THE STRUCTURE OF A HG(II) COMPLEX 03.5 - 5OF ADENINE N(1)-OXIDE. By <u>M. Damodara Poojary</u> and H. Manohar, Department of Inorganic and Physical Chemistry, Indian Institute of Science, Bangalore 560012, India.

The structure basically consists of a dimeric unit with a centre of symmetry involving two $HgCl_2$ and two adenine N(1)-oxide molecules.

Adenine N(1)-oxide acts as a bridging bi-dentate ligand coordinating through N(7) and O(1). The chloride ions link up metal ions to give a polymeric structure. Mercury has a distorted square-pyramidal coordination geometry with O(1) occupying the axial position. Three chloride ions and N(7) atom constitute the base. Hg(II) is indirectly linked to the N(6) atom through a N-H....Cl hydrogen bond.

On the basis of the present structural obser-vations it is suggested that during Hg-DNA complexation, Hg interacts directly with N(6) by releasing a proton at pH \sim 9.0, thus resulting in a chelate. Earlier models, on the other hand, have implicated N(3) and O(4) atoms of thymidine as binding sites for Hg in DNA. Interesting features of the structure in relation to Hg-DNA interaction will be discussed.

Crystal data : $Hg(C_5H_5N_5O)Cl_2$, monoclinic, $P2_1/n_0 = 6.685(1), b = 11.798(2),$

c = 10.155(1)Å, β = 100.22(1)°, Z = 4, 1912 observed reflections measured by diffractometer, R = 0.074.

03.5-6 X-RAY STRUCTURE OF (AQUO) (5'-PHOS-PHOPYRIDOXYLIDENEGLYCINATO) COPPER(II) TRI-HYDRATE. By S.P.Sudhakara Rao and <u>H.Manohar</u>, Department of Inorganic and Physical Chemistry, Indian Institute of Science, Bangalore 560012, India and R.Bau, Department of Chemistry, University of Southern California,Los Angeles, California 90089-1062, U.S.A.

The role of metal ions in nonenzymatic pyridoxal-dependent reactions of *A*-amino acids pyridoxal-dependent reactions of α -amino acids has been examined by a number of studies in solution. Metal-pyridoxylideneamino acid com-plex is an intermediate in such a reaction, the metal serving as a trap for the preformed Schiff base of pyridoxal and amino acid. The title compound crystallizes in space group P1 with a=8.394, b=9.819, c=10.884Å, α =101.85, β =97.48, γ =112.60° and Z=2. The structure was solved by heavy atom method and refined was solved by heavy atom method and refined by least squares to R=0.064 for 1464 observed reflections.

Cu(II) ion has a square pyramidal coordination geometry. Three donor atoms from the terdentate Schiff base ligand (phenolic O, imine N and carboxylate O) and a water O constitute the base (M-L distances: 1.9-2.0Å). The axial site is occupied by a phosphate 0 of a neighbouring molecule $(M-O = 2.31\text{\AA})$, resulting in a polymeric structure. Unlike in bis(pyridoxylideneglycinato)nickel(II) complex (Rao and Manohar, XV Natl. Seminar on Crystallography, Bangalore, 1984), the ligand is nearly planar. These observations can be correlated with the lower and higher activi-ties of Ni(II) and Cu(II) ions in nonenzymatic reactions.

03.5-7 CRYSTAL AND MOLECULAR STRUCTURE OF 7-BENZYLAMINO-2-METHYLMERCAPTO THIAZOLO[5,4-d] PYRIMIDINE.By A.De, R.Ghosh and P.Roychowdhury, Department of Physics (X-ray Lab.), University College of Science; 92, A.P.C.Road; Calcutta-700009, INDÍA.

The crystal structure of the compound has been determined from three dimensional X-ray diffraction data for 2478 unique reflections, collected on an Enraf-Monius CAD-4 diffractoradiation with a crystal of dimensions 0.40X0.28X0.13 mm.The crystal data are: $C_{13}H_{12}N_4S_2, M_r = 288.399, monoclinic, space group$ $\beta_{2_1/a,a=8.745(2),b=13.230(8),c=11.728(2)8}, \beta_{2_1/a,a=8.745(2),v=1353.58^3,z=4,p_m=1.43,p_x=1.42}$ Mg m⁻³, $\mathcal{A} = 0.37$ mm⁻¹ for MoK₂ ($\lambda = 0.7107$ Å). The positions of the sulpher atoms were located from a Patterson synthesis and its phases were used to compute a difference Fourier synthesis which enabled the location Fourier synthesis which enabled the location of the other non-hydrogen atoms.All the hydrogen atoms were located from a subsequent difference Fourier synthesis.The structure was refined by the full-matrix least-squares (Busing,Martin and Levy, 1962) to a R of 0.035 for 1788 'observed' reflections [$I \ge 36(I)$] with all hydrogen atom parameters included in the final cycles of refinement.

03.5-8 STUDIES ON NUCLEOBASES AND THEIR DERIVATIVES: CRYSTAL STRUCTURES OF 1-((S)-2,3-DIHYDROKYPROPYL)URACIL(C7H10N204),

5-BROMO, 1-((S)-2,3-DIHYDROXYPROPYL) URACIL ($C_7H_9N_2O_4Br$) AND ARPRINOCID (9 - (2- CHLORO-6-FLUOROBENZYL) ADENINE, C12H9N5FC1) By

<u>P. Thomas Muthiah</u> and S.K. Mazundar, Crysta-llography Division, Saha Institute of Nuclear Physics, Bidhannagar, Calcutta-64, India.

l-((S)-2,3-dihydroxypropyl)uracil (structure I) crystallizes in the space group P2₁2₁2₁ with a = 4.719(1), b = 8.882(1), c= 19.768(3) A° and z = 4; the structure was solved by direct method and refined to R=0.036. 5-Bromo, 1-((S)-2,3-dihydroxypropyl) uracil (structure II) crystallizes in the space group P1 with a = 6.175(r)b = 7.453(2), c = $5.205(1)Å_{3}^{\prime}=100.00$ (2), $\beta = 102.58(3)$, $\gamma = 83.74(3)^{\circ}$ and z = 1. The structure was solved by heavy atom method and refined to R=0.050. The propyl side chain is extended in these two compounds and the related al inhatic nucleoside analogues related aliphatic nucleoside analogues -9 ((S)2,3-dihydroxypropyl) adenine (SDPA) and 9 ((RS)2,3-dihydroxypropyl) adenine, (RSDPA). (G.S.D. King and L.Sengier, J.Chem.Res. (M), 1501 (1981)). The two hydroxy groups are tran with respect to each other in structures II trans and in SDPA, but gauche in structure I and and in SDFA, but gauche in structure I and RSDFA. In structure I, the plane defined by $C(1^{i})$, $C(2^{i})$ and $C(3^{i})$ also contains $O(3^{i})$ and makes an angle of 70.7° with the uracil plane. In structure II, the plane defined by $C(1^{i})$, $C(2^{i})$ and $C(3^{i})$ makes an angle of 97.3° with the uracil and the two hydroxy oxygen lies on