Functional Annotation and Three-Dimensional Structure of Dr0930 from *Deinococcus radiodurans*, a Close Relative of Phosphotriesterase in the Amidohydrolase Superfamily^{†‡}

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ABSTRACT: Dr0930, a member of the amidohydrolase superfamily in Deinococcus radiodurans, was cloned, expressed, and purified to homogeneity. The enzyme crystallized in the space group $P3_121$, and the structure was determined to a resolution of 2.1 Å. The protein folds as a $(\beta/\alpha)_{\gamma}\beta$ -barrel, and a binuclear metal center is found at the C-terminal end of the β -barrel. The purified protein contains a mixture of zinc and iron and is intensely purple at high concentrations. The purple color was determined to be due to a charge transfer complex between iron in the β -metal position and Tyr-97. Mutation of Tyr-97 to phenylalanine or complexation of the metal center with manganese abolished the absorbance in the visible region of the spectrum. Computational docking was used to predict potential substrates for this previously unannotated protein. The enzyme was found to catalyze the hydrolysis of δ - and γ -lactones with an alkyl substitution at the carbon adjacent to the ring oxygen. The best substrate was δ -nonanoic lactone with a k_{cat}/K_m of 1.6×10^6 M⁻¹ s⁻¹. Dr0930 was also found to catalyze the very slow hydrolysis of paraoxon with values of k_{cat} and k_{cat}/K_m of 0.07 min⁻¹ and 0.8 M⁻¹ s⁻¹, respectively. The amino acid sequence identity to the phosphotriesterase (PTE) from *Pseudomonas* diminuta is \sim 30%. The eight substrate specificity loops were transplanted from PTE to Dr0930, but no phosphotriesterase activity could be detected in the chimeric PTE-Dr0930 hybrid. Mutation of Phe-26 and Cys-72 in Dr0930 to residues found in the active site of PTE enhanced the kinetic constants for the hydrolysis of paraoxon. The F26G/C72I mutant catalyzed the hydrolysis of paraoxon with a k_{cat} of 1.14 min⁻¹, an increase of 16-fold over the wild-type enzyme. These results support previous proposals that phosphotriesterase activity evolved from an ancestral parent enzyme possessing lactonase activity.

The amidohydrolase superfamily $(AHS)^1$ of enzymes has been shown to catalyze the hydrolysis of a broad spectrum

of amide and ester bonds within carboxylate and phosphate substrates (1). In addition to hydrolytic reactions, members of the AHS have also been shown to catalyze decarboxylation and isomerization reactions (2, 3). All members of the AHS possess a distorted (β/α)₈-barrel structural fold and an active site at the C-terminal end of the central β -barrel with a monoor binuclear metal center (1). The metal center functions to activate solvent water for nucleophilic attack and to stabilize the tetrahedral or trigonal bipyramidal transition state (1). The AHS was first identified by Sander and Holm, who recognized the structural similarities among adenosine deaminase, urease, and phosphotriesterase (PTE) (4). Since the initial discovery in 1997 more than 30 different reactions have been shown to be catalyzed by members of the AHS in prokaryotic and eukaryotic organisms (1, 5).

Many of the enzymes within the AHS have an *unknown*, *uncertain*, or *incorrect* functional annotation. It has been estimated that the total number of reactions catalyzed by members of the AHS will ultimately be greater than 100 (5). We have focused our attention on the elucidation of function for those members of the AHS whose substrate and reaction profiles are unknown. We are also interested in the

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¹ Abbreviations: AHS, amidohydrolase superfamily; ATCC, American Type Culture Collection; IPTG, isopropyl D-thiogalactopyranoside; PHP, phosphotriesterase homology protein; AhIA, *N*-acylhomoserine lactone acylase from *Rhodococcus erythropolis*; PTE, phosphotriesterase; Sso2552, the lactonase also known as SsoPox from *Sulfolobus solfataricus*; PPH, putative parathion hydrolase from *Mycobacterium tuberculosis*; ICP-MS, inductively coupled plasma emission mass spectrometry.

establishment of the structural basis for the molecular evolution of new catalytic activities from existing protein templates. Newly identified reactions catalyzed by members of the AHS include the deamination of *N*-formimino-Lglutamate and *S*-adenosylhomocysteine (6, 7). The identification of the reaction specificities for members of the AHS of unknown function can be addressed via substrate screening of compound libraries (6), genomic and operon context for bacterial enzymes (6, 8), and computational docking of potential substrates to three-dimensional structures (7, 9). Computational docking of high-energy tetrahedral intermediates was effectively used in the identification of the substrate profile for Tm0936 from *Thermotoga maritima* (7).

The large number of reactions catalyzed by members of the AHS suggests that the active site structure perched at the C-terminal end of the β -barrel is amenable to the efficient evolution of new enzymes with modified substrate profiles. One interesting example is that of the bacterial phosphotriesterase (10). The PTE from Pseudomonas diminuta has been shown to catalyze the hydrolysis of organophosphate triesters such as the insecticide paraoxon with values of k_{cat}/K_m that approach the diffusion limit of $10^8 \text{ M}^{-1} \text{ s}^{-1}$ (11). The high turnover rates are unusual since organophosphate triesters are not natural products and these compounds have been introduced into the environment only since the middle of the last century. The PTE from P. diminuta has been shown to possess a weak ability to hydrolyze cyclic carboxylate esters (12). It has been suggested that the present-day PTE has evolved from an enzyme whose physiological role was the hydrolysis of lactones (12).

In this paper we have determined the structural and catalytic properties of Dr0930 from *Deinococcus radiodurans*. This enzyme has a sequence identity to the PTE from *P. diminuta* of approximately 30%. The enzyme has been crystallized and the structure determined to a resolution of 2.1 Å. Computational docking of high-energy intermediates and substrate screening have been used to demonstrate that this enzyme efficiently catalyzes the hydrolysis of γ - and δ -lactones with values of k_{cat}/K_m that exceed 10⁶ M⁻¹ s⁻¹. The phosphotriesterase activity is very weak, but this promiscuous activity can be enhanced by over 1 order of magnitude with a relatively small number of changes to active site residues.

MATERIALS AND METHODS

Materials. All chemicals were obtained from Sigma or Aldrich. The genomic DNA from *D. radiodurans* R1 was purchased from ATCC. *Escherichia coli* BL21(DE3) and XL1-blue cells were obtained from Stratagene. The expression vector pET-30a(+) and *platinum pfx* DNA polymerase were purchased from Invitrogen. The genomic DNA purification kit was obtained from Promega. Oligonucleotide synthesis and DNA sequencing were performed by the Gene Technology Laboratory of Texas A&M University.

Cloning of Gene for Dr0930. The gene for Dr0930 from *D. radiodurans* was amplified from the genomic DNA using the primer pair 5'-AGAACTTC<u>CATATG</u>ACGGCACA-GACGGTGACGGGCG-3' (forward) and 5'-ACG<u>GAAT-TCTTACCCGAACAGCCGGGCCGGGGTTG-3'</u> (reverse). The forward primer contained an *NdeI* restriction site, and the reverse primer introduced an *Eco*RI restriction site. The

PCR product was gel purified, digested with *NdeI* and *Eco*RI, and cloned into the expression vector pET-30a(+) at the *NdeI* and *Eco*RI sites. The fidelity of the insert was verified by direct DNA sequencing. The mutants Y28F, Y97F, and Y98F were constructed using the QuikChange mutagenesis kit from Stratagene. The mutants F26G, C72I, F26G/C72I, Y98F, F26G/C72I/Y97W/Y98F, G207D/R228H/L231H, and F26G/C72I/Y97W/Y98F/G207D/R228H/L231H were constructed using the method of overlap extension (*13*). The coding regions of all expression plasmids were completely sequenced.

Protein Purification. The plasmid containing the gene for Dr0930 was transformed into BL21(DE3) electrocompetent cells. A single colony was used to inoculate a 5 mL overnight culture of LB medium supplemented with 50 µg/mL kanamycin. The 5 mL overnight culture was then used to inoculate 1.0 L of LB medium with 50 µg/mL kanamycin. Cells were grown at 30 °C, and gene expression was induced by addition of 0.5 mM isopropyl D-thiogalactopyranoside (IPTG) and 1.0 mM metal salt (ZnAc₂, MnCl₂, or CoCl₂) when the OD_{600} of the culture reached 0.6. When $MnCl_2$ or CoCl₂ was added to the growth medium, 0.1 mM 2',2bipyridal was added to chelate the iron in the growth medium as soon as the OD_{600} of the culture was approximately 0.1. The cells were harvested by centrifugation 18 h after addition of IPTG. About 10 g of cells was obtained from 3 L of culture. For purification, 10 g of cells was resuspended in 50 mL of 50 mM Hepes, pH 7.5, and lysed by sonication (5 s pulses for 30 min at 0 °C) in an ice bath. After centrifugation, the nucleic acids were removed by adding 2% (w/v) protamine sulfate dropwise, and then the protein was precipitated from the supernatant solution by slowly adding ground ammonium sulfate until 60% saturation. The precipitated protein from the ammonium sulfate step was resuspended in a minimum quantity of 50 mM Hepes buffer, pH 7.5, filtered with a 0.2 μ m membrane, and loaded onto a Superdex 200 gel filtration column (Amersham Pharmacia). The fractions containing the recombinant protein were pooled based on the results from PhastGel electrophoresis. The protein solution was loaded onto a Resource Q anionexchange column (Amersham Pharmacia) and eluted with a linear gradient of NaCl in 20 mM Hepes, pH 7.5. Dr0930 eluted from the column at approximately 150 mM NaCl. The purity of the protein was verified by SDS-PAGE.

Metal Analysis. The protein concentration was determined spectrophotometrically at 280 nm using a 1 cm quartz cuvette in a SPECTRAmax-340 UV—vis or CARY 100 spectrophotometer. The extinction coefficient at 280 nm used for calculating the concentration of Dr0930 was $28710 \text{ M}^{-1} \text{ cm}^{-1}$ (*14*). The metal content of the purified protein was determined by inductively coupled plasma emission mass spectrometry (ICP-MS). Before ICP-MS measurements, the protein samples were treated with concentrated nitric acid overnight and then diluted with distilled H₂O until the final concentration of nitric acid was 1%.

Analytical Ultracentrifugation. Protein samples for analytical ultracentrifugation were prepared in 50 mM Tris, pH 7.5. The experiments were performed using a Beckman model Optima XL-A analytical ultracentrifuge equipped with an AN60 rotor. Sedimentation velocity experiments used a two-channel aluminum centerpiece containing a 400 μ L reference and a 380 μ L sample. Preliminary sedimentation velocity experiments were carried out at six different concentrations:

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26.1, 13.1, 6.5, 2.6, 1.3, and 0.64 μ M. The samples were subjected to ultracentrifugation at 40000 rpm at 25 °C. Absorbance data were collected at 280 nm for the first three concentrations and at 230 nm for the three lower concentrations. Approximately 13 scans were collected and analyzed using Ultrascan 9.9 version for Windows (15, 16). Under these conditions, there was no dependence of sedimentation as a function of protein concentration. To determine the molecular weight of the protein, another sedimentation velocity experiment was carried out at a speed of 45000 rpm at 20 °C. Absorbance data were collected using a radial step size of 0.003 cm at 280 nm at three different concentrations: 23.4, 15.1, and 6.7 µM. Approximately 70 scans for each cell were collected and used for analysis. The sedimentation coefficient was determined to be 4.7 ± 0.1 S using the van Holde-Weischet analysis from Ultrascan. The average molecular mass was calculated to be 69 ± 5 kDa using 2-D spectrum analysis from Ultrascan (17, 18). This molecular mass corresponds to a homodimer since the molecular mass of a single subunit is 34.7 kDa.

Kinetic Measurements and Data Analysis. The kinetic measurements were performed with a SPECTRAmax-340 plate reader or a CARY 100 spectrophotometer. The hydrolysis of lactones was monitored using a pH-sensitive colorimetric assay (19). Protons released from the carboxylate product were measured using the pH indicator cresol purple. The reactions were performed in 2.5 mM Bicine, pH 8.3, containing 0.2 M NaCl, 0.01-6 mM lactone substrate, 0.1 mM cresol purple, and various amounts of Dr0930. The final concentration of DMSO was 1.4%. The change in absorbance at 577 nm was monitored in a 96-well UV-visible plate. The conversion factor (Δ OD/mol of H⁺) was obtained from an acetic acid titration curve at 577 nm ($\epsilon = 1.88 \times 10^3$ M^{-1} , 1.4% DMSO). The protein was stored in 50 mM Hepes, pH 7.5, and exchanged with 10 mM Bicine, pH 8.3, with a PD-10 column before use. The lactones were dissolved in DMSO and then diluted to 2.5 mM Bicine, pH 8.3. A background rate was observed in the absence of substrate due to acidification by atmospheric CO₂. The background rate (5-10 mOD/min) was independent of substrate concentration and subtracted from the initial rates. Measurement of the phosphotriesterase activity was conducted in 50 mM Ches, pH 9.0, using paraoxon as the substrate. The reaction was monitored spectrophotometrically by following the formation of *p*-nitrophenol at 400 nm.

Kinetic parameters were obtained by fitting the initial rates directly to eq 1, where ν is the initial velocity, E_t is the enzyme concentration, k_{cat} is the turnover number, [A] is the substrate concentration, and K_m is the Michaelis constant.

$$\nu/E_{\rm t} = k_{\rm cat}[{\rm A}]/(K_{\rm m} + [{\rm A}])$$
 (1)

Crystallization and Data Collection. Crystals of Zn²⁺bound Dr0930 were grown by vapor diffusion at room temperature. The protein solution contained Dr0930 (11.2 mg/mL) in 20 mM Hepes, pH 7.5, and 0.5 mM ZnCl₂; the precipitant contained 1.2 M (NH₄)₂SO₄, 0.1 M Tris, pH 8.5, and 0.2 M Li₂SO₄. Crystals appeared in 4 days and exhibited diffraction consistent with the space group $P3_121$, with one molecule of Dr0930 per asymmetric unit. These crystals contain 60% solvent and are characterized by a V_m of 3.1 Å³/Da, both of which are on the high side of the distribution exhibited by protein crystals. Prior to data collection, the

Table 1:	Data	Collection	and	Refinement	Statistics
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	native Dr0930
Data Collection	
beamline wavelength (Å) spacegroup no. of molecules in au	NSLS X4A 0.97915 <i>P</i> 3 ₁ 21 1
Unit Cell Paramete	rs
	61.505 205.94 25-2.1 (2.18-2.1) 26992 98.7 (98.2) 0.041 (0.103) 45.4 (33.1)
Refinement	
resolution R_{cryst} R_{free} rmsd for bonds (Å) rmsd for angles (deg) no. of protein atoms no. of waters no. of ions PDB entry	25.0-2.1 0.232 (0.315) 0.277 (0.352) 0.005 1.4 2440 384 2 Zn ²⁺ 3FDK

 $^{\it a}$ Numbers in parentheses indicate values for the highest resolution shell.

crystals were transferred to cryoprotectant solutions composed of their mother liquors supplemented with 20% glycerol. After incubation for ~15 s, the crystals were flashcooled to 100 K in a nitrogen stream. Data for the Zn^{2+} bound Dr0930 were collected to a resolution of 2.1 Å at the NSLS X4A beamline (Brookhaven National Laboratory) on an ADSC CCD detector. Intensities were integrated and scaled with DENZO and SCALEPACK (20). Data collection statistics are given in Table 1.

Structure Determination and Refinement. The structure of Zn^{2+} -bound Dr0930 was determined by molecular replacement with PHASER (21), using the coordinates of the phosphotriesterase from *P. diminuta* (PDB file 1HZY) as the search model (30.8% sequence identity). Iterative cycles of automatic model rebuilding with ARP (22), manual rebuilding with TOM (23), and refinement with CNS (24) were performed. The model was refined at 2.1 Å with an R_{cryst} of 0.232 and an R_{free} of 0.277. The final structure contained protein residues 2–323, 384 water molecules, and 2 well-defined Zn^{2+} ions. Representative electron density of the active site of Dr0930 liganded with two Zn^{2+} ions is shown in Figure 1. Final crystallographic refinement statistics are provided in Table 1.

Docking of High-Energy Intermediates. In parallel with the experimental elucidation of the catalytic function of Dr0930, docking calculations were carried out in order to identify potential substrates. These calculations employed the Kyoto Encyclopedia of Genes and Genomes (KEGG) LIGAND library of biogenic compounds (25, 26), a database of small molecules that are found in nature. This database consists of diverse molecules, often with unrelated chemical functionalities; it was not tailored to Dr0930 except to the extent that only molecules bearing functionalities known to be acted on by members of the amidohydrolase superfamily were considered, but this still included about 40% of all the molecules in KEGG. In the version of KEGG used here, the molecules had been modified to resemble the tetrahedral or



FIGURE 1: Representative electron density for the active site of Dr0930 contoured at 1.5σ . Electron density shown was calculated to 2.1 Å resolution with the coefficients $(2F_o - F_c)$. The metal ions are depicted as blue spheres, and the bridging hydroxide is shown as a green sphere.

trigonal bipyramidal intermediates that are observed along the reaction coordinate of the enzymatic reactions that the AHS is known to catalyze. It has been shown previously that the high-energy intermediate (HEI) states of potential substrates are preferentially recognized by members of the AHS and that docking calculations utilizing them are more likely to yield the correct substrate (7, 9). The library of HEIs specific to reactions of the amidohydrolase superfamily was used (7, 9). Briefly, all molecules of the KEGG library that contained one or more of the 19 functionalities that are involved in any of the known amidohydrolase reactions were converted to their respective high-energy forms. As most of the AHS reactions are nucleophilic substitutions involving an attack by a hydroxide leading to a tetrahedral intermediate state, all possible stereoisomers corresponding to attacks from the re and si faces were generated. In addition, the protonated and unprotonated forms of the leaving groups were prepared. This protocol led to approximately 22500 different highenergy forms of the 4207 molecules in KEGG that featured a reactive group recognized by an AHS enzyme. Library generation and docking were done in an unbiased and general way, as we did not know what the substrates for the enzyme might be. We note that the molecule ultimately found to have the highest turnover rate, 11, was not part of the original KEGG library, and it was thus individually converted to a HEI, applying the same settings as for the entire library, so that we could model its enzyme-bound configuration by docking.

Docking was carried out with DOCK 3.5.54 as described previously (9), rank-ordering the molecules by a physicsbased score consisting of van der Waals, electrostatic, and solvation terms. After docking, poses were filtered based on a distance criterion to ensure that the reactive group was within 3.5 Å of the catalytic nucleophile and thus in a potentially catalytically competent geometry. The top 500 poses according to the score were inspected visually to ascertain this.

Construction, Expression, and Purification of PTE-Dr0930 Chimera. The binuclear metal centers in Dr0930 from D. radiodurans and PTE from P. diminuta are essentially identical. To identify the structural determinants for the differences in substrate specificities between these two enzymes, a structure-based sequence alignment was performed using DaliLite. The structure-based sequence alignment was used to identify the precise locations of the eight specificity loops at the C-terminal ends of the central β -strands in each protein. A chimeric protein was designed that substituted the eight specificity loops in Dr0930 with the corresponding loops from PTE. The gene for the chimeric Dr0930-PTE was chemically synthesized by Mr. Gene (http:// mrgene.com) and optimized for expression in E. coli with respect to codon usage and overall GC content. An amount of 5 μ g of lyophilized plasmid DNA was supplied by the manufacturer with the synthesized fragment inserted between KpnI and SacI restriction sites in the vector pGA18 with ampicillin resistance. The KpnI restriction site at the 5'-end of the fragment is followed by an EcoRI and a NdeI restriction site which comprises the ATG codon of the gene. At the 3'-end of the gene, two stop codons were inserted followed by *HindIII* and *SacI* restriction sites. The designed gene carried internal restriction sites embedding loops 7 (restriction sites NcoI and AclI) and 8 (restriction sites TspRI and AatII).

The PTE-Dr0930 chimeric gene was inserted into pET-21a(+) and pET-28a(+) vectors, using the restriction sites NdeI and HindIII. To analyze the solubility of the protein, different expression cells (T7 Express, T7 Express Rosetta, BL21(DE3) RIPL, ArcticExpress, and ArcticExpress RIL) were transformed with these vectors carrying the synthetic gene. The chimeric protein was expressed in an analytical scale with 1.0 mM ZnCl₂ in the culture medium overnight at different temperatures (13, 20, 30, and 37 °C). All conditions yielded only insoluble protein. It was possible to purify the chimeric protein from the insoluble fraction of the T7 Express cells (transformed with pET-21a(+) containing the synthetic gene and grown in 250 mL of ZnCl₂-lacking LB medium) by refolding it in the presence of ZnCl₂. To solubilize the protein, the pellet was washed with 10 mM phosphate, pH 7.5, then dissolved in 20 mL of 6 M guanidinium chloride (GdmCl) and 10 mM phosphate, pH 7.5, and incubated for 30 min at room temperature with continuous stirring. Subsequently, 20 mL of 1 M GdmCl and 10 mM phosphate, pH 7.5, was added, and the solution was incubated for another 30 min at room temperature with continuous stirring. After centrifugation at 13000 rpm, the supernatant solution was transferred to a new reaction vessel and mixed with 70 mL of 2 M GdmCl and 10 mM phosphate, pH 7.5. This solution was dialyzed three times against 5 L of 50 mM phosphate, pH 7.5, 50 mM bicarbonate, and 100 µM ZnCl₂.

RESULTS

Cloning, Expression, and Purification of Dr0930. The gene for Dr0930 from D. radiodurans was successfully cloned and inserted into the vector pET-30a(+) for expression in E. coli. The protein expresses well in E. coli after induction with IPTG in the presence of divalent metal ions (ZnAc₂, MnCl₂, or CoCl₂) in the culture medium. A total yield of 30-125 mg of homogeneous protein was obtained from 1-3L of LB medium. The protein purity was estimated to be greater than 95% as judged by PhastGel electrophoresis. N-Terminal amino acid sequence analysis for the first six amino acid residues gave the sequence as MTAQTV, which



FIGURE 2: Visible absorption spectrum for Dr0930 at a subunit concentration of 450 μ M.

is consistent with the first six amino acid residues as deduced from the DNA sequence. The enzyme expressed in the presence of 1.0 mM ZnAc₂ in the growth medium was found to contain 1.2 equiv of zinc and 0.4 equiv of iron per subunit. The concentrated enzyme was intensely purple in color. Shown in Figure 2 is the visible absorption spectrum of the protein at a concentration of 450 μ M. The absorption maximum is at 550 nm with a molar extinction coefficient of 455 M⁻¹ cm⁻¹.

Structure of Dr0930. Dr0930 adopts an $(\beta/\alpha)_8$ structural fold. The protein monomer is composed of a $(\beta/\alpha)_7\beta$ -barrel with N-terminal and C-terminal extensions of the chain from both sides of the barrel. The following chain segments are included in the eight β -strands of the barrel: $\beta 1$ (residues 17-26), $\beta 2$ (61-66), $\beta 3$ (89-94), $\beta 4$ (141-146), $\beta 5$ (173-177), $\beta 6$ (199-202), $\beta 7$ (223-226), and $\beta 8$ (259-269). The N-terminal extension of the chain includes a β -loop (3-10). The C-terminal extension includes three distorted α -helices (278-284, 291-303, and 307-322). The last α -helix closes the entrance to the barrel from the N-terminal side. Two long segments between strands $\beta 1$ and $\beta 2$ and between strands $\beta 3$ and $\beta 4$ of the barrel contain two helices each. The remaining segments between the β -strands of the barrel each contain a single helix.

The active site is located at the C-terminal end of the barrel and is open to bulk solvent. Two metal ions are bound in the active site; the most deeply buried and most solvent exposed cations are denoted as α and β , respectively. The α -cation is coordinated by His-21 and His-23 from strand β 1, Asp-264 from strand β 8, and the carboxylated Lys-143 from strand β 4 (Figure 3). The β -cation is coordinated by His-176 from strand β 5, His-204 from strand β 6, and the carboxylated Lys-143. A bridging hydroxide ion serves as an additional ligand for both cations. The interatomic distance between the two metal ions is 3.4 Å, with distances to the bridging hydroxide of 1.9 and 2.1 Å. No other water molecules at distances less than 3.8 Å from both Zn²⁺ ions or from the bridging hydroxide were detected.

The Dr0930 monomers are related by a crystallographic 2-fold axis and form an apparent dimer in the current crystal structure (Figure 4), which buries a total surface area of 2835 Å² at the interface. Loop 1 (after strand β 1) and loop 3 (after strand β 3) of each molecule make extensive contacts with the 2-fold-related molecule. For example, Thr-36 OG1 (loop



FIGURE 3: Model for the active site of Dr0930. The α -metal ion (left) and the β -metal ion (right) are shown as yellow spheres, and the bridging hydroxide is shown as a red sphere.



FIGURE 4: Ribbon representation of the Dr0930 dimer. The molecules are oriented with the 2-fold axis perpendicular to the plane of the page. The metal ions at each active site are shown as red spheres. There are extensive intermolecular interactions formed by loops L1 and L3 from each monomer.

1) forms hydrogen bonds to Ala-133* N and Glu-127* OE2 of the adjacent molecule. Thr-105, Phe-109, and Leu-113 (loop 3) participate in various hydrophobic interactions with the residues of the adjacent molecule Pro-29*, Tyr-28*, Trp-269*, and Pro-274*, some of which are not far from the entrance to the active site of the symmetry mate. There are also some polar interactions between residues in loop 3 and the residues of the symmetry-related molecule: Thr-105 OG1-Glu-101* O, Tyr-106 OH-Asp 34* OD2, Tyr-106

OH-Arg 272* N, Arg 110 NH1-Pro 273* O, and Arg 110 NH1-Arg 272* O.

The phenolic oxygens of Tyr-28, Tyr-97, and Tyr-98 are 8.93, 3.48, and 8.15 Å away, respectively, from the β -cation. To determine if the absorbance at 550 nm of the wild-type enzyme is due to a charge transfer complex between any of these tyrosines and the β -cation, they were individually mutated to phenylalanine. The purified Y28F mutant contained 1.1 equiv of Zn and 0.4 equiv of iron, and the protein absorbed at 550 nm with a molar extinction coefficient of 490 M⁻¹ cm⁻¹. The Y97F mutant contained 1.4 equiv of zinc and 0.3 equiv of iron, but no absorbance could be detected at 550 nm at concentrations up to 5 mg/mL. The Y98F mutant adsorbed at 550 nm with an extinction coefficient of 460 M⁻¹ cm⁻¹. This mutant contained 0.9 equiv of zinc and 0.2 equiv of iron. Enzyme expressed in the presence of the iron chelator 2',2-bipyridal and 1.0 mM Mn²⁺ contained less than 0.1 equiv of iron and was transparent in the visible spectrum (data not shown).

Docking Calculations. The initial attempt to dock the highenergy intermediate (HEI) version of the KEGG database to the X-ray structure of Dr0930 failed as few molecules were docked in a pose that was even remotely close to the metal ions, a necessary condition for successful turnover. Upon inspection of the binding site it became clear that the side chain of Arg-228 was blocking access to the catalytic center. Consequently, the side chain of this residue was rotated 180° around the $C_{\gamma}-C_{\delta}$ bond and minimized with CHARMM (27), using the CHARMm22 force field (28). This conformation is lower in energy compared to the original one, which is due, in part, to the decreased distance between Arg-228 and the side chain of Asp-206, resulting in a more favorable electrostatic interaction. In the second virtual screen of the KEGG HEI database, in which DOCK found poses that placed the reactive center, represented by the high-energy intermediate, within the distance of 3.5 Å for 1821 molecules, a substantial enrichment of lactones was observed. Among the top 100 docking-ranked KEGG metabolites, there were 55 high-energy intermediates that were docked in a catalytically competent manner, 21 of which were lactones (including three in the top ten). Consistent with this ranking, lactones were identified as preferred substrates of Dr0930 by experimental assay. Among the high-ranking lactones that were turned over were molecules ranking 173 (compound 6), 325 (compound 9), 349 (compound 1), and 417 (compound 13). After visual inspection of the 500 highest ranking molecules, lactones ranking 89 and 216 were also suggested based on their docking prediction and tested experimentally but were not hydrolyzed measurably by the enzyme. In addition to these lactones, nine other high-ranking molecules, including several lactams, were tested after they had been selected as potential substrates because of their docking ranks and predicted geometries, but none of them was turned over. Subsequently, the substrate with what turned out to be the highest turnover rate, 11, was docked into the structure of Dr0930, mostly to investigate its proposed structure and stereochemical preferences. This compound had a docking score that, had it been included in the initial KEGG-derived HEI database, would have ranked at 72 overall (see Materials and Methods). Compound 11 is a δ -lactone; the corresponding γ -lactone, 6, ranks at 173.



FIGURE 5: Computed binding mode of compound 11 (golden carbons). Metal ions are depicted in light blue. The dashed lines indicate the distances between the terminal methyl group of compound 11 and selected residues within the active site. Additional details are presented in the text.

To investigate stereochemical preferences, all possible enantiomer combinations of compounds **6** and **11** were compared directly. As the carbon atom at the base of the hydrophobic tail is chiral, this comparison included four different species: *re-R*, *re-S*, *si-R*, and *si-S*. For both molecules, docking vastly prefers an orientation that corresponds to a hydrophilic attack from the *si* face (Figure 5 depicts the highest ranking pose of **11**). Among the top 50 poses (i.e., orientations and conformations), 46 are such that they could be attacked by a hydroxyl anion positioned between the metal ions. Of these, 20 are *si-R*; 12 are *si-S*; 3 are *re-R*; and 11 are *re-S*.

We also looked at the predictions for the *N*-acylhomoserine lactones in detail. It has been demonstrated that this class of molecules is turned over by Sso2522, and it was an open question whether the same is true for Dr0930. Although there were 7 *N*-acylhomoserine lactones in the docked library, none of them was placed in a catalytically competent pose. The docking of the organophosphate triesters was also investigated. In total, six such molecules can be found in the database, but only one docked in a sensible pose. This molecule had a favorable score, placing it at rank 5. However, we were unable to demonstrate that this compound, 2,2-chlorovinyl dimethyl phosphate (dichlorvos) is a substrate for Dr0930. The value of k_{cat} is less than $4 \times 10^{-3} \text{ s}^{-1}$.

Substrate Specificity and Kinetic Measurements. The catalytic activity of Dr0930 was initially examined for the hydrolysis of organophosphate triesters, diesters, and monoesters. No catalytic activity could be detected with ethyl 4-nitrophenyl phosphate or 4-nitrophenyl phosphate. Slow hydrolysis could be detected with the organophosphate triester, paraoxon (diethyl 4-nitrophenyl phosphate), and the values of k_{cat} , K_m , and k_{cat}/K_m were determined to be 0.072 min⁻¹, 1.4 mM, and 0.83 M⁻¹ s⁻¹, respectively. These results indicated that the native substrate profile of this enzyme does not involve phosphate esters. No activity could be detected with phenyl acetate, but significant hydrolytic activity could be detected with a number of lactones identified in the initial screening protocols. We measured the catalytic activity of Dr0930 with an array of lactones and found that this enzyme

Scheme 1



can efficiently hydrolyze γ - and δ -lactones. The structures of these compounds are given in Scheme 1. The kinetic parameters for the Zn-, Mn-, and Co-substituted forms of Dr0930 are presented in Table 2, and the kinetic constants for the three tyrosine to phenylalanine mutants are given in Table 3. The best substrates identified to date are γ - or δ -lactones with an alkyl substituent attached to the carbon adjacent to the ring oxygen. For Mn/Mn-Dr0930 the values of k_{cat} and k_{cat}/K_m for δ -decanoic lactone are 460 s⁻¹ and $1.3 \times 10^6 \text{ M}^{-1} \text{ s}^{-1}$, respectively. No activity could be detected with compounds **14–23** with turnover numbers greater than 0.01 s^{-1} . The structures for additional compounds (**S24–S37**) that were found to be inactive with the Zn- and Mnsubstituted forms of Dr0930 are presented in the Supporting Information as Figure S1.

Loop Transplantation from PTE to Dr0930. The binuclear metal center within the active site of Dr0930 is identical with the one found in the active site of the phosphotriesterase from P. diminuta. Shown in Figure 6 is a structural superposition between these two proteins. It is clear from this comparison that the central β -barrel and interconnecting α -helices superimpose quite nicely and that the major structural differences between these two enzymes lie predominantly within the composition and conformation of the eight $\beta \alpha$ loops that interconnect the eight β -strands and the eight α -helices. However, Dr0930 catalyzes the hydrolysis of paraoxon more than a million-fold more slowly than does PTE from *P. diminuta*. Since the substrate specificity of β/α barrel proteins is apparently dictated by the residues located at the C-terminal end of the β -strands and the $\beta\alpha$ loops, a chimeric protein was designed where the eight $\beta \alpha$ loops found close to the active site of PTE replace the corresponding loops in Dr0930. This was done to determine whether the inherent structural and dynamic information embedded within the eight loops in PTE is sufficient to increase the promiscuous triesterase activity in Dr0930. A sequence alignment based upon the structural alignment between PTE and Dr0930 is presented in Figure 7. Shown in red are the specific residues for the 8 $\beta\alpha$ loops of PTE that were swapped for the eight corresponding loops of Dr0930 (shown in blue).

The gene for the chimeric PTE-Dr0930 protein was synthesized, and attempts were made to express the protein under a variety of conditions. However, in every case the protein was found to be insoluble. The protein was solubilized by the addition of 6 M GdmCl, which was then removed stepwise by dialysis against 50 mM potassium phosphate, pH 7.5, containing 100 μ M ZnCl₂ and 50 mM bicarbonate, to induce folding. A total of 3.6 mg of the soluble chimeric protein was isolated that was judged 80% pure by SDS gel electrophoresis. However, no phosphotriesterase activity was detectable for the PTE-Dr0930 chimera ($k_{cat} < 10^{-4} \text{ s}^{-1}$).

Redesign of Dr0930 Active Site. The three-dimensional structure of PTE with a bound substrate analogue has been used to identify those residues that come together to form the substrate binding pocket in the active site (29). The 12 most significant residues include G60, I106, W131, F132, H254, H257, L271, L303, F306, S308, Y309, and M317. The corresponding residues within the active site of Dr0930 are F26, C72, Y97, Y98, R228, L231, V235, I266, W269, R272, P273, and P282, respectively. In addition, H254 in PTE is hydrogen bonded to D233, and the corresponding residue in Dr0930 is G207. In an attempt to enhance the promiscuous triesterase activity of Dr0930, some of these residues were replaced with the corresponding residues found within the active site of PTE. These substitutions were done singularly and in combination with one another. The mutant forms of Dr0930 were purified to homogeneity, and their activity toward the hydrolysis of paraoxon was measured at pH 9.0. The results are summarized in Table 4. The most active mutant was F26G/C72I with a k_{cat} of 1.14 min⁻¹ and $k_{\text{cat}}/K_{\text{m}}$ of 7.7 M⁻¹ s⁻¹. The turnover number of this mutant is enhanced 16-fold relative to the wild-type enzyme.

DISCUSSION

Purification and Properties. Dr0930 from D. radiodurans was cloned and purified to homogeneity. The purified protein was found to contain a mixture of zinc and iron when expressed in a medium that was supplemented with added zinc. At high concentrations the protein was a deep purple in color with an absorption maximum at 550 nm. This property is reminiscent of the optical properties for purple acid phosphatase or uteroferrin (30). Those proteins contain a binuclear iron center, and the visible spectrum has been proposed to be due to a charge transfer complex between a tyrosine and one of the two irons within the binuclear metal center (31). For uteroferrin the extinction coefficient for the Fe^{3+}/Fe^{2+} complex is approximately 4000 M⁻¹ cm⁻¹ (30). This is considerably higher than the value of 455 M^{-1} cm⁻¹ measured for Dr0930, but the Fe content is less than 1 equiv and the distribution of the iron between the two sites is not known. Nevertheless, the X-ray structure determination of Dr0930 showed that the phenolic oxygen of Tyr-97 is 3.5 Å away from the β -metal ion. Mutation of Tyr-97 to a phenylalanine resulted in a protein that was transparent in the visible region. In addition, expression of Dr0930 in a medium where the iron content was reduced with an ironspecific chelator in the presence of added Mn²⁺ resulted in

Table 2: Kinet	ic Parameters	for Zn/Zn-,	Mn/Mn-, an	nd Co/Co-Dr0930	at pH 8.3
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		Zn/Zn-Dr0930			Mn/Mn-Dr0930			Co/Co-Dr0930		
compd	dock rank	$k_{\rm cat}~({\rm s}^{-1})$	$K_{\rm m}~({\rm mM})$	$k_{\rm cat}/K_{\rm m}~({\rm M}^{-1}~{\rm s}^{-1})$	$k_{\rm cat}~({\rm s}^{-1})$	$K_{\rm m}~({\rm mM})$	$k_{\rm cat}/K_{\rm m} \ ({\rm M}^{-1} \ {\rm s}^{-1})$	$k_{\rm cat}~({\rm s}^{-1})$	$K_{\rm m}~({\rm mM})$	$k_{\rm cat}/K_{\rm m} \ ({\rm M}^{-1} \ {\rm s}^{-1})$
1	349	4.3 ± 0.3	4.6 ± 0.6	$(9.4 \pm 1.3) \times 10^2$						
(<i>R</i> / <i>S</i>)-2	nl^a	7.4 ± 0.2	1.9 ± 0.1	$(3.9 \pm 0.2) \times 10^3$	155 ± 5	8.4 ± 0.4	$(1.8 \pm 0.2) \times 10^4$			
(R)- 2	nl	8.1 ± 0.1	1.5 ± 0.1	$(5.5 \pm 0.2) \times 10^3$	163 ± 3	6.5 ± 0.2	$(2.5 \pm 0.1) \times 10^4$			
(R/S)- 3	nl	5.2 ± 0.1	0.75 ± 0.05	$(6.9 \pm 0.5) \times 10^3$	260 ± 9	3.4 ± 0.2	$(7.7 \pm 0.5) \times 10^4$			
(R/S)-4	172	5.0 ± 0.1	0.13 ± 0.01	$(3.8 \pm 0.3) \times 10^4$	220 ± 4	0.34 ± 0.02	$(6.4 \pm 0.4) \times 10^5$			
(R/S)- 5	nl	2.0 ± 0.04	0.063 ± 0.007	$(3.2 \pm 0.4) \times 10^4$	212 ± 12	0.57 ± 0.07	$(3.7 \pm 0.2) \times 10^5$			
(<i>R/S</i>)-6	173	5.8 ± 0.2	0.14 ± 0.02	$(4.1 \pm 0.1) \times 10^4$	95 ± 4	0.39 ± 0.06	$(2.5 \pm 0.2) \times 10^5$			
(S)- 6	173	14 ± 1	0.15 ± 0.02	$(9.3 \pm 1) \times 10^4$	230 ± 12	0.58 ± 0.11	$(4.0 \pm 0.3) \times 10^5$	70 ± 2	0.09 ± 0.1	$(8 \pm 1) \times 10^{5}$
(<i>R/S</i>)-7	nl	2.1 ± 0.1	0.14 ± 0.02	$(1.5 \pm 0.2) \times 10^4$	100 ± 4	0.34 ± 0.04	$(2.9 \pm 0.4) \times 10^5$			
(S)- 8	nl	1.9 ± 0.1	0.35 ± 0.06	$(5.4 \pm 0.9) \times 10^3$	87 ± 2	0.65 ± 0.05	$(1.3 \pm 0.1) \times 10^5$	17 ± 1	0.70 ± 0.06	$(2.4 \pm 0.2) \times 10^4$
9	325	48 ± 3	3.0 ± 0.3	$(1.6 \pm 0.2) \times 10^4$	93 ± 3	3.3 ± 0.2	$(2.8 \pm 0.2) \times 10^4$			
(<i>R/S</i>)-10	nl	32 ± 1	1.8 ± 0.1	$(1.8 \pm 0.1) \times 10^4$	88 ± 7	2.3 ± 0.4	$(3.8 \pm 0.7) \times 10^4$			
(<i>R/S</i>)-11	72	42 ± 1	0.10 ± 0.02	$(4.2 \pm 0.7) \times 10^5$	203 ± 5	0.13 ± 0.01	$(1.6 \pm 0.1) \times 10^{6}$	180 ± 5	0.09 ± 0.01	$(1.8 \pm 0.2) \times 10^{6}$
(<i>R/S</i>)-12	nl	16 ± 1	0.08 ± 0.01	$(2.0 \pm 0.2) \times 10^5$	460 ± 12	0.37 ± 0.04	$(1.3 \pm 0.1) \times 10^{6}$	180 ± 7	0.19 ± 0.03	$(9.5 \pm 1.6) \times 10^5$
13	417	ND	ND	59 ± 4	2.4 ± 0.3	3.5 ± 0.8	$(6.9 \pm 1.7) \times 10^2$			
^a Not i	n original K	EGG library	/.							

Table 3: Properties of Tyrosine to Phenylalanine Mutants of Dr0930^a

		metal content				
protein	color	zinc	iron	$k_{\rm cat} ({\rm s}^{-1})$	$K_{\rm m}~({\rm mM})$	$k_{\rm cat}/K_{\rm m}~({\rm M}^{-1}~{\rm s}^{-1})$
wild type	purple	1.2	0.4	42 ± 1	0.10 ± 0.02	$(4.2 \pm 0.7) \times 10^{4}$
Y28F	purple	1.1	0.4	26 ± 1	0.10 ± 0.02	$(2.7 \pm 0.4) \times 10$
Y97F	colorless	1.4	0.3	0.33 ± 0.01	0.20 ± 0.02	$(1.7 \pm 0.1) \times 10$
Y98F	purple	0.9	0.2	7.0 ± 0.1	0.40 ± 0.02	$(1.8 \pm 0.2) \times 10$

^{*a*} Hydrolysis of compound **11** at pH 8.3.



FIGURE 6: Structural superposition between Dr0930 and PTE. The eight $\beta \alpha$ loops from PTE are depicted in red whereas the eight $\beta \alpha$ loops from Dr0930 are depicted in blue. The central β -barrel, the surrounding α -helices, and the two zinc atoms for both proteins are depicted in shades of gray.

protein that was transparent at 550 nm. Therefore, the intense purple color of the wild-type Dr0930 is consistent with a charge transfer complex between Tyr-97 and iron bound to the β -metal position within the binuclear metal center. To the best of our knowledge the only other member of the AHS that absorbs in the visible region of the spectrum is the H55C mutant of Co/Co-PTE, which is bright green in color (32).

X-ray Structure Determination. The most similar protein structure to Dr0930 is Sso2522 from *Sulfolobus solfataricus*

(PDB code 2VC5) with a DALI (33) score of 42.5 and 28% overall sequence identity. The next most similar structural homologues for Dr0930 are the bacterial phosphotriesterases from P. diminuta (PDB code 1HZY) and Agrobacterium radiobacter (PDB code 2R1N) and the PTE homology protein from E. coli (PDB code 1BF6), with DALI scores of \sim 41.0. The structures of Dr0930 and Sso2522 can be superimposed with an rmsd of 1.61 Å for 298 Ca pairs. Superposition of Dr0930 with PTE from P. diminuta (PDB code 1HZY) results in an rmsd of 1.32 Å for 276 Cα pairs, and superposition with 2R1N results in an rmsd of 1.46 Å for 283 Ca pairs. The superposition of Dr0930 with the PTE from P. diminuta is shown in Figure 6. The major differences involve the length and/or the conformation of the $\beta\alpha$ loops. However, the two metal atoms, the bridging hydroxide, and the direct metal ligands (His-21, His-23, His-176, His-204, Asp-264, and the carboxylated Lys-143 in Dr0930) are conserved between the two proteins and superimpose well. However, the terminal water molecule bound to the most solvent exposed β -metal ion of PTE is absent in Dr0930.

The subunit organization of the Dr0930 dimer is very similar to that found in the crystal structures of the phosphotriesterases from *P. diminuta* (34) and *A. radiobacter* (35) and the lactonase Sso2522 from *S. solfataricus* (36). The dimer interfaces are all formed by interactions between the protruding L1 and L3 loops of each molecule and the protein chain of the adjacent molecule. These dimers are distinguished by slight differences in the relative orientation of their monomers. For example, the dimers of Dr0930 and *P. diminuta* phosphotriesterase (PDB code 1HZY) superimpose with an rmsd of 2.5 Å for 568 C α pairs.

Docking Calculations. Three lessons emerge from the *in* silico substrate predictions for this enzyme. First, docking of high-energy intermediates enriched the lactone functional-

		HZI	H23 100p	T			
Dr0930	TAQTVTGAV	/AAAQLGATLP H F	HVIFGYPGY.AG	DVTLGPFD	HAAALASCT	ETARALLARGI	QTVVD <mark>AT</mark>
PTE	.DRINTVRGPI	TISEAGFTLT H F	HICGSSAGF1RA	WP EFFGS	<mark>SRK</mark> ALAEKAV	RGLRRARAAGV	RTIVD <mark>VS</mark>
SsoPox	MRIPLVGKdSI	ESKDIGFTLI H E	HLRVFSEAVRQQ	WPHLY.N	JEDEEFRNAV.	NEVKRAMQFGV	KTIVDPT
PHP		FDPT.GYTLA H E	HLHIDLSGFK	.NNVD.C	CrldOYAFIC	OEMNDLMTRGV	RNVIEMT
PPH	VPELNTARGPI	DTADLGVTLM H F	HVFIMTTEIAON	YPEAWGI	DEDKRVAGAL	ARLGELKARGV	DTIVDLT
AhlA	MSSVOTVRGPV	/dtadlgkvlm h f	HVFVLGEELRON	YPDYPEPWI	DEEVRVADAV	AKLTEVKSRGI	STIVDPT
	~		~				
	100p 2		¥97	100p 3			
Dr0930	PNDCGRNPAFI	REVSEATGLOII	CATGFYYEGEGA	TTYFKFRAS	SLGD AESE	IYEMMRTEVTE	GIAGTGI
PTE	TEDIGROVSLI	AEVSRAADVHTV	AATGLWE DP	PLSMRLR.	SVEE	LTOFFLRETOY	GIEDTGI
SsoPox	VMGLGRDTRFM	EKVVKATGINLV	AGTGTYTYT, DL	PFYFLNR	STDE	TADLETHDIKE	GTOGTLN
рнр	NRYMGRNAOFM	ILDVMRETGINVV	ACTGYYODA FF	PEHVATR	SVOE	LAOEMVDETEO	GIDGTEL
 ррн	VIGLORVIPRI	ARVAAATELNIV	VATGLYTYN DV	PEVENYLG	CAOLDGPET	MTDMFVRDIEH	GTADTGT
 Ahla	VIGLGRYIPRI	VRINEOVDLNIT	VATGIYTYN DI	PFOFHYTGI	CLIFDGPEP	MVELEVKDIRE	GTAGTGV
	VIGLORITIKI						01110101
	K143	100p 4		H176 la	000 5		H204
Dr0930	RAGVIKLASS	Rdat TPYEOLFF	RAAARVORETGV	PTTT HTOE	LOOGPOOAE	LUTSUGADPAR	TMTGHMD
PTE	RAGITKVATT	G KATPFOELVI	KAAARASLATGV	PVTT H TAAS	SORDGE OOAA	TFESEGUSPSR	VCTGHSD
Scopor	KAGEVKTAAD	EPGITKDVEKVI	RAAATANKETKV	PTTTHSNA	INNTGLEOOR	TLTEEGVDPGK	TLIGHLG
DHD	KAGTTARIGTS	EGKITPLEEKVE	TAAALAHNOTGR	PISTHTSES	TMGLEOLA	LTOPHGADI CK	VTVGHCD
DDU	KAGILKCATDE	DC LTDCVFDVI		DIGT H THA(TENEECVDL.QD	WYTCHCC
7511 7617	DAGELECATES					VIAFECADUTE	WICHEG
AIIIA	KASPURCATE	IEG-DIEGVERVE			II BOULVAOK	VDAGEOADDIK	V V 101100
	1000 6		loon	7			D264
Dr0930	CNTDPA YHRET	U.RHGUSTAFDRT	GLOG	MUGT P	DAERLSVIT	TLLGEGYADRI.	LLSHDST
PTE	DTDDLSYLTAI	AARGYLTGLDHT	PHSATGLEDNAS	ADALLGIRS	WOTRALLIK	ALTDOGYMKOT	
SsoPox	DTDNTDYTKKI	ADKGSFTGLDRY	GLD	LFL	VDKRNETTI.	RLIKDGYSDKT	MISHDYC
рнр	I'KDNI'DNII'KM	ITDLGAYVOFDTT	GKN	SYYI	PDEKRTAMLH	ALRDRGLLNRV	MLSMDIT
 РРН	DSTDVGYLEEI	TAAGSYLGMDRE	GVD	VISI	PFODRVNTVA	RMCERGHADKM	VLSHDAC
Ahla	DTTDLDYLCAI	ADAGSLLGMDRE	GLE	PFLS	SFEDRVNTVV	EMARRGYAEKM	VLAHDAS
	511515110iii						
		100p 8					
Dr0930	WHWLGRPPAIE	PEAALPAVKD	.WHPLHISDDIL	PDLRRRGI	reeovgomtv	GNPARLEG	
PTE	FGFS.SYVTN.	IMDVMDRVN	IPDGMAFIPLRVI	PFLREKGVI	POETLAGITV	TNPARFLSPTL	RAS
SsoPox	CTTD. WGTA.	KPEYKPK, LAPF	W.SITLIFEDTI	PFLKRNGVN	JEEVIATIFK	ENPKKFFS	
PHP		.RRSH1kANGGY	GYDYLLTTFI	POLROSGES	SOADVDVMLR	ENPSOFFO	
PPH	CYFDA LF	EELVPVAMPNW	. HYLHIHNDVT	PALKOHGV	DEOLHTMI V	DNPRRIFEROG	GYO
Ahla	CFIDY	FPSAAREVALE	NWNYTHTSDDVT	PALLERGY	CEDOAKAWI'A	DNPRRYFES	x
min	·· · · · · · · · · · · · · · · · · · ·			. • • • • • • • • • • • • • • • • • • •			

FIGURE 7: Structure-based sequence alignment between PTE (gil54778533) and Dr0930 (gil15805954). The specific amino acids of PTE that replaced those of Dr0930 in the PTE-Dr0930 chimera are shown in red and blue, respectively. Identical residues between PTE and Dr0930 are shown in light gray. Residues involved in metal complexation are printed in bold and labeled. The relationships of these sequences to those found in Sso2522 (also known as SsoPox; gil15899258), PTE homology protein (PHP; gil16131257), putative parathion hydrolase from *M. tuberculosis* (PPH; gil15839610), and *N*-acylhomoserine lactone acylase from *R. erythoropolis* (AhlA; gil46947344) are shown for comparison.

		metal content				
protein	color	zinc	iron	k_{cat} (\min^{-1})	K _m (mM)	$k_{\text{cat}}/K_{\text{m}}$ (M ⁻¹ s ⁻¹)
wild type	purple	0.71	0.52	0.072	1.4	0.83
F26G	black	0.85	0.34	0.55	8.3	1.1
C72I	purple	0.61	0.87	0.023	1.4	0.29
F26G/C72I	purple	0.64	0.81	1.14	2.7	7.7
Y98F	purple	1.17	0.38	0.06	1.8	0.55
F26G/C72I/Y97W/Y98F	yellow	0.41	0.66			0.91 ^b
G207D/R228H/L231H	purple	1.29	0.47	nd ^c	nd	nd
F26G/C72I/Y97W/Y98F/ G207D/R228H/L231H	yellow	0.62	0.27	nd	nd	nd

up to a concentration of paraoxon of 3 mM. c nd = not detectable.

ity that was ultimately found to be recognized by Dr0930. Second, as compelling as this enrichment was, it was insufficient in itself to identify those lactones best recognized by the enzyme. This has two origins. The very best ranking lactones by docking that we were actually able to obtain and test (e.g., those ranked 30 and 148) were not hydrolyzed by the enzyme; these are false positives, as are the high-ranking lactams and the organophosphate triester that were tested but were not substrates. Dr0930 represents an enzyme where substrate recognition goes beyond physical complementarity of the enzyme for an HEI and where the cost of forming the HEI from the ground state also plays an important role; this energy difference between ground and excited states presumably distinguishes many of the lactams from the lactones and many of the stabilized lactones from those aliphatic δ -lactones that are among the best substrates. Conversely, the very best lactones that are commercially available, and that are recognized by the enzyme, are simply not found in KEGG. These are false negatives, but their absence reflects not so much on docking as on the library of molecules that are docked. Finally, the structural predictions allowed us to investigate the stereochemical preference of the reaction.

Substrate Specificity. Dr0930 has been shown to hydrolyze lactones but has a very weak phosphotriesterase activity. The best substrates identified to date are the γ - and δ -lactones with a hydrophobic chain attached to the carbon adjacent to the ring oxygen. Compound **12** has a value of k_{cat} that exceeds 400 s⁻¹ and a k_{cat}/K_m that is greater than 10⁶ M⁻¹ s⁻¹. These values are in excess of those previously reported for other lactonases in the amidohydrolase superfamily:



FIGURE 8: Facial orientations of the amide or ester bonds of substrates or inhibitors in the active sites of Sso2522 (32), dihydroorotase (33), and isoaspartyl dipeptidase (34) plus the proposed facial selectivity for Dr0930. The indicated face of the trigonal functional group to be hydrolyzed is directed toward the nucleophilic hydroxide in the ligand-bound complexes determined from X-ray crystallography. The bond to be cleaved is shown in bold.

AhlA, PPH, and Sso2522 (12). However, we were unable to find any activity toward the hydrolysis of the *N*-acyl derivatives of homoserine lactone (compounds **21** and **22**). This is perhaps not too surprising since the hydrophobic tails in these compounds extend from different parts of the lactone ring and the unsubstituted lactones (compounds **1** and **9**) are poorer substrates than those with a hydrophobic extension at the carbon adjacent to the ring oxygen (Table 2). However, AhlA, PPH, and Sso2522 have all been reported to catalyze the hydrolysis of lactones such as **6** and **11** in addition to *N*-acylhomoserine lactones such as **22** (12). The reason for this difference in specificity is not at all clear.

The structure of Sso2522 (also known as SsoPox) from S. solfataricus complexed with the inhibitor N-decanoyl-Lhomocysteine thiolactone has been determined (PDB code 2VC7) (36). This enzyme, which possesses lactonase and a promiscuous phosphotriesterase activity, contains the same binuclear metal center as the bacterial phosphotriesterases and Dr0930 except that iron is bound to the α -site and cobalt is bound to the β -site (36). In the inhibitor complex of Sso2522, the carbonyl oxygen of the lactone ring interacts with the β -metal and Tyr-97. The sulfur in the lactone ring interacts weakly at a distance of 3.2 Å with the iron in the α -metal position. In the Sso2522 structure, the lactone ring is also contacted by the side chains of Arg-223, Leu-72, Val-27, and Ile-261. In Dr0930, these residues are replaced by Arg-228, Cys-72, Phe-26, and Trp-269. In Sso2522 a long hydrophobic channel accommodates the aliphatic tail of the N-acylhomocysteine inhibitor and is composed of the side chains of Phe-229, Leu-228, Leu-226, Trp-278, Thr-265, Ala-275, and Leu-274 (36). The long hydrophobic channel accommodating the substrate aliphatic chain in Sso2522 is also present in the Dr0930 structure. The corresponding residues in Dr0930 are Val-235, Met-234, Leu-231, Trp-287, Trp-269, Val-284, and Ala-283.

It is interesting to note that in the structure of Sso2522 with the *N*-acylhomocysteine lactone inhibitor, the bridging hydroxide is poised to attack the *re* face of the thiolactone ring (shown in Figure 8). This facial selectivity is at odds with that observed for other members of the amidohydrolase superfamily with bound substrates or inhibitors.² For example, in the complex of dihydroorotase (DHO) with dihydroorotate, the bridging hydroxide is positioned to attack the *re* face of the amide bond (ref *37*; PDB code 1J79). The same facial selectivity is observed in the complex of isoaspartyl dipeptidase (IAD) with the substrate β -Asp-His

(ref 38; PDB code 1YBQ). This selectivity is further supported by the orientation of a methyl phosphonate transition state inhibitor in the active site of N-acetyl-Dglucosamine 6-phosphate deacetylase (NagA) (PDB code 2P53) (39). The facial selectivity observed in DHO, IAD, and NagA optimally positions the invariant aspartate residue from the end of β -strand 8 to transfer the lone proton from the bridging hydroxide to the leaving group during the transformation from substrate to the hydrolyzed products. In the catalytic mechanism proposed for Sso2522 it has been suggested that Cys-258 may function to protonate the leaving group hydroxyl based upon the orientation of the Nacylhomocysteine lactone within the active site (36). This conclusion is unlikely to be true since this residue is not conserved within those members of the AHS that have been demonstrated to possess lactonase activity (see Figure 7). It appears that the homocysteine lactone is binding in an unproductive orientation in the structure of Sso2522, and this may explain why no turnover is observed for this compound.

Mechanism of Action. A catalytic reaction mechanism may be proposed for the hydrolysis of lactones by Dr0930. In this mechanism the carbonyl oxygen would be polarized by interaction with the β -metal ion. In simple lactones, such as depicted by **11**, the *si* face of the lactone ring (see Figure 8) would be orientated toward the hydroxide that bridges the two metal ions. Nucleophilic attack by the bridging hydroxide would initiate the formation of a tetrahedral intermediate and transfer of the proton from the hydroxide to Asp-264. The intermediate would collapse with subsequent transfer of the proton to the leaving group alcohol.

Enhancement of Phosphotriesterase Activity in Dr0930. Two approaches were taken in an attempt to enhance the phosphotriesterase activity of Dr0930. The first attempt involved the transplantation of the eight specificity loops from the PTE of P. diminuta for the eight corresponding loops within the active site of Dr0930. The number of residues within each of these eight loops differed between PTE and Dr0930, but the three-dimensional position for the starting and ending residues within each of these eight loops is the same between these two proteins (Figure 7), and thus it was anticipated that the chimeric protein would be structurally very similar to PTE. However, the chimeric protein was largely insoluble, and the hydrolysis of paraoxon could not be detected with the protein that could be made soluble through refolding. There are thus some rather subtle issues that govern the folding of the chimeric protein that are not well understood. Nevertheless, the construction of rationally placed site-directed mutations within the active site of Dr0930 resulted in the enhancement of the paraoxonase activity by a factor of 16 with only two amino acid changes. These initial results suggest that further improvements in the expression of phosphotriesterase activity within Dr0930 should be possible.

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SUPPORTING INFORMATION AVAILABLE

The structures of compounds found not to be substrates for Dr0930 (Figure S1). This material is available free of charge via the Internet at http://pubs.acs.org.

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