02.12-1 NEUTRON DIFFRACTION STUDY OF THE INTERACTION OF ETHANOL WITH LYSOZYME. By M.S. Lehmann, S.A. Mason and G.J. McIntyre, Institut Laue-Langevin, Grenoble Cedex, France.

The study was undertaken to estimate experimentally both the relative number and specific locations of the hydrophobic interactions in the lysozyme structure. The goal was to assess the usefulness of neutron diffraction techniques for the examination of very small deuterated molecules in the protein solvent.

A non-deuterated triclinic crystal of hen egg-white lysozyme (Nisonoff, J.M., Brik, L.C. & Jensen, L.H. (1975), Acta Cryst. A31, 16) was soaked at pH 4.6 in a solution containing 25% C2D5OH. Reflections were measured in 9 days to 200 Å on the neutron diffractometer D6, and consisted of all data to 2.6 Å and the 70% strongest to 2.0 Å; in all 6047. The wavelength was 1.675 Å and the cell is 27.32 Å, 32.158 Å, 34.274 Å, 68.24°, 108.63° and 111.80°. Ethanol molecules were located after preliminary refinements, and 16 ordered water molecules from the deuterated structure were included. Each ethanol molecule was described as two hard spheres of radii 1 Å separated by 2 Å. The Hendrickson-Konnert restrained refinement program was used, and the final R value was 0.097. Including the flat solvent did not improve the R factor significantly, but it led to a very good agreement between the thermal parameters for this structure and the previous high-resolution neutron studies of the triclinic lysozyme structure (Nisonoff, S.A., Bentley, C.A. and McIntyre, G.J. (1982) Brookhaven Symposium 32, in press).

In all 15 ethanol sites were found with occupation between 0.0 and 0.2 and in every case there was clear evidence of hydrophobic interactions. Comparison of the experimentally determined local hydrophobicity of the surface with calculation of accessible surface, and with the water structure in non-alcoholic lysozyme will be given.

02.12-2 ON THE CONSERVATION OF PROTEIN-SOLVENT INTERACTIONS IN IMMUNOGLOBULIN VARYING DOMAINS. By S. Swaminathan, W. Furey, C.S. Yoo, B.C. Wang and M. Sax, Biocrystallography Laboratory, P.O. Box 12055, VA Medical Center, Pittsburgh, PA 15240 and the Department of Crystallography, University of Pittsburgh, Pittsburgh, PA 15260.

A detailed study of the water structure of Bence-Jones protein Rhe was made at 1.8 Å resolution (Furey, Wang, Yoo and Sax (1983), J. Mol. Biol. 167, 651-665). Several tightly bound water molecules making at least two hydrogen bonds with the protein were found; in some cases the water molecules were located in a cavity and were completely surrounded by residues of non-hypervariable regions. We suggest that these waters may be in an ordered region and should be considered an integral part of the protein itself.

As a test of this idea, we have undertaken a study to see whether these structural water molecules are present (or can be accommodated) in other crystal structures of V domains. We found that the structural waters in the non-hypervariable regions of Rhe could indeed be accommodated in the corresponding regions of other V domains. The method and results will be presented.

* Deceased, August 31, 1983

02.12-3 ENVIRONMENTAL EFFECTS ON WATER-MEDIATED TRANSFORMATIONS IN THE CRYSTALS OF RIBONUCLEASE A. By D.M. Salamone, R. Kodandapani and M. Vidyasagar, Molecular Biophysics Unit, Indian Institute of Science, Bangalore 560012, India.

We have earlier shown that a new monoclinic form of ribonuclease A, prepared by acetone diffusion into a protein solution in triethylamine, undergoes a reversible transformation, as evidenced by changes in the diffraction pattern and the cell dimensions, when the relative humidity around the crystal is reduced to 93% (Curr. Sci. (1984) in press). Subsequent studies indicate that the well-known crystal form of ribonuclease A grown from 55% 2-methylpentan-2,4-diol (MPD) in phosphate buffer does not transform even when the relative humidity is reduced beyond 75% whereas the same form grown from aqueous ethanol transforms between relative humidities of 93% and 90%. The new crystal form, after being soaked in the phosphate buffer-MPD mixture, transforms only when the relative humidity is reduced to about 75%. It, however, readily transforms at a relative humidity of 90%, when aqueous ethanol is used for soaking. Further experiments on water-mediated transformation of different forms of ribonuclease A and other proteins, under different environmental conditions are in progress. The results obtained so far suggest that environmental factors significantly influence these transformations which could well involve well-defined conformational transitions as a function of hydration, in addition to changes in crystal packing.

02.12-4 ANALYSIS AND PREDICTION OF SOLVENT STRUCTURES IN PROTEIN CRYSTALS. By J. Noutl, A.R. Sialecki and M.N.G. James, SRC Group in Protein Structure and Function, Biochemistry Department, University of Alberta, Edmonton, Alberta, Canada.

The refined high resolution structures of eight enzymes and enzyme inhibitor complexes obtained in this laboratory have provided the basis for an analysis of ordered solvent structure in protein crystals. The results of this analysis have been used to develop an algorithm for the prediction of ordered water molecule locations on the surface of globular protein molecules.

In agreement with other workers, we find a monolayer of ordered solvent covering nearly all of the accessible surface of these proteins, with approximately 15% of the observed solvent molecules forming part of a second layer. The molecules exhibit order and occupancy, with a strong correlation between these properties and the degree of interaction with the protein surface, as assessed in both electrostatic and van der Waals terms.

Although all these crystals are grown in the presence of high concentrations of salt, we find only three unique ordered ion positions in the entire set. All three are involved in inter-molecular interactions. We conclude that the large disordered solvent regions in these crystals offer an energetically more attractive environment for this type of ion. Thus, binding is only observed in very specific niches, or when the number of ions needed to maintain electrical neutrality is too high for them all to be accommodated in the disordered region.

The ordered solvent structure often forms a bridge between adjacent protein molecule surfaces, and there are rather few direct protein-protein contacts. Possible interpretations of such bridge solvent structures are that they increase the stability of the crystal