01.1-7 DATA COLLECTION FACILITY FOR STUDY OF BIOLOGICAL MACROMOLECULES USING SYNCHROTRON RADIATION. By R.N. Sweet, J.B. Hastings, W.C. Thomson, D.S. Wise, and B.F. Schoenborn, Department of Biology and National Synchrotron Light Source, Brookhaven National Laboratory, Upton, NY 11973.

A data collection facility is being constructed at the National Synchrotron Light Source (NSLS) x-ray storage ring. Characteristics of this facility are as follows. X-ray Source: The NSLS storage ring is designed to be operated at 2.5 GeV, 500 mA, with a critical energy of 4.2 keV for the bending magnet source. The electron beam dimensions will be only a few tenths of a millimeter at the tangent point. Monochromatization: Double single crystal monochromator of Ge or Si. Will be automated for rapid selection of wavelength for use of resonant scattering phenomena. Beam focussing: A single cylindrically cut (radius = 27 mm) longitudinally bent (radius = 3 m) mirror lies 9 m from the electron beam tangent point, 9 m from specimen. Collimation: The natural dimensions of the focussed beam will be 0.2 - 0.3 mm. Slits are used to limit the beam to smaller dimensions. The natural crossfire of the beam will be 0.25 mm in the vertical and 3 mm in the horizontal. Detector: X-ray film will be mounted either in flat cassettes with the beam normal to the plane (12.5 cm square) or in cylindrical cassettes with the axis of 10 cm radius curvature lying normal to the beam (12.5 x 18 cm). Detector distance: Flat film may be as close as 5.5 cm, cylindrical film as close as 10 cm. Data processing: Both film-scanning hardware and rotation-film scanning software are available within the Biology Department. This facility may be used by visitors to assess data quality, but most data processing will be done in visitors' home laboratories.

01.2-2 CRYSTAL STRUCTURE OF AZOTOBACTER CYTOCHROME C5 AT 3.5 Å. By G.B. Dustin, D.C. Carter, and S. O'Donnell, Dept. of Crystallography, University of Pittsburgh, Pittsburgh, PA 15260 USA.

The crystal structure of the dimeric cytochrome c5 from Azotobacter vinelandii has been solved from a 2.5 Å resolution MIR map with figure of merit 0.71. The space group is C2 with a = 45.73, b = 37.56, c = 42.55 Å, β = 111.3°, where the dimer is superposed on the crystallographic two-fold axis. The iodine derivative prepared with N-iodosuccinimide has two sites 1.7 and 2.0 Å from the 3' and 5' carbons of tyrosine 18. The platinum derivative prepared with [Pt(H2)2] has three sites at the dimer interface. A model comprising 576 atoms was built into the MIR map and has been partially refined to R = 0.36 with 2.5 Å data.

The heme ligands are cysteines 19 and 22, and the Fe ligands are the heme, histidine 23 and methionine 65. The horne planes of the dimer form a 67° angle and the Fe-Fe distance is 16.4 Å across the dimer interface, which is comprised of hydrophobic contacts. Dimerization partially occludes the heme edge normally exposed to solvent in bacterial c-type cytochromes. However, both propionate groups of the heme are exposed to solvent on a face of the molecule containing an arginine and several lysines, and there is a marked clustering of aspartates and a glutamate on the opposite solvent-exposed face of the molecule. The overall polypeptide fold is homologous to cytochrome c51 (B.E. Dickerson, et al.), but the molecule is more compact, has a higher a-helical content, and contains a disulfide between cysteines 69 and 72.

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Lipoamidedehydrogenase is an FAD containing enzyme which is part of the pyruvatedehydrogenase and a-ketoglu-taric acid dehydrogenase complex. In A. vinelandii it is a dimer with Mw = 112,000.

It catalyses the reoxidation of dihydrolipoic acid which is covalently bound in amide linkage to the core enzyme of the complex: NAD+ + dihydrolipoamide + NADH + H+ + lipoamide

It crystallizes as thin plates from solutions containing 10 mg/ml protein in 0.1 M K-phosphate pH 7.5, 0.1 M EDTA and 20% PEG 4000 in spacegroup P212121, with a = 64.0 Å, b = 83.8 and c = 193 Å and one dimer per asymmetric unit. The plates are enlarged by seeding the small crystals up to 1.0 x 0.6 x 0.2 mm.

The crystals are quite sensitive to radiation damage. The effective lifetime of the crystals could be prolonged with synchrotron radiation. In combination with cooling down to -10° C the resolution was increased up to 2.4 Å. Therefore a native dataset will be collected at DESY, HMBL at Hamburg using synchrotron radiation.

Lipoamidedehydrogenase displays extensive homology with Glutathione reductase. The three-dimensional structures of the latter is known. Elucidation of the structure of lipoamidedehydrogenase will be essential to explain the functional differences between the enzyme and Glutathione reductase.

01.2-3 THE USE OF OPTIMISED ANOMALOUS DISPERSION IN PROTEIN CRYSTALLOGRAPHY FOR HEAVY ATOM LOCATION AND REFINEMENT, AND REFLECTION PHASING. By M.Z. Papiz1, J.R. Belliiwell2, and D.J. Cruickshank2.

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The wavelength tunability of synchrotron x-radiation allows the scattering of specific atoms to be varied according to the standard relation for the atomic scattering factor:

\[ f = f_0 + f'(\lambda) + f''(\lambda) \]

In protein crystallography the use of wavelengths close to the absorption edges of bound metal atoms allow the different values of \( f' \) and \( f'' \) coefficients to be optimised and varied in a controlled way. The changes induced in the diffraction pattern can be used to locate the anomously scattering atom and to phase each reflection. In the case of two different metal atoms, eg Mn and Ca, the different values of \( f'' \) for a given \( \lambda \) for each element allow the specific identification of the atoms bound to the protein and a better discrimination than would be available simply from the different atomic numbers of each atom.

To illustrate the accuracy of data required for the methods, results will be discussed on Pseudomonas aeruginosa cyt. c5 (Sawyer, Harding, Gould, Dames, Papiz and Belliwell (Abstracts this meeting)) and Mn + Carpea lectin (Einspahr, Suguna, Suddath, Kilis, Belliwell and Papiz (Acta Cryst. submitted)) using a variety of wavelengths provided by the experimental workstation for protein crystallography on beam line 7 of the Daresbury SRS.
The strategies adopted in the refinement of occupancies, temperature factors and coordinates of the anomalously scattering data will be considered. Attention will be given to the expected statistical distribution of Bijvoet differences.

**01. DETERMINATION OF MACROMOLECULAR STRUCTURES**

The errors were calculated using phases computed from the refined protein structure (Furey, et al., J. Mol. Biol., 197, 1981) and the heavy atom parameters previously reported. More results and discussions will be presented.

**01.2-4 DIRECT DETERMINATION OF SAS PHASE INFORMATION**

By D.S.C. Yang, Z.B. Xu, W. Furey Jr. and B.C. Wang, Biocrystallography Laboratory, P.O. Box 12055, VA Medical Center, Pittsburgh, PA 15240, U.S.A. and the Department of Crystallography, University of Pittsburgh, Pittsburgh, PA 15260, U.S.A.

When dealing with Single-wavelength Anomalous Scattering data (SAS), the SAS-vector $F_{SAS}$ is defined as the projection of the protein structure factor vector $F_p$ onto the anomalous component of the anomalous scatterer's contribution to the total structure factor. The associated phase angle, $\phi_{SAS}$ is numerically equal to the average value of a pair of true and false phase solutions in the conventional method. Recently we found a relationship for identifying triplet phase invariants with values near $\pi/2$ or $-\pi/2$ linking these SAS phases. That is

$$\phi_{SAS} = \frac{\phi_{PS} + \phi_{PS} + \phi_{PS}}{3} \approx \frac{\pi}{2} \text{ or } -\frac{\pi}{2}$$

where $S$ denotes SAS phases and $\Delta F_p$ denotes $|F_p-F_h|$. Equation (1) is very similar to the rule $R_{ANO,1}$ of Karle (Paper 12, Int. Sch. on Cryst. Comp., Japan, 1983). However, the above relationship is considerably more reliable than $R_{ANO,1}$ which links protein phases. A calculation with experimentally obtained anomalous scattering data, collected from an Au-derivative of Bence Jones Protein Rhe (Wang, et al., J.M.B. 129, 657, 1970) produced the following results:

| #Ref. of largest $|\Delta F|$ | #Triplets | Ave. Error ($^\circ$) | $R_{ANO,1}$ | Equation 1 |
|---|---|---|---|---|
| 50 | 42 | 68.9 | 23.3 | |
| 100 | 343 | 65.9 | 35.7 | |
| 150 | 1062 | 67.0 | 38.9 | |
| 200 | 2476 | 71.6 | 41.8 | |
| 300 | 5000 | 71.7 | 45.3 | |
| 500 | 5000 | 71.7 | 40.1 | |

The errors were calculated using phases computed from the refined protein structure (Furey, et al., J.M.B. 197, 1981) and the heavy atom parameters previously reported. More results and discussions will be presented.

**01.2-5 PROGRESS REPORT ON THE STRUCTURE DETERMINATION OF Cd, Zn METALLOTHIONEIN**

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Single crystals of Cd, Zn metallothionein (isofom II) from rat liver were grown by seeding using sodium formate as a precipitant (Melts, et al., J.B.C. 256, 6255, 1983). The unit cell is tetragonal with space group $P_{4}2_{1}2_{1}2$ (or $P_{4}2_{1}2_{1}2$) and cell constants $a = b = 31.0, c = 120.04 \AA$. There is one molecule per asymmetric unit. Assays of the single crystals are consistent with those of the "isolated" protein which contains 5 Cd and 2 Zn per metallothionein molecule. Native data to 2.3 \AA resolution has been collected by oscillation photography with a rotating anode x-ray source. The merging $R$ factor (based on $F$) is 0.032 for reflections equivalent by point group symmetry and 0.041 for Bijvoet pairs. Data to 2.3 \AA resolution were also collected for a potential isomorphous derivative (Tungsten). The $R$ factor between the native and derivative data is 0.139. In addition to the traditional isomorphous replacement method, we are trying to develop protein phases from the values of 3-phase structure invariants estimated by the direct methods of Hauptman (Acta Cryst. A38, 289, 1982 and Acta Cryst. A38, 632, 1982). Results of the study will be presented. This work is supported by NIH grant GM-32913.

**01.2-6 DIRECT DETERMINATION OF SIR PHASE INFORMATION BY AN EXTENSION OF KARLE’S RULE**

By Z.B. Xu, D.S.C. Yang, W. Furey Jr., M. Sax, J. Rose and B.C. Wang, Biocrystallography Laboratory, P.O. Box 12055, VA Medical Center, Pittsburgh, PA 15240, U.S.A. and the Department of Crystallography, University of Pittsburgh, Pittsburgh, PA 15260, U.S.A.

Karle (Acta Cryst. A39, 800, 1983) recently introduced a simple rule for identifying triplet phase invariants with values near 0 or $\pi$ from single isomorphous replacement (SIR) data. The rule states that:

$$\phi_{SIR} \times \phi_{SIR} \times \phi_{SIR} = \frac{\phi_{SIR} \times \phi_{SIR} \times \phi_{SIR}}{0}$$

where $\Delta F = |F(h,k,l) - F(h,k,l)|$, $F_{SIR}$ and $F_{SIR}$ are structure factor amplitudes for the derivative and native data respectively. The rule is valid for protein phases only when reflections with the largest $|\Delta F|$ in the data set are used.

Recently we applied Karle’s rule to Au-SIR data of Bence Jones Protein Rhe (Wang, et al., J. Mol. Biol., 129, 657, 1979). Using 280 reflections with the largest $|\Delta F|$ values (top 11%) and by means of symbolic addition we obtained all 280 individual phases. When the phases were compared with those computed from the refined protein structure (Furey, et al., J. Mol. Biol., 197, 1981) we found average phase errors of 2.8° and 34.3° for the 131 centric and 149 acentric reflections respectively. However when the new phases were compared with the SIR phases (calculated from refined heavy atom positions) the average phase errors were 4.1° and 16.7° respectively for centric and acentric reflections. This observation and other considerations led us to discover that the approximations used in Karle’s derivation fit well with the idea of assuming an artificial structure factor vector which is the projection of the $F_p$ vector onto the $F_h$ vector.