

derivatives and performing SAD experiments on the MX beamlines at the ESRF in Grenoble. Biophysical and biochemical studies have also been performed in order to dissect the role of RecN and its various domains so as to propose a much more accurate mechanism of DSB recognition.

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Keywords: DNA repair, homologous recombination, RecN

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Structural insight into modes of substrate selectivity and catalysis in the shikimate dehydrogenase superfamily

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Shikimate dehydrogenase (SDH) catalyzes the reversible, NADPH-dependant reduction of dehydroshikimate to shikimate, a key step in the biosynthesis of the aromatic amino acids in plants, fungi and bacteria. The absence of the enzyme in animals makes it an attractive target for antibiotics. SDH belongs to an enzyme superfamily, the members of which utilize a common structural scaffold to catalyze reactions involving a range of substrates. We are exploring the diverse substrate preferences of the members of the SDH superfamily by x-ray crystallographic analysis of the enzymes. Our structural characterization of two SDH homologs has identified a complement of active site residues that appear to be important determinants of substrate preference. We investigate the biochemical role of these residues by site-directed mutagenesis. We further explore the significance of these residues by attempting to reengineer the substrate preference of one SDH homolog. In addition, we show by mutagenesis and kinetic analysis that an invariant pair of ionizable active site residues, a lysine and an aspartate, act as a catalytic dyad in two functionally distinct SDH homologs, providing evidence for a conserved catalytic mechanism across the SDH superfamily. Structural and mechanistic characterization of the members of the SDH superfamily will aid in the rational design of drugs targeting the enzymes.

Keywords: Dehydrogenase, enzyme structure, enzyme catalysis

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Crystal structure of Cytosine Deaminase complexed with a mimic of the tetrahedral intermediate

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Cytosine Deaminase from *E. coli* is a member of the amidohydrolase superfamily. The crystal structure of the zinc-activated enzyme was solved in the presence of a mimic of the tetrahedral intermediate. This compound inhibits the deamination of cytosine with K_i of 52 nM. The

zinc and iron containing enzymes were characterized to determine the effect of the divalent cations on activation of the hydrolytic water.

Mutation of Gln-156 decreases the catalytic activity by more than 5 orders of magnitude, supporting its role in substrate binding. Mutations of Glu-217, Asp-313, and His-246 significantly decrease catalytic activity, supporting the role of these three residues in activation of the hydrolytic water molecule and facilitation of proton transfer reactions.

A chemical mechanism for substrate deamination by cytosine deaminase is proposed.

Keywords: enzyme, crystal, structure

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Crystal structure of flavin reductase from *Rhizobium* sp. strain MTP-10005

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Resorcinol hydroxylase from *Rhizobium* sp. strain MTP-10005 is a two-component enzyme system. The small component (GraD) is an oxidoreductase containing a flavin molecule as a cofactor. GraD catalyzes the NADH-dependent reduction of free FAD according to a ping-pong bisubstrate-biproduct mechanism. The reduced FAD is then used by the large component GraA to hydroxylate resorcinol to hydroxyquinol.

GraD was crystallized at 293 K by the sitting-drop vapour-diffusion method using a precipitant solution containing 13 - 14% (w/v) PEG 2000, 6 - 9% (v/v) 2-propanol, 100 mM sodium citrate pH 5.6, 100 mM DTT and 200 μ M FAD. The approximate dimension of the obtained crystals was $0.1 \times 0.1 \times 0.15$ mm³. The crystal diffracted to 1.8 Å and belongs to space group $P4_12_12$ with unit cell parameters of $a = b = 77.7$ Å and $c = 124.2$ Å. The asymmetric unit contains two molecules of GraD with a corresponding crystal volume per protein mass (V_M) of 2.35 Å³/Da and a solvent content of 47.6% by volume. The crystal structure has been determined by molecular replacement and refined at 1.8 Å resolution. The current model was refined to an R -factor of 16.1% ($R_{\text{free}} = 19.2\%$). GraD exists as a homodimer, and each monomer was found to contain an FAD.

Keywords: flavin, reductase, protein crystallography

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Unexpected reactions resulting from mutating catalytic residues in an amidase

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Nitrilase superfamily amidases catalyze the conversion of various amides to their corresponding acids and ammonia using highly conserved Cys, Glu, Glu, Lys (CEEK) catalytic residues. They find applications as potential biocatalysts in the fine chemical industry; as tools for drug synthesis; while those from prokaryotic organisms are attractive drug targets. Although the catalytic mechanism for these

enzymes has been proposed, details of the reaction process are still not clear. This paper presents findings from a study that is aimed at understanding the mechanism of action in amidases, particularly the contribution of each of the catalytic residues in the reaction process.

The four catalytic residues (C165, E61, E139 and K131) have been mutated individually in the *Nesterenkonia* sp. amidase (NitN). In each case, the mutant was reacted with a range of amide substrates and the results analyzed by mass spectroscopy and X-ray crystallography. Mutation of the catalytic cysteine has allowed visualization of non-covalently bound substrates in the active site. However, unexpected reactions resulting in unusual adducts at the reactive cysteine have been observed with the mutants of the two glutamates and the lysine.

Mutation of the two active site glutamates and the lysine resulted in unstable mutants that lacked biochemical activity. While the reaction of E61Q/L mutants with some amide substrates resulted in thioester acyl-enzyme intermediates, 'abnormal' reactions have been observed with fluoroacetamide (FAE) and acrylamide (ACR). These include an S_N2 reaction which displaces the fluorine of FAE and a Michael addition of ACR at the catalytic cysteine, both of which have been visualized in the E61L/Q mutants. E139Q mutant does not exhibit any reactions with most of the amide substrates except with ACR where a Michael addition is observed. A cacodylate adduct at the catalytic cysteine has been visualized in the E139Q active site. The K131Q/H mutants showed no reactions, adducts or intermediates with the tested amide substrates, but a crystal structure of K131Q has revealed an adipamide reaction intermediate covalently attached to the catalytic cysteine. The trapped intermediate is likely to be of an *E.coli* metabolite that reacted with the mutants during protein expression.

The occurrence of unusual reactions with the NitN mutants highlights the importance of the catalytic residues (particularly the glutamates) in ensuring optimal positioning of the substrates in the active site and has also given insights into the critical balance of forces necessary for enzymatic catalysis in amidases.

Keywords: catalysis, mechanism, adduct

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Structural investigation of *Pseudomonas aeruginosa* glyoxalase I enzymes

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Methylglyoxal is a naturally produced cytotoxic electrophile. The toxicity of methylglyoxal is thought to be due to attack of nucleophilic centres of macromolecules including DNA, RNA and proteins. The glyoxalase enzyme system serves to detoxify methylglyoxal via a two-step pathway dependent on cellular glutathione. The first step is the conversion of the non-enzymatically formed glutathione-methylglyoxal hemithioacetal to the corresponding thioester by the metalloenzyme Glyoxalase I. Glyoxalase II cleaves the thioester to produce D-lactate and regenerate glutathione.

Pseudomonas aeruginosa is a multi-host, opportunistic pathogen. It is implicated in the pathology of pneumonia, septic shock, and other types of infection. *P. aeruginosa* uniquely possesses three separate enzymes with glyoxalase I activity, GloA1, GloA2 and GloA3. Other organisms only contain a single glyoxalase I enzyme. Interestingly, GloA1 and GloA2 contain a Ni²⁺ catalytic centre and are inactive when bound to Zn²⁺ unlike GloA3, which is active when bound to Zn²⁺ [1]. Having more than one enzyme with different metal centres may equip the bacteria with greater chemical flexibility allowing the microbe to

inhabit more environments, i.e. areas that are deficient in Zn²⁺ but not Ni²⁺. The presence of three different glyoxalase I proteins within the same organism provides the unique opportunity to compare both metal ion activation classes of this enzyme. X-ray crystallographic studies can provide detailed structural information about how the protein binds to the metal ion as well as metal ion-substrate interactions.

We report the crystal structures of GloA2 with Ni²⁺ and with Zn²⁺ bound at the active site. Anomalous data were collected to confirm the identity of the metal ion. The coordination geometries of the Zn²⁺ and Ni²⁺ centered *P. aeruginosa* GloA2 enzyme are almost identical as seen in the *E. coli* glyoxalase I enzyme [2], which is also only active with an octahedrally coordinated metal group [3].

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Keywords: glyoxalase I, metalloenzyme, coordination

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Structural basis for increased thermal stability of adenylate kinase variants

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Engineering proteins for higher thermal stability is an important and difficult challenge. We describe a comprehensive approach to mutate proteins to be more stable, and report structural analysis of the redesigned protein variants.

First, we identified mutations for thermal stabilization of our model, adenylate kinase based on a variety of experimental and computational techniques. Then, we designed variants by combining the individual stabilizing mutations together. In the experiment using differential scanning calorimetry, the adenylate kinase variants displayed considerable increases in their thermal stabilities.

We determined crystal structures of the variants to confirm the structural basis for their thermal stabilization. The structures showed that the resulting variants have mutation(s) for extra electrostatic interactions by newly added ion pairs, additional hydrophobic interaction and optimized local structural entropy suggesting their contribution to thermal stabilization of the variants.

Keywords: thermostable, mutagenesis, enzyme

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Structure of the β -galactosidase from *Kluyveromyces lactis*

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β -galactosidase (EC 3.2.1.23) is the enzyme responsible of the hydrolysis of the disaccharide lactose into glucose and galactose. It is