{31}$ Few details concerning its preparation and properties are available.

2. Phenyl

The reaction of LiPh with CrCl₃ in ether¹¹² simply gives metathesis in contrast to reduction to Cr(0) usually observed with Grignard reagents. The orange crystalline product, isolated in high yield, has the formula $(LIPh)_{3}CrPh_{3}(ether)_{2.5}$. It is also formed starting with CrBr₃, CrCl₃(py)₃, Cr(acac)₃, and Cr(ll) salts, disproportionation occurring in the latter in-

Figure 6. Partial schematic structure of $[Li_2Cr(C_4H_8)_2(Et_2O)_2]_2$.

Figure 7. A schematic drawing of $Na₂CrPh₅(ether)₃(THF)$.

stance.¹¹³ Hydrolysis with water gives 6 mol of benzene but some biphenyl is formed on reaction with iodine:

(LiPh)₃CrPh₃(ether)_{2.5} + 4.5l₂ \rightarrow 3Lil + Crl₃ + 3Phl + 1.5Ph₂

Somewhat surprising also is its reaction with molecular hydrogen to give postulated Li₃CrHPh₅ and Li₃Cr₂H₃Ph₆. Its molecular weight in benzene is close to theory initially, but ether is gradually lost and $(Liph)_{3}CrPh_{3}$ can be isolated as a yellow powder. The magnetic moment (3.61) is characteristic of octahedral Cr(lll) and its structure is probably similar to that of $Li_3CrMe_6(diox)_3$ (cf. the structure of $Na_2CrPh_5(ether)_3(THF)$ below).

Lower members of the M_{x-3} CrPh_xS_y family ($x = 4, 5$, or 6, $M = Na$ or Li, $S = an$ ether) have also been studied. The blue-green product resulting from the reaction of NaPh with CrCl₃ is Na₂CrPh₅(ether)₃,¹¹⁴ analogous to the lithium derivative reported several years earlier.¹¹⁵ (Yellow Na₃CrPh₆ is unstable toward loss of NaPh.) It is monomeric in ether (ebullioscopy), has a magnetic moment of 3.66, and gives olivegreen Na₂CrPh₅(bpy)₂ (μ = 3.90) with bipyridyl. A crystal structure¹¹⁶ of Na₂CrPh₅(ether)₃(THF) reveals a greatly distorted trigonal-bipyramidal geometry about Cr with each Na interacting with two phenyl rings (Figure 7).

The lowest members ($x = 4$, M = Li, Na) are prepared by redistribution reactions¹¹⁷ (ether molecules omitted):
 $4Na_2CrPh_5 + CrCl_3 \rightarrow 5NaCrPh_4 + 3NaCl$

$$
4Na_2CrPh_5 + CrCl_3 \rightarrow 5NaCrPh_4 + 3NaCl
$$

 $2Li_3CrPh_6 + CrCl_3 \rightarrow 3LiCrPh_4 + 3LiCl$

Each can be characterized as its blood-red dimethoxyethane adduct, MCrPh₄(dme)₄ (μ = 3.85 for M = Na, 3.65 for M = Li). The magnetic moments appear to rule out formulation as $[M(dme)₄]$ ⁺[CrPh₄]⁻; octahedral Cr(III) with two dme ligands and two bridging phenyl groups to $Na(dme)_2$ is more plausible.

All members of the Cr(III) class react with CrCI $_3$ to yield

Cr(II) complexes under proper conditions:¹¹⁸
\n
$$
4Na_2CrPh_5(ether)_3 + CrCl_3(THF)_3 \rightarrow
$$
\n
$$
5NaCrPh_3(ether)_{1.5} + \frac{5}{2}Ph_2 + 3NaCl
$$
\n
$$
2Li_3CrPh_6(ether)_{2.5} + CrCl_3(THF)_3 \rightarrow
$$

 $3LiCrPh₃(ether)_{1.5} + $\frac{3}{2}$ Ph₂ + 3LiCl$

These are brown-black, soluble in THF, benzene, diglyme, and diethyl ether, and react characteristically with H_2O , $HgCl_2$ and I_2 . Their reduced magnetic moments (ca. 0.6-0.8) and dimeric character (molecular weight studies) indicate they probably are structurally similar to $[Li₂CrMe₄(THF)₂]$ ₂.

Finally, a monomeric, anionic Cr(ll)-phenyl complex has been isolated: 50

$$
\text{Cr}(\text{CHPh}_2)\text{Ph}_2 \text{ (in situ)} + 2\text{LiPh}_{\overline{\text{THF}-\text{ether}}}^{\overline{\text{THF}-\text{ether}}} \text{Li}_2\text{Cr}(\text{CHPh}_2)\text{Ph}_4
$$

-0.5Ph}_2\text{CHCHPh}_2
Li₂\text{CFPh}_4(\text{THF})_4

It can also be prepared from $CrCl₂(THF)₂$. The yellow crystalline complex is stable in an inert atmosphere to approximately 90'. Its magnetic moment (4.73) is characteristic of a high-spin $d⁴$ species and suggests it is not a member of the dimeric class, $[L₂CrR₄S₂]₂$, but the sole member of a third class of "anionic" organochromium species. Apparently the large phenyl groups prevent dimerization.

A molybdenum species, (LiPh)₃MoPh₃(ether)₃, is formed in 20% yield on reaction of MoCl₅ with 10 mol of LiPh in ether at -30° (ref 119). The yellow crystals, obtained from a dark red reaction mixture, dissolve in ether, benzene, THF, or butyl ether to again give a red color which is believed that of a species formed by **loss** of LiPh. Hydrolysis produced ether and benzene in a 1:2 ratio while thermal decomposition left a black, crystalline, ether-soluble residue which was not identi-

\n
$$
\text{field. Its "anomalous" reaction with HgCl}_2, \text{viz.}
$$
\n

\n\n $\text{(LiPh)}_3\text{MoPh}_3 + 6\text{HgCl}_2 \rightarrow 3\text{PhHgCl} + 1.5\text{C}_6\text{H}_5 - \text{C}_6\text{H}_5 + \text{MoCl}_3 + 3\text{LiCl}$ \n

was taken as evidence against the "Li₃MoPh₆(ether)₃" formulation suggested by other authors.¹²⁰

An apparently analogous tungsten complex was prepared from WBr₅ and excess LiPh in ether at $0-10^{\circ}$ (ref 121). It is a finely crystalline, black powder which gives deep violet benzene and ether solutions. Reformulation as (LiPh)4WPh₂(eth $er)$ ₄ was suggested¹²² based on titration¹²³ and flame photometry.¹²⁴ It reacts with 2 mol of hydrogen reportedly to $give WH(LiPh)₄. ^{125,126}$

Anionic vanadium complexes have also been reported. For $example, ¹²⁷$

$$
e,121
$$

10LiPh + VCl₃(THF)₃ \longrightarrow (LiPh)₄VPh₂(ether)₄

As for Cr, the Grignard reagent gives zerovalent species.¹²⁸ The permanganate-colored crystalline product $(\mu = 3.85)$, soluble in ether and benzene, was later prepared from VC_{D2}Ph and 7LiPh in 66% yield.¹²⁹ With only 2 mol of LiPh in the latter reaction, (LiPh)VPh₂(ether) is formed. A third complex was also reported:

$$
V\text{Cp}_2 + 7\text{LiPh} \rightarrow 2\text{LiCp} + (\text{LiPh})_5\text{VPh}_2\text{(ether)}_5
$$

Two niobium complexes have been reported: 130, 131

NbBr₅ + 8LiPh
$$
\longrightarrow
$$
 (LiPh)₃NbPh₄(ether)₃ ($\mu = 1.66$)
\nNbBr₅ + 9LiPh \longrightarrow (LiPh)₄NbPh₂(ether)_{3.5} ($\mu = 1.81$)
\n \longrightarrow 24 h

Both are black-violet, thermally unstable, and soluble in benzene or dioxane; they do not conduct in dioxane. Niobium oxidation states $4+$ and $2+$, respectively, but no definite structures, were proposed.

Black-violet $(LIPh)_{4}$ TaPh₂(ether)_{3,5} was prepared in an identical manner^{132a} (μ = 1.68). A bright green intermediate, $(LiPh)_{3}TaPh_{3}(ether)_{2.5-3}$, was also isolated.

In another report^{132b} the authors draw a distinction between magnetically "normal" and magnetically "abnormal" complexes of the type (LiPh)₄MPh_x(ether)_v on the basis of oxidation products with oxygen. The complexes of Nb, Ta, and W give triphenylene while pure LiPh and those of V and Cr do not. They inferred the former contain ortho-disubstituted phenyl groups, a hypothesis supported by the 'H NMR spectrum of the W complex in which an A_2B_2 pattern characteristic of an ortho-disubstituted phenyl group appears. They suggested the Nb, Ta, and W complexes therefore be reformulated as $M(IV)$ species, $(LiPh)_{4}M(C_{6}H_{4})_{2}$. The manner in which phenylene coordinates was not discussed, but it could conceivably bridge two metals as in Ni₂(o -C₆H₄)₂(PEt₃)₄.¹³³

The only truly anionic phenyl complex is $[TaPh_6]^-$ (ref 134). Addition of 5 mol of LiPh (17.8% in a benzene:ether solvent, $77:23$) to TaCl₅ in benzene at ca. 20° followed by filtration, and addition of tetrahydrofuran gave crystals of orange, light-sensitive $[Li(THF)_4]^+$ $TaPh_6]^-.$ Other reactant ratios or temperatures gave lower yields *(0;* none at all) though this could at least partially be a function of product solubility. Salts containing $[NEt₄]⁺$ and $[AsPh₄]⁺$ were prepared by metathesis reactions. All have 'H NMR spectra consistent with their formulation and conduct in tetrahydrofuran or acetonitrile. Isolation of $\text{[TaPh}_6]^+$ is surprising since the metal is commonly reduced in systems of this type. Reactant ratio, order of addition of reagents, and mixed solvent are probably all crucial in preventing reduction. $\lceil \text{TaPh}_6 \rceil$ probably forms directly but disproportionation of a reduced species is not unreasonable, especially in view of the low isolated yield.

WH(LiPh)₄,^{125,126} Li₃CrHPh₅,¹¹³ and Li₃Cr₂H₃Ph₆¹¹³ are the postulated products of the reaction of molecular hydrogen with the corresponding anionic phenyl complexes (vide supra). Perhaps the best characterized complexes of this type are diamagnetic Li₂MoPh₂H₂^{110b} and Li₄Mo- $Ph_2H_2Me_2$ (diox)₂^{110c}:

 $5Li_3M_0Ph_6 + M_0Cl_3$ (THF)₃ \rightarrow 6Li₂MoPh₂H₂ + $3C_6H_5-C_6H_5 + 3LiCl + 12''C_6H_4$

 $Li_2M_0Ph_2H_2 + 2LiMe-diox \rightarrow Li_4MoPh_2H_2Me_2(diox)_2$

Apparently ortho-hydrogen abstraction (see section V1.A) gives dehydrobenzene which can be trapped by anthracene to give triptycene.^{110b} Both complexes were characterized by hydrolyses and by 'H NMR in which the peaks at *7* 8.05 and 8.70, respectively, were assigned to the proposed bridging hydride ligands. Further investigation of these species, including an x-ray structure, would be relevant to understanding the peculiarities of MPh_x species, their mode of decomposition, and metal reduction.

No group 4 or 7 phenyl complexes analogous to those described in this section have been reported. Ether-insoluble LiMnPh₃ (as well as the analogous p -tolyl and p -anisyl complexes) was reported in an early work,³¹ but preparation and properties have not been described in detail.

3. *Other Alkyls*

Dark blue $[Cr(CH_2CMe_3)_4]^-$ (ref 29, 83) and blue-green $[Cr(CH₂SiMe₃)₄]⁻$ (ref 83, 88) are the products of reaction of $CrCl₃(THF)₃$ in tetrahydrofuran with the appropriate lithium reagent. They are stable only in THF, and attempts to isolate crystalline salts by addition of large cations failed. They are

readily oxidized in solution by air to $CrR₄$ from which they may be regenerated polarographically or with sodium amalgam. Half-wave reduction potentials with $[NBu_4]^{+1-}$ as the supporting electrolyte in ethanol are -1.28 V for $[Cr(CH_2SiMe_3)_4]^-$ and -1.65 V for $[Cr(CH_2CMe_3)_4]^-$ vs. the saturated calomel electrode. Electronic properties suggest that these are two of the few examples of tetrahedral Cr(1ll).

A chromium complex containing the bidentate alkyl - $(CH₂)₄$ - has been prepared: 135

$$
c_2a -
$$
 has been prepared:¹³⁵
CrCl₃ + 3LiC₄H₈Li-dioxane $\xrightarrow{\text{other}}$ Li₃Cr(C₄H₈)₃(diox)_{2.5}
(μ = 3.9)

It is believed analogous to $Li₃CrMe₆(diox)₃$ as suggested by physical properties and decomposition to $[Li_2Cr(C_4H_8)_2$ (eth $er)_2$ ₂. A structural determination of the latter showed it to be a member of the $[Li_2$ CrR₄S₂ $]_2$ family (S = solvent; see section II.B. 1). A preparation employing $LiC₅H₁₀Li$ gave another member of this family, $[Li_2Cr(C_5H_{10})_2(LiBr)(ether)_2]_2$, directly.

Other anionic phenyl chromium species have been isolated with pyrrolyl, diphenylamido, or dicyclohexylphosphido ligands¹³⁶ (eq 2). The first is in the MCrR₄ class (M = e.g., Li); **tic phenyl chromium species have**
 yl, diphenylamido, or dicyclohexy

2). The first is in the MCrR₄ class (
 $\begin{aligned}\n+ \text{LiC}_4H_4N &\longrightarrow \text{LiCrPh}_3(C_4H_4N)T\n\end{aligned}$

gands¹³⁶ (eq 2). The first is in the MCrR₄ class (M = e.g., Li);
\n
$$
+ \text{ LiC}_4H_4N \longrightarrow \text{LiCrPh}_3(C_4H_4N)(THF)_3
$$
\n
$$
(\mu = 3.78)
$$
\n
$$
\text{CrPh}_3(THF)_3 + 2\text{KNPh}_2 \cdot 3\text{diox} \xrightarrow{-\frac{1}{2}Ph_2} K_2\text{CrPh}_2(\text{NPh}_2)_2(THF)_2
$$
\n
$$
(\mu = 4.67) \qquad (2)
$$
\n+ 2\text{LiPCy}_2 \xrightarrow{-\frac{1}{2}Ph_2} \text{Li}_2\text{CrPh}_2(\text{PCy}_2)_2(THF)_4\n
$$
(\mu = 4.77)
$$

the second and third are in the M_2 CrR₄ class. Though paramagnetic, the latter two gave 'H NMR spectra with phenyl signals at *7* 2.72 and 2.75, respectively. No conclusions concerning the stability of the Cr(1ll) species, or, alternatively, the ease with which they are reduced to Cr(ll), could be drawn.

Finally, there are the poorly characterized, polymeric $LIMnBu₃$ and $LIMnEt₃$ species.³¹

C. Cationic

No simple cationic complexes, $[MR_x]^+$, have been reported, only cationic chromium complexes containing donor ligands, such as water (see section **V.E)** and amines. The latter are of the type cis - $[CrR_2(L-L)_2]$ ⁺ (R = Ph, ^{137, 138} CH_2SiMe_3 , ^{139, 140} anisyl, ^{138b, 141, 142} tolyl, ^{138b} and L-L = bpy, or $R = Ph$ and $L-L = phen.137$ It is also suspected that products with the composition WMeCI₅L₃ (L = e.g., py) are ionic, $i.e., [WMeCl₄L₃]+Cl^-$ (ref 143).

Orange-yellow $[CrPh_2(bpy)_2]$ ⁺ I^- is slightly soluble in alcohol, THF, and acetone, and thermally stable (dec **230').** Salts containing other anions $(X^- = CI^-, BT^-, NO_3^-, ClO_4^-,$ $[BPh_4]^-$ and $[BH_4]^-$) may be prepared by reaction with AgX. Significant bond lengths from three crystal structures are shown in Table I. The results indicate little correlation between the Cr-C bond length and the compound's "stability". **40**

D. Chelating Alkyls and Aryls

Several workers have prepared complexes containing alkyl or aryl ligands which chelate to the metal via a substituent donor. Their extraordinary thermal stability (the last decomposes at 350') may be attributed to the fact that the ligands are sterically bulky, stereochemically rigid, and chelate strongly. Attempts to prepare Ti and V species containing the $C_5H_4NC_6H_4$ ligand gave complexes which decomposed readily at room temperature.

Ill. Halide-Alkyl Complexes

Halide-alkyl complexes, MR_vX_z (X = halide), are usually prepared with a Grignard or lithium reagent deficiency or with milder alkylating agents like Zn, Hg, B, AI, Sn, or Pb alkyls. It is sometimes stated they are thermally less stable than binary alkyls, but the reverse in probably more often the case. They are best classified and discussed according to the metal.

A. Group 4

TiCl4 with Me₂AICI: In 1959 Beermann and Bestian¹⁴⁹ reported the reaction of

$$
1 \text{ Me}_2\text{AICI:}
$$
\n
$$
1 \text{ Me}_2\text{AICI:}
$$
\n
$$
1 \text{ NHe}_2\text{CI:}
$$
\n
$$
1 \text{ NHe}_2\text{CI:}
$$
\n
$$
1 \text{ NHe}_2\text{CI:}
$$
\n
$$
1 \text{ NHe}_3 + \text{ AIMe}_2\text{CI:}
$$

Addition of excess NaCl effectively removed AIMeCl₂ as $Na^{+}[AlMeCl₃]⁻$. TiMeCl₃ has also been prepared from reaction of TiCl4 with MeMgCl suspended in hexane¹⁵⁰ or with $ZnMe₂$.¹⁵¹ It also forms smoothly from any of the polymethylated Ti species and the appropriate quantity of TiCl4.

Methyltitanium trichloride is a deep violet solid which is stable at room temperature and melts at 28-29° to give a yellow oil. It gives yellow solutions in saturated or aromatic hydrocarbons and is monomeric in benzene. Its red ether solutions probably contain TiMeCl₃(ether),. Surprisingly, Ti-MeCl₃ reportedly decomposes more rapidly in ether than in hydrocarbons, a reversal of the usual behavior for alkyls.¹⁵² Decomposition in chlorinated solvents gives methyl chloride, in deuterated hydrocarbons only $CH₄$. 153

Black-violet TiMeBr₃ was prepared by the ZnMe₂ route^{151,152} and has properties analogous to TiMeCl₃. It has also been prepared admixed with TiMe₂Br₂ from TiBr₄ and MeMgBr suspended in benzene at 0-10° (ref 150). It is less stable than the trichloride, an observation which appears general for many halide-alkyl complexes.

Further alkylation of TiMeCl₃ with AlMe₃¹⁴⁹ or ZnMe₂¹⁵¹ gives black TiMe₂Cl₂. It is markedly less stable than TiMeCl₃ and decomposes at ca. -10° to TiCI₂. Addition of dioxane to a yellow hexane solution gives a yellow dioxane adduct. 'H NMR studies of methyl exchange between ZnMe₂ and Ti- $Me₂Cl₂$ and between Ti(CD₃)Cl₃ and TiMe₂Cl₂ have been described, **154** but the system's complexity makes interpretation difficult.

Berthold and Groh¹⁸ briefly mentioned TiMe₃I, but no further details have appeared.

The TiMe_n X_{4-n} ($n = 1, 2, 3, 4$) system is the only one in which at least one representative species has been isolated for each value of *n.* Thermal stability of the pure (solventfree) complexes decreases in the order $n = 1 > 2 > 3 > 4$ and $X = CI > Br > I$. More information on their solid-state structure could shed some light on the relationship of thermal stability to molecular structure. For example, TiMeCI₃ may be dimeric in the solid state as evidenced by its deep violet color, though it is monomeric and yellow in benzene.

Characteristically, complexes containing alkyl groups other than methyl are markedly less stable. The TiRCl3 species have been prepared where $R = e^{\frac{1}{2}(149.151)}$ propyl,¹⁵¹ isobutyl,¹⁴⁹ and pentyl¹⁵¹ employing alkylaluminum or zinc reagents but are stable only at low temperatures. Only blue to blue-violet bipyridyl adducts, TiRCl₃(bpy), were isolated in low yield.¹⁵¹ Yet Bawn and Gladstone report¹⁵⁵ that TiEtBr₃ (and TiMeCI₃) can be prepared from PbR₄ and TiX₄ (R = Me or Et, $X = C1$ or Br) in heptane or toluene. Pure TiEtCl₃, a violet solid which melts to a red liquid at room temperature, was obtained in 40-50% yield by vacuum distillation. In the presence of lead compounds it decomposes readily to C_2H_6 , C_2H_5Cl , and Ti(IV) species. The zinc route also gives TiEtCl₃, as well as TiPrCI₃ and Ti(n -C₅H₁₁)CI₃.¹⁵²

Phenyltitanium halides are prepared by arylation with diphenylzinc: 151,152

$$
TIX4 + 0.5ZnPh2 \xrightarrow{\text{120} \atop \text{pentane}} TIPhX3 + 0.5ZnX2
$$

(X = CI, Br)

The reaction between $TiCl₄$ and LiPh in toluene at -50° also gives TiPhCI₃, but separation of lithium chloride is troublesome. Red-green dichroic TiPhCl₃ has properties analogous to TiMeCl₃ but is less thermally stable, suggesting that aryl complexes are not necessarily more stable than the corresponding alkyls, especially if the alkyl has no β hydrogen.

Several zirconium(IV) species can be prepared employing zinc reagents. $152,156$ in contrast to the titanium system, zinc dialkyls ($R = Me$, Et, Pr) do not react with ZrX_4 ($X = Cl$, Br) in hexane. In ether they do, but only to give nearly insoluble **sol**ids containing zinc. However, in toluene at -10 to 0° brickred solutions of $ZrMeX_3$ (X = CI, Br) result. Addition of diethyl

ether gives stable, isolable diethyl etherates, $ZrMeX₃(ether)₂$. In pyridine at **0'** alkylation proceeds further to give brown solutions which probably contain the adducts, $ZnR_2X_2(py)_2$. These could not be isolated but addition of bipyridyl gave isolable $ZrR_2X_2(bpy)$. In no solvent did ZrX_4 react with SnR₄, consistent with the finding that $TiMeCl₃$ and $SnCl₄$ give $TiCl₄$ and 0.25 SnMe₄. Though $ZrR_2X_2(bpy)$, which is soluble in tetrahydrofuran and pyridine, is believed to be monomeric, the authors suggested $ZrRX_3$ (ether)₂ is not, since it is only slightly soluble in ether. The authors also argue that ligand free alkyl complexes are more likely to be dimeric for zirconium than titanium.

What is believed to be $ZrPhCl₃(THF)₃$ has been prepared from PhMgCl and $ZrCl₄$ in toluene-THF.¹⁵⁷ Several adducts were isolated but extreme sensitivity to water and oxygen and low solubility made characterization difficult.

Titanium and zirconium benzyl complexes have been prepared⁶⁵ from M(CH₂Ph)₄ in toluene at -10° (eq 3). All are

solubility made characterization difficult.
\n
$$
um
$$
 and zirconium benzyl complexes have been pre-
\nfrom M(CH₂Ph)₄ in tolerance at -10° (eq 3). All are
\nM(CH₂Ph)₄ + HX \longrightarrow M(CH₂Ph)₃X
\nMX₄
\n WX_4
\n $WCH_2Ph)_{2}X_2$ Ti(CH₂Ph)₃F
\nM = Ti; X = Cl, Br, I
\n $M = Zr; X = Cl$

rather unstable in solution and monomeric in benzene with the exception of $Ti(CH_2Ph)_3F$ whose apparent molecular weight is concentration dependent.

Partial alkylation of Ti(lll) has been less successful. Treatment of TiCl₃(THF)₃ with 2 mol of LiPh in ether at -50° gives red-violet TiPh₂Cl(ether)₃ solutions.²⁷ Red crystals were isolated at temperatures below **-50'.** In pyridine an essentially identical procedure gives isolable, deep blue $TiRCI₂(py)₃$ species $(R = Me$ or Ph) which are "fairly stable" at room temperature under nitrogen.¹⁵⁸ Kühlein and Clauss¹⁵⁹ prepared TiMeCl₂ at -78° in aliphatic hydrocarbons by a less direct approach:

 2 TiMeCl₃ + Hg(SiMe₃)₂ \rightarrow Hg + 2TiMeCl₂ + 2SiMe₃Cl

Isolated adducts are listed in Table II. Six-coordinate species are by far the most common. Five-coordinate species probably form in solution with some monodentate ligands, but isolation is rarely possible. At low temperatures the ¹H NMR spectra of TiMeCl₃(L-L') (L-L' = a chelating ligand) suggest a static mer configuration;¹⁶⁰ if the two ends of L-L' are different, the preferred donor trans to Me is $0 >$ $N > S$.¹⁶¹ in general the finding that TiR₄L₂ stability varies according to how firmly L is bound also holds true for $MR_nX_{4-n}L_2$ (M = Ti or Zr).

B. Group 5

The best known alkyl complexes of the vanadium triad are niobium and tantalum methyl complexes. In 1964 Juvinall¹⁶³ reported the low-yield preparation of $NbMe₃Cl₂$ and $TaMe₃Cl₂$ by alkylation with dimethylzinc in pentane. The volatile yellow complexes were trapped at -36° during solvent removal. Actually the reactions are quite clean and the products easily isolable.^{164,165} Two are obtained virtually pure in high yield: NbCl₅ + ZnMe₂ $\frac{1}{2}$ and $\frac{1}{2$

$$
1^{164,165} \text{Two are obtained virtually pure in high-yie-164.165} \text{Two are obtained virtually pure in high-yie-164.165} \times 2 \times 24 \text{ h}
$$
\n
$$
1^{164,165} \text{Two are obtained virtually pure in high-yie-164.165} \times 2 \times 1^{164,165} \text{Two are obtained virtually pure in high-yie-164.166} \times 2 \times 1^{164,165} \text{Two are obtained with the initial point is 1.52 nC12 and 1.52 nC12 is 1.52 nC12 is 1.52 nC12 and 1.52 nC1
$$

Traces of NbMeCl₄ or TaMe₂Cl₃ respectively can be converted to the more highly methylated complexes by addition of

TABLE **II.** Halide-Alkyl Adducts of the Group 4 Metals

M	R	X	L_n	Ref
Τi	Me	Cl ₃	2CH ₃ CN, 2py, 2THF, 2Me ₂ S, 2THF, 2(1,4-thioxane)	162
			dme, 2,5-dithiahexane, dppe	160, 162
			bpy	151, 162
			$\mathsf{TMEDA}, o\text{-}(Me, N), C_{\kappa}H_{4}, o\text{-}C_{\kappa}H_{4}(OCH_{3})(NMe_{2})$	160
			CH ₃ OCH ₃ CH ₃ NMe ₂ , CH ₃ OCH ₃ CH ₂ SMe ₁ $CH3SCH2CH2NMe2$, $o-C6H4(CH2NMe2)(NMe2)$	161a
			THF, α -picoline, 1.4-dithiane	162
			Dioxane, PPh,	149, 162
			CH, OCH, CH, CH=CH,	161 _b
Τi	Et, Pr, pentyl	Cl ₃	bpy	151
Ti	Ph	CI ₃	2ether, 2py, bpy	151
Zr	Me,	CI, Br,	bpy	156
Zr	Et,	Cl ₂ , Br ₂	bpy	156
Zr	Me	$CI3$, Br ₃	2ether	156

TABLE **111.** Physical Characteristics and Spectroscopic Data for MMenCI,-, (M = Nb or Ta, *n* = **1, 2, 3)**

 a RT = room temperature. b in benzene, c Some slight shifts occur upon lowering the temperature. All Nb species exhibit broad peaks at room temperature which sharpen on cooling. This behavior is more likely due to

the appropriate quantity of $ZnMe₂$ while $NbMe₃Cl₂$ may similarly be prepared quantitatively from NbMe₂Cl₃ (ref 95). Alkylation does not proceed beyond the trimethyl stage with $ZnMe₂$ though methyl group exchange between NbMe₃Cl₂ and $ZnMe₂$ in CCI₄ is rapid on the NMR time scale.¹⁶³ Monomethyl species form in about **95%** purity in situ in aromatic hydrocarbons from either the di- or trimethyl species and the appropriate quantity of MCI_5 . Since MCI_5 is nearly insoluble in pentane, MMeCl₄ cannot form completely.^{164,165} Both NbMeCI4 and TaMeCI4 recently have been prepared and isolated from MCl₅ and 0.5HgMe₂ or 1.0SnMe₄ in CH₂Cl₂ at -35° (ref 166). Similar reactions with NbBr₅ were slower, required more alkylating agent, and gave markedly less stable NbMeBr₄ with difficulty. [Similarly, TaMe₃Br₂ (from TaBr₅ and $ZnMe₂$) decomposes slowly in the solid state at -35° to yield explosive, shock- or temperature-sensitive products.⁹⁵] Finally, $TaMe₂Cl₃$ has only been prepared admixed with Ta- $MeCl₄$ and/or TaMe₃ $Cl₂$. The compounds are listed in Table III (MR4CI species are unknown). The more volatile di- and trimethyl complexes are probably monomeric while the MMeCI4 species are nearly dimeric in benzene.

A large number of adducts have been prepared with group **5** and 6 donor ligand^."^-'^^ Three types have been isolated: dimers containing a bridging ligand, e.g., $(NbMeCl₄)₂(diox);$ normal six-coordinate species, e.g., NbMe₂Cl₃(PPh₃); and seven-coordinate species containing a chelating ligand, e.g., $TaMe₃Cl₂(bpy)$. The six-coordinate species are assumed octahedral. A crystal structure¹⁶⁸ of TaMe₃Cl₂(bpy) reveals a distorted capped trigonal prism with two of the methyl-metal

bond distances equal to 2.24 and 2.16 A. The third methyl and a chloride are disordered. Solution studies were hampered by redistribution reactions to give adducts of other MMe_nCl_{5-n} species and a tendency to dissociate a chloride ion as evidenced by conductivity measurements in dichloromethane. Attempted adduct formation with tertiary amines gave metal reduction (common for early transition metals) while oxygen was slowly abstracted from triphenylphosphine oxide, trimethylphosphine oxide, or $(Me_2N)_3P=O$.¹⁶⁶ Abstraction from the very stable P^V oxides illustrates niobium and tantalum's high oxygen affinity.

Several derivatives of TaMe₃CI₂ $[TaCp_2Me_3, ^{95}$ TaMe₃- $(BH_4)_2$ ⁹⁵ TaMe₃(C₈H₈) (see section V.D.), TaMe₃- $(BH_2(pz)_2)_2$, ^{169a} and TaMe₃(acac)₂^{169a}] have been prepared straightforwardly.

Vanadium methyl and ethyl complexes have been mentioned briefly. Reaction of BMe₃ with VCI₄ yields VMe₂CI₂ and VMeCl₃; with BEt₃, VEt₂Cl₂ results.¹⁷⁰ VR₂Cl₂ (R = Me, Et) may also be prepared employing $ZnR₂$ and several adducts are known.¹⁵² Surprisingly, bright violet VMe₂Cl₂ reportedly does not dissolve in aliphatic hydrocarbons and decomposes at 150- 170'.

The only other group 5 complexes are Ta(CH₂CMe₃)₃Cl₂,⁸⁴ $M(CH_2SiMe_3)_3Cl_2$ (M = Nb, ^{169b} Ta, ^{169c}), M(CH₂SiMe₃)₂Cl₃ (M = Nb or Ta),^{169b,c} and Ta(CH₂SiMe₃)Cl₄.^{169c} The first is formed from TaCl₅ and 3Me_3 CCH₂MgCI in ether⁸⁴ or 1.5Zn(CH₂CMe₃)₂ in pentane.⁹⁵ The latter were prepared in ether from MCI_5 and $Zn(CH_2SiMe_3)_2$. All are thermally stable and most sublime in vacuo without decomposition at ca. 100'. Several recent attempts to prepare Nb and Ta benzyl complexes were reportedly unsuccessful.^{72b}

C. Group 6

in group 6. Phenylchromium halides have received greatest attention

employing organoaluminum compounds in THF:17' A series of CrRCl₂(THF)₃ complexes has been prepared

Phenyichromium halides have received greatest attention
in group 6.
A series of CrRCl₂(THF)₃ complexes has been prepare
emplying organoaluminum compounds in THF:¹⁷¹
CrCl₃(THF)₃ + AIR₃ (or AIR₂OEt)
$$
\xrightarrow{\text{THF}}
$$
CrRCl₂(THF)₃ + traces of CrCl₂(THF)_{1 or 2}
R = Me. Et, Pr, or Bu^{*i*}

Yields can be high (79% for $R = Et$ using AIEt₂(OEt)). Thermal stabilities decrease in the expected order (Me $> Et$ > Pr $>$ Bu^{\prime}), decomposition giving alkanes, alkenes, and alkyl coupling products. Decomposition of CrMeCl₂(THF)₃ in CDCl₃ $(90^{\circ}, 1 \text{ h})$ gave only traces of CH₃D. No evidence for chromium-hydrido species was found. Pyridine, acetonitrile, and bipyridyl adducts were also isolated.

Several of these compounds had been prepared earlier by other methods, for example, $CrMeCl₂(THF)₃$ from $CrCl₃$ and MeMgCl in THF.¹⁷² A structure of one, Cr(p- $_{\text{tol}$)CI₂(THF)₃,^{173,174} reveals a *mer* geometry and an interesting Cr-O bond lengthening trans to the p -tolyl group. The

authors suggested that a similar "trans effect" might be an important factor in the conversion of triarylchromium(III) compounds to $bis(\pi\text{-}arene)$ chromium(0) or chromium(1) species; that is, loss of a trans ligand may be the initial step. Structural studies of $CrPh_2Cl(THF)_{3}$ (vide infra) and CrPh₃(THF)₃ would provide interesting comparisons.

A benzyl complex, $Cr(CH_2Ph)Cl_2(THF)_3$, was believed to form at -20° in tetrahydrofuran¹⁷⁵ but only bibenzyl and $CrCl₂(THF)₂$ could be found after warming to 20^o. Hydrolysis at low temperature gave $[Cr(CH_2Ph)(H_2O)_5]^2$ ⁺ (ref 176). A pyridine adduct was prepared by oxidation of CrCl₂(py)₂ with PhCH₂CI:^{176,177}

$$
2CrCl2(py)2 + PhCH2Cl \xrightarrow{py} CrCl3(py)3 + Cr(CH2Ph)Cl2py3
$$

Red-brown $Cr(CH_2Ph)Cl_2(py)_3$ decomposes at 40-60° in benzene or pyridine to give bibenzyl and Cr(ll) salts, but the solid may be stored for months under dry nitrogen at 5-10°. It reacts with perchloric acid in water to give $[Cr(CH₂Ph)(H₂O)₅]$ ²⁺ and with 1,3-cyclohexadiene to give benzene, toluene, and bibenzyl. Both Cr(CH₂Ph)₂CI(THF), and Cr(CH2Ph)a(THF), are unstable in tetrahydrofuran. **175**

Steric factors must be responsible for coordination of only two THF molecules in Cr(CHPh₂)Cl₂(THF)₂:50,178

ne, toluene, and bibenzyl. Both Cr(CH₂Ph)₂Cl(THF)_x
2Ph)₃(THF)_x are unstable in tetrahydrofuran.¹⁷⁵
ric factors must be responsible for coordination of
4F molecules in Cr(CHPh₂)Cl₂(THF)₂^{50,178}
CrCl₃ + Ph₂CHLi(diox)₂
$$
\xrightarrow{\text{THF}}
$$
 Cr(CHPh₂)Cl₂(THF)₂

The bright blue complex has a stability comparable to CrMeCI₂(THF)₃. Decomposition in THF-hexane, benzene, or ether gives CrCl₂(THF)₂ and Ph₂CHCHPh₂; CrCl₂(donor) forms in the presence of bpy, 2py, or dme (=donor). Complexes having only one halide, or none, could not be isolated.

Only one example of a CrR₂CI species is reported. A redistribution reaction between $[CrPh_6]^{3-}$ or $[CrPh_5]^{2-}$ and $CrCl₃(THF)₃$ in dme gives $CrPh₂Cl(dme)_{1.5}$.¹⁷⁹ This completes the Cr(III) series, CrPhCl₂ to $[CrPh_6]^{3-}$ (solvents omitted).

Alkylation of WCI₆ with a variety of alkylating agents gives

green, thermally unstable WRCl₅.^{143,152,180a,181}
\nWCl₆ + 0.5MMe₂
$$
\xrightarrow{-35^{\circ}}
$$
 WMeCl₅ + 0.5MCl₂
\n[M = Zn (in ether) or Hg (in CH₂Cl₂)]
\nWCl₆ + 0.5ZnPh₂ $\xrightarrow{\text{other}}$ WPhCl₅ (incomplete)
\nWCl₆ + SnR₄ $\xrightarrow{\text{hexane}}$ WRCl₅ + SnR₃Cl
\n(R = Me, Et, Bu, or Ph)
\nWCl₆ + $\frac{1}{3}$ BR₃ $\xrightarrow{\text{other}}$ WRCl₅ + $\frac{1}{3}$ BCl₃
\n(R = Me, Et, Bu)

Traces of Zn, Sn, or B possibly affect stability; i.e., one report¹⁴³ claims WMeCl₅ decomposes at -15^o, another¹⁸¹ at 65° . Authors noted that (i) WCI₆ and ZnR₂ (R = Et or Bu) in ether at -60° give WCI₄ which may be isolated as WCI₄L₂ (L = THF or py); (ii) WCl₆ and SnR₄ do not react in ether; and (iii) WCl₆ and ZnR₂ do not react in hexane. WPhCl₅ is apparently not well characterized. A recent report^{180b} claims that diamagnetic WPhCl₃ (fine brown crystals) is the only pure product:

Figure 11.10.3 (The Brown of galaxies) is the only
\nuluct:

\n
$$
WCI_6 + 2SnPh_4 \xrightarrow{\text{refluxing}} WPhCl_3 + 2SnPh_3Cl + PhCl
$$
\npentane

Some adducts, WMeCl₅(py)₂, WMeCl₅(bpy), and WEtCl₅(bpy), appear to be eight-coordinate while conductivity measurements suggest WMeCI₅L₃ (L = NHEt₂ and py) is ionic, i.e., $[WMeCl₄]⁺Cl₋$.

Extension of these studies to molybdenum was unsuccessful.¹⁸² No Mo(V) species were isolated employing MoCl₅ and $ZnMe₂$, SnMe₄, or ZnEt₂ in hexane, ether, tetrahydrofuran, or pyridine. In ether, MoCl₄ and $0.5ZnMe₂$ gave MoMeCl₃(ether)₂ (insoluble, μ = 2.0) but in tetrahydrofuran or pyridine only Mod_{3L_3} (L = THF or py). Methane and methyl chloride were commonly observed products. Reaction of MoCl₅ and AIPh₃ gave similar results.¹⁸³

D. Anionic Complexes

Anionic (or cationic) complexes should be isolable. Simple

halide addition to niobium¹⁶⁵ and titanium^{184,185} gives anions:
\n
$$
NEt_4Cl + NbMe_2Cl_3 \xrightarrow{CH_3CH} [NEt_4]^+[NbMe_2Cl_4]^-
$$
\n
$$
NEt_4X + TiMeX_3 \xrightarrow{CH_2Cl_2} [NEt_4]^+[Ti_2Me_2X_7]^-(purple)
$$
\n
$$
[NEt_4]^+{}_2[Ti_2Me_2X_8]^2-(brown)
$$
\nor
$$
[NEt_4]^+{}_2[TiMeX_5]^2-(blue)
$$

The dimers are believed analogous to $[Ti_2Cl_9]^-$ and $[Ti₂Cl₁₀]²$ which have (respectively) doubly and triply bridging chlorides. Interestingly, they are stable at room temperature only a short time while the mononuclear Ti anion is indefinitely stable. The latter is also stable toward oxygen. Reexamination of systems in which only neutral species were sought could be fruitful. For example, $VMe₂Cl₂$ and ZnMe₂ give Zn $\text{[VMe}_{4}\text{Cl}_{2}\text{]}$ -OEt₂ and LiMe and VCI₄ give Li₂[V- $Me₄Cl₂$]. ¹⁵²

IV. Cyclopentadienyl-Alkyl Complexes

It is now believed that cyclopentadienyl "stabilizes" the metal-alkyl bond simply by occupying coordination sites which would otherwise be utilized in decomposition just as amines and phosphines promote stability by adduct formation. Readily available early metal cyclopentadienyl complexes have long been used to prepare many exemplary alkyl complexes.

A. Groups 4 and 5

Two groups prepared TiCp₂R₂ (R = Me, Ph, m-tolyl, and p-dimethylaminophenyl) approximately 20 years ago.¹⁸⁶⁻¹⁸⁸ Other members of this class are $ZrCp_2Me_2$, 189a, 195c $HfCp_2Me_2$,^{195c} MCp₂Ph₂ (M = Zr or Hf),^{195c} MC_{P2}(CH₂Ph)₂ (M $=$ Ti, Zr),¹⁹⁰⁻¹⁹² and MCp₂(CH₂SiMe₃)₂ (M = Ti, Zr, Hf).^{86,87,89,193,194} The related π -indenyl species, M(lnd)₂R₂ (M $=$ Ti, Zr, Hf, R = Me; M = Ti, Zr, R = Ph) have also been prepared.^{195c} Several in which R_2 is a chelate are also known:

All were prepared from MCD_2Cl_2 and a lithium or Grignard reagent. Photolysis of MCp₂Me₂ (M = Ti, Zr, Hf) in the presence of PhC=CPh also reportedly gives MCp₂(C₄Ph₄).^{195b} Thermal stabilities are variable, approximately $R = Me <$ CH_2Ph < Ph < $CH_2SiMe₃$ < chelates and Ti < Zr. For example, orange-yellow TiCp₂Me₂ blackens in the solid state at 40' and decomposes at ca. **0'** in solution, while colorless $ZrCp_2Me_2$ sublimes with little decomposition at 100°. An exception is $TiCp₂(C₄H₈)$ which must be prepared and purified below -30° . In this case a carbon-carbon bond can cleave to yield TiCp₂(C₂H₄)₂, but β -hydrogen abstraction probably predominates. Interestingly, $Ti(n^5-C_5Me_5)$ ₂Me₂ is relatively stable,¹⁹⁸ suggesting that a bimolecular reaction or one involving hydrogen abstraction from the π -C₅H₅ ring may be one pathway by which MCp₂R₂ species decompose. Decomposition of several $TiCp_2R_2$ species $(R = \text{aryl or benzyl})$ has been studied recently.^{199a} The structure of TiCp₂Ph₂ is pseudo-tetrahedral with a mean Ti-C bond length of 2.272 **A, 199b**

Several MCp₂RX (M = Ti or Zr, X = a halide) complexes are also known. Yellow ZrCp2MeCl was prepared from $(ZrCp₂CI)₂O$ and trimethylaluminum²⁰⁰ and in the following manner: 189a

 $4ZrCp_2Me_2 + 2PbCl_2 \rightarrow 4ZrCp_2Me(Cl) + Pb + PbMe_4$

The corresponding ethyl complex, ZrCp₂(Et)Cl, is reportedly stable only at 0° (ref 201). Those with $R =$ cyclohexyl, 4-

$$
ZrCp_2Cl_2 \xrightarrow{AIE1_3} ZrCp_2(Et)Cl (27\%)
$$

vinylbiphenyl, and 2-methyl-2-butyl are also only marginally stable;^{189b} the carbon to which the metal is attached in the last two is uncertain.

Addition of 1 mol of RMgX to TiCp₂Cl₂ (R = Me, Et, Pr, Bu, Bu', n-C₅H₁₁, CH₂CMe₃, Ph, CH₂Ph, and CH₂CH₂Ph,²⁰² and $CH₂SiMe₃⁸⁷$) gives the corresponding TiCp₂RCI complexes. Their thermal stability appears greater than that for MCD_2R_2 . Interestingly, TiCp₂RCI apparently ionizes, even in nonpolar solvents, if R is CH₂CMe₃, Ph, or CH₂Ph, to give [TiCp₂S, R]⁺ $(S = solvent)$. In acetonitrile treatment with AgNO₃ gave AgCI, but no organometallic products could be isolated.

Compounds containing one cyclopentadienyl ring, TiCpMe₃, ^{194,203} TiCp(CH₂SiMe₃)₃, ^{194,204} and TiCpPh₃, ²⁰⁵ are known. As might be expected, they are somewhat less stable than the corresponding $TiCp₂R₂$ species. $TiCpPh₃$ decomposes readily in ether to give biphenyl and TiCpPh(ether)₂ and gives TiCpPh(NH₃)₂ with NH₃.

Titanium(III) compounds, $TiCp_2R$, are obtained from TiCp₂Cl and RMgX in ether $(X = CI \text{ or } BT; R = Ph, o, m, and)$ p -tolyl, 2,6-Me₂C₆H₃, 2,4,6-Me₃C₆H₂, or CH₂Ph).²⁰⁶ The green, monomeric complexes have a magnetic moment close to the spin-only value for one unpaired electron (1.58 for R = Ph and 1.66 for R = 2,6-xylyl). Their thermal stabilities vary in the order R = p-tolyl \sim m-tolyl \leq Ph \leq CH₂Ph \leq ties vary in the order R = p-tolyl \sim m-tolyl \lt Ph \lt CH₂Ph \lt o-tolyl \lt 2,6-xylyl \sim mesityl.²⁰⁷ Attempts to prepare alkyl derivatives with $R = Me$, Et, Prⁱ, or CMe₃ were unsuccessful though evidence for TiCp₂Me by electrolytic reduction^{208a} or on reaction of TiCp₂Cl with MeMgX^{208b} exists. TiCp₂-(cH2SiMe~)~~~ and TiCpz[CH(SiMe3)2] **lola** are also known. The aryl complexes react with molecular nitrogen to give deep blue diamagnetic $(TICp_2R)_2N_2$,²⁰⁹ but TiCp₂(CH₂SiMe₃) does not.

Only the analogous vanadium(III) complexes $(R =$ Ph, 210,211 Me, 212 or CH₂Ph²¹²) were isolated starting with $VCD₂Cl₂$ or $VCD₂Cl$. That steric factors do not allow $VCD₂R$ to form is suggested by the following: (1) VCD_2 (C=CPh)₂ can be prepared, presumably since the alkynyl ligand has minimal steric requirements; (2) the allyl ligand in $VCD₂(C₃H₅)$ is σ -bonded, not π -bonded as in the corresponding titanium and niobium complexes; and (3) the nitrogen ligand in $VCD₂(o Me₂NCH₂C₆H₄$) is not coordinated as in the corresponding Ti complex.¹⁴⁷ Interestingly, while VCp₂R (R = Me or CH₂Ph) reacts with CO to give VCp₂(CO)(COR), VCp₂Ph yields $VCD(C_5H_5Ph)(CO)_2^{213}$ (cf. Ph migration to C_8H_8 , section V.D, and Et to η^5 -C₅H₅²¹⁴).

ESR evidence suggests that addition of a large excess of RMgX (R = Et, Me, Prⁱ) to TiCp₂Cl gives $[TiCp_2R_2]^-$ (ref 208b). Surprisingly, not only $[TiCp_2Me_2]^-$, but $[TiCp_2Et_2]^$ was proposed to be stable for 1-2 h at room temperature in THF under the experimental conditions employed.

The only known TiCpRR' species are the following:^{79,215}
TiCp₂Ph₂ + LiCH₂Ph \rightarrow TiCpPh(CH₂Ph)

 $TiCp_2Ph_2 + LiCH_2Ph \rightarrow TiCpPh(CH_2Ph)$
 $TiCp_2(CH_2Ph)_2 + LiCH_2Ph \rightarrow TiCp(CH_2Ph)_2$

Addition of LiPh to TiCp₂Ph₂ reportedly¹⁸⁷ gives $[TiCp_2Ph_3]^{-}$, which decomposes 216 via intermediate TiCpPh₂.

Paramagnetic $NbCp_2R_2$ (R = Ph,^{217,218} Me,²¹⁹ and CH $_2$ SiMe $_3$ ⁸⁶) and TaCp $_2$ Me $_2$ ²¹⁹ are more stable than diamagnetic TiCp₂R₂, suggesting that an unpaired electron (if this were the only criterion) does not markedly decrease stability. At -45° NbCp₂Ph₂ reacts normally with HCI in ether to give $NbCp_2Cl_2$ and 2 mol of benzene, but, curiously, neither product is found when the reaction is carried out at room temperature.218

Finally, two rather unusual compounds with the $MCp₂$ basic unit are known. The first is $ZrCp₂(H)Me²²⁰$ which was prepared from $ZrCp_2$ MeCl and $LiAlH_4$ or $LiAl(OCMe_3)_3H$ in tetrahydrofuran. Its insolubility in common organic solvents

Figure 8. The structure of $Nb(\eta^5-C_5H_5)_2(C_2H_4)(C_2H_5)$.

and broad $\nu_{\text{M-H}}$ infrared band at low frequencies (1500, 1310; $\nu_{\text{M-D}}$ = 1090, 965) suggest a polymeric structure which presumably does not allow reductive elimination to give methane and ZrCp₂. The second is $NbCp_2Et(C_2H_4):^{221}$

$$
NbCp_2H_3 \xrightarrow[-1/2]{C_2H_4} NbCp_2(H)(C_2H_4) \xleftarrow[-1/2]{C_2H_4} NbCp_2Et(C_2H_4)
$$

The crystal structure²²² shows that the Et and C_2H_4 carbon atoms all lie in a plane which passes through Nb and bisects the Cp-Nb-Cp angle (Figure 8). On loss of ethylene β -hydrogen abstraction regenerates NbCp₂(H)(C₂H₄).

6. Groups 6 and 7

Many MCp(CO)₃R (M = Cr, Mo, or W), CrCp(NO)₂R, and MoCp₂(NO)R complexes have been prepared, most often by the salt method;²²³ for example

the salt method;²²³ for example
0.5[MoCp(CO)₃]
$$
_{2}
$$

Na⁺[MoCp(CO)₃]⁻ \rightarrow MoCp(CO)₃R

Exotic preparative schemes have also been used
MoCp(CO)₃H + CH₂N₂ - MoCp(CO)₃Me (r

$$
MoCp(CO)3H + CH2N2 \rightarrow MoCp(CO)3Me (ref 188)
$$

 $\text{MoCp(CO)}_2H + \text{CH}_2\text{N}_2 \rightarrow \text{MoCp(CO)}_3\text{Me}$ (ref 188)
[ReCp(CO)₂(NO)]⁺ + NaBH₄ - > ReCp(CO)(NO)Me (ref 224)

as well as straightforward methods

$$
CrCp(NO)2I + MeMgl \rightarrow CpCr(NO)2Me (ref 188)
$$

Variations in R and other ligands (for example, tripyrazolylborate²²⁵ for Cp or tertiary phosphines for CO) are many, and a full discussion of their preparation and chemistry is not possible in this article [see ref 226-228 and earlier reviews].

The system based on MCD_2Cl_2 (M = Mo or W) has yielded

many new species, the chemistry of which is developing rapidly:^{214,229-231}

\n
$$
MoCp_2Cl_2 + AICl_2Et \rightarrow MoCp_2ClEt
$$
\n
$$
MOp_2(C_2H_3R) \xrightarrow{RY} [MOp_2(C_2H_3R)R']^+
$$
\n
$$
(M = Mo, R = H, R' = Me; M = W, R = H, Me,
$$
\n
$$
(R' = Me \text{ or } CH_2Ph; X = \text{halide})
$$
\n
$$
[MOp_2(H)(C_2H_3R)]^+ \xrightarrow{PPh_3} [MOp_2(PPh_3)(CH_2CH_2R)]^+
$$

$$
(R = H, M = Mo; R = Me, M = W)
$$

\n
$$
WCp_2H_2 + C_6H_6 \xrightarrow{h\nu} WCp_2Ph(H)
$$

\n
$$
WCp_2H_2 + mesitylene \xrightarrow{h\nu} WCp_2(CH_2C_6H_3Me_2)_2
$$

\n
$$
[MoCp_2(C_2H_4)Me] \xrightarrow{PPhMe_2} [MoCp_2Me(PPhMe_2)]^+
$$

\n
$$
(olefin)Pl^+ = proice-almost-octorin]V, bovo=0, atvortive
$$

PPhMep

 $[MCp₂(olefin)R]$ ⁺ species almost certainly have a structure analogous to $NbCp_2Et(C_2H_4)$ (vide supra). The crystal structure^{229c} of WCp₂(CH₂-3,5-Me₂C₆H₃)₂ reveals a M-C bond length of 2.28 and 2.29 **A** and Cp-W-Cp angle of 75'. The authors propose that it and $WCD_2(CH_2\text{-}p\text{-}C_6H_4Me)_2$ form by insertion of W into a methyl C-H bond.

V. Other Alkyl Complexes

Since it is now believed that the transition metal to carbon bond is not inherently weak (see section VI.E), the role of other ligands in determining the stability and reactivity of a given complex will be examined more closely in the future. Several classes of complexes containing an alkyl or aryl ligand are discussed below according to the type of "ancillary" ligand.

A. Alkoxides

In 1952 Herman and Nelson reported²³²⁻²³⁵ the first stable complex containing a carbon-titanium bond, $TIPh(OPr')_3$ (eq. 4). It can be stored indefinitely in the dark at 10° but decom-

$$
Ti(OPri)4 + Liph/LiBr
$$

$$
TiPh(OPri)3-LiOPri-LiBr-ether
$$

$$
0.25TiCl4
$$

$$
0.25TiCl4 + LiCl + LiBr + ether
$$
 (4)

(53%)

poses rapidly at its melting point (88-90') to give biphenyl and, at ca. 200°, also a 1:l mixture of saturated and unsaturated C₄ hydrocarbons. The stability of TiPh(OPr \dot{v}_{3} and analogous compounds was proposed to be primarily a function of the substituent groups' electronegativity. A stability order for TiR_nX_{4-n} complexes was suggested: R = Bu < Me < ethynyl

< p-anisyl < Ph < α -naphthyl < indenyl; X = OPr' \sim OPr \sim \leq p-anisyl \leq Ph \leq α -naphthyl \leq indenyl; $X =$ OPr' \sim OPr \sim OBu \sim OBu' $>$ OMe \sim Cl $>$ F; and $n = 1 > 2 > 3 > 4$. The structure of TiPh(OPr \dot{v}_{3} is unknown, but a dimeric formulation with two bridging alkoxide groups is plausible.

The analogous methyl complex, a distillable **(50',** 0.01 mm), viscous, yellow liquid is prepared from Ti(OPr \dot{O}_3 Cl and LiMe in ether at -50° (ref 236, 237). Nonvolatile TiMe(OEt)₃ was prepared similarly. Addition of a second LiMe to TiMe(0- Pr')₃ gives TiMe₂(OPr['])₂. Both the mono- and dimethyl compounds undergo redistribution reactions with TiMeCl₃:

2Time(OR)₃ + TimeCl₃
$$
\xrightarrow{CH_2Cl_2}
$$
 3Time(OR)₂Cl
85%
Time(OR)₃ + 2TimeCl₃ \longrightarrow 3Time(OR)Cl₂
80%
(also TiCl₄ + Time₂(OR)₂) (R = Pr⁷ or Et)

All may not be monomeric; for example, TiMe(OPr¹)₂CI is soluble in pentane, melts at 62-64', and sublimes in vacuo, while TiMe(OEt)Cl₂ is only slightly soluble in toluene. The apparent molecular weight (362) of TiMe(OPr \dot{v}_{3} in benzene indicates some tendency to dimerize.

One member of the TiR_n(OR')_{4-n} family, Ti(CH₂Ph)₂(OEt)₂,

which is known to be dimeric in benzene,⁶⁵ decomposes least rapidly of the series $Ti(CH_2Ph)_3F$ > $Ti(CH_2Ph)_3Cl$ > $Ti(CH_2Ph)_2Br_2$ > $Ti(CH_2Ph)_3Br$ > $Ti(CH_2Ph)_4$ > $Ti(CH_2-P)_2Br_2$ $Ph_2(OEt)_2$. Generalization is difficult since Ti $(CH_2Ph)_3F$ is also strongly associated in benzene. An attempt to correlate structure with stability would be valuable but appears premature.

A titanium complex containing the chelating 2-methyl-2,4-

pentanediolate ligand can be isolated in good yield:²³⁸

\n
$$
TiCl_{2}(C_{6}H_{12}O_{2}) + 2MeMgl/ether \xrightarrow{TID^{o}} TiMe_{2}(C_{6}H_{12}O_{2})
$$

A dimeric structure was suggested by molecular weight and confirmed by x-ray studies.²³⁹ Each titanium has a trigonalbipyramidal geometry with two methyl groups located in the trigonal plane (Figure 9). The Ti-O(bridging) bond lengths are longer (1.97-2.06 **A)** than the Ti-O(termina1) (1.77 and 1.78 **A),** while the Ti-carbon distances vary from 2.11 to 2.19 **A.** The compound decomposes slowly at room temperature in the solid state or in solution.

In addition to the direct method, alkoxides, in principle, may be prepared by reaction of metal alkyls with alcohol or

oxygen (see section VII.C). Ether cleavage is more unusual:³⁷

\n
$$
[ZrPh_2(\text{ether})]_2 \xrightarrow{\text{heat}} 2ZrPh(OEt) + C_6H_6 +
$$

\n
$$
0.5 \text{biphenyl} + 0.5C_2H_6 + 1.5C_2H_4
$$

Alkoxide ligands can take part in decomposition reactions, though most likely only under extreme conditions, for example, in the high temperature decomposition of TiPh(OPr¹)₃.²³²

B. Oxo

Allowing VOCl₃ to react with a deficiency of Grignard reagent or passing V(CH₂SiMe₃)₄ through a cellulose column gives lemon-yellow $V(O)(CH₂SiMe₃)₃$ which is relatively air stable, sublimes in vacuo (70°), and is light sensitive.⁸⁸ It does not react with water, alcohols, carbon monoxide, tertiary phosphines, or primary amines under mild conditions.

The reaction of V(O)Cl₃ with HgPh₂^{240,241} or ZnPh₂,²⁴⁰ but not SnPh₄, in aliphatic hydrocarbons at -25° gives red, crystalline V(O)PhCI₂. It is stable in hydrocarbons at \leq -25° but at ca. -10° gives biphenyl, chlorobenzene, and a trace of benzene in what is believed to be a concerted, *nonhomolytic* decomposition²⁴¹ (eq 5). With Hg(C₆H₅D)₂ (92.8% monodeut-
2V(O)PhCl₂ ------> V(O)Ph₂Cl + V(O)Cl₃

$$
2V(O)PhCl2 \longrightarrow V(O)Cl2 + V(O)Cl3
$$

\n
$$
Ph2 + V(O)Cl3
$$

\n
$$
Ph2 + V(O)Cl3
$$

erated), dilabeled biphenyl results. Oxygen "insertion" into the metal-phenyl bond may occur since hydrolysis of the product prepared with 0.5 mol of HgPh₂ gives a 75% yield of phenol. Surprisingly, electron donors *destabilize* V(O)PhCl₂; in ether it decomposes completely in minutes at -50° . However, isolation of $V(O)PhCl₂(bpy)$ was successful (dec 14°). Reaction of $V(O)Cl_3$ with SnMe₄ did not occur, while $ZnMe_2$ gave a product from which $ZnCl₂$ could not be removed.²⁴⁰

Reaction of $V(O)(OPr')_3$ with HgPh₂ failed to give $V(O)$ -Ph(OPr j_2 in either ether or benzene²⁴² but in pentane ZnMe₂ reacted smoothly:

accied smoothing:

\n
$$
V(O)(OPT')_3 + ZnMe_2 \xrightarrow{pentane} V(O)Me(OPr')_2 + ZnMe(OPr')
$$

The red-brown viscous product can be distilled in vacuo **(50',** \leq 2 mm) but decomposes rapidly at 70°. No adducts (e.g., with bipyridyl) could be isolated. The parent ion was the highest peak in the mass spectrum.

Methylmagnesium iodide reduces $NbOCI₃$ to $NbOCI₂$ in

Figure 9. Schematic drawing of $[\text{TIME}_2(\text{C}_6\text{H}_{12}\text{O}_2)]_2$.

ether but not in toluene-ether mixtures (ca. 2:1) at -15° .²⁴³ Only NbMeOCI₂ adducts, e.g., NbMeOCI₂(OPMe₃)₂, which was also prepared from NbMeCl₄ and 3OPMe₃, could be isolated. No reaction was observed between $NbOCl₃$ and $HgMe₂$ or SnMe4 under a variety of experimental conditions.

The tungsten complexes, $WCD_2(O)R_2 (R = Me, Et, Ph, or$ CH₂Ph), were prepared from WCp₂OCl₂ and RMgl in ether or $tetrahydrofuran.²⁴⁴$ They decompose on attempted sublimation at ca. 60'. The phenyl and benzyl compounds are moderately stable in air.

At -78° in CH₂Cl₂, W(O)Cl₄ and HgMe₂ give W(O)Me- $Cl₃$.¹⁸¹ It is markedly more unstable than WMeCl₅ and decomposes at ca. -10° . The HMPA adduct, W(O)MeCl₃(HM-PA), was also obtained from reaction of WMeCl₅ with HMPA.

Finally, $Re(O)(PPh₃)₂Cl₃$ or $Re(O)Cl₄$ reacts with methyllithium to give thermally stable, red-purple $Re(O)Me₄$ (mp 44°).245 Its mass spectrum shows the parent ion and its infrared spectrum the characteristic strong Re-O stretch at 1016 cm⁻¹. Its ESR spectrum at -175° suggests a tetragonal-pyramidal structure. It is thermally stable in the vapor state to ca. 150 $^{\circ}$ and does not react with water, alcohols, H₂, CO, or SO_2 under mild conditions. With I_2 it gives CH₃I quantitatively and both *02* and NO react rapidly to give as yet uncharacterized products.

C. Dialkylamido

A series of complexes, $Ti(NR_2)_3(R')$ (R = Me, R' = Me, CD₃, Et; R = Et, R^{\prime} = Me, CD₃, Pr, Pr^{\prime}, Bu, CMe₃ and Ph), were prepared from Ti(NR₂)₃X and R'MgX or LiR' (X = a halogen).²⁴⁶ They all melt below room temperature, sublime easily in vacuo, and decompose around 70'. **All** were characterized unambiguously by ¹H NMR. Note that R' can be isopropyl or even tert-butyl, ligands which are normally susceptible to β -hydrogen abstraction. The bulky NR₂ ligands apparently discourage both decomposition and adduct formation with typical donor ligands. Decomposition²⁴⁷ gives methane and diamagnetic Ti(IV) residues:

$$
Ti[NMe2]3(CD3) \rightarrow
$$

30% CH2D2, 30% CHD3, 30% CD4 4- Ti(lV)

Ti $[NEt₂]$ ₃(CD₃) \rightarrow

50% CH₂D₂, 20% CH₃D, 20% CHD₃ + Ti(IV)

the authors proposed abstraction from the dialkylamido group by an "ionic" rather than radical mechanism.

D. Cyclooctatetraene

Complexes of the type $Zr(C_8H_8)R_2$ (R = Me, Et) have been prepared from $Zr(C_8H_8)Cl_2$.²⁴⁸ The dimethyl derivative forms a yellow dietherate in which the ether may be bound to the metal. If C_8H_8 is planar, the adducts are monomeric, 18-electron species. The methyl compound decomposes to give methane and the ethyl to give 67% C₂H₆, 14% C₂H₄, and 10% of a mixture of butane and butenes (455.5 ratio) in 4 h at 60'.

Niobium and tantalum complexes containing both cyclooctatetraene and alkyls are also known:249

$$
MCI_{5} + \frac{1.2[K^{+}]_{2}[C_{8}H_{8}]^{2-\text{ volume}}}{2. \text{LiR}} M(C_{8}H_{8})_{2}R
$$

\n
$$
(M = Nb \text{ or } Ta, R = Me, \text{ or } Ph)
$$

\n
$$
MRCl_{4} + 2[K^{+}]_{2}[C_{8}H_{8}]^{2-\frac{\text{toluene}}{-78^{\circ}}} M(C_{8}H_{8})_{2}R
$$

\n
$$
(M = Nb, R = Me; M = Ta, R = Me \text{ or } Ph)
$$

\n
$$
TaMe_{3}Cl_{2} + [K^{+}]_{2}[C_{8}H_{8}]^{2-\frac{\text{toluene}}{-78^{\circ}}} Ta(C_{8}H_{8})Me_{3}
$$

The red-brown, crystalline $M(C_8H_8)_2R$ complexes are moderately soluble in tetrahydrofuran and dichloromethane and do not sublime without decomposition while $Ta(C_8H_8)Me_3$ is blue, sublimes nicely at 100 $^{\circ}$ and 1 μ , and is soluble in aromatic hydrocarbons. In $M(C_8H_8)_2R$ one C_8H_8 ring is believed planar (η^8) , the second diene-like (η^4) . Monomeric $Ta(C_8H_8)Me_3$ most likely has a planar C_8H_8 ring with three methyl groups disposed in a tripod-like fashion. When $M(C_8H_8)_2$ Ph reacts with dmpe or diars, the Ph ligand migrates to a C_8H_8 ring to give complexes containing a 2,3,4,5,6- η -en $do-8$ -phenylbicyclo $[5.1.0]$ octadienyl ligand, M(C₈H₈)- $(C_8H_8Ph)(L-L)$ (L-L = dmpe or diars).

E. Aquo

Benzyl chloride^{250,251} or radicals generated from $PhCH_2CMe_2OOH^{252}$ oxidize $Cr(ClO_4)_2$ to give $[Cr(H_2O)_5$ - $(CH_2Ph)^{2+}$ in aqueous solution. The former, in which benzyl radicals are postulated, resembles formation of $[Co(CH₂Ph)(CN)₅]$ ³⁻ from benzyl bromide and $[Co(CN)₅]$ ³⁻ (ref **253).**

$$
PhCH2Cl + Cr2+ \rightarrow CrCl2+ + PhCh2.
$$

$$
PhCH2· + Cr2+ \rightarrow Cr(CH2Ph)2+
$$

The analogous methyl complex was first postulated as a minor product of Cr^{2+} oxidation with Me₃COOH^{254,255} but later was similarly prepared and isolated in perchloric acid by ion-exchange chromatography. 256 it may also be obtained by methyl group transfer from an organocobaloxime²⁵⁷ or cor-

$$
10^{258}
$$
 complex to aqueous Cr²⁺:
\n
$$
C_0R(Hdmg)_2(H_2O) + Cr^{2+} + 2H^+ \rightarrow
$$
\n
$$
C_0^{2+} + [Cr(H_2O)_5(R)]^{2+} + 2H_2dmg
$$

(dmg = dimethylglyoxime)

It is relatively stable $(t_{1/2}$ = several hours at 0°) in perchloric acid as are its Et and Pr analogs.

The striking fact, of course, is that the metal-carbon bond survives in an aquo complex and is cleaved only relatively slowly even in aqueous acid²⁵⁹ (the rate is dependent on $[H^+]$). The fact that it is dicationic and/or that Cr(III) substitution processes are slow must account for its stability in aqueous acid.

F. Carbonyl

One of the few group 7 alkyl species, $M(CO)_{5}R$ $[M = Mn,$ Re], is prepared by alkylation of $[M(CO)_5]^-$ (ref 223) or by decarbonylation of RCOM(CO)₅.²⁶⁰ Like most "18-electron" monoalkyl complexes containing relatively nonlabile ligands, B

R can vary widely. They are probably best classed as carbonyls and will not be discussed here (see ref 226, 227, 261).

VI. Decomposition

Transition-metal alkyl decomposition modes have been discussed frequently and studied specifically for metals outside groups 4-7 (see ref 276). However, no unified theory of decomposition, or even a clear understanding of specific cases (with one or two exceptions), has resulted. Wilkinson,⁸³ Baird,²⁷⁶ Braterman and Cross,²⁷⁷⁻²⁷⁹ and Mingos,²⁸⁰ among others, discuss decomposition broadly. We intend to present what we feel are the most pertinent results which support postulated decomposition pathways. Much mechanistic information derives from studies of platinum alkyls, which are experimentally convenient because of stability to air, but the results probably apply equally well to early transition metal alkyls. Ti,1920262-268 **V** 269.270 Cr,270-275 and Mn,32 in addition to

A. &Hydrogen Abstraction

Recognition that many transition metal alkyl derivatives decompose by abstraction of a β -hydrogen atom has had substantial practical consequences. The number of isolable alkyl compounds is growing rapidly through use of alkyls lacking a β -hydrogen as described in preceding sections. The very success of this approach carries with it the danger that β abstraction will be overemphasized.

 β -Hydrogen abstraction or elimination can describe more than one reaction, the overall results of which may be identical. The best documented and the one usually meant by " β hydrogen elimination" is intramolecular β -hydrogen abstraction by the metal:

H I CHR M-CH2-CH,R - M-11 - "decomposition" CH2

The best-studied example is $Pt(PPh₃)₂(n$

the best-studied example is
$$
Pt(PPh3)2(n-C4H9)2
$$
²⁸¹
\n $Ph3P$
\n $Pt4H9$
\n $Pt(PPh3)2 + n-C4H10 + 1-C4H8$
\n $Ph3P$

The following are pertinent:

(1) Decomposition is a first-order intramolecular process.

(2) Addition of PPh₃ decreases the rate; presumably PPh₃ dissociation is inhibited.

(3) Deuterium in the 1, 1-C4D2H7 derivative is extensively scrambled in the 1-butene product.

(4) Added 1-butene does not exchange during the decomposition.

An important point is that β -hydrogen abstraction from polyalkyl complexes can give equal amounts of alkane and al-

kene, a phenomenon once thought characteristic of homolytic M-C bond cleavage. Butene presumably arises by β -abstraction to give A, from which it dissociates, while butane forms by reductive elimination (section V1.D) from the hydrido-butyl compound, B. Presumably these principles can be applied to many systems, though rarely can decomposition be ascribed to this specific mode, excluding all others.

In the early transition metals, a nice example of β -hydrogen abstraction is the thermolysis of $NbCp_2Et(C_2H_4)$ to give isolable NbCp₂(C₂H₄)(H).²²¹ With only a single alkyl group on the metal, alkane should not form unless an intermolecular reaction occurs:

$$
M-H + M-R \rightarrow 2M + RH
$$

A β -hydrogen may also be abstracted from an aryl ligand. In TiCp₂Ph₂ or its C₆D₅ analog either Ti or phenyl abstracts an ortho hydrogen to give benzene^{282a}

$$
Cp_2Ti(C_6D_5)_2 \xrightarrow{C_6H_6, 80^\circ} C_6D_6 (1 mol)
$$

Evidence for a phenylene complex, "Cp₂Ti(C₆H₄)", consists of in situ reactions with $CO₂$ or PhC=CPh (eq 6). Phenylene complexes have also been postulated in $Cr,13$ W, Nb, and Ta,^{132b} and Mo^{110b} systems.

As mentioned above, a β hydrogen can be abstracted directly by another alkyl:

$$
M \times_{CH_2CH_2R}^{CH_2CH_2H} \longrightarrow M \longrightarrow \underset{CHR}{CH_1} + CH_3CH_2R
$$

Of course, the overall result is identical with abstraction by the metal.

Finally, intermolecular abstractions of both types are possible (eq **7).** Again they cannot be distinguished from each other, but in theory can be distinguished from intramolecular abstraction since the formal oxidation state of the metal is reduced by only one.

B. a-Hydrogen Abstraction

Much less well recognized is a process in which an α -hydrogen of an alkyl ligand is abstracted to give an alkylidene, or carbene, complex. A substantial amount of information indicates this is also a significant decomposition mode.

As for β -abstraction, α -abstraction can occur in four ways. Intramolecular abstraction by the metal has been suggested by several authors.^{230,265,283-287} This process, M-CHR₂ \rightarrow $M(H)(CR₂)$, is more likely to occur in complexes with less than 18 valence electrons since the number must increase by two; the relationship to formal metal oxidation state is less clear. What is perhaps the first evidence of intermolecular α -hydrogen abstraction, photolysis of CD₃TiCl₃ in hydrocarbons to give CD4 and a series of perdeuterioalkanes up to C_9D_{20} , 153 could involve a transient CD_2 =M complex. However, photolysis of metal alkyls is, in general, even less well understood than thermolysis.

The only documented example is an apparent intramolecular α -abstraction by another alkyl⁹⁴ (eq 8 and 9). The neounderstood than thermolysis.
The only documented example is an apparent
lar α -abstraction by another alkyl⁹⁴ (eq 8 a
Ta(CH₂CMe₃)₃Cl₂ + 2LiCH₂CMe₃

$$
Ta(CH2CMe3)3Cl2 + 2LiCH2CMe3 \longrightarrow
$$

\n
$$
Ta(CH2CMe3)5 + 2LiCl (8)
$$

\n
$$
Ta(CH2CMe3)5 \longrightarrow (Me3CCH2)3Ta = C
$$

\n
$$
CH2CMe3 + CMe4
$$

\n
$$
CMe3 (9)
$$

pentylidene ligand probably forms by abstraction of a neopentyl α -proton by a neighboring neopentyl group in the sterically crowded penta(neopentyl) intermediate. An analogous α -abstraction may occur in the thermal decomposition of Ti(CH2Ph)4 in which 2.66 mol of toluene is generated per mole of complex,65 exactly the amount predicted by redistribution of all α hydrogens to give toluene and "Ph C(Ti)₃".

The dimeric species, ${M[CH_2SiMe_3]_2[CSiMe_3]}_2$ (M = Nb or Ta), $83,84$ possibly form from incipient M[CH₂SiMe₃]₃-[CHSiMe₃] by a second α -hydrogen abstraction from CHSiMe₃ by CH₂SiMe₃. Intermolecular reactions, possibly including the Grignard reagent, are also likely.84 In a related vein, toluene is produced on addition of AI(CH₂Ph)₃ to Ti(CH₂Ph)₄.⁶⁵ Interestingly, however, alkylidene α hydrogens are possibly more easily abstracted as suggested by the reaction of Ta(CH₂CMe₃)₃(CHCMe₃) with LiR-L_x to give Ta-**(CH2CMe3)3[C(CMe3)(Li*Lx)] .94396**

Carbenes may be formed in olefin disproportionation sysvia equilibrium 10 has been proposed.²⁸⁶⁻²⁸⁹

terms, possibly by
$$
\alpha
$$
-hydrogen abstraction. Disproportionation via equilibrium 10 has been proposed.²⁸⁶⁻²⁸⁹

\n
$$
M = CR'_{2}
$$

\n
$$
CR_{2} \longrightarrow M - \parallel C
$$

\n
$$
CR_{2} \longrightarrow M - \parallel C
$$

\n
$$
CR_{2}
$$

C. Ligand Hydrogen Abstraction

Several examples demonstrate that nonalkyl ligands donate hydrogen in a decomposition reaction in which alkane is produced. For example, $Ti(\eta^5-C_5Me_5)Me_2$ decomposes as shown in eq 11.¹⁹⁸ Apparently α -abstraction to give Ti(η^5 -

 C_5Me_5)₂(CH₂) and methane is unfavorable. Perhaps the cyclopentadienyl's methyl protons are more acidic than the metal's methyl protons. Similar examples are (i) H abstraction from η^5 -C₅H₅ in UCp₃R decomposition,²⁹⁰ (ii) abstraction from the ethyl group of a diethylamido ligand in Ti($NR₂$)₃R' species, 246 and (iii) abstraction of ortho hydrogen in a PPh₃ ligand in $M(PPh_3)_3Me$ (M = Rh²⁹¹ or Ir^{292}) to give $M(C_6H_4PPh_2)(PPh_3)_2$ and CH₄ (see also ref 16a).

D. Reductive Elimination (Coupling)

Conceptually one of the simplest decomposition modes of a polyalkyl complex is alkyl coupling. It is relatively uncommon but well established for the polymethyl derivatives, *fac-* $[PHMe₃L(L')₂]$ ⁺ (L = tertiary phosphine),^{293a} PdMe₂(P- $Et₃_{2}$,^{293b} and AuMe₃(PPh₃),²⁹⁴ which cleanly eliminate ethane. In the first, two methyl groups trans to L' couple more readily if L' has a high "trans-influence", and $L = L'$. In the second, evidence suggests the two methyl groups must be cis to couple.

Coupling to give biphenyl is common for polyphenyl derivatives like CrPh₃(S)_x (S = solvent). Much less commonly, polybenzyl compounds give bibenzyl. Evidence suggests that aryl coupling in early metal compounds is not necessarily analogous to the cis reductive elimination of methyl groups and may be much more complex.¹³

At some point during metal alkyl decomposition, an alkyl and hydride ligand are likely to be on the same metal. Their coupling is common, for example, as the final step in hydrogenation of activated olefins by dihydride complexes. A wellstudied example is hydrogenation of activated olefins by $MoCD₂H₂.²⁹⁵$ The intermediate hydrido-alkyl complex can sometimes be isolated and the reductive elimination reaction studied as a separate step (eq 12).

E. Homolytic M-C Bond Cleavage

Though metal alkyls like PbMe4 decompose at high temperatures to give free alkyl radicals,²⁹⁶ evidence that early transition metal alkyls decompose similarly at low temperatures is slight. Detection of coupling products is inadequate because they may arise in other ways. For example, decomposition of copper and silver propenyl complexes gives hexadienes with retention of configuration, suggesting the absence of free propenyl radicals.^{297,298a} Yet free radical decomposition may take place under certain circumstances as in chlorinated solvents (see, for example, ref 299a). Some Cu^{298b} and Ag alkyls also decompose by homolytic M-C bond cleavage, though probably not exclusively. The only satisfactory answer to this dilemma is to directly observe carbon based radicals or radical pairs (by ESR or CIDNP, respectively) on decomposition of a well-characterized metal alkyl. (The attempt to observe a benzyl radical on decomposition of $Zr(CH_2Ph)_4$ was unsuccessful.⁷³) Experiments with radical trapping agents^{299b} (see, for example, ref 299c) are less convincing since it is difficult to exclude the possibility that the trapping agent reacts with the metal complex and, in effect, initiates homolytic cleavage.^{299d,e}

The postulate that the dissociation energy of the transition metal to carbon bond is significantly lower than that of a main group element is unsupported by estimates obtained *so* far $[M-Me$ and M-Ph (M = Pt, Ti)^{83,276,279}]. Metal-carbon bond force constants in TiMeCl₃300 and TiMe₄301 do not differ significantly from those in analogous main group compounds. Metal-carbon bond length also does not correlate well with stability.^{140,226} A large number of early transition metal peralkyl complexes are now available and should permit resolution of this issue, for example, by calorimetric studies. Some of the first related experiments of this type have shown that the M-CH₃ dissociation energy in M(CH₃)(CO)₅ (M = Mn, Re) may, in fact, be on the order of that of M-C0.299f

VI/. Reactions of Alkyl Complexes

Many transition metal alkyl reactions are similar to those of main group metal alkyls with some exceptions such as reaction with N_2 or SO_2 . We survey briefly the common transition metal alkyl reactions^{2,6,11} and emphasize examples within the early metal category. Fundamental reactions of metal alkyls from a mechanistic viewpoint can also be found in a recent book by Heck.^{16b}

A. Electrophiles

Protonic reagents generally readily cleave the metal-alkyl bond to yield the alkane. In a few cases, e.g., $Cr(CMe₃)₄$, Ti(1adamantyl)₄, and $[Cr(H₂O)₅R]²⁺$, strong acids are required. In principle the proton may attack the metal, with RH formation by reductive elimination, or attack the alkyl carbon atom directly. A means of distinguishing between the two is not readily apparent. The former, in which the metal's *formal* oxidation state increases by two, does not seem possible for d^0 complexes such as WMe $_6$ or Ti(CH₂CMe₃)₄.

Lewis acids like BR₃, AIR₃, ZnR₂, or even MgX₂ (X = a halide) are often present after alkyl preparation or are added to "activate" the metal alkyl. Reactions between aluminum alkyls and transition metal halides have been extensively studied, yet most details remain obscure. A recent example is the reaction between $M(CH_2Ph)_4$ and $Al(CH_2Ph)_3$ (M = Zr, Ti).⁶⁵ When $M = Zr$ a red crystalline product results which may contain bridging CH₂Ph groups. When $M = Ti$, however, no stable product was isolated, and toluene formed with time. Compare these results with the following proposal: 302

$$
Cp_2TiCl_2 + 2AIMe_3 \xrightarrow{\text{115}^\circ} Cp_2Ti(CH_2AIMeCl)_2 + 2CH_4
$$

Since alkyls with few d electrons, like TiMe₄(d^0), are also good Lewis acids, similar reactions (intermolecular α -hydrogen abstraction) may initiate decomposition: 265

$$
2\text{Time}_4 \rightarrow \text{CH}_4 + \text{``Me}_3\text{Ti''} + \text{``Time}_3(\text{CH}_2)\text{''}
$$

Mechanistic details are not obvious.

B. **Hydrogen**

tion of hydrogen with a transition metal alkyl:
 $R_xM + H_2 \rightarrow R_xMH_2$ "Oxidative addition" is an attractive primary step for reac-

$$
R_xM + H_2 \rightarrow R_xMH_2
$$

Reducive elimination gives RH:

\n
$$
R_x M H_2 \rightarrow RH + R_{x-1} M H \rightarrow R_{x-2} M + RH
$$

Oxidative addition to d^0 complexes in this strict sense, however, seems impossible since higher oxidation states are unknown. Two reasonable alternatives are: (i) a catalytic amount of a lower valent metal complex forms by reductive elimination of R-R followed by addition of H_2 and redistribution, viz.

$$
R_2M \rightarrow R-R + M
$$

$$
M + H_2 \rightleftharpoons M H_2
$$

$$
R_2M + MH_2 \rightleftharpoons 2RMH
$$

$$
2RMH \rightarrow 2RH + M
$$

and (ii) hydrogen addition to M-R proceeds via a transition state in which only one end of the H₂ molecule is attached to the metal, viz.

Compare (ii) with the known reaction of LiPh with H_2 to give LiH and benzene. The reaction of hydrogen with the $Li₃MPh₆$ complexes almost certainly falls into this category. It is surprising in any case that the $M(CH_2Ph)_4$ complexes (M = Ti or Zr) "catalyze" olefin hydrogenation under conditions where they do not react with hydrogen.⁶⁵ However, since they also react with olefins, species which react more readily with hydrogen may form initially.

Hydrogen reactions are preparatively valuable in several instances. Hydrogen (1 atm) reacts with $TiCp₂Me₂$ in solution to give "titanocene", $Ti_2(C_{10}H_8)Cp_2H_2$.³⁰³ In contrast, ZrCp₂Me₂ gives a dimeric crimson species postulated to be $(ZrCp_2Me)_2$, ^{189a} while Ti(η^5 -C₅Me₅)₂Me₂ is stable toward H₂ under moderate conditions.¹⁹⁸ At 500 atm hydrogen reacts with MMe₅(dmpe) ($M = Nb$ or Ta) in THF in the presence of dmpe to give $MH₅(dmpe)₂.⁹⁵$

C. Oxygen

The reaction of oxygen with metal alkyls to give the corresponding alkoxides has been known for many years. Brindley and Hodgson³⁰⁴ recently studied its reaction with some selected metal alkyls, MR₄ (M = Ti or Zr, R = CH₂SiMe₃; M = Zr, $R = CH_2Ph$) and M_2R_6 (M = Mo, R = CH₂SiMe₃ or $CH₂CMe₃; M = W, R = CH₂Ph$. In hydrocarbons at 20 and -74' each rapidly absorbed 0.5 mol of *O2* per alkyl group. They postulated a radical chain mechanism propagated by alkylperoxy radicals:

cosulated a radical chain incoHamilton property, and is

\n
$$
R_nM + Q_2 \rightarrow [R_nM \cdot Q_2] \rightarrow R \cdot (initialion)
$$
\n
$$
R \cdot + Q_2 \xrightarrow{\text{fast}} RO_2 \cdot
$$
\n
$$
RO_2 \cdot + R_nM \rightarrow RO_2MR_{n-1} + R \cdot (propagation)
$$

Peroxy transition metal alkyls were difficult to detect since they were rapidly converted to alkoxides. Nevertheless at small amounts of peroxide. At -74° Zr(CH₂Ph)₄ and W_2 (CH₂Ph)₆ gave 0.6 and 0.8 mol of peroxide per mole of metal, respectively. On hydrolysis the latter gave benzyl hydroperoxide. Only traces of peroxides were observed for the CH₂SiMe₃ and CH₂CMe₃ species. Phenothiazine retarded oxidation, consistent with the proposed free radical mechanism. The authors concluded that the hindered approach of peroxy radicals to vacant metal orbitals significantly slowed the reaction rate. 20° $Mo_2[CH_2CMe_3]_6$, $W_2(CH_2Ph)_6$, and $Zr(CH_2Ph)_4$ gave

D. Halogens

carbon bond: Chlorine, bromine, and iodine commonly cleave the metal- $M-R + X_2 \rightarrow MX + RX$

$$
M - R + X_2 \rightarrow MX + RX
$$

In one case, 305 Mn(CO)₅R, the configuration at carbon is reportedly retained, suggesting an initial oxidative addition of Br₂ to the metal followed by reductive elimination of RBr.

In contrast, Br₂ may attack the α carbon of R in $[Cr(H₂O₅)₅(R)]²⁺$ directly to give a transition state of the "open" variety since kinetically stable $[Cr(H₂O)₅Br]²⁺$ was not observed.³⁰⁶ The bimolecular rate constants varied significantly with R, decreasing in the sequence Me $>$ Pr $>$ Et $>$ $CH₂CMe₃$. Similar kinetic studies of the reaction of $[Cr(p RC_6H_4)(H_2O)_5$ ²⁺ (R = Me, H, Br, CF₃, CN) with Br₂ and I_2 provided further support for the open transition state.³⁰⁷

Several authors ${}^{308-310}$ found predominant inversion of configuration at carbon in other systems. It is possible that at least two cleavage mechanisms operate, the relative amount of each depending on the metal, halogen, ligands, and experimental conditons.

The possibility of free radical pathways should not be ignored. Certain alkyl halides oxidatively "add" to $Ir(1)^{311}$ or **Pt(0)312** by a free radical mechanism. A similar cleavage mechanism (by X_2) could be proposed. Interestingly, iodine reportedly does not react with $M(CH_2SiMe_3)_4$ (M = Ti and Zr) in the absence of oxygen.304 Possibly a free radical pathway initiated by peroxyalkyl radicals predominates.

E. Carbon Dioxide and Carbon Disulfide

In the preparation of transition metal alkyls, $CO₂$ often is added at low temperatures to destroy excess Grignard or lithium reagent. A misconception that metal alkyls do not react with $CO₂$ has therefore arisen. While it is probably true that most transition metal alkyls do not react with CO₂ as readily as do Grignard reagents, several, especially the more highly alkylated ones, do react under mild conditions. For example, $CO₂$ reacts with M(CH₂Ph)₄ (M = Ti or Zr) at room temperature and atmospheric pressure $(Zr > Ti)^{.65}$ Hydrolysis yields phenylacetic acid and tribenzylcarbinol in approximately equal amounts. In a second example $CO₂$ apparently reacts with the decomposition product^{282b} (eq 13). Similar CS_2 "in-

sertions" into a metal-alkyl bond are known as well as a crystal structure of Re(C0)4(SSCPh) in which the dithio ligand is bidentate. 313 Recent articles $313,314$ provide further examples, details, and discussion of $CO₂$ and $CS₂$ reactions.

F. Carbon Monoxide

Migration of an alkyl ligand to a CO ligand to give the acyl with retention of configuration at the alkyl carbon^{310b} is now a well-known phenomenon, 260 migrations in M(CO)₅R, FeCpLL'R (e.g., $L = CO$, $L' = PPh_3$), and MCp(CO)₃R (M = Mo or W) being studied most thoroughly. Several early metal examples follow.

The reaction of CO with $TiCp_2(X)(R)$ (R = Me, CH_2Ph , X = CI; $R = Et$, $X = Ci$, I) probably proceeds via TiCp₂(X)(R)(CO) followed by migration of R to CO to give TiCp₂(X)(COR) ($v_{C=0}$) $1600-1625$ cm⁻¹).³¹⁵ Decarbonylation of TiCp₂-(CI)(COCH₂Ph) gave TiCp₂(CI)(CH₂Ph) essentially quantitatively at 40° in heptane in 0.5 h.

A similar reaction occurs between $NbCp_2Et(C_2H_4)$ and CO to give $NbCp_2(CO)(COEt)^{316}$ at high temperature and pressure, but TaCp₂Me(CO) is unchanged at 80° in benzene under 1 atm of CO after 1 day.95

Acyl-alkyl complexes appear unstable with respect to further alkyl attack on the acyl:

$$
Ticp_2(CH_2Ph)_2 \xrightarrow{CO_2} Ticp_2(CO)_2 + PhCH_2CCH_2Ph \t (ref 190)
$$
\n
$$
Ticp_2Ph_2 \xrightarrow{CO_2} Ticp_2(CO)_2 + PhCPh \t (ref 282c)
$$
\n
$$
Cp_2T_1 \xrightarrow{CO_2} Ticp_2(CO)_2 + \t D^2
$$
\n
$$
(ref 196)
$$

0 c_{p₂Ti(CI)(CCH₃) + MeMgCl $\frac{\text{co}}{\text{C}_{\text{MgCl}_2}}$ **C**₂} I1 [TaPhJ - - inter alia PhCCH,Ph + Ph,COH

$$
TiCP2(CO)2 + CH3CCH3 (ref 315)
$$

0

0

When more than two alkyl ligands are present, further reaction of the incipient ketone might be expected as in^{134}

$$
[\text{TaPh}_6] \text{-} \xrightarrow{CO} \text{hydrolysis} \text{inter alia PhCCH}_2\text{Ph} + \text{Ph}_3\text{COH}
$$

The coordinated or free diketone, PhCOCOPh, is apparently reduced on hydrolysis. Products before hydrolysis may be complex; e.g., $Ta(C_8H_8)Me_3$ and $TaMe_5(dmpe)$ react with excess CO to give products with no infrared absorption characteristic of $C=O$ or $C=O$.⁹⁵ Metal-oxygen bonds may be present as in the reaction³¹⁷ of Zr(CH₂Ph)₄ and CO where Zr-0-C-Zr bonds had to be postulated; the authors could find no evidence for ZrC=O or ZrCOR.

G. Nitric Oxide

An alkyl group also readily migrates to an NO ligand. NO (2 mol) reacts with $ZrCp_2Me_2$ or $ZrCp_2(Cl)(Me)$ to give monomeric species containing the N-methyl-N-nitrosohydroxylaminato ligand, $ZrCp_2[ON(Me)N=O]$ (CI or Me).^{189a} No further reaction occurred in several hours at higher temperatures. A similar but not monomeric product formed on reaction of 2 mol of NO with $TiCp₂Me₂$.

Hexamethyltungsten reacts with 4 mol of NO to give yellow WMe₄[ON(Me)N==0]₂ in essentially quantitative yield.²⁴ The crystal structure shows an eight-coordinate geometry somewhere between square-antiprismatic and dodecahedral³¹⁸ (Figure 10). The four oxygen atoms lie nearly in a plane while the four methyl groups do not; the CH₃'-W-CH₃' angle is 128.9', the CH3-W-CH3 angle 100.2'. Its **'H** NMR spectrum in CD_2Cl_2 at room temperature shows two sharp singlets in a 4:2 ratio at τ 8.42 and 5.97, respectively, suggesting nonrigid behavior. Below **-50'** the *7* 8.42 peak becomes two at *7* 9.65 and 7.59. The highest observable peak in the mass spectrum was WMe₃[ON(Me)N= O_2^+ .

NO (4 mol) reacts with MMe_xCl_{5-x} ($x = 2$ or 3, $M = Nb$ or Ta) to give seven-coordinate $MMe_{x-2}Cl_{5-x}[ON-$ (Me)N= O_2 .³¹⁹ The structure of the tantalum complex where $x = 3$ is a slightly distorted pentagonal bipyramid with the two chelating ligands and a methyl group in the pentagonal plane. The tantalum-carbon bond length, 2.248 **A,** is not unusual.

H. Olefins and Acetylenes

The reactions of chromium alkyl and aryl species, some prepared in situ, with olefins, $272,273,320$ dienes, $273,320,321$ diphenylacetylene,^{284,285,322-324} 2-butyne,³²⁵⁻³²⁹ and acetylene⁵⁴ have been studied extensively. Olefins and dienes are isomerized and hydrogenated. With acetylenes, cyclic and linear condensation products in addition to products resulting from hydrogenation and alkyl or aryl transfer are found; e.g., see eq 14.^{284c,323} Methyl complexes of Ti, Zr, V, Nb, Mo, and

Figure 10. Schematic drawing of WMe₄[ON(Me)N=0]₂.

Mn give similar results 322 while phenyl groups transfer from TiPh₄ to PhC= CPh^{330} or CrPh₃(THF)₃ to HC= CH^{54} In fact,

such findings are observed rather generally.^{328,331} The interesting feature in this context is that the products incorporate methylene (formally), which is one reason why authors have postulated chromium-methylene complexes.324 In one instance, $Fe(\eta^5-C_5H_5)(CO)_2R$, dimethyl acetylenedicarboxylate inserts into the Fe-R bond with retention of configuration at carbon.^{310b}

Whitesides and Ehmann³²⁸ studied the stoichiometric reaction of 2-butyne with Cr $Ph_3(THF)_3$, which yields, inter alia, **1,2,3,4-tetramethyInaphthalene.** They excluded a free or metal-complexed **tetramethylcyclobutadiene** intermediate based on

CrPh3(THF), + CD3CsCCH3 &:: *0 0*

Further labeling and kinetic isotope experiments and examination of relative yields of **1,2,3,4,5-pentamethyInaphthalene** and **1,2,3,4,6-pentamethyinaphthalene** employing CrR₃(THF)₃ $(R = \sigma, p$, or m-tolyl) "demonstrates that a long-lived arynechromium complex is also not an intermediate in these cyclizations". Numerous other results led to the proposal (solvents omitted) in eq 15. The "chromocycle" intermediate is

similar to the product of the reaction of $TiCp_2Ph_2$ with PhC $=$ CPh.^{282c} In the reaction of 2-butyne with Cr(o -anisyl)₃, a related chromocycle forms by cleavage of methoxy from the ring. $325,326$

Alkyl ligand transfer to an olefin (or olefin "insertion") is comparatively difficult to study since it yields another alkyl ligand which similarly reacts with more olefin. If the first transfer occurs, then subsequent transfers are also likely. Alkyl chain growth in this manner is usually postulated in olefin polymerization by a transition metal catalyst, but details are still unclear. For example, the olefin may not need to π -bond to the metal before the alkyl transfers. Group 4 metal catalysts are most active, usually only in the presence of a cocatalyst like an aluminum alkyl, the function of which is uncertain. Recently, however, ethylene has been polymerized with $M(CH_2Ph)_4$ (M = Ti, Zr) and CrRCl₂(THF)₃ (R = Me, Et, etc.) in the absence of cocatalysts 171 (see also ref 56b). Perhaps the alkyl-olefin complex most similar to a postulated intermediate in an active system is $NbCp_2Et(C_2H_4),^{221}$ but it only loses ethylene and gives $NbCp_2(H)(C_2H_4)$ on heating. Alkyl transfer to other unsaturated hydrocarbons like Cp (ref 213, 214) or $C_8H_8^{249}$ are probably poor models for the general reaction. Better models might be based on deVries' observation¹⁵³ that isobutylene reacts with TiMeCI₃ to give a product which on hydrolysis yields some neopentane. A second isobutylene "insertion" is probably sterically unfavorable.

Transfer of an aryl group to an olefin is basically different since an alkyl complex is formed which usually does not readily react with more olefin. In the case of Pd, ^{16b} rapid β hydrogen elimination gives a new olefin, the aryl-substituted form of the starting olefin.

Two groups report the use of tungsten alkyls as olefin disproportionation catalysts. Hexamethyltungsten on alumina converts propylene to a mixture of ethylene and bu t enes^{332,333} while tetrabenzyltungsten with AICI₃ in benzene converts 2-pentene into a mixture of 2-butene and 3-hexene.^{72a} In each case the alkyl complex almost certainly does not survive and the true catalyst is probably more closely related to group 6 disproportionation catalysts prepared in other ways (see, for example, ref 334).

1. Ketones

Since many main group alkyls like ZnEt₂, Hg(p -tolyl)₂, and $PbEt₄$ do not readily react with the carbonyl function like a Grignard reagent, a wide range of reactivities between transition metal alkyls and (e.g.) ketones should be expected. The Gilman test, 335 in which a metal alkyl adds to Michler's ketone, $(p-Me_2NC_6H_4)_2CO$, sometimes fails. For example, it is negative for TiMeCl₃¹⁸ and CrPh₂Cl.¹⁷⁹ It is positive for binary alkyls except where steric crowding prevents ready addition. Though other factors or side reactions may conceivably be involved, the test results should indicate whether a given alkyl or aryl complex will react with ketones in general. Qualitatively, one might expect that the more powerful the alkylating agent which is required to prepare a given alkyl, the more readily the alkyl will react with the carbonyl function.

A study of the reaction of methylniobium and methyltantalum chlorides, primarily NbMe₂Cl₃, with ketones and aldehydes has recently appeared.^{336a} Bulky ketones like Me-COCMe3 behave as donor ligands while a methyl group migrates to acetone or benzaldehyde to give an alkoxy ligand.

J. Other Reactions

A few other reactions should be mentioned.

Isocyanates "insert" into the metal-methyl bond in MMe_xCl_{5-x} (x = 1, 2, 3; M = Nb, Ta) to give acetamido complexes. $336b$ With isonitriles, complexes containing -C-(Me)= NR result. Each is a fairly general reaction. $337-339$ When $x = 2$ and R = Me or Ph in RNCO, only $MCI₃[N(R)C(O)Me]₂$ species could be isolated in pure form while both $MMeCl₂[N(R)C(O)Me]₂$ and $MCl₂[N(R)C(O)Me]₃$ result when $x = 3$. In all cases the possibility that the substrate inserted into the M-CI rather than M-Me bond could not be eliminated. Niobium complexes are considerably more unstable than those of tantalum and decompose near **25'.**

lsothiocyanates react similarly to give products of the type $MCI_4[N(R)C(S)Me]$, $TaMeCl_3[N(R)C(S)Me]$, and $NbCl_3$ - $[N(Me)C(S)Me]_2$ (R = Me or Ph; M = Nb or Ta).³⁴⁰ Thiocyanates, however, form simple adducts such as $MMe₂Cl₃(NCSMe)₂$. A crystal structure of NbCl₃[N(Me)- $C(S)$ Me]₂ shows two bidentate *N*-methylthioacetamide groups in the pentagonal plane of a pentagonal bipyramidal structure.

Carbodiimides, RN==C=NR (R = Prⁱ, Cy, or p-tolyl), react with MMe_xCl_{5-x} (M = Nb or Ta) to give complexes containing the RN- \dot{C} (Me)=NR ligand which, in at least one case, Ta- $MeCl₂[CyNC(Me)=-NCy]₂$, is bidentate^{336c} as shown by x-ray studies. The reactivity sequence follows the order $MMeCl₄$ $MMe₂Cl₃$ > $MMe₃Cl₂$, an order which holds also for the reactions discussed above.

Sulfur dioxide often reacts with group 4 metal alkyls to give 0-bonded sulfinates in contrast to the more common **S**sulfinates formed with group 8 metal alkyls.^{341,342} With TiCp₂R₂ and TiCp₂RX (R = alkyl or aryl; X = halide), monomeric $TiCp_2(O_2SMe)_2$, $TiCp_2(O_2SPh)_2$, and $TiCp_2Cl(O_2SMe)$ were formed.³⁴³ In the latter, ¹H NMR evidence suggested isomers based on the mode of $O₂$ SMe bonding. The reaction of SO₂ with ZrCp₂MeCl did not give the expected product but one with the stoichiometry of a bis- $SO₂$ adduct postulated to be $ZrCp(O_2SC_5H_5)(O_2SMe)Cl.$ Sulfur dioxide reacts with TiMeX₃ ($X = Br$, CI), but the products have not been well characterized. 185

Zr(CH2Ph)4 reacts with *SOz,* PhNCO, and MeNCS to give Zr (CH₂Ph)(O₂SCH₂Ph)₃(O-sulfinate), Zr [N(Ph)COCH₂Ph]₄, and $Zr[N(Me)CSCH_2Ph]_4$, respectively.⁷⁷ It reacts with NO to give mixtures of the expected $ON(CH_2Ph)N=O$ complexes.

An intriguing reaction between molecular nitrogen and a phenyl ligand has been reported³⁴⁴ (eq 16). Tich in the expected ON(CH₂Ph)N=0 complexes.

The expected ON(CH₂Ph)N=0 complexes.

An intriguing reaction between molecular nitrogen and a

henyl ligand has been reported³⁴⁴ (eq 16).

TiCp₂Ph₂ + 5LiPh $\frac{N_2}{N$

N2

0.03:O. 17 mol per Ti at 1 atm

0.15:0.65 mol per Ti at 100 atm

A TiN=NPh intermediate was postulated to account for these results. In the corresponding o-tolyl system, a mixture of all three isomeric toluidines resulted, which suggests isomerization possibly via benzyne-like intermediates during the course of the reaction (cf. ref 282a). A later study³³⁵ showed LiPh was not necessary.

IX. Addendum

In the period following completion of the manuscript, many significant findings have been reported. Perhaps the most interesting are reports by Schwartz et al. and by Schrock on the use of early transition metal alkyl compounds as potential reagents for organic synthesis.

Schwartz and coworkers³⁴⁵ have recently developed synthetically useful reactions based on zirconium alkyls of the type ZrCp₂(R)CI. These are prepared by addition of olefins to ZrCp₂(H)CI, a reaction which was briefly explored by Wailes, Weigold, and Bell.^{189b} The Zr atom migrates to the least hindered accessible position along the alkyl chain, probably by a series of Zr-H additions and eliminations. Cleavage reactions with HX, X_2 ($X = Br$, I), or acyl halides, or "insertions" with CO or *02,* thereby provide routes to alkyl halides, aldehydes, esters, acids, ketones, or alcohols. When R was fhreo- or erythro-CHDCHDCMe₃, the configuration at the α -carbon atom was retained on cleavage with Br₂ or on "insertion" of CO or **SO2,** while some racemization occurred on reaction with O₂.^{345c} Oxidation of the zirconium alkyl derivatives with tert-butyl hydroperoxide followed by hydrolysis gave primary alcohols in high yield.345d

Similar interest pertains to alkylidene complexes of tantalum which are structural analogs of phosphorus ylides. The readily available⁹⁴ (neopentyl)₃Ta= $CHCMe₃$ has been studied most thoroughly.^{346a} it reacts with C=O functions to give olefins in much the same way as do phosphorus ylides, but the scope of the Ta=C reactions appears to be broader.

Simple methylene complexes of tantalum have now been prepared. The reaction of TaMe₃Cl₂ with 1 or 2 mol of TiCp in toluene gave trimethyl compounds, TaCpMe₃Cl and TaCp₂Me₃, respectively. In contrast similar reactions between Ta(CH2CMe3)2C13 or Ta(CH2Ph)3C12 and 2 mol of TlCp provided alkylidene compounds (see section VI.B), TaCp₂(CHC-Me₃)Ci, and TaCp₂(CHPh)(CH₂Ph), respectively.^{346b} Treatment of TaCp₂Me₃ with CPh₃⁺BF₄⁻ gave $[TaCp_2Me_2]$ ⁺BF₄⁻, which, on deprotonation with base, gave the first definitive example of a transition metal-methylene complex, TaCp₂(CH₂)(CH₃). It is stable at room temperature but slowly decomposes in apparently a bimolecular fashion to give $TaCD_2(CH_3)$ (CH₂) and the fragment, "TaCp₂(CH₃)", which reacts with L (L = C₂H₄, CO, PR₃) to give TaCp₂(CH₃)L. Its reactions with CH₃I and AIMe₃ suggest that the methylene carbon atom is nucleophilic. A crystal structure³⁴⁷ showed the Ta-methylene bond length to be 2.026 (10) **A** while 'H NMR studies showed ΔG^{\ddagger} for the barrier to rotation about the Ta= CH_2 bond to be \geq 21.4 kcal mol⁻¹. In contrast, the ΔG^+ values for carbene ligand rotation in $TaCp₂(CHCMe₃)Cl$ and $TaCD₂(CHPh)(CH₂Ph)$ are 16.8 \pm 0.1 and 19.3 \pm 0.1 kcal $mol⁻¹$, respectively.

Lappert, Patil, and Pedley³⁴⁸ have measured the heats of alcoholysis of some titanium and zirconium complexes, MR4 $(R = CH₂SiMe₃, CH₂CMe₃, or CH₂Ph), M(NR'₂)₄ (R = CH₃ or$ C₂H₅), and MCI₄, and several hafnium analogs. The calculated "mean bond energies", $E(M-C)$, which varied from ca. 45 to 75 kcal mol^{-1}, can be taken as further evidence that the transition metal to carbon bond is not necessarily weak (see section VI.E). Close comparisons of $E(M-C)$ values, however, can be misleading because an alternative description of a particular M-C bond strength, the mean bond dissociation energy, \bar{D} (M-C), can be quite different from a particular \bar{E} (M-C), as in the case of $R = CH₂Ph$.

Galyer, Wilkinson, and Lloyd³⁴⁹ found the photoelectron spectrum of WMe₆ and TaMe₅ to be consistent with O_h and *D3,,* structures, respectively, in the gas phase. The spectrum reported by Cradock and Savage²⁶ is believed not to be that of WMe₆.

Several new rhenium alkyls have been reported.³⁵⁰ Volatile, red ReOMe₄ is the product of a complex reaction between ReOCl₃(PPh₃)₂ and methyllithium in diethyl ether in the presence of traces of molecular oxygen. An analogous reaction employing Me₃SiCH₂MgCl gave blue ReO(CH₂SiMe₃)₄ and red $\text{Re}_2\text{O}_3(\text{CH}_2\text{SiMe}_3)_6$. All are thermally stable and fairly unreactive toward water, dilute mineral acids, alcohols, H_2 , CO_2 , CO, and *SO2.* The ESR and electronic spectra of the monomers are consistent with square-pyramidal geometries. 351

A brief communication described the isolation of thermally stable MnR₂ complexes in which $R = CH_2CMe_3$, CH_2SiMe_3 , and CH₂CMe₂Ph.³⁵² They are apparently oxidized by molecular oxygen to give relatively unstable, green MR₄ complexes analogous to $Mn(1-norborn)l₄.⁹⁸ A uranium complex,$ **Li2U(CH2SiMe3)6(TMEDA)7,** was also described. It is the first uranium alkyl containing more than one metal to carbon σ bond.

The preparation and properties of $W(CH_2Ph)_4$ ^{72a} have now been described more fully.³⁵³ Red-brown W(CH₂Ph)₄ is best prepared from WCI4 and $Mg(CH_2Ph)_2$ in diethyl ether. It is sparingly soluble in common solvents and stable at room temperature. The authors also describe the preparation of olive-green W(CH₂Ph)CI₅ from WCI₆ and B(CH₂Ph)₃ in ether at -15 to -30° . Both form sparingly soluble bipyridyl adducts.

An expanded version of the original report⁷¹ on isolable "V(CH₂Ph)₄" has also appeared.³⁵⁴ The black, low-melting crystals have the composition $V(CH_2Ph)_4(\text{ether})_n(\text{dioxane})_m$ where $n \ge 1$ and $m \ge 2$. The ESR signal at 77^oK occurs at g $=$ 1.965, in contrast to earlier results,⁷⁰ and no signal was found in ether or as a solid at 0°. "V(CH₂Ph)₄" decomposes at 90° in ether to give 3.3-3.7 mol of toluene per V (cf. $Ti(CH_2Ph)_4$ decomposition, section II.A.3) and a dark residue which gives $V(a\text{cetate})_3$, H₂, and toluene on treatment with acetic acid.

Fröhlich and coworkers have briefly described the preparation of yellow $Zr(CH_2Ph)_2(NPh_2)_2^{355}$ and paramagnetic $(\mu =$ 1.54) $V(NEt₂)₃(C₂H₅)^{.356}$ The former was prepared by addition of 1-2 mol of $H N Ph₂$ to $Zr(CH₂Ph)₄$ and the latter by addition of LiC₂H₅ to V(NEt₂)₃CI in heptane. V(NEt₂)₃(C₂H₅), a green liquid, appears to be thermally more stable than its Ti analog (see section V.C); it decomposes at ca. 1 **15'.**

The chemistry of organochromium compounds has been discussed thoroughly in a recent book by Sneeden. 357

X. References

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