PROGRESS IN NIOBIUM AND TANTALUM COORDINATION CHEMISTRY

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1. INTRODUCTION

The material covered by this review was mainly found in Chemical Abstracts, Volumes 102-107, i. e. the years 1985, 1986 and 1987. Thus, the majority of the papers mentioned has been published in previous years, i.e. approximately between 1984 and mid-1987. A few newer references are included. The review can be considered as a continuation of previous reviews in this series [1-3] and of other reviews on niobium, tantalum and in some cases vanadium coordination chemistry [4-9].

The papers are referred in the following sections mainly according to the formal oxidation state of the metal, ranging from (V) to below (0). In some cases it is not trivial to divide after the oxidation state. Within a section, the material is further sub-divided according to the kind of ligands bound to the metal. The basic order of sections is such: halogens, halogens mixed with non-halogens, no halogens.

Short modern reviews on basic niobium and tantalum chemistry can be found in many textbooks on inorganic chemistry, e.g. [7, 10-11]. Niobium and tantalum compounds are of importance in several areas of scientific interest, but some of these areas are of less importance from the point of view of a coordination chemist. Hence, papers dealing with subjects such as the structural chemistry of hydrides of Nb and Ta metals, or the catalytic activity of e.g. Nb205 or Nb02 supported catalysts [12], are not referenced. Likewise, papers dealing with e.g. sputtering for the preparation of thin-film optical devices, or the characterization of nonlinear optical crystals of doped compounds (typically LiNbO3 and KTaO3), or the preparation of ceramics (e.g. NbC-NbO-NbN sintered phases), alloys, superconductors or magnetic materials, and the properties of such materials, are generally not referenced. The new research-field on charge density waves in e.g. niobium and tantalum sulphides is covered by referring to some of the most important new papers. Extraction chemistry on niobium and tantalum is shortly included (the extraction of niobium from ore or scrap to ferroniobium, niobium complexes and finally high-purity niobium metal has recently been reviewed [13-15].

Traditionally, coordination chemistry has been concerned with molecular species, but in recent years it has become increasingly linked with solid state chemistry. Structural results are now often described in terms of polyhedra, see Table 1, instead of sometimes rather mysterious close-packed arrays, as discussed in a recent account on the relation between coordination chemistry and the solid state [16]. Thus, the dramatic increase in the reseach for new materials with interesting properties is reflected in the large number of solid state structural results referred in this review.

Table 1. Oxidation states and common stereochemistry of Niobium and Tantalum compounds [7], [10-11].*

Oxidation state	Electronic state	Coordination number	Polyhedron	Examples
M(V)	ď	4	Tetrahedron	[NbO4] ³⁻ , [NbO{N(SiMe ₃) ₂ } ₃]
		5	Trigonal bipyramid	[MCl5](vapour), [TaMe5],
			Distorted square pyramid	$[M{NMe_2}_5]$, $[NbOCl_4]$, $[NbSCl_3(Ph_3PS)]$
		6	Octahedron	[TaF ₆] ^{Γ} , [Nb ₂ Cl ₁₀], [TaOCl ₃], [NbAlCl ₈], [Nb(SO ₄) ₆] ^{Γ} , [NbCl ₅ ·OPCl ₃], [Nb(OMe) ₅] ₂ , [MCl ₃ (OR) ₂] ₂ , [TaCl ₅ ·S(CH ₃) ₂], NaNbO ₃ (perovskite)
			Trigonal prism	[Nb(S2C6H4)3]
		7	Distorted pentagonal bipyramid	[NbOF6] ³⁻ , [Nb(O ₂)F5] ²⁻ , [NbO(C ₂ O ₄)3] ³⁻ , [NbO(S ₂ CNE ₂) ₃], [Ta=S(S ₂ CNE ₂) ₃], [Ta(Me ₂)(S ₂ CNMe ₂) ₃], [Ta(Me ₃)(S ₂ CNR ₂) ₂]
			Capped trigonal prism	[TaF ₇] ² -
		8	Bicapped trigonal prism or square antiprism	$[TaF_8]^{3-}$, $[Ta(PS_4)S_2]$
			Dodecahedron	$[M(O_2)_4]^{3-}$, $[Ta(S_2CNMe_2)_4]^+$, $[Nb(O_2)_2(C_2O_4)_2]^3$
		9	?	$[(\eta^5-C_5H_5)_2T_8H_3], [\{NbCl_2(\eta^5-C_5H_5)_2\}_2O]$
		10	?	[NbCl5(O2CCOOH)5]
M(IV)	dl	4	D _{2d} Tetrahedron	[Nb(NEt ₂) ₄]
		5	Trigonal bipyramid	[NbH ₂ (OSiBu ^t ₃) ₂] ₂
		6	Octahedron	$[NbCl_6]^{2-}$, $[TaCl_4(py)_2]$
		7	Distorted pentagonal bipyramid	[NbF ₇] ³ -
			Capped octahedron	[NbCl4(PMe3)3]
		8	Dodecahedron	[Nb(CN)8] ⁴⁻
			Square antiprism	[Nb2Ci8(PMe3)4]
		high	?	[(η ⁵ -C ₅ H ₅) ₂ NbMe ₂]
M(III)	d ²	6	Octahedron	$[Nb_2Clg]^{3-}, \ [NbCl_3(py)_3], \ [Ta_2Cl_4(\mu\text{-}Cl_2)(SMe_2)_4]$
			Trigonal Prism	LiNbO ₂
		7	Complex	[TaCl3(CO)(PMe2Ph)3], [TaH(PPh3)2(dmpe)2]
	2	8	Dodecahedron	[Nb(CN)8] ⁵ -
M(II)	d ³	4	Square	NbO
		6	Octahedron	[NbCl ₂ (PMe ₃) ₄]
			Trigonal prism	NbS
		8	Dodecahedron	[TaH2Cl ₂ (PMc ₃) ₄]
1.00	d ⁴	_	Square antiprism	[TaH2Cl2(dmpe)2]
M(I)	a٠	5	Pentagonal bipyramid	[TaL ₂ R(C ₂ H ₄) ₂]
		6	?	$[Nb(\eta^4-C_4H_6)_2(\eta^3-Meallyl)]$
		7	Capped octahedron	[TaH(CO) ₂ (dmpe) ₂], [Ta(CO) ₃ (PMe ₃) ₄] ⁺
			Capped trigonal prism	[TaX(CO)2(dmpe)2], [NbCl(CO)3(PMe3)3]
	.e		'4-legged pianostool'	[Nb(CO)4(C5H5)]
M(0)	d ₅	6	Octahedron	[Nb(dmpe)3]
M(-I)	d6	6	Octahedron	[Nb(CO)6]-
M(-III)	d ⁸	5	Trigonal bipyramid	[Ta(CO) ₅] ³ -
		6	?	$[M(CO)_3(C_5H_5)]^{2}$

^{*} For meaning of ligand names, see list of abbreviations.

2. Nb(V) and Ta(V) compounds (d⁰)

2.1 Nb(V) and Ta(V) complexes containing only halides

The NbF5 and TaF5 *mono*meric molecules have been studied in the gas phase by means of electron diffraction [17]. A D_{3h} symmetry is preferable to describe the structure of the molecules. In the solid state MF_5 (M = Nb, Ta) exists as tetrameric molecules, M_4F_{20} [18, 19]. Solid Ta₄F₂₀ was studied by ¹⁸¹Ta NQR spectroscopy at low temperature, and evidence was found for the presence of hexadecapole interactions obtained (I = 7/2 for ¹⁸¹Ta). The Ta₄F₂₀ entity is rhombohedrally distorted [19].

Thermodynamic data for gaseous tantalum fluorides, TaF_5 , Ta_2F_{10} , and Ta_4F_{20} were reviewed and data were given also for a large number of liquid and solid tantalum halide and oxyhalide compounds [20]. The molecular constants and thermodynamical characteristics of the NbF₆⁻ and TaF_6 ⁻ ions were determined in vapours over the oxyfluorides between 730 and 960 K [21]. The electronic structure of the TaF_6 ⁻ anion was calculated by the discrete-variation $X\alpha$ method in an expanded basis of numerical Hartree-Fock functions [22]. Graphite can accomodate trigonal-*bi*pyramidal NbF₅ molecules and octahedral NbF₆⁻ ions, forming intercalation compounds [23].

Thermodynamic data for gaseous tantalum halides, TaCl₅, TaBr₅ and TaI₅ were reviewed and data were given also for a large number of liquid and solid tantalum halide and oxyhalide compounds [20]. Enthalpy contents and molar heat capacities of NbCl₅ and TaCl₅ were measured as function of temperatures [24]. The stability of NbCl₅ gas in contact with metallic niobium was studied by DTA and TG methods in order to find the optimum conditions for the synthesis of lower chlorides [25]. The reduction to NbCl₄ starts at *ca.* 205 °C. NbCl₅ does not exhibit fluorinating behaviour towards *poly*fluorinated *cyclo*alkenes at 150 °C [26]. Experimental hints for the preparation of solid NbCl₅ was described [27]. The heat capacity of niobium *penta*chloride was measured from 8 to 315 K and thermodymanic data evaluated [28].

The IR spectra of Ar and N₂ matrices trapping vapours coming from solid NbX₅ (X = Cl and Br) have been investigated [29]. The vapours contain *monomeric* NbX₅, *dimeric* Nb₂X₁₀ and also *poly*meric species. Assignments of the spectra were given.

NbCl₅ and NbBr₅ molecules of D_{3h} symmetry have been subjected to vibrational normal coordinate analyses using Wilson's GF method. Assignments of observed vibrational frequencies and calculated potential energy distribution values for each vibration are given [36].

Solid NbCl₅ and TaCl₅ compounds have been reexamined by X-ray diffraction and IR spectroscopy [37]. NbCl₅ was studied by 93 Nb and 35 Cl nuclear quadrupole resonance spectroscopy in the temperature range 4.2 - 440 K [38]. Catalytic properties of NbCl₅ and TaCl₅ on the preparation of triphenylbenzene was reported [39,40]. The sublimation pressure of solid NbBr₅, NbI₅ and TaI₅ has been determined by a spectrophotometric method, as function of temperature [41]. The melting point of NbI₅ was 673 ± 1 K, and the vapour pressure of the melt was given versus temperature [41]. The photoelectron spectrum (AlK α , HeI) of NbI₅ was studied in an attempt to find the relation between the energy of the niobium core levels and the oxidation number [42].

Both NbF₅ and TaF₅ have been intercalated into graphite in the presence of chlorine, and it has been thought to happen via the formation of complex chloride-fluoride ions NbF₅Cl⁻ and TaF₅Cl⁻. This idea has found support by the characterization (by X-ray diffraction and mass spectroscopy) of intercalates formed via reaction between HgF₂ and NbF₅ or TaF₅. The NbF₆⁻ and TaF₆⁻ ions were identified by Raman spectroscopy [43].

Not just TaF₆⁻ but a mixture of the ions TaF₆⁻, TaF₇²⁻, TaF₈³⁻ in mutual exchange of F-seems to exist in a saturated solution of K₂TaF₇ dissolved in 48 % HF, according to ¹⁸¹Ta NMR data [44]. Also, the ¹⁸¹Ta NMR spectrum of the [TaCl₆]⁻ ion in solution was obtained and interpreted in terms of a perfect octahedral symmetry: Tantalum has a large quadrupolar moment and, without the cubic environment, the NMR band would be much broader than found [44]. The TaF₆⁻ anion has been identified in dry methylene chloride solution by means of ¹⁹F NMR spectroscopy [18].

Vibrational spectra were studied of niobium and tantalum complexes in hydrofluoric acid [45]. For high concentrations of HF and metal, TaF₆-, TaF₇²-, NbF₆- and NbOF₅²- ions exist in the solutions. According to the interpretation of the spectra, dilution leads to a relative increase in TaF₇²- and NbOF₅²- concentrations [45]. Raman spectra of K₂NbF₇, K₂TaF₇ and CsTaF₆ have been published in connection with extraction experiments [46].

Alkali metal hexafluorotantalates, $A[TaF_6]$ with A = Li, Na, K, Rb, or Cs, were prepared and characterized by IR and X-ray diffraction. Crystal symmetries and lattice constants were determined [47]. Based on coordination geometry, a classification of fluorides and oxyfluorides of niobium or tantalum, i.e. the compounds $AMF_{6-2X}O_X$ with A = Li, Na, K, Rb, Cs and M = Nb or Ta, has been attempted [48]. Low-temperature phase transitions in KNbF₆ and KTaF₆ were determined by the ¹⁹F NMR relaxation method and related to rotation of the MF₆-octahedral anions [49].

The lightly blue silver(II) hexafluoro niobate and tantalate Jahn-Teller-compounds, $Ag[MTaF_6]_2$ with M = Nb or Ta (Ta is paramagnetic), were prepared from oxides in an autoclave at ca. 380 °C and under a F₂-pressure of ca. 3 kbar. Single crystal needles were characterized by X-ray diffraction (layer structure with cumulated AgF_6 and MF_6 octahedra) [50].

Methods of preparing shiny crystals of the new compounds $Hg_3[MF_6]$ and $Hg_{3-x}[MF_6]$, M = Nb and Ta, and x small, have been described [51]. These compounds are exceptional in containing electrically conducting either layers or linear chains of mercury atoms in between layers of $[NbF_6]^-$ and $[TaF_6]^-$ octahedra. A transformation between the two kinds of structures takes place in contact with liquid SO_2 or liquid AsF_3 at temperatures from room temperature to $120 \, ^{\circ}C$ [51].

The compounds K[NbCl6] and K[TaCl6] have been studied [52] to elucidate the mechanism of their superionic conduction. Methods applied involved ac-conductivity, differential scanning calorimetry, and powder neutron diffraction. Crystal symmetries and phase relations versus temperature were reported [52]. The electrical conductivity of NbCl5-NaCl, NbCl5-KCl, TaCl5-NaCl and TaCl5-KCl binary melt systems were measured to determine the phase diagrams [53]. Enthalpy contents and molar heat capacities of RbNbCl6, CsNbCl6, RbTaCl6 and CsTaCl6 were measured as function of temperatures, and all these compounds undergo allotropic solid-solid transformations as well as fusion, and enthalpies and entropies associated with these transformations have been evaluated [24].

New pyridinium salts, $(py-H...py)^+[MCl_6]^-$ and $(py-H...py)^+[MX_5Y]^-$, with M = Nb, Ta; X = F, Cl; Y = Cl, Br, and $py = C_5H_5N$ (pyridine), have been characterized [54]. Also, new acidic hydrogen-bound etherate salts, $(Et_2O-H...OEt_2)^+[MCl_6]^-$ and $(Et_2O-H...OEt_2)^+[MX_5Y]^-$, with M = Nb, Ta; X = F, Cl; Y = Cl, Br and $Et_2O = diethylether$, have been prepared [54]. Furthermore, acidic ethylenediamine salts, $H[MCl_6] \cdot n(en)$ and $H[MX_5Y] \cdot n(en)$, with M = Nb, Ta; X = F, Cl; Y = Cl, Br; n = 2, 3 or 4 and $en = H_2NCH_2CH_2NH_2$ (ethylenediamine), were obtained [54]. The compounds were characterized by chemical analysis and IR spectra.

Electron-transfer-induced ligand-exchange reactions have been discovered by electrochemical methods (cyclic voltammetry and constant potential electrolysis) applied on niobium chlorides (Et₄N)[NbCl₆], (Et₄N)₂[NbCl₆], [NbCl₅], and [NbCl₄(NCMe)₂], in "superdry" acetonitrile (NCMe). Two one-electron reduction waves were observed, corresponding to consecutive reductions to Nb(IV) and Nb(III) compounds [55].

Cyclic voltammograms have been obtained on NbCl₅ dissolved in a room temperature molten salt liquid consisting of 49.0:51.0 mol % AlCl₃-(1-methyl-3-ethylimidazolium chloride), proving that the *dimeric* niobium oxide chloride species (formed by reaction with the usually present oxide impurities) can be converted to the "oxide-free" [NbCl₆]⁻ species by a treatment of the melt with COCl₂ [56].

Crystalline NbAlCl₈ was prepared directly from Al₂Cl₆ and Nb₂Cl₁₀: The crystal structure consists of an AlCl₄ tetrahedron sharing an edge with an NbCl₆ octahedron [57], see Figure 1.

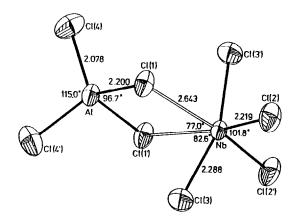


Figure 1. Structure of the [NbAlCl8] molecule. Reproduced with permission from [57], B. Krebs, H. Janssen, N. J. Bjerrum, R. W. Berg and G. N. Papatheodorou, *Inorg. Chem.* 23 (1984) 164.

Two salts, [PCl4][NbCl6] and [PCl4][NbCl6]·SOCl2, of the tetrachlorophosphonium and hexachloroniobate ions have been prepared from PCl5 and NbCl5 in thionyl chloride solution [58]. Their structures were studied by vibrational spectroscopy.

The *penta*chlorides MCl_5 (M = Nb, Ta) were found to be reactive with an excess of *tri*thiazyl chloride, (NSCl)₃ in CCl₄ suspension to give the ionic compounds, [S₄N₄Cl][MCl_6]. Furthermore, with SCl₂, the ionic species [N(SCl)₂][MCl_6] can be obtained. The products were characterized by chemical analysis and IR-spectra [59].

The electrochemical behaviour of Ta(V) was studied at 400-550 °C in mixed melts containing LiCl-KCl-TaCl₅ [60]. By reduction, the anionic [TaCl₆] complex most probably forms a stable and soluble Ta(III) complex, which in a second step can be further reduced to the metal. By titration of such melts with solid KF it was shown that in mixed chloride-fluoride melts, primarily [TaF₇]²- complexes are present, but also small amounts of [TaF₆] and [TaClF₆]²-coexisted with them [60].

The crystal structure of K_2TaF_7 has been determined by using single crystal X-ray diffraction data [61]. The TaF_7^{2-} polyhedra may be described as monocapped trigonal prisms (distorted from true mm or C_{2v} symmetry) with the capping atom located on one of the rectangular faces, see Figure 2. The Ta-F bond lengths varied from 1.918 to 1.975 Å. Raman spectra showed bands at 645 cm⁻¹ (Ta-F stretching) and 395 and 280 cm⁻¹ (Ta-F bending). The previously unreported luminescense properties of crystals of K_2TaF_7 as well as of K_2NbF_7 have been reported [61-62]. The emission spectrum of crystalline K_2NbF_7 showed a long

progression in a Nb-F deformational mode at 290 cm⁻¹ [62]. The mechanism of electrochemical reduction of K₂NbF₇ and K₂TaF₇ on a nickel electrode in LiF-NaF molten mixtures was studied, and stable NbNi₃ and TaNi₃ phases identified [63].

Figure 2. Structure of the [TaF7]²⁻ ion in the potassium salt. Reproduced with permission from [61], C. C. Torardi, L. H. Brixner, and G. Blasse, *J. Solid State Chem.* 67 (1987) 21.

The reaction of Ta₂O₅ with NH₄HF₂ and alkaline earth fluorides on heating was shown to give CaTaF₇, BaTaF₇, SrTaF₇, Ba₃(TaF₈)₂, Ta₂O₃F₆, and Ba(TaF₆)₂, and the stability of these and analogous compounds was discussed [64].

The hydrazinium heptafluorotantalate monohydrate compound, (N₂H₆)TaF₇·H₂O, decomposes in four steps through the intermediates (N₂H₆)TaF₇, (N₂H₅)TaF₆ and (NH₄)TaF₆, according to results obtained by thermogravimetry, differential thermal analysis, X-ray powder diffraction, chemical analysis and vibrational spectra [65].

An organic molecular charge-transfer salt, (BEDT-TTF) $_3$ Ta $_2$ F $_{11}$ has been prepared electrochemically (BEDT-TTF is a short name for bis-ethylenedithiolotetrathiafulvalene) [66]. This salt contains the Ta $_2$ F $_{11}$ anion with a unique linear fluorine bridge between two tantalum atoms, according to a crystal structure determination [66]. The salt exhibits an optically enhanced magnetic phase transition which was detected by ESR absorption spectroscopy [67]. The Ta $_2$ F $_{11}$ anion has also been been identified in dry methylene chloride solution by means of $_1$ F NMR spectroscopy [18] and in the crystal structure solution of (Hg $_4$ ²⁺)(Ta $_2$ F $_{11}$ ⁻) [68].

2.2 Nb(V) and Ta(V) oxyhalide complexes

2.2.1 Oxyfluoride complexes

Molecular compounds NbOF₃, TaOF₃, NbO₂F and TaO₂F were formed by the reaction between NbF₅ or TaF₅ and silica under various conditions [69]. The compounds were identified by X-ray diffraction and mass spectroscopy [69]. Thermodynamic data for gaseous tantalum oxyhalides, TaOF₃ and TaO₂F were reviewed and data are also given for a large number of liquid and solid tantalum halide and oxyhalide compounds [20]. Chemical and structural data were given for NbOF₃ and TaOF₃ [69]. A powder sample of NbO₂F was investigated by ⁹³Nb NMR two-dimensional spectroscopy [70].

"Non-stoichiometric" niobium oxide fluorides, Nb₃₁O₇₇F, Nb₃₄O₈₄F₂, Nb₅₉O₁₄₇F, and Nb₆₅O₁₆₁F₃ were prepared by reacting Nb₂O₅ with Nb₃O₇F at 1200 °C [71]. According to high-resolution electron microscopy and computer image simulations, these compounds have block-structures, like *e.g.H*-Nb₂O₅ [71].

The molecular constants and thermodynamical characteristics of the NbOF₄⁻ ion were determined in vapours over niobiumoxyfluorides between 730 and 960 K [21]. Based on the coordination geometry, development of a classification system has been attempted for the fluorides and oxyfluorides of niobium or tantalum, i.e. the compounds $AMF_{6-2x}O_x$ with A = Li, Na, K, Rb, Cs and M = Nb or Ta, and x = an integer [48].

The crystal structures of sodium *penta*fluoro oxy-niobates(V), Na₂[NbOF₅] and α -Na₃[NbOF₆] were solved [72-72A], see Figure 3. Na₂[NbOF₅] contained discrete [NbOF₅]²⁻ ions of slightly distorted octahedral configuration [72]. The O-Nb-F equatorial angles were larger than 90°, being on the average 97° \pm 2°, and the symmetry of the ion was found to be approximately C_{4v} . In Na₃[NbOF₆], the [NbOF₆]³⁻ ion with pentagonal *bipy-ramidal* configuration was determined, adding diversity to the earlier found *monocapped* octahedral geometry of the ion [72A].

Unexpectedly, crystal structure solutions of $(Hg_3)[(MF_5)_2SO_4]$ salts, with M = Nb or Ta, have proven that ions $[(MF_5)_2SO_4]^{2-}$ with a MF_5 -O-SO₂-O- MF_5 bonding geometry have a significant stability [68].

¹⁹F NMR spectroscopy was used to study the oxyfluoride complexes present in acidic aqueous solutions and in acetonitrile [73]. The *trans*-[NbOF₄(OH₂)]⁻ anion and the [NbOF₅]²⁻ anion were detected.

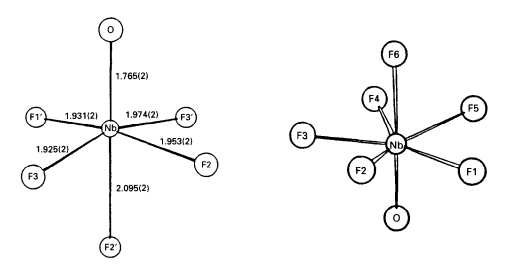


Figure 3. The [NbOF₅]²⁻ and [NbOF₆]³⁻ anions as found in the crystal structures of the sodium salts. Reproduced with permission from [72-72A], R. Stomberg, *Acta Chem. Scand.* A 38 (1984) 603 and A 37 (1983) 453.

 $[MOF_5]^{2-}$ anions (M= Nb and Ta) have been identified by IR spectroscopy as corrosion products, formed by the attack of HF-DMF-H₂O₂ solutions on the metal [74]. The Raman spectrum of K₂NbOF₅ has been published in connection with extraction experiments [46]. The preparation of alkali metal *mono*oxofluoroniobates, ANbOF₄ and A2NbOF₅ (A= Li, Na, K, Rb and Cs), in molten NH₄HF₂ was described [75]. IR spectra indicate ANbOF₄ to have a chain structure and A2NbOF₅ a discrete-ion structure. X-ray data were given for LiNbOF₄ and NaNbOF₄ [75]. In the KF-Nb₂O₅ system, three intermediate tetragonal-tungsten-bronze phases, KNb₂O₅F, KNb₄O₁O_F and KNb₆O₁₅F were identified [76]. Also, unit cell dimensions of K₂NbOF₅, K₂TaOF₅, K₃NbO₂F₄ and K₃TaO₂F₄ were determined by X-ray powder diffraction [77]. The compounds were prepared by heating the corresponding fluoro-*per*oxy-metallates [77].

The reaction between Nb₂O₅ and CoO in molten NH₄HF₂ has been studied by DTA, IR spectroscopy and X-ray diffraction [78]. CoNbOF₅ was formed. At high temperatues it pyrolysed to Co₂NbO₃F₃ giving off NbF₅ and HF vapours [78].

An attempt has been made [79], to make it possible beforehand to distinguish, among niobium and tantalum oxide fluorides, those compounds which will have a coordination type of structure (containing isolated octahedra): The ions present must be of a similar size; otherwise island,

chain, layer, or framework type structures seem to be the result [79]. New niobium and tantalum oxyfluoride double salts, Li₄NbO₄F and Li₄TaO₄F, have been prepared and characterized by X-ray powder diffracion and infrared spectroscopy [80-82]. When passing from LiMF₆ complex salts to LiMO₃ through oxy-fluorides, a transition from compounds of an island-type through chain and laminated ones to framework structures was observed [80]. The reaction between NbO₂F and alkali metal carbonates was studied by DTA, DGA and X-ray analysis; M_4 NbO₄F and M_2 NbO₃F (M = Li, Na, K, Rb and Cs) were formed [83]. The reaction between TaO₂F and lithium, sodium and potassium carbonates was studied by DTA, DGA and X-ray analysis; Li₄TaO₄F, Na_{0.9}5TaO_{2.9}5F_{0.05}, Na₂Ta₂O₅F₂, K₂TaO₃F and K₆Ta_{6+x}O₁₅-xF_{6+7x} (x is small) were the main components formed [84].

By heating Nb₂O₅ and NbO₂F with Cu₂O it was possible to obtain red crystals of Cu_{0.6}Nb₆O_{14.6}F_{1.4}, suitable for X-ray structure solution [85]. The structure is of the same type as that of LiNb₆O₁₅F, H-LiTa₃O₈ and Cu_{0.8}Ta₃O₈, containing pentagonal columns with seven-coordinated niobium atoms in NbX₇ pentagonal *bipyramids* (X = F or O), sharing edges with five NbX₆ octahedra (see Figure 4). These building units are further linked *via* cornersharing.

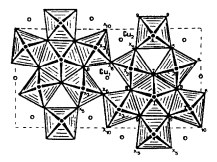


Figure 4. Projected structure of Cu_{0.6}Nb₆O_{14.6}F_{1.4}. Niobium is displaced from the center of the (hatched) octahedra. Reproduced with permission from [85], P. N. Wa Llunga, *Acta Chem. Scand.* A38 (1984) 641.

Within the system PbO-PbF₂- M_2 O₅-MO₂F (M = Nb, Ta), several crystal structures have been obtained:

- a) Pb₂Nb₃O₇F₅ is built of slabs of ReO₃ structure type of net composition Nb₃O₇F₃₂ 2 -, separated by layers of Pb₂F₂ 2 +, and related to the Bi₄Ta₃O₁₂ structure [86].
- b) The Pb₃Nb₄O₁₂F₂ and Pb₃Ta₄O₁₂F₂ structures contain corner-sharing $M(O,F)_6$ octahedra, related to the pyrochlore, fluorite and β -Na₂Ta₂O₅F₂ structures [87].

- c) The Pb3Ta5O9F13 structure can be described as made up from slabs of ReO3 structure type and bands of red PbO2 structure type [88].
- d) The Pb₁₂Ta₉O₂₀F₂₉ structure contains columns of ReO₃ structure type, 3 times 3 octahedra wide, separated by layers of Pb₂F₂²⁺, and related to the layered Bi₂O₂²⁺ perovskites (Aurivillius phases) [89].
- e) The $Pb_X M(O,F)_{3+x/2}$ (x = ca. 0.25, M = Nb, Ta) structures which are disordered superstructures of the hexagonal tungsten bronze type [90], see Figure 5.

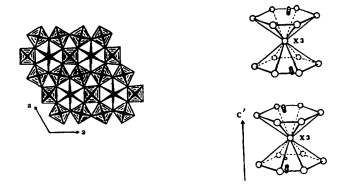


Figure 5. Structure of $Pb_X M(O,F)_{3+x/2}$, with x = ca. 0.25 and M = Nb, Ta. The Pb atoms are located in the hexagonal tunnels (left). A view of a suggested Pb-X-Pb arrangement (X is F or O) in the tunnels (right). Reproduced with permission from [90], Ö. Sävborg, J. Solid State Chem. 57 (1985) 160.

2.2.2 Other Oxyhalide complexes

Thermodynamic data for gaseous tantalum oxyhalides, TaOCl₃, TaO₂Cl, TaOBr₃, TaO₂Br, TaOI₃ and TaO₂I were reviewed and data given also for a large number of liquid and solid tantalum halide and oxyhalide compounds [20].

The IR spectra of Ar and N_2 matrices trapping vapours coming from solid NbOX₃ and NbX₅/Nb₂O₅ mixtures (X = Cl, Br and I) have been investigated [29]. The vapours contain monomeric NbOX₃ and also polymeric species. Assignments of the spectra were given.

The He I α photoelectron spectrum of gaseous *mono*meric NbOCl₃ was assigned using the new semi-empirical molecular-orbital calculation technique SCC-X α [30]. Molecular orbital

correlation diagrams are given and bonding modes of NbOCl₃ are discussed in comparison with similar molecules like POCl₃ and VOCl₃ [30]. The sublimation pressure and the standard entropy of NbOCl₃ was redetermined [31].

NbOCl₃ and TaOCl₃ powders have been examined by X-ray diffraction; the two compounds were found to be *iso*-structural and indexed powder diagrams were given [32]. The sublimation pressure of solid NbOCl₃ has been determined by the transportation method, as a function of temperature [33]. Thermodynamic functions for solid and gaseous NbOCl₃ were given [33]. Compounds like NbO₂Cl Nb₃O₇Cl, Nb₅O₁₁Cl₃, NbO₂Br, TaOCl₃, TaO₂Cl, Ta₃O₇Cl, TaO₂Br (some of which are new) have been made and studied [34]. The structure of NbOBr₃ was investigated by electron diffraction [35].

Niobium *penta*chloride reacts with *tri*chloromethane, forming $NO^+[NbOCl_4]^-$ and CCl_4 [91]. The $[M_2OCl_{10}]^{2-}$ anions, M = Nb or Ta, were found in crystal structures [92-93], in the case of Nb for the first time. Both of the ions consist of two $MOCl_5$ octahedra sharing the oxygen apex and with linear M-O-M bonding arrangements and inversion symmetries.

Lightly red NH₄[NbOBr₄] is formed when a mixture of NbOBr₃ and NH₄Br is heated in a sealed ampoule at 400 °C. X-ray data and vibrational spectra of NH₄[NbOBr₄] have been reported [94, 94a].

Potentiometric investigations of niobium(V) and tantalum(V) oxychloro complex formation were done in NaCl-AlCl₃ melts at 175 °C [95]. Results were explained by a two-equilibria model:

with pK values of 2.2 and 3.95 for M = Nb, and 2.74 and 4.52 for M = Ta, respectively.

The X-ray structure of prepared Bi₄NbO₈Cl single crystals was solved and consists of [NbO₄] layers composed of much distorted octahedra which are canted with respect to one another. From powder data it was concluded that Bi₄TaO₈Cl and Bi₄NbO₈Br crystallize with the same structure [96].

Luminescence properties of La₃TaO₄Cl₆ crystals have been reported [97]. The structure is similar to that of Pr₃NbO₄Cl₆ [98] and contains infinite linear chains of corner-sharing TaO₅ trigonal *bi*pyramids with the equatorial oxygens corner-sharing the oxygens of the LaCl₇O₂ *poly*hedra. Crystals in which lanthanum(III) was partly replaced with indium or other rare earth ions (Sm, Eu, Tb, Dy and Tm) have been prepared and their peculiar luminescence properties examined [99].

Iso-structural $Ln_3MO_5XCl_3$ crystals have been prepared and their crystal data reported, with Ln = La, Ce, Pr, Nd, Th; M = Nb, Ta; X = O, OH, F [100]. The single crystal structures of La₂ThTaO₆Cl₃ and Ce₃TaO₆Cl₃ [100], as well as of La₃TaO₅(OH)Cl₃ [101] were solved. The structures consist of TaO₆ polyhedra with unusual trigonally prismatic environments for Ta, see Figure 6.

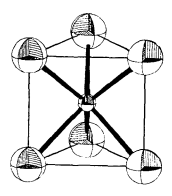


Figure 6. The unusual trigonally prismatic coordination polyhedron of TaO₆ in the structure of La₂ThTaO₆Cl₃. Ta-O bond length = 1.984 Å. Reproduced with permission from [100], U. Schaffrath and R. Gruehn, J. Less-Comm. Met. 137 (1988) 61.

2.3 Nb(V) and Ta(V) peroxyhalide and peroxy-pseudohalide complexes

The penta fluoro-peroxy-niobate(V) ion, $[NbF_5O_2]^{2-}$, 1, which is known [102] to be pentagonal bipyramidally seven-coordinate, has been investigated in the sodium and potassium salts

$$\begin{bmatrix} F & F' & O \\ F'' & Nb & O \\ F'' & F' & O \end{bmatrix}$$

by means of Raman and IR spectroscopic techniques [103-105, 106]. For the sodium salt, bands observed at 955, 880 and 900 cm⁻¹ were assigned to \vee (O-O), \vee _S(Nb-O) and \vee _{aS}(Nb-O), respectively, based on C_{2v} symmetry, but the assignments were questioned because of probable

impurities present in the sample. A diagram shows the π -bonding involved in between NbF5 and the *per*oxide ion. (The work by Nour *et al.* has been published three times in virtually identical forms [103-105]).

Mono-peroxy-chloroniobates $A_2[Nb(O_2)Cl_5]$ with A = K, NH₄ or Et₄N were found to react with HF in CH₃CN giving $A_2[Nb(O_2)F_5] \cdot n$ HF, with n = 2-5 [107]. In the presence of various acido ligands, $A_2[Nb(O_2)F_3L_2]$ (A = K, Cs, NH₄ or Et₄N; L = NCS, OAc, CF₃CO₂, HCO₂) and $A_3[Nb(O_2)F_2L_2]$ ($L' = SO_4$ and C₂O₄) were formed. The complexes were characterized by IR spectroscopy [107].

The di-peroxy-compound K₃[Ta(O₂)₂F₄] was shown to undergo UV-photodecomposition [108]. The kinetics of oxygen evolution obey a parabolic rate equation, indicating a monoexcitation process for the photolysis [108]. The activation energy was determined and the decomposition given as

$$K_3[Ta(O_2)_2F_4](s) \longrightarrow K_3[TaO_2F_4](s) + O_2(g)$$

according to chemical analyses, IR spectroscopy and thermogravimetry [109].

2.4 Nb(V) and Ta(V) chalcogeno-halide complexes

Niobium sulphide *tri*chloride, NbSCl₃, has been obtained by reacting NbCl₅ with hexamethyldisilthiane, Me₃Si-S-SiMe₃, in dichlormethane at room temperature [110]. The reaction is fast and gives a precipitate which does not react further. The NbSCl₃ product was characterized by IR spectroscopy as a coordination polymer containing Nb-S-Nb bonds, in contrast to other kinds of NbSCl₃ which contain Nb=S bonds [110].

In the solid state at 100 °C and under vacuum, Nb₂Cl₁₀ reacts with B₂S₃ to form NbSCl₃ [111]. In CH₂Cl₂ solution, NbSCl₃ and P(Ph)₄Cl form crystals of [P(Ph)₄][NbSCl₄], which contain quadratic-pyramidal [NbSCl₄] ions with a 2.09 Å long Nb=S bond, according to the X-ray structure determination [111], see Figure 7. An improved synthetic procedure to obtain [P(Ph)₄][NbSCl₄] has recently been described [112]. IR spectra of the crystals are discussed in reference [111]. By hydrolysis of [P(Ph)₄][NbSCl₄] in the presence of POCl₃ in CH₂Cl₂, [P(Ph)₄]₂[NbOCl₄(O₂PCl₂)]·CH₂Cl₂ is formed [113].

Presumably, solid NbS₂Cl has been obtained from NbS₂Cl₂ by electrochemical oxidation at 160-200 °C in a cell using a NaAlCl₄ molten salt as electrolyte [114]. The structure of NbSBr₃ was investigated by electron diffraction [35]. The reactivity of NbSBr₃ and NbSeBr₃ with respect to the formation of complexes was studied [115].

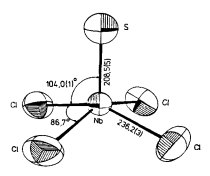


Figure 7. The X-ray structure of the quadratic-pyramidal [NbSCl4] ion in [P(Ph)4][NbSCl4]. Reproduced with permission from [111], U. Müller and P. Klingelhöfer, Z. Anorg. Allgem. Chem. 510 (1984) 109.

The preparation of NbSCl₃ and some thiohalogeno compounds containing [NbSCl₅]²⁻, [TaSCl₅]²⁻ and [NbSBr₅]²⁻ will soon be described [115A]. The crystal structures of $(PPh_4)_2[NbSCl_5]\cdot 2(CH_2Cl_2)$ and $NEt_4[NbCl_6]$ were solved [115A].

2.5 Nb(V) and Ta(V) halide complexes with O-donor ligands

2.5.1 Fluoride complexes with O-donor ligands

According to 19 F NMR spectral data, tantalum*penta*fluoride TaF5 reacts with O,O-diphenyl-N-benzoylamidophosphate, (PhO)₂P(O)NHC(O)Ph, in methylene chloride solution to form essentially a dimeric complex cation [F5Ta-O-P+(OPh)₂-NH-C+(Ph)-O-TaF5] and *mono*meric complex cations, [F5Ta-OP+(OPh)₂NHC(O)(Ph)] and [F₄Ta{-OP+(OPh)₂NHC(O)(Ph)}₂], balanced by TaF6⁻ anions [116]. The *mono*dentate *mono*meric cation forms [F₅Ta{-OP(OPh)₂=NC(O)(Ph)}] by splitting off one proton from the NH group [116].

2.5.2 Chloride complexes with O-donor ligands

The [NbOCl₄(THF)]⁻ (THF = tetrahydrofuran) anion has been characterized by spectroscopy and crystallography [117]. It has two oxygens in trans positions and four chlorines in a nearly octahedral configuration.

By hydrolysis of [P(Ph)4][NbSCl4] [111] in the presence of POCl3 in CH₂Cl₂ solution, [P(Ph)4]₂[NbOCl₄(O₂PCl₂)]·CH₂Cl₂ was formed [113]. Attempts to prepare the compound directly from Nb₂Cl₁₀, POCl₃, P(Ph)₄Cl and H₂0 yielded crystals of [P(Ph)₄][NbOCl₄(OH₂)] [113]. Crystal structures of both compounds, determined by means of X-ray diffraction data, show quadratic-pyramidal [NbOCl₄]⁻ ions, to which a molecule of either H₂O or a [PO₂Cl₂]⁻ ion is attached in *trans*-position to the O atom, see Figure 8. IR spectra of the crystals are discussed in reference [113].

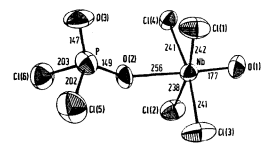


Figure 8. The X-ray structure of the [NbOCl₄(O₂PCl₂)]²⁻ ion in its [P(Ph)₄]⁺ salt. Distances in pm. Reproduced with permission from [113] P. Klingelhöfer and U. Müller, Z. Anorg. Allgem. Chem. 516 (1984) 85.

The tri-alkoxy di-chloride niobium(V) complexes, Nb(OR)₃Cl₂ (where $R = CH_3$, C_2H_5 , i-C₃H₇, and n-C₄H₉) have been prepared as lightly coloured compounds [118-119].

Reactions of $M(OPh)_5$ (M = Nb or Ta) with acetyl or benzoyl chlorides (AcCl or BzCl) in 1:1 or 1:2 molar ratio results in the replacement of phenoxy groups with chloride [120]. On heating, in molar ratio 1 to 3, however, compounds $MCl_3(OPh)_2RCO_2Ph$ and in molar ratio 1 to higher than 4, $MOCl_3RCO_2Ph$ were formed, with $R = CH_3$ (methyl) or C_6H_5 (phenyl) [120].

Complex formation among NbCl₅ and oxygen-containing donors D in organic solvents was studied by titration methods [121]. In the non-solvating solvent benzene, the complexes NbCl₅D and NbCl₅D₂ were progressively formed, with $D = (C_4H_9O)_3PO$ and (iso- $C_5H_{11})_3PO$. In butyl acetate, in which 1:1 solvates with NbCl₅ already occur, the addition of oxygen donor molecules D resulted in the progressive formation of NbCl₅(CH₃COOC₄H₉)D, [NbCl₄(CH₃COOC₄H₉)D₂]Cl and [NbCl₄D₃]Cl complexes, with $D = (C_4H_9O)_3PO$, (iso- $C_5H_{11}O$ ₂(CH₃)PO, (C₈H₁₇)₂SO, (C₆H₁₃)₂SO and (iso-C₅H₁₁)₃PO. These D ligands are listed here in the order of increasing oxygen donor power [121].

NbOCl₃ reacts with acetamide ($L = H_3CO-NH_2$) in CH₃CN to give actamidates, 1:1 electrolytes NbOCl₃ L_n (n = 2.5, 3, 4, 6, 7), in which L is O-bonded [122]. In ethanol the compounds undergo alcoholysis to give NbOCl₂(OEt) L_4 (EtOH).

Complexes $[TaCl_nL_{5-n}]$ with n=2 and 3 and L=2,6-di-t-butylphenoxide have been obtained by letting TaCl₅ react with lithium 2,6-di-t-butylphenoxide in benzene [123]. The compounds were characterized by elemental analysis, IR, and 1H and ^{13}C NMR spectra. The X-ray crystal structure was determined for the case of $[TaCl_2(2,6$ -di-t-butylphenoxide)₃]. This complex is a monomer with a square pyramidal coordination geometry around Ta. The phenyl rings (see Figure 9) are oriented such as to minimise interactions between 2- and 6- substituents. A comparison of the new structure with that of $[TaCl_3(2,6$ -di-t-butylphenoxide)₂] [124], is given in reference [123].

Figure 9. The coordination in square pyramidal [TaCl₂(2,6-di-t-butylphenoxide)₃], adapted after reference [123].

Lithium 2,6-di-isopropylphenoxide reacts with TaCl5 to give [TaCl2(diethylether)(2,6-di-isopropylphenoxide)3], which on recrystallization in air gives the binuclear [Ta2Cl(μ -Cl)2(2,6-di-isopropylphenoxide)5(μ -O)] [123]. The X-ray crystal structure was solved, to give the structure shown in Figure 10. Also, in this structure the phenyl rings are oriented such as to minimise interactions between 2- and 6- substituents.

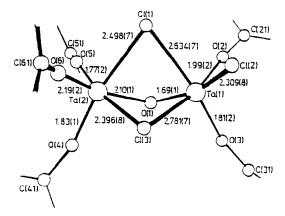


Figure 10. Inner coordination geometry from the crystal structure of dinuclear [Ta₂Cl(μ-Cl)₂(2,6-di-isopropylphenoxide)₅(μ-O)]. Reproduced by permission from [123], G. R. Clark, A. J. Nielson and C. E. F. Rickard, *Polyhedron* **6** (1987) 1765.

A series of chloro-chlorophenoxy salicylaldehyde complexes, NbCl_n(OC₆H₄Cl)_{4-n}·(sal) with n=0 to 3 can be prepared [125] by refluxing niobium *penta*chloride in the presence of o-chlorphenol and salicylaldehyde (salH). The compounds were characterized by several methods, and seem to contain octahedrally coordinated *mono*meric niobium. In one case, for NbCl₂(OC₆H₄Cl)·2(sal), evidence for a dimeric structure (see Figure 11), with eight-coordinated Nb was presented [125].

A series of chloro-chlorophenoxy α -hydroxyketone complexes of Nb(V), [NbCl_n(OC₆H₄Cl)_{4-n}(L)] with n=0 to 3 and LH = benzoin (benzH) or 2-hydroxyacetophenone (hapH) was prepared by refluxing niobium *penta*chloride in benzene in the presence of o-chlorophenol and HL in predetermined molar ratios [126]. Complexes of the type [NbCl₂(OC₆H₄Cl)(L)₂] were likewise obtained. Structures of the complexes were assigned on the basis of elemental analysis, conductance, cryoscopic, magnetic and IR spectral studies. In the case of [NbCl_n(OC₆H₄Cl)_{4-n}(L)] they seem to contain octahedrally coordinated *mono*meric niobium. In the case of [NbCl₂(OC₆H₄Cl)(L)₂], a dimeric structure (see Figure 11) with eight-coordinated Nb was assumed [126].

$$\begin{array}{c|cccc}
C_6H_4CI & & & \\
O & CI & CI & O \\
O & Nb & O & & \\
O & CI & CI & O \\
O & CI & CI & O
\end{array}$$

$$L = O O$$

Figure 11. The estimated structure of NbCl₂(OC₆H₄Cl)(L)₂ dimers with eight-coordinated Nb. Chelate ligands (L) are salicylaldehyde [125], benzoin or 2-hydroxyacetophenone [126]. Adapted after [125] and [126], S. C. Chaudhry. J. Gupta and S. Mehta, Indian J. Chem. 23 A (1984) 1036, and 24 A (1985) 521.

The complexation reaction of 5,5'-methylene-bis-salicylaldehyde (MBS) with TaCl₅ was studied under various conditions (in DMF solution in dry nitrogen or ambient air or in oxalate solution) [127]. The 1:1 compounds obtained were diamagnetic and insoluble in common organic solvents. Probable structures (see Figure 12) were proposed on the basis of elemental analyses, magnetic measurements and infrared spectra [127].

Figure 12. Probable structures of tantalum MBS complexes [TaCl₃C₁₅H₁₀O₄] and [TaClC₁₅H₁₀O₅] with five and seven-coordinated Ta. Reproduced with permission from [127], S. Chomal, A. S. Aazmi and G. C. Shivahare, *Acta Chim. Hung.* 122 (1986) 127.

The preparation of *hetero*cyclic carboxylates of niobium(V) and tantalum(V) in dry benzene has been considered for the acids HL = HTCA = 2-thiophenecarboxylic acid and HL = HTAA = 2-thiopheneacetic acid:

Complexes of the type $MCl_{5-n}L_n$ were obtained, with M = Nb, Ta, n = 1-3 and L = TCA or TAA [128]. The complexes were characterized on basis of chemical analyses as well as IR and ¹H NMR data. A tentative seven-coordinated structure was assigned to the MCl_3L_2 complexes (2), with M = Nb or Ta and L = TCA or TAA [128].

$$\begin{bmatrix} R & & CI & & 2 \\ & & & CI & & 2 \end{bmatrix}$$

White crystalline compounds of the general formula NbCl₄(OOCR) and TaCl₄(OOCR) can be obtained from the *penta*chlorides and an equimolar amount of the carboxylic acids, RCOOH with R = H, CH₃, C₂H₅, n-C₃H₇, CH₂Cl, CHCl₂ and CCl₃. With carboxylic acid anhydrides (RCO)₂O with $R = CH_3$ and C₂H₅, brown *poly*meric compounds NbOCl(OOCR)₂ and TaOCl(OOCR)₂ were obtained. The compounds were characterized by chemical analysis and IR spectroscopy [129].

According to results [130] obtained by IR spectrophotometric, conductimetric and potentiometric methods and chemical analysis, the 7-nitroso-8-hydroxyquinoline-5-sulphonic acid, 3, has an ability to coordinate to Nb(V) forming green chelates such as [(C9N2O5SH4)Nb(H2O)2(Cl)2]Cl and Na[(C9N2O5SH4)2Nb(H2O)2(Cl)2]. The ligand is bonded to Nb through the oxygens of nitroso and hydroxy groups. Stability constants are reported [130].

Tantalum(V) complexes with citric and *tri*hydroxyglutaric acids have been studied [131].

Niobium(V) complexes with bis- β -diketones (L) were studied [132]. When L is 2-phenyl-1,1,3,3-tetra-acetylpropane (PTAP) or 2-phenyl-1,3-di-acetyl-1,3-di-benzoylpropane (PDADBP), 1:1 complexes (NbOCl(C₁₇H₁₈O₄) and NbOCl(C₂₇H₂₂O₄)) were obtained, according to analytic data and molecular weight determinations, together with IR, UV, 1 H-NMR and magnetic measurements. *Tri*meric structures of the type shown below, 4, were assumed [132].

2.5.3 Bromide complexes with O-donor ligands

New *tri*bromooxide Nb(V) complexes of the type NbOBr $_3$ ·2L, with $L = \text{Ph}_3\text{PO}$, CH $_3\text{CN}$, (CH $_3$) $_2\text{SO}$, ((CH $_3$) $_2\text{NO}$) $_2\text{CO}$ and C $_4$ H $_8$ S, and NbOBr $_3$ ·Et $_2$ S have been prepared [133].

2.6 Nb(V) and Ta(V) halide complexes with N-donor ligands

Novel compounds with the general formula $A_3[MNX_5]$ (M = Nb, Ta; A = K, Cs, Ph_4P , TEBA = triethylbenzylammonium; X = F, Cl) have been synthesized by heating dry mixtures of MCl_5 , NH_4Cl and ACl in an argon atmosphere [134]. There are no previous report on nitrido-complexes of niobium or tantalum containing a metal-nitrogen tripple bond, 5.

The new compounds were characterized by chemical analysis and by physico-chemical methods; the IR spectra contain a strong narrow band in the range 1000-1050 cm⁻¹ due to the *M*-N triple bond stretching [134].

Previously, the synthesis of (NH₄)₃[Ta₂NBr₁₀] by ammonolysis of TaBr₅ with NH₄Br under HBr formation has been described [135]. This unusual reaction in sealed glass ampoule at ca. 400 °C has been used to prepare also the compounds (NH₄)₃[Nb₂NBr₁₀] and (NH₄)₃[Ta₂NI₁₀] [94]. The X-ray crystal structures have been solved [94], showing linear symmetrical Nb=N=Nb bridges (see Figure 13) that also occur in the other compounds. The vibrational spectra of these red μ -nitrido halides were interpreted to contain $\vee_{as}(M_2N)$ at ca. 950 cm⁻¹ (IR) and $\vee_s(M_2N)$ at ca. 215 cm⁻¹ (Raman), the frequencies depending on the particular salt [94].

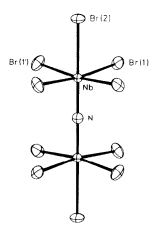


Figure 13. Structure of the [Nb₂NBr₁₀]³⁻ anion. The Nb-N bond distance is 1.845(2) Å. Reproduced with permission from [94], M. Hörner, K.-P. Frank and J. Strähle, Z. Naturforsch., 41 b (1986) 423.

The properties of Nb₆Br₉N₇ were studied and reactions leading to Nb₆Br₄(NH)₁₁(NH₂)₄ (with ammonia) and to Na₄Nb₆($C \equiv C$)₄(N)₈NH and Na₅Nb₆($C \equiv C$)₅(N)₅(NH)₅ (with sodium acetylide) were described [136]. Pyrolysis of these compounds gave Nb₄N₃ and NbC products.

Ammonolysis of NbBr₅ have been studied in order to obtain precursor compounds (e.g. NbBr(NH₂)₂NH) that can be thermally converted into advanced ceramic materials such as nitrides or carbonitrides [136]. One reaction was represented by

room temperature

6 NbBr₅ + 46 NH₃
$$\xrightarrow{}$$
 $\xrightarrow{}$ $\xrightarrow{}$ 220 °C

6 [NbBr₃(NH₂)₂·2NH₄Br·3.66NH₃] $\xrightarrow{}$ $\xrightarrow{}$ $\xrightarrow{}$ $\xrightarrow{}$ $\xrightarrow{}$ $\xrightarrow{}$ $\xrightarrow{}$ $\xrightarrow{}$ Nb₆Br₉N₇ + 21 NH₄Br + 18 NH₃

According to 19 F NMR spectra, tantalum *penta*fluoride reacts with *tri*-ethylamin (L) in methylene chloride solution to form a *mono*dentate N-donor complex [TaF₅(L)] [116], and with picolinic acid (LH) in acetonitrile solution to form *mono*dentate N-donor 1:1 and 1:2 six-coordinated complexes [TaF₅(LH)] and [TaF₄(LH)₂] [137]. The picolinic acid only coordinates *mono*dentately, see Figure 14.

Figure 14. Structure of $[TaF_5(LH)]$ for LH = picolinic acid [137].

Further molecular complexes of the general formula $[TaF_5L]$, where L is a N-donor ligand have been synthesized and characterized. For L = hexaphenoxycyclophosphazene $(N_3P_3(OPh)_6)$, the coordination takes place between Ta and one of the N-atoms of the N_3P_3 heterocyclic ring, according to NMR and IR spectroscopic results [18]. The complex is shown below, **6**.

The pentachlorides MCl_5 (M = Nb, Ta) were found to be reactive with trithiazyl chloride, (NSCl)₃ in CCl₄ suspension to give the donor-acceptor complexes $MCl_5 \cdot NSCl$ and $(MCl_5)_2 \cdot N_2S_2$. The products were characterized by chemical analysis and IR-spectra [59].

Tantalum *penta*chloride reacts with *bis(tri*methylsilyl)carbo*di*imide (CH₃)₃SiNCNSi(CH₃)₃ to give Si(CH₃)₃Cl and [Cl₄TaNCNSi(CH₃)₃]₂ [138]. IR and Raman spectra of the white compound, 7, were interpreted [138] to give a planar Ta₂N₂ structure ($R = \text{CNSi}(\text{CH}_3)_3$):

$$\begin{array}{c|cccc}
CI & R & CI \\
CI & N & CI \\
Ta & N & CI \\
CI & CI & CI
\end{array}$$

The *penta*chlorides of niobium and tantalum react with N(trimethylsilyl)-t-butylamine to give $[M(NCMe_3)Cl_3(NH_2CMe_3)]$ and with primary amines to form $[M(NR)(NHR)Cl_2(NH_2R)]$ which contain imido, amido and amine ligands coordinated to the same metal centre (M = Nb, Ta) [139-140].

In benzene, [Nb(NCMe₃)Cl₃(NH₂CMe₃)]₂ reacts with 4-picoline (pic) to give [Nb(NCMe₃)Cl₃(pic)₂], with 2,2'-bipyridyl(bipy) to give [Nb(NCMe₃)Cl₃(bipy)]·1/12 C₆H₆ and with N,N,N',N'-tetramethylethylenediamine (tmed) to give [Nb(NCMe₃)Cl₃(tmed)]. The tantalum complexes [Ta(NCMe₃)-Cl₃(bipy)₂]·1/3 C₆H₆, [Ta(NCMe₃)Cl₃(tmed)]·1/6 C₆H₆, [Ta(NCMe₃)(NHCMe₃)Cl₂(bipy)]·1/2 C₆H₆, [Ta(NEt)Cl₃(NH₂Et)]₂ and [Ta(NCHMe₂)(NHCHMe₂)Cl₂(bipy)] were also prepared. The complexes were characterized by melting points, chemical analyses and IR and NMR spectra, and suggestions of their geometry given [141]. The crystal structure of [TaCl(μ-Cl)(NBu^t)(NHBu^t)(NH₂Bu^t)]₂, a Ta(V) complex containing terminal imido, amido and amino ligands, was determined [139] (Bu^t is tert-butylamine), see Figure 15.

A methoxynitrene complex of tantalum(V), $[Cl_3(bpy)Ta \equiv NOMe]$, has been prepared and characterized [142]. In the presence of 2,2'-bipyridine (bpy), TaCl₅ reacted with O-methyl-hydroxylamine to form a red *monomer* complex, exhibiting an X-ray crystal structure with a distorted octahedrally coordinated tantalum atom [142], see Figure 16. The tantalum-nitrogen bond has a distance of 1.744 Å, and it gives an IR absorption band at 950 cm⁻¹.

It was shown that TaCl₅·(CH₃CN) and TaCl₃[O₂S(N:C(CH₃)Cl)₂] formed when SO₂ was passed through a solution of TaCl₅ in CH₃CN; and also isomerization and hydrolyzation reactions were observed, giving [TaOCl{N:C(CH₃)Cl}{OS(O)NHC(O)(CH₃)}] [143].

The preparation of organo-imido complexes of Nb and Ta from reactions of the *penta*halides with amines were studied [140] to obtain simpler and better synthetic procedures. Previously, *monoalkylamido* complexes $Ta(NHR)_2Cl_2(NH_2R)$ with R = Me, Et, Pr^n and Bu^n [144] and

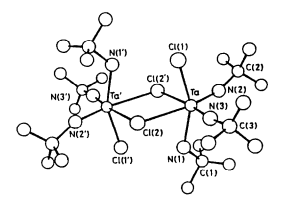


Figure 15. Crystal structure of the binuclear [TaCl(μ-Cl)(NBu^t)(NHBu^t)]2 complex. Reproduced with permission from [139], T. C. Jones, A. J. Nielson and C. E. F. Rickard, J. Chem. Soc. Chem. Comm. 1984 (1984) 205.

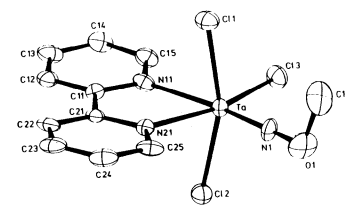


Figure 16. Structure of the [Cl₃(bpy)Ta = NOMe] complex. The Nb-N triple bond distance is 1.744 Å and the Ta-N-O angle 1740 (nearly linear). Reproduced with permission from [142], K. Haug., W. Hiller and J. Strähle, Z. Anorg. Allg. Chem. 533 (1986) 49.

Nb(NHR)₃Cl₂ with R = Me [145] have been claimed on basis of analytical data. Me₃SiNHCMe₃ and MCl₅ (M = Nb, Ta) react in dry benzene solution to form the imido complexes $M(\text{NCMe}_3)\text{Cl}_3(\text{NH}_2\text{CMe}_3)$ and these and other compounds were characterized by IR-, ¹H and ¹³C NMR-spectroscopy [140]. The compounds are expected to have octahedral

coordination, like what was the case for the one structure which has been solved, see Figure 17. Note the linearity of the Ta \equiv N - C bonding system. Ta-N_{imido} and Ta-N_{amido} bond lengths were 1.70 and 2.28 Å, respectively.

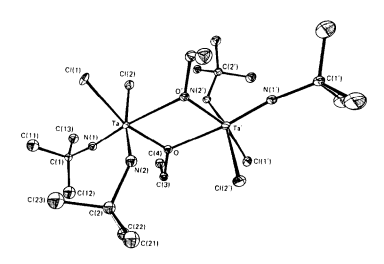


Figure 17. The molecular structure of white [Ta(NCMe₃)(μ-OEt)Cl₂(NH₂CMe₃)]₂. Hydrogen atoms have been omitted. Reproduced with permission from [140], P. A. Bates, A. J. Nielson and J. M. Waters, *Polyhedron* 4 (1985) 1391.

The reaction of TaCl₅ with Me₃SiNHAr (NAr = N-2,4-C₆H₃-(Prⁱ)₂) in the presence of donor solvents L provides the imido complexes [Ta(NAr)Cl₃L₂] in high yield (L = tetrahydrofuran, dimethoxyethane, pyridine, tetrahydrothiophene, etc.) [470]. The properties of these imido complexes and their reactions with alkynes were studied. The crystal structure was solved for the complex [Ta(NAr)(O-2,6-C₆H₃Me₂)Cl₂(py)₂], which turned out to confirm the octahedral geometry predicted by NMR data. The bonding in the complexes are discussed [470].

2.7 Nb(V) and Ta(V) oxyhalide complexes with N-donor ligands

NbOCl₃ reacts with acetoxime (HL, (H₃C)₂C=NOH) in CH₃CN to give oximates, NbOCl₃(HL)₂, NbOCl₂L(HL)₂ and NbOCl₂(HL), in which L^- is N-bonded [122].

Niobium(V) in hydrochloric and sulphuric acid media, in the presence of an excess of chloride or thiocyanate ions, reacts with 3-hydroxy-2-methyl-1-phenyl-4-pyridone (HR) to give complexes, Nb(OH)₃ClR or Nb(OH)₃(NCS)R, which are extractable into chloroform [146]. The

experimental conditions for quantitative extraction of niobium(V) into the organic phase are given, permitting a separation from zirconium(IV) and hafnium(IV). The identity of the complexes was determined spectrophotometrically.

Niobium *penta*chloride reacts with *tri*chloromethane, forming CCl₄ and NO⁺[NbOCl₄]⁻, which latter complex in acetonitril solution reacts with *tri*phenylmethylphosphonium chloride to form PPh₃Me[NbOCl₄(CH₃CN)] [91]. The crystal structure solution of this complex (and the IR and Raman vibrational spectra) show that the niobium is octahedrally coordinated with O and N-bonded CH₃CN in *trans* positions to each other, see Figure 18.

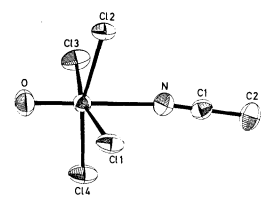


Figure 18. The crystal structure of the PPh₃Me[NbOCl₄(CH₃CN)] complex. Reproduced with permission from [91], W. Hiller, J. Strähle, H. Prinz and K. Dehnicke, Z. Naturforsch. 39 b (1984) 107.

Substituted 2-hydroxy-benzalbenzoyl hydrazones (L) react with NbCl₅ forming complexes of the type Nb(L)₂Cl [147]. Infrared spectra were interpreted to indicate (I) enolisation of the C=O followed by deprotonation and complexation with Nb, and (I) azomethine nitrogen complexation with Nb and (I) deprotonation of phenolic OH and complexation with Nb. The central metal ion acquires a coordination number of seven, see Figure 19 (I).

Tantalum(V)-[bis-(8-hydroxyquinoline)] coordination compounds [TaCl3(BHQ)], [TaOCl(BHQ)] and [TaO(HBHQ)3], with $H_2(BHQ) = bis$ -(8-hydroxyquinoline) = $C_{18}H_{10}(OH)_2N_2$, are yellow and insoluble in common organic solvents [148]. Probable structures of the three complexes are given, based on chemical analyses, magnetic measurements, IR spectra and thermograms. The O and N atoms of ligands coordinate to tantalum, forming chelate rings and giving tantalum coordination numbers of 7, 6 and 7, respectively [148].

Dihydroxychalcones, i.e. substituted 2,2'-dihydroxy-benzalacetophenones (L), react with NbCl5 forming complexes of the type Nb(L)3Cl2 [147]. Infrared spectra were interpreted to indicate complexation to niobium in such a way that the metal acquires a coordination number of eight, see Figure 19 (B).

$$R \xrightarrow{CH=N-N=C} CH = N-N=C$$

$$O = C$$

$$CH = N-N=C$$

$$CH = N-N=C$$

$$CH = CH(C_6H_4OH)$$

Figure 19. The estimated structure of A: benzalbenzoyl hydrazone *mono*mer complexes Nb(L)₂Cl with seven-coordination; B: dihydroxychalcone monomer complexes Nb(L)₃Cl₂ with eight-coordination. Reproduced with permission from [147], N.S. Biradar, V.L. Roddabasanagoudar and T.M. Aminabhavi, *Indian J. Chem.* 24 A (1985) 703.

In dimethylformamide (DMF), niobiumpentachloride and 1-phenyl-3-methyl-4-(2-methoxybenzeneazo)-5-pyrazolone (PMMBP, an azo-dye) were found to react in a ratio of 1:2 [149]. Based on chemical analysis and IR-spectra the following possible structure, 8, was proposed for the complex.

The complexation reaction of 5,5'-methylene-bis-salicylaldoxime (MBSO) with TaCl₅ was studied under various conditions (in DMF solution in dry nitrogen or ambient air or in oxalate solution) [127]. The 1:1 compounds obtained were diamagnetic and insoluble in common organic solvents. Probable structures (see Figure 20) were proposed on the basis of elemental analysis, magnetic measurements and infrared spectra [127].

$$\begin{bmatrix}
HO & H & H & OH \\
I & I & I & OH \\
N=C & C=N & O \\
O & CH_2 & O & I \\
C & CH_2 & O & I \\
O & CH_2 & O & O \\
O & CH_2 & O & O \\
O & O & O$$

Figure 20. Probable structures of tantalum MBSO complexes [TaCl₃C₁₅H₁₂O₄N₂] and [TaClC₁₅H₁₂O₅N₂] with five and seven-coordinated Ta. Reproduced with permission from [127], S. Chomal, A. S. Aazmi and G. C. Shivahare, *Acta Chim. Hung.* **122** (1986) 127.

The tantalum(V) coordination polymers of 5,5'-methylene-bis-salicylaldehyde-aniline Schiff-base (MBSASB), 5,5'-methylene-bis-salicylaldehyde-o-phenylenediamine Schiff-base (MBSOPDSB) or 5,5'-methylene-bis- salicylaldehyde-triethylenetetramine Schiff-base (MBSTETASB) have been prepared and characterized [150]. The Schiff-bases are double-acting bidentate ligands which coordinate to tantalum via nitrogens and oxygens on both sides. Tantalum is in turn 6 or 7 coordinated and form polymeric chains:

$$\cdots$$
- Schiff-base - $\text{TaO}_{\chi}\text{Cl}_{\gamma}$ - Schiff-base - $\text{TaO}_{\chi}\text{Cl}_{\gamma}$ - \cdots

2.8 Nb(V) and Ta(V) thiohalide complexes with N-donor ligands

NbSBr₃ reacts with ligands to give NbSBr₃·2L (for ligands $L = \text{CH}_3\text{CN}$, py, 1,4-thioxane, tetrahydrothiophene), NbSBr₃·3py, and NbSBr₃·bpy (py = pyridine, bpy = 2,2'-bipyridine) [115]. NbSeBr₃ reacts only with CH₃CN to give NbSeBr₃·2CH₃CN. The complexes were

characterized by IR, ¹H NMR, and diffuse reflectance spectra and electrical conductivity [115].

Complexation of thiopicolinamides TPMA and TPA with niobium(V) was studied in dry DMF-methanol medium (TPMA = thiopicolinmethylamide and TPA = thio-picolinanilide) [151]. Insoluble nine-coordinated bi-nuclear [Nb₂Cl₄L₆] compounds were formed, having two Cl bridges and three pairs of NS coordinated chelate rings, according to chemical analyses and IR-spectra, see Figure 21.

Figure 21. Proposed structures of thiopicolinanilide and thiopicolinmethylamide nio-bium(V) complexes. Reproduced with permission from [151], R. K. Sharma and G. C. Shivahare, *J. Indian Chem. Soc.* 62 (1985) 18.

2.9 Nb(V) and Ta(V) halide complexes with C-donor and other ligands

Because of the short Nb-C and Ta-C bond lengths observed experimentally, the complexes in this section can be taken to be in the oxidation state +V.

The d^{O} - d^{O} ditantalum complex [TaCl₂(SMe₂)(N-C₆H₅)]₂(μ -Cl)₂ can be prepared from the reaction of Ta₂Cl₆(Me₂S)₃ and azo-benzene in benzene-toluene solution (like the *iso*-structural niobium compound) [152]. Single crystal X-ray data show a *di*nuclear compound, adopting a distorted edge-sharing *bi*-octahedral structure, see Figure 22. The Ta=N-Ph distance is 1.747(8) Å. IR data are given [152].

After the first characterization in 1980 of a η^2 -alkyne complex (the anion [TaCl₄(py)(PhCCPh)]⁻ with py = pyridine and PhCCPh = diphenylacetylene), several similar complexes have now been described. By use of NbCl₄(THF)₂ (THF = tetrahydrofuran) as a starting material, the following compounds have been obtained and characterized by means of their crystal structures (PhCCMe = 1-phenyl-1-propyne): NbCl₃(PhCCPh) [153],

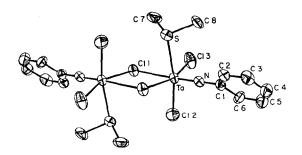


Figure 22. Drawing of the [TaCl₂(SMe₂)(N-C₆H₅)]₂(μ-Cl)₂ molecule. Reproduced with permission from [152], J. A. M. Canich, F. A. Cotton, S. A. Duraj and W. J. Roth, *Polyhedron* 5 (1986) 895.

and NbCl₃(PhCCPh)(THF)₂ [154], NbCl₃(PhCCMe)(THF)₂ [154], [Mg₂Cl₃(THF)₆][NbCl₄(PhCCPh)(THF)]·0.5(THF) [154], and Nb₂OCl₄₃(PhCCPh)(THF)₄ [154]. The structures are illustrated in Figures 23 and 24. It is evident that a sevenfold coordination around the metal atom is preferred. Note also the threefold coordination around the Cl₁ ligands in Figure 23.

Silaacyl derivatives of tantalum have been discovered: $(Cp^*)Cl_3Ta[\eta^2 - COSiMe_3]$, with $Cp^* = \eta^s - C_5Me_5$ and Me = methyl [155-156], $(Cp^*)Cl_3Ta[\eta^2 - OCLSiMe_3]$ where the Lewis donor L binds to the η^2 -silaacyl carbon atom [156], $(Cp^*)Cl_3Ta\{\eta^2 - OC(SiMe_3)[P(OMe)_3]\}$ [157] and $(Cp^*)Cl_2Ta(\eta^4 - OC(SiMe_3)P(OMe)_2O$ [157]. X-ray crystal structure, as well as spectroscopic and reactivity data of these six and seven coordinate complexes are given.

2.10 Nb(V) and Ta(V) pseudo-halide complexes with oxy-donors

Tri-alkoxy di-pseudohalide niobium(V) complexes, Nb(OR)3(Ps)2 (where $R = CH_3$, C₂H₅, i-C₃H₇, n-C₄H₉ and $Ps = CN^-$, NCO⁻, NCS⁻ and N₃⁻) have been prepared using chloride/pseudohalide exchange in hot dry tetrahydrofuran solution. The sixteen light-yellow to dark-brown compounds were characterized as Nb(OR)3(Ps)2 on the basis of chemical analyses and IR spectral data [118].

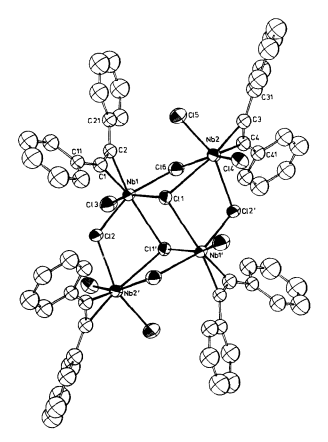


Figure 23. Structure of the tetrameric alkyne niobium(V) complex: (NbCl₃(PhCCPh))₄ [153]. Reproduced with permission from [153], E. Hey, F. Weller and K. Dehnicke, Z. Anorg. Allgem. Chem. 514 (1984) 25.

Also, belonging to the above series, seven tri-phenoxy di-pseudohalide niobium(V) complexes, Nb(OR)₃(Ps)₂ (where $R = C_6H_5$ (phenyl) and $Ps = CN^-$, NCO $^-$, NCS $^-$, N₃ $^-$, NH₂ $^-$ and C9H₆NO $^-$ (oxine)) were prepared using chloride/pseudohalide exchange in hot dry acetonitrile or tetrahydrofuran solution [158]. These brown compounds were characterized as Nb(OR)₃(Ps)₂. Analytical data, conductance, and IR and pronton NMR spectral data were reported [158]. A monomeric trigonal bipyramidal structure, 9, was assumed:

$$Ps$$
 OR
 Ps
 OR
 Ps

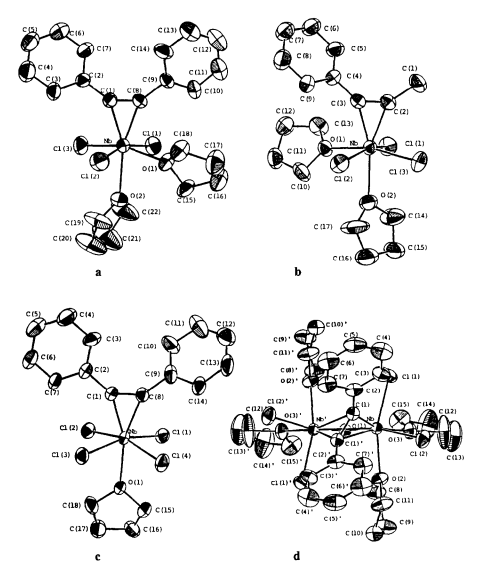


Figure 24. of niobium(V) complexes: Structures new alkyne (a) NbCl3(PhCCMe)(THF)2, NbCl3(PhCCPh)(THF)2, (b) = (c) [Mg2Cl3(THF)6][NbCl4(PhCCPh)(THF)]·0.5(THF), (d) Nb2OCl43(PhCCPh)(THF)4 [154]. The crystal structures a. Reproduced with permission from [154], F. A. Cotton and M. Shang, Inorg. Chem. 29 (1990) 508.

Reactions of acetyl *iso*thiocyanate, CH₃CONCS, with *iso*propoxides of Nb(V) and Ta(V) in dry benzene give products of the type $M(OPr^i)_{5-x}(NCS)_x$, x=1 to 4 [159]. On the basis of IR spectra and molecular weight determinations, it was concluded that the *mono*- and *di*-thiocyanato derivatives are *di*meric with double alkoxy bridging; no information is given on the higher thiocyanates.

The tri-alkoxy (or tri-phenoxy) di-chloride niobium(V) complexes, Nb(OR)₃Cl₂, where (OR)₃ = (OCH₃)₃, (OC₂H₅)₃, (i-OC₃H₇)₃, (n-OC₄H₉)₃, (OC₆H₅)₃, [(OC₆H₄O) + (OC₆H₄OH)], [(O₂C₆H₃OH) + (OC₆H₃(OH)₂)], react with NaNO₂, AgNO₂ or AgNO₃ in dry tetrahydrofuran forming Nb(OR)₃L₂ with L = NO₂, ONO⁻ or ONO₂- [119]. The 21 complexes formed were characterized by chemical analysis and IR spectroscopy [119].

New *pseudo*halide, oxine and amido derivates of niobium(V) *di*chloro *tri*-catecholoate or *tri*-pyrogallolate complexes, [Nb(Ps)₂(X)], where $Ps = CN^-$, NCO $^-$, NCS $^-$, N3 $^-$, NH2 $^-$ and C9H6NO $^-$ (oxine) and X = (OC6H4O)(OC5H4OH) (*tri*-catecholate) or (OC6H3O-HO)(OC5H3(OH)2) (*tri*-pyrogallolate), were isolated and characterized by chemical analysis and IR spectra [160]. Also, these 12 coloured solids are *mono*meric with an assumed trigonal *bi*pyramidal coordination geometry.

2.11 Oxides of niobium(V) and tantalum(V)

Niobium and tantalum oxides are numerous, and structurally very complicated and elaborate. Columbite-tantalite series of minerals of general composition (Fe,Mn)(Nb,Ta)₂O₆ and pyrochlore minerals are in this respect no exception. The pentoxides and many other phases are built of MO₆ octahedra sharing edges and corners, but this can be (and is) done in an almost unlimited number of ways (block structures). By partial oxygen loss various subvalent phases are formed, e.g.Nb₂2O₅4, NbO₂, NbO and so on.

The presence of the molecules Nb₂O₄, Nb₂O₅, Nb₄O₉, and Nb₄O₁₀ were identified in the gas phase over lithium niobate [161]. The molecules were characterized thermochemically (atomization energies and standard heats of formation) [161].

Defect structures in non-stoichiometric Nb₂O₅ was treated statistically [162]. At high temperatures and pressures, Nb₂O₅ and Ta₂O₅ solids may be reduced by Ta metal foil to form NbO₂ and Ta₀O₂ phases [163]. Shock-induced phases in Nb₂O₅ were studied by X-ray diffraction analysis and high-resolution electron microscopy [164-165].

Recrystallization experiments in closed silica glass ampoules with H-Nb₂O₅/NbCl₅ mixtures gave crystals of N-Nb₂O₅, B-Nb₂O₅ and T-Nb₂O₅ depending on circumstances [166].

The specific heat of the high temperature modification of α "-Ta₂O₅ were determined at 10-320 K [167]. Thermodynamic properties were reported. A phase transition was observed at 215.3 K [168]. The thermodynamic properties of rhombic β -Ta₂O₅ were redetermined by the e.m.f. method at high temperatures [169], and below room temperature by the method of adiabatic calorimetry [170]. The dependence of the electronic structure of Ta₂O₅ on coordination number and symmetry of nearest coordination sphere around Ta was determined [171]. A high temperature phase transition in Ta₂O₅ were found [172].

X-ray photoelectron spectroscopic analysis of 4 keV Ar⁺-bombarded Nb₂O₅ and NaNbO₃ has revealed the formation of lower-valence oxides and even metallic niobium [173].

The structure of Nb₂O₅ phases [174] and the many Nb₂O₅, LiNbO₃ and NaNbO₃ polymorphs have been described, *e.g.* in relation to the phases within the Li₂O-Nb₂O₅ system [175-177], the Li₂O-Nb₂O₅-Ta₂O₅ system [178].

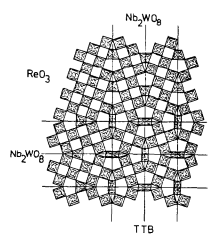


Figure 25. Twin planes relating the structure of ReO₃-type to the framework structures of Nb₂WO₈ and the tetragonal tungsten bronze (TTB), Na_xWO₃ [179]. Reproduced with permission from *Chemica Scripta* **26** (1986) 547, article by B.-O. Marinder.

The NaNbO₃-Nb₂O₅-WO₃ system was reviewed [179]; very complicated structures occur: Na₂Nb₄O₁₁, NaNb₃O₈, Na₂Nb₈O₂₁, NaNb₇O₁₈, NaNb₁₃O₃₃ [180], WNb₁₂O₃₃, W₃Nb₁₄O₄₄, W₄Nb₂₆O₇₇, Nb₂WO₈, Nb₂₆W₂₆O₁₄₃ [181-182], just to mention a few. Also,

Na₂Ta₄O₁₁ and Ag₂Ta₄O₁₁ were characterized [183-184]. Many of these structures are based on cumulated blocks of octahedra like in ReO₃, or related to the tetragonal tungsten bronze (TTB) structure, known from *e.g.* Na_xWO₃. One typical example is shown in Figure 25 for the case of Nb₂WO₈ [179].

A powder sample of LiNbO₃ was investigated by ⁹³Nb NMR two-dimensional spectroscopy [70]. LiTaO₃ single crystals were studied by Raman spectroscopy in ferro- and *para*-electric phases and extensive assignments of observed bands were given [185]. A lithium insertion compound (Li,Cu)TaO₃ with the LiNbO₃ structure was prepared [471]. AgNbO₃ single crystals were studied by Raman spectroscopy at different temperatures, revealing a complex sequence of phases [186].

Polymorphism at high temperatures was studied in the oxides Li₃NbO₄ and Li₃TaO₄ having nearly rock-salt structure [81], and these salts were characterized by X-ray powder diffracion and infrared spectroscopy [80, 82]. Comments were given on the formation of LiNb₃O₈ [176]. The binary systems Li₂O-M₂O₅ were reinvestigated and phases like *e.g.* Li₁6M₄O₁₈ (*M* = Nb or Ta) [177], Li₂Nb₆O₁₆ [187] and Li₂Nb₂8O₇₁ [188] found. Compound formation among Li₃NbO₄ and Li₂WO₄ or LiNb₃O₈ and TiO₂ was studied in the solid state [189-190]. The reaction of Ta₂O₅ and Li₂CO₃ was studied by mass spectra, thermogravimetry, and X-ray phase analysis to establish the mechanism of the LiTa₃O₈ formation [191]. If tantalic acid was used in stead of Ta₂O₅, similar results were obtained [192]. The *ortho*tantalate Li₃TaO₄ has been protolysed to form Li_{1.2}H_{1.8}TaO₄ and Li_{1.1}H_{1.9}TaO₄ having rock-salt related structures [193]. Proton motion in solid HNbO₃ and HTaO₃ (obtained by H⁺-Li⁺ ion exchange) was determined by proton NMR relaxation [194]. The electric conductivity of HNbO₃ powder was measured by impedance spectroscopy [195]. LiNbO₃ and LiTaO₃ phases have been substituted in part with the pair Cu(II)-Ti(IV) [196]. Ferroelectric tungsten-bronze single crystals K₃Li₂Nb₅O₁₅ have been grown [197].

NaTaO₃ and Na₂Ta₄O₁₁ phases were prepared by decomposition of pyrochlores obtained under hydrothermal conditions [183].

To the already known structures of crystals formed from mixtures of potassium oxide and niobium pentoxide, (e.g. KNbO3, KNb3O8 and K8Nb18O49 and the long list of non-stoi-chiometric K2O:Nb2O5 mixtures), K3Nb7O19 has now been added. It forms white triclinic crystals with a structure, solved by X-ray diffraction methods, consisting of edge-shared pairs of octahedra. As Figure 26 shows, the pairs of octahedra are corner-shared to one another to form double strings seven pairs long, and the strings are corner-shared to other strings forming tunnels [198]. A high-resolution electron microscopic study of the KNbO3-Nb2O5 system has confirmed the existence of layer structures K4Nb6O17 and L-KNb3O8, three tetragonal tungsten-bronze related structures and a block structure KNb13O33 [199].

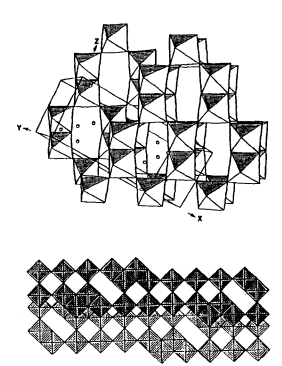


Figure 26. View of the structure of K₃Nb₇O₁₉ showing one of the sets of tunnels. Open circles represent K in the unit cell. Also shown is an idealized diagram of the block structure. Reproduced with permission from [198], G. D. Fallon, B. M. Gatehouse and L. Guddat, J. Solid State Chem. 61 (1986) 181.

The compounds LiNbWO₆ and LiTaWO₆ have been protolysed to form HNbWO₆ and HTaWO₆ in hot aqueous H₂SO₄ [200-201]. The reaction is accompanied by a structural transformation from the rutile to the ReO₃ structure. Compounds were characterized by chemical analysis, X-ray powder diffraction, thermogravimetry and IR spectroscopy [200].

Methods for synthesis of antiferroelectric bismuth orthotantalate, BiTaO₄, [202] and diclectric antimony orthotantalate, SbTaO₄ [203], were studied.

In the Cu_2O - Ta_2O_5 system, $Cu_5Ta_{11}O_{30}$ and $Cu_3Ta_7O_{19}$ phases were prepared *in vacuo* at high temperatures and studied by X-ray diffraction [204]. They have similar structures, based on layers of α - U_3O_8 type (containing pentagonal *bi*pyramids) and layers composed of octahedra, only differing in the sequence of these layers [205].

The CaTa₂O₆ and CaTa₄O₁₁ phases were prepared at high temperature and studied by high-resolution transmission electron microscopy. The resulting structure was compared with computer simulated images [205]. In CaTa₂O₆ tantalum is octahedrally, in CaTa₄O₁₁ pentagonal *bi*pyramidally and octahedrally coordinated [205]. X-ray powder diffraction data were given for *mono*clinic 2TiO₂·5Nb₂O₅, prepared by simultaneous hydrolysis of titanium and niobium alkoxides [206].

The crystal structure has been found for a new kind of lamellar niobate, Ba(Nb₃O₈)₂·H₂O. The layers are similar to those of KNb₃O₈, but one layer out of two is displaced [207].

Iron niobate crystals form an ordered super-structure of α -PbO₂, the Columbite-type, FeNb₂O₆, see Figure 27(c), according to a neutron powder diffraction stydy (Rietveld profile-refinement method) [209]. During preparation, it appears to be essential to control the partial pressure of oxygen; low pressures favour the formation of *tri*-rutile structure phases with high content of Fe²⁺, according to X-ray and Mössbauer data [210]. The *tri*-rutile FeNb₂O₆ structure was solved by single crystal X-ray diffraction and found to consist of chains of MO₆ octahedra (M = Fe or Nb, disordered), in which each octahedron shares a pair of opposite edges [211]. The *tri*-rutile structure was confirmed for FeTa₂O₆ in an investigation covering X-ray and neutron diffraction, magnetic susceptibility, heat capacity and Mössbauer techniques [212]. The magnetic structures of this Fe(II)-compound was also described. Iron tantalates FeTaO₄, FeTa₂O₆ and Fe₄Ta₂O₉ have a considerable range of non-stoichiometry [213].

Crystals of Sr₄FeTaO₈ and Sr₄CoTaO₈ with K₂NiF₄ structure were examined by X-ray and electron diffraction and magnetic susceptibility measurements [214]. The crystals were unstable in air. The structures and electronic spin states were discussed [214]. Rutile phases related to CrNbO₄ (which contains Nb(V)) have been prepared and characterized in terms of the structural, magnetic and electronic properties [215]. Thermal interaction between Ta₂O₅ and Co₃O₄ at high temperatures, led to the formation of CoTa₂O₆ and other phases [216].

Single crystals of NiTa₂O₆ were prepared and investigated by X-ray methods [208]. The structural relationships were systematised between the different ordered and disordered AB₂O₆ and ABO₄ oxo-metallates, e.g. the α -PbO₂ and Columbite structures, see Figure 27. Mixed phases, Zn(Nb_xTa_{1-x})₂O₆ are known also [217].

In the CaO-Nb₂O₅-P₂O₅ system, CaNb₂O₆, Ca₃Nb₂O₈, β-Nb₂O₅, NbO₂ and NbO(PO₄) are formed at 1400 °C [218-219], and in the BaO-Nb₂O₅-P₂O₅ system, BaNb₂P₂O₁₁ is formed [220]. By heating, the latter compound decomposes into oxides and NbOPO₄. The crystal structures were solved for BaNb₂P₂O₁₁ (isomorphous with BaNb₂V₂O₁₁) and NbPO₅ (NbO₆ octahedra joined to PO₄ tetrahedra, forming a framework) [220]. Other phosphates are mentioned in a later section.

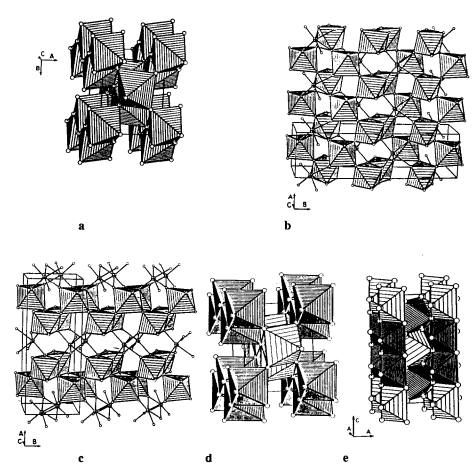


Figure 27. Crystal structures of (a): α-PbO₂, (b): ZnTa₂O₆ in the α-PbO₂ type structure with Zn²⁺ and Ta⁵⁺ octahedra (the latter are hatched), (c): ZnNb₂O₆ belonging to the Columbite type structure, with Nb5⁺ octahedra hatched, (d): FeNbO₄ in the Wolframite type structure, with Fe³⁺ and Nb⁵⁺ differently hatched and (e) NiTa₂O₆ in the rutile SnO₂ type structure, with Ta⁵⁺ densely hatched. Reproduced with permission from [208], H. Müller-Buschbaum and R. Wichmann, Z. Anorg. Allgem. Chem. 536 (1986) 15.

Compounds formed in the BaO-Nb₂O₅, BaO-Nb₂O₅-TiO₂, BaO-Nb₂O₅-V₂O₅, CaO-Nb₂O₅-P₂O₅ and BaO-Nb₂O₅-P₂O₅ systems have been studied by X-ray phase and structure analyses [218, 220-223]. In the BaO-Nb₂O₅ system, Ba₄Nb₂O₉ and Ba₅Nb₄O₁₅ were identified [221]. Compounds Ba₆Ti₁4Nb₂O₃₉, Ba₁4Ti₄0Nb₂O₉₉, Ba₁0Ti₂8Nb₂O₇₂

and $Ba_18Ti_54Nb_2O_{132}$ were isolated and their structures studied [222]. In the $BaO-Nb_2O_5-V_2O_5$ system, at 700-850 ^{O}C tetragonal $Ba_4Nb_2V_2O_{14}$ and rhombohedric $BaNb_2V_2O_{11}$ were formed, among others [223]. The crystal structure of the latter was determined. It contained NbO_6 octahedra linked through vertices, forming ReO_3 -type infinite layers of composition $Nb_2O_6^{2-}$ [223].

Compounds formed in the ZnO-Nb₂O₅ and ZnO-Nb₂O₅- A_2 O systems (A = Na, K) have been studied by X-ray phase analyses [224]. In the ZnO-Nb₂O₅ system, Zn₃Nb₂O₈, ZnNb₂O₆ and other phases were identified [224]. Compounds K₃Zn_{0.33}Nb_{7.67}O₂₁, K₂Zn₂Nb₄O₁₃ and K₂M₃Nb₃O₁₃ where M is Al, Cr, Ga, Fe, Sc, Y, or In were isolated and their properties studied [224].

Compounds formed in the PbO-Nb₂O₅-K₂O system have been studied by thermal and X-ray phase analyses, and a compound, $K_{2\chi}Pb_{1-\chi}Nb_2O_6$, was found [225]. The formation of sodium and potassium lead niobates, Na₂Pb₄Nb₁₀O₃₀ and K₂Pb₄Nb₁₀O₃₀, by the method of heating mixtures of Na₂CO₃ or K₂CO₃, Nb₂O₅ and PbO powders were studied by thermogravimetry and X-ray diffraction [226]. The K₂Pb₄Nb₁₀O₃₀ phase melts congruently at 1600 K [227].

A series of oxides, $AM_3O_9(M_2O_4)_n$, has been made and characterized by X-ray and electron diffraction. The series was derived from the rutile structure by chemical twinning (e.g., A = Na, K; M = Ba, Fe, Ti, Nb; n = 2 and 4) [228]. The *penta*valent oxidation state of niobium is counterbalanced by the presence of *tri*-valent ions (e.g., Ti(III), Fe(III)) in the rutile structure.

A topotactic dehydration of the lamellar oxide $HK_2Ti_5NbO_{14} \cdot H_2O$ to the oxide $K_4Ti_{10}Nb_2O_{27}$ was performed, and the process studied by electron and X-ray diffraction [229]. The structural model of $K_4Ti_{10}Nb_2O_{27}$ involved intergrowth of $K_3Ti_5NbO_{14}$ layers with the $K_2Ti_6O_{13}$ tunnel structure [229].

The compound LiNbGeO₅, recently prepared [230], have been found to give a strong blue luminescence when irradiated with UV light at low temperatures [231]. The crystal structure is similar to that of sillimanite (Al₂SiO₅) with octahedrally coordinated Nb+ 5 , distorted in a way similar to that of α -NbPO₅, i.e. with one shorter and one longer Nb-O distance (1.78 and 2.32 Å, respectively). The Raman spectrum of LiNbGeO₅ was reported [231].

The formation of calcium pyroniobate, Ca₂Nb₂O₇ (as well as Sr₂Nb₂O₇ and Ca₂Ta₂O₇), in a sodium chloride/sulphate melt at 800-850 °C was reported [232]. Substitutions in Ca₂Nb₂O₇, (Ca_X M_{1-X})₂(Ti_{1-X}Nb_X)₂O₇, with pyrochlore and perovskite-type layer structures, were also studied for M = La, Pr, Nd, Sm, Dy, Bi and 0 < x < 1 [233].

The self-diffusion of ⁸⁹Sr, ¹⁴⁰Ba and ⁹⁵Nb cations in strontium and barium niobates BaNb₂O₆, Sr₂Nb₂O₇, Sr₆Nb₂O₇ and Sr₅Nb₄O₁₅ has been studied in an effort to understand the solid-state high-temperature reactive-diffusion processes between SrCO₃

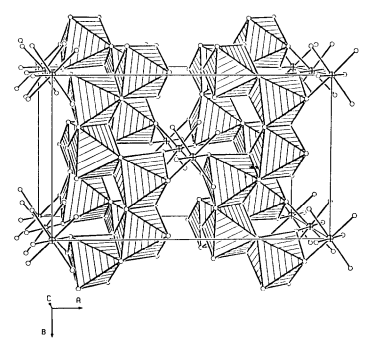


Figure 28. Crystal structure of Zn₄Ta₂O₉, showing edge-linked TaO₆ and Ta_{0.5}Zn_{0.5}O₆ octahedra (hatched) and chains of TaO₆ octahedra (not hatched). Reproduced with permission from [242], M. Waburg and H. Müller-Buschbaum, *Z. Anorg. Allgem. Chem.* **522** (1985) 137.

(SrO) and Nb₂O₅ [234]. The X-ray crystal structure of SrNb₂O₆ and SrNb₆O₁₆ was solved by the Rietveld powder diffraction profile technique [235]. The crystallization of BaNb₂O₆ was described in [236].

The luminescence in Sr₂Nb₂O₇ and Sr₂Ta₂O₇ were compared with other luminescing perovskite-like niobates and tantalates with structures based on corner-sharing NbO₆ and TaO₆ octahedra, and with the associated electronic delocalization playing an important role for the luminescense [237]. The X-ray powder diffraction diagrams of TiNb₂O₇, ZrNb₂O₇ and mixed phases thereof were given in [238].

The X-ray single crystal structure of Ni₄Nb₂O₉ was shown to consist of a three-dimensional framework of NiO₆-octahedra, between which isolated pairs of NbO₆-double octahedra [Nb₂O₉] are deposited [239]. The structure is much more complicated (480 atoms *per* cell) than those of similar compounds with Co, Fe, Mn or Mg substituting Ni; compounds which also have cumulated NbO₆-double octahedra.

Mixed oxides Nb₂Te₂O₉ and Ta₂Te₂O₉ were prepared by solid state reactions between oxides TeO₃ and Nb₂O₅ or Ta₂O₅. Crystal data, IR spectra and thermal decomposition processes were given [240]. The crystal structure of Ta₂Te₂O₉ was solved [241]. It contains two different kinds of TaO₆ octahedra, sharing oxygens with each other and with TeO₄ tetrahedra.

A new compound, Zn₄Ta₂O₉, was synthesized by high temperature reaction. The structure was solved and found [242] to consist of a Zn/O-network with incorporated one-dimensional TaO₆-chains, see Figure 28.

Also Sr₄Ta₂O₉ has been studied and it seems to have a complex perovskite super-structure which manifests itself on X-ray diffraction photographs and electron micrographs [243].

Single crystals of *mono*clinic Ta₂Te₂O₉ were grown from Ta₂O₅-TeO₂ melts [244]. The solved X-ray crystal structure showed infinite puckered layers of composition $(Te_4O_{12})_n$ alternating with layers of nearly regular TaO₆ octahedra sharing corners. The Ta-O bond distances ranged between 1.88 and 2.07 Å [244].

New complex F_{d3m} -pyrochlore $A_2(M'_{1-x}M''_{x})_2O_7$ type oxides have been made, with A=Y and $M'_{1-x}M''_{x}=Mn_2/3^2+Nb_4/3^5+$, $Mn_2/3^2+Ta_4/3^5+$, Mn_3+Nb_5+ or Mn_3+Ta_5+ . Oxidation states were obtained from the measured magnetic susceptibility, and hexagonally indexed powder X-ray diffraction patterns were given [245]. Bronzoid and pyrochlore phases $A_xNb_xW_{1-x}O_3$ have been characterized in the KNbO₃-WO₃ and CsNbO₃-WO₃ systems (A = K, Cs; x < 0.5) [246]. Pyrochlore type compounds $KMWO_6\cdot H_2O$ (M=Nb, Ta), or perhaps more correctly (KH₂O)(MW)O₆, were synthesized and characterized by thermogravimetry, differential scanning calorimetry and X-ray powder diffraction [247].

Complex oxides of the type $La(M'_{0.67}M_{0.33})O_3$, for M' = Mn, Co, Ni and Mg and M = Nb and Ta, with a perovskite structure, have been made and studied by X-ray diffraction [248].

The system La₂O₃-Li₂O-Nb₂O₅ was studied, and a new phase, La₃Li₇Nb₂O₁₃, was found [249]. X-ray diffraction diagrams of this compound and of La₂LiNbO₆ are given [249-250]. In the system La₂O₃-Li₂O-Ta₂O₅, two new phases, La₂LiTaO₆ and La₃Li₇Ta₂O₁₃, were identified by X-ray diffraction [250-251]. La₂LiNbO₆ and La₂LiTaO₆ are ordered *ortho*-rhombic perovskites, whose vibrational spectra are known [252].

In the system Bi₂O₃-Nb₂O₅, complicated ordered solid solution structures, being related to those of fluorite, pyrochlore and perovskite, were studied by high-resolution electron microscopy and electron diffraction in order to elucidate structural principles [253]. Very impressive pyrochlore-like units of cumulated octahedra were found [254]. A high resolution electron micrograph of Bi₄Nb₂O₁₁ was given in [255]. The systems Bi₂O₃-GeO₂-Nb₂O₅ and Bi₂O₃-GeO₂-Ta₂O₅ were studied separately [256].

In the BaO- R_2 O₃-Nb₂O₅ systems (R = trivalent Sc, In, Lu, Yb, Tm, etc.), a new 18R stacking polytype Ba₆R_{0.5}Nb_{4.5}O₁₈ has been prepared at ca. 1200 °C [257]. The stability of the new polytype decreases with increasing radius of the R ion (for Y, Tb, Gd, and Eu), but for R = Sm and Nd no 18R phase was observed. For R = Er, Ho, and Dy there was a small contamination with Ba₅Nb₄O₁₅. Unit cell data are given, based on X-ray diffraction [257]. Also, BaO- R_2 O₃-Ta₂O₅ systems [258] and systems containing titania have been studied. Perovskite-type structures were found, like e.g. Ba₈Ti₃ M_4 O₂₄ with M = Nb or Ta [259] and Ba₇Nb₄Ti₂O₂₁ [260]. A yellow 11H hexagonal perovskite Ba₁₁Re₇/₄Nb₇/₄W₇/₂O₃₂ has been characterized [261].

Lead zink niobate, Pb(Zn_{1/3}Nb_{2/3})O₃, is an excellent ferroelectric perovskite. It has been obtained by a sol-gel process involving the complex Zn[Nb(OEt)₆]₂, followed by an *in situ* sintering process [262]. Crystal growth and properties of Pb₂FeNbO₆ [263] and Pb₂ScTaO₆ [264] were studied.

The Nb-O bond lengths in NdNbO₄ and niobium-vanadium mixed NdNb_{1-x}V_xO₄ crystals of Fergusonite structure, see Figure 29, were studied by extended X-ray absorption (EXAFS) [265]. The Nb-O distances, 1.89 ± 0.03 Å, were unexpectedly almost independent of composition and essentially unchanged from the values exhibited in NdNbO₄.

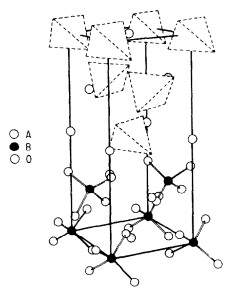


Figure 29. Tetragonal structure of NdNb_{1-x}V_xO₄ crystals. A = Nd, B = Nb or V. Reproduced with permission from [265], G. S. Knapp, M. V. Nevitt, A. T. Aldred and T. K. Klippert, J. Phys. Chem. Solids 46 (1985) 1321.

The high-melting fluorite-related rare earth tantalum oxides $3R_2O_3$ ·Ta₂O₅ or R_3 TaO₇, R = rare earth = La, Pr, Nd, Sm, Eu, Gd, Tb, Dy, Ho, Er, Yb have had their melting (solidification) points redetermined (range ca. 1900-2500 °C), and the correlation of the temperatures with rare earth ionic radii and crystal structures was examined [266]. Temperature and treatment dependent polymorphism in these oxides was studied by X-ray powder diffraction [267-268].

KNbUO₆, RbNbUO₆, CsNbUO₆ and TlNb₂UO_{11.5} single crystals were made from U₃O₈, Nb₂O₅ and the *mono*valent carbonates [269-270]. Their X-ray crystal structures were solved and show complicated systems of UO₇ and NbO₅ (or NbO₆) *poly*hedra sharing edges and corners and with the *mono*valent ions in tunnels or between slabs [269-270], see Figure 30.

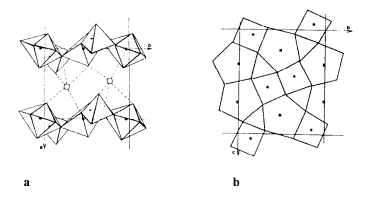


Figure 30. The CsNbUO₆ structure seen in two projections, showing (a) the layered structure and (b) the NbO₅ and UO₇ polyhedral network within a layer.Reproduced with permission from [270], M. Gasperin, Acta Cryst. C43 (1987) 404.

The compounds UNb₂O₇, UNb₄O₁₂ and UNb₃O₁₀ were examined by magnetic susceptibility, XPS and ESR methods and all were found to contain *penta*valent niobium [271].

The series of perovskite compounds $A^{I}A^{II}_{2}Nb_{3}O_{10}$ with $A^{I} = H$, Na, K, Rb, Cs and $A^{II} = Ca$ have been made and characterized structurally by X-ray diffraction [272].

Plate-like green crystals of α -PrNb₃O₉ have been obtained hydrothermally [273]. The structure was solved and shown to consist of double and single zig-zag chains of corner-shared distorted NbO₆-octahedra which sometimes also share edges, see Figure 31.

LaTa₃O₉ crystals were obtained and studied by X-ray and electron diffraction and electron microscopy [274]. The structure is containing ribbons of pentagonal TaO₇-bipyramids in one direction, connected by planes of TaO₆-octahedra. The tunnels formed in this way are occupied by La atoms [274].

Ternary perovskite oxides La(Mn2/3Nb1/3)O3, La(Co2/3Nb1/3)O3, La(Ni2/3Nb1/3)O3, La(Mn2/3Ta1/3)O3, La(Co2/3Ta1/3)O3 and La(Ni2/3Ta1/3)O3 have been made. IR spectra and X-ray diffraction crystal structures of these highly ordered compounds were studied [275]. Also, the crystal structure of La3(Ga5.5Nb0.5)O14 was solved [276].

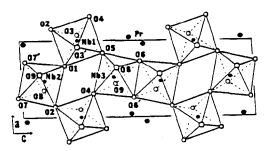


Figure 31. Projection of the view along α-PrNb₃O₉ structure showing connected NbO₆-octahedra and Pr³⁺ ions (small filled circles) in tunnels. Reproduced with permission from [273], C. C. Torardi, L. H. Brixner and C. M. Foris, *J. Solid State Chem.* 58 (1985) 204.

Thermal stability of rare-earth element niobates was studied in the high-temperature region [277]. Thermodynamic data, melting points and electric conductivity data were given for the following compounds: La₃NbO₇, Nd₃NbO₇, LaNbO₄, NdNbO₄, EuNbO₄, TbNbO₄, YNbO₄, NdNb₃O₉ [277].

The crystal structures of Sr_2RNbO_6 (R = Sm and Tm) have been determined and were found to be of a distorted cryolite-type containing octahedra sharing corners [278].

New hexagonal perovskite-like compounds $Sr_3LnNb_3O_{12}$ (with Ln = La, Pr or Nd) and $Sr_3LaTa_3O_{12}$ and $Sr_3PrTa_3O_{12}$ have been prepared by long-time sintering of mixtures of lanthanide oxides, niobium or tantalum oxides and strontium carbonate at 1350 °C [279]. The unit-cell parameters of the single-phase materials were determined [279]. Also, transparent $Ba_3LaNb_3O_{12}$ single crystals have been grown [280].

The crystal structure of (Cs_{0.75}K_{0.25})(Nb,Ti)U₂O₁₁ was shown to contain niobium-titanium pairs in *tri*angular *bi*pyramidal coordination joined by a corner [281].

High resolution electron microscopy was used to study the structures of K₇Nb₁₅W₁₃O₈₀ and Bi₄Nb₅O₁₈F [282]. The Fourier-processed images gave atomic coordinates near those of the refined X-ray structures.

Solid state reactions have lead to single crystals of new kinds of tetragonal tungsten bronze-types whose X-ray structures could be solved: The Ba₆FeNb₉O₃₀ structure has a statistical distribution of Fe³⁺ and Nb⁵⁺ in the corner-linked octahedral framework with *pentagonal* tunnels [283]. SrNb₆O₁₆ is *iso*structural with NaNb₆O₁₅F according to results found by the Rietveld powder diffraction profile technique [235]. A new compound, Ba₄FeTa₁₀O₃₀ with a tunnel structure is also related to NaNb₆O₁₅F [284]. The complicated structures of *e.g.* Ba₄CoTa₁₀O₃₀ [286], Ba₄MgTa₁₀O₃₀ and Ba₄NiTa₁₀O₃₀ have been determined [285]. The Ba₄CoTa₁₀O₃₀ structure conforms to the network of the Na₂Nb₁₂O₃₀F₂ type [286].

The PbO-Nb₂O₅-WO₃ system contains solid solutions of composition PbNb₂O₆·xWO₃ and Pb₄Nb₄O₁₁·xWO₃, both with the high temperature PbNb₂O₆ tetragonal tungsten bronze-type structure [287].

Heat treatments of mixtures of Ta₂O₅, WO₃ and W metal at high temperatures resulted in the formation of Ta₂WO₈ [288-289]. The structural properties of this and other bronzes, containing pentagonal tunnels, were studied by X-ray diffraction and by optical and electron microscopic methods [288-289].

Phase equilibria at 1300 °C have been studied for the SrO-La₂O₃-Nb₂O₅ system [290]. An indexed X-ray pattern was given for the Sr₃La₂Nb₁₂O₃₆ phase [290]. Analogous ternary lanthanide phases $Sr_3Ln_2Nb_{12}O_{36}$ with Ln = La - Lu were studied as well, and the crystal structure of $Sr_3Nd_2Nb_{12}O_{36}$ (tetragonal tungsten-bronze type) was solved [291].

In the system Ta₂O₅-La₂O₃, a new compound, LaTa₇O₁₉, has been made [292]. High resolution transmission electron microscopy and X-ray diffraction methods were used to study its crystal structure [292]. It can be considered to be built from two kinds of coordination *poly*hedra, octahedral TaO₆ and double layers of pentagonal TaO₇-bipyramids.

The products of the reaction between Eu_2O_3 and Nb_2O_5 in the presence of reducing agents like EuO, Nb, NbO, and NbO_2 have been characterized by X-ray L_{III} absorption spectroscopy [293]. The oxidation states were found to be Eu^3+ and Nb^5+ in Eu_3NbO_7 , whereas Eu_3NbO_6 contains also Eu^2+ . On the other hand, $EuNbO_3$ clearly was proven to contain Eu^2+ and Nb^4+ [293]. In the presence of SrO, $Sr_6Nb_2O_{11}$ and Eu_3NbO_6 may be formed, but a number of previously described Eu-Nb-O phases were proved to be non-existing [294].

Single crystals of $Sr_6Nb_34O_91$ were prepared by CO_2 -laser technique [295]. The structure, solved by X-ray diffraction techniques, consists of a network of corner-sharing NbO₆-octahedra similar to the known $[M_{10}O_{30}]^{12}$ - tetragonal tungsten-bronze (see Figure 32). The tunnels of the network are partly filled by Sr^2 +, Nb⁵+ and O^2 --ions.

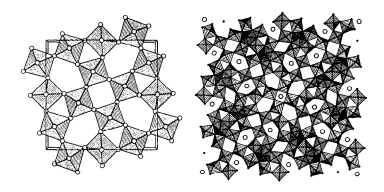


Figure 32. Perspective drawing of the $[M_{10}O_{30}]^{12}$ - tetragonal tungsten bronze network (a) and the new $Sr_6Nb_34O_{91}$ structure (b), both viewed along the direction [001]. Reproduced with permission from [295], K. Schückel and H. Müller-Buschbaum, *Rev. Chim. Minérale* 23 (1986) 154 and Gauthier-Villars, Publisher (Paris).

The compound La₃Ga_{5.5}Nb_{0.5}O₁₄, having the Ca₃Ga₂Ge₄O₁₄ structure, has been prepared during a study of the phase relations in the La₂O₃ - Ga₂O₃ - Nb₂O₅ system [296]. Also, the compounds GaNbO₄ and GaNb₁₁O₂₉ were formed.

In the systems R_2O_3 - V_2O_5 - Ta_2O_5 , with R = a rare earth metal (Y, La, Ce, Pr, Nd, Sm, and Eu), new compounds of the type $RVTa_2O_9$ have been discovered and characterized by their X-ray powder diffraction diagrams, and in some cases, IR spectra were given [297-299].

2.12 Heteropolycompounds of Nb(V) and Ta(V)

The structures, properties and uses of metal-oxygen "heteropoly" clusters in aqueous solutions and in solids were reviewed [300]. A polyeder model illustrating the discussed kind of structures obtainable with niobium or tantalum is given in Figure 33. The $[Mn^{IV}O_6Nb_{12}O_{32}]^{12}$ - anion depicted can be thought of as containing two $[Nb_6O_{19}]^{8}$ - basic building units, known e.g. from the $[HNb_6O_{19}]^{7}$ - anion [300].

A new niobium-vanadium *iso*polycompound has been prepared and assigned the formula {Na₅H[NbV₅O₁₈]}₂ [301].

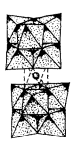


Figure 33. The *hetero*poly anion [Mn^{IV}O₆Nb₁₂O₃₂]¹²-with octahedrally coordinated Mn⁴⁺. Reproduced with permission from [300], H.-J. Lunk, S. Schönherr and V. I. Spitzin, *Z. f. Chem.* 27 (1987) 157.

2.13 Perchlorates, sulphates, phosphates and other oxy-complexes of Nb(V) and Ta(V)

Anhydrous Niobium(V) perchlorate, Nb(ClO₄)₅, and the perchlorato-niobates, Cs[Nb(ClO₄)₆] and Cs₂[Nb(ClO₄)₇], were prepared and characterized by chemical analysis, X-ray powder diagrams and IR-spectra [302]. Nb(ClO₄)₅ is a crystalline hygroscopic substance, which decomposes endothermally at about 70 °C, forming NbO(ClO₄)₃ and Cl₂O₇. Above 115 °C, NbO(ClO₄)₃ is transformed exothermally to NbO₂(ClO₄) with the liberation of Cl₂ and O₂. The reaction of Nb(ClO₄)₅ with CsClO₄ in HClO₄ gives the perchloratoniobates Cs[Nb(ClO₄)₆] and Cs₂[Nb(ClO₄)₇] [302]. The IR-spectrum of Nb(ClO₄)₅ contains sets of bands characteristic of unidentate and bidentate perchlorato-groups, whereas the complexes in Cs[Nb(ClO₄)₆] and Cs₂[Nb(ClO₄)₇] are unidentately bound [302].

Anhydrous Tantalum(V) *perchlorate*, Ta(ClO₄)5, and the *perchlorato*-tantalates, Cs[Ta(ClO₄)6] and Cs₂[Ta(ClO₄)7], have been prepared and characterized similarly [303]. Ta(ClO₄)5 is a colourless, crystalline, hygroscopic substance, which looses Cl₂O₇ at about 90 °C, to give TaO(ClO₄)3. Above 130-140 °C, TaO(ClO₄)3 is transformed exothermally to TaO₂(ClO₄) with the liberation of Cl₂ and O₂. The reaction of Ta(ClO₄)5 with CsClO₄ in HClO₄ gives the *perchloratotantalates* Cs[Ta(ClO₄)6] and Cs₂[Ta(ClO₄)7] [303]. Ta(ClO₄)5 gives sets of IR bands characteristic of *unidentate* and *bidentate perchlorato-groups*, whereas Cs[Ta(ClO₄)6] and Cs₂[Ta(ClO₄)7] are *unidentate* [303].

Electrode reaction equilibria of niobium complexes were studied by potentiometry and polarography in *per*chlorate and sulphate acidic solutions [304-305]. One- and two-electron reactions:

$$Nb(V) + e^{-} = Nb(IV)$$

$$Nb(V) + 2e^{-} = Nb(III)$$

occurred at the mercury electrode, Nb(V) probably being NbO(OH) $_2^+$, NbO(OH) $_2$ (SO₄) $_-$, NbOSO₄ $_+$ and NbO(SO₄) $_2^-$, and Nb(III) being Nb³⁺ and NbSO⁴⁺, respectively. Equilibria kinetics were studied [304-305].

The complexation reaction of Nb(V) with SO₄²- and HSO₄- was studied using UV spectroscopy; it was found that freshly prepared solutions are in general non-equilibrium ones [306].

The crystal structures were solved for K₇Nb(SO₄)₆ and K₇Ta(SO₄)₆, prepared by precipitation from solutions of Nb₂O₅ or Ta₂O₅ in molten K₂S₂O₇. The coordination polyhedron consisted of six *uni*dentate sulphato ligands [307], see Figure 34.

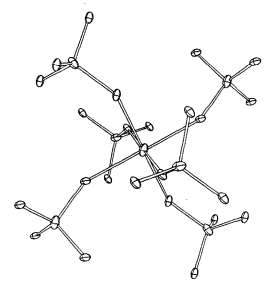


Figure 34. The structure of [Nb(SO₄)₆]⁷⁻ and [Ta(SO₄)₆]⁷⁻ ions with six *uni*dentate sulphato ligands [307-308]. Reproduced with permission from F. Borup and R. W. Berg.

The crystal structure of β -NbPO5 (or NbO(PO4)) contains PO4 tetrahedra and strands of two NbO6 octahedra, being connected in such a way that a ReO3-bronze type of structure with pentagonal tunnels is formed [309]. New allotropic forms of niobium and tantalum phosphates, NbPO5 and TaPO5 [310], and hydrates thereof NbOPO4·nH2O [311] were prepared and characterized by spectroscopy and by structural methods.

Preparation techniques for *ortho*tantalates $ATa(PO_4)_2$, with A = an alkali metal, have been developed [312]. Thermally induced solid state decomposition reactions, X-ray diffraction diagrams and IR-spectra were given [312]. In the crystal structure of RbTa(PO₄)₂, the tantalum atoms are in octahedral coordination with six PO₄ groups [313]. Infinite chains are formed by the TaO₆ and PO₄ groups. The results agree with earlier studies on HTa(PO₄)₂ and CsTa(PO₄)₂.

Sodium niobium phosphate Na₄Nb(PO₄)₃ exists in both crystalline and vitreous states [314]. Lattice data, EPR and optical absorption spectra were obtained. Vitrification seems to favour the formation of a niobyl group [314].

The crystal structure of niobyl tetraphosphate, $(NbO)_2P_4O_{13}$, has been solved, showing the Nb atom in octahedral coordination with five O atoms from different $P_4O_{13}^{6-}$ ions and one separate O atom [315].

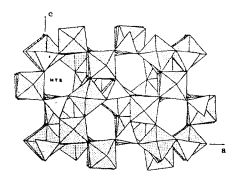
Niobium(V)-containing phosphates with the general formula $Cu_xNb_{1-x}Ti_{1+x}(PO_4)_3$ ($0 \le x \le 1$) were made by sintering TiP_2O_7 , Cu and oxides at high temperatures [316]. These compounds were recently shown to belong to the Nasicon type of structures, $NaA_2(PO_4)_3$ with A = Ti, Ge, Ce or analogous silicates. In such structures, three-dimensional lattice networks of cumulated octahedra and tetrahedra exist, forming large cavities, which are empty in $NbTi(PO_4)_3$ and in the new compounds [316]. Niobium is octahedrally coordinated to six oxygens.

The large family of niobium phosphate bronzes, $A_{4-x}Nb_6P_4O_{26}$ ($0 \le x \le 2$, A = K, Rb, Ba), has been studied, including $K_3Nb_6P_4O_{26}$ characterized by a mixed framework formed of corner-sharing NbO₆ octahedra and PO₄ tetrahedra and with K^+ located in between in intersecting tunnels [317], see Figure 35.

The crystal structure of K₅[Nb₆Zr₂P₅O₃₄] has been solved [318]. It contains NbO₆ and ZrO₆ octahedra sharing corners and PO₄ tetrahedra sharing corners with octahedra to form a three-dimensional network with potassium ions situated in interconnected channel-like cavities [318].

Also, the structure of newly prepared crystals of CaNb₂O(P₄O₁₃)(P₂O₇) has been resolved [319]. The main outstanding feature of this compound is the coexistence in the same lattice

of two kinds of phosphate anions with different degrees of condensation: a tetrameric one: $(P_4O_{13})^{6-}$ and a dimeric linear one: $(P_2O_7)^{4-}$. Niobium atoms are in two octahedral NbO6 coordination arrangements, one of which is nearly regular and another one less regular [319].



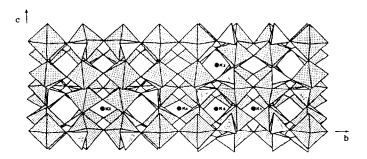


Figure 35. Projections onto (010) and (100) of the K₃Nb₆P₄O₂₆ framework structure [317]. Reproduced with permission from [317], M. M. Borel, A. Grandin, A. Benabbas, A. Leclaire and B. Raveau, *Mater. Res. Bull.* 24 (1989) 1485.

The 11-tungstoniobophosphoric acid, H₄(PNbW₁₁O₄₀), xH₂0, was prepared and its UV absorption and IR spectral proterties examined [320].

The role of Nb₂O₅ and Ta₂O₅ in glass-forming systems were discussed in some detail [321-323]. Raman spectra of some NbO₆ containing glasses and crystals of LiNbO₃ or MgNb₂O₆ were compared [321,323]. NbO₆ and TaO₆ octahedra are capable of forming continuous networks in Cs₂O-Nb₂O₅-Al₂O₃ and P₂O₅-BaO-Ta₂O₅-Al₂O₃ system glasses [322-323]. Other phosphates are mentioned in the section on oxides.

2.14 Intercalation of molecules into Nb(V) and Ta(V) oxides

Many niobates and tantalates are well-suited frameworks for intercalation: Intercalation of primary monoamines $NH_2C_nH_{2n+1}$ with n ranging from 1 to 9 [324] and of diamines $H_2N-(CH_2)_n-NH_2$, with n ranging from two to ten [325], can be done into the lamellar niobate $HNb_3O_8\cdot H_2O$ with interesting results. The intercalated products can be dehydrated or hydrated at room temperature. If hydrated, they can - easily and reversibly - be dehydrated to compounds such as $e.g.[H_3N-(CH_2)_n-NH_3]_{0.5}[Nb_3O_8]$. Crystallographic investigations have revealed that the diamines tend to orient themselves transverse to the Nb_3O_8 layers, forming, for the larger values of n, dense organic layers [325]. A typical structure is shown in Figure 36. Similar results were found for the monoamines [324].

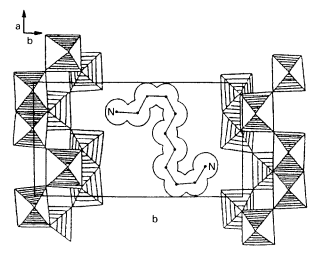


Figure 36. Structure of [H₃N-(CH₂)9-NH₃]_{0.5}[Nb₃O₈] (n = 9). The Nb₃O₈ layers and the organic layer are distinctly shown. Reproduced with permission from [325], R. Nedjar, M. M. Borel and B. Raveau, J. Solid State Chem. 71 (1987) 451.

The intercalation properties of HTiNbO₅ in the presence secondary and tertiary amines was also studied [326-327]. The TiNbO₅ layers, though different from the Nb₃O₈ layers of Figure 36, are able to form similar kinds of hydro-amine oxide layer compounds.

The layered solid acid $HCa_2Nb_3O_{10}$ can be intercalated with organic primary amines $C_nH_{2n+1}NH_2$, n = 1 to 16 [328].

2.15 Peroxy-niobates(V) and -tantalates(V)

The reactivity of the niobium *tetraper*oxo complex ion, $[Nb(O_2)_4]^{3-}$, was studied theoretically, using the multiple scattering X- α model as a basis for the calculation of molecular electrostatic potentials and electron deformation densities [329]. The electronic ground state configuration of $[Nb(O_2)_4]^{3-}$ is given as ${}^1A_1(2a_2)^2(2b_1)^2(8e)^4$ [330].

2.16 Nb(V) and Ta(V) sulphides and related S- and Se-containing compounds

Amorphous niobium sulphide, Nb₂S₅, was prepared by letting hexamethyldisilathiane (CH₃)₃Si-S-Si(CH₃)₃ react with NbCl₅ in acetonitrile at room temperature [331]. Acetonitrile was used because of high solubility of the intermediate NbSCl₃, and in this way a pure product could be obtained [331].

The structural chemistry of binary and ternary selenides of niobium and tantalum has been extensively studied. The four tetrachalcogenometalates K_3MX_4 (M=Nb, Ta, X=S, Se) have been obtained and characterized by e.g. single crystal X-ray diffraction methods [332]. They all contain discrete tetrahedral MX_4^{3-} ions. Single crystals of Tl_3MX_4 (M=Nb, Ta, X=S, Se) were synthesized and characterized by DTA, X-ray diffraction and infrared photographs [333]. Such crystals contain isolated, parallel-oriented MX_4 tetrahedra with metals M forming a body-centered cubic lattice. The ionicity of the chemical bonding in Tl_3MS_4 (M=Nb or Ta) was estimated to be intermediate between (I), a crystal consisting of ionicly bonded Tl^+ and MS_4^{3-} , and (I), an all covalent crystal. The bond ionicity increases from Nb to Ta [334-335]. Also, for I03 Cu3TaS4 and I13TaX4 (I13TaX4 (I

Discrete anions Ta₂S₁₁⁴⁻ and Nb₄Se₂₂⁶⁻ of the metals in oxidation state +5 have been discovered in the newly solved crystal structures of the salts K₄Ta₂S₁₁ and K₃Nb₂Se₁₁ [472].

Black platy hexagonal crystals of approximate composition Ba₂NbS₄(S₂)_{0.5} have been prepared from BaS, Nb and S at 1000 °C [339]. The solved X-ray crystal structure shows that Nb atoms occupy octahedral interstices in between layers of disordered S and S₂ ions [339]. Related compounds Nb₂PdSe₆, Nb₂Pd_{0.71}Se₅, Nb₃Pd_{0.72}Se₇, Ta₂NiS₅ and Ta₂NiSe₅ have been prepared and examined [16, 340-341].

In the Ba-Ta-S system, unstoichiometric compounds with tantalum vacancies exist. Hence, Ta can have a formal charge of +5, even when the formula looks otherwise. Thus, BaTaS3 is BaTa0.8S3. The same seems to be the case for BaTa2S5, Ba3Ta2S8, Ba9Ta4S20 and now Ba16.5Ta9S39, all of which have been characterized by means of their crystal structures [342].

Compounds Ta_2MSe_7 (M = Ni, Pt) have also been prepared and shown [343-344] to contain Ta(V)-centered bicapped trigonal prisms of Se^{2-} ions and Se-Se bonds analogous to $TaSe_3$. In addition to this, the structure also contains chains of Ta-centered octahedra and Pt- or Ni-centered square pyramids. The metallic versus non-metallic properties were examined by electronic structure calculations.

Following the earlier synthesis (and structure evaluation) of Nb₂Pd₃Se₈ [345], now also Ta₂Ni₃S₈ [346], Ta₂Pd₃Se₈ [347], Ta₂Pt₃Se₈ [346], Co₂Nb₄PdSe₁₂ [346], Co₂Ta₄PdSe₁₂ [347-348], [Co_{1.5}Pt_{0.5}]Ta₆PtSe₁₆ [349] and Nb₂Pd_{0.71}Se₅ [347] single crystal fibers or platelets have been prepared, by the method of long-time heating of the elements in closed quartz cells, placed in a temperature gradient and with bromine as transporting agent. The crystal structures have been solved and a review on these compounds and their structural chemistry has been published [16].

The basic Nb₂Pd₃Se₈ channel type structure, like that of Ta₂Pd₃Se₈, see Figure 37, contains niobium atoms in a trigonal prismatic environment of Se atoms and two types of Pd atoms square planar and square pyramidal - each coordinated by Se atoms [345]. The structural results are consistent with the simple valence description: Nb(V), Pd(II) and Se(-II). The other structures contain columns of Nb or Ta in somewhat similar octahedral and trigonal-prismatic sites, surrounded by shared sulphur or selenium atoms. Ta₂Ni₃S₈ can be characterized as a semiconductor [346], whereas Nb₂Pd_{0.71}Se₅ is a metallic conductor along the needle axis [347].

Ta₂P₂S₁₁ is a new phase of the Ta-P-S system [350]. Single crystal X-ray diffraction studies on Ta₂P₂S₁₁, obtained from elements at high temperature, yielded a structure based on bipolyhedral [Ta₂S₁₁] with hepta-coordinated tantalum(V) and tetrahedral [PS₄]. The [Ta₂S₁₁] units are formed of two distorted [TaS₇] groups sharing a triangular face made of mono- and disulphide anions (in some positions disordered). The [Ta₂S₁₁] units are shown in Figure 38.

The framework of the $[Ta_2S_{11}]$ units leaves three kinds of empty tunnels in the structure. The compound can be formulated as $Ta(V)_2P(V)_2S(-II)_9(S(-I)_2)$ and it is a *diamagnetic* semi-conductor [350].

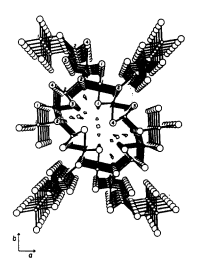


Figure 37. Perspective view along [001] of the structure of Ta₂Pd₃Se₈. Ta, Pd and Se atoms are small filled circles, small open circles, and large open circles, respectively. Reproduced with permission from [347], D. A. Keszler, J. A. Ibers, S. Maoyu and L. Jiaxi, *J. Solid State Chem.* 57 (1985) 68.

A new tunnel structure compound with inserted polymeric sulphur was found by X-ray diffraction study on single crystals of $Ta_4P_4S_29$, obtained from the elements by tempering under vacuum [351]. The structure is based on bi-prismatic, bicapped [Ta_2S_{12}] units including sulphur pairs, bonded to each other through [PS_4] tetrahedra sharing sulphurs with [Ta_2S_{12}]. The [Ta_2S_{12}] unit is shown in Figure 39. The framework leaves large tunnels, in which (S_{10}) sulphur-chains are found to be inserted. The compound can be formulated as $Ta(V)_4P(V)_4S(-II)_16(S(-I)_2)_4S(0)_5$ and it is accordingly a diamagnetic semiconductor [351].

2.17 Nb(V) and Ta(V) complexes with O-donor ligands

2.17.1 Nb(V) and Ta(V) alkoxides

The complex salt Li[Nb(OEt)6] was obtained as a result of crystallization from an ethanolic solution of lithium ethoxide and niobium ethoxide [352]. The crystal structure was recently determined [353]. A partial hydrolysis product, [Li2Nb2(OEt)10(OH)2], has a structure which was solved by single crystal X-ray diffraction analysis, see Figure 40. The structural unit is based on the *tetra*nuclear Li2Nb2 framework, similar to the [Ti(OMe)4]4 and [W(OEt)4]4

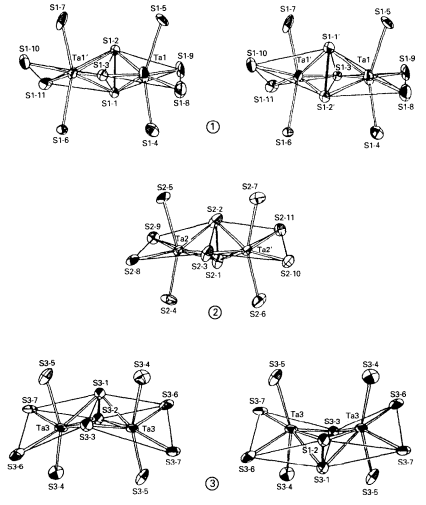
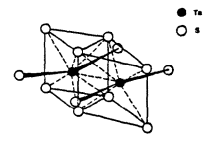


Figure 38. The three different [Ta₂S₁₁] bipolyhedral units encountered in Ta₂P₂S₁₁. (1) and (3) have two arrangements due to disorder in the shared sulphur triangle. Reproduced with permission from [350], M. Evain, S. Lee, M. Queignec and R. Brec, J. Solid State Chem. 71 (1987) 139.

tetramers; however, two of the four [MO₆] octahedrons are replaced by [LiO₄] tetrahedra. The interest in Li[Nb(OEt)₆] is large, partly because of its possible application as a precursor-material for the preparation of LiNbO₃ optical components [353].

Zn[Nb(OEt)6]2, was obtained from treating K[Nb(OEt)6] with ZnCl₂ [262]. K[Nb(OEt)6] was obtained from dissolving potassium into dry ethanol (EtOH) and letting it react with [Nb(OEt)5]. From Zn[Nb(OEt)6]2, a lead zink niobate, Pb(Zn_{1/3}Nb_{2/3})O₃, has been obtained by a sol-gel process [262].



Perspective drawing of the [Ta₂S₁₂] bicapped biprismatic unit of the basic tunnel structure of Ta₄P₄S₂₉. Reproduced with permission from [351], M. Evain, M. Queignec, R. Brec and J. Rouxel, J. Solid State Chem. 56 (1985) 148.

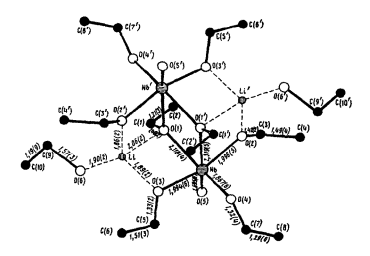


Figure 40. Structure of [Li₂Nb₂(OEt)₁₀(OH)₂]. Reproduced with permission from [352], A. I. Yanovskii, E. P. Turevskaya, N. Ya. Turova and Yu. T. Struchkov, Sov. J. Coord. Chem. 11 (1985) 63

The 3-methyl-1-pentene-3-oxy derivates, $M(OC(C_2H_5)(CH_3)CH:CH_2)_6$, with M = Nb or Ta, have been prepared [354]. They were characterized by elemental analysis, mol weight determinations, and IR and ¹H NMR spectral studies, and found to be *mono*meric colourless complexes, soluble in common organic solvents [354].

The penta-iso-propyloxide complex [Nb(OPr i)5] reacts with [La(OPr i)3] in alcoholic solution forming crystalline alkoxyniobates with formula [LaNb2(OPr i)13]. The structure of the complexes was discussed in terms of the so-called "coordination polymerism" concept [355].

Iso-propyloxide complexes $K[M(OPr^i)_6]$ (M = Nb, Ta) react with $CrCl_3 \cdot 3THF$ (THF = tetrahydrofuran) in benzene in molar ratios 3:1 yielding soluble complexes of the type $Cr[M(OPr^i)_6]_3$ [356]. On heating under vacuum the green complexes tend to disproportionate into $Cr(OPr^i)_3$ and $M(OPr^i)_5$. A number of bimetallic alkoxides ($Cr[M(OR)_6]_3$ with R = Me, Et, Pr^i , and/or Am^i in varying combinations, and $Cr[M(OPr^i)_4(acac)_2]_3$ with acac = acetylacetonate) have also been synthesized by alcoholysis of $Cr[M(OPr^i)_6]_3$ with methanol, ethanol and t-amyl alcohol. The compounds were characterized by chemical analysis, by infrared, visible and electron spin resonance spectroscopy, and by magnetic susceptibility measurements [356]. This and the replaceability of only twelve out of eighteen isopropoxy groups were taken as evidence for the following structure, 10:

$$\begin{array}{c|c}
CHMe_{2} \\
OR \\
OR \\
OR \\
OR \\
CHMe_{2}
\end{array}$$

In the acetylacetonate complexes, $Cr[M(OPr^i)_4(acac)_2]_3$, niobium or tantalum achieve a coordination number of eight (acac is bidentate) [356].

A new class of bimetallic isopropoxides of Ni(II) with the general formula Ni[$M(OPr^i)_6$]2, with M = Nb or Ta, have been synthesized and characterized by elemental analyses, molecular weight determinations, IR and VIS spectroscopy, in addition to magnetic susceptibility measurements and alcohol exchange reaction studies [357]. An octahedral geometry was assigned to Ni(II) primary bimetallic alkoxides (Ni[$M(OR)_6$]2, M = Nb or Ta, R = Me, Et, Pr^n or Bu^n), while in secondary derivatives, (e.g. Ni[$M(OR)_6$]2, M = Nb or Ta, and $R = Pr^i$), an equilibrium between octahedral and tetrahedral forms seems to exist. The tetrahedral and octahedral structures deduced were 11 and 12 [357].

Bimetallic alkoxides of copper(II) of the type $Cu[M(OPr^i)_6]_2$ (M = Nb or Ta) have been obtained, by letting $CuCl_2$ interact with $K[M(OPr^i)_6]$ in the molar ratio 1:2 in benzene solution [358]. These blue-green complexes undergo facile alcohol interchange with primary alcohols such as methanol, n-propanol and n-butanol, forming complexes of the type $Cu[M(OR)_6]_2$, R = Me, Et, Pr^n and Bu^n . With t-butanol, only $Cu[Ta(OPr^i)_2(OBu^i)_4]_2$ was obtained. Infrared, electronic and electron spin resonance spectral and magnetic susceptibility measurements indicated a distorted octahedral D_{4h} geometry, 13, for Cu(II) [358].

Novel *ter*metallic *iso*propoxides $[(Pr^iO)_4M(\mu\text{-}OPr^i)_2 \text{ Be}(\mu\text{-}OPr^i)_2\text{Al}(OPr^i)_2]$, M = Nb(V) and Ta(V) have been synthesized from $Pr^iOBe(\mu\text{-}OPr^i)_2\text{Al}(OPr^i)_2$ and $M(OPr^i)_5$ and characterized by IR, NMR and mass spectroscopy [359]. The complexes have the structure shown in Figure 41, and they are soluble in anhydrous common organic solvents and can be volatilized without noticeable disproportionation.

A mixed alkoxide-aryloxide of tantalum, $Ta_2(OC_6H_3Pr^i_2)_4(OCH_3)_6$ has been isolated as white crystals, whose structure has been solved [360]. The structure, 14, consists of an octahedral arrangement of oxygen donor atoms around two tantalum metal centers in an edge shared bi-octahedron with two methoxide ligands (R = the 2, 6-di-isopropylphenoxy group) [360].

Some ethoxy-complexes are mentioned in the section on carboxylates.

Figure 41. Structure of *ter*metallic *iso*propoxides $[(Pr^iO)_4M(\mu\text{-}OPr^i)_2Be(\mu\text{-}OPr^i)_2Al(OPr^i)_2]$, M = Nb(V) and Ta(V). Reproduced with permission from [359], M. Aggrawal and R.C. Mehrotra, *Polyhedron* 4 (1985) 845.

2.17.2 Nb(V) and Ta(V) alkenoxides

4-pentene-1-oxy derivatives of niobium and tantalum, $M(OCH_2CH_2CH_2CH=CH_2)_5$, have been prepared in dry benzene and characterized on the basis of elemental analyses, molecular weight determinations, IR and PMR spectral studies [361]. The presence of double bonds does not seem to have any marked effect on the properties of these compounds when compared with the corresponding alkoxides.

An X-ray study of the alkylidene complex Ta(CHCMe₃)(S-2,4,6-C₆H₂-Prⁱ₃)₃(SEt₂) have suggested why the complex is inactive for metathesis of ordinary olefins [362].

2.17.3 Nb(V) and Ta(V) carboxylates

White crystalline compounds of the general formula $NbO_2(OOCR)$ and $TaO_2(OOCR)$ can be obtained from the *penta*chlorides and an excess of the carboxylic acids, RCOOH with R = H, CH_3 , C_2H_5 , $n-C_3H_7$, CH_2Cl , $CHCl_2$ and CCl_3 . The compounds were characterized by chemical analysis and IR spectroscopy [129].

A tris-complex of niobium(V), NbOL₃, with L = the pyridine-2-caboxylate anion, 15, as a bidentate ligand, was prepared by controlled oxidation of the tetrakis-niobium(IV) complex Nb(L)₄. The compound was characterized by electronic and IR spectra [363].

The preparation of *hetero*cyclic carboxylates of niobium(V) and tantalum(V) in dry benzene has been considered for the acids HL = HTCA = 2-thiophenecarboxylic acid and HL = HTAA = 2-thiopheneacetic acid.

The ethoxy complexes of the type $M(OEt)_{5-n}L_n$ were obtained, with M = Nb, Ta, n = 1 to 3 and L = TCA or TAA [128]. The complexes were characterized on basis of chemical analysis as well as IR and ¹H NMR data. A tentative seven-coordinated structure, 16, was assigned to the $M(OEt)_3L_2$ complexes:

$$\begin{bmatrix} R & O \\ O \\ O \end{bmatrix}_{2} & OR & 16 \end{bmatrix}$$

 $M(OEt)_3L_2$, with M = Nb or Ta and L = TCA or TAA [128].

A study of the mechanism of formation of mixed complexes of niobium(V) with 4-(2-pyridylazo)-resorcinol (PAR or H_2R) and hydroxy-acid ions (A^{2-} = oxalate and tartrate) was done [364]. The reaction was represented by a ligand-ligand replacement reaction and a ligand addition reaction:

$$NbOA_{i}^{(3-2i)+} + H_{2}R --> [NbOA(HR)_{2}]^{-} + (i-1)A^{2-} + 2H^{+}$$

 $NbO^{3+} + A^{2-} + H_{2}R --> [NbOA(HR)_{2}]^{-} + 2H^{+}$

When the hydroxyacid residue A^{2-} is citrate, *tri*hydroxy-glutarate or mucate, the replacement of the residues by PAR anions were not possible due to steric hindrance [364]. In this case the mixed complexes can be made by the addition reaction [364].

The niobium(V)-4-(2-pyridylazo)-resorcinol (PAR) - tartaric acid - sodium chlorate complex system exhibits a catalytic polarographic wave which can be applied to sensitive determination of minute amounts of niobium in geological samples [365]. Further properties of 4-(2-pyridylazo)-resorcinol (PAR) complexes are described in the section on extraction.

A new seven coordinate *mono*meric complex, the oxo-tris-(tropolonato)niobium(V) monohydrate, NbO T_3 ·H₂O ($T = O_2H_5C_7$, the tropolonato seven-sided-ring group) was synthesized and its X-ray crystal structure determined [366]. The distorted pentagonal-bipy-ramid configuration of oxygen atoms around niobium is shown in Figure 42. The terminal O atom occupies an axial position with a short Nb-O bond (1.71 Å). The analogous compounds NbS T_3 and NbSe T_3 have been obtained by reaction of NbO T_3 with [(Me₃Si)₂Y] (Y = S or Se).

The complexation reaction of 5,5'-methylene-bis-salicylaldehyde (MBS) with TaCl₅ was studied under various conditions (in DMF solution in dry nitrogen or ambient air or in oxalate solution) [127]. The 1:3 compound obtained was diamagnetic and insoluble in common organic solvents. A probable structure (see Figure 43) was proposed on the basis of elemental analysis, magnetic measurements and infrared spectra [127].

2.18 Nb(V) and Ta(V) complexes with S- and Se-donor ligands

By mixing NbCl₅, LiSCH₂CH₂SLi (lithium 1,2-ethane *di*thiolate) and (Et)₄NCl (tetraethyl ammonium chloride) in molar ratios 1:3:1in acetonitrile it was possible [367] to prepare crystals of the composition [(Et)₄N][Nb(SCH₂CH₂S)₃]. TaCl₅ formed a similar compound. The anion

Figure 42. X-ray crystal structure of oxo-tris-(tropolonato) niobium(V) monohydrate, [NbO(O₂H₅C₇)₃]·H₂O. The distorted pentagonal-bipyramid configuration is easily seen. Reproduced with permission from [366], M. G. B. Drew, D. A. Rice and D. M. Williams, Inorg. Chim. Acta 118 (1986) 165.

Figure 43. Probable structure of tantalum MBS complex [TaC45H33O13] with seven-coordinated Ta. Reproduced with permission from [127], S. Chomal, A. S. Aazmi and G. C. Shivahare, *Acta Chim. Hung.* 122 (1986) 127.

of the niobium complex was studied by X-ray crystal structure determination [367]. It has three chelate rings arranged like a propeller (C_3 -symmetry, absolute configuration \land , see Figure 44).

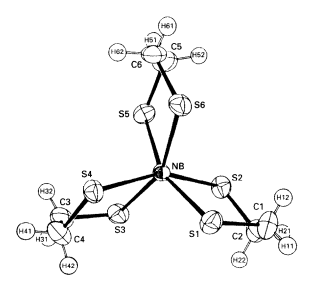


Figure 44. Structure of the anion △-[Nb(SCH₂CH₂S)₃]⁻. Reproduced with permission from [367], K. Tatsumi, Y. Sekiguchi, A. Nakamura, R. E. Cramer and J. J. Rupp, *Angew. Chem.* 98 (1986) 95.

Also, the synthesis and IR and Raman spectroscopic studies of $[A][M(SCH_2CH_2CH_2S)_3]$ and $[A][M(SCH=CHS)_3]$ with $A = Ph_4P$, Et₄N and M = Nb, Ta and the crystal structure of $[Ph_4P][Nb(SCH_2CH_2CH_2S)_3]$ have been presented [473]. Electronic spectra and structure/bonding properties were discussed in relation to extended Hückel calculations [473].

High-coordination-number diethyl-dithiocarbamate compounds have been obtained by letting the pentahalides react with Na(S₂CNEt₂) [368]. The following compounds were identified by means of their crystal structures: [Nb(S₂CNEt₂)₄]Br, [Nb(S₂CNEt₂)₃S] and [Ta(S₂CNEt₂)₃(S₂)]. The (S₂CNEt₂) ligands are all chelating the metals. The first of these compounds is eight-coordinate with mmmm dodecahedral symmetry, the next one is seven-coordinate with an axial multiple Nb=S bond and a pentagonal bipyramidal molecular geometry and the third one is eight-coordinate with the metal in a distorted dodecahedral environment [368].

The coordination chemistry of *polynuclear* transition metal complexes with sulphur ligands has been reviewed recently [368A].

2.19 Nb(V) and Ta(V) complexes with O- and S-donor ligands

Polysulphide anions S_X^{2-} , with x=2 or more, have been incorporated into a red niobium(V) dinuclear complex salt, $(PPh_4)(NH_4)[Nb_2(OMe)_2(S_2)_3(S_5)O]$, see Figure 45. The ligands around Nb form a pentagonal bipyramid, in which the bridging oxygroup and the oxygen atoms

of the methoxy groups are on axial positions and the S atoms of the S_5^{2-} and non-equivalent S_2^{2-} ligands are equatorially positioned. Preparative details, IR and Raman data are given [369].

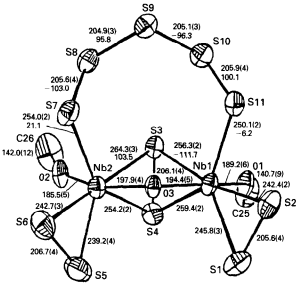


Figure 45. Structure of the anion [Nb₂(OMe)₂(S₂)₃(S₅)O]²⁻. Reproduced with permission from [369], A. Müller, J. Schimanski, M. Römer, H. Bögge, F.-W. Baumann, W. Eltzner, E. Krickemeyer and U. Billerbeck, *Chimia* 39 (1985) 25.

The new seven coordinate *monomeric* complexes NbS T_3 and NbSe T_3 have been obtained by reaction of NbO T_3 with [(Me₃Si)₂Y] ($T = O_2H_5C_7$, the tropolonato seven-sided-ring group and Y = S or Se). X-ray crystal structures were determined [366], see the analogous oxide shown in Figure 42.

The Nb=S group has been examined in the *tris*-N,N-diethyl dithiocarbamato compound NbS(S₂CNEt₂)₃ [370-371]. The X-ray structure solution of the yellow crystals showed two independent molecules with pentagonal bipyramidal coordination, in which the terminal sulphide atom occupies an axial position and the other six positions are taken by sulphur of dithiocarbamate [370]. The results compare well with previously solved structures of NbO(S₂CNEt₂)₃ [372] and TaS(S₂CNEt₂)₃ [373].

Dithiolato dinitrogen (or hydrazido (4-)) tantalum(V) complexes, $[Ta(SAr)_3(THF)]_2(\mu-N_2)$ with $Ar = 2,6-C_6H_3(Pr^i)_2$, $2,4,6-C_6H_2(Pr^i)_3$ and phenoxide analogues (S replaced by O) have been made $(pr^i = isopropyl)$ [374]. Also, crystal structural comparisons have been made of

[Ta(S-2,6-C₆H₃(Prⁱ)₂)₃(THF)]₂(μ -N₂) and [Ta(O-2,6-C₆H₃(Prⁱ)₂)₃(THF)]₂(μ -N₂). The compounds contain nearly linear Ta=N-N=Ta cores and the Ta coordination is five-coordinate [374].

The preparation of $(PPh_4)[NbS(SPh)_4]$ was described, and the crystal structure of the compound was solved, giving a square-pyramidal geometry at the metal with a multiply-bonded sulphur at the apex (Nb-S = 2.171(2) Å) [112].

2.20 Nb(V) and Ta(V) oxynitrides and complexes with O- and N-donors

New oxynitrides $ANbO_2N$ and $ATaO_2N$ with A = Ba, Sr, Ca were prepared and shown to have a ABO₃ perovskite type structure [375].

Ta(OPr i)5- $_x$ (SB) $_x$ coordination compounds of tantalum(V) have been synthesized in boiling anhydrous benzene solution, OPr i being the isopropoxide radical, x = 0, 1, 2 or 3 and SBH is aldimines of the type o-HO-C6H4-CH:N-C6H4-CH3-o, o-HO-C6H4-CH:N-C4H9-n, C_6 H5-CH:N-NH-CONH2 or C_6 H5-CH:N-NH-CSNH2 [376-377]. The aldimines were obtained by condensation of salicylaldehyde with o-methylaniline or n-butylamine and of benzaldehyde with semicarbacide hydrochloride or thiosemicarbacide. The complexes were fully characterized by their analytical data, conductances, magnetic data, IR, VIS and PMR spectra and molecular weights determined by the cryoscopic method in benzene. The complexes are all monomeric, with coordination numbers six, seven and eight for the x = 1-, 2- and 3-derivatives, respectively, with the aldimines acting as bidentate ligands, using O, N or S atoms for chelating to tantalum [377].

Complexation of niobium(V) and tantalum(V) with 4-oximino-3-methyl-2-pyrazolin-5-one takes place via oximino N and pyrazolone carbonyl O, resulting in the *bi*nuclear compounds $LM(O)(\mu-O)_2M(O)L$, (M = Nb or Ta, HL = 4-oximino-3-methyl-2-pyrazolin-5-one), according to elemental analysis, magnetic and IR spectral methods [378].

The complexation reaction of 5,5'-methylene-bis-salicylaldoxime (MBSO) with TaCl₅ was studied under various conditions (in DMF solution in dry nitrogen or ambient air or in oxalate solution) [127]. The 1:3 compound obtained was diamagnetic and insoluble in common organic solvents. A probable structure (see Figure 46) was proposed on the basis of elemental analysis, magnetic measurements and infrared spectra [127].

Niobium and tantalum *penta*ethoxides react with *mono*functional benzoyl hydrazones, BHyH [379], or *bi*functional benzoyl hydrazones, BH₂ [380], in refluxing benzene to give *mono*meric red or brown products of the type $M(OEt)_{5-X}(BHy)_X$, x = 1, 2 or 3 [379], or yellow to orange $M(OEt)_{5-2X}(B)_X$, x = 1 or 2 and x = Nb, Ta, respectively [380]. The *bi*functional benzoyl

Figure 46. Probable structure of tantalum MBSO complex, [TaC45H39O13N6], with seven-coordinated Ta. Reproduced with permission from [127], S. Chomal, A. S. Aazmi and G. C. Shivahare, *Acta Chim. Hung.* 122 (1986) 127.

hydrazones are *tri*dentate [380]. The benzoyl hydrazones employed were prepared by condensing an amine with an aldehyde or ketone, *e.g.* benzaldehyde, acetophenone and furfural. On the basis of elemental analysis, IR and NMR spectra and molecular weight determinations, it was concluded that the *mono* functional hydrazones behave like *bi* dentate ligands coordinating to the metal through deprotonated enolic and azamethine groups [379]. The following 6, 7 and 8 coordinate structures, 17,18 and 19, were tentatively assigned to the complexes.

Niobium(V) Schiff-base complexes of the type Nb(OEt) $_{5-x}(L)_x$ where x = 1 or 2 and LH is the Schiff-base salicylidene-2-aminpyridine, 20, have been isolated as liquids from niobium pentaethoxide /Schiff-base mixtures in dry benzene [381].

The Schiff-base salicylidene-2-aminopyridine.

The compounds were *mono*meric in boiling benzene. IR spectra indicated coordination through azomethine nitrogen and phenolic oxygen. Also tantalum alkoxides are known to react with certain Schiff-bases [382].

Niobium(V) forms a ternary complex [Nb(BPHA)₂(DCTA)] (BPHA = N-benzoyl-N-phenylhydroxylamine and DCTA = 1,2-diamino-cyclohexane-tetraacetic acid) which can be used for analytical purposes (polarographic determination of traces of Nb) [383].

A comparative characterization was given of glycinato-niobates and glycinato-tantalates, synthesized in basic media [384]. Both the carboxylate and the amino groups of the glycinic acid were coordinated to the metal, according to IR-spectral evidence [384].

The synthesis has been performed and the crystal structure solved of Nb(OAr-2,6-Ph₂)₃(NMe)(HNMe₂) and the corresponding Ta methylimide complex (OAr-2,6-Ph₂ is 2,6-diphenylphenoxide) [385].

2.21 Nb(V) and Ta(V) complexes with As-ligands

A new kind of complex anion, [NbAsg]³-, see Figure 47, was discovered in crystals containing also rubidium and a rubidium cryptate. The new ion of symmetry $\overline{8}2m - D_{4d}$ participates in the formation of one-dimensional chains with stacked crown-forming (As⁻)g polyanions containing Nb⁵+ in their centers separated by Rb⁺ cations. Some data on the crystal structure and the molecular orbital diagram are reported [386].

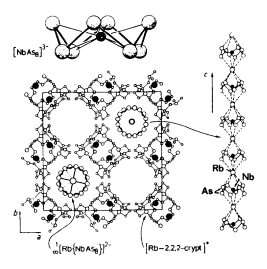


Figure 47. Structure of the new complex anion, [NbAs8]³⁻, and the crystal in which it was found. Reproduced with permission from [386], H.-G. von Schnering, J. Wolf, D. Weber, R. Ramirez and T. Meyer, *Angew. Chem.* 98 (1986) 372.

2.22 Nb(V) and Ta(V) carbides and complexes with C-donor ligands

Niobium carbide NbC has unusual properties such as being hard and brittle. Its electronic structure has been studied by high-energy electron energy-loss spectroscopy [387].

The reaction between the Nb(V) cyclopentadienyl derivate $[(\eta^5-t\text{-BuC}_5H_4)_2\text{Nb}]BH_4$ and sulphur in toluene solution has been studied [388]. In this way, a new Nb(V) compound, $\{(Cp')_2\text{Nb}(\eta^2-S_2)\}_2(\mu-S_5)(Cp'=\eta^5-t\text{-BuC}_5H_4=tert\text{-butyl-substitued cyclopentadienyl})$, has been synthesized and its crystal structure determined, see Figure 48.

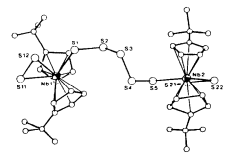


Figure 48. Structure of (t-BuC₅H₄)₄Nb₂S₉. Reproduced with permission from [388], H. Brunner, U. Klement, J. Wachter, M. Tsunoda, J.-C. Leblanc and C. Moise, *Inorg. Chem.* **29** (1990) 584.

2.23 Biological complexes of Nb(V) and Ta(V)

Chelate macrocyclic complexes of the extended π -bonding type have been little studied in the period. Acid-base properties of the *meso*-nitrogen atoms of a substituted tetrazaporphyrin metallate(V) ring system (*tetra*-4-*tert*-butylphtalocyanine, OTaPc') were studied in anhydrous acetic and sulphuric acids, for the case of niobium [389] and for tantalum [390]. The kinetic stabilities (stepwise protonation constants [389]) were determined by visible absorption spectroscopy. Niobium and tantalum phtalocyanine complexes belong to the group of stable phtalocyanines in proton-donor media [390].

The syntheseis, properties and crystal structures of some phtalocyaninato complexes have recently been reported: The *dinuclear* Br3⁻ salt μ-bromo-μ-*di*oxo-*bis*(phtalocyaninato-niobium(V))*tri*bromide [390A], and the *mono*nuclear *tri*chloro(phtalocyaninato)Tantal(V) [390B], both having coordination numbers of seven.

Complexes [Nb(por)(O)(OAc)] and [Nb(por)(X)3], (in which por is a porphyrinate²- anion: oep = octaethylporphyrinate, tpp = tetraphenylporphyrinate or tptp = tetraparatolylporphyrinate and X = Cl or Br), have been prepared and characterized on the basis of IR, UV-VIS, electrochemical and EPR data [391].

Ascidians (sea squirts) are able to extract niobium from sea-water containing 0.1 ppb of Nb, but the complexes responsible for the binding of niobium are not known. Recently, it was shown [392] in vitro that amino-acids like glycine, alanine, serine, tyrosine, cysteine cannot be responsible, but certain α -hydroxy carboxylic acids (lactic, malic, glycollic, and tartaric acids) are able to candidate, and suggestions are made for the mechanism [392].

2.24 Solvent extraction of Nb(V) and Ta(V)

The extraction of niobium from ores and the refinements needed to obtain high-purity niobium complexes and niobium metal have been reviewed [13-14]. The distribution coefficients for extraction of various elements into the commonly used solvent methyl-isobutyl-ketone (MIBK) are given, according to reactions similar to e.g.

4 HF + NbF₇²⁻ + 3 MIBK
$$\longrightarrow$$
 H₂NbF₇·3 (MIBK) + 2 HF₂⁻

Separation coefficients of niobium and tantalum in various ketones and other details are also given [14-15]. In the extraction/separation process, MIBK can be replaced by *tri*-butyl phosphate (TBP) forming a soluble complex H₂NbF₇·3(TBP) [15].

The extraction of niobium(V) and tantalum(V) from >4 molar sulphuric acid solutions by di-2-ethylhexylhydrogen phosphate (D₂EHHP) has been studied by chemical analysis, infrared spectroscopy and other methods [393]. The extracted compounds did not contain any sulphate group, and corresponded to the formulae NbO(D₂EHHP)₃ and TaO(D₂EHHP)₃, tending to polymerize in the organic phase, perhaps forming $MO(D_2EHHP)_3(H-D_2EHHP)_3$, M = Nb or Ta [393].

Niobium(V) in hydrochloric and sulphuric acid media, in the presence of an excess of chloride or thiocyanate ions, reacts with 3-hydroxy-2-methyl-1-phenyl-4-pyridone (HR) to give complexes, Nb(OH)₃ClR or Nb(OH)₃(NCS)R, which are extractable into chloroform [146]. The experimental conditions for quantitative extraction of niobium(V) into the organic phase are given, permitting a separation from zirconium(IV) and hafnium(IV) [394]. The identity of the complexes was determined spectrophotometrically [146].

A spectrophotometric study was made on the extraction of mixed complexes of niobium(V) and tantalum(V) with 4-(2-pyridylazo)-resorcinol (PAR or H₂R) and hydroxy-acids (oxalate, tartrate, citrate, trihydroxy-glutarate and mucate) by a chloroform solutions of tri-n-octylamine

(TOA) [395]. The Nb(V)-PAR-oxalate, Nb(V)-PAR-tartrate and Ta(V)-PAR-oxalate compounds go into the basic organic phase, probably forming the singly charged anions [NbO(C_2O_4)(HR)₂]⁻, [NbO($C_4H_4O_6$)(HR)₂]⁻ and [Ta(OH)₂(C_2O_4)(HR)₂]⁻, respectively [364, 395].

Re-extraction of NbOF $_5^{2-}$ and TaF $_7^{2-}$ from solvents with C=O, S=O and P=O functional groups has been studied by means of chemical analysis and Raman spectroscopy [46].

3. Niobium(V) and (IV) mixed complexes (d^0 and d^1)

The knowledge of the complicated phase relationships within the system NbO₂-Nb₂O₅, some of which have already been mentioned, has been extended through the preparation of 1-2 mm large crystals of Nb₂O₅₁, Nb₄7O₁₁₆, Nb₂5O₆₂ and Nb₅3O₁₃₂, all belonging to the NbO_{χ} type with 2.4 < χ < 2.5 [396]. Many systems with NbO₂-Nb₂O₅ and a third metal oxide have been studied. Nb₁₂O₂₉ has been modified to form V₃Nb₉O₂₉, the block-structure of which was studied by electron microscopy [397]. A new kind of block structure was found in the crystal structure of V₈Nb₅O₂₉ [398]. Lithium metal has been inserted into V₃Nb₉O₂₉ forming e.g.Li_{13.6}V₃Nb₉O₂₉ [399].

Non-stoichiometric phases $Hf_2Nb_20O_{51}$, $Zr_3Nb_44O_{116}$, $Hf_3Nb_44O_{116}$, β - $ZrNb_24O_{62}$ and β - $HfNb_24O_{62}$ have block structures, according to high resolution transmission electron microscopy [400].

A new mixed-valence oxoniobate compound, Sr₅Nb⁴+₃Nb⁵+₂O₁₆, was synthesized from Sr and Nb oxides in a hydrogen plasma at about 2000 °C. The complicated crystal structure was solved and found [401] to contain four different kinds of NbO₆ octahedra (two Nb(IV) and two Nb(V)) and one relatively strongly distorted NbO₄ tetrahedron, see Figure 49. Hence, niobium(IV) then can be octahedral as well as tetrahedral coordinated within the same framework.

Another new mixed-valence oxy-niobate compound, $Sr_7Nb_6O_{21}$, has been prepared [402]. The method involved a mixture of 4 Sr_2O_3 , 2 Nb and 1 Nb_2O_5 which was pressed and heated to about 2500 $^{\rm O}$ C by means of the thermal power of a CO_2 laser. Single crystals produced were investigated by X-ray diffraction techniques and proved to have a structure consisting of layers of perovskite-type, with a width of about 13 Å, and containing 6 layers of corner-linked NbO₆ octahedra and Sr_2^{2+} ions in between, see Figure 50. The three different kinds of niobium atoms all have trigonal symmetry with three long Nb - O bonds (2.09 - 2.18 Å) and three short ones (1.85 - 1.96 Å). The formula $Sr_7Nb_6O_{21}$ can be given as $Sr_{1.167}Nb_2^{+4}O_{0.333}Nb_2^{+5}O_{0.667}O_{3.5}$, showing the close relationship of the structure to the already known perovskites $CaNbO_{3.5}$ and $Ba_{1.25}TaO_{3.75}$ [402].

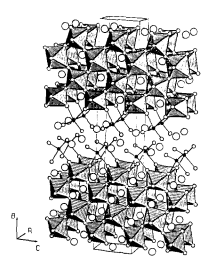


Figure 49. Structure of the mixed-valence oxy-niobate compound, Sr₅Nb⁴⁺₃Nb⁵⁺₂O₁₆, showing four kinds of NbO₆ octahedra (hatched) and one kind of deformed NbO₄ tetrahedra (small filled circles). Reproduced with permission from [401], K. Schückel and H. Müller-Buschbaum, Z. Anorg. Allgem. Chem. 528 (1985) 91.

4. Nb(IV) and Ta(IV) compounds (d1)

Low valent complexes of niobium and tantalum were described in a 275 pages long thesis [403].

4.1 Nb(IV) and Ta(IV) halide complexes

Thermodynamic data for gaseous tantalum halides, TaF4, TaCl4, TaBr4, and TaI4 were reviewed, and data given also for a large number of liquid and solid tantalum halide and oxyhalide compounds [20]. Crystals of NbCl4 have been obtained by chemical vapour transportation [404].

The vapour pressures of solid NbCl₄ and NbBr₄ have been determined by a spectrophotometric method, as function of temperature near 600 K [405]. NbCl₄ is *mono*meric in the gaseous state. Gas phase visible/near-IR spectra of NbCl₄ and NbBr₄ are given [405]. The stability of gaseous NbCl₄ was studied by tensimetry, DTA and TG methods [25, 406], and the enthalpies of formation of solid NbCl₄, NbBr₄ and NbI₄ were given [407-408]. A possible



Figure 50. Unit cell of the mixed-valence oxyniobate compound, Sr₇Nb⁴⁺₂Nb⁵⁺₄O₂₁, showing a perovskite layer of corner-linked NbO₆ octahedra (hatched) and the beginning of the next layers. Strontium ions (in the cavities between octahedra) are not shown. Reproduced with permission from [402], K. Schückel and H. Müller-Buschbaum, Z. Anorg. Allgem. Chem. 523 (1985) 69.

disproportionation of NbCl₄ into NbCl₃ and NbCl₅ occurred at ca. 320 °C [406]. The photoelectron spectrum (AlK α , HeI) of NbI₄ (or rather Nb₂I₈) was studied in an attempt to find the relation between the energy of the niobium core levels and the oxidation number [42].

The magnetic properties of crystals of Li₂NbF₆ and Na₂NbF₆ were determined from 4.2 to 293 K [409]. The crystals exhibit *para*magnetic behaviour as a result of the presence of Nb(IV) cations at the centers of independent octahedra, which are distorted from the ideal octahedral O_h -symmetry. The standard enthalpy of formation was determined for the Rb₂TaBr₆ crystal and for the TaBr₆²⁻ ion in the gas state [410]. The ferri-magnetic behaviour of trigonal MnNbF₆ was studied in order to obtain information on the Mn²⁺-F⁻-Nb⁴⁺ super-exchange interaction in this structure of corner-sharing NbF₆ and MnF₆ octahedra [411].

Niobium and tantalum in K_2NbF_7 and K_2TaF_7 will be partly reduced to oxidation state IV after being bombarded in a 4 keV Ar^+ ion beam [412].

A ternary chloride of tin(II) and tantalum(IV), Sn[Ta₂Cl₉], has been prepared as a green diamagnetic solid [413]. The X-ray crystal structure of this compound has several surprising features: discrete face-sharing bioctahedral [Ta₂Cl₉] units that show a chiral distortion from D_{3h} symmetry, see Figure 51.

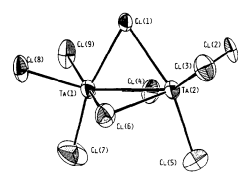


Figure 51. Structure of the [Ta₂Cl₉] unit as found in the Sn²⁺ salt. Reproduced with permission from [413], F. A. Cotton, E. Babaian-Kibala, L. R. Falvello and M. Shang, *Inorg. Chem.* **29** (1990) 2591.

4.2 Nb(IV) and Ta(IV) oxyhalide complexes

Thermodynamic data for gaseous tantalum oxyhalides, TaOF₂, TaOCl₂, TaOBr₂, and TaOI₂ were reviewed and data given also for a large number of liquid and solid tantalum halide and oxyhalide compounds [20]. Solid NbOF₂ has been synthesized and found to crystallize in space group *Pm3m* with one formula in the cell. Nb is on an octahedral site. Visible and UV spectra and magnetic behaviour between 77 and 300 K are reported [414].

NbOCl₂ and TaOCl₂ powders have also been examined by X-ray diffraction; the two compounds were found to be *iso*-structural and indexed powder diagrams were given, as well as structural and IR spectral data for the grey-green niobium compound [32]. NbOBr₂, TaOBr₂ and TaOI₂ crystals have further been prepared and examined by X-ray diffraction [34].

4.3 Nb(IV) and Ta(IV) halide complexes with O-donors

Dinuclear niobium(IV) complexes, Nb₂Cl₄(OMe)₄(MeOH)₂·2MeOH and Nb₂Cl₄(O-Me)₄(CH₃CN)₂, have been prepared and studied by X-ray crystallography and proton NMR

spectroscopy [415]. The complexes have an edge-sharing bioctahedral geometry and possess a single Nb-Nb bond and two bridging methoxy groups, see Figure 52. Axial and equatorial terminal ligands seem to interchange easily [415].

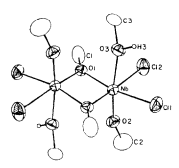


Figure 52. Structure of the centrosymmetric *di*nuclear niobium(IV) molecule, Nb₂Cl₄(OMe)₄(MeOH)₂. Reproduced with permission from [415], F. A. Cotton, M. P. Diebold and W. J. Roth, *Inorg. Chem.* **26** (1987) 3319.

The compound trans-NbCl₂(Bu^tC(O)CHC(O)Bu^t)₂ or, in short notation, trans-NbCl₂(dpm)₂ with Hdpm = dipivaloylmethane = 2,2,6,6-tetramethylheptane-3,5-dione, has been prepared in two polymorphs [416]. The X-ray crystal structures have been solved, and they show the same kind of molecule, see Figure 53. The UV-visible spectrum for this octahedral or rather D_2h -symmetry niobium⁴⁺ chromophore (the d^1 electron system) shows a doublet band at 588.4 and 626.7 nm. High formal oxidation states of the metal in many cases lead to charge transfer transitions obscuring the d-d transitions. By combining VIS and EPR spectral measurements with a molecular orbital calculation (the Fenske-Hall procedure), a reasonable and consistent picture of the electronic structure of the molecule was obtained [416]. The doublet in the spectrum was assigned to the d-d transitions yz --> xz and $yz --> x^2 - y^2$ (at ca. 16000 and 17000 cm⁻¹). The EPR spectrum with the expected 10-line signal, with < g> = 1.930 and with $< A_{iso}> = 149$ G also were explainable.

The new *bi*nuclear Nb(IV) compound, Nb₂Cl₂(OEt)(O₂CMe)₅, has been shown to have a crystal structure in which two (η^2 -CH₃COO)₂Nb groups are joined by μ -OC₂H₅, two μ -Cl atoms and μ 2- η^2 -CH₃CO₂. In this complex the Nb-Nb distance indicates the presence of a single bond [117].

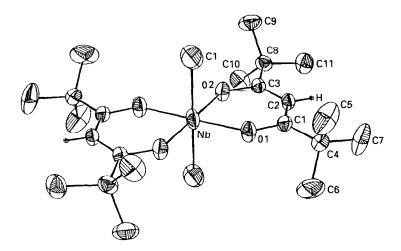


Figure 53. Structure of *trans*-NbCl₂(dpm)₂ with Hdpm = dipivaloylmethane in its *monoc*linic phase. Reproduced with permission from [416], F. A. Cotton, M. P. Diebold and W. J. Roth, *Polyhedron* 4 (1985) 1485.

4.4 Other Nb(IV) and Ta(IV) halide complexes

Solid NbS₂Cl₂ was obtained by letting Nb react with S₂Cl₂ [114]. It contains Nb₂(S₂)₂⁴⁺ groups linked with one another by Cl⁻ ions in a layered lattice [417] and is unstable when heated [418]. Its electronic, vibrational, resonance Raman spectra and the crystal valence force field were reported [419]. The compound underwent oxidation forming NbS₂Cl at 160-200 °C in an electrochemical cell using a NaAlCl₄ molten salt as an electrolyte [114]. It can be intercalated with lithium to form Li_{0.7}NbS₂Cl₂ [420].

The tantalum dimer $Ta_2Cl_6(SMe_2)_3$ has been found to react with the bulky chelating ligand 1,3-bis(disopropylphosphino)propane (dippp) to form [(dippp)TaHCl]₂(μ -S)(μ -H)₂ [474]. The bonding was deduced form spectroscopic data and the solved X-ray structure.

Dark brown paramagnetic, triangulo niobium(IV) clusters, (Cp')₃Nb₃Cl₃(μ_2 -Cl)₃(μ_3 -O)(μ_3 -X), with Cp' = C₅H₄Me (methylcyclopentadienyl) and X = OH or Cl have been prepared. X-ray structure determinations show that these clusters contain a Nb₃ triangle bridged by Cl atoms along each edge and capped by a μ_3 -oxo ligand and a μ_3 -X ligand on opposite sides of the Nb₃ plane. Terminal Cp' and Cl ligands complete the coordination sphere about each Nb atom. Bonding in the clusters is discussed, based on their 10-line ESR spectra (the three d^1 electrons form an S = 1/2 system, coupled to one Nb nucleus with I = 9/2) [421-423].

Tantalum(IV) complexes, (Cp')₂Ta₂X₄(μ_2 -X)₂, with Cp' = e.g. C₅H₄Me (methylcyclopentadienyl) and X = Cl or Br have been prepared. X-ray structures were determined, including their reactivity as precursors to the doubly bonded (Cp')₂Ta₂(μ -X)₄ tantalum(III) complexes. Bonding in the clusters is discussed [475].

4.5 Nb(IV) and Ta(IV) oxides

By use of mass spectrometry and a Knudsen cell, the gaseous molecule NbO₂ has been characterized thermodynamically with respect to dissociation stability [424]. Thermodynamic data for gaseous tantalum(IV) oxide, TaO₂, were reviewed and data given also for a large number of liquid and solid tantalum halide and oxyhalide compounds [20].

Solid NbO₂ was studied theoretically by molecular orbital model calculations on small clusters in order to explain the photo-electron spectra [425]. MO level schemes and atomic charges of [NbO₅]⁶- and [NbO₆]⁸- have been given [425]. Thermogravimetric measurements have shown that solid NbO₂ is a metal-deficient oxide, whose main defects are neutral niobium vacancies in the rutile structure [426]. Rutile Nb_{0.94}O₂ was prepared by shock reduction of Nb₂O₅, like previously rutile TaO₂ was prepared from Ta₂O₅ [427]. Recovered products were examined by X-ray powder diffraction analysis [427]. TaO₂ is unstable, but can be studied in (Ti,Ta)O₂ rutile solid solutions. Crystal field splittings for Ta have been studied in such phases [428].

A new mixed-oxidation-state compound NaNb(II)Nb(IV)₂O₅F has been prepared. Single crystals are *dia*magnetic and black with metallic luster. The structure was determined and it shows short Nb-Nb distances [429].

The compound $(Ti^{3+})_2(Ti^{4+})_{1-x}(Nb^{4+})_xO_5$ with x=0.075 was examined magnetically versus temperature [430]. Nb(IV)-Nb(IV) pairs were discovered in new mixed valence oxides $KTi_{2-x}Nb_{5+x}O_{17}$ rutile crystals $(0 \le x \le 1.75)$ [431].

The product EuNbO3 of the reaction between EuO and NbO2 has been characterized by X-ray $L_{\rm III}$ absorption spectroscopy. The oxidation states were proven to be Eu²⁺ and Nb⁴⁺ [293].

4.6 Nb(IV) and Ta(IV) complexes with O-donors

A red *tetrakis*-complex of niobium(IV), NbL₄, with L = the pyridine-2-caboxylate anion, 15, was prepared in dry benzene. The compound was analysed and characterized as a non-ionic *mono*mer Nb(IV) complex by iodine titration, by molecular weight and conductivity determinations. Electronic and IR spectra as well as magnetic susceptibility were reported [363].

Crystallographic analyses of two salts, $K_2(H_3NCH_2CH_2NH_3)[Nb(C_2O_4)_4]\cdot 4H_2O$ and $K_4[Nb(C_2O_4)_4]\cdot 3H_2O$, have shown that the *tetrakis*(oxalato)niobium(IV) anion, $Nb(C_2O_4)_4^{4-}$, can be either square *anti*prismatic or dodecahedral in the solid state [432].

The new niobium(IV) nonamethoxide anion, $[Nb_2(OMe)_9]^-$, has been characterized in three X-ray crystallographic structure solutions; it has a confacial *bi*octahedral geometry of virtual D_{3h} symmetry with Nb-Nb bonding and three O-bridging methoxy groups [433].

4.7 Nb(IV) and Ta(IV) sulphides and selenides

The layered compounds MX_2 , with M = Nb, Ta and X = S, Se tend to be of interest in many contexts. The M atoms and the chalcogen atoms are in hexagonal close-packed layers; layers of M are sandwiched between layers of chalcogen, see Figure 54. Nb and Ta atoms display both octahedral and trigonal-prismatic coordination [16].

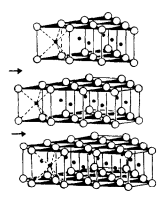


Figure 54. Schematic representation of the MX₂ type structure. Intercalation can take place as indicated by arrows. The drawing is adapted with permission by [347], D. D. Keszler, J. A. Ibers, S. Maoyu and L. Jiaxi, *J. Solid State Chem.* 57 (1985) 68.

Phases NbS_x with x = ca. 1.5 or 1.7 are obtained often when preparing NbS₂ from Nb₂O₅ and H₂S at 700 °C [434]. NbS_xSe_{2-x} layer crystals, with different *polytype*-structures depending on x, for $0 \le x \le 2$, were prepared and studied by X-ray diffraction [435].

The properties of NbS₂ were studied by X-ray photoelectron spectroscopy and by magnetic susceptibility [436]. The kinetics of the oxidation of NbS₂ to Nb₂O₅ has been studied thermogravimetrically as a function of the partial pressure of oxygen [437].

Inner-shell-electron energy-loss spectroscopy [438], scanning tunneling microscopy [439] and 181 Ta Mössbauer spectroscopy [440] were used to obtain information on electronic structure and atomic arrangements of 2*H*-TaS₂. In Li intercalated 2*H*-TaS₂, the electronic configuration of Ta changes form $Ta^{4+}(5d^{1})$ toward $Ta^{3+}(5d^{2})$ [440]. Electronic structures of other 2*H*-TaS₂ intercalates are given in [441]. Preparational details for 2*H*-TaS₂ intercalates have been reported in [442] and 181 Ta Mössbauer spectra given in [443]. The superstructure in 4*Hb*-TaS₂ was studied by high resolution electron microscopy [444].

A phase transition in 1*T*-TaS₂ involving charge density waves has been studied by X-ray scattering [445-446], electron diffraction [447], ¹⁸¹Ta NMR and NQR spectroscopy [448], magnetic methods [449], thermal methods [450], and high resolution electron and scanning tunneling microscopy [439,451-452]. The transition is hydrogen impurity dependent [453-454]. The special interest of the study is that the phase transition which clearly could be followed on precession photographs does not involve the underlying crystalline structure, only the waves. Similar waves occur in 1*T*-TaSe₂, see *e.g.* [455].

It has been shown that silver ions can be intercalated between the layers in 2H- MS_2 [456] (M = Nb, Ta). In such crystals, grown using vapour transport with chlorine as transport agent, short Ag-Nb or Ag-Ta distances, determined by X-ray diffraction methods, suggest interaction between the 5s-orbital of silver and the lowest unoccupied (or partly occupied) d_z 2-orbital of the trigonal-prismatically coordinated transition metal. The stability of intercalated phases Ag_xMS_2 was studied [456] by EMF measurements and compared with the similar alkali metal intercalates. Also, lithium, copper, silver, iron, tin, zink and many others [457] have been intercalated into NbS2 [458] [459-464] and NbSe2 [465-466] and into molybdenum substituted niobium dichalcogenides [467].

TaS₂ has been intercalated with hydrogen [468], lithium [441-442, 469, 476], sodium [442-443], lead [441], tin [441] and substituted with molybdenum [477], and with rare earth metals (La and Ce), forming several new phases with interesting superconductivity [478]. TaS₂ can be intercalated with ammonia [479-480], pyridine [481-482] and with metalhydroxides [483].

Co-intercalation of alkali metal hydroxides and water molecules into NbS₂ and TaS₂ can be made, according to chemical analyses, X-ray diffraction and NMR spectroscopic investigations [484].

Crystals of Nb₂Se₉ have been re-examined by structure solution and classified as containing Nb(IV); hence the formula can be specified as $2\text{Nb}^{4+2}(\text{Se}_2)^{2-}(\text{Se}_5)^{4-}$, in accordance with the one-dimensional semiconducting properties. The structure is based on chains of niobium in bicapped trigonal prismatic coordination with selenium and has short Nb-Nb bonds [485].

Also, the CuTaS3 crystal structure solution has given the valence description as Cu²⁺Ta⁴⁺3S²⁻ [485]. A new ternary chalcogenide, Ta₂Cu_{0.80}S₆, has been prepared and its crystal structure solved [486]. It contains S₂²⁻ pairs and the Ta atoms must then have a formal valency of IV. It shows a close relation to the CuTaS3 structure, containing units of infinite chains of CuS4 tetrahedra and TaS₆ octahedra [486].

The compounds Nb₂PdSe₆, Nb₂Pd_{0.71}Se₅, Nb₃Pd_{0.72}Se₇, Ta₂NiS₅ and Ta₂NiSe₅ have been prepared and examined [16, 340-341]. They contain new kinds of layered and channel structures, see *e.g.* Figure 55. They have interesting electric and magnetic properties, and exhibit structural phase transitions. Ta₂NiS₅ have been intercalated topotactically with hydrazine, 1,2-diamino-ethane and 1,3-diamino-propane and the inter-layer distance enlargements studied by X-ray diffraction [487].

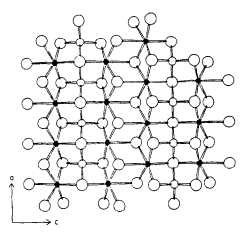


Figure 55. The *ortho*rhombic *Cmcm* Ta₂NiS₅ structure in the (a,c) plane. Octahedral tantalum and tetrahedral nickel are shown, surrounded by sulphur (or selenium) in layers three atoms thick. Reproduced with permission from [340], F. J. Di Salvo, C. H. Chen, R. M. Fleming, J. V. Waszczak, R. G. Dunn, S. A. Sunshine and J. A. Ibers, *J. Less-Comm. Met.* 116 (1986) 51.

Furthermore, it was found that e.g.the reaction of Ta₂NiSe₅ and Ta₂PdSe₆ with n-butyllithium affords the compounds Li₂Ta₂NiSe₅ and LiTa₂PdSe₆ which have layered structures [488].

A paramagnetic compound, NbNiTe5, with metallic conductivity and a new kind of layer structure has been prepared [489]. Each layer consists of bicapped trigonal prismatic niobium atoms and octahedral nickel atoms coordinated by tellurium atoms, see Figure 56. For comparison, note the structure of Ta2NiS5 and Ta2NiSe5, Figure 55.

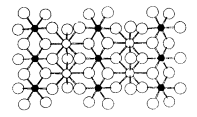


Figure 56. View of one layer of the layer structure of NbNiTe₅, showing the coordination around the Nb atoms (small open circles) and Ni atoms (small filled atoms). Reproduced with permission from [489], E. W. Liimatta and J. A. Ibers, *J. Solid State Chem.* 71 (1987) 384.

In the compounds NbTe₂ and TaTe₂ the metals occupy octahedral positions [16]. NbTe₂ reacts with PCl₅, giving NbCl₅ and PCl₃ [490].

4.8 Other Nb(IV) and Ta(IV) complexes

Complexes [Nb(por)(O)] and [Nb(por)(X)₂] (in which por is a porphyrinate²⁻ anion: oep = octaethylporphyrinate, tpp = tetraphenylporphyrinate or tptp = tetraparatolylporphyrinate and X = Cl or Br) have been prepared, by reduction of [Nb(por)(X)₃] or [Nb(por)(O)(OAc)] [391]. The compounds were characterized on the basis of IR, UV-VIS, electrochemical and EPR data [391]. The [Nb(por)(X)₂] complexes have axial symmetry [391].

4.9 Nb(IV) and Ta(IV) complexes with N- and C-donors

Ammonolysis of tantalum alkyls has been studied in order to obtain a safe procedure for preparing solid state materials. In this way cubic TaN and a *tri*meric nitride, $((Cp')MeTaN)_3$ $(Cp' = methylcyclopentadienyl and Me = methyl) were obtained. The crystal structure of the latter compound was solved, see Figure 57, and other properties elucidated. The compound can be reduced to form <math>[K(nEt_2O)]^+[((Cp')MeTaN)_3]^-$. Molecular orbital diagrams are given [491].

The formation of vacancies in niobium carbides has been studied theoretically [492].

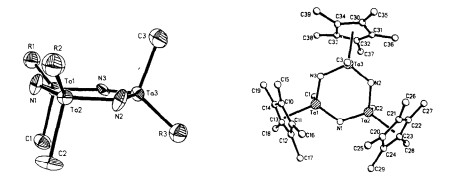


Figure 57. Crystal structure of the *tri*meric nitride molecule, ((Cp')MeTaN)₃ (Cp' = methylcyclopentadienyl and Me = methyl) [491]. Reproduced with permission from [491], M. M. B. Holl, M. Kersting, B. D. Pendley, and P. T. Wolczanski, *Inorg. Chem.* 29 (1990) 1518.

5. Mixed Nb(IV) and Nb(III) complexes (d^1 and d^2)

A new mixed valence (Nb(III) and Nb(IV)) compound of unknown structure, $(C_5Me_5)_3Nb_3S_7$), has been obtained by letting the Nb(I) cyclopentadienyl derivate (η^5 -C5Me5)Nb(CO)4 and sulphur react in tetrahydrofuran solution under irradiation [388].

6. Nb(III) and Ta(III) complexes (d²)

6.1 Nb(III) and Ta(III) halides and halide complexes

Thermodynamic data for gaseous tantalum halides, TaF3, TaCl3, TaBr3, and TaI3 were reviewed and data given also for a large number of liquid and solid tantalum halide and oxyhalide compounds [20].

The thermodynamic properties of crystalline TaCl₃ were studied, and the composition of the vapour phase in equilibrium with it was determined. The vapour contained TaCl₅, TaCl₄, TaCl₃, TaCl₂, TaCl, Ta, Cl, and Cl₂. The heats, free energies and entropies of sublimation were determined [493].

It has been shown that electrochemically formed [TaCl₄]⁻ ions, in LiCl-KCl-TaCl₅ molten mixtures at 400-550 °C, by fluoride ion addition are unstable and disproportionate to primarily [TaF₇]²- and metallic tantalum [60].

6.2 Nb(III) and Ta(III) chalcogenohalide complexes

Thermodynamic data for gaseous tantalum(III) oxyhalides, TaOF, TaOCl, TaOBr, and TaOI were reviewed and data given also for a large number of liquid and solid tantalum halide and oxyhalide compounds [20].

The selenoiodide NbSeI has been prepared and classified as Nb₄Se₄I₄, a cubic cluster compound containing tetrahedral Nb₄Se₄ units with niobium-niobium bonding, according to the results of a crystal structure solution [494]. The structure has a close relation to the new GaNb₄Se₈ structures, see Figure 69. Nb₄Se₄I₄ can be intercalated with lithium to form Li_{2.8}Nb₄Se₄I₄ [420].

6.3 Nb(III) and Ta(III) halides with O- and chalcogen-donors

With Nb₂Cl₆(THT)₃ as starting material (THT = tetrahydrothiophene), dimeric Nb(III) alkoxide compounds, Nb₂Cl₅(OR)(HOR)₄ (R = Me, Et, Prⁱ), have been made [495]. According to crystallographic examination, these compounds have edge-sharing bioctahedral geometries in which the Nb atoms are bridged by one chloride and one alkoxide ligand. Nb₂Cl₅(OMe)(MeOH)₄ has a short Nb-Nb distance, indicative of a metal-metal double bond [495]. For R = Et and Prⁱ, the compounds are unstable in THF, forming tetranuclear Nb(IV) complexes, [Nb₂OCl₄(OR)₂(THF)₂]₂ [495], see Figure 58. The compounds have been further characterized in solution by proton NMR spectroscopy [495].

A dinuclear Nb(III) complex salt, (NMe₄)[Nb₂Cl₂(THT)(CH₃CO₂)₅]·CH₂Cl₂ with THT = tetrahydrothiophene has been characterized by NMR, IR, cyclic voltammetry and X-ray crystallography [496]. The Nb-Nb distance and the diamagnetism indicated a double metal-metal bond. Acetate is coordinated in three different modes: Bridging bidentate, chelating, and bridging uni-dentate, see Figure 59.

Niobium and tantalum face-sharing dimers $M_2\text{Cl}_6(\text{SMe}_2)_3$ (M=Nb or Ta, and with two bridging Cl) have been found to react with dimethylsulphide, SMe₂, to form edge-sharing bioctahedra $M_2\text{Cl}_6(\text{SMe}_2)_4$ [497]. The crystal structures were compared to those of similar complexes in which dimethylsulphides were replaced by 3,6-dithiaoctane, in an attempt to understand the M-M bonding schemes in these complexes [497]. Also, single crystal structures of di- μ -chloro-tetrachloro-bis--(tetrahydrofuran)-[μ -(dimethylsulphide)]-di-niobium(III),

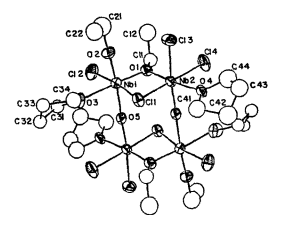


Figure 58. Crystal structure of the tetranuclear [Nb2OCl4(OEt)2(THF)2]2 molecule [495]. Reproduced with permission from [495], F. A. Cotton, M. P. Diebold and W. J. Roth, *Inorg. Chem.* 26 (1987) 3323.

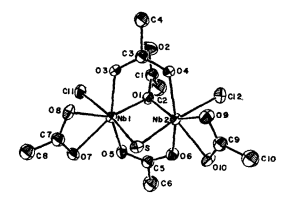


Figure 59. View of the [Nb₂Cl₂(THT)(CH₃CO₂)₅]⁻ anion. Carbon atoms in the bridging THT ligand have been omitted for clarity. Reproduced with permission from [496], F. A. Cotton, M. P. Diebold, M. Matusz and W. J. Roth, *Inorg. Chim. Acta* 112 (1986) 147.

[Nb₂Cl₆(C₄H₈O)₂(C₂H₆S)], and the analogous tantalum compound have been solved [498]. The coordination geometry can here be described as confacial-bioctahedral, see Figure 60, with a double metal-to-metal bond and a coordination number of 7 around the metal.

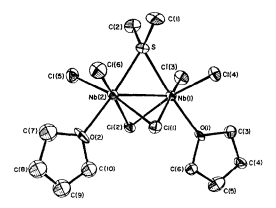


Figure 60. View of the [NbCl₂(THF)]₂(μ-Cl)₂(μ-SMe₂) molecule. Reproduced with permission from [498], F. A. Cotton, S. A. Duraj and W. J. Roth, *Acta Cryst.* C41 (1985) 878.

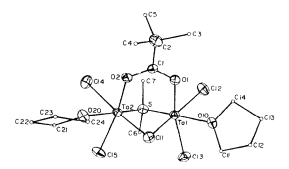


Figure 61. Structure view of the [TaCl₂(THF)]₂(μ-SMe₂)-(μ-O₂CC₄H₉) molecule. Reproduced with permission from [499], F. A. Cotton, M. P. Diebold, S. A. Duraj and W. J. Roth, *Polyhedron* 4 (1985) 1479.

A binuclear tantalum(III) complex with a bridging carboxylato ligand, Ta₂Cl₅(O₂CC₄H₉)(SMe₂)(THF)₂, has been obtained by letting Ta₂Cl₆(SMe₂)₃ react with C₄H₉CO₂Li [499]. The complex is a dimer with a distorted octahedra-sharing-an-edge geometry and some Ta=Ta bonding, see Figure 61. It is interesting to note that the existence

of this green compound proves that low-valent tantalum can be stable in contact with oxygen-donor ligands, and moreover that this is the case even when the *dimer* contains a bridging *bidentate* ligand.

For the crystal structure of di- μ -chloro-tetrachloro-bis-(tetrahydrofuran)-[μ -(dimethylsulphide)]-di-tantalum(III), [Ta₂Cl₆(C₄H₈O)₂(C₂H₆S)], see the analogous niobium compound and Figure 60 [498].

6.4 Nb(III) and Ta(III) halides with N- and P-donors

Relativistic MS $X\alpha$ molecular orbital calculations have been performed under $C_{2\nu}$ symmetry on di- μ -chloro-tetrachloro-tetra-ammine tantalum(III) compounds [Ta₂Cl₆(NH₃)₄] and [Ta₂Cl₆(NH₃)₄] [500]. These complexes were used as models for Ta₂Cl₆(γ -pic)₄ and short-lived anion species obtained from this [501] by pulse radiolysis reduction (γ -pic = γ -picoline = 4-methylpyridine). Calculated electronic transitions compared to experimental absorption spectra have lead to the conclusion that the transient Ta dimer has a formal bond order of 2.5 or 1.5 (electronic configuration $\sigma^2 \pi^2 \delta^*$) [135,500].

Bis-nitrogen-chelated, edge-sharing bioctahedral molecules $[M_2Cl_6(Et_2NCH_2CH_2NEt_2)_2]$ (M = Nb, Ta) with two bridging chlorides and four uni-planar terminal chlorides have been isolated, and their crystal structures determined [502]. These molecules have five membered M-N-C-C-N rings and are in the achiral meso conformation of C_{2h} symmetry with one ring of λ and the other ring of δ conformation, and the twofold axis passing along the metal-metal vector. For the phosphorous substituted analogous molecules $[M_2Cl_6(Et_2PCH_2CH_2PEt_2)_2]$ (M = Nb, Ta) chiral conformations of D_2 symmetry have been isolated [502-503], see Figure 63.

Ions Nb(III) and Ta(III) in the presence of hydroxide form binuclear complexes with nitrogen, in which the N₂ molecule is significantly activated and can be converted to hydrazine and ammonia [504].

A dinuclear organoimido niobium(III) complex, Nb₂Cl₆(NPh)₂(SMe₂)₂, has been synthesized by adding triphenylazide to a solution of the metal-metal double bonded dimer Nb₂Cl₆(SMe₂)₃ in toluene at 0 °C [505], in a simpler way than previously [506]. The compound is air-sensitive and can be described as [NbCl₂(NPh)(SMe₂)]₂(μ-Cl)₂ (see the analogous tantalum compound depicted in Figure 22). Its reactivity was investigated, see reference [505].

An exeptionally reactive 16 electron tantalum(III) *mono*mer, TaClH₂(PMe₃)₄, has been made and characterized with respect to its neutron structure, IR-, ¹H- and ³¹P[¹H] NMR spectra, and reactivity [507]. In the solid state, the coordination polyhedron is a slightly distorted pentagonal *bi*pyramid with two axial phosphines and the hydride ligands in *cis* coordination, see Figure 62.

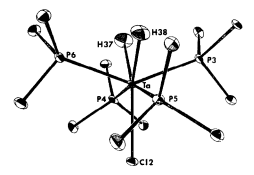


Figure 62. The neutron diffraction structure of the Ta(III) compound TaClH₂(PMe₃)₄ [507]. Methyl hydrogens have been omitted for clarity. Reproduced with permission from [507], M. L. Luetkins, Jr., M. D, Hopkins, A. J. Schultz, J. M. Williams, K. C. Fair, F. K. Ross, J. C. Huffman and A. P. Sattelberger, *Inorg. Chem.* 26 (1987) 2430.

TaClH₂(PMe₃)₄ reacts with 1,2-bis(dimethylphosphino)ethane (Me₂PCH₂CH₂PMe₂ = dmpe) to give TaClH₂(dmpe)₂, with ethylene to give TaCl(C₂H₄)(PMe₃)₄, with dinitrogen to give the μ -dinitrogen complex [TaClH₂(PMe₃)₃]₂(μ -N₂) and with carbonmonoxide to give TaCl(CO)₃(PMe₃)₃ [507]. The crystal structure of TaCl(CO)₃(PMe₃)₃ has been solved based on X-ray data [507].

Bis-phosphorous-chelated, edge-sharing bioctahedral molecules $[Ta_2Cl_6(Et_2PCH_2PEt_2)_2]$ with two bridging chlorides and four uni-planar terminal chlorides have been isolated, in diastereomeric forms, and their crystal structures determined [503]. The difference between these molecules is in the conformations of the five membered Ta-P-C-C-P rings. Form I is the achiral meso compound of C_{2h} symmetry with one ring of λ and the other ring of δ conformation, and the twofold axis passing along the metal-metal vector. Form 2 is chiral of D_2 symmetry, with both rings in λ conformation and the twofold axes intersecting at the midpoint of the Ta-Ta bond. Form 2 has also been isolated for the case of niobium [502-503], see Figure 63.

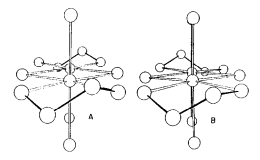


Figure 63. The structure of the two different ligand conformations in the *dinuclear* [Nb₂Cl₆(Et₂NCH₂CH₂NEt₂)₂] (A) and [Nb₂Cl₆(Et₂PCH₂CH₂PEt₂)₂] (B) molecules [502]. Reproduced with permission from [502], J. A. Canich and F. A. Cotton, *Inorg. Chem.* **26** (1987) 4236.

Discrete *tri*nuclear complexes of niobium(III) and tantalum(III) have been described: $(PEt_3H)^+[Nb_3Cl_{10}(PEt_3)_3]$ and $(PEt_3H)^+[Ta_3Cl_{10}(PEt_3)_3]$ [508]. A related compound, $Nb_3Cl_7(PMe_2Ph)_6$, with a nonintegral oxidation state was also described. The structures have been established by X-ray crystallography, see Figure 64. In each case there is an $M_3(\mu_3-Cl)(\mu_2-Cl)_3$ core surrounded by the remaining nine ligands. Six and eight electrons are available for M-M bonding, according to Fenske-Hall molecular orbital calculations [508].

Finally, a discrete tetranuclear tetrahydrothiophene ethylene-bis(diphenylphosphine) complex of tantalum(III) has been described: [Ta₂Cl₄(μ-Cl)₂(μ-SC₄H₈)]₂(μ-Ph₂PCH₂CH₂PPh₂)₂·C₇H₈ [509]. The structure consists of two confacial bi-octahedral units of [Ta₂Cl₄(μ-Cl)₂(μ-SC₄H₈)] (with Ta=Ta double bonds, see Figure 60), bridged by two Ph₂PCH₂CH₂PPh₂ ligands to form the unusual tetrameric units. The structure was established by X-ray crystallography on the red crystals [509].

6.5 Nb(III) and Ta(III) halides with C-donors

The reaction between niobium(V)chloride and phenyl-acetylene was investigated in carbon tetrachloride [510]. Two mols of phenylacetylene were found to react with one mol NbCl5 forming first a five-coordinated complex (Figure 65, I) which in turn reacts with more

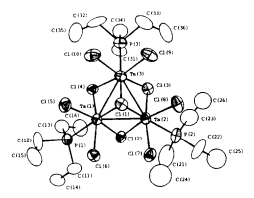


Figure 64. The structure of the [Ta₃(μ₂-Cl)₃Cl₆(PMe₃)₃]⁻ ion [508]. Reproduced with permission from [508], F. A. Cotton, M. P. Diebold, X. Feng and Wieslaw J. Roth, *Inorg. Chem.* 27 (1988) 3413.

phenyl-acetylene to form an unstable π -bonded complex (Figure 65, II), and finally catalytically produces NbCl₃ and oligomerized (mainly *tri*merization) of the phenylacetylene.

NbCl₅ + 3C₆H₅C=CH

$$Cl$$
 Cl
 $CHCl$
 Ph
 $CHCl$
 Ph
 $CHCl$
 $CHCl$
 Ph
 $CHCl$
 $CHCl$

Figure Cyclotrimerization of phenylacetylene by NbCl₅. Adapted 65. from [510] C. J. du Toit, J. A. K. du Plessis and G. Lachmann, S. Afr. J. Chem. 38 (1985) 195.

A tantalum(III) species, probably $Cl_3Ta[(C_6H_5)C=C(H)Cl]_2$, 21, is formed during the TaCl₅-catalysed cyclotrimerization of phenyl-acetylene $C_6H_5C\equiv CH$ [40]. The complex 21 shown above is assumed to react with more phenylacetylene forming triphenylbenzene and TaCl₃ [40].

A new unique dinuclear Niobium(III) cation, $[Nb_2Cl_3\{(PhC)_4\}(THF)_4]^+$, has been prepared and characterized by solving the X-ray crystal structure of a salt [511]. The cation has a short (2.695 Å) Nb=Nb double bond and bridging as well as terminal chlorine atoms. A C-shaped PhC-C(Ph)C(Ph)-CPh chain clasps the Nb=Nb bond. The structure is shown in Figure 66.

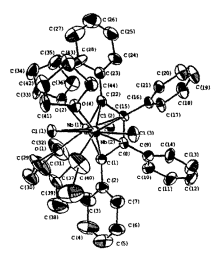


Figure 66. The structure of the unique dinuclear Nb(III) cation, [Nb₂Cl₃{(PhC)₄}(THF)₄]⁺ [511]. Four tetrahydrofuran groups (left side) and four C-phenyl groups (right side) are shown in full. Reproduced with permission from [511], F. A. Cotton and M. Shang, *Inorg. Chem.* 29 (1990) 2619.

Another new unique tetranuclear Nb(III) anion, [Nb₄OCl₈{(PhC)₄}₂]²⁻, has been prepared and characterized by solving the X-ray crystal structures of several different salts [511-512]. The anion consists of a planar, rectangular Nb₄ group with an oxygen atom at the center. The long Nb··Nb edges are doubly bridged by Cl atoms, and there is one terminal Cl atom on each

Nb atom. A C-shaped PhC-C(Ph)C(Ph)-CPh chain clasps each short Nb··Nb edge. The mode of bonding is shown in Figure 67. The oxidation state of Nb is most likely to be +III, and hence two Nb=Nb double bonds can be formed pertetramer unit, in accordance with the short (2.611 Å) Nb-Nb distance. The central oxygen atom in this anion has an environment that is without precedent. Note the relation between the two ions in Figures 67 and 66.

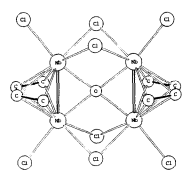


Figure 67. The structure of the unique tetranuclear Nb(III) anion, [Nb4OCl8{(PhC)4}2]²⁻, as found in several salts [511-512]. Phenyl groups are omitted for clarity. Reproduced with permission from [511], F. A. Cotton and M. Shang, *Inorg. Chem.* **29** (1990) 2619.

Finally, a hexanuclear Nb(III) compound of composition (Bu) $_3$ Sn[Nb₆Cl₁₉(PhC)₁₂]·C₇H₈, has been made and characterized by its X-ray crystal structure [513]. The hexanuclear entity consists of a central triangular Nb₃(μ ₃-Cl)(μ -Cl)₃ cluster with a Cl₃NbCl₂(c-C₄Ph₄) unit of roughly octahedral shape (if c-C₄Ph₄ is considered as one ligand) attached to each central Nb atom by three mutually cis Cl atoms. The Nb-Nb distances in the central cluster (2.87 Å) are indicative of Nb-Nb single bonds and there is no further Nb-Nb bonding. The mode of bonding is shown in Figure 68. The oxidation state of Nb is considered to be +III.

6.6 Nb(III) and Ta(III) oxides, sulphides and selenides

Electrode reaction equilibria of involving niobium(III) complexes were studied potentiometrically and polarographically in perchlorate and sulphate acidic solutions [304-305]. A two electron reaction:

$$Nb^{5+} + 2e^{-} = Nb^{3+}$$

occurred at the mercury electrode, the Nb(III) complexes probably being Nb^{3+} , $NbSO_4^+$ or $Nb(OH)_2^+$. Equilibrium constants were given [304]. An intermediate Nb(IV) complex might also be formed [305].

Several bis-oxo-capped M_3X_{17} clusters (M = Nb and X = O) have now been obtained and their crystal structures solved [117]. These complexes have a core of three Nb atoms and two oxygens forming a triangular bipyramid. Each niobium are further bound to five outer-sphere ligands. The structures have been solved for the cases of $[\text{Nb}_3\text{O}_2(\text{SO}_4)_6(\text{H}_2\text{O})_3]^{5-}$, $[\text{Nb}_3\text{O}_2(\text{O}_2\text{CBu}^1)_6(\text{THF})_3]^+$, and $[\text{Nb}_3\text{O}_2(\text{O}_2\text{CCH}_3)_6(\text{THF})_3]^+$. In all these compounds, six chelating groups provide twelve ligands, and three monodentate groups provide the last three oxygens [117].

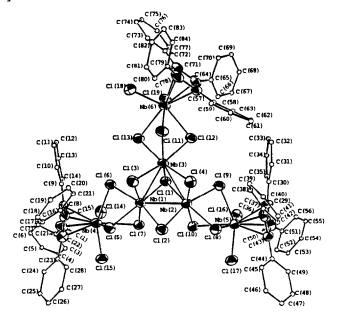


Figure 68. The unique hexanuclear Nb(III) compound (Bu)₃Sn[Nb₆Cl₁₉(PhC)₁₂]·C₇H₈, characterized by its X-ray crystal structure [513]. The entity consists of a central triangular Nb₃(μ₃-Cl)(μ-Cl)₃ cluster with three roughly octahedral Cl₃NbCl₂(c-C₄Ph₄) units attached to central Nb atoms by Cl atoms. Reproduced with permission from [513], F. A. Cotton and M. Shang, *Inorg. Chem.* 29 (1990) 2614.

The preparation and crystal structure of Nb₂S₃ are discussed in [514].

New GaNb₄S₈ and GaNb₄Se₈ cluster compounds of Niobium(III), have been prepared by heating the elements in closed silica tubes [494]. According to the results of crystal structure solutions, the compounds contain tetrahedral Nb₄S₄ or Nb₄Se₄ units with a Nb₄ cluster as the core and with niobium-niobium bonding, see Figure 69. The structure has a close relation to the AB₂X₄ spinel structures and the new Nb₄Se₄I₄ structure [494].

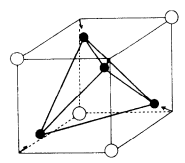


Figure 69. New Nb4Se₄I₄, GaNb₄S₈ and GaNb₄Se₈ cluster compounds of Niobium(III) contain tetrahedral Nb₄S₄ or Nb₄Se₄ units with a Nb₄ core. Filled circles = Nb, open circles = S or Se. Reproduced with permission from [494], H. B. Yaich, J. C. Jegaden, M. Potel, M. Sergent, A. K. Rastogi and R. Tournier, J. Less-Comm. Met. 102 (1984) 9.

6.7 Nb(III) and Ta(III) nitrides

In the vapour phase, NbN, molecules have been characterized by spectroscopy (molecular constants for upper $^3\Phi$ and ground $^3\Delta$ states, obtained from vibrational and rotational spectra) [515-516].

Solid niobium nitride, NbN, has unusual properties such as being hard, brittle and a super-conductor below 16 K. Its electronic structure has been studied by high energy electron energy loss spectroscopy [387].

The formation of vacancies in niobium nitrides has been studied theoretically [492]. Non-stoichiometric compact cubic δ -NbN_{1-x} phases with small x values were prepared and investigated [517]. Energy band structure studies have been performed on NbN single crystals

using the linearized augmented-plane-wave method [518].

6.8 Nb(III) and Ta(III) C-donor complexes

The reaction between the Nb(III) cyclopentadienyl derivate $(\eta^5-C_5Me_5)_2NbBH_4$ and sulphur has been studied [388].

7. Niobium and tantalum clusters of oxidation state < III

The electronic bonding in Nb₆Cl₁₄, and, in general, in Nb₆Cl₁₂ L_6^{2+} clusters (with L as a ligand) was described using graph theory, by means of 76 valence electrons including 16 in eight face-localized three-center bonds [519].

In acidic aqueous solution, green $[Ta_6Cl_{12}]^{2+}$ cluster ions can be oxidized by nitrate, bromate, dichromate or Ce^{4+} to form $[Ta_6Cl_{12}]^{3+}$ and $[Ta_6Cl_{12}]^{4+}$ clusters in separate potentiometric steps, before the formation of Ta_2O_5 , xH_2O starts [520]. $[Ta_6Br_{12}]^{2+}$, $[Nb_6Cl_{12}]^{2+}$ and $[Nb_6Br_{12}]^{2+}$ behave in a rather similar way [520-521].

The ligand replacement reactions in (pyH)₂[Ta₆Br₁₂]Cl₆ at elevated temperature were followed by chemical analyses in steps; the chloride atoms penetrate into the interior of the cluster gradually, forming the ions [Ta₆Br₆Cl₆]⁴⁺, [Ta₆Br₂Cl₁₀]⁴⁺ and [Ta₆Cl₁₂]⁴⁺ in agreement with theoretical predictions [522].

The mixed oxidation state cluster ions $[Nb_6Cl_{12}]^{2+,3+}$ and $[Ta_6Cl_{12}]^{2+,3+}$ have been used for incorporation into Na-montmorillonite as a first step in obtaining new oxide materials by calcination [523].

Dark red solid cluster hydroxides $M_2[\text{Ta}_6\text{Cl}_{12}](O\text{H})_6 \cdot n\text{H}_2O$ with M = Na, K, Rb, (CH₃)₄N and (C₂H₅)₄N and n = 12 to 20 were prepared, and their stability towards air oxidation in alkaline methanol-water solution examined [524]. Insoluble [Ta₆Cl₁₂](OH)₄·10H₂O and soluble clusters [Ta₆Cl₁₂]X₄· $n\text{H}_2\text{O}$ and [Ta₆Cl₁₂]X₆²⁻, X = Cl, Br are the products of acidification with HCl and HBr [524]. The are no evidence that the OH⁻ ions enter into the inside coordination sphere of the cluster unit.

A new dodeca- μ -bromo-hexabromo-octahedro-hexaniobate(4-) salt, KGd[Nb₆Cl₁₈], containing (Nb₆Cl₁₂)Cl₆ clusters has been studied by crystal structure solution [525]. The average oxidation state of Nb is 7/3. The found geometry of the cluster is shown in Figure 70. A similar structure was found for the case of K₄[(Nb₆Br₁₂)Br₆] [526], isotypic with the previously found K₄[(Nb₆Cl₁₂)Cl₆] [527].

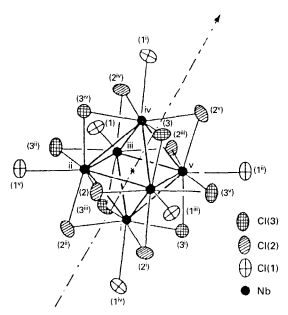


Figure 70. Structure of octahedral (Nb₆Cl₁₂)Cl₆ clusters surrounded by K⁺ and Gd³⁺ in the trigonal crystal. Reproduced with permission from [525], S. Ihmaïne, C. Perrin and M. Sergent, *Acta Cryst.* C43 (1987) 813.

Other salts, $R[Nb_6Cl_{18}]$, $R[Ta_6Cl_{18}]$, $MR[Nb_6Cl_{18}]$ and $MR[Ta_6Cl_{18}]$ with M=a mono valent ion and R=a rare earth, have been made [528]. All of these compounds crystallize in the trigonal system (rhombohedral space group $R\overline{3}$), except when M=Cs, for which cases the lattices are hexagonal (space group $P\overline{3} lc$) [528].

Alkoxides of the hexanuclear niobium and tantalum cluster units have now been made and studied: $[M_6X_{12}](OMe)_2 \cdot 4MeOH$ and $M'_2[Ta_6Cl_{12}](OMe)_6 \cdot 6MeOH$, M = Nb, Ta, X = Cl, Br, Me = methyl and M' = alkali metal [529]. Physical and spectral data are reported.

The photoelectron spectra (AlK α , HeI) of Nb₃I₈ (with trigonal Nb₃ groups) and the Nb₆I₈ cluster compounds Nb₆I₁₁, HNb₆I₁₁, Nb₆I₈(CH₃NH₂)₆ and Nb₆I₈(C₃H₇NH₂)₆ were studied in an attempt to find the relation between the binding energy of the niobium core levels and the oxidation number [42].

Clusters with chloro and trialkylphosphine ligands, $[(M_6Cl_{12})Cl_2\{P(R)_3\}4]^{n+}$, M = Nb, Ta, $R = C_2H_5$, C_3H_7 and C_4H_9 , and n = 0, 1 or 2 have been isolated in cis and trans isomers by column chromatography; structure determinations have shown that bond distances between the metals in the cluster depend not only on the oxidation state of the cluster but also on the kind of terminal ligands on each metal, see Figure 71. Electronic spectra of the charged clusters (n = 1 or 2) show bands indicating splittings of a metal cluster orbital $(t_{1u} \text{ or } t_{2g})$ due to lowering of the symmetry from O_h [530].

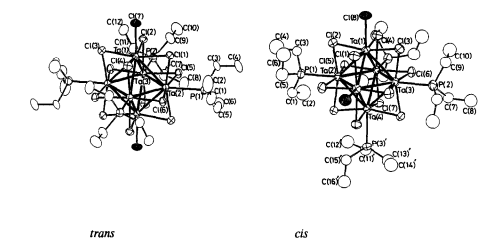


Figure 71. Structure of octahedral *trans* and *cis* isomers of [(Ta₆Cl₁₂)Cl₂{P(C₂H₅)₃}4]⁺ ions (from BF₄⁻ salts), isolated by column chromatography. Reproduced with permission from [530], H. Imoto, S. Hayakawa, N. Morita and T. Saito, *Inorg. Chem.* 29 (1990) 2007.

A new mixed-oxidation-state compound NaNb(II)Nb(IV)₂O₅F has been prepared. Single crystals are *dia*magnetic and black with metallic luster. The structure was determined and it shows short Nb-Nb distances [429].

Green crystals of Nb₃O₈ was prepared and their properties investigated [531]. The relation between Nb₃Cl₈ and other discrete *tri*nuclear niobium and tantalum clusters have been discussed [508].

Dark green single crystals of Mg₃Nb₆O₁₁ (with Nb of mean oxidation state 2.667) were prepared by heating a pellet containing MgO, Nb, Nb₂O₅ (15:14:8) at 1550 °C in a Nb-container. The X-ray structure was solved and found to contain clusters of Nb₆ octahedra, arranged in combination with other atoms to form cubic close-packed layers [532]. A similar structure was found for Mn₃Nb₆O₁₁ crystals, prepared in a somewhat different way [532].

Cluster compounds $Na_3Al_2Nb_34O_{64}$ and $Na(Si,Nb)Nb_{10}O_{19}$ with niobium in different oxidation states (+5, +4, +2.5 and +5, +4, +2.67, respectively) have been made and their crystal structures obtained by X-ray diffraction [533]. In both compounds, characteristic building units of Nb_6O_{12} and Nb_2O_{10} groups as well as isolated NbO_6 octahedra are found

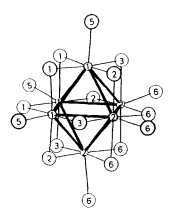


Figure 72. Structure of octahedral Nb₆O₁₂O₆ clusters found in Na(Si,Nb)Nb₁₀O₁₉. Large and small circles denote oxygen and niobium atoms.Reproduced with permission from [533], J. Köhler and A. Simon, Z. Anorg. Allgem. Chem. 553 (1987) 106.

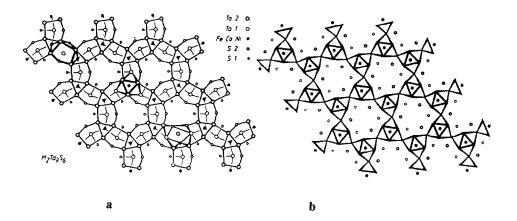


Figure 73. Projection of the M_2 Ta9S6 structure with M = Fe, Co, Ni down the c-axis, with emphasis on (a): the coordination polyhedra and (b): the condensed tantalum clusters, centered on the 3d metal. Reproduced with permission from [535], B. Harbrecht and H. F. Franzen, J. Less-Comm. Met. 113 (1985) 349.

in layers. The Nb₆ units are close-packed and surrounded by oxygen, forming a Nb₆O₁₂O₆ unit (see Figure 72), closely analogous with the Nb₆Cl₁₂Cl₆ unit known from *e.g.* K₄Nb₆Cl₁₂Cl₆. The stability of the compounds are discussed [533].

New metal-rich sulphides Ta₂S, Ta₆S, M_x Ta_{6-x}S (M = V, Cr; x = ca. 1), Fe₂Ta₉S₆, Ni₂Ta₉S₆ and Co₂Ta₉S₆ have been prepared through high-temperature techniques [534-537]. The hexagonal crystal M_2 Ta₉S₆ (M = Fe, Ni, Co) structures determined from X-ray data exhibit condensed tetrakaidecahedral tantalum clusters, accomodating the transition metal, surrounded by sulphur and having a channel structure, see Figure 73. Metal-metal interactions and the presence of S-lined channels in the M_2 Ta₉S₆ (M = Fe, Ni, Co) structures were studied in connection with extended Hückel electron structure calculations [537]. The Ta₆S and M_x Ta_{6-x}S crystals contain pentagonal *anti*prismatic tantalum columns surrounded by sulphur, see Figure 74 [536-536A].

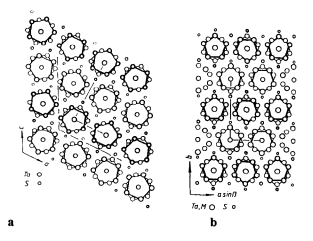


Figure 74. Projection of (a) Ta₆S (high temperature modification) and (b) M_x Ta_{6-x}S (M = V, Cr; x = ca. 1) structures, viewed with emphasis on the pentagonal antiprismatic metal columns (Ta, M, large circles). Reproduced with permission from [536], B. Harbrecht and H. F. Franzen, Z. Anorg. Allgem. Chem. 551 (1987) 74.

The technique of heating mixtures of metal sulphides or selenides in a tungsten Knudsen cell has been used for preparing new metal-rich transition metal sulphides and selenides. The following compounds have been made and their structures have been examined: Nb₁₄S₅, Nb₂₁S₈, Nb₂Se, Ta₂Se [536B], Ta₆S [536A], Ta₂S, Fe₂Nb₉S₆ and M'_2 Ta₉S₆ and M'_2 Ta₁₁Se₈ [563C] (M' = Fe, Co, Ni), as well as Ta_{6-x} M_x S. The structures can be thought of as modified metals with extended metal-metal bonding [538], and the oxidation state is not always known.

8. Nb(II) and Ta(II) complexes (d³)

Thermodynamic data for gaseous tantalum(II) halides, TaF₂, TaCl₂, TaBr₂, TaI₂ and TaO were reviewed and data given also for a large number of liquid and solid tantalum halide and oxyhalide compounds [20].

The thermodynamic properties of crystalline TaCl₂ were studied, and the composition of the vapour phase in equilibrium with it determined. The vapour contained TaCl₅, TaCl₄, TaCl₃, TaCl₂, TaCl, Ta, Cl and Cl₂. The heats, free energies and entropies of sublimation were determined [493].

Niobium and tantalum in K_2NbF_7 and K_2TaF_7 are reduced partly to oxidation state II after being bombarded by a 4 keV Ar^+ ion beam [412].

The He-I photoelectron spectra and first ionization energies of NbO and TaO molecules in the vapour phase have been obtained and assigned with the aid of Hartree-Fock-Slater calculations [539]. The optical spectrum of the gaseous free radical molecule NbO have been examined [540], and unusual intensity features have been found in its magnetic hyperfine structure [541]. By use of mass spectrometry and a Knudsen cell, the gaseous molecule NbO has been characterized thermodynamically with respect to dissociation stability [424]. Also, the NbO molecule was studied in argon and krypton matrix-isolation experiments (absorption and magnetic circular dichroism) at low temperatures [168]. Solid NbO, having a much discussed vacant NaCl type of structure [542], was shown to disportionate into NbO₂ and Nb at high temperatures and pressures [543].

The formation of vacancies in niobium oxide Nb₃O₃ has been studied theoretically [492].

The formation of chemical bonds in the hexagonal close-packed niobium carbide Nb₂C was studied by MO LCAO cluster calculations, and the results compared with the X-ray emission spectra [544].

The Nb(II) cyclopentadienyl compound, [(Cp)4Nb2(S)2] has been studied [545].

9. Nb(I) and Ta(I) complexes (d4)

Thermodynamic data for gaseous tantalum(I) halides, TaF, TaCl, TaBr, and TaI were reviewed and data given also for a large number of liquid and solid tantalum halide and oxyhalide compounds [20].

Nb(I) or Ta(I) complexes can be made from hexacarbonyl metalates of niobium(-I) or tantalum(-I), Na[M(CO)6] or Na[M(CO)6]·xTHF with M = Nb or Ta, x = ca. 0.5 to 5 and THF = tetrahydrofuran, by exclusively a two-electron transfer process using suitable agents such as CuCl, FeCl3, or bis(acetylacetonato)nickel(II) [546]. The metallate(I)complexes were Na[M2(μ -X)3(CO)8] with X = Cl, Br, I, the 1,2-bis(diphenylphosphino)ethane complexes [M(acac)(CO)3(Ph2P-CH2CH2-PPh2)], the cyclohexylisocyanide complexes [M(acac)(CO)3(CyNC)2], and [M(acac)(CO)4(THF)] [546]. The latter tantalum compound was studied by X-ray structure determination; tantalum(I) was hepta-coordinated, being surrounded by the bidentate acetylacetonato ligand, by four carbon monoxide groups and by the oxygen atom of the tetrahydrofuran ligand, see Figure 75. The THF ligand can easily be substituted together with one of the carbonyl groups [546].

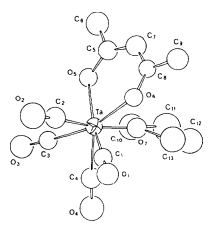


Figure 75. The [Ta(acac)(CO)₄(THF)] crystal structure. The tantalum(I) was found to be *hepta*coordinated, being surrounded by the *bidentate* acetylacetonato ligand, the *mono*dentate tetrahydrofuran ligand and the four CO molecules. Reproduced with permission from [546], F. Calderazzo, G. Pampaloni and P. F. Zanazzi, *Chem. Ber.* 119 (1986) 2796.

The reaction between the Nb(I) cyclopentadienyl carbonyl derivate (η^5 -C₅Me₅)Nb(CO)₄ and sulphur in tetrahydrofuran solution under irradiation has been studied [388]. In this way, a new mixed valence (Nb(III) and Nb(IV)) compound of unknown structure, (C₅Me₅)₃Nb₃S₇), has been obtained and some of its properties have been determined [388].

The structures of niobocene carbonyl chloride, π -(C₅H₅)₂Nb(CO)Cl, and niobocene carbonyl hydride, π -(C₅H₅)₂Nb(CO)H, were reported [547-548]. Both compounds have a wedged sandwich geometry with four-coordinate Nb(I) and with an eclipsed conformation of the cyclo*penta*dienyl (Cp) rings, see Figure 76.

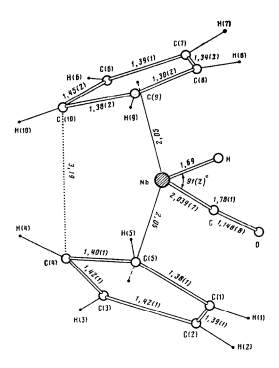


Figure 76. Structure view of the niobocene carbonyl hydride π-(C₅H₅)₂Nb(CO)H molecule. Niobocene carbonyl chloride π-(C₅H₅)₂Nb(CO)CI has a closely similar structure. Reproduced with permission from [547], A. S. Antsyshkina, L. M. Dikareva, M. A. Porai-Koshits, V. N. Ostrikova, Yu. V. Skripkin, A. A. Pasynskii, O. G. Volkov and V. T. Kalinnikov, *Russ. J. Coord. Chem.* 10 (1984) 872, *Engl. Transl.* of *Koord. Khim.* 10 (1984) 1560.

Gas phase reactions of Nb⁺ and Ta⁺ with alkanes and alkenes in a mass spectrometer have been reported. A long list of reaction products were identified, including the first *mononuclear* niobocene Nb(Cp)₂⁺ structure (Cp = cyclopentadienyl [549].

10. Nb(0) and Ta(0) complexes and alloys (d⁵)

Small clusters of niobium Nb_{χ} (x = ca. 5 - 10) were made and studied with a time-of-flight mass spectrometer [550]. Spectroscopic (photodetachment) data were given for Nb_2^+ and Nb_4^- [550].

The photoelectron spectrum (Al- K_{α} , HeI) of Nb was studied in an attempt to find the relation between the energy of the niobium core levels and the oxidation number [42].

Niobium and tantalum in K_2NbF_7 and K_2TaF_7 are partly reduced to oxidation state 0 after being bombarded by a 4 keV Ar^+ ion beam [412].

The formation of chemical bonds in the hexagonal close-packed niobium carbide Nb₂C and nitride Nb₂N was studied by MO LCAO cluster calculations, and the results compared with the X-ray emission spectra [544].

Enthalpies of formation and atomization for intermetallic compounds in the Al-Ta system (e.g. for TaAl₃, Ta₂Al₃, Ta₂Al and Ta₄Al) have been determined and compared with similar sulphur compounds [551].

Vapour phase reductions of NbCl₅ and GeCl₄ mixtures by hydrogen were studied at high temperatures. Under certain conditions the deposited products formed were super-conducting Nb₃Ge alloys [552].

In NbTe₄ and TaTe₄, the metals are in a square *anti*prism of Te atoms [16]. In the compounds NbTe₂ and TaTe₂ the metals occupy octahedral positions [16].

The Nb-Cr phase diagram was reviewed [553]. The intermediate compound Cr₂Nb has high and low temperature phases, whose structures and thermodynamics are given [553]. It was shown by self-consistent orbital calculations that the unknown compounds NbCe and NbLa should not be stable at ordinary pressure [554].

Crystal structure and magnetic properties of the *para*magnetic compounds NbCoSn, NbNiSn, TaCoSn and TaNiSn have been obtained [555]. Finally, NbNiTe₅ contains Nb in a *bi*capped trigonal prism of Te atoms [16].

11. Niobium and Tantalum in oxidation states below 0

Hexacarbonylmetallates of niobium(-I) or tantalum(-I), $Na[M(CO)_6]$ and $Na[M(CO)_6] \cdot x$ THF with M = Nb or Ta, x = ca. 0.5 to 5 and THF = tetrahydrofuran, have been prepared [556]. These compounds can be oxidized by suitable agents, exclusively by a two-electron transfer process to Nb(I) or Ta(I) complexes [546].

A new, safer synthetic procedure to carbonylate NbCl₅ forming the [Nb(CO)₆]⁻ anion has been found [557]. Dry pyridine was used as the solvent, and magnesium/zink as the reducing agent at room temperature and at atmospheric pressure of carbon monoxide. The hexa-carbonyl niobium(V) complex ion was isolated as the yellow-orange, tetrahydrofuran-stabilized

sodium salt [Na(C4H8O)][Nb(CO)6]. Other salts, such as the brick-red nickel-phenanthroline salt, [Ni(phen)3][Nb(CO)6], and the yellow μ -nitrido-bis(triphenyl-phosphorous[1+]) salt, [(PPh3)2N][Nb(CO)6] have been prepared. According to X-ray diffraction methods applied on the latter salt, the [Nb(CO)6]⁻ ion has an almost perfect octahedral geometry [557].

The ⁹³NMR spectrum of the [Nb(CO)₆]⁻ ion has been reported and the dependence upon *iso*topic replacement of ¹²C¹⁶O ligands by ¹³C¹⁶O and ¹²C¹⁸O studied. Also, IR and Raman spectra of the hexacarconyl anion were reported for the solid (Et₄N)[Nb(CO)₆] [558]. Furthermore, the ¹⁸¹Ta NMR spectrum of the [Ta(CO)₆]⁻ ion in solution was obtained and interpreted in terms of a perfect octahedral symmetry: Tantalum has a large quadrupolar moment and without the cubic environment the NMR band would be much broader than found [44].

The *penta*carbonylniobate(-III) and the corresponding tantalate *tri*anion form the lowest known oxidation states of Nb and Ta. The synthesis and chemistry of some $[M(CO)_5]^{3-}$ complexes, M = Nb and Ta, have been described in a dissertation [559].

12. Compounds of unknown oxidation state

Some compounds have already been referred which have uncertain oxidation states. Additional information is given here.

By use of mass spectrometry and a Knudsen cell, gaseous molecules EuNbO₃, EuNbO₂, EuNb₂O₆ have been characterized thermodynamically and shown to have a relatively high stability [424].

Quasi-one-dimensional conductors have been of much interest to physicists, because of their interesting properties in relation to charge density wave-type instabilities [560]. Some of the papers do certainly not contain coordination chemical information and are not referred.

In NbS₃ the structural unit is the *bic*apped trigonal prism [561]. When heated, NbS₃ looses one sulphur atom for every S₂ pair, leading to formation of a non-stoichiometric NbS₂ structural type [561]. A monoclinic NbS₃ phase can be obtained at high pressures and temperatures [562]. Stacking disoder in NbS₃ was studied by diffuse X-ray scattering [563]. Electronic, vibrational and resonance Raman spectra were obtained of the layered semiconducting NbS₃ compound [564].

The NbSe₃ structure contains selenium in tetrahedral and octahedral cavities; the octahedral cavities are large enough to contain also lithium atoms without significant deformation of the lattice. Such lithium intercalated phases have been prepared by treating NbX₃ (X = S or Se) with *n*-butyl-lithium in hexane. Li_XNbS₃ and Li_XNbSe₃ phases with x = 1 - 3 were characterized

by chemical analysis, X-ray powder diffraction and differential thermal analysis [562-565]; they might have a potential application as cathode material in lithium secondary batteries [514]. Optical investigations have been performed on the one-dimensional chain compounds NbSe₃ [566-567].

Thermal conductivity of whiskers of *ortho*rhombic NbS₃ and TaS₃ was studied versus temperature in the range 10-250 K, without any sharp anomaly being detected [568]. Monoclinic NbS₃ was prepared under high pressure, and it can be intercalated with lithium to form Li_xNbS₃ (x = 2-3) in batteries [569].

Charge density waves were studied in *ortho*rhombic [567,570-571] and monoclinic [572] phases of TaS₃. Unstoichiometric Ta₃S_{1,8} has been prepared and its structure studied [573].

A comparative study was done on the layered MX_3 trichalcogenides, TaSe₃, TaS₃ and NbSe₃ [574]. Broken X-X bonds of equilateral-like MX_3 chains and short intra- and interlayer $X\cdots X$ contacts were found to be crucial for the semi-metallic properties of TaSe₃ and for the charge density wave phenomena of NbSe₃ and TaS₃, and qualitative agreement was obtained between experimental observations and band electronic structure calculations [574].

The main representative, $(NbSe_4)_3I$, of an unusual series of one-dimensional compounds, the $(MX_4)_nY$ type phases with M = Nb, Ta; X = S, Se; Y = halogen, - has had its structure thoroughly characterized by single crystal X-ray work [575]. Formally, the compond can be written as $2Nb^4 + Nb^5 + 6Se_2^2 - I^-$. The $(NbSe_4)_3I$ structure consists of essentially one-dimensional $NbSe_4$ -chains separated by iodine ions (Nb-Nb) inter-chain separation is $< 6.7 \, \text{Å}$). The coordination of the eight selenium(-I) ions is *anti*-prismatic as well for the Nb(V) as for the Nb(IV) ions, see Figure 77.

Optical investigations have been performed on the one-dimensional chain compound (TaSe₄)₂I [566-567]. The low temperature IR spectrum and other properties were recorded [576-579]. Magnetic susceptibility and other properties of quasi-one-dimensional (NbSe₄)₂I and (TaSe₄)₂I were reported [580-581]. Thermal properties of the one dimensional (NbSe₄)₃I were studied near its structural phase transition at 247 K [582]. Raman scattering spectra from (NbSe₄)_{10/3}I and (NbSe₄)₃I were recorded as a function of temperature in an effort to understand the nature of the phase transitions [583]. Also, ultrasonic properties [584], far-IR reflectance spectra [585] and the X-ray crystal structure of (NbSe₄)₃I at 30 K [585] were determined, shedding light on the Nb-Nb chain interaction. ⁹³Nb NMR results on (NbSe₄)₃I was discussed in terms of a charge density wave transition at 274 K [586]. At room temperature, Raman spectra of (NbSe₄)₃I and (TaSe₄)₃I single crystals are now known [587]. Other properties of (NbSe₄)₃I and (TaSe₄)₃I are reported in [588-589]. (NbSe₄)₃I can be intercalated with lithium to form Li₇.5(NbSe₄)₃I [420]. By passing through a second order displacive phase transition at 274 K, the Nb-Nb in-chain separations in (NbSe₄)₃I changed from *tri*plets of two-long-one-short to one-long-one-mean-one-short (ranging from 3.31 to 3.06 Å) [575].

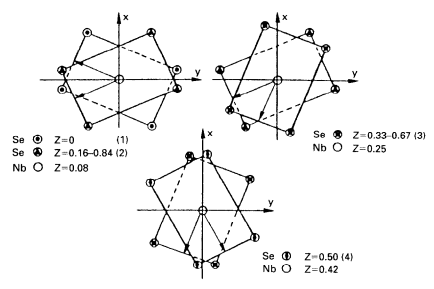


Figure 77. Three rectangular [NbSeg] anti-prisms encountered in the mixed valence (NbSe4)3I structure. Reproduced with permission from [575], P. Gressier, L. Guémas, A. Meerschaut, Mater. Res. Bull. 20 (1985) 539.

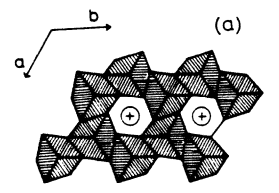


Figure 78. The structure of Nb₃Se₄, projection along the hexagonal *c*-axis. Empty channels between the edge and face-sharing NbSe₆ octahedra can easily take up *e.g.*alkali metals. After [591], T. Ohtani and S Onoue, *J. Solid State Chem.* **59** (1985) 324 and references there.

New $A_xNb_6Se_8$ phases with the Nb₃Te₄ structure have been prepared by molten salt ion exchange (A = Na, K, Rb, Cu, Ag, Zn, Cd, Pb and $0 \le x \le 1$) [590]. The A ions enter topotactically into the empty channels of the three-dimensional network structure of Nb₃Se₄, with little effect on the density, see Figure 78.

The charge-density-wave phase transition at 110 K in the hexagonal one-dimensional conductor Nb₃Te₄ was studied by X-ray diffraction and other methods [592].

The magnetic properties of NbUS₃ and TaUS₃ near the Curie points were investigated [593-594]. Also, the space groups and cell constants of NbUS₃ and TaUS₃ structures were given [594].

Needle crystals of Nb₄Se₁₆Br₂ and Ta₄Se₁₆Br₂ have been prepared from the elements contained in an evacuated sealed pyrex tube placed in a 500-460 °C gradient [560]. The crystal structures have been solved. The structures are based partly on the presence of waved [MSe₄] chains. Along the chains, four niobium or tantalum atoms in a near eight-coordinated *anti*-prismatic environment are found, forming groups of M₄Se₁₆ with short (ca. 3.1 Å) bonds [560]. Also, chains were found in the solved structure of Nb₆Se₂OBr₆ [595].

Crystals of M'Nb₃Se₁₀, with M'=e.g.Fe, have been the subject for numerous studies on charge density wave phenomena. Part of the selenium can be substituted with sulphur, and also some of the niobium can be exchanged, e.g. with iron [596]. The compounds are structurally characterized by the presence of two distinct chains, (I) a trigonal prismatic [NbSe₃] chain similar to that exhibiting the shortest Se-Se pairing in NbSe₃, and (2) a double chain of edge-shared [(Fe,Nb)Se₆] octahedra. Structure determinations, and magnetic and ESR measurements are reported of some of the non-stoichiometric compounds [596].

Nb₂Pd_{0.71}Se₅ single crystal fibers have been obtained by the method of long-time heating of the elements in a closed quartz cell, placed in a temperature gradient and with bromine as transporting agent [347]. The structure was solved based on X-ray diffraction data. It was found to be related to the basic Nb₂Pd₃Se₈ channel type structure, see Figure 37. The Nb₂Pd_{0.71}Se₅ structure is laminar and contains slabs of [Nb₄PdSe₁₀]. Nb atoms are six and seven coordinated by Se in edge-sharing trigonal prisms and *mono*capped trigonal-prismatic sites. Nb₂Pd_{0.71}Se₅ is a metallic conductor along the needle axis, consistent with the non-stoichiometric composition and the non-integral formal oxidation state of the Nb atoms [347].

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