PS07.01.08 SYNTHESES OF [IrCl₆]²⁻ {[HP-*i*-Pr₃]+}₂,1 AND [Ir₂Cl₈(P-*i*Pr₃)]²⁻ {[HP-*i*-Pr₃]+}₂, 2, TWO POSSIBLE STARTING MATERIALS FOR THE DIRECT SYNTHESIS OF Irv(H)₂(Cl)₃(P-*i*-Pr₃)₂,3¹.CRYSTAL STRUCTURES OF 1, 2 AND 3¹. P. Mura, M. Allevi, A. Ettorre, Istituto di Strutturistica Chimica "G. Giacomello", CNR, C.P. N° 10, 00016 Monterotondo Stazione - Roma - Italy

Very few iridium(V) (d4) complexes are known2. To our knowledge phosphine complex IrV(H)2(Cl)3(P-i-Pr3)2, 31 is the first Ir(V) compound in which hydride and chloride ligands are present. We obtained complex 3 as by-product in the syntesis of the IrIII(H)(Cl)₂(P-i-Pr₃)₂,4³. Despite our efforts, the direct synthesis of 3 was unsuccessfull. The syntheses of $[IrCl_6]^2 + \{[HP-i-Pr_3]^+\}_2, 1 \text{ and } [Ir_2Cl_8(P-i-Pr_3)]^2 + \{[HPi-Pr_3]^+\}_2, 1 \text{$ 2 may give us two possible starting materials to obtain 3. In fact complexes 1 and 2 are also two transient intermediates in the synthesis of 4 and probably, before, of 3: reductive elimination of a molecule of HCl by 3 may give rise to 43. Crystallografic data: For [IrCl₆]²- {[HP-i-Pr₃]+}₂, 1 (at 298 K), a=9.314(1) Å, b=15.415(2) Å, c=10.073(1) Å, $\beta=95.38(1)^{\circ}$ in space group P 21/n. For [Ir2Cl8(Pi-Pr3)]2- {[HP-i-Pr3]+}2, 2 (at 298 K), a = 16.360(4)Å, b = 13.967(2)Å, c = 18.603(3)Å, $\beta = 93.74(2)^{\circ}$, in space group P $2_1/c$. For IrV(H)₂(Cl)₃(P-*i*-Pr₃)₂, 3 (at 298 K), a = 8.761(2) Å, b= 8.823(1) Å, c = 9.808(2) Å, $\alpha = 100.66(1)^{\circ}\beta = 91.20(1)^{\circ}$, $\gamma = 119.72(1)^{\circ}$ in space group P 1.

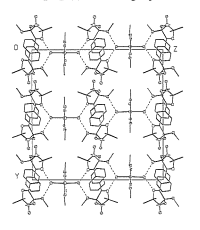
(1) Presented in part to the XVI Congress of IUCr, 1993, Beijing, China, Collected Abstract p. 240.

(2) M. Loza, J. W. Faller, R. H. Crabtree, Inorg. Chem., 1995, 34, 2937 and reference therein.

(3) D. Capitani, P. Mura submitted to Inorg. Chem.

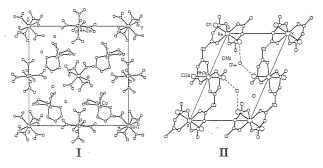
PS07.01.09 CRYSTAL STRUCTURE OF NICKEL(II) AND COBALT(II) ADDUCTS WITH 3-IMIDAZOLINE NITROXIDE. N.V. Pervukhina, A.B.Burdukov, V.I. Ovcharenko, Institute of Inorganic Chemistry Sib. Branch RAS, Novosibirsk, Russia, International Tomography Center Sib. Branch RAS, Novosibirsk, Russia

The crystal and molecular structures of novel compound Ni(hfac)₂(H₂O)₂L₂ (hfac=hexafluoroacetylacetonato-, L=2,2-dimethyl-5,5-dimethoxy-4-phenyl-3-imidazoline-1-oxyl) have been determined by single-crystal X-ray analysis ('SYNTEX P2₁', MoK, /2-scan). The coordination polyhedron of the nickel atom is formed by four hfac oxygen atoms (Ni-O 2.009(2)-2.036(2) Å) and two water oxygens (Ni-O 2.075(3) and 2.029(3) Å) which comprise almost regular octahedron. The nitroxide moieties L do not participate in coordination but form the H-bonds with coordinated water molecules (N•••H 2.01, 1.93 Å and O(CH₃)•••H 2.00, 1.97 Å), therefore joining Ni(hfac)₂(H₂O)₂L₂ are orthorhombic, space group P2₁2₁2₁, with 8 molecules on unit cell of dimensions a=11.625(3) Å, b=19.467(7), c=19.547(8) Å, V=4423.5(3) Å³, D =1.513 g•cm⁻³, 1272 unique I_{hkl} >2 σ_{I} , R=0.0458). The complex Cb(hfac)₂(H₂O)₂L₂ is isostructural to Ni(hfac)₂(H₂O)₂L₂ according to powder diffraction data.



PS07.01.10 'CLOSEST PACKINGS' LAW IN CRYSTAL STRUCTURES OF NEW Rh(III) COMPLEXES. N.V. Podberezskaya, G.V. Romanenko, Institute of Inorganic Chemistry Sib. Branch RAS, Novosibirsk, Russia

Crystal structures of two novel coordination Rh(III) compounds were determined (SYNTEX P21, MoK, /2-scan, 3°250) : cubic $[Co(en)_3][Rh(NO_2)_6]$ •3H₂O, en=C₂H₈N₂ (I) (a=16.540(5) Å, V=4525(2) Å³, Pa3, Z=8, D_c=1.974 g cm⁻³, 589 unique I_{hkl} > $2\sigma_I$, R=0.0905) and triclinic Na[Rh(acac)_2Cl_2]•H_2O, $acac=C_5H_7O_2^{-1}$ (II) (a = 12.616(6), b = 8.411(4), c = 8.499(4) Å, ∞ =108.37(4), β= 95.47(4), γ= 110.45(4)°, V = 780.3 Å³, P1(-), Z = 2, D_c =1.758 g cm⁻³, 1200 unique I_{hkl}>2 σ_I , R=0.0463). In I, the joint packing of the complex cations, anions and water molecules presents the stacking after the three-layered closest packing with the combination of NaCl and CaF₂ structure types; in **II**, the complex anions produce the stacking after the hexagonal closest packing with Na+ and H2O located in its cavities. The presence of organic ligands in the composition of the complexes does not contradict our understanding of packing rules for inorganic coordination compounds in the description of complex cations and anions as the quasi-spheric particles of the Ref radius.



PS07.01.11 BULK HELICOIDS IN CRYSTAL STRUCTURES OF RARE-EARTH AND SODIUM TETRAKISPIVALOYLTRI-FLUOROACETONATES, NaLn(PTA)₄. T.M. Polyanskaya, G.V. Romanenko, N.V. Podberezskaya, Institute of Inorganic Chemistry Sib. Branch RAS, Novosibirsk, Russia

We have carried out an X-ray study of the volatile NaLn(PTA)4 (Gd-Lu) complexes. The parameters for Er are as follows: a=13.114, c=21.004 Å, sp. gr. P3₁, Z=3. In the structure, each PTA ligand performs a tridentate function, forming a six-membered chelate cycle with the Er atom, and, besides, coordinates the neighboring Na atom by the additional Na-O bond. The coordination polyhedron of each metal atom includes the O atoms of four PTA ligands. The Na atom bridges two nonequidistant Er atoms (3.833 and 3.893 Å). The M-O bond strengths are much the same (the mean Er-O and Na-O bond values 2.325 and 2.396 Å). Four vertices of the eight-vertex Na polyhedron are occupied by the F atoms (the mean Na-F 2.678 Å) of the same PTA ligands which are shared by Na and Er. The trans-edge-sharing eight-vertex polyhedra about Er and Na form the array with extended structure which runs toward a crystallographic 31 axis, but the Er and Na ions are not sited on the axis and the extended array has a 'helical' structure. The independent link of the array is equal to two polyhedra : the Na1 atom, nearest to Er, supposedly converts into a gaseous phase.

