

03-Crystallography of Biological Macromolecules

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The $\alpha_2(\text{Mg}^{2+})\beta_2(\text{Fe}^{2+})$ shows a very low O_2 affinity. It is constrained more strongly to the deoxy quaternary structure than native deoxy hemoglobin (Minagawa, 1990, M.thesis, Osaka Univ.). Crystallization of deoxy $\alpha_2(\text{Mg}^{2+})\beta_2(\text{Fe}^{2+})$ was carried out in ammonium phosphate/sulphate buffer (pH=6.5) according to the method described by Perutz (*J.Crystal Growth*, 1968, 2, 54-56). Crystals were grown from solution whose final ammonium phosphate/sulphate buffer concentration was 2.4-2.5M. Crystals of $\alpha_2(\text{Mg}^{2+})\beta_2(\text{Fe-CO})$ were grown at room temperature using the batch method previously described by Ward (*J.Mol.Biol.*, 1975, 98, 237-256). The best crystals were grown from 20-25% PEG.1000(w/v). X-ray diffraction data of $\alpha_2(\text{Mg}^{2+})\beta_2(\text{Fe}^{2+})$ and $\alpha_2(\text{Mg}^{2+})\beta_2(\text{Fe-CO})$ crystals were collected to about 1.7Å and 2.0Å resolution, respectively, using synchrotron radiation at the Photon Factory (wavelength=1.04Å, I=341-327mA, E=2.5GeV). The refinement of the $\alpha_2(\text{Mg}^{2+})\beta_2(\text{Fe}^{2+})$ structure was carried out with the restrained least-squares program PROLSQ of Hendrickson and Konnert (*Methods Enzymol.*, 1985, 62, 241-246). An R-factor of 17.1% was obtained for the data between 10.0 and 1.7Å with magnitudes greater than 3.0σ . We are refining the structure of $\alpha_2(\text{Mg}^{2+})\beta_2(\text{Fe-CO})$ crystal, and refined structures will be presented in the conference.

Crystal	Total No. of reflections	Unique reflections	R-merge (%)	Space Group
$\alpha_2(\text{Mg}^{2+})\beta_2(\text{Fe}^{2+})$	209163	90366	6.87	$P2_1$
$\alpha_2(\text{Mg}^{2+})\beta_2(\text{Fe-CO})$	150689	44289	7.58	$P2_12_12_1$

PS-03.04.21 THE CRYSTALLOGRAPHIC STRUCTURE OF BAR-HEADED GOOSE METHEMGLOBIN. By Hua Jing, Guangying Lu,* Ziqian Hua and Xiaocheng Gu Department of Biology, Peking University, Beijing 100871, P.R.China

The protein under investigation is the methemoglobin of Bar-headed Goose, a bird which can fly over the summit of Mt.Everest(8848m) during its normal migration between Tibet and India. In order to elucidate the molecular mechanism of high altitude respiration, the structure at high resolution of Bar-headed Goose hemoglobin is necessary. We present here the method to prepare the crystal of methemoglobin, the way to solve phase problem using molecular replacement method and the procedure of structure refinement.

Crystallization of methemoglobin was carried out at 18°C by the hanging drop method. The protein was precipitated by polyethylene glycol(PEG) at optimum pH 7.8. After one week, crystals could be found in some drops. The crystal belongs to space group $P4_22_12_1$, with 4 molecules in the unit cell of dimension: a=b=81.6Å, c=107.2Å. Crystallographic data were collected on Area Detector to resolution 2.3Å with Rmerge 7.21. The ratio of the observed reflections to the number of theoretical reflections is 94.6%, to resolution 2.3Å.

The initial phases of measured reflections were solved by molecular replacement method. In order to find out the possible orientation of search model in the unit cell and to locate it correctly in the cell, an automatic processing program system

AUTOMR(Yoshiki Matsuura, *Appl.Cryst.*, 1991, 24, 1063-1066) was used to carry out these calculations. The 2.0Å structure of horse hemoglobin from Protein Data Bank(PDB) was used as the search model. It has 70% homology with Bar-headed Goose hemoglobin. Firstly, the search model molecule was put in a cubic P1 cell with cell dimension as large as 100Å and the structure factor was calculated by conventional algorithm to the 4Å resolution. Secondly, the rotation function was calculated and a number of highest independent peaks were listed and put into next calculation. Thirdly, translation function was calculated to find the correct position of rotated search model. In this step, only 100 to 300 reflections at low resolution were used to define correct solutions. Finally, rigid-body refinement was carried out to refine the rotation and translation parameters. The correct solution is usually obtained as a clear minimum of R value in the listing. In our calculation, the minimum R value was 0.428, which corresponds to the angular position (49.48, 270.66, 125.47) in spherical polar coordinates and translation position(0.00, 40.25, 21.83). By plotting the main chain models in the unit cell, a reasonable molecular packing was observed.

From the correctly rotated and translated model of horse methemoglobin, we calculated the initial phases of measured reflections and the initial electron density map. The residues of horse hemoglobin were substituted by that of Bar-headed Goose hemoglobin with FRODO program. Phases of the substituted structure was then calculated and input into XPLOR program together with the measured intensity of reflections. In XPLOR program, simulating annealing method was applied to refine the structure. The final crystallographic R value is 0.245 to 2.5Å resolution. Further refinement of the structure with FRODO program is still in progress. We hope that the mechanism of high altitude respiration could be partly explained after the structure at high resolution is solved.

PS-03.04.22 STRUCTURE OF THE RECOMBINANT N-TERMINAL LOBE OF HUMAN LACTOFERRIN AT 2.0Å RESOLUTION. By Catherine L. Day, Bryan F. Anderson, John W. Tweedie and Edward N. Baker,* Department of Chemistry and Biochemistry, Massey University, Palmerston North, New Zealand.

The three-dimensional structure of the N-terminal half-molecule of human lactoferrin, Lf_N , prepared by recombinant DNA methods, has been determined at 2.0Å resolution. The protein is in its iron-bound form and is deglycosylated. X-ray diffraction data were obtained by diffractometry to 3.2Å resolution and synchrotron data collection, using Weissenberg photography with imaging plates, to 1.8Å resolution. The structure was solved by molecular replacement using as search model the N-lobe (residues 1-333) of native diferric human lactoferrin (Lf) (Anderson, Baker, Norris, Rice & Baker, *J. Mol. Biol.*, 1989, 209, 711-734). Restrained least squares refinement (program TNT) has resulted in a model structure with R = 0.184 for all data (34180 reflections) in the resolution range 8.0 to 2.0Å. The model comprises 2490 protein atoms (residues 4-327), 1 Fe^{3+} ion, 1 CO_3^{2-} ion and 180 water molecules.

The structure of Lf_N is essentially the same as that of the N-lobe of intact Lf, being folded into two similar $\alpha\beta$ domains, with the Fe^{3+} and CO_3^{2-} ions bound in a specific site in the interdomain cleft. These details are not affected by either deglycosylation or expression from cloned DNA. The C-terminus has unfolded, however; whereas in Lf residues 321-333 form a helix crossing between the domains at the back of the iron site, in Lf_N 321-326 are extended, forming a third interdomain β -strand, and 327-333 are disordered. The conformational change is attributed to the loss of stabilising interactions from the C-lobe, and is mediated by Gly residues at 321 and 323. The conformational change also causes Lf_N to release iron more readily by increased solvent exposure of the hinge region at the back of the iron site. The metal and anion binding geometry, hydrogen bonding in the molecule, and solvent structure have been analysed. Two cis-proline residues and a clear example of amino-aromatic hydrogen bonding have been found.