## 03-Crystallography of Biological Macromolecules

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CRYSTAL STRUCTURES OF LVS49 PHOSPHOLIPASES A2 FROM BOTHROPS ASPER AND BOTHROPS GODMANI VENOMS by R.K.Arni<sup>1\*</sup>, R.Ward<sup>2</sup> and J.M.Gutierrez3, 1Department of Physics, UNESP-IBILCE, Sao Jose do Rio Preto-SP, Brazil. <sup>2</sup>EMBL, Heidelberg, FRG. <sup>3</sup>Instituto Clodomiro Picado, Costa Rica.

Recent sequence analysis of phospholipases A2 has indicated the existence of a variant whose characteristic feature is the complete absence of catalytic activity (Maraganore et al., 1987, J.Biol.Chem., 259, 13839) as a result of the substitution of Asp49 by Lys49 in the calcium binding loop. These proteins have been shown to cause liposome leakage by a novel calcium independent process (Rufini et al., 1992, Biochem., 31,12424). Crystals of B.asper PLA are orthorhombic, space group P2(1)2(1)2(1), a=50.2, b=67.8 and c=88.0A, contain a dimer in the asymmetric unit and diffract to 2.8A. Crystals of B.godmani PLA are tetragonal, space group P4(1)2(1)2, a=b=60.6, c=84.7A, Contain a monomer in the asymmetric unit and diffract to 2.6A. The structures have been determined by molecular replacement and refined by simulated annealing (X-PLOR, Brunger, A.T., 1988, J. Mol. Biol., 203,803) and conventional least-squares methods. The results of the structure determinations and structural implications determinations and structural implications of the substitution of Asp49 by Lys49 will be presented.

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PS-03.07.22 X-RAY CRYSTALLOGRAPHIC ANALYSIS OF THE PHOTOSENSITIVE NITRILE HYDRATASE. J. Honda\*, N. Kamiya, T Nagamune, K. Kiribuchi, H. Sasabe, H. Iwasaki and I. Endo, The Institute of Physical and Chemical Research (RIKEN), Saitama 351-01, Japan.

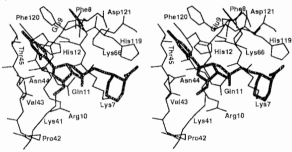
The nitrile hydratase (NHase) hydrates various nitrile compounds to the corresponding amides (Asano, Y. et al.(1980). Agric. Biol. Chem. 44, 2251-2252). The NHase from bacteria Rhodoceccus has active and inactive forms, and the inactive form is readily converted to active form by light irradiation (Nagamune, T. et al.(1990). Biochem. Biophys. Res. Commun. 168, 437-442). This photosensitive NHase consists of 2 subunits αβ each with molecular weights of 22,787 (206 residues) and 23,428 (212 residues) respectively, and it has 2 iron atoms per molecule and a quinône-like colactor (Nagamune, T. et al.(1991). J. Mel. Biol. 220, 221-222). In order to understand its enzymatic mechanism and ultimately the photoactivation mechanism, we have performed an X-ray crystallographic analysis of this NHase. The inactive form of NHase, which is a more stable form of the enzyme, was purified from Rhadococcus, and crystallized by vapor-diffusion method. The crystal belongs to orthorhombic system with space group 121212, cell dimensions of a=117.4 Å, b=145.7 Å, c=52.1 Å and V<sub>m</sub> value of 2.4Å<sup>3</sup>/Da with 2 molecules per asymmetric unit as described previously (ibid.). The intensity data of the inactive NHase crystal was collected using Weissenberg camera and Imaging Plate (Sakabe, N. (1991). Nuclear Instr. Methods Phys. Res. A303, 448-463) at beamline 6A2 in the Photon Factory, Kes. Ag. The data of up to 2.2 Å resolution was collected from only one crystal with a total of 155,481 observed reflections and 33,167 independent reflections, processed using WEIS (Higashi, T. (1989). J. Appl. Crystallogr. 22, 9-18) which was 72% of the total reflections measurable at this resolution. The data gave an overall Rmerge ( $\Sigma 11-<1>1/\Sigma$  I) of 6.8%. The rotation function in polar angles was calculated using POLARREN (by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated a non-tractallographic 2 (and prize with the careful by W. Kabsch) which indicated with the careful by W. Kabsch) which crystallographic 2-fold axis parallel to the a-b plane at  $\phi$ =45°. This result crystanographic 2-ford dats parametro the are plane at 0==0. This result is consistent with the image of transmission electron microscopy of NHase microcrystals which reveals the molecular packing of the NHase. A survey of heavy atom derivatives is currently in progress.

We are grateful to Prof. N. Sakabe and Dr. A. Nakagawa (KEK) for their help at the Photon Factory. We are also grateful to Dr. Y. Sugawara (EEL PN) for her help in computer works.

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PS-03.07.23 MONOCHROMATIC X-RAY STRUCTURE ANALYSIS OF RIBONUCLEASE A AS A PRELIMINARY TO TIME-RESOLVED LAUE ANALYSIS. By T. Nonaka\*, E. Tanno and Y. Mitsui, Department of BioEngineering, Nagaoka University of Technology, Kamitomioka, Nagaoka, Niigata, Japan.

We plan to study dynamics of reaction mechanism of bovine pancreatic ribonuclease A (EC 3.1.27.5), which cleaves singlestranded RNA, by time-resolved Laue method. substantial fraction of unique reflections in a single shot using a Laue camera developed by Dr. N. Watanabe of Photon Factory at Tsukuba, Japan, high symmetry crystal form is preferable. We have Japan, high symmetry crystal form is preferable. We have crystallized intact ribonuclease A (Sigma Type XII-A) by a vapor diffusion method to yield trigonal  $P3_221$  (a = b = 64.19 Å, c = 64.96diffusion method to yield trigonial  $r_{3221}$  (a = b = 04.19 A,  $\gamma = 120$ °) crystals. Monochromatic X-ray diffraction data were collected at 15 °C using a Rigaku R-axis IIc imaging plate detector, using one crystal. The data set with intensity  $|F| \ge 1$   $\sigma$  is 75.7 % complete to 1.7 Å resolution with an R<sub>merge</sub> of 0.049. The coordinate set of a semisynthetic ribonuclease A (Martin, P. D. et al., (1987). J. Biol. Chem. 262, 15930-15938; Brookhaven Protein Data Bank code 1SRN) was used as a search model for rotation and translation search. The catalytic center of the current model is shown below with a proposed caged substrate (an o-nitrobenzyl derivative of cyclic cytidine 2',3'-monophosphate) superimposed.



PS-03.07.24 THE 3-D X-RAY STRUCTURES OF (1-3,1-4) -B- GLUCANASE AND (1-3,1-3) -B-GLUCANASE FROM BARLEY GRAIN TO 2.6A RESOLUTION. By J.N. Varghese\*, T.P.J. Garrett, L. Chena P.B.Hoja and G.B. Finchera, Biomolecular Research Institute, Parkville, Victoria, Australia 3052, a La Trobe University, Bundoora, Victoria, Australia 3083.

The most important enzymes in the depolymerization of the walls of the starchy endosperm in germinating barley are the (1-3,1-4) -B-glucan 4-glucanohydrolases. These enzymes catalyse the hydrolysis of (1-4) -B-glycosyl linkages in (1-3,1-4) -Bglucans, only where the glycosyl residue is preceded by a (1-3) linked glycosyl residue. The (1-3,1-3) -\(\beta\)-glucanase is also expressed at relative high levels in germinating barley, but the functional significance is unclear and could be related to its antifungal properties.

The 3-D structure of the (1-3,1-4) glucanase has been determined by M.I.R. methods to 2.6 A, using three heavy atom derivatives and anomalous dispersion. The 3-D structure of the (1-3,1-3) glucanase which has 50% sequence homology with the (1-3,1-4) glucanase, was solved by molecular replacement using a 3A MIR electron density of the (1-3,1-4) enzyme, and subsequent non-crystallographic symmetry averaging (there are two independent images in the asymmetric unit.