

An Organotransition-Metal Complex with Pentagonal-Pyramidal Structure**

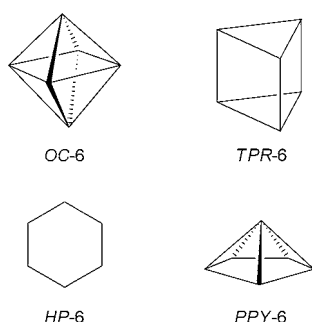
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Dedicated to Professor Pablo J. Alonso on the occasion of his 60th birthday

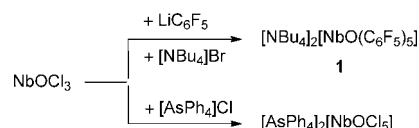
It was Alfred Werner in the late 19th century,^[1] who first suggested that six ligands should adopt an octahedral (*OC*-6) geometry around a given central atom. This geometrical arrangement minimizes repulsive effects between six ligands (LCP model) as well as between six valence-electron pairs (VSEPR model).^[2] It is no wonder, then, that the many myriads of six-coordinate compounds prepared ever since, almost invariably show an *OC*-6 geometry. There are, however, some examples of alternative geometries currently known (Scheme 1).^[3] Thus, a number of compounds containing dithiolato ligands^[4] $[M(S_2C_2R_2)_3]^{q-}$ as well as a few homoleptic organo-transition-metal complexes^[5] $[MR_6]^{q-}$ that show trigonal-prismatic (*TPR*-6) structures.^[6] Hexagonal-planar (*HP*-6) structures have been found in $[Ni(EtBu)_6]$ compounds ($E = P, As$),^[7] where the nickel center is hosted by a cyclic hexadentate framework. Pentagonal-pyramidal (*PPY*-6) structures have been found mainly in post-transition-element compounds^[8–10] including the fascinating species

$[XeOF_5]^-$.^[8a,b,10] The *PPY*-6 geometry is less frequent in transition-metal compounds, where it has been found mainly associated to either pentadentate ligands^[11] or to bidentate ligands of the peroxo type^[12] as in $CrO(O_2)_2py$ (py = pyridine).^[13] Here we report on what we consider to be the first organo-transition-metal complex with *PPY*-6 structure.^[14]

The oxohalide compound $NbOCl_3$ reacts with LiC_6F_5 in Et_2O , in the presence of NBu_4Br , to give organoniobium(v) derivative $[NBu_4]_2[NbO(C_6F_5)_5]$ (**1**) in reasonable yield (Scheme 2 and see the Supporting Information for further details). Compound **1** was isolated as a white, air-sensitive solid, and is best kept at $-30^\circ C$ to avoid thermal degradation. The IR spectrum of **1** shows a sharp signal at 986 cm^{-1} , which was assigned to the $\nu(NbO)$ mode, as well as strong bands at $1495, 953\text{ (C-F)}$, and 746 cm^{-1} (X -sensitive mode), which are characteristic of metal-bound C_6F_5 groups.^[15]



Scheme 1. Idealized geometries in six-coordinate compounds.^[3]



Scheme 2. Synthetic procedures leading to $[ER_4]^+$ salts of the six-coordinate anions $[NbOX_5]^{2-}$ ($X = Cl, C_6F_5$) starting from the same precursor species.

The structure of **1**-*n*-hexane was established by single-crystal X-ray diffraction methods. The anion $[NbO(C_6F_5)_5]^{2-}$ (Figure 1)^[16] exhibits a *PPY*-6 structure^[17] according to the low continuous-shape measure (CShM) value for that geometry: $S(PPY-6) = 0.76$.^[18] The oxo ligand is located in the apical position and is tightly bound to the niobium center. The observed Nb–O bond length ($168.5(4)\text{ pm}$) coincides well with the sum of the covalent radii corresponding to a triple bond: $r_3(Nb) + r_3(O) = 169\text{ pm}$.^[19] The *ipso* carbon atoms of the C_6F_5 groups define a nearly regular pentagonal base with long Nb–C bonds ($235.6(3)\text{ pm}$ average value). The helical arrangement of all five C_6F_5 rings around the Nb–O axis (tilt angles of approximately 75° with respect to the basal plane) makes the $[NbO(C_6F_5)_5]^{2-}$ anion chiral and both *C* and *A* configurations were present in the crystal. This arrangement is related to that found in the seven-coordinate species $[ZrF_2(C_6F_5)_5]^{3-}$, which has a pentagonal-bipyramidal (*PBPY*-7) structure with tilt angles of approximately 65° with respect to the equatorial plane.^[20] It is interesting to note that, in the latter compound, the zirconium atom lies in the equatorial

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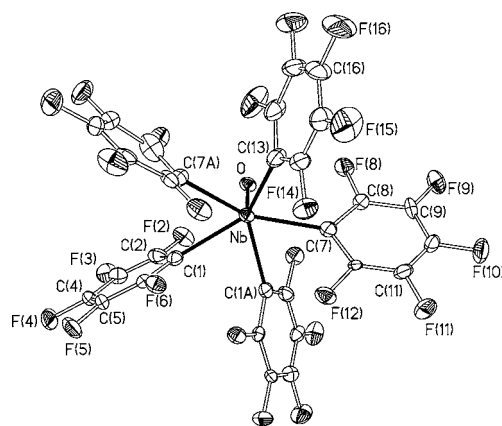
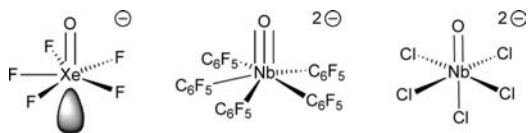


Figure 1. Displacement-ellipsoid diagram (50% probability) of the $[\text{NbO}(\text{C}_6\text{F}_5)_5]^{2-}$ anion as found in single crystals of **1**·*n*-hexane. Selected bond lengths [pm] and angles [°] with estimated standard deviations: Nb–O 168.5(4), average Nb–C 235.6(3), average C–Nb–O 101.4(2), average contiguous C–Nb–C 70.4(1).

plane defined by the five donor carbon atoms as expected for a *PBPY*-7 geometry. Even in the formally *PPY*-6 xenon oxyfluoride derivative $\text{NO}[\text{XeOF}_5]$ (Scheme 3) the xenon atom is only very slightly above the pentagonal plane (2.5(1) pm).^[8a] This arrangement can be attributed to the presence of a stereochemically active lone valence-electron pair on the xenon atom in the hypervalent $[\text{XeOF}_5]^-$ anion, which can thus be considered as a virtual seven-coordinate species with a pseudo-pentagonal-bipyramidal structure.^[8b] On the other hand, the niobium atom in the hypovalent d^0 species **1**, which has no lone valence-electron pairs on the metal, is located 47 pm above the basal plane.



Scheme 3. Structural comparison between related $[\text{AOX}_5]^{9-}$ species.

The unusual *PPY*-6 geometry found for the niobium center in **1** is in sharp contrast (Scheme 3) to the common *OC*-6 structure previously established for the homologous derivative $[\text{AsPh}_4][\text{NbOCl}_5]$.^[21] The latter compound is formed following ligand addition to the same oxohalide precursor, NbOCl_3 (Scheme 2). The structural difference is particularly conspicuous considering that Cl and C_6F_5 ligands are thought to have similar electronic effects as substituents.^[22] It has been suggested, however, that electron-poor metal compounds containing $(\sigma + \pi)$ -donor ligands usually follow the VSEPR model, while those containing mainly σ -donor ligands are more prone to adopt non-VSEPR structures.^[23] The different coordination polyhedra found for the isoelectronic $[\text{NbOX}_5]^{2-}$ species (Scheme 3) are in keeping with this proposal, because halides ($\text{X} = \text{Cl}$) belong to the first ligand category and σ -organo groups ($\text{X} = \text{C}_6\text{F}_5$) belong to the

second. The oxo ligand in $[\text{NbOCl}_5]^{2-}$ exerts a considerable *trans* influence on the axial chloride ligand, as the Nb–Cl_{ax} distance (256.5(4) pm) is significantly longer than the average Nb–Cl_{eq} (237.9(4) pm).^[24] The oxo ligand in $[\text{NbO}(\text{C}_6\text{F}_5)_5]^{2-}$ is *trans* to a vacant site, which shows no affinity for ligands such as NCMe, CNtBu, and OCMe_2 . Although it is sensible to assume that the *ortho*-fluorine substituents would shield this vacant site on the metal, there is no evidence to support the existence of any *o*-F...Nb interaction (see below). We believe that the marked *trans*-influence of the triply bonded oxo ligand^[25] in **1** together with the mainly σ -donor character of the C_6F_5 ligands are responsible for the unusual geometry observed. Steric effects might also apply, but it has been experimentally established that five C_6F_5 groups are indeed able to adopt a square-pyramidal (*SPY*-5) arrangement even around a light metal atom as in the homoleptic organochromium(III) derivative $[\text{NBu}_4][\text{Cr}(\text{C}_6\text{F}_5)_5]$.^[26]

The ^{19}F NMR spectrum of **1** in CD_2Cl_2 solution at 183 K (Figure 2b) is in keeping with the solid-state structure. Just one signal is observed for the *para*-fluorine substituents ($\delta_{\text{F}} = -165.7$ ppm) suggesting chemical equivalence of all five C_6F_5

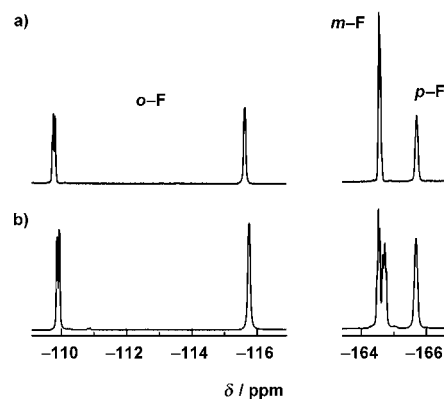


Figure 2. ^{19}F NMR spectrum (470.385 MHz) of **1** in CD_2Cl_2 solution: a) at 293 K and b) at 183 K.

groups. The fact that two signals are observed for the *ortho*-fluorine substituents ($\delta_{\text{F}} = -109.9$ and -115.7 ppm) denotes hindered rotation around the Nb–C bond.^[27] The sharply different chemical shifts found for the two kinds of *ortho*-fluorine substituents are in agreement with their substantially different chemical environments. Likewise, two signals are also observed for the *meta*-fluorine substituents ($\delta_{\text{F}} = -164.5$ and -164.7 ppm). When the temperature is raised to 293 K (Figure 2a), the signals corresponding to the *ortho*- and *para*-fluorine substituents maintain their described positions, while those corresponding to the *meta*-fluorine substituents merge into a single signal at the midpoint between them ($\delta_{\text{F}} = -164.6$ ppm). This behavior suggests that a fluxional process takes place upon raising the temperature. Unfortunately, a thorough study of this incipient dynamic process was precluded by the thermal instability of the sample; compound **1** readily decomposes in solution above 293 K.

The *PPY*-6 geometry is also energetically favored in the gas phase, according to structural optimizations on the $[\text{NbO}(\text{C}_6\text{F}_5)_5]^{2-}$ anion by density functional theory (DFT)

methods (see the Supporting Information for further details). Three bonding orbitals are found to be involved in the Nb–O interaction: $1\sigma + 2\pi$ (Figure S9). The molecular orbital corresponding to the σ bond arises from the overlap of a mainly d_{z^2} orbital (a'_1 symmetry) with the $O(p_z)$ orbital. Interaction between the (d_{xz}, d_{yz}) degenerate orbitals of the $Nb(C_6F_5)_5$ fragment (e''_1 symmetry) with the $O(p_x, p_y)$ orbitals of the oxo ligand gives rise to two orthogonal π bonds, which are also degenerate in energy. The triple bond character is in keeping with the short Nb–O bond length and the high *trans*-influence exerted by the strongly bound oxo ligand. No real minimum was found for an *OC*-6 geometric isomer. In the latter polytope, the marked *trans*-influence of the oxo ligand was evidenced by the substantial lengthening of the axial Nb–C bond (up to 268 pm). Another higher-energy polytope (152 kJ mol^{−1} above the *PPY*-6 one) was found to be a seven-coordinate species in which the axial C_6F_5 group shows a marked lateral swing to enable an additional *o*-F–Nb bond (Figure S7). The existence of even secondary *o*-F...Nb interactions in the *PPY*-6 polytope can be ruled out based on structural and electronic grounds. Thus, the moderate swing observed for the C_6F_5 groups in this case (α_2 and α_3 angles in Figure S6) tends to bring the *ortho*-fluorine substituents farther away from the niobium center (see α_2 and α_3 values given in Table S1). Moreover, a topological analysis of the quantum charge-density function (ρ) calculated for the *PPY*-6 polytope (DFT optimized geometry) using the quantum theory of “Atoms In Molecules” (QTAIM) has not afforded the location of bond critical points linking any of the *ortho*-fluorine substituents to the niobium center.

In summary, the six-coordinate organoniobium(v) compound $[NBu_4]_2[NbO(C_6F_5)_5]$ (**1**) is, to the best of our knowledge, the first organotransition-metal complex for which a *PPY*-6 structure has been established (X-ray diffraction). This unusual geometry is not just a solid-state effect, as it seems to be preserved in solution at low temperature (¹⁹F NMR spectroscopy) and it is the most energetically favored geometry in the gas phase as well (DFT calculations). Species with the *PPY*-6 geometry had been previously suggested as potential intermediates in ligand rearrangement processes operating in six-coordinate compounds,^[28] but later rejected because they were estimated to lie too high in the energy profile.^[29] The results herein demonstrate that the *PPY*-6 geometry can indeed be stabilized to such an extent as to become the ground-state structure. This opens up the possibility for *PPY*-6 derivatives to be seriously considered as valid intermediates or transition states in mechanistic pathways, especially in processes involving mainly σ -donor ligands.

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