2. Niobium and Tantalum 1993

Terence P. Kee

CONTENTS

INT	RODUCTION		17
2.1	NIOBIUM(V	7) AND TANTALUM(V)	18
	2.1.1	Complexes with group 17 donor ligands	18
	2.1.2	Complexes with group 16 donor ligands	20
	2.1.3	Complexes with group 15 donor ligands	25
2.2	LOWER OX	IDATION STATES OF NIOBIUM AND TANTALUM	29
	2.2.1	Complexes with group 17 donor ligands	29
	2.2.2	Complexes with group 16 donor ligands	29
	2.2.3	Complexes with group 15 donor ligands	32
REF	ERENCES		3.5

INTRODUCTION

This review is intended to cover the literature on the coordination chemistry of niobium and tantalum through the year 1993 as reflected in *Chemical Abstracts* volumes 118, 119 and 120. To keep the review within manageable length, we have chosen to limit the study, in a similar fashion to the 1992 review, to low nuclearity inorganic coordination systems only; consequently several areas have had to be omitted. Specifically, organic/organometallic systems have been omitted unless there was good reason on account of the non-carbon based components of the coordination sphere, high-nuclearity cluster chemistry (including polyoxometallates) and solid state and materials science have been excluded unless again there was a strong reason to include them from an inorganic/coordination chemistry viewpoint.

The layout of the review focuses on the different oxidation states of niobium and tantalum which range from +5 down to -2. However, most compounds in oxidation states lower than +5 are found to be stabilised by carbon-based organic ligands and consequently, many such compounds lie outside the scope of the present review. It has proven more efficient, from a presentational point of view, to separate the +5 oxidation state from the others since most coordination chemical studies have inevitably dealt with the former state. Within each main section the sub-sections are organised according to the nature of the coordinated ligands; halogens from group 17, chalcogens from group 16, pnictogens from group 15 and ligands with other donor atoms. Within these broad boundaries however, there are areas of overlap where ligands from more than one group are present; but these species will be covered in at least one of the relevant sections.

The readers' attention is directed to a comprehensive review of the coordination chemistry of niobium and tantalum covering the years 1985-1987 especially for areas of cluster chemistry, solid state and materials related science [1] and to the companion reviews of niobium and tantalum coordination chemistry for 1992 [2].

2.1 NIOBIUM(V) AND TANTALUM(V)

2.1.1 Complexes with group 17 donor ligands

As in many previous studies, the pentahalides of niobium and tantalum have found use as precursors to other derivatives of d^0 niobium and tantalum salts via reaction with suitably donating ligands. Thus, the reaction of (EtO)₂P(=S)(OMe) (L) and Ph₂P(=S)(OEt) (L') with MF₅ (M = Nb, Ta) has been studied by ¹⁹F and ¹H NMR spectroscopy in dichloromethane (CH₂Cl₂) and acetonitrile (MeCN) solvents. The first stage of reaction occurs with coordination of the ligands to the MF₅ through the sulfur atoms with formation of octahedral MF₅L. After this, thione—thiol isomerisation of the coordinated ligands occurs as a result of which the oxygen atom becomes the donor atom. The rate of isomerisation involving the methoxy group is greater than that involving the ethoxy group [3]. A seven-coordinate intermediate is formed in which the ligands are coordinated to the central metal through sulfur and alkoxy oxygen atoms. If the solvent is not rigorously purified to remove dissolved oxygen, then besides the thione—thiol isomerisation, oxidation of the ligand occurs with separation of elemental sulfur. From a comparison of ¹H NMR chemical shifts of the uncoordinated and coordinated ligands and the values of the shift of the absorption bands of v(P=O) in the IR spectra, conclusions are drawn on the greater acceptor capacity of NbF₅ with respect to TaF₅ toward the phosphoryl-containing ligands [3].

Along similar lines to the work above, highly pure Nb(dpm)₂Cl₃ [Hdpm = dipivaloylmethane] has been synthesised by the reaction of Hdpm with NbCl₅ using anhydrous solvents under an atmosphere of highly pure argon gas [4]. The formation of Nb(dpm)₂Cl₃ was confirmed using elemental analysis, FT—IR, ¹H and ¹³C NMR and thermogravimetric-differential thermal analysis (TG—DTA). Vaporisation of impurities was not found to occur in the TG experiments. Nb(dpm)₂Cl₃ vaporised at 180—340°C without residue and is a promising new source material of metalloorganic chemical vapour deposition (MOCVD) for the preparation of photo-integrated circuit thin films [4].

The reaction of [Ph₂P(≈O)]₂C=CH₂ (L) with PF₅ and TaF₅ has been studied by ³¹P and ¹⁹F NMR spectroscopies. The compounds PF₅.L, [PF₅]₂.L, [PF₄.L][PF₆], TaF₅.L, [TaF₄.L]⁺, mer-Ta(O)F₃.L and [TaF₃.L]₂O were reported to be formed [5].

Spectroscopic studies on molten salt solutions continue to be a major feature of investigation for group 5 metal halides. Raman spectra have been obtained at temperatures in the range 375-650 K and pressures of up to four atmospheres from GaCl₃—NbCl₅ and GaCl₃—TaCl₅ binary mixtures in the liquid and vapour states. The data indicate formation of NbGaCl₈ and TaGaCl₈ liquid and vapour dinuclear addition complexes. The spectra were interpreted in terms of a C₂v configuration for the MGaCl₈ (M = Nb, Ta) molecules consisting of an MCl₆ octahedron sharing

2.1.2 Complexes with group 16 donor ligands

Since both niobium and tantalum are highly oxophilic metals, there is always a significant literature devoted to compounds in which these metals are bound to oxygen and to a lesser degree, sulfur. In most cases, the metal alkoxo and thiolato functions [M—XR] and the oxo or thio functions [M=X] (X = O, S) form the basic structural motifs with the ethereal dative bond [M—OR₂] being a feature of more loosely bound ether molecules as found in precursor compounds.

The aryloxo compound TaCl₂(O-4-¹BuC₆H₄)₃ was synthesised by reacting TaCl₅ with HO-4-¹BuC₆H₄ in a 1:3 molar ratio in dry C₆H₆ under reflux and was characterised by elemental analysis, conductance, IR, NMR and mass spectral studies. The reaction of TaCl₂(O-4-¹BuC₆H₄)₃ with equirnolar amounts of hydroxy-containing substrates having labile protons (L) such as benzoin, 2-hydroxyacetophenone, salicylaldehyde in 1:1 molar ratio afforded the six-coordinate compounds TaCl₂(O-4-¹BuC₆H₄)₃,L [9].

Aryloxo compounds have also been obtained from the reaction of dimethylamido complexes of titanium, tungsten and niobium with HO-2,4-¹Bu-6-Ph-C₆H₂. Three new mixed amido-aryloxo compounds were isolated [Ti(O-2,4-¹Bu-6-Ph-C₆H₂)₂(NMe₂)₂], [Nb(O-2,4-¹Bu-6-Ph-C₆H₂)₂(NMe₂)₃] (3) and [W₂(O-2,4-¹Bu-6-Ph-C₆H₂)₂(NMe₂)₄] and structurally characterised.

Compound 3 possesses a monomeric distorted trigonal bipyramidal geometry in which two of the amido ligands occupy mutually trans positions. The distortion from idealised geometry results from the opening up of the angle between the two formally equatorial aryloxo ligands to $144.9(1)^{\circ}$ which is presumably a reflection of the steric interactions between these two bulky ligands. Indeed, the extent of this distortion is such that the molecule is beginning to approach a square-pyramidal geometry. The conformation of the amido groups in 3 is interesting since it can shed light on the possible contributions to $p\pi$ - $d\pi$ bonding between the planar nitrogen atoms and the metal. The π -bonding of three NMe₂ groups arranged in a T-shape about a d^0 metal centre, as in 3, will be maximised with all three NC₂ propellers orientated mutually perpendicular thus allowing each nitrogen atom to donate into a different available metal d-orbital. However, in 3 this is not the case as the two mutually trans amides are parallel and the resulting competition for d-orbitals results in a significant lengthening of the Nb—N axial bonds [2.032(3) and 2.036(3)Å] over the corresponding equatorial distance [1.94(3)Å]. It is possible that steric effects are responsible for this or electronic effects due to competitive p-bonding between the metal and the aryloxo oxygen atoms.

Heating of these compounds for extended periods did not lead to cyclometallation of either the arylor tert-butyl group of the aryloxo ligand [10].

The use of metal compounds containing sterically demanding aryloxo ligands as catalysts for the polymerisation of unsaturated organic substrates has been examined by the group of Nakamura [11]. Thus, the system comprising WCl_n(O-2,6-Me₂C₆H₃)_{6-n} and EtMgBr (n = 1-4) has been examined as a catalyst for the polymerisation of 1-alkynes in toluene solvent at -20°C to +60°C for 24 h. The system WCl₄(O-2,6-Me₂C₆H₃)₂/Et₃Al polymerised ¹BuCCH to a very high molecular weight polymer ($M_n > 2 \times 10^6$) with a narrow polydispersity index (M_w/M_n ca. 1.2). By increasing the number of aryloxo ligands, less bulky 1-alkynes were polymerised to high molecular weight polymers. Thus, 1-butyne gave an orange polymer with $M_n = 9.4 \times 10^4$ and $M_w/M_n = 3.5$. Similar results were obtained when the corresponding niobium and tantalum systems were examined [11].

A variety of other mixed metal and mixed alkoxo compounds have been prepared by the Mehrotra group including XSn{Al(OiPr)4}, XSn{Zr2(OiPr)9}, XSn{M(OiPr)6}, $\{A(O^iPr)_A\}Sn\{Zr_2(O^iPr)_O\}$ and $\{Nb(O^iPr)_A\}Sn\{Ta(O^iPr)_A\}$ $\{X=CI,O^iPr,O^iBu;M=Nb,Ta\}$. These compounds were characterised by elemental analysis, molecular weight measurements and multinuclear (¹H, ¹³C, ²⁷Al and ¹¹⁹Sn) NMR analyses [12]. In a similar fashion, the compounds $Cl_{4-n}Sn\{M(O^{i}Pr)_{6}\}_{n}$ (M = Nb, Ta; n = 1-3) have been synthesised by the reactions of $SnCl_{4}$ with an appropriate salt K{M(OiPr)6} in the necessary molar ratios. These chloro precursors were then used to obtain $(RO)_2Sn\{M(O^iPr)_6\}_2$ (R = Me, iPr), $\{Ta(O^iPr)_6\}_2Sn\{Nb(O^iPr)_6\}_2$ and {Al(OiPr)4}4ClSn{Nb(OiPr)6}2 by halide atom replacement with the appropriate potassium alkoxide salt. Characterisation of these compounds was provided by elemental and alkoxo group analysis, IR, molecular weights and multinuclear (¹H, ¹³C, ²⁷Al and ¹¹⁹Sn) NMR spectroscopic analyses [13]. A variety of other mixed metal alkoxo systems based on tert-butoxide have been characterised using the same techniques by the same research group. Some examples include FeCl2{Al(OBu¹)4}2, Fe(OBu¹)2{Al(OBu¹)4}2, Fe{Al(OBu¹)4}{Nb(O¹Pr)6}2 and Fe[Al(OBu^t)4]2[Nb(OⁱPr)6] which were synthesised by metathetical reactions. Alcoholysis of Fe(Al(OBu¹)4)2{Nb(O¹Pr)6} with an excess of MeOH and ¹BuOH produces Fe[Al(OMe)₄]₂[Nb(OMe)₆] and Fe[Al(OBut)₄]₂[Nb(O i Pr)₂(OBut)₄] respectively [14].

$$MCl_5 + PPh_3 + R^1R^2CO \longrightarrow [R^1R^2C(PPh_3)OMCl_4]Cl$$
 (2)
 $(M = Nb, Ta)$

The formation of the [M—O] bond via the interaction of metal halides and tertiary phosphines with carbonyl substrates has been reported [15]. This reaction (2) is a form of phosphonium ion formation and bears a remarkable resemblance to the Pudovik and Abramov reactions of organophosphorus chemistry [16]. Thus, treatment of aldehydes and ketones with one mole equivalent of a Lewis acid such as TiCl₄, NbCl₅ or TaCl₅ and one mole equivalent of PPh₃ affords the derivatives [R¹CH(PPh₂R²)OMCl_n]Cl and [R¹R²C(PPh₂R²)OMCl_n]Cl (M = Nb, Ta; n = 4; R¹ = various organic groups; R² = Ph) as air and moisture sensitive solids [15]. On the basis of NMR spectroscopic studies, some of the adducts obtained from aldehydes exist in solution as

mixtures of isomers or oligomers whereas the ketone derivatives are in equilibrium with free ketone and NbCls.PPha.

As well as alcohols, carboxylic acids have been used as a versatile source of the [M—O] bond. Thus, it was found that the products from the interaction of MCl₅ (M = Nb, Ta) with RCO₂H (R = Ph, 4-MeC₆H₄, 4-FC₆H₄) included {[TaCl₂(μ-O)(O₂CC₆H₄Me-4)]₄) (4) which contains a [Ta₄O₄] core, [NbCl₃(O₂CR)₂(μ-O)] and [TaCl₄(O₂CR)] [17]. In a similar fashion, reaction of NbCl₅ with aryl carboxylic acids yields the oxo-niobium carboxylates [NbCl₃(O₂CR)₂(μ-O)] [R = Ph (5), 4-MeC₆H₄, 2-MeC₆H₄, 4-FC₆H₄, 4-ClC₆H₄, 2-FC₆H₄, 2-ClC₆H₄, 2-BrC₆H₄, 2,6-Me₂C₆H₃] [18]. A single crystal X-ray analysis of the benzoate derivative (5) shows a μ-oxo-bis(μ-carboxylato)dimetal core in which each niobium atom is in a distorted octahedral environment, the other three sites on each metal being occupied by chlorine atoms. Spectroscopic data suggest similar structures for the remaining derivatives.

(4) $R = 4 - MeC_6H_4$

The preparation and crystal structure of Nb(HTRENCAM).DMSO [H₆TRENCAM = hexadentate tris(catecholamide) (6)] has been reported, revealing a cavitand-like arrangement of the three catecholamide groups with the protonated non-binding nitrogen atom directed inwards to the niobium atom. The geometry around the niobium atom (shaded) resembles a distorted mono-capped trigonal antiprism (7) [19].

The use of sterically demanding alkoxo ligands has continued to provide a rich seam of chemistry, especially in the field of low coordinate metal species containing d-electrons. Such lowcoordinate metal centres are particularly susceptible towards oxidation addition and cyclisation reactions with enormous potential for the manipulation of both saturated and unsaturated organic substrates and indeed, as model systems for the investigation of mechanistic features of important industrial reactions such as the Fischer-Tropsch process. The reduction of Ta(SILOX)3Cl2 (SILOX = Bu₃SiO-) with Na/Hg in the under H₂ afforded Ta(SILOX)₃H₂ [(8); 43%]. Compound (8) underwent a thermal cyclometallation to TaH(SILOX)2OSilBu2CMe2CH2 (9) but was reconstituted by treatment with H₂ (6 days, 3 atm). Treatment of (8) with C₂H₄, neat CCl₄ and MeI in diethylether generated Ta(SILOX)3HEt (63%), Ta(SILOX)3HCl (63%) and Ta(SILOX)3HI (62%) respectively. Reduction of Ta(SILOX)3HI with Na/Hg in thf produced a ring-opened thf compound [(SILOX)3TaH]2[\mu:\pi-(CH2)3O] (58\pi). Photolysis of Ta(SILOX)2Cl2CH2Ph gave [(SILOX)2TaCl]2(µ-H)2, C7Hg and a trace of dibenzyl. Reduction of Ta(SILOX)2Cl3 with Na/Hg under 1 atm. H2 over the course of 15 days yielded an unbridged D2d dimer [(SILOX)2TaH2]2 (83%) which possessed a [Ta—Ta] single bond with a length of 2,720(4)Å by single crystal X-ray diffraction [20]. Subsequent exposure of [(SILOX)₂TaH₂]₂ to 2 equivalents of HCl, 1 equiv. O₂ and 1 equiv. Me₃NO provided [(SILOX)₂TaCl]₂(μ-H)₂ (78%), [(SILOX)₂TaH]₂(μ-O)₂ [(10); 95%] and $[(SILOX)_2TaH]_2(\mu-H)_2(\mu-O) [(11); 67\%)]$. Derivatisation of (10) with C₂H₄ gave $\{(SILOX)_2TaEi\}_2(\mu-O)_2$ (39%). Compound (11) exists as a mixture of two C_2 symmetric isomers; the hydrides of one isomer exchanged with an energy barrier of ΔG ca. 8 kcal mol-1 whilst those of the other isomer exchanged coincidentally with isomeric interconversion (an energy barrier of ΔG ca. 11 kcal mol-1 was measured). Dihydride (8) undergoes σ-bond metathesis with D₂ forming initially Ta(SILOX)3HD and the exchange of (8) with hydrogen was measured directly using spin saturation transfer techniques. Carbonylation of (8) and Ta(SILOX)₃HEt afforded η²-aldehyde complexes of the general form Ta(SILOX)a(n-RCHO), Treatment of [(SILOX)aTaCl]a(u-H)a with CO yielded [(SILOX)₂TaCl]₂(u-H)(u:n²:n²-CHO) [(12); 56%)] whilst exposure of dimeric [(SILOX):TaHolo to 1 equivalent of CO afforded [(SILOX):TaHlo(u-CHo)(u-O) [(13); 67%). Subsequently, {(SILOX): TaH\: (u.m²:m²-CHO)(u.m¹:m²-CH2O)|Ta(SILOX)::] (14) was isolated in 55% yield and converted to [Ta(SILOX)212(u-O)2(n-CHMe) [(15); 61%)]. The sequence [(SILOX)₂TaH₂]₂ + CO going to (13) then (14) then (15) exhibits the critical bond breaking and the C-H and C-C bond making events of the Fischer-Tropsch (F-T) process. Extended exposure of [(SILOX)₂TaH₂]₂ or (14) to 1 atm. CO provided [(SILOX)₂Ta]₂(μ:η¹:η¹-CH=CHO)(μ : η^1 : η^2 -CH₂O)(μ -O) (16; 50%). Carbonylation of [(SILOX)₂TaH]₂(μ -O)₂ (10) generated [(SILOX)₂Ta]₂(μ-O)₂(μ-CH₂O) as the major product (70-90%), while treatment of (11) with CO afforded first [(SILOX)₂TaH]₂(μ-O)₂[TaMe(SILOX)] (90%) and then [(SILOX)₂Ta]₂(μ-O)₂(μ-MeCHO) (90%). ¹³CO labelling studies were used to follow the (13)-(15)-(16) conversions, thus providing a basis for mechanistic interpretation. Dimeric structures allow oxygenated fragments to remain coordinated to two tantalum atoms throughout the sequence. Insertion into [Ta-H] bonds may initiate each carbonylation process. Stereochemical consequences of SILOX ligation are discussed in relation to the structures and dynamics of the dinuclears, while the electrophilic tantalum centres are important in H/D exchange and carbonylation chemistry. The carbonylation chemistry underscores three critical points regarding the F-T process: (i) hydride transfer to CO is a reasonable alternative to CO dissociation; (ii) adsorbed hydrocarbyl and oxygenate fragments are related by reversible C-O bond breaking and bond making events; (iii) oxygenate and hydrocarbyl adsorbates can be removed protolytically, akin to hydrogenation [20].

Metal-oxo species are again well represented. CsBCO₆ (B = Sb, Nb, Ta; C = Te, Mo, W, U) have been prepared from CsNO₃ and the respective metal oxides or UO₂(NO₃)₂.6H₂O by the ceramic method. All of the compounds have a perovskite structure except CsNbUO₆ which is monoclinic, space group $P2_1/c$, Z = 4 [21].

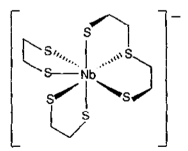
A number of derivatives in which the metal contains a formal {M=O} unit have been reported. Thus Nb(O)Cl₃.L (L = PMe₃, PMePh₂, PPh₃, PEt₃), Nb(O)Cl₃.(Ph₂PCH₂CH₂PPh₂) and Nb(O)Cl₃(PMe₂Ph)₃ were prepared by reaction of either [Nb(O)Cl₃]_n or Nb(O)Cl₃(thf)₂ with a stoichiometric amount of the phosphine in CH₂Cl₂ solvent. The influence of the different ligand fields on the IR stretching frequency of the metal-oxo unit was discussed in terms of the geometry of the complex and the stereo-electronic properties of the attendant ligands. Thus, the more phosphines attached to the metal and the more electron-donating the phosphorus centre, so the lower the [Nb=O] stretching frequency [22]. On the basis of similarities in the IR and other spectral data, it was envisaged that Nb(O)Cl₃(PMe₂Ph)₃ possessed a similar monocapped trigonal anti-prismatic geometry to Nb(O)Cl₃(PMe₃)₃ but there was no evidence for the former existing in different, apparently isomeric, forms unlike the latter.

Moving to ligands based on sulfur donor atoms, aspects of the coordination chemistry of dithiophosphates have been examined. Thus, $M(O^iPr)_3[S_2P(OR)_2]_2$ (M = Nb, Ta; R = Pr, iPr , iBu , Ph) were prepared by reacting $MCl_2(O^iPr)_3$ with the requisite molar ratio of Na or NH4[S₂P(OR)₂] in thf solvent. The monomeric products are viscous liquids which are soluble in common organic

solvents and highly susceptible to hydrolysis; they have been characterised by IR and NMR spectroscopies which indicate a didentate mode of attachment of the dithiophosphato groups to the metal [23].

A number of transition metal complexes containing ligands based on the 1-substituted-2-tetrazoline-5-thione scaffold (17) have been reported including complexes of niobium and tantalum. These complexes have been characterised using elemental analysis, conductance measurements, IR and UV-VIS spectral studies. The Ph, 3-MeC₆H₄, 4-MeC₆H₄ and 4-ClC₆H₄ derivatives of 1-substituted-2-tetrazoline-5-thione are coordinated through the thione tautomeric form with VO₂+, MoO₂²⁺, WO₂²⁺ and ZrO²⁺ species but interact via the thiol form in the cases of Nb⁵⁺ and Ta⁵⁺ resulting in deprotonation of the thiol. The metal-ligand vibrations in the far-IR region have also been examined [24].

A facile [C—S] bond cleavage reaction has been reported to take place in the reaction between NbCl₅ and 3 equivalents of Li(TPDT) (H₂TPDT = 3-thiapentane-1,5-pentanedithiol) followed by a cation exchange with Ph₄PBr to afford, (Ph₄P)[Nb(S)(EDT)(TPDT)], (Ph₄P)[Nb(EDT)₂(TPDT)] [(18); H₂EDT = 1,2-ethanedithiol) and (Ph₄P)[Nb(O)(EDT)(TPDT)] [25]. The crystal structure of (18) is also reported (monoclinic, space group $P2_1/c$, a = 14.031(2), b = 9.140(2), c = 28.030(4) Å; $\beta = 107.28(1)^{\circ}$ and Z = 4) and is found to contain a seven coordinate mixed dithiolate complex anion (shown below), in which the primary coordination sphere of the niobium centre is composed solely of sulfur donor atoms with a distorted pentagonal-bipyramidal geometry.



(18) Hydrogens omitted for clarity

2.1.3 Complexes with group 15 donor ligands

Because of the highly electrophilic nature of niobium and tantalum, most studies in this field have focused on ligands containing electronegative nitrogen donor atoms.

The complex Ta(=CH^tBu)Cl₃(thf)₂ is one such precursor compound which adds iminoboranes ^tBuB=NMes in a 1:2 molar ratio to afford the cyclobutane compound (19) [26].

As in previous years, amido [M—NR₂] and imido [M=NR] compounds have formed the bulk of niobium and tantalum systems studied with group 15 donor ligands.

The reaction of Cp*TaCl₄ (Cp* = C₅Me₅) with 2 equivalents of LiNHAr (Ar = 2,6-iPr₂C₆H₃) in the solvent affords the imido compound Cp*Ta(NAr)Cl₂. Upon reaction of Cp*TaCl₄ with 4 equivalents of LiNHAr in diethylether solvent the isolated product is [Cp*Ta(NAr)₂Cl][LiOEl₂] (20) [27]. The corresponding the adduct is obtained when reaction is performed in the solvent or when Cp*Ta(NAr)Cl₂ is reacted with 2 equivalents of LiNHAr in the solvent. A single crystal X-ray diffraction study on (20) reveals a [LiOEl₂]+ moiety bridging the imido nitrogens. In solution, the two tantalum-bound imido groups are equivalent with restricted rotation about the [Ta—N—Cipso] bonds implying an intact [LiOEl₂]+ bridge. The compound Cp*Ta(NAr)(NEl₂)Cl (prepared from Ta(NAr)(NEl₂)Cl₂(thf)₂ and LiCp* reacts slowly with LiNHAr in refluxing the to afford [Cp*Ta(NAr)₂Cl][Li(thf)].

Multidentate ligands have proved a popular theme for study. Thus, it has been reported by Schrock and co-workers that TaCl₅ reacts with Li₃(N₃N) [(N₃N) = (Me₃SiNCH₂CH₂CH₂)₃N] to give Ta(N₃N)Cl₂ which reacted with LiPHR (R = Ph, Cy, 'Bu) in Et₂O to give Ta(=PR)(N₃N) (21). Aldehydes react with (21) to afford Ta(=O)(N₃N) and RP=CHR' (R = R' = tBu; R = Ph, 'Bu, R' = ferrocenyl, Fc; R = R' = Ph) [28]. The phosphaalkene 'BuP=CHFc dimerises to the corresponding

1,3-diphosphetan. Cyclopentadiene adds to unstable PhP=CHFc to afford the Diels-Alder adduct. Compound (21) (R = cyclohexyl) has been examined by X-ray crystallography and is found to be orthorhombic, space group $Pna2_1$, with Z=4, a=19.754(2), b=11.862(1), c=12.993(1) Å.

Me₃Si
$$R_3$$
 SiMe₃ R_3 SiMe₃ R_3 SiMe₃ R_3 SiMe₃ R_4 S

Along similar lines, Arnold and co-workers have independently examined the same systems in relation to their interactions with group 16 donor ligands. Thus, $TaCl_5$ reacted with $(HN)_3N$ in Et_2O to give $Ta(N_3N)Cl_2$, which reacted subsequently with $(thf)_2LiESi(SiMe_3)_3$ (E = S, Se, Te) at $-78^{\circ}C$ to afford $Ta(=E)(N_3N)$. Both $Ta(N_3N)Cl_2$ and $Ta(=E)(N_3N)$ were characterised by IR, Raman, ${}^{1}H$, ${}^{13}C\{{}^{1}H\}$, ${}^{77}Se\{{}^{1}H\}$, ${}^{125}Te\{{}^{1}H\}$ NMR and mass spectroscopic techniques. $Ta(=Se)(N_3N)$ (22) is orthorhombic and $Ta(=Te)(N_3N)$ (23) monoclinic, both compounds possessing the same distorted trigonal bipyramidal geometry as compound (21) with the selenium and tellurium atoms in axial positions with respect to the tertiary nitrogen atoms [29].

The 1:1 reaction of TaCl₂[2,6-(Me₂NCH₂)₂C₆H₃](CH⁴Bu) with ⁴BuCH₂ZnCl affords the dimetallic compound (24) in high yield. The five-coordinate tantalum metal centre has a coordination geometry between trigonal bipyramidal and square pyramidal, whilst the zinc centre has a distorted tetrahedral ligand array. The alkylidync function and the aryl ipso-carbon bridge between the zinc and tantalum centres. The p-orbital at C_{ipso} of the aryldiamine ligand forms a 2-electron bond to the zinc centre, thus functioning as an 8-electron donor. The reaction of (24) with tmen at

60°C leads to elimination of ZnCl₃ and the formation of (25); which results from an alkylidyne-mediated activation reaction of a methyl C—H bond in a dimethylamino group [30]. In the presence of $[Pd(2-Me_2NCH_2C_6H_4)(\mu-X)]_2$ (X = Cl, I), the reaction of (25) with trnen appears to generate $TaCl_2[2-Me_2NCH_2C_6H_3](C'Bu)[2,6-(Me_2NCH_2)_2C_6H_3]$ and as such, is an unanticipated, new palladium-mediated alkylidyne functionalisation reaction.

The addition of TaCl₂Me₃ to 2 equivalents of LiNHSiR₃ (R = ¹Bu) in hexane solvent results in the liberation of methane and isolation of Ta(=NSiR3)(NHSiR3)Me2 (26) in 63% yield [31]. Thermolysis of (26) in benzene resulted in disproportionation but in pyridine and thf solvents, Ta(=NSi^tR₃)₂L₂Me (L = pyridine 68%; thf 13%) and methane were produced. The bis-adduct is considered to form via a 1.2-elimination of methane from Ta(=NSi¹R3)(NHSiR3)Me₂py which is obtained from (26) and pyridine at 25°C, ¹H and ¹³C NMR spectra of the bis-adduct indicated equivalent pyridine ligands but a single crystal X-ray structure determination revealed trigonal bipyramidal pseudo-Cs stereoisomer with an axial methyl group and equatorial imido ligands that reflects the steric requirements of the bulky R3SiN= units. Rather long [Ta=N] distances (1.810(13) and 1.819(13)Å1 support electronic arguments suggesting that the imides donate a maximum of 6-electrons to the metal centre, Addition of TaCls to 4 equivalents of LiNHSiR3 in Et2O at -78°C afforded Ta(=NSi¹R₃)(NHSiR₃)₂Cl and R₃SiNH₂. Alkylation of Ta(=NSi¹R₃)(NHSiR₃)₂Cl with AlMe₃ in hexane, PhLi in Et₂O/hexane, PhCH₂K in toluene and RCH₂Li in Et₂O provided Ta(=NSi⁴R₃)(NHSiR₃)₂R (R = Me 78%; Ph 64%; CH₂Ph 51%; CH₂R 39%). Addition of R3SiNH2 or R3SiOH to (26) yielded Ta(=NSiIR3)(NHSiR3)2Me or Ta(=NSi¹R₃)(OSiR₃)₂Me (52%) and methane. Thermolysis of Ta(=NSi¹R₃)(NHSiR₃)₂R effected 1,2-RH elimination to form transient Ta(=NSiR3)(NHSiR3)2 which is capable of adding one C-H bond across one imide linkage. Ground state information was obtained via the approach to equilibrium of Ta(=NSi¹R₃)(NHSiR₃)₂Ph and CH₄ but observation of a ditantalum species $[(R_3SiNH)_2Ta(=NSiR_3)]_2(\mu:\eta^1,\eta^1-1,4-C_6H_4)$ complicated the measurements. Simulation of the approach to equilibrium yielded rate constants consistent with the previously measured 1,2-RHelimination rates and showed that Ta(=NSi¹R₃)(NHSiR₃)₂R possessed relatively similar ground state free energies. Equilibration of Ta(=NSi'R3)(NHSiR3)2CH2Ph to the aryl complex Ta(=NSi4R3)(NHSiR3)2C6H4Me in toluene at 182° gave similar results. The data portray differing 1,2-RH-elimination rates that result from significant transition state energy differences, ruling out a late transition state despite a rough correlation of rate with the C-H bond strength of the eliminated alkane/arene. The implications of these measurements, including the possibility of d⁰-alkane or arene complexes as intermediates and differences in [Ta-C] bond strengths are discussed in great detail [31].

Among the studies with niobium(V) and tantalum(V) and phosphorus donor ligands other than the phosphinidine species (21), it has been reported that Ta(OAr)₂Cl₂(H)(PMe₂Ph)₂ (HOAr = 2,6-diisopropylphenol) reacts with 1, 2 or 3 equivalents of organic isocyanides to produce a sequence of organometallic products resulting from the initial insertion into the [Ta—H] bond and subsequent coupling reactions [32]. These products contain a number of unusual nitrogen based ligands as illustrated in Scheme 1.

2.2 LOWER OXIDATION STATES OF NIOBIUM AND TANTALUM

2.2.1 Complexes with group 17 donor ligands

The study of solutions of $[(Ta_6X_{12})(H_2O)_4X_2].4H_2O$ (X = Cl, Br) in methanol-H₄/methanol-D₄ (1:5 v/v) by ¹H NMR spectroscopy has been reported. Within the region 3.7—3.8 ppm, the spectra reveal a complex multiplet characteristic of methyl groups in coordinated methanol [33]. The ten resonances observed in the bromo system can be assigned to the nine species, including cis and trans isomers, which can be formed in the substitution series $[(Ta_6X_{12})(H_2O)_{6-x}(MeOH)_x]^{2+}$ with 1 < x < 6. A model for predicting the chemical shift and intensity of each methyl resonance in the series was derived for the case where H₂O and MeOH compete on an equal basis for the six terminal coordination sites of the Ta_6Br_{12} cluster unit. The ¹H NMR spectrum of $[(Ta_6Cl_{12})(H_2O)_4Cl_2].4H_2O$ in methanol changes slowly with time and achieves the equilibrium condition after ca. three days. This spectrum is more complicated because of competition of Cl⁻, along with H₂O and MeOH, for the terminal coordination sites. This is supported by specific conductance data in methanol and quantitative detection of free chloride ions using ³⁵Cl NMR spectroscopy [33].

2.2.2 Complexes with group 16 donor ligands

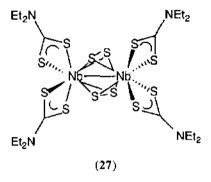
Since a decrease in metal oxidation state commonly results in a lowered electrophilicity, the lower oxidation states of niobium and tantalum tend to be stabilised more by the lower congeners of group 16 rather than by oxygen. Nevertheless, an oxochloroniobium(IV) complex has been isolated

by zinc reduction of NbCl₅ in concentrated HCl solution. The product, an extremely moisture and oxygen sensitive intense blue solid, was assigned a dimeric structure (H₃O)+₂[Nb₂O₂Cl₆] based on elemental analysis, IR, UV-VIS spectroscopy as well as magnetic and conductimetric measurements [34].

In the area of surface oxo-bound organometallics, the reduced metal species has been shown to have important catalytic applications. New niobium dimers on four different kinds of silica (SiO₂), prepared by the reaction between [Nb(η^5 -C₅H₅)(H)- μ -(η^5 ; η^1 -C₅H₄)]₂ and surface hydroxo functions followed by chemical treatments, have been found to have different structures with different bond distances and coordination numbers associated with the [Nb—Nb] bond [35]. These results have been gained by EXAFS analysis. Subsequently, it was found that these niobium dimers are active in the dehydration of ethanol; indeed, the activity appears to be selective for this alcohol. The rate constant for ethene formation was found to be influenced by adsorbate-adsorbate or niobium-niobium interactions. The ratio of rate constants for intra- and intermolecular dehydration was also controlled by changing the niobium structures. On a SiO₂ surface, the direct niobium-niobium bond was not observed while the dimeric structure with bridging oxygen was formed after exposure to ethanol at 523 K. Probably, on this catalyst, ethanol dehydration proceeds in conjunction with the formation and breaking cycles of the niobium-niobium bond [35, 36].

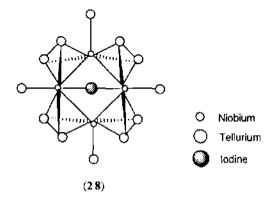
Turning to sulfur chemistry, tantalum sulfide cluster ions ($Ta_nS_m^+$; n < 9; m < 30) have been produced by direct laser ablation of a mixture of tantalum and sulfur powders, and have been studied with a tandem time-of-flight mass spectrometry. The tantalum sulfide clusters in maximum abundance were found to be $[Ta_nS_{2n+7}]^+$ (n = 1-9) detected in the first stage mass spectra. The main dissociation path of the UV-photolysis (248 nm) of tantalum sulfide ions is a sequential S_2 loss. For cluster ions with n > 3, the photolysis yields $Ta_3S_4^+$ or $Ta_4S_6^+$ as the major product [37].

Continuing on the theme of sulfur clusters, the dimeric species $Cs_4Nb_2S_4(NCS)_8$ has been reacted with NaS_2CNEt_2 in water to afford the compound $Nb_2S_4(S_2CNEt_2)_4$ (27) which has been shown to crystallise in the monoclinic system, space group C2/m, a = 21.181(8), b = 6.958(1), c = 16.623(6) Å; $\beta = 133.95(2)^{\circ}$; Z = 2 [38]. The compound contains the $[Nb_2(\mu^2-S_2)_2]^{4+}$ core in which the metal ions are in the formal +4 oxidation state, the remaining valency resulting in the formation of a [Nb-Nb] bond with a distance of 2.890(2) Å. The dithiocarbamate ligands, $[S_2CNEt_2]$ coordinate in a didentate fashion and along with the S_2^{2-} ligands form a distorted square antiprismatic geometrical arrangement around each niobium atom.



A multi-unit system based on niobium sulfide clusters and containing [MS4]3-; FeCl2; NaSEt (M = V, Nb) has been shown to undergo heteronuclear self-assembly in MeCN solvent to afford the clusters, [V2FexSg(SEt)o]3-, [Nb2FexSg(SEt)o]3- and [Nb2FexSg(SEt)o]5- which were isolated as EtaNH+ salts [39]. These clusters have the triply-bridged double cubane stereochemistry $\{|MFe_3S_4(SEt)_3|_2(u-SEt)_3\}^{3-}$, as demonstrated for the M = V cluster by single-crystal X-ray diffraction. These clusters, together with others previously reported, form a set [MoFeaSx(SEt)o]3whose members (M = V, Nb, Mo, W, Re) possess the same charge, identical ligands and essentially congruent structures with only small metric differences. This condition permits the examination of changes in heterometal on cluster properties. Redox potentials for the reversible couples $\{M_2Fe_6S_8(SEt)_9\}^{3-/4}$ and $\{M_2Fe_6S_8(SEt)_9\}^{4-/5}$ decrease in the order M = V < Nb = Mo < W = MoRe. Reductions become more difficult as the ferrous character of the core increases. Redox reactions are accompanied by changes in electron density that largely occur in Fe₃S₄ cluster portions. The concept of cuboidal Fe₃S₄ as a cluster ligand is considered in terms of isomer shifts of heterometal clusters. The shifts of the majority of such clusters are close to those protein-bound [Fe₃S₄]^{Q/1}-. providing an apparent definition of oxidation states. Isotropic shifts of ligands at iron sites provide a convenient indicator of the ground spin state of the clusters. Cubane-type MFe₃S₄ clusters are of renewed interest in view of the occurrence of the cuboidal cluster fragment MoFe₃S₃ in the co-factor of nitrogenase. The possible relation of these results to the FeMo-co-factor and the FeV-co-factor of nitrogenase is noted [39]. If the co-factor of vanadium containing nitrogenase has an analogous structure, properties derived from the heterometal cuboidal fragment may be influenced by vanadium in a manner similar to that for the cubane-type clusters.

After sulfur, tellurium ligands have been best represented in the lower valent coordination chemistry of niobium and tantalum. Thus, a new infinite-chain niobium telluride iodide has been prepared by reaction of the elements at 893 K. Nb4(Te2)4Te4I (28) is monoclinic, space group C2/c, with a = 21.957(5), b = 6.147(3), c = 19.925(4)Å and $\beta = 122.40(1)^\circ$. Compound (28) represents a new, one-dimensional structure type. The structure consists of infinite chains of Nb4(Te2)4Te4I which are formed by the tetra-nuclear butterfly cluster units shown below and in which iodine atoms bridge different cluster units [40].



The complex $[Ta(Te_2)_2]_4[PtI_6]I_2$ (29) was synthesised from the elements at 680°C and its crystal structure determined; space group P1, a = 16.831(1), b = 17.740(1), c = 19.018(1)Å and $\alpha = 93.57$, $\beta = 94.08$, $\gamma = 102.09$ °. The structure of this quasi-one-dimensional compound consists of $[Ta(Te_2)_2]$ chains separated by $[PtI_6]$ groups and iodine atoms [41] such that the tantalum atoms adopt a square prismatic geometry as illustrated below. A formal oxidation state assignment $[(Ta^{5+})(Te_2^{2-})_2]_4[(Pt^{4+})(I^-)_6](I^-)_2$ indicates that semi-conducting properties are to be expected for this material

2.2.3 Complexes with group 15 donor ligands

Among the lower oxidation states, the most common have been +3 or +4 with nitrogen or phosphorus donor ligands. Thus, the single crystal X-ray analysis of TaCl4py2 [(30); py = C5H5N] has been reported [42]. The compound was found to be monomeric with a distorted octahedral geometry in which the two pyridine ligands occupy mutually *trans* positions. It is suggested that the observed structure may be due to a combination of steric and electronic factors.

Lower valent complexes of tantalum have been demonstrated to be useful models for homogeneous hydrogenation catalysts. It has been demonstrated that an $\eta^1(N) - \eta^2(N,C)$ bonding rearrangement of quinoline occurs upon reducing quinoline adducts of d^0 -Ta(V) to the formal d^2 -Ta(III) oxidation state and that $\eta^2(N,C)$ coordination allows selective hydrogenation of the heterocyclic ring. In addition, the $\eta^2(N,C)$ coordination mode renders the C_{α} of a substituted pyridine substrate susceptible to nucleophilic attack resulting in C-N bond cleavage. This demonstration of C-N bond scission affords a more fundamental understanding of one way to

cleave C—N bonds in hydrogenation catalysis. One feature of this reduced oxidation state tantalum system is that, in contrast to osmium η^2 -pyridine systems such as $[(\eta^2-C,C-2,6-Me_2C_5H_3N)Os(NH_3)_5]^{2+}$, the tantalum system binds heterocycles only in the $\eta^2(N,C)$ mode in the d^2 -electronic configuration, thereby disrupting the aromaticity of the heterocyclic ring only [43].

The syntheses and crystal structures of a number of monomeric tantalum(IV) amido halide and dimeric tantalum(IV) imido halide complexes have been reported by Hoffman and co-workers [44]. Ta[N(SiMe3)2]2Cl₃ was reduced by Na/Hg to afford Ta[N(SiMe3)2]2Cl₂ (31). Ta(NSiMe₃)[N(SiMe₃)2]Cl₂ was reduced under analogous conditions to give {Ta(μ-NSiMe₃)[N(SiMe₃)2]Cl₂ (32). Both compounds (31) and (32) have been characterised by single crystal X-ray diffraction analyses where it was found that (31) possesses a severely distorted tetrahedral geometry whilst (32) has an edge-shared tetrahedral structure with a tantalum-tantalum separation of 2.621(1)Å [44].

$$(Me_3Si)_2N$$
 $(Me_3Si)_2N$ $(Me_3Si)_2N$ $(Me_3Si)_2N$ $(Me_3Si)_2N$ $(Me_3Si)_2N$ $(SiMe_3)_2N$ $(SiMe_3)_2N$

A number of other amido complexes of group 5 metals in lower oxidation states have been reported by the same group, including Nb[N(SiMe₃)₂]₂Cl₂ and Nb(NPh)₄, both of which possess tetrahedral geometries, the former being isostructural to the tantalum analogue (31) [45].

Treatment of Tp'NbCl2(PhCCR') [R' = Me, Et; Tp' = hydrido tris(3,5dimethylpyrazolylborato)) with NaOMe gives good yields of the corresponding chiral chloromethoxo derivatives, Tp'NbCl(OMe)(PhCCR'). The methyl and phenyl-methoxo complexes, Tp'NbR"(OMe)(PhCCR') (R' = Me, R" = Me; R' = Et, R" = Me; R' = Et, R" = Ph) are obtained in high yields upon reaction of either Tp'NbCl(OMe)(PhCCR') (R' = Me, Et) with MeLi or PhLi respectively [46]. Due to the resultant chirality at the niobium atom, the ethyl groups in these complexes give rise to AMX3 type ¹H NMR spin systems although in some of the complexes, deceptively simple spectra can be observed regardless of solvent or temperature even though rotation about the [Nb-Ph] bond can be frozen out. The complexes, Tp'NbR"(OMe)(PhCCR') (R' = Me, R" = Me; R' = Et, R" = Me; R' = Et, R" = Ph) react with CO to form five-membered oxoniobacycles Tp'Nb(OMe)[η^2 -C(Ph)C(R)C(Me)O] [R = Me (33); Et] and Tp'Nb(OMe)[η^2 -C(Ph)C(Et)C(Ph)O] respectively from regioselective coupling of the coordinated alkyne with an acyl or benzoyl group. A single crystal X-ray diffraction study has confirmed the structure of (33), wherein the metallacycle is folded in an envelope conformation about the Ca-O axis with the Ca-Nb-O triangle constituting the flap. Examination of metric parameters within the ring and comparison with related complexes indicate that an analogy between the oxoniobacycle and the bonding description for a metallacyclopentatriene is appropriate [46].

Along similar lines, deprotonation with "BuLi in truen at the propargylic site of the coordinated alkyne in Tp'NbCl(X)(PhCCMe) (R = H, X = Cl, OMe; Tp' = as above) followed by reaction with MeI or PhCH₂Br affords the alkylated alkyne compounds Tp'NbCl(X)(PhCCCH₂R) (R = Me, X = Cl, OMe; R = CH₂Ph, X = Cl, OMe) in high yield. The addition of PhCH₂Br to deprotonated Tp'NbCl(OMe)(PhCCCH₂Me) affords a 4:1 diastereoisomeric mixture of Tp'NbCl(OMe)(PhCCCHMe(CH₂Ph)) whereas a 1:6 ratio is obtained under the same conditions starting from Tp'NbCl(OMe)[PhCCCH₂(CH₂Ph)] and MeI [47].

Moving on to phosphorus-bound systems, two examples of a new type of face-sharing bioctahedral niobium complex that contain three bridging chloro ligands and two terminal phosphines in a syn configuration have been reported by Cotton and co-workers [48]; (Bu4N)[Nb2Cl7(PEt3)2] (34), the isostructural complex (Bu4N)[Nb2Cl7(PMe3)2].thf (35) and the related triangular (Me3PH)(Bu4N)[Nb3Cl10(PMe3)3(BF3)].thf (36) (the relevant metal containing anions are illustrated below). Compounds (34) and (35) have effective magnetic moments of 0.84 and 0.99 BM respectively at room temperature. Pathways of interconverting edge-shared bioctahedral [Nb2Cl6(PEt3)4] to the face-sharing [Nb2Cl7(PEt3)2]—and finally to the triangular trinuclear [Nb3Cl10(PEt3)3]—and even larger clusters is discussed [48].

The monomeric, paramagnetic tantalum(II) complex, TaBr₂(PMe₂Ph)₄ (37) has been isolated from the reduction of TaBr₅ with LiBEt₃H in toluene solvent in the presence of PMe₂Ph and characterised by single crystal X-ray diffraction techniques [49]. The coordination polyhedron

is essentially distorted octahedral with a mutually cis arrangement of bromo ligands. However, the structure is severely distorted towards a bicapped tetrahedron with the angle Br-Ta-Br = 84.23(4)°.

The phosphido-phosphinidene coordination mode of a P_2 ligand has been realised in the trinuclear complex [(TaL)₃(P₄)(P₂)] (L = 1,3- 4 Bu₂C₆H₃) prepared by reaction of TaL(CO)₄ with white phosphorus (P₄) in decalin solvent at 190°C for 4 h. Structural proof was obtained by single-crystal X-ray diffraction of the product of the reaction of [(TaL)₃(P₄)(P₂)] with Fe₂(CO)₉, namely [(TaL)₃[P₄{Fe(CO)₄}(P₂)] (38) [50].

The kinetics of the reaction of [Ta(COSiPr¹3)(CO)(dmpe)₂], (dmpe = 1,2-dimethylphosphinoethane) with Me₃SiCl in the total afford [Ta(Me₃SiOCCOSiPr¹3)(CO)(dmpe)₂Cl] have been studied using stopped flow spectrophotometric techniques [51]. The rate is greatly accelerated by the addition of the salts, [(n-pentyl)₄N]Cl and [(n-butyl)₄N][BPh₄] due to the increased ionic strength of the solution. The reaction is first order in both [Ta(COSiPr¹3)(CO)(dmpe)₂] and Me₃SiCl in the presence of excess [(n-butyl)₄N][BPh₄]. At 22(1)°C, a second order rate constant of 1.71(4) M⁻¹s⁻¹ was obtained. These results were interpreted in terms of initial electrophilic attack of the silyl reagent at the oxygen atom of the carbonyl ligand in [Ta(COSiPr¹3)(CO)(dmpe)₂]. In order to model the transition state or intermediate of this rate-determining step in the [C—C] bond-forming reaction, AlEt₃ was added to the siloxocarbene complex [Ta(COSiBulPh₂)(CO)(dmpe)₂]; spectroscopic and structural characterisation of this complex revealed that coupling does not necessarily occur upon addition of a Lewis acid; adduct formation appears to be the initial event resulting in the formation of [Ta(COSiBulPh₂)(COAlEt₃)(dmpe)₂]. However, further discussion of the organometallic aspects of this work lies outside the scope of this review.

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3. Scandium 1993

Catherine E. Housecroft

CONTENTS

INT	RODUCTION	37
	SOLID STATE COMPOUNDS	
	FULLERENE DERIVATIVES	
3.3	COMPLEXES WITH CARBABORANE LIGANDS	38
3.4	COMPLEXES WITH HALIDE LIGANDS	38
	COMPLEXES WITH GROUP 16 DONOR LIGANDS	
3.6	COMPLEXES WITH GROUP 15 DONOR LIGANDS	40
RFF	FRENCES	41

INTRODUCTION

This review surveys the coordination chemistry of scandium reported during the year 1993 and follows a similar format to the 1992 review [1]. The literature has been searched by using Current Contents and the Cambridge Crystallographic Data Base. Structural figures have been redrawn using coordinates taken from the latter, implemented through the ETH, Zürich [2]. Organometallic complexes have, in general, been excluded, although in section 3.2 some fullerene chemistry is described.

A review which includes discussions of zero oxidation state scandium compounds has appeared [3].

3.1 SOLID STATE COMPOUNDS

The reaction of elemental scandium with Sc₂O₃ and iodine in the solid state has been studied by DTA. The product is ScOI, and the structure of this compound has been determined by X-ray crystallography [4]. The crystal structure of CsScS₃ has also been elucidated [5].

3.2 FULLERENE DERIVATIVES

Members of a series of endohedral scandafullerenes have been characterised by mass spectrometric and ESR spectral data. In the same study, Shinohara et al. have shown that Sc@C82 can be separated from Sc3@C82 by using liquid chromatography with an ethanol-deactivated silicagel column [6]. The same workers have reported that Sc2@C74, Sc2@C82 and Sc2@C84 may be