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Soft diphosphine and diarsine complexes of niobium(v) and tantalum(v) fluorides: synthesis, properties, structures and comparisons with the corresponding chlorides†

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The reactions of the soft diphosphines o-C₆H₄(PMe₂)₂, Me₂P(CH₂)₂PMe₂, Et₂P(CH₂)₂PEt₂ or $o-C_6H_4(PPh_2)_2$ with NbF₅ or TaF₅ in anhydrous MeCN solution produce [MF₄(diphosphine)₂][MF₆] (M = Nb or Ta), which have been characterised by microanalysis, IR, ¹H, ¹⁹F(¹H), ³¹P(¹H) and ⁹³Nb NMR spectroscopy. X-ray crystal structures are reported for the isomorphous $[MF_4{O-C_6H_4(PMe_2)_2}][MF_6]$, which confirm the presence of eight-coordinate (distorted dodecahedral) cations. The corresponding reactions using $o-C_6H_4(AsMe_2)_2$ produced [MF₄($o-C_6H_4(AsMe_2)_2$)₂][MF₆] which were similarly characterised, including by the X-ray structure of $[NbF_4(o-C_6H_4(AsMe_2)_2)_2][NbF_6]$. These are very rare examples of arsine complexes of high valent metal fluorides. The chloro complexes [NbCl₄(o-C₆H₄(PMe₂)₂)₂]Cl, [TaCl₄(o-C₆H₄(PMe₂)₂)₂][TaCl₆], [NbCl₄{Me₂P(CH₂)₂PMe₂}₂][NbCl₆] and [MCl₄{o-C₆H₄(AsMe₂)₂}₂][MCl₆] were prepared and their structural and spectroscopic properties compared with the fluoride analogues. Attempts to prepare diphosphine complexes of NbOF3 were unsuccessful, but the NbOCl3 complexes, [{{Me₂P-(CH₂)₂PMe₂<math>NbOCl₃₂ $\{\mu-Me₂P(CH₂)₂PMe₂\}$ $[(o-C_6H_4(PMe_2)_2)NbOCl_3(\mu-O)NbCl_3(CH_3CN)(o-C_6H_4(PMe_2)_2)NbOCl_3(\mu-O)NbCl_3(CH_3CN)(o-C_6H_4(PMe_2)_2)NbOCl_3(\mu-O)NbCl_3(CH_3CN)(o-C_6H_4(PMe_2)_2)NbOCl_3(\mu-O)NbCl_3(CH_3CN)(o-C_6H_4(PMe_2)_2)NbOCl_3(\mu-O)NbCl_3(CH_3CN)(o-C_6H_4(PMe_2)_2)NbOCl_3(\mu-O)NbCl_3(CH_3CN)(o-C_6H_4(PMe_2)_2)NbOCl_3(\mu-O)NbCl_3(CH_3CN)(o-C_6H_4(PMe_2)_2)NbOCl_3(\mu-O)NbCl_3(CH_3CN)(o-C_6H_4(PMe_2)_2)NbOCl_3(\mu-O)NbCl_3(CH_3CN)(o-C_6H_4(PMe_2)_2)NbOCl_3(\mu-O)NbCl_3(H_4(PMe_2)_2)NbOCl_3(\mu-O)NbCl_3(H_4(PMe_2)_2)NbOCl_3(H_4(PMe_2)_2)N$ $C_6H_4(PMe_2)_2$] were obtained. X-Ray structures are also reported for $[NbCl_4\{o-C_6H_4(PMe_2)_2\}_2]Cl$, $[NbCl_4(o-C_6H_4(AsMe_2)_2]_2][NbCl_5(OEt)], [NbCl_4(o-C_6H_4(PMe_2)_2)_2][NbOCl_4(CH_3CN)], [\{\{Me_2P(CH_2)_2PMe_2\}_2\}_2][NbCl_5(OEt)], [NbCl_4(o-C_6H_4(PMe_2)_2)_2][NbOCl_4(CH_3CN)], [\{\{Me_2P(CH_2)_2PMe_2\}_2\}_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2\}_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2\}_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2\}_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2\}_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2\}_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2\}_2PMe_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2\}_2PMe_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2\}_2PMe_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2]_2PMe_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2]_2PMe_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2]_2PMe_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2]_2PMe_2][NbOCl_4(CH_3CN)], [\{Me_2P(CH_2)_2PMe_2]_2PMe_2[NBOC]_2PMe_$ $NbOCl_3/\mu-Me_2P(CH_2)_2PMe_2$ and $[\{o-C_6H_4(PMe_2)_2\}NbOCl_3(\mu-O)NbCl_3(CH_3CN)\{o-C_6H_4(PMe_2)_2\}]$.

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Introduction

Niobium(v) and tantalum(v) fluorides are white crystalline solids which have a tetranuclear structure in the solid state, composed of six-coordinate metal centres at the corners of a square, with single fluoride bridges along each edge. They are very strong Lewis acids and have been explored as constituents of superacid media, although they are weaker than SbF_5 or AsF_5 in this respect. Despite the strong fluoride bridges, they dissolve easily in many O- or N-donor solvents and in halocarbon solvents in the presence of hard donor ligands. A wide

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range of adducts with O-donor Lewis bases have been thoroughly characterised, and in addition to simple adduct formation, C-O, C-H or C-C bond cleavage, rearrangements and polymerisation reactions have been observed.3-5 A smaller range of N-donor adducts are also known.3-5 As would be expected for very hard Lewis acids, complexes with soft Lewis bases are rare. We have reported extremely moisture sensitive thioether complexes of types $[MF_5(SR_2)]$ (M = Nb or Ta), $[MF_4(SR_2)_4][MF_6]$ and $[MF_4\{RS(CH_2)SR\}_2][MF_6]$; the last two types contain eight-coordinate cations.^{6,7} Unstable complexes with selenoethers also form, but these decompose in a few hours with fluorination of the ligand.^{6,7} There appear to be no characterised examples of these pentafluorides complexed to neutral phosphorus or arsenic ligands. The heavier halides MX_5 (X = Cl or Br) are also strong Lewis acids which form many complexes with both hard and soft donor ligands, including P- or As-donor ligands.8 It is notable that hydrolysis of the adducts with the heavier halides, or sometimes O-abstraction from neutral ligands, results in oxide-halide complexes, 5,8,9 whereas similar O/F exchange does not occur with the pentafluorides.^{3,4} Complexes of NbOF₃ were reported very recently, formed by reaction of MF5, hard donor ligands including OPR₃, OSMe₂, 2,2'-bipyridyl or 1,10-phenanthroline,

Dalton Transactions Paper

and the siloxane (Me₃Si)₂O (HMDSO),⁵ although the tantalum analogues remain unknown. Here we report the synthesis of complexes of NbF5 and TaF5 with diphosphine and diarsine ligands, attempts to make NbOF3 adducts, and comparable data on complexes with NbCl₅, TaCl₅ and NbOCl₃.

Results and discussion

[MF₄(diphosphine)₂][MF₆] complexes

Initial attempts to form complexes of alkyldiphosphines (L-L) $(L-L = o-C_6H_4(PMe_2)_2, Me_2P(CH_2)_2PMe_2 \text{ or } Et_2P(CH_2)_2PEt_2)$ used reaction of the appropriate MF5 with the diphosphine in anhydrous CH2Cl2 solution, similar conditions to those used to make thioether complexes, 6,7 and diphosphine complexes of TiF₄ from [TiF₄(MeCN)₂].¹⁰ However, this approach failed; the major products were phosphonium salts of the well known [MF₆] anions, and some other unidentified species. It is likely that the CH₂Cl₂ is activated towards reaction with the phosphines by the strong Lewis acidity of the pentafluorides. In contrast, reaction of MF₅ with the diphosphines in anhydrous MeCN gave colourless solutions, which on concentration or precipitation with anhydrous diethyl ether, deposited [MF₄-(L-L)₂ [MF₆] as white microcrystalline powders, except for [TaF₄{Et₂P(CH₂)₂PEt₂}₂][TaF₆], which was a waxy solid. The complexes [MF₄{o-C₆H₄(PPh₂)₂}₂][MF₆] formed similarly, but PMe₃ did not react cleanly with the MF₅ in anhydrous MeCN, the isolated products containing substantial amounts of coordinated nitrile (IR and ¹H NMR spectroscopic evidence) as well as some PMe3. The [MF5]4 dissolve in MeCN with formation of [MF₄(MeCN)₄][MF₆], ^{4g} which may be intermediates in the reactions; however since [MF₄(RS(CH₂)₂SR)₂][MF₆] form from reaction of MF₅ with RS(CH₂)₂SR in CH₂Cl₂,^{6,7} a donor solvent is not essential to break up the tetrameric pentafluorides. It seems that the soft PMe3 cannot compete successfully with the hard MeCN for the metal centres, at least when the nitrile is present in large excess as solvent (whilst use of chlorocarbon solvents is ruled out by the reactivity described above). Note that PMe3 also failed to cleanly displace MeCN from [TiF₄(MeCN)₂].¹⁰ Several chloride analogues, [NbCl₄{o- $C_6H_4(PMe_2)_2\}_2$ Cl, $[TaCl_4\{o-C_6H_4(PMe_2)_2\}_2][TaCl_6]$ and $[NbCl_4-C_6H_4(PMe_2)_2]_2$ {Me₂P(CH₂)₂PMe₂}₂[[NbCl₆] were made for comparison purposes from reaction of the corresponding MCl₅ with the diphosphine in anhydrous MeCN. The structure of [TaCl₄{Me₂P-(CH₂)₂PMe₂}₂]⁺ has also been reported. The niobium chloro complexes are orange, while those with tantalum are white or pale yellow.

The diphosphine complexes of the metal fluorides are moisture sensitive solids, modestly soluble in anhydrous MeCN, in which they retain their integrity (NMR evidence see below), although they decompose in CH₂Cl₂, and are very readily hydrolysed in solution. Hydrolysis produces a mixture of free diphosphine and the phosphonium hexafluorometallate(v), e.g. $[Me_2P(CH_2)_2PMe_2H][NbF_6]$ from $[NbF_4][Me_2P-1][NbF_6]$ (CH₂)₂PMe₂}₂[NbF₆], identified by their characteristic NMR spectra. The chloro-complexes are less moisture sensitive and

generally poorly soluble in non- or weakly coordinating solvents.

It is convenient to discuss the X-ray structures first and then interpret the spectroscopy and reactions in terms of the complex units present. The two diphosphine complexes $[MF_4\{o-C_6H_4(PMe_2)_2\}_2][MF_6]$ (M = Nb or Ta) are isomorphous (Table 1) and contain eight coordinate distorted dodecahedral cations and the familiar octahedral anions. The cation geometries (Fig. 1 and 2) show essentially identical M-F and M-P distances for the two metal centres, with the former slightly longer (by ~ 0.07 Å) than those found in the [MF₆]⁻ anions. In part this can be attributed to the increase in coordination number, but it is notable that the d(M-F) in the cations are longer (by ~0.02 Å) than those found in the eightcoordinate cations in [MF₄{RS(CH₂)₂SR}₂]⁺ or [MF₄{MeO-(CH₂)₂OMe₂]⁺, ^{4b,6,7} possibly indicating some steric crowding by the phosphorus centres which carry three substituents, as against two in the Group 16 donor ligands. Comparison of the cation geometry in [NbCl₄{o-C₆H₄(PMe₂)₂}₂]Cl (Fig. 3) with those in the fluorides shows d(Nb-P) has increased by $\sim 0.04 \text{ Å}$, which could be due to steric crowding, but may also reflect weaker Lewis acidity of the tetrachloroniobium(v) centre compared to the fluoride analogue.

The IR spectra of the [MF₄(L-L)₂][MF₆] show the presence of the L-L, the absence of phosphine oxide groups, and strong overlapping features in the range 620-550 cm⁻¹ assigned as terminal M-F stretching vibrations.⁵⁻⁷ The ¹H NMR spectra (Experimental section) show the expected resonances for the neutral ligand, shifted to high frequency on coordination. In the diphosphine complexes ²J_{PH} couplings were usually not clearly resolved. Multinuclear NMR spectra (19F, 31P) are much more informative. The ¹⁹F{¹H} NMR spectra in MeCN solution at ambient temperatures show the characteristic 10 line multiplet at $\delta = +103$ ppm for $[NbF_6]^-$ and a singlet at $\delta = +38$ ppm for [TaF₆]^{-.5,6} The ¹⁹F{¹H} resonances of the diphosphine containing cations appear as broad lines to low frequency of the resonances for the corresponding anions (complexes with N-, S- or O-donor ligands usually have resonances to high frequency of the corresponding $[MF_6]^-$ anion^{4–7}). Under higher resolution, binomial quintet couplings are apparent on the cation resonances, which are assigned as ${}^2J_{\rm PF}\sim 40$ --60 Hz (Table 2) due to coupling with the four equivalent phosphorus centres (Fig. 4). In most cases the couplings are clearly resolved, although in $[NbF_4\{o-C_6H_4(PMe_2)_2\}_2][NbF_6]$ and $[NbF_4\{o-C_6H_4(PPh_2)_2\}_2][NbF_6]$ they appear as shoulders on a single broad resonance. Apart from small temperature drifts, the ¹⁹F{¹H} change little on cooling the solutions to 223 K, showing exchange processes are slow even at room temperature. This contrasts with the thioether complexes, 6,7 which showed only a single very broad resonance at room temperature due to rapid dissociative ligand exchange, and which exhibited separate resonances for cation and anion only at low temperatures. The ³¹P{¹H} NMR spectra of the diphosphine complexes each show a single resonance, with very large high frequency coordination shifts (Table 2). In some cases under high resolution these resonances show binomial quintet

Table 1 X-ray crystallographic data^a

In Garage			
Compound	$[\mathrm{NbF}_4[\mathrm{o\text{-}C_6H_4}(\mathrm{PMe_2})_2\}_2][\mathrm{NbF_6}]$	${ m [TaF_4[o-C_6H_4(PMe_2)_2\}_2]}$ ${ m [TaF_6]}$	$[\mathrm{NbF}_4\{o\text{-}\mathrm{C}_6\mathrm{H}_4(\mathrm{AsMe}_2)_2\}_2][\mathrm{NbF}_6]$
Formula M Crystal system Space group (no.) a/k b/k c/k a/c b/k a/c a/c b/k a/c a/c b/k a/c a	$C_{20}H_{32}F_{10}Nb_{2}P_{4}$ 772.16 Triclinic $P\bar{1}$ (no. 2) 12.246(4) 12.325(4) 12.458(4) 11.026(2) 110.266(2) 101.266(2) 1467.5(6) 2 1.072 768 12.736 0.0520 5703 333,0 0.0639, 0.1839 0.0694, 0.1862	$C_{20}H_{32}F_{10}P_4Ta_2$ 948.24 Triclinic $P\bar{1}$ (no. 2) $12.253(2)$ $12.348(2)$ $12.348(2)$ $12.481(2)$ $114.629(2)$ $110.486(3)$ $101.38(3)$ $1471.6(4)$ 2 7.720 896 $14 828$ 0.0457 5150 $333,0$ $0.0577,0.1840$ $0.06257,0.1857$	$C_{20}H_{32}AS_4F_{10}Nb_2$ 947.96 Orthorhombic $Fddd$ (no. 70) $12.989(4)$ $21.329(6)$ $22.051(6)$ 90 90 $6109(3)$ 8 5.121 3.648 5.121 3.648 5.121 3.648 5.121 3.648 5.121 3.648 5.121 6.0273 6.00573 6.00573 6.00573 6.00573
Compound	$[\{\{Me_2P(CH_2)_2PMe_2\}NbOCI_3\}_2-\mu-\{Me_2P(CH_2)_2PMe_2\}]$	$[\mathrm{NbCl}_4[o\text{-}\mathrm{C}_6\mathrm{H}_4(\mathrm{PMe}_2)_2]_2]\mathrm{Cl}\text{-}\mathrm{CH}_3\mathrm{CN}$	$[\{o\text{-}C_6H_4(PMe_2)_2\}NbOCl_3\text{-}\mu\text{-}O\text{-}NbCl_3(CH_3CN)\{o\text{-}C_6H_4(PMe_2)_2\}]$
Formula M Crystal system Space group (no.) a/ \mathring{A} c/\mathring{A} c/\mathring{A} d/\mathring{A}	C ₁₈ H ₄₈ Cl ₆ Nb ₂ O ₂ P ₆ 880.90 Monoclinic P ₂₁ /c (no. 14) 9.096(5) 10.987(6) 18.064(11) 90 90.158(15) 90 1805.3(18) 2 1.362 825 825 825 0.0550 344 160, 0 0.0577, 0.1332 0.0699, 0.1442	C ₂₂ H ₃₅ Cl ₅ NNbP ₄ 707.55 Monoclinic C2/m (no. 12) 21.764(6) 10.178(2) 14.770(4) 90 109.431(9) 90 3085.6(13) 4 1.044 1.044 1.044 1.044 1.044 1.044 1.044 1.043 0.0280 3.191 180, 3 0.0327, 0.0650 0.0362, 0.0663	$C_{22}H_{35}Cl_6NNb_2O_2P_4$ 867.91 Monoclinic $P2_1/c$ (no. 14) 11.845(8) 33.518(19) 8.596(6) 90 92.35(3) 9 3410(4) 4 1.352 1736 1736 1736 1756 1756 17515

Paper

Fig. 1 The cation of [NbF₄{o-C₆H₄(PMe₂)₂}₂][NbF₆], ellipsoids are drawn at the 40% level. Selected bond lengths (Å) and angles (°): Nb1–F3 = 1.952(4), Nb1–F2 = 1.952(5), Nb1–F1 = 1.955(4), Nb1–F4 = 1.960(4), Nb1–P1 = 2.637(2), Nb1–P4 = 2.642(2), Nb1–P2 = 2.651(2), Nb1–P3 = 2.654(2), P1–Nb1–P2 = 71.35(7), P4–Nb1–P3 = 71.38(7), F3–Nb1–F2 = 95.29(19), F3–Nb1–F1 = 92.86(19), F2–Nb1–F4 = 92.78(19), F1–Nb1–F4 = 95.35(19).

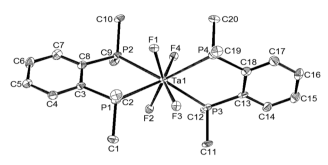


Fig. 2 The cation of $[TaF_4\{o-C_6H_4(PMe_2)_2\}_2]TaF_6]$, ellipsoids are drawn at the 40% level. Selected bond lengths (Å) and angles (°): Ta1-F2=1.944(9), Ta1-F3=1.950(10), Ta1-F1=1.955(9), Ta1-F4=1.970(9), Ta1-P1=2.645(4), Ta1-P4=2.657(4), Ta1-P3=2.658(4), Ta1-P2=2.664(4), P1-Ta1-P2=71.61(13), P4-Ta1-P3=71.53(13), F2-Ta1-F3=95.5(4), F3-Ta1-F1=93.0(4), F2-Ta1-F4=92.9(4), F1-Ta1-F4=96.0(4).

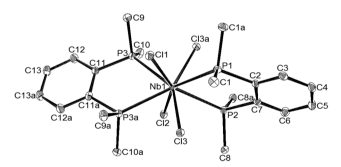


Fig. 3 The cation of $[NbCl_4\{o-C_6H_4(PMe_2)_2\}_2]Cl$, ellipsoids are drawn at the 40% level. Selected bond lengths (Å) and angles (°): Nb1-Cl1=2.4309(9), Nb1-Cl3=2.4341(6), Nb1-Cl2=2.4424(10), Nb1-P3=2.6833(7), Nb1-P1=2.6843(10), Nb1-P2=2.6869(10), P3-Nb1-P3=2.32(3), P1-Nb1-P2=73.189(3), Cl1-Nb1-Cl3=94.99(2), Cl3-Nb1-Cl2=95.029(14). Symmetry operator a=(x,-y,z).

patterns due to ${}^2J_{PF}$, although these were poorly resolved in the spectra of several of the niobium cations.

The 93 Nb NMR spectra (93 Nb: 100% abundance, I = 9/2, $\Xi = 24.44$ MHz, $Q = -0.2 \times 10^{-28}$ m², $D_c = 2740$) of [NbF₄(L-L)₂]-[NbF₆], show the characteristic binomial septet at $\delta \sim -1550$ ppm

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Complex	$\delta^{(^{19}\mathrm{F}\{^{1}\mathrm{H}\})^a/\mathrm{ppm}}$	$\delta(^{31}\mathrm{P}\{^1\mathrm{H}\})^a/\mathrm{ppm}$	$\Delta P (\delta(\text{complex}) - \delta(\text{ligand}))$	Diffuse reflectance UV/Vis data ^c /cm ⁻¹
[NbF ₄ {o-C ₆ H ₄ (PMc ₂) ₂ } ₂][NbF ₆] [NbF ₄ {o-C ₆ H ₄ (AsMc ₂) ₂ } ₂][NbF ₆] [NbF ₄ {Mc ₂ P(CH ₂) ₂ PMc ₂ } ₂][NbF ₆] [NbF ₄ {Mc ₂ P(CH ₂) ₂ PBt ₂ } ₂][NbF ₆] [NbF ₄ {o-C ₆ H ₄ (PPh ₂) ₂ } ₂][NbF ₆] [TaF ₄ {o-C ₆ H ₄ (PMc ₂) ₂ } ₂][TaF ₆] [TaF ₄ {CC ₆ H ₄ (ASMc ₂) ₂ } ₂][TaF ₆] [TaF ₄ {Mc ₂ P(CH ₂) ₂ PMc ₂ } ₂][TaF ₆] [TaF ₄ {CC ₆ H ₄ (PPh ₂) ₂ } ₂][TaF ₆] [TaF ₄ {CC ₆ H ₄ (PPh ₂) ₂ } ₂][TaF ₆] [NbC ₄ {O-C ₆ H ₄ (PPh ₂) ₂ } ₂][TaF ₆] [NbC ₄ {O-C ₆ H ₄ (PMc ₂) ₂] ₂][TaG ₆] [TaCl ₄ {o-C ₆ H ₄ (PMc ₂) ₂] ₂][TaCl ₆]	102.3 (10 lines, ${}^{1}{}_{NbF} = 335 \mathrm{Hz}$, [6F]), -7.8 (quintet, ${}^{2}{}_{PF} = 47 \mathrm{Hz}$, [4F]) 103.6 (10 lines ${}^{1}{}_{NbF} = 335 \mathrm{Hz}$, [6F]), $+27.1$ (8, [4F]) ^b 102.0 (10 lines ${}^{1}{}_{NbF} = 335 \mathrm{Hz}$, [6F]), -10.9 (quintet ${}^{2}{}_{PF} = 50 \mathrm{Hz}$, [4F]) 102.4 (10 lines ${}^{1}{}_{NbF} = 335 \mathrm{Hz}$, [6F]), $+3.3$ (quintet, ${}^{2}{}_{PF} = 45 \mathrm{Hz}$, [4F]) 102.8 (10 lines, ${}^{1}{}_{NbF} = 335 \mathrm{Hz}$, [6F]), $54.8 (\mathrm{br} \mathrm{m}$, [4F]) 37.4 (8, [6F]), -39.8 (quintet ${}^{2}{}_{PF} = 60 \mathrm{Hz}$, [4F]) 39.5 (8, [6F]), $-28.0 (\mathrm{s}, [4F])^b$ 38.3 (6, [6F]), $-25.7 (\mathrm{quintet}, {}^{2}{}_{PF} = 55 \mathrm{Hz}$, [4F]) 38.6 (8, [6F]), $16.8 (\mathrm{quintet}, {}^{2}{}_{PF} = 57 \mathrm{Hz}$, [4F]) $-25.7 (\mathrm{quintet}, {}^{2}{}_{PF} = 57 \mathrm{Hz}$, [4F]) $-25.7 (\mathrm{quintet}, {}^{2}{}_{PF} = 57 \mathrm{Hz}$, [4F])	38.9 (br, m)	94 84 69 53 90 84 73 73 52 103 113	28 330, 32 900 23 400(sh), 28 000 30 300(sh), 33 550 28 000, 32 900 27 300, 33 300 33 000. 23 400(sh), 27 000 33 800 30 300(sh), 33 000 25 980, 32 330 27 550, 31 250

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^a Recorded in MeCN-CD₃CN solution 295 K unless indicated otherwise. ^b In CH₂Cl₂-CD₂Cl₂ solution 295 K. ^c Ligand→metal charge transfer bands.

Dalton Transactions Paper

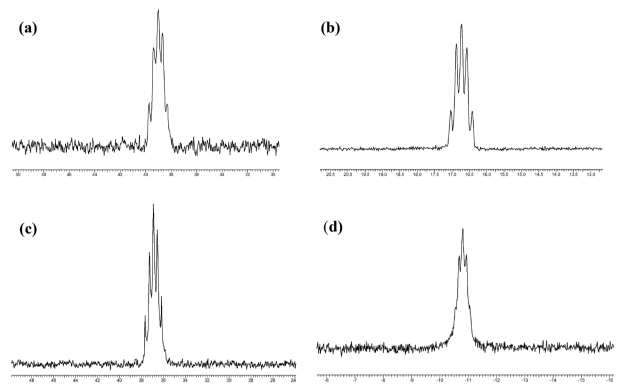


Fig. 4 NMR spectra of the cations: (a) $^{31}P\{^{1}H\}$ of $[TaF_{4}\{o-C_{6}H_{4}(PMe_{2})_{2}\}_{2}]^{+}$; (b) $^{19}F\{^{1}H\}$ of $[TaF_{4}\{o-C_{6}H_{4}(PMe_{2})_{2}\}_{2}]^{+}$; (c) $^{31}P\{^{1}H\}$ of $[NbF_{4}\{Me_{2}P-(CH_{2})_{2}PMe_{2}\}_{2}]^{+}$; (d) $^{19}F\{^{1}H\}$ of $[NbF_{4}\{Me_{2}P-(CH_{2})_{2}PMe_{2}\}_{2}]^{+}$; (d) $^{19}F\{^{1}H\}$ of $[NbF_{4}\{Me_{2}P-(CH_{2})_{2}PMe_{2}\}_{2}]^{+}$; (e) $^{19}F\{^{1}H\}$ of $[NbF_{4}\{Me_{2}P-(CH_{2})_{2}PMe_{2}\}_{2}]^{+}$; (e) $^{19}F\{^{1}H\}$ of $[NbF_{4}\{Me_{2}P-(CH_{2})_{2}PMe_{2}\}_{2}]^{+}$; (f) $^{19}F\{^{1}H\}$ of $[NbF_{4}\{Me_{2}P-(CH_{2})_{2}PMe_{2}\}_{2}]^{+}$; (g) $^{19}F\{^{1}H\}$ of $[NbF_{4}\{Me_{2}P-(CH_{2})_{2}PMe_{2}\}_{2}]^{+}$; (h) $^{19}F\{^{1}H\}$ of $[NbF_{4}\{Me_{2}P-(CH_{2})_{2}PMe_{2}]^{+}$; (h) $^{19}F\{^{1}H\}$ of $[NbF_{4}\{Me_{2}P-(CH_{2})_{2}PMe_{2}]^{+}$

for the anion,⁵ but for L-L = o-C₆H₄(PMe₂)₂ or Me₂P- $(CH_2)_2$ PMe₂, very broad features at $\delta \sim -1100$ ppm $(W_{1/2} \sim 5000$ Hz) were observed in the room temperature spectra, which are tentatively assigned to the dodecahedral cations; these resonances were lost on cooling the solutions. The complexes are very easily hydrolysed in solution, and trace water results first in the loss of the coupling patterns on the cation resonances, and then complete loss of the cation resonance, although the resonances of the water stable [MF₆] remain. The diffuse reflectance UV/Vis spectra of the [MF₄(diphosphine)₂][MF₆] show several broad features in the range 27 000-33 000 cm⁻¹, which for these d^0 complexes can be assigned as $P(\sigma) \rightarrow M(d)$ charge transfer transitions since the $F(\pi) \rightarrow M(d)$ charge transfer bands are expected to occur in the far-UV.12 For those phosphines containing aromatic groups, there are also $\pi \rightarrow \pi^*$ transitions in the near-UV region. Comparisons with the corresponding [MCl₄(diphosphine)₂][MCl₆] (Table 2) show that the $P(\sigma) \rightarrow M(d)$ charge transfer bands occur at higher energy in the fluorides, an effect observed in other systems, 7,10 and expected due to the strong M-F bonding which raises the energy of the metal d-orbitals. The $Cl(\pi) \rightarrow M(d)$ charge transfer bands are observed in the near ultraviolet region.¹³ From the data reported (Experimental section) it seems the $Cl(\pi) \rightarrow M(d)$ transitions in the dodecahedral cations occur at rather lower energy than in the octahedral anions.‡

$[MF_4{o-C_6H_4(AsMe_2)_2}_2][MF_6]$

The reaction of o-C₆H₄(AsMe₂)₂ with both MF₅ reagents in anhydrous MeCN produced [MF₄{o-C₆H₄(AsMe₂)₂}₂][MF₆] as cream powders. These are very rare examples of diarsine complexes with high valent fluorides.3 TiF4 does not complex with o-C₆H₄(AsMe₂)₂, ¹⁰ and the complexes with SnF₄ or GeF₄ were too unstable to isolate. 14,15 In contrast to the stability of the diphosphine complexes, the diarsines in [MF₄{o- $C_6H_4(AsMe_2)_2$ [MF₆] are partially displaced on dissolution of the complexes in MeCN (NMR evidence), but they are stable for some hours in anhydrous CH2Cl2 solution, although slow reaction with the solvent occurs over several days. The successful isolation of the diarsine complexes from MeCN results from their deposition as the least soluble species from an exchanging mixture of complex, diarsine and nitrile adducts of MF₅ in solution, whilst their relative stability to CH₂Cl₂, which contrasts with that of the phosphines, is due to the lower nucleophilicity of the arsenic centres. The [NbF₄{o- $C_6H_4(AsMe_2)_2$ [NbF₆] is not isomorphous with the diphosphine analogue, but shows a similar eight-coordinate cation and six-coordinate anion geometry (Fig. 5).

The d(Nb-F) in the diarsine complex is slightly longer than those in the diphosphine analogue by ~ 0.04 Å. The complexes of this diarsine with NbCl₅ and TaCl₅ of type [MCl₄{o-C₆H₄(AsMe₂)₂}₂][MCl₆]§, were reported many years ago, ¹⁶ and

[‡] In D_{2d} symmetry the metal d-orbitals split $a_1 + b_1 + b_2 + e$, but a more detailed assignment is not possible on the limited data available.

[§] Originally formulated as seven-coordinate monomers.

Paper

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Fig. 5 View of the cation of $[NbF_4{o-C_6H_4(AsMe_2)_2}_2][NbF_6]$. Ellipsoids are drawn at the 40% level. Selected bond lengths (Å) and angles (°): Nb1-F1 = 1.991(4), Nb1-As1 = 2.7518(8), As1-Nb1-As1c = 71.17(3), F1-Nb1-F1a = 95.6(2), F1-Nb1-F1c 148.3(2), F1-Nb1-F1b = 92.9(2). Nb1lies at the intersection of 3 orthogonal 2-fold rotation axes. Symmetry operators a = (x, 5/4 - y, 1/4 - z), b = (1/4 - x, y, 1/4 - z), c = (1/4 - x, y, 1/4 - z)5/4 - y, z).

the X-ray structures are available for [TaCl₄{o-C₆H₄(AsMe₂)₂}₂]- $[TaCl_5(OEt)]^{17}$ and for the $[NbCl_4\{o-C_6H_4(AsMe_2)_2\}_2]^+$ various anions - [NbOCl₄]⁻, [NbO₂Cl₃]²⁻ (ref. 17) and $[NbCl_5(OEt)]^-$ (see ESI†). The d(Nb-As) distances in these various salts are not significantly different to those in [NbF₄{o- $C_6H_4(AsMe_2)_2$ [NbF₆]. The ¹⁹F(¹H) NMR resonances of the cations in [MF₄{o-C₆H₄(AsMe₂)₂}₂][MF₆] are broad singlets to high frequency of the diphosphine analogues (Table 2). In the UV/Vis spectrum the $As(\sigma)\rightarrow M(d)$ in $[MF_4(diarsine)_2][MF_6]$ occur at lower energy (~23 000-28 000 cm⁻¹) than for the corresponding transitions in the diphosphines, as expected given the lower electronegativity of As. 18

NbOF₃ and NbOCl₃ complexes

We recently reported that reaction of NbF5 with a range of N-(2,2'-bipy, 1,10-phen, tmeda) or O-donor (R₃PO, Ph₂P(O)CH₂P-(O)PPh2, dmso) ligands in MeCN-CH2Cl2 solution, followed by addition of the siloxane, HMDSO, produced white [NbOF₃L₂] $(L = R_3PO, dmso)$ or $[NbOF_3(L'-L')]$ (L'-L' = 2,2'-bipy, 1,10-bipy)phen, tmeda or Ph₂P(O)CH₂P(O)PPh₂), although weaker donor ether, nitrile or thioether complexes did not form.⁵ In the present work attempts to use the same approach failed to yield NbOF₃-diphosphine adducts, with $[NbF_4(L-L)_2][NbF_6]$ (L-L = $Me_2P(CH_2)_2PMe_2$ or $o-C_6H_4(PMe_2)_2$) solutions in MeCN being unchanged after 24 h. from addition of HMDSO, as shown by ¹⁹F and ³¹P NMR spectroscopy. Similarly, MeCN solutions of [NbOF₃L₂] (L = dmso or Ph₃PO) did not react with o-C₆H₄(PMe₂)₂ or Me₂P(CH₂)₂PMe₂. We have reported elsewhere 19 that alkyl diphosphines fail to displace the hard O-donor from $[ZrF_4L_2]$ (L = dmso or dmf), and it is plausible that in the NbOF3 systems, the hard Nb centre prefers the O-donor to the softer phosphorus. However, the failure of the siloxane to perform O/F exchange was unexpected and the reason for this is not clear.

Tertiary phosphine complexes of NbOCl3 include both sixcoordinate [NbOCl₃(PR₃)₂] (R = Me, Ph; R₃ = MePh₂) and seven-coordinate [NbOCl₃(PMe₃)₃].^{20,21} In the present work, reaction of NbCl₅ with HMDSO in MeCN, which forms [NbOCl₃(MeCN)₂] in situ, followed by addition of one mol. equivalent of Me₂P(CH₂)₂PMe₂, afforded white [(NbOCl₃)₂-

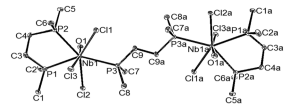


Fig. 6 View of the dimer [{{Me₂P(CH₂)₂PMe₂}NbOCl₃}₂{µ-Me₂P-(CH₂)₂PMe₂}]. Ellipsoids are drawn at the 40% probability level. Selected bond lengths (Å) and angles (°): Nb1-O1 = 1.724(4), Nb1-Cl1 = 2.506(2), Nb1-Cl2 = 2.517(2), Nb1-Cl3 = 2.605(2), Nb1-P1 = 2.6716(19), Nb1-P2= 2.677(2), Nb1-P3 = 2.760(2), O1-Nb1-Cl1 = 97.14(15), O1-Nb1-Cl2 = 94.74(15), Cl1-Nb1-Cl2 = 145.83(6), O1-Nb1-Cl3 = 167.75(15), Cl1-Nb1-Cl3 = 86.36(6), Cl2-Nb1-Cl3 = 88.74(6), O1-Nb1-P1 = 91.43(15), O1-Nb1-P2 = 85.03(15), O1-Nb1-P3 = 94.84(15), P1-Nb1-P2 = 94.84(15)71.04(6). Symmetry operator a = (-x, -y, -z).

{Me₂P(CH₂)₂PMe₂}₃]. The structure determined from colourless crystals grown from CH2Cl2 solution, showed this to be the symmetric dimer [{{Me₂P(CH₂)₂PMe₂}NbOCl₃}₂{μ-Me₂P-(CH₂)₂PMe₂}] (Fig. 6), containing seven-coordinate niobium.

The bond lengths are similar to those in [NbOCl₃(PMe₃)₃]²⁰ but the geometry is quite different, due to the presence of the five-membered chelate ring, with P1-Nb1-P2 = 71.04(6)°, whereas in [NbOCl₃(PMe₃)₃] the P-Nb-P angles are all >112°. The $\nu(Nb=O)$ of 939 cm⁻¹ in [{{Me₂P(CH₂)₂PMe₂}NbOCl₃}₂-{μ-Me₂P(CH₂)₂PMe₂}] contrasts with that of 882 cm⁻¹ reported for [NbOCl₃(PMe₃)₃].^{20,21} The dimer structure is retained in solution, shown most clearly by the ¹H and ³¹P{¹H} NMR spectra, which distinguish the bridging and chelating diphosphines. The phosphorus resonance [2P] of the bridging ligand has a coordination shift (Δ) of +47, whereas the more intense resonance [4P] has $\Delta = +80$, indicative of a five-membered chelate ring.22

The reaction of NbCl₅ and HMDSO in MeCN followed by addition of o-C₆H₄(PMe₂)₂, gave a white precipitate identified as $[{o-C_6H_4(PMe_2)_2}\NbOCl_3(\mu-O)\NbCl_3(CH_3CN){o-C_6H_4(PMe_2)_2}],$ and red-orange [NbCl₄{o-C₆H₄(PMe₂)₂}₂][NbOCl₄(CH₃CN)] was isolated from the filtrate. The structure of the latter is described in ESI.†

The former contains seven-coordinate niobium centres, but in contrast to [(NbOCl₃)₂{Me₂P(CH₂)₂PMe₂}₃], the two niobium centres have different ligand donor sets (P2O2Cl3 and P2NOCl3) and are linked by a near-linear (177°) oxido-bridge (Fig. 7). It may be that the rigid o-C₆H₄(PMe₂)₂, which is pre-organised for chelation, is disfavoured as a bridge in this case and the oxido-bridge is formed instead. The CSD contains only a single example of o-C₆H₄(PMe₂)₂ coordinated as a bridging ligand, in $[(Cp*IrC1_2)_2\{\mu-o-C_6H_4(PMe_2)_2\}].^{23}$

Conclusions

A new series of complexes of niobium(v) and tantalum(v) fluoride with soft, neutral phosphorus or arsenic donor ligands has been prepared and characterised. All are of type [MF₄(L-L)₂]-[MF₆] containing eight-coordinate cations, which seem to be **Dalton Transactions**

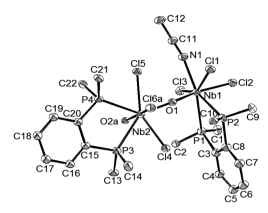


Fig. 7 The structure of $[{o-C_6H_4(PMe_2)_2}]NbOCl_3(\mu-O)NbCl_3(CH_3CN){o-Particles}$ C₆H₄(PMe₂)₂)], with ellipsoids drawn at the 40% probability level. There is disorder around Nb2, involving O2a and Cl6a, and only the major component is shown. Selected bond lengths (Å) and angles (°): Cl1-Nb1 = 2.451(4), Cl2-Nb1 = 2.507(4), Cl3-Nb1 = 2.497(4), Cl4-Nb2 = 2.489(4), Cl5-Nb2 = 2.478(4), N1-Nb1 = 2.268(13), Nb1-O1 = 1.757(10), Nb1-P2 = 2.658(4), Nb1-P1 = 2.668(5), Nb2-O2A = 1.843(13), Nb2-O1= 2.197(10), Nb2-Cl6A = 2.479(5), Nb2-P4 = 2.652(5), Nb2-P3 = 2.197(10)2.671(4), Nb1-O1-Nb2 = 177.5(6), P2-Nb1-P1 = 71.43(13), P4-Nb2-P3= 71.92(13).

the preferred structural units for many complexes of these two M(v) fluorides.³ Despite the extreme hard/soft Lewis acid/base combinations, they are robust complexes with no evidence for decomposition in the solid state if protected from moisture, and the NMR spectra show no significant dissociation of the ligands in inert solvents at ambient temperatures. These properties contrast with those of complexes containing other soft donors, e.g. the rapid decomposition of the selenoether complexes with fluorination at selenium, and the extensive dissociation of the thioether adducts in solution at ambient temperatures.^{6,7} The properties also contrast with the very unstable phosphine complexes of TiF4, and the absence of any arsine ligand adducts with TiF4, GeF4 or SnF4. 10,14,15 Comparisons with the corresponding complexes of Nb or Ta pentachlorides show considerable similarities, both in composition and properties, which contrasts with the behaviour of Group 4 tetrahalides, where there is little in common between the chemistries of ZrF4 or HfF4 (which do not form complexes with soft donor ligands),19 and the six- or eightcoordinate complexes formed the heavier tetrahalides.²⁴ The coordination chemistry of TiF4 is also very different from those of TiX_4 (X = Cl, Br or I). ^{10,25} The coordination chemistry of VF₄ merits thorough study, and on the basis of the results reported herein, it is possible that even VF₅ may form neutral ligand complexes under appropriate conditions.³

Experimental

Infrared spectra were recorded as Nujol mulls between CsI plates using a Perkin Elmer Spectrum 100 over the range 4000-200 cm⁻¹. ¹H NMR spectra were recorded from CD₂Cl₂ or CD₃CN solutions using a Bruker DPX400 spectrometer. ¹⁹F{¹H}, ³¹P{¹H} and ⁹³Nb NMR spectra were recorded using a Bruker DPX400 spectrometer and are referenced to external CFCl₃, external 85% H₃PO₄, and [Et₄N][NbCl₆] in MeCN respectively. UV/visible spectra were recorded from solid samples diluted with BaSO₄ using the diffuse reflectance attachment of a Perkin-Elmer Lambda 19 spectrometer. Microanalyses on new complexes were undertaken by London Metropolitan University. Preparations used standard Schlenk and glove box techniques under a N2 atmosphere with rigorous exclusion of moisture. Solvents were dried by distillation from CaH₂ (CH₂Cl₂ or CH₃CN) or Na/benzophenone ketyl (diethyl ether). Hexamethyldisiloxane was dried over molecular sieves. Anhydrous NbX_5 and TaX_5 (X = F or Cl), PMe_3 , Me_2P - $(CH_2)_2PMe_2$, $Et_2P(CH_2)_2PEt_2$ and $o-C_6H_4(PPh_2)_2$ were obtained from Aldrich, Apollo or Strem and used as received. o-C₆H₄(PMe₂)₂ and o-C₆H₄(AsMe₂)₂ were made by literature methods.26,27

 $[NbF_4{o-C_6H_4(PMe_2)_2}_2][NbF_6]: NbF_5 (0.18 g, 1.0 mmol) was$ dissolved in anhydrous MeCN (10 mL) and o-C₆H₄(PMe₂)₂ (0.20 g, 1.0 mmol) added. A clear solution formed which deposited a cream precipitate after ~20 min. After stirring for 1 h the precipitate was filtered off, rinsed with MeCN (2 mL) and dried in vacuo. White powder. Yield: 0.30 g, 79%. Anal: required for C₂₀H₃₂F₁₀Nb₂P₄ (772.2): C, 31.1; H, 4.2. Found: C, 31.0; H, 4.0%. ¹H NMR (CD₃CN, 295 K): $\delta = 1.70$ (br s, [12H]), 7.82 (0 [2H]), 7.95 (s, [2H]). 19 F{ 1 H} NMR (CD₃CN, 298 K): δ = 102.3 ([6F], 10 lines, ${}^{1}J_{NbF} = 335 \text{ Hz}$, $[NbF_{6}]^{-}$), -7.8 (quintet, $^{2}J_{PF}$ = 47 Hz, [4F], cation); (243 K) 102.3 ([6F], 10 lines, $^{1}J_{NbF}$ = 335 Hz), -10.8 (br s, [4F]). $^{31}P\{^{1}H\}$ NMR (CD₃CN, 298 K): δ = 38.9 (br m); (233 K) 39.5 (br m). 93 Nb NMR (CD₃CN, 298 K): δ = ~-1110 (vbr, cation), -1553 (septet, [NbF₆]⁻); (233 K): -1554 (septet). IR (Nujol)/cm⁻¹: 611 (vbr), 588 (sh) (NbF). UV/Vis (dr)/ cm⁻¹: 28 330, 32 900.

 $[NbF_4\{o-C_6H_4(AsMe_2)_2\}_2][NbF_6]: NbF_5 (0.18 g, 1.0 mmol)$ dissolved in anhydrous MeCN (15 mL) and o-C₆H₄(AsMe₂)₂ (0.29 g, 1.0 mmol) added. A clear yellow solution formed which deposited a pale yellow-cream precipitate after ~20 min. After stirring for 1 h the solution was concentrated to ~5 mL, the precipitate was filtered off, rinsed with MeCN (2 mL) and dried in vacuo. Yellow powder. Yield: 0.40 g, 85%. Anal: required for C₂₀H₃₂As₄F₁₀Nb₂ (948.0): C, 25.3; H, 5.4. Found: C, 25.4; H, 5.3%. ¹H NMR (CD_2Cl_2 , 293 K): δ = 1.63 (s, [12H]), 7.58 (s, [2H]), 7.67 (s, [2H]); (243 K): 1.63 (s, [12H]), 7.56 (s, [2H]), 7.64 (s, [2H]). $^{19}F\{^{1}H\}$ NMR (CD₂Cl₂, 293 K): δ = 103.6 ([6F], 10 lines ${}^{1}J_{NbF} = 335 \text{ Hz}$, [NbF₆]⁻), 27.1 (s, [4F], cation); (233 K): 103.6 (10 lines), 24.6 (s). 93 Nb NMR (CD₂Cl₂, 293 K): δ = -1549 (septet, [NbF₆]⁻). IR (Nujol)/cm⁻¹: 620 (sh), 610 (sbr), 586 (sh) (NbF). UV/Vis (dr)/cm⁻¹: 23 400 (sh), 28 000, 32 050.

 $[NbF_4\{Me_2P(CH_2)_2PMe_2\}_2][NbF_6]: NbF_5 (0.18 g, 1.0 mmol)$ was dissolved in anhydrous MeCN (15 mL) and Me2P-(CH₂)₂PMe₂ (0.15 g, 1.0 mmol) added. A clear solution formed which deposited a white powder on stirring. After stirring for 1 h the solution was concentrated to ~5 mL, and dry diethyl ether (5 mL) added. The precipitate was filtered off, rinsed with MeCN (2 mL) and dried in vacuo. White powder. Yield: 0.25 g, 76%. Anal: required for $C_{12}H_{32}F_{10}Nb_2P_4$ (676.1): C, **Paper**

21.3; H, 4.8. Found: C, 21.1; H, 4.9%. ¹H NMR (CD₃CN, 293 K): δ = 1.94 (s, [12H]), 2.33 (s, [4H]); (233 K): 1.96 (s, [12H]), 2.33 (s, [4H]). ¹⁹F{¹H} NMR (CD₃CN, 293 K): δ = 102.0 ([6F], 10 lines $^{1}J_{\text{NbF}}$ = 335 Hz, [NbF₆]⁻), -10.9 (quintet $^{2}J_{\text{PF}}$ = 60 Hz, [4F], cation); (233 K): 102.7 ([6F], 10 lines, $^{1}J_{\text{NbF}}$ = 335 Hz,), -13.8 (quintet, [4F]). ³¹P{¹H} NMR (CD₃CN, 293 K): δ = 36.9 (quintet, $^{2}J_{\text{PF}}$ = 60 Hz); (233 K): 38.0 (quintet). ⁹³Nb NMR (CD₃CN, 293 K): δ = -1062 (vbr, s, cation), -1553 (septet, [NbF₆]⁻); (233 K): -1555 (septet). IR (Nujol)/cm⁻¹: 614 (sh), 573 (sbr), 554 (sh) (NbF). UV/Vis (dr)/cm⁻¹: 30 300 (sh), 33 550.

 $[NbF_4[Et_2P(CH_2)_2PEt_2]_2[NbF_6]$: was made similarly. The complex is more soluble in MeCN and was isolated by removing the MeCN in vacuo and stirring the waxy white residue with dry diethyl ether (10 mL) for 2 h, after which the powder was filtered off and dried. White powder. Yield: 78%. Anal: required for C₂₀H₄₈F₁₀Nb₂P₄ (788.3): C, 30.5; H, 6.1. Found: C, 30.4; H, 6.1%. ¹H NMR (CD₃CN, 293 K): δ = 1.13 (br s, [12H]), 1.89 (m, [8H]), 2.15 (m, [4H]); (233 K): 1.07 (m, [12H]), 1.85 (m, [8H]), 2.16 (m, [4H]). $^{19}F\{^{1}H\}$ NMR (CD₃CN, 293 K): δ = 102.4 ([6F], 10 lines, ${}^{1}J_{NbF} = 335 \text{ Hz}$, $[NbF_{6}]^{-}$), +3.3 (quintet, ${}^{2}J_{PF} =$ 45 Hz, [4F], cation); (CD₃CN, 243 K): 101.9 (10 lines), +0.04 (quintet, ${}^{2}J_{PF}$ = 45 Hz, [4F]). ${}^{31}P\{{}^{1}H\}$ NMR (CD₃CN, 293 K): δ = 50.3 (br s); (233 K): 50.0 (br m). 93 Nb NMR (CD₃CN, 293 K): δ = -1553 (septet, [NbF₆]⁻); (233 K): -1555 (septet). IR (Nujol)/ cm⁻¹: 630 (sh), 611 (sbr) (NbF). UV/Vis (dr)/cm⁻¹: 28 000, 32 900.

 $[NbF_4]$ $\{o-C_6H_4(PPh_2)_2\}_2[NbF_6]$: Powdered $o-C_6H_4(PPh_2)_2$ (0.40 g, 0.9 mmol) was suspended in dry MeCN (20 mL) and the mixture stirred vigorously. Powdered NbF₅ (0.16 g, 0.9 mmol) was added and after ~5 min a clear solution was produced. After a further 3 h the solution was concentrated to ~5 mL in vacuo, when a pale cream solid deposited. This was filtered off, rinsed with MeCN (1 mL) and dried in vacuo. Yield: 0.39 g, 70%. Anal: required for C₆₀H₄₈F₁₀Nb₂P₄ (1268.7): C, 56.8; H, 3.8. Found: C, 56.9; H, 3.6%. ¹H NMR (CD₂Cl₂ 293 K): $\delta = 7.65 - 7.23$ (m). ¹⁹F{¹H} NMR (CD₂Cl₂, 293 K): $\delta = 102.8$ (10 lines, ${}^{1}J_{\text{NbF}} = 335 \text{ Hz}$, [6F], [NbF₆]⁻) 54.8 (br m, [4F], cation); $(CD_2Cl_2, 243 \text{ K})$: $\delta = 102.9 \text{ (10 lines. [6F])}, 48.8 \text{ (br s, [4F])}.$ ³¹P{¹H} NMR (CD₂Cl₂, 293 K): δ = 40.3 (vbr); (233 K): 41.4 (quintet, ${}^{2}J_{PF}$ = 43 Hz). ${}^{93}Nb$ NMR (CD₂Cl₂, 293 K): δ = -1549 (septet, [NbF₆]⁻). IR (Nujol)/cm⁻¹: 630 (s br), 602 (s br) (NbF). $UV/Vis (dr)/cm^{-1}$: 27 300, 33 300.

[TaF₄{o-C₆H₄(PMe₂)₂}₂][TaF₆]: TaF₅ (0.28 g, 1.0 mmol) was dissolved in anhydrous MeCN (10 mL) and o-C₆H₄(PMe₂)₂ (0.20 g, 1.0 mmol) added. A clear solution formed which was stirred for 1 h, concentrated to ~5 mL and the precipitate filtered off, rinsed with MeCN (2 mL) and dried *in vacuo*. White powder. Yield: 0.36 g, 75%. Anal: required for C₂₀H₃₂F₁₀P₄Ta₂ (948.3): C, 25.3; H, 3.4. Found: C, 25.2; H, 3.3%. ¹H NMR (CD₃CN, 295 K): δ = 1.70 (br s, [12H]), 7.79 (s, [2H]), 7.92 (s, [2H]); (243 K): 1.68 (s, [12H]), 7.77 (s, [2H]), 7.92 (s, [2H]). ¹⁹F {¹H} NMR (CD₃CN, 298 K): δ = 37.4 ([6F], [TaF₆]⁻), -39.8 (quintet, ${}^2J_{\rm PF}$ = 60 Hz, [4F], cation); (243 K): 38.3 (s, [6F]), -42.3 (m, [4F]). ³¹P{¹H} NMR (CD₃CN, 298 K): δ = 35.4 (quintet, ${}^2J_{\rm PF}$ = 60 Hz); (233 K): 36.1 (br s). IR (Nujol)/cm⁻¹: 621 (sh), 596 (sh), 572 (vs) (TaF). UV/Vis (dr)/cm⁻¹: 33 000.

[TaF₄{*o*-C₆H₄(AsMe₂)₂}₂][TaF₆]: TaF₅ (0.28 g, 1.0 mmol) was dissolved in anhydrous MeCN (10 mL) and *o*-C₆H₄(AsMe₂)₂ (0.29 g, 1.0 mmol) added. A clear yellow solution formed which deposited a small amount of pale cream precipitate after ~20 min. After stirring for 1 h the solution was concentrated to ~5 mL, dry diethyl ether (5 mL) added, and the precipitate filtered off, rinsed with MeCN (2 mL) and dried *in vacuo*. Yellow powder. Yield: 0.40 g, 85%. Anal: required for C₂₀H₃₂As₄F₁₀Ta₂ (1124.0): C, 21.4; H, 2.9. Found: C, 21.3; H, 2.8%. ¹H NMR (CD₂Cl₂, 293 K): δ = 1.63 (s, [12H]), 7.59 (s, [2H]), 7.64 (s, [2H]); (243 K): 1.60 (s, [12H]), 7.55 (s, [2H]), 7.60 (s, [2H]). ¹⁹F₁¹H₁ NMR (CD₂Cl₂, 293 K): δ = 39.5 (s, [6F], [TaF₆]⁻), -28.0 (s, [4F], cation); (243 K): 39.7 (s, [6F]), -29.0 (s, [4F]). IR (Nujol)/cm⁻¹: 616 (sh), 578 (br) (TaF). UV/Vis (dr)/cm⁻¹: 23 400 (sh), 27 000, 30 600.

[TaF₄{Me₂P(CH₂)₂PMe₂}₂][TaF₆]: was made similarly to the niobium analogue. White microcrystalline solid. Yield: 83%. Anal: required for C₁₂H₃₂F₁₀P₄Ta₂ (852.2): C, 16.9; H, 3.8. Found: C, 16.8; H, 3.8%. ¹H NMR (CD₃CN, 293 K): δ = 1.81 (s, [12H]), 2.10 (s, [4H]); (233 K): 1.82 (s, [12H]), 2.10 (s, [4H]). ¹⁹F{¹H} NMR (CD₃CN, 293 K): δ = 38.3 (s, [6F], [TaF₆]⁻), -40.8 (quintet, ²J_{PF} = 60 Hz, cation); (CD₃CN, 243 K): 38.8 (s, [6F]), -43.6 (quintet, [4F]). ³¹P{¹H} NMR (CD₃CN, 293 K): δ = 36.9 (quintet, ²J_{PF} = 60 Hz); (233 K): 37.9 (quintet). IR (Nujol)/cm⁻¹: 613 (sh), 575 (vbr, s) (TaF). UV/Vis (dr)/cm⁻¹: 33 400.

[TaF₄{Et₂P(CH₂)₂PEt₂}₂][TaF₆]: was made similarly to the niobium analogue. White waxy solid. Yield: 65%. ¹H NMR (CD₃CN, 293 K): δ = 1.17 (br s, [12H]), 1.80 (br s, [8H]), 2.15 (s, [4H]); (233 K): 1.08 (s, [12H]), 1.84 (s, [8H]), 2.12 (s [4H]). ¹⁹F {¹H} NMR (CD₃CN, 293 K): δ = 38.8 (s, [6F], [TaF₆]⁻), -25.7 (quintet, ²J_{PF} = 55 Hz, [4F], cation); (CD₃CN, 243 K): 38.9 (s, [6F]), -28.8 (quintet, [4F]). ³¹P{¹H} NMR (CD₃CN, 293 K): δ = 45.0 (quintet, ²J_{PF} = 55 Hz); (233 K): 45.3 (quintet). IR (Nujol)/cm⁻¹: 615 (sh), 587 (vbr, s) (TaF). UV/Vis (dr)/cm⁻¹: 33 800.

[TaF₄{*o*-C₆H₄(PPh₂)₂}₂][TaF₆]: was made similarly to the niobium analogue using *o*-C₆H₄(PPh₂)₂ (0.40 g, 0.9 mmol) and TaF₅ (0.26 g, 0.9 mmol). Yield: 0.48 g, 73%. Anal: required for C₆₀H₄₈F₁₀P₄Ta₂ (1444.8): C, 49.9; H, 3.6. Found: C, 49.8; H, 3.5%. ¹H NMR (CD₂Cl₂, 293 K): δ = 7.77–7.23 (m). ¹⁹F₁H NMR (CD₂Cl₂, 293 K): δ = 38.6 (s, [6F], [TaF₆]⁻), 16.8 (quintet, ²*J*_{PF} = 57 Hz, cation); (243 K): δ = 38.7 (s), 12.3 (br s). ³¹P₁H NMR (CD₂Cl₂, 293 K): δ = 38.6 (br); (233 K): 39.0 (quintet, ²*J*_{PF} = 57 Hz). IR (Nujol)/cm⁻¹ 583 (s, br), 545 (s, br) (TaF). UV/Vis (dr)/cm⁻¹: 30 300, 33 000.

[NbCl₄{o-C₆H₄(AsMe₂)₂}₂][NbCl₆]: Made as described.¹⁶ Orange-red powder. ¹H NMR (CD₂Cl₂, 293 K): δ = 1.92 (s, [12H]), 7.78 (s, [2H]), 7.90 (s, [2H]). ⁹³Nb NMR (CD₂Cl₂, 293 K): δ = +6.2 (s, [NbCl₆]⁻). IR (Nujol)/cm⁻¹: 324 (vs) (NbCl). UV/Vis (dr)/cm⁻¹: 20 350, 29 760.

[TaCl₄{o-C₆H₄(AsMe₂)₂}₂][TaCl₆]: Made as described.¹⁶ Deep yellow powder. ¹H NMR (CD₂Cl₂, 293 K): δ = 2.00 (s, [12H]), 7.80 (s, [2H]), 7.86 (s, [2H]). IR (Nujol)/cm⁻¹ 323 (vs), 303 (s) (TaCl). UV/Vis (dr)/cm⁻¹: 23 530, 31 250.

[NbCl₄{o-C₆H₄(PMe₂)₂}₂]Cl: NbCl₅ (0.135 g, 0.50 mmol) was dissolved in acetonitrile (5 mL) whilst stirring giving a bright yellow-green solution. o-C₆H₄(PMe₂)₂ (0.20 g, 1.0 mmol) was

Dalton Transactions Paper

added slowly to the solution, which resulted in rapid formation of a red-brown precipitate. The reaction was left to stir another 10 min. The solution was filtered, the precipitate was washed with small amount of dichloromethane and dried in vacuo. Yield: 0.280 g, 83%. Red-orange single crystals of [NbCl₄(o-C₆H₄(PMe₂)₂)₂]Cl were grown from a saturated dichloromethane solution cooled in the freezer. Anal: required for C₂₀H₃₂Cl₅NbP₄ (666.5): C, 36.0; H, 4.8. Found: C, 35.9; H, 4.7%. ¹H NMR (CD₂Cl₂, 295 K): $\delta = 1.96$ (s, [12H]), 7.70–7.62 (m, [4H]). ${}^{31}P{}^{1}H$ NMR (CH₂Cl₂-CD₂Cl₂, 298 K): δ = 58.4 (s). IR (Nujol)/cm⁻¹: 320 (s), 293 (s) (NbCl). UV/Vis (dr)/cm⁻¹: 23 000, 34 500.

 $[NbCl_4\{Me_2P(CH_2)_2PMe_2\}_2][NbCl_6]$: NbCl₅ (0.135)0.50 mmol) was dissolved in acetonitrile (5 mL) whilst stirring giving a bright yellow-green solution. Me₂P(CH₂)₂PMe₂ (0.075 g, 0.50 mmol) in 3 mL acetonitrile was added to the solution which resulted in rapid formation of red-brown precipitate. The reaction was left to stir 10 min. The solution was filtered, the precipitate washed with small amount of dichloromethane and dried in vacuo. Yield: 0.130 g, 65%. Anal: required for $C_{12}H_{32}Cl_{10}Nb_2P_4$ (840.6): C, 17.1; H, 3.8. Found: C, 17.3; H, 3.9%. ¹H NMR (CD₂Cl₂, 293 K): δ = 1.73 (br s, [12H]), 2.01 (br, [4H]). ³¹P{¹H} NMR (CH₂Cl₂-CD₂Cl₂, 298 K): $\delta = 56.3(s)$. ⁹³Nb NMR (CH₂Cl₂-CD₂Cl₂, 298 K): $\delta = 6.1(s)$. IR (Nujol)/cm⁻¹: 303 (s, vbr) (NbCl). UV/Vis (dr)/cm⁻¹: 25 980, 32 330.

 $[TaCl_4]$ $[o-C_6H_4]$ $[PMe_2]_2$ $[TaCl_6]$: To a suspension of $TaCl_5$ (0.090 g, 0.25 mmol) in acetonitrile (5 mL) was added o-C₆H₄(PMe₂)₂ (0.050 g, 0.25 mmol) with stirring which resulted in a formation of a white precipitate. The reaction was left to stir for 30 min, and then filtered. The white solid was washed with small amount of dichloromethane and dried. Yield: 0.10 g, 66%. Anal: required for C₂₀H₃₂Cl₁₀P₄Ta₂ (1112.8): C, 21.6; H, 2.9. Found: C, 21.6; H, 3.0%. ¹H NMR (CD₂Cl₂, 295 K): $\delta = 1.71$ (s, [12H]), 7.75 (br, [4H]). ³¹P{¹H} NMR CH₂Cl₂- CD_2Cl_2 , 298 K): $\delta = 44.6$ (br s). IR (Nujol/cm⁻¹): 324 (s), 279 (s) (TaCl). $UV/Vis (dr)/cm^{-1}$: 27 550, 31 250.

 $[NbCl_4\{o-C_6H_4(PMe_2)_2\}_2][NbOCl_4(CH_3CN)]$ and $[\{o-C_6H_4-C_6H_4\}_2][NbOCl_4(CH_3CN)]$ $(PMe_2)_2$ $NbOCl_3(\mu-O)NbCl_3(CH_3CN)$ $\{o-C_6H_4(PMe_2)_2\}$ $\}:$ (0.135 g, 0.50 mmol) was dissolved in acetonitrile (5 mL) giving a bright yellow-green solution. HMDSO (0.10 g, 0.60 mmol) was added. The mixture was left to stir under nitrogen for 30 min during which time the solution turned very pale yellow. o-C₆H₄(PMe₂)₂ (0.10 g, 0.50 mmol) in 3 mL acetonitrile was added slowly to the solution which resulted in formation of white precipitate immediately with a solution colour change to red-orange. The reaction was left to stir for another 5 min and then the white precipitate filtered off. The filtrate was refrigerated for several days to give red-orange crystals of $[NbCl_4\{o-C_6H_4(PMe_2)_2\}_2][NbOCl_4(CH_3CN)]$ Yield: 0.023 g, 10%. Anal: required for C₂₂H₃₅Cl₈NNb₂OP₄ (922.8): C, 28.6; H, 3.8; N, 1.5. Found: C, 28.6; H, 3.8; N, 1.6. ¹H NMR $(CD_2Cl_2, 295 \text{ K}): \delta = 1.98 \text{ (s, [3H], MeCN)}, 2.11 \text{ (s, [24H])},$ 7.73–7.87 (m, [8H], aromatic H). ${}^{31}P{}^{1}H{}$ NMR (CH₂Cl₂-CDCl₃, 298 K): $\delta = 55.2$ (s). IR (Nujol/cm⁻¹): 2305 (vw), 2279 (vw) (MeCN), 947 (s, NbO), 320 (vbr), 289 (s) (NbCl). The white pre-

cipitate was washed with small amount of dichloromethane and dried. Colourless single crystals of [{o-C₆H₄(PMe₂)₂}- $NbOCl_3(\mu-O)NbCl_3(CH_3CN)\{o-C_6H_4(PMe_2)_2\}$] were grown from a saturated dichloromethane solution in the freezer. Yield: 0.045 g, 12%. Anal: required for C₂₂H₃₅Cl₆NNb₂O₃P₄ (884.0): C, 29.9; H, 4.0, N, 1.6. Found: C, 30.5; H, 3.9; N, 1.6. ¹H NMR $(CD_2Cl_2, 295 \text{ K})$: $\delta = 1.58 \text{ (s, [6H], PMe}_2), 1.62 \text{ (s, [12H], PMe}_2),$ 1.77 (s, [6H], PMe₂), 1.98 (s, [3H], MeCN), 7.57–7.68 (m, [8H]). ³¹P{¹H} NMR (CH₂Cl₂-CDCl₃, 298 K): δ = 42.9 (s, [P]), 38.3 (br, [2P]), 36.4 (s, [P]). IR (Nujol/cm⁻¹): 943 (s, NbO), 824(s) (Nb-O-Nb), 355, 304 (s, NbCl).

 $[(\{Me_2P(CH_2)_2PMe_2\}NbOCl_3)_2(\mu-Me_2P(CH_2)_2PMe_2)]: NbCl_5$ (0.135 g, 0.50 mmol) was dissolved in acetonitrile (5 mL) whilst stirring giving a bright vellow-green solution. HMDSO (0.10 g, 0.60 mmol) was added. The mixture was left to stir under nitrogen for 30 min during which time the solution turned very pale yellow. Me₂P(CH₂)₂PMe₂ (0.075 g, 0.50 mmol) in 5 mL of dichloromethane was added to the solution which resulted in formation of white precipitate immediately. The reaction was left to stir another 10 min and then filtered. The precipitate was washed with a small amount of dichloromethane and dried in vacuo. Yield: 0.088 g, 76%. Colourless single crystals of $[\{Me_2P(CH_2)_2PMe_2\}NbOCl_3\}_2(\mu-Me_2P-$ (CH₂)₂PMe₂)] were grown from saturated dichloromethane solution in the freezer. Anal: required for C₁₈H₄₈Cl₆Nb₂O₂P₆ (880.9): C, 24.5; H, 5.5. Found: C, 24.5; H, 5.5%. ¹H NMR $(CD_2Cl_2, 295 \text{ K}): \delta = 1.52 \text{ (s, [12H], PMe}_2), 1.77 \text{ (s, [24H], PMe}_2),$ 2.14-2.38 (m, [12H], CH₂). ³¹P{¹H} NMR CH₂Cl₂-CD₂Cl₂, 298 K): $\delta = 33.3$ (s, [4P]), 0.7 (s, [2P]). IR (Nujol)/cm⁻¹: 939 (m, NbO), 321(w), 288(m) (NbCl).

X-ray experimental

Details of the crystallographic data collection and refinement parameters are given in Table 1. Crystals suitable for single crystal X-ray analysis were obtained as described above. Data collections used a Rigaku AFC12 goniometer equipped with an enhanced sensitivity (HG) Saturn724+ detector mounted at the window of an FR-E+ SuperBright molybdenum ($\lambda = 0.71073 \text{ Å}$) rotating anode generator with VHF Varimax optics (70 µm focus) with the crystal held at 100 K (N2 cryostream). Structure solution and refinements were performed with either SHELX (S/L)97 or SHELX(S/L)2013²⁸ and were straightforward except as detailed below. H atoms bonded to C were placed in calculated positions using the default C-H distance and refined using a riding model. The anion in [NbF₄{o-C₆H₄(AsMe₂)₂}₂]-[NbF₆] was disordered which was satisfactorily modeled with a number of partially occupied F positions and thermal restraints. The $[{o-C_6H_4(PMe_2)_2}\NbOCl_3-\mu$ parameter O-NbCl₃(CH₃CN){o-C₆H₄(PMe₂)₂}] crystals were weakly diffracting with poorly defined reflection profiles and it was necessary to employ global restraints to the atomic displacement parameters. The apical chlorine and oxygen atoms coordinated to Nb2 are modeled as disordered (ca. 80/20) over the 2 positions using thermal parameter constraints and distance restraints (Nb-Cl = 2.47, Nb-O 1.8).

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Paper

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