MS07.00.05 MINERALOMIMETIC CRYSTAL STRUCTURES OF CN-LINKED CADMIUM COMPLEXES. T. Iwamoto, Department of Chemistry, College of Arts and Sciences, The University of Tokyo, Komaba, Meguro, Tokyo 153, Japan

Polymeric structures of cadmium cyanide and polycyanopolycadmates give a number of CN-linked mineralomimetic crystal structures like pyroxene, clays, H- and Lcristobalite, H-tridymite, zeolites, rutile, beryl, pyrite, etc., with or without guest molecules accommodated in inter- or intralayer space in 2D structures, or in cavities in 3D structures. The Cd atoms in the mineralomimetic structures take the positions corresponding not only to tetrahedral Si but also octahedral Fe, Mg, Ti, etc. The Cd-CN-Cd span longer than Si-O-Si or Si-O-M generates a more voluminous void space in the mineralomimetic latticework than in real minerals so that guest molecule or bulky secondary ligand is required to stabilise the latticework structure. In other words the guest or the ligand plays a role of template to stabilise the mineralomimetic structure. The zeolite-like 3D host of [Cd<sub>3</sub>(CN)<sub>7</sub>]- gives isomorphous clathrates of C<sub>6</sub>H<sub>6</sub>, PhMe, o- and m-C<sub>6</sub>H<sub>4</sub>Me<sub>2</sub>, and PhEt of Pnam space group but the clathrates belong to P6<sub>3</sub>/mmc and Pa for p-C<sub>6</sub>H<sub>4</sub>Me<sub>2</sub>. Involvement of imidazole (imH) as a secondary ligand at octahedral or tetrahedral Cd in the host structures affords zeolite-like C<sub>6</sub>H<sub>6</sub> and PhMe, beryl-like m-C<sub>6</sub>H<sub>4</sub>Me<sub>2</sub>, rutile-like pC<sub>6</sub>H<sub>4</sub>Me<sub>2</sub>, and clay-like PhEt guest clathrates. Silica-like Cd(CN)2 host gives polymorphs similar to H- and L-cristobalite and H-tridymite depending upon the geometry of guest molecule. A pyrite(FeS2)-like array of octahedral Cd2+ and [Cu2(CN)7]5- isostructural to pyrosilicate-like [Cd2(CN)7]3is observed in  $H_3[Cd\{Cu_2(CN)_7\}]-14H_2O$  together with a rutile(TiO<sub>2</sub>)-like array of hydrogen-bonded water molecules, 12H<sub>2</sub>O ring like Ti4+ and 2H2O like O2-.

MS07.00.06 REVOLUTIONARY ADVANCES IN CRYSTALLOGRAPHIC ANALYSES OF HIGH-NUCLEARITY TRANSITION METAL CLUSTERS: FROM POINT-DETECTOR TO CCD AREA-DETECTOR SYSTEMS. Lawrence F. Dahl<sup>1</sup>, Masaki Kawano<sup>1</sup>, Jeffrey W. Bacon<sup>1</sup>, John M. Bemis<sup>1</sup>, Peter D. Mlynek<sup>1</sup>, Nguyet T. Tran<sup>1</sup>, Charles F. Campana<sup>2</sup>, <sup>1</sup>Department of Chemistry, University of Wisconsin-Madison, Madison, WI 53706 <sup>2</sup>Siemens Analytical X-Ray Instruments, 6300 Enterprise Lane, Madison, WI 53719

The availability of the SMART charge-coupled-device (CCD) area detector system approximately two years ago for collecting single-crystal X-ray diffraction data with MoK $\alpha$  radiation has proven to be crucial for our current research on giant-sized metallic and bimetallic clusters. This lecture will illustrate: (1) the much greater capability of the CCD system relative to a point-detector system in determining both the molecular architectures and stoichiometries of a number of geometrically unprecedented close-packed bimetallic Ni/Pd and Ni/Cu clusters, especially for weakly diffracting and/or small crystals; and (2) crystallographic problems that bedevil modern X-ray diffraction analysis of large metal clusters.

PS07.00.07 EFFECT OF ASCORBIC ACID AS REDUCING AGENT IN THE PREPARATION OF 1:1 CHLORO(L)Cul COMPLEXES(L:PYRIDINE, QUINOLINE, ACRIDINE). A. Alemi, S. Shirinevare, Dept. of inorganic. Chem., Tabriz Univ. Tabriz-Iran

Some  $Cu^{II}$   $L_2C1_2(L:Acridine, pyridine and other Heterocyclic Amines) complexes were prepared by Dubsky et co-worker(I) and also PETER et al (2) were studied some characters and structure of these type complexes. Investigation of the Reaction between ascorbic acid and copper(II) chlorid in the presence of some heterocylic amines (pyridine, quinoline, acridine) shows that, the product of reactions are in the form of <math>Cu^{IC}$  L complexes, and

also the reactions of  $Cu^{II}Cl2L2$  complexes with ascorbic acid shows the same product, as we were found previously. In this study we were used elementry analysis, IR, FT-IR, NMR, XRD and also conductivity and magnetic succeptibilty measurement of the products were examined. To summarize,the Role of ascorbic acid in two different reaction process is, no any ligand interaction with complexes. It should be noted that the action of ascorbic acid with Cu(II) complexe with transformation of then to Cu(I) complexe is a new method of  $Cu^{II}$  complexe preparation from  $Cu^{II}$  complexe without any ligand exchange.

1)Dubsky,J. V. et al colln. Czech. Chem. Commun, 1(1929)528 2)PETER, C. et al Aust. J. Chem 1989,42,115-36

PS07.00.08 RHENIUM(V)-OXO COMPLEXES WITH IMI-DAZOLES: PRIME CANDIDATES FOR DISORDER, SU-PERSTRUCTURE AND COUNTERION SUBSTITUTION. Suzanne Bélanger and André L. Beauchamp, Département de chimie, Université de Montréal, C.P. 6128, Succ. centre-ville, Montréal, Québec, H3C 3J7, Canada

Rhenium(V) dioxo complexes show interesting optical properties and a potential for applications as radiopharmaceuticals. To understand these properties, compounds with various imidazoles were prepared. These compounds proved to be a prolific source of challenging crystallographic problems.

A *trans* octahedral species containing an axial O=Re=O unit and four equatorial imidazoles can possess a 4-fold axis, but symmetry is often lowered by the poorly scattering imidazoles adopting different orientations. In addition, instability generates  $ReO_4^-$  with liberation of protons. As a result, partial anion substitution by  $ReO_4^-$  and mixed  $[ReO_2L_4]$ +/ $[ReO(OH)L_4]^2$ + phases are not uncommon.

These features lead to space group ambiguity, disorder and extremely unstable refinement conditions. The correct structure is reached only after the unit cell, Laue symmetry and systematic absences have been carefully checked. Refinement in all possible space groups has to be considered and it must be ascertained that disorder is not an artefact due crystal twinning. The examples to be discussed include the oxobridged dimer  $[Re_2O_3(BiimH_2)_4]Cl_4,\ a\ stoichiometric\ hemiprotonated\ phase <math display="inline">[ReO_2(BimH)_4][ReO(OH)(BimH)_4](ReO_4)_3$  and two polymorphs of the oxo-methoxo compound  $[ReO(OCH_3)(1-MeIm)_4](BPh_4)_2.$ 

 $(BiimH_2 = bi-imidazole, BimH = benzimidazole, 1-MeIm = 1-methylimidazole).$ 

PS07.00.09 STRUCTURAL INVESTIGATIONS OF BIMETALLIC Sb3+ ETHOXIDES AND A COMPARISON TO RELATED STRUCTURES. U. Bemm and R. Norrestam, Department of Structural Chemistry, Stockholm University. S-106 91 Stockholm, Sweden.

By an systematic investigation of the Ni2+ and Mn2+ antimony ethoxide systems [1] several species are found. Seven of these compounds have been structurally investigated, e.g. Ni<sub>6</sub>Sb<sub>4</sub>O<sub>4</sub>(OEt)<sub>16</sub>(HOEt)<sub>4</sub> (I) [2], Mn<sub>7</sub>Sb<sub>4</sub>O<sub>4</sub>(OEt)<sub>18</sub>(HOEt)<sub>2</sub> (II) [4] and  $Mn_8Sb_4O_4(OEt)_{20}$  (III) [3], by single-crystal X-ray diffraction methods. Some of the species are structurally quite similar and have been investigated for further structural relationships, see e.g. figure below. The obtained metal and oxygen atom core fragments of the investigated structures have also been compared to other alkoxides and related compounds found with the Cambridge Structural Database System, CSDS. 90 structures were retrieved from the database and the investigation has resulted in a simple classification of the  $M_xO_y$  type, where the core fragment with the relation be further described  $[\mu_3,\!\mu_4,\!\mu_5][m_1,\!m_2,\!m_3,\!m_4,\!m_5,\!m_6].$   $\mu_a$  and  $m_b$  are the coordinations around the oxygen atoms and the metal atoms, respectively. 90%