See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/45582086

Structural Determinants for the Stereoselective Hydrolysis of Chiral Substrates by Phosphotriesterase

ARTICLE *in* BIOCHEMISTRY · SEPTEMBER 2010

Impact Factor: 3.02 \cdot DOI: 10.1021/bi101058z \cdot Source: PubMed

CITATIONS

CITATION

9

READS

14

7 AUTHORS, INCLUDING:



Yubo Fan

The Methodist Hospital System

43 PUBLICATIONS 1,133 CITATIONS

SEE PROFILE



Yi Qin Gao

Peking University

98 PUBLICATIONS 2,014 CITATIONS

SEE PROFILE



Lijiang Yang

Peking University

39 PUBLICATIONS 822 CITATIONS

SEE PROFILE



Frank Raushel

Texas A&M University

312 PUBLICATIONS 9,399 CITATIONS

SEE PROFILE



Biochemistry. Author manuscript; available in PMC 2011 September 21.

Published in final edited form as:

Biochemistry. 2010 September 21; 49(37): 7988–7997. doi:10.1021/bi101058z.

Structural Determinants for the Stereoselective Hydrolysis of Chiral Substrates by Phosphotriesterase[†]

Ping-Chuan Tsai \S , Yubo Fan \S , Jungwook Kim ‡ , Lijiang Yang \S , ψ , Steven C. Almo ‡,* , Yi Qin Gao \S , * , ψ , and Frank M. Raushel \S , *

- § Department of Chemistry, Texas A&M University, College Station, TX 77843-3255
- [‡] Albert Einstein College of Medicine, 1300 Morris Park Avenue, Bronx, New York 10461

Abstract

Wild-type phosphotriesterase (PTE) preferentially hydrolyzes the R_p-enantiomers of the nerve agents sarin (GB) and cyclosarin (GF) and their chromophoric analogues. The active site of PTE can be subdivided into three binding pockets that have been denoted as the small, large and leaving group pockets based on high resolution crystal structures. The sizes and shapes of these pockets dictate the substrate specificity and the stereoselectivity for catalysis. Mutants of PTE have been prepared that exhibit substantial changes in substrate specificity and the ability to differentiate between chiral substrates. For example, the G60A is stereoselective for the hydrolysis of the Rp-enantiomer of the chromophoric analogues of sarin and cyclosarin whereas the H254G/ H257W/L303T (GWT) mutant reverses the stereoselectivity for the enantiomers of these two compounds. Molecular dynamics simulations and high resolution X-ray structures identified the correlations between structural changes in the active site and the experimentally determined kinetic parameters for substrate hydrolysis. New high resolution structures were determined for the H257Y/L303T (YT), I106G/F132G/H257Y (GGY) and H254Q/H257F (QF) mutants of PTE. Molecular dynamics calculations were conducted using the S_p- and R_p-enantiomers of the analogues for sarin and cyclosarin for the wild-type PTE and the G60A, YT, GGY, QF, and GWT mutants. The experimental stereoselectivity correlated nicely with the difference in the computed angle of attack for the nucleophilic hydroxide relative to the phenolic leaving group of the substrate.

Phosphotriesterase (PTE1), isolated originally from soil microbes, catalyzes the hydrolysis of a wide range of organophosphate esters, including agricultural insecticides and chemical warfare agents (1,2). The X-ray crystal structure of [Zn²+/Zn²+]-PTE reveals a homodimeric (β/α)₈-barrel structural fold with a binuclear metal center in the active site (3). The two zinc ions are bridged by a hydroxide and a carbamate functional group, formed by the reaction of CO₂ with the ϵ -amino group from an active site lysine residue (4). X-ray crystal structures from the Holden laboratory, determined in the presence of inhibitors, have identified three

SUPPORTING INFORMATION AVAILABLE

The MD simulations for the G60A, H257Y/L303T, H254G/H257W/L303T, I106G/F132G/H257Y mutants are presented in the supporting information in Tables S1-S5. This information is available free of charge via the Internet at http://pubs.acs.org.

[†]This work was supported in part by the National Institutes of Health (GM 68550). The X-ray coordinates and structure factors for the QF, GGY, and YT mutants of PTE have been deposited in the Protein Data Bank (PDB accession codes: 2OQL, 2O4M, and 2OB3).

^{*}To whom correspondence may be sent: (FMR) telephone: (979)-845-3373; fax: (979)-845-9452; raushel@tamu.edu. (YQG) telephone: 86-10-67253815; gaoyq@pku.edu.cn. (SCA) telephone: (718) 430-2746; fax: (718)-430-8565; almo@aecom.yu.edu.

VCurrent address: Beijing National Laboratory of Molecular Sciences and College of Chemistry and Molecular Engineering, Peking University, Beijing, 100871, China.

¹Abbreviations: PTE, phosphotriesterase; GWT, the H254G/H257W/L303T mutant of PTE; YT, the H257Y/L303T mutant of PTE; GGY, the I106G/F132G/H257Y mutant of PTE; and QF, the H254Q/H257F mutant of PTE.

binding pockets that facilitate the association of substrates with the PTE active site (3). These sub-sites have previously been denoted as the *small*, *large* and *leaving group* pockets and are defined by the space enclosed by the side chains of Gly-60, Ile-106, Leu-303 and Ser-308; side chains of His-254, His-257, Leu-271 and Met-317; and side chains of Trp-131, Phe-132, Phe-306, and Tyr-309, respectively (3). A three dimensional representation of the PTE active site is presented in Figure 1.

Wild-type PTE is stereoselective for the hydrolysis of chiral organophosphorus esters (5–8). The degree of stereoselectivity for the wild-type enzyme depends on the size of the substituents attached to the central phosphorus core. For example, the value of $k_{\text{cat}}/K_{\text{m}}$ for wild-type PTE with the R_p-enantiomer of compound 1 (a chromophoric analogue of the nerve agent sarin or GB) is 22-fold larger than that for the corresponding S_p-enantiomer (9). The enantiomeric preference increases to 760 for the hydrolysis of the R_p- and S_penantiomers of compound 2, (an analogue for the nerve agent cyclosarin or GF). The mutation of Gly-60 to alanine results in a substantial enhancement in the relative stereoselectivity for the R_p-enantiomers of compounds 1 and 2 (9). Rational engineering of PTE, through enlargement of the small pocket and reduction of the large pocket, can invert the inherent enantioselectivity of the wild-type enzyme for the hydrolysis of chiral organophosphonates (10). The relative values of $k_{\text{cat}}/K_{\text{m}}$ for the R_{p} - and S_{p} -enantiomers of compounds 1 and 2 (Scheme 1) are graphically illustrated in Figure 2 for specific mutants of PTE. Wild-type PTE and the G60A mutant preferentially hydrolyze the R_p-enantiomers of compounds 1 and 2 when compared to the corresponding S_p-enantiomers, whereas H254G/ H257W/L303T (GWT), H257Y/L303T (YT) and I106G/F132G/H257Y (GGY) prefer the S_p-enantiomers. The H254Q/H257F (QF) mutant is not stereoselective for the hydrolysis of compound 1 but this mutant enzyme preferentially hydrolyzes the R_p-enantiomer of compound 2.

The stereoselective properties of PTE can be manipulated through the mutation of residues in the active site (8–11). What is needed is a physical explanation that is able to correlate changes in the observed kinetic constants with specific changes in the three-dimensional structure. Molecular dynamics simulations have been utilized to probe for correlated changes in three-dimensional structure and catalysis. For example, the wild-type lipase from *Pseudomonas aeruginosa* has no preference for the hydrolysis of either the R- or S-enantiomers of 2-methyl-decanoic acid esters. However, lipase variants, mutated through directed evolution, were discovered to have an increased chiral selectivity towards the S-enantiomer and molecular dynamics simulations have been conducted on the wild-type and mutant enzymes (12). The active site of epoxide hydrolase from *Aspergillus niger* has been redesigned and the catalytic preference for the S-enantiomer of glycidyl phenyl ether increased from 5- to 115-fold. The wild-type enzyme and a series of mutant enzymes with enhanced enantioselectivity were investigated using molecular dynamics simulations and molecular docking techniques (13).

The X-ray crystal structures of wild-type PTE and five mutants have been determined to high resolution. These structures were used as the starting point for molecular dynamics simulations with the R_p - and the S_p -enantiomers of compounds 1 and 2 using the AMBER suite of programs. The binding poses of each enantiomer within these proteins support the experimentally observed stereoselectivity for substrate hydrolysis (9). These efforts demonstrate that MD simulations can facilitate the design of modified enzymes with enhanced catalytic activities for specific substrates.

Materials and Methods

Protein Purification, Crystallization and X-ray Structure Determination

The QF, GGY, and YT mutants of PTE were expressed and purified to homogeneity as described previously (8,9). The QF mutant was crystallized by sitting-drop vapor diffusion at 21 °C after mixing 1.0 µL of protein with 1.0 µL of the reservoir solution (0.1 M Bis-Tris, pH 6.5, and 20% PEG MME 5000) and equilibrating over 0.1 mL of the reservoir solution. X-ray data were collected on an R-Axis IV⁺⁺ image plate detector using CuKα radiation from a Rigaku RU-H3R X-ray generator and processed using HKL2000 (14). The QF crystal exhibited diffraction consistent with the space group P₁. There was a single homodimer of QF in the asymmetric unit. The GGY mutant was crystallized by hangingdrop vapor diffusion at 21 °C by mixing 1.0 μL of protein with 1.0 μL of reservoir solution (0.2 M zinc acetate, 0.1 M sodium cacodylate, pH 6.5, and 12% PEG MME) and equilibrating over 1.0 mL of reservoir solution. X-ray data were collected on the CCD detector ADSC QUANTUM 210 at the NSLS beam line X6A. The crystal exhibited diffraction consistent with space group P1 with two homodimers in the asymmetric unit. The YT mutant was crystallized by hanging-drop vapor diffusion at 21 °C by mixing 1.0 µL of protein with 1.0 µL of reservoir solution (0.1 M Bis-Tris pH 6.5, and 12% PEG MME 5000) and equilibrating over 1.0 mL of reservoir solution. X-ray data were collected on the CCD detector ADSC QUANTUM 315 at the NSLS beam line X29A. The crystal exhibited diffraction consistent with space group P1 with one homodimer in the asymmetric unit.

The structures of the three mutants were determined by molecular replacement using the program MOLREP with PDB code: 1P6B as the initial search model (15). Solvent water molecules were built using Arp/wArp (16). All subsequent model building and refinement was carried out with Coot (17) and REFMAC5 (18) for the QF and GGY mutants or SHELXL (19) for the YT mutant. The final models were refined to 1.80, 1.64 and 1.04 Å for the QF, GGY, and YT mutants, respectively. The data collection and refinement statistics are provided in Table 1.

Computational Details

Chain A in the X-ray crystal structures of PTE, including the wild-type protein (PDB code: 1EZ2), G60A (PDB code: 3CS2), YT (PDB code: 2OB3), GWT (PDB code: 1P6C), QF (PDB code: 2OQL) and GGY (PDB code: 2O4M) mutants were used for the molecule dynamics simulations with the R_{p} - and S_{p} -enantiomers of 1 and 2 in the active site. The conformations of the amino acids side chains with the highest occupancy were used for the simulations if there were more than one available in the crystal structure. All waters interacting with chain A were kept for the simulations. For the two structures (PDB codes: 3CS2 and 2O4M) where cacodylate was bound in the active site instead of the bridging hydroxide, the active sites were aligned to that of the wild-type PTE (PDB code: 1EZ2), the cacodylate was removed and hydroxide was added between the two Zn²⁺ ions accordingly. The Co²⁺ ions in the active site of G60A in the structure depicted by PDB: 3CS2 were replaced by Zn²⁺ ions in the computation. The substitution of Co²⁺ in the active site of PTE for Zn²⁺ does not significantly affect the kinetic constants or the overall structure of the active site (2). The geometries of the analogues were fully optimized at the level of B3LYP(20-22)/6-31++G(d',p') (23-25) and the charge distribution on each atom in these molecules was computed using the RESP-fit method at the level of B3LYP/cc-pVTZ (26) with the solvent effect ($\varepsilon = 4$) included using the polarizable continuum model (PCM) (27– 29). The force field for Zn²⁺ and the charge distributions for the four Zn-coordinating histidines (His-55, His-57, His-201 and His-230), the carboxylated lysine (Lys-169) and hydroxide were developed by Pang (30). The side chains of the Zn-coordinating histidines were fully deprotonated and modeled as imidazolates (30). His-254 was assigned as the

protonated form while all of the remaining histidines were kept neutral with N^ϵ protonated. The aspartate and glutamate residues (Asp-100, Asp-105, Asp-253 and Glu-56) were kept neutral to compensate for the deprotonation of the Zn-coordinating histidines. In the mutant enzymes, the corresponding histidines were kept in the same protonation states as those in the wild-type enzyme except for mutations to His-254 and His-257. The net negative charge was neutralized by added Na^+ ions. Explicit TIP3P waters (31) were added as a truncated octahedral water box with a 10-Å buffer.

All operations were processed using the AMBER 9 Leap module (32), which led to a total of 72 systems (6 enzymes with 2 chiral substrates and 3 different initial poses for each enantiomer) solvated with 7400 to 8000 water molecules. The systems were further extended using the periodic boundary condition. The Amber force field 99 (33) with the reoptimized parameters for the peptide backbone (34) was utilized for all amino acid residues and the hydroxide ion, while the general AMBER force field was used for the substrates. MD simulations were conducted at 300 K and 1 atm. The SHAKE algorithm (35) was used to constrain all bonds involving hydrogens. A 10.0 Å cutoff was applied for nonbonding interactions. The Particle Mesh Ewald method was employed to treat the long-range electrostatic interactions (36,37).

MD trajectories were obtained on these structures using the AMBER suite of programs. Two steps of minimization were conducted before MD simulations. During the minimization, the systems were first optimized for 1000 cycles while the backbone of the protein was frozen with a force constant of 500.0 kcal/mol/Å². The systems were further optimized for 2500 cycles without restraints, followed by a 20-ps MD simulation of heating the system from 0 to 300 K, with a force constant of 10.0 kcal/mol/Å² upon all peptide backbone atoms. During the MD simulations, the distance between the oxygen of the P=O moiety and Zn_B was kept at 2.2 Å with a force constant of 100.0 kcal/mol/Å². This restraint allows the substrates to interact with the residues in the three subsites and for the phosphonate core to bind directly to Zn_{β} prior to hydrolysis. In addition, for many of the disfavored binding states, the substrates would dissociate into the bulk solution in the absence of this restraint due to large unfavorable steric effects. Since the ligands were manually added to the active sites many disfavored interactions to the protein overwhelmed the binding between the phosphonate core and Zn_B in the initial geometries not only for the favored structures but also for the disfavored. Therefore, it would not be possible to compare the disfavored complexes to the more favored states. At a distance (d_{ZnO}) of 2.2 Å, all of the steric effects between the substituents from the substrates are similar to those that would occur prior to hydrolysis and thus the overall comparison of the various structures becomes more reasonable. The binding energies computed with this restraint are not very meaningful because the energy contributions from the Zn₈-O distance restraint vary for the mutants and the initial binding poses.

In the first nanosecond of relaxation, the angle for O-P-X, where O is the oxygen of the bridging hydroxide, P is the phosphorus of the phosphonate ester and X is one of the three possible singly bonded atoms attached directly to the phosphorus core, was set to 180° with a restraining force constant of $100.0 \, \text{kcal/mol/degree}^2$. The restraining force was released after the relaxation. The equilibrium structure without the angle restraint should be similar to the bound substrate complex prior to attack by the bridging hydroxide. The productive conformations in our simulations are close to the near attack conformers (NAC) defined by Bruice (38). The angles for O-P-X are defined as A_L , A_A and A_M where X is the phenolate oxygen in the *p*-acetophenolate leaving group, the alkoxide oxygen of the isopropyl or cyclohexyl substituent and the methyl carbon, respectively, for compounds 1 and 2. The distance between $Z_{n\beta}$ and the phosphoryl oxygen of the substrate is defined as d_{ZnO} while the distance between the phosphorus and the bridging hydroxide oxygen is defined as d_{PO} .

Data were collected every 0.5 ps in continuous trajectories until the system reached equilibrium. The lengths of the trajectories varied from 10 to 25 ns. The last 5-ns of the equilibrated trajectories were utilized for structural analyses.

Results and Discussion

Three-Dimensional Structures of QF, GGY, and YT

The three dimensional structures of QF, GGY, and YT are nearly identical to that of the wild-type enzyme. In the QF and YT structures, one homodimer was found in the asymmetric unit and two metal atoms are located in the active site, bridged by a water molecule or hydroxide. The crystals of GGY contained two homodimers in the asymmetric unit, but cacodylate is bound between the two active site metal ions instead of hydroxide in all four subunits (data not shown). Cacodylate was included in the crystallization solution and the binding mode for this compound in the active site is similar to that observed previously in the structure of the G60A mutant (39). Crystals of the YT mutant exhibited outstanding X-ray diffraction; the data were collected and processed to 1.04 Å. The high resolution data enabled anisotropic temperature factor refinements and yielded a model with R_{work}/R_{free} of 0.105/0.127. An example of the $2F_o - F_c$ electron density map around the binuclear metal center is shown in Figure 3.

Comparison of Active Site Structures

The active site structures of wild-type PTE and the five mutant enzymes (G60A, YT, GWT, QF, and GGY) that have been determined by X-ray crystallography are shown in Figures 4a–f. Superposition of the $C\alpha$ atoms of the five mutant enzymes with those of wild-type PTE, gives an RMSD for G60A, YT, GWT, GGY and QF of 0.36, 0.51, 0.42, 0.36 and 0.55 Å, respectively. The mutations that have been introduced into the active site of PTE do not, therefore, significantly change the backbone conformation of the protein, including those regions near the active site. The crystal structures of PTE reveal that the small binding pocket is the most isolated from the external environment compared to the large and the leaving group pockets. The large binding pocket is formed by His-254, His-257, Leu-271 and Met-317, but overall, this pocket is accessible to solvent molecules. The leaving group pocket is even more exposed to solvent than the large binding pocket.

The impact of the mutations on the structure of the substrate binding pocket is readily detectable in some instances, as in the structure of GGY (Figure 4c). The replacement of Ile-106 with glycine clearly enlarges the small binding pocket compared to that of the wild-type enzyme, consistent with the observed kinetic results demonstrating that GGY favors the hydrolysis of the S_p -enantiomers over the R_p -enantiomers. For G60A, the methyl substitution reduced the size of the small binding pocket (Figure 4b) and the preference for the R_p -enantiomer is enhanced relative to the wild-type enzyme. In some cases, however, the structural changes are somewhat ambiguous, as observed with H254Q of the QF mutant (Figure 4f). The net effect of H254G in the GWT mutant is not intuitively clear, since the mutation actually increases the size of the large binding pocket (Figure 4e). The GWT mutant exhibits an inverted stereoselectivity relative to the wild-type enzyme, and hydrolyzes S_p -1 and S_p -2 more effectively than the two R_p -enantiomers. The replacement of the adjacent His-257 to tryptophan may have compensated for the changes caused by the H254G mutation.

Mechanism of Action for Substrate Hydrolysis by PTE

A comprehensive reaction mechanism for the hydrolysis of organophosphates by PTE has been proposed (40). In this mechanism the substrate binds in the active site via an initial interaction of the phosphoryl oxygen with M_{β} . The bridging hydroxide subsequently attacks

the phosphorus center and the leaving group departs via the formation/breakdown of a trigonal bipyramidal transition state. This mechanism is consistent with the net inversion of stereochemistry with chiral phosphate substrates (41). X-ray crystal structures of bound substrate analogues show that the phosphoryl oxygen electrostatically interacts with M_{β} (42). In addition, the kinetic constants for the hydrolysis of phosphate and thiophosphate esters indicate that the P=O bond is polarized in the transition state by a direct interaction with M_{β} (43). Finally, the three-dimensional structure of diethyl phosphate bound to PTE shows that the hydrolysis product bridges the two divalent cations in the active site and displaces the bridging hydroxide (39). Based on these experimental observations, the optimal substrate binding pose is one where the angle of attack by the bridging hydroxide, relative to the leaving group phenol, is ~180°, and the phosphoryl oxygen is positioned to interact with M_{β} . In this pose the angle from the attacking hydroxide to the other two substituents is ~70° (180° - 110°).

Initial Geometries of Bound Substrates for MD Simulations

Models of the productive binding modes for the R_p - and S_p -enantiomers of $\bf 1$ and $\bf 2$ with PTE are graphically depicted in Figure 5 and designated as R_L and S_L , respectively. In these initial poses, the phenolate substituent is placed in the leaving group pocket while the methyl and O-alkyl groups are positioned in either the small or large pocket, depending on the absolute stereochemistry of the substrate. The four nonproductive binding modes include orientations where either the methyl or O-alkyl group is positioned within the leaving group pocket at an initial angle of ~180° relative to the attacking hydroxide. These poses have been designated as R_A , R_M , S_A , or S_M , depending on which one of the substituents is positioned within the leaving group pocket. Since compounds $\bf 1$ and $\bf 2$ are chiral, there are a total of 12 initial poses for each of the six enzymes to be evaluated.

At the termination of the simulations, the angles from each of the three substituents to the attacking hydroxide, and the distances from phosphorus to the hydroxide and from M_{β} to the phosphoryl oxygen were averaged. The distances (d_{PO} and d_{ZnO}) and angles (A_A , A_L , and A_M) are defined as shown in Scheme 2. A_L represents the angle from the attacking hydroxide to the leaving group phenol through the phosphorus. A_A is the angle from the hydroxide to the O-alkyl group and A_M is the angle to the methyl substituent. The value of d_{PO} is defined as the distance between the bridging hydroxide and the phosphorus atom of the substrate. The value of d_{ZnO} is the distance between the phosphoryl oxygen and M_{β} .

The values of A_L for the three initial binding poses for R_p-1 with wild-type PTE are plotted as a function of time in Figure 6. During the first 1.0 ns, A_L oscillates within a small range due to the initial restraint on this parameter. After 1.0 ns this restraint was removed and all three substituents within a given substrate are allowed to rotate freely. As shown in Figure 6, the leaving group quickly changes its orientation after the release of the initial restraint. The equilibrium position for each of the starting poses is reached in a relatively short period of time. For the R_L and R_M poses, the relative orientation of the substrate in the active site is quite stable and the phenolic group remains in either the leaving group pocket (for the R_L pose) or large pocket (for the R_M pose). However, for the R_A pose the leaving group repositions from the small pocket, after the initial restraint is released, to an orientation that is ~111° relative to the attacking hydroxide. In this structure the leaving group moves to the large pocket although the phosphonate core does not rotate significantly relative to the initial geometry. Snapshots selected from the trajectories during the last 5-ns of simulations were used to estimate the average deviations from the initial poses. The angles (A_I) and distances (d_{PO}) in these snapshots were subsequently used to assess the relative reactivities of the various complexes.

Molecular Dynamic Simulations of Wild-type PTE

The structure parameters for the binding of compounds 1 and 2 to wild-type PTE are listed in Table 2. Representative images of the enzyme-substrate complexes after the MD simulations are shown in Figure 7a, b for R_p -1 and S_p -1 and Figure 7c, d for R_p -2 and S_p -2. With R_p -1, for the initial pose R_A the leaving group is not in an appropriate orientation relative to the attacking hydroxide with a value of A_L = 111° and thus displacement of the leaving group cannot readily occur. From the three initial binding complexes, R_L has the best final orientation with a value of A_L = 160°, which is the closest to 180° among all poses. In R_A and R_M the leaving group is unproductively positioned at 111o and 74°, respectively. In addition, the distance between the bridging hydroxide and phosphorus, d_{PO} , is the shortest for A_L , with a value of 3.4 Å. Therefore, the initial binding pose of R_L is the most optimal in leading to productive hydrolysis of R_p -1.

The equilibrium orientation of the leaving group starting from S_A is not in an appropriate position for hydrolysis with a value of $A_L = 68^\circ$. For either S_L or S_M , the equilibrium value for A_L is ~140–142°, which deviates from the ideal angle of ~180°. Since the bulky phenolate group is too large to fit properly in the small binding pocket, a conformational change was observed in the MD simulations for S_M . The phenolate group rotates out of the small binding pocket and complexes in the active site in a binding mode that is very similar to S_L , although not identical. These results are consistent with the higher reactivity of wild-type PTE for R_p -1 relative to S_p -1. The attack angle (161° vs 140°) and shorter distance between the hydroxide and phosphorus (3.4 Å vs 3.8 Å) favors R_L over S_L and thus the R_p -enantiomer of 1 is a better substrate than is the Sp-enantiomer.

For the R_p and S_p -enantiomers of 2, the equilibrium value of A_L (156°) for R_L is greater than that for S_L (143°) and the distance between hydroxide and the phosphorus center (d_{PO}) is shorter by 0.3 Å (3.5 vs. 3.8 Å). Therefore, the values for A_L and d_{PO} are more optimal for the hydrolysis of R_p -2 compared to S_p -2. for an in-line S_N 2-like reaction mechanism. In the MD simulations the conformational change from S_M to S_L was not observed in the analysis of S_p -2. The larger size of the cyclohexyl group over the isopropyl group apparently prevents the S_M conformation from rotating to an S_L -like conformation during the simulation.

Molecular Dynamics Simulations with the G60A Mutant

During the simulations, the initial R_A pose changes to an R_L -like one because of the enhanced mismatch of the bulky leaving group and the reduced size of the small pocket. For the R_L and S_L poses, the angle A_L for the R_p -enantiomer is greater than that for the S_p -enantiomer by 40° (Table 3). The distance d_{PO} for the R_p -enantiomer is shorter than that for the S_p -enantiomer by 0.6 Å. These results are fully consistent with the observed enhancement in the ratio of hydrolysis of the R_p - and S_p -enantiomers relative to the wild type enzyme. G60A catalyzes the hydrolysis of the R_p -enantiomer 84-fold faster than the S_p -enantiomer.

For compound 2 the initial R_L pose is the only one that results in a suitable angle of attack A_L (162°) for the bridging hydroxide (Table 4). In addition, the distance (d_{PO}) between the bridging hydroxide and the phosphorus center is the shortest for this pose. The A_L angles for the five other initial poses are all smaller than 103° (see supplementary Tables S1–S5). The reduced small pocket strongly disfavors the binding of either the p-acetylphenol leaving group or the cyclohexyl substituent. Due to steric repulsion between the cyclohexyl group and the small pocket, the initial S_L pose changes to the S_A pose during the MD simulation. The large difference in the computed value of A_L for R_L relative to S_L (84°) is consistent

with the substantial difference in rates for the hydrolysis of the R_p - and S_p -enantiomers of compound 2 by the G60A mutant.

Molecular Dynamic Simulations with GWT, YT, and GGY

The GWT, YT and GGY mutants favor the hydrolysis of the S_p -enantiomer of compound 1. The stereoselectivity of these mutants is thus inverted relative to the wild type enzyme. Consistent with these results, the computed values for $\Delta A_L(R_p-S_p)$ with compound 1 are reversed relative to wild-type PTE, although the values (-2, -6, and 0 for GWT, YT and GGY, respectively), are not significantly different from one another given the computational uncertainties associate with these values (Table 3). For the GWT and YT mutants, the values of d_{PO} for the S_p -enantiomer are shorter than those for the R_p -enantiomer by 0.2 and 0.3 Å, respectively. However, $\Delta d_{PO}(R_p-S_p)$ for the GGY mutant is essentially zero.

The GGY, YT, and GWT mutants prefer to hydrolyze the S_p -enantiomer of compound 2. The values for $\Delta A_L(R_p-S_p)$ with GGY and YT are -14° and -2° , respectively. Moreover, d_{PO} for the S_p -enantiomers is 0.4 Å shorter for the GGY mutant and 0.1 Å shorter for the YT mutant (Table 4). However, for the GWT mutant, these two parameters favor the R_p enantiomer by 2° and 0.3 Å, respectively.

Molecular Dynamic Simulations with QF

No significant stereoselectivity towards compound 1 was observed for QF. However, this mutant preferably hydrolyzes the R_p -enantiomer of compound 2, relative to the S_p -enantiomer by a factor of 16. From the simulations with compound 1 bound to QF, small differences were found for both $\Delta A_L(R_p\text{-}S_p)$ and $\Delta d_{PO}(R_p\text{-}S_p)$ of 8° and -0.1 Å, respectively. However, for compound 2 a favored $\Delta A_L(R_p\text{-}S_p)$ of 16° and $\Delta d_{PO}(R_p\text{-}S_p)$ of -0.5 Å for the $R_p\text{-}enantiomer$ were obtained (Table 4).

Correlation between A_L and Reactivity

The analysis of the MD results has focused on the attack angle (A_L) and the distance between the hydroxide and the phosphorus center of the substrate (d_{PO}). The strongest correlation with the observed stereoselectivity is the attack angle A_L . Shown in Figure 8 is a plot of the relative reactivity for hydrolysis of the two enantiomers of compounds 1 and 2 with the value of $\Delta A_L(R_p\text{-}S_p)$, where the maximum possible value for $\Delta A_L(R_p\text{-}S_p)$ is ~110°. The largest deviations from this correlation occurs for the hydrolysis of compound 2 with the wild type and the GWT mutant. For compound 2 the bulky cyclohexyl group complicates the interactions between the substrate and the active site. Therefore, the conformational degrees of freedom are likely to be more complex and may involve concerted motions of substrate and nearby residues. The overall efficiency of hydrolysis of compound 2 is significantly lower than that of compound 1 (as judged by the absolute value of k_{cat}/K_m) indicating that the size of the cyclohexyl group might be too large to fit optimally in the active site.

The values of $\Delta d_{PO}(R_p-S_p)$ are not directly related to the chiral preferences observed experimentally but appear to be a consequence of $\Delta A_L(R_p-S_p)$, resulting in an enhancement in selectivity. For compound 1, the more optimal the value of A_L , the shorter is the value of d_{PO} . However, this relationship does not hold for compound 2.

Summary

Wild-type PTE preferentially hydrolyzes the less toxic R_p -enantiomers of the nerve agents GB and GF (9) and the chromophoric analogues (compounds 1 and 2) of these nerve agents. Mutations localized to the active site of PTE have been identified where the stereoselectivity is either enhanced or inverted (8,9,11). It has been assumed that the enhancement in

reactivity for the hydrolysis of the initially slower S_p -enantiomers derives from the optimization of the active site for the specific compound to be hydrolyzed. For example, the methyl and isopropyl groups in the R_p -enantiomer of compound 1 fit reasonably well in the small and large pockets when the phosphoryl oxygen interacts with Zn_β and the leaving group points outwards toward solution. Since the S_p -enantiomer of compound 1 is turned over at a rate 1/20 that of the R_p -enantiomer, the orientation and/or conformational dynamics of PTE bound with the S_p -1 must be inferior to that for R_p -1. The relatively straightforward strategy of resizing the large and small pockets using rational design and semi-random mutagenesis achieved the desired results in the enhancement or inversion of stereoselectivity exhibited by the wild-type enzyme (8,10,11,44). The G60A mutant suppresses the rate of hydrolysis of the slow S_p -enantiomers of 1 and 2 through the substitution of a methyl group into the small pocket within the active site. In the GWT mutant the stereoselectivity is apparently reversed by reducing the size of the large pocket and expanding the size of the small pocket.

Based on the MD simulations, the orientation of the leaving group is the determining factor for chiral selectivity. The optimization of the substrate within the active site can be estimated by the angle of the attacking hydroxide toward the phosphorus center relative to the leaving group phenol. A direct correlation was found for the experimentally determined stereoselectivity and the calculated attack angle (Figure 8). For these compounds the stereoselectivity varies over a range of six orders of magnitude. The logarithm of $^{R/S}(k_{cat}/K_m)$ correlates positively with the difference in the angle of attack on the phosphorus center by the bridging hydroxide, $\Delta A_L(R_p\mbox{-}S_p)$. The correlation was superior for the enantiomers of 1 relative to 2 which implies that the more bulky cyclohexyl substituent in 2 may induce some more complex binding interactions. To address these issues, longer time-scale simulations based on enhanced sampling techniques and additional initial binding poses can be tested.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

We thank Jennifer Cummings for help in the preparation of the figures for this paper. All simulations were conducted with the supercomputing facilities--Brazos and Medusa (NSF CHE-0541587) at Texas A & M University.

References

- Dumas DP, Durst HD, Landis WG, Raushel FM, Wild JR. Inactivation of organophosphorus nerve agents by the phosphotriesterase from *Pseudomonas diminuta*. Arch Biochem Biophys. 1990; 277:155–159. [PubMed: 2154956]
- Omburo GA, Kuo JM, Mullins LS, Raushel FM. Characterization of the zinc binding site of bacterial phosphotriesterase. J Biol Chem. 1992; 267:13278–13283. [PubMed: 1320014]
- 3. Benning MM, Hong SB, Raushel FM, Holden HM. The binding of substrate analogs to phosphotriesterase. J Biol Chem. 2000; 275:30556–30560. [PubMed: 10871616]
- Benning MM, Kuo JM, Raushel FM, Holden HM. Three-dimensional structure of phosphotriesterase: an enzyme capable of detoxifying organophosphate nerve agents. Biochemistry. 1994; 33:15001–15007. [PubMed: 7999757]
- 5. Hong SB, Raushel FM. Stereochemical preferences for chiral substrates by the bacterial phosphotriesterase. Chem Biol Interact. 1999; 120:225–234. [PubMed: 10421456]

 Chen-Goodspeed M, Sogorb M, Raushel FM. Determining reactivity and stereospecificity of phosphotriesterase from *Pseudonomas diminuta* by site-specific mutagenesis. FASEB J. 1999; 13:A1446–A1446.

- 7. Chen-Goodspeed M, Sogorb MA, Wu FY, Hong SB, Raushel FM. Structural determinants of the substrate and stereochemical specificity of phosphotriesterase. Biochemistry. 2001; 40:1325–1331. [PubMed: 11170459]
- Chen-Goodspeed M, Sogorb MA, Wu FY, Raushel FM. Enhancement, relaxation, and reversal of the stereoselectivity for phosphotriesterase by rational evolution of active site residues. Biochemistry. 2001; 40:1332–1339. [PubMed: 11170460]
- 9. Tsai PC, Bigley A, Li Y, Ghanem E, Cadieux CL, Kasten SA, Reeves TE, Cerasoli DM, Raushel FM. Stereoselective hydrolysis of organophosphate nerve agents by the bacterial phosphotriesterase. Biochemistry. 2010; 49:0000–0000.
- Hill CM, Li WS, Thoden JB, Holden HM, Raushel FM. Enhanced degradation of chemical warfare agents through molecular engineering of the phosphotriesterase active site. J Am Chem Soc. 2003; 125:8990–8991. [PubMed: 15369336]
- Li WS, Lum KT, Chen-Goodspeed M, Sogorb MA, Raushel FM. Stereoselective detoxification of chiral sarin and soman analogues by phosphotriesterase. Bioorganic & Medicinal Chemistry. 2001; 9:2083–2091. [PubMed: 11504644]
- Bocola M, Otte N, Jaeger KE, Reetz MT, Thiel W. Learning from directed evolution: theoretical investigations into cooperative mutations in lipase enantioselectivity. Chembiochem. 2004; 5:214– 223. [PubMed: 14760743]
- Reetz MT, Bocola M, Wang LW, Sanchis J, Cronin A, Arand M, Zou J, Archelas A, Bottalla AL, Naworyta A, Mowbray SL. Directed evolution of an enantioselective epoxide hydrolase: uncovering the source of enantioselectivity at each evolutionary stage. J Am Chem Soc. 2009; 131:7334–7343. [PubMed: 19469578]
- 14. Otwinowski, Z.; Minor, W. Macromolecular Crystallography, part A. In: Carter, CW., Jr; Sweet, RM., editors. Methods in Enzymology. 1997. p. 307-326.
- 15. Vagin A, Teplyakov A. MOLREP: An automated program for molecular replacement. J Appl Crystallogr. 1997; 30:1022–1025.
- Lamzin, VS.; Wilson, KS.; Perrakis, A. Crystallography of biological macromolecules. In: Rossmann, MG.; Arnold, E., editors. International Tables for Crystallography. Kluwer Academic Publishers; Dordrecht, The Netherlands: 2001. p. 720-722.
- 17. Emsley P, Cowtan K. Coot: model-building tools for molecular graphics. Acta Crystallogr D Biol Crystallogr. 2004; 60:2126–2132. [PubMed: 15572765]
- 18. Murshudov GN, Vagin AA, Dodson EJ. Refinement of macromolecular structures by the maximum-likelihood method. Acta Crystallogr D Biol Crystallogr. 1997; 53:240–255. [PubMed: 15299926]
- Sheldrick GM, Schneider TR. SHELXL: high-resolution refinement. Methods Enzymol. 1997;
 277:319–343. [PubMed: 18488315]
- Becke AD. Density-Functional Exchange-Energy Approximation with Correct Asymptotic-Behavior. Phys Rev A. 1988; 38:3098–3100. [PubMed: 9900728]
- 21. Becke AD. A New Mixing of Hartree-Fock and Local Density-Functional Theories. J Chem Phys. 1993; 98:1372–1377.
- 22. Lee CT, Yang WT, Parr RG. Development of the Colle-Salvetti Correlation-Energy Formula into a Functional of the Electron-Density. Physical Review B. 1988; 37:785–789.
- Krishnan R, Binkley JS, Seeger R, Pople JA. Self-Consistent Molecular-Orbital Methods .20. Basis Set for Correlated Wave-Functions. J Chem Phys. 1980; 72
- 24. Clark T, Chandrasekhar J, Spitznagel GW, Schleyer PV. Efficient Diffuse Function-Augmented Basis-Sets for Anion Calculations .3. The 3–21+G Basis Set for 1st-Row Elements, Li-F. J Comput Chem. 1983; 4
- Hehre WJ, Ditchfie R, Pople JA. Self-Consistent Molecular-Orbital Methods. 12 Further Extensions of Gaussian-Type Basis Sets for Use in Molecular-Orbital Studies of Organic-Molecules. J Chem Phys. 1972; 56:2257–2261.

26. Kendall RA, Dunning TH, Harrison RJ. Electron-Affinities of the 1st-Row Atoms Revisited - Systematic Basis-Sets and Wave-Functions. J Chem Phys. 1992; 96:6796–6806.

- 27. Cances E, Mennucci B, Tomasi J. A new integral equation formalism for the polarizable continuum model: Theoretical background and applications to isotropic and anisotropic dielectrics. J Chem Phys. 1997; 107:3032–3041.
- 28. Cossi M, Barone V, Mennucci B, Tomasi J. Ab initio study of ionic solutions by a polarizable continuum dielectric model. Chem Phys Lett. 1998; 286:253–260.
- 29. Mennucci B, Cances E, Tomasi J. Evaluation of solvent effects in isotropic and anisotropic dielectrics and in ionic solutions with a unified integral equation method: Theoretical bases, computational implementation, and numerical applications. J Phys Chem B. 1997; 101:10506–10517.
- Pang YP. Successful molecular dynamics simulation of two zinc complexes bridged by a hydroxide in phosphotriesterase using the cationic dummy atom method. Proteins. 2001; 45:183– 189. [PubMed: 11599021]
- Jorgensen WL, Chandrasekhar J, Madura JD, Impey RW, Klein ML. Comparison of Simple Potential Functions for Simulating Liquid Water. J Chem Phys. 1983; 79:926–935.
- 32. Case DAADT, Cheatham I, TE, Simmerling CL, Wang J, Duke RE, Luo R, Merz KM, Pearlman DA, Crowley M, Walker RC, Zhang W, Wang B, Hayik S, Roitberg A, Seabra G, Wong KF, Paesani F, Wu X, Brozell S, Tsui V, Gohlke H, Yang L, Tan C, Mongan J, Hornak V, Cui G, Beroza P, Mathews DH, Schafmeister C, Ross WS, AKP. AMBER. 2006:9.
- 33. Wang JM, Cieplak P, Kollman PA. How well does a restrained electrostatic potential (RESP) model perform in calculating conformational energies of organic and biological molecules? J Comput Chem. 2000; 21:1049–1074.
- 34. Simmerling C, Strockbine B, Roitberg AE. All-atom structure prediction and folding simulations of a stable protein. J Am Chem Soc. 2002; 124:11258–11259. [PubMed: 12236726]
- 35. van Gunsteren WF, Berendsen HJC. Algorithms for Macromolecular Dynamics and Constraint Dynamics. Mol Phys. 1977; 34:1311–1327.
- Darden T, York D, Pedersen L. Particle Mesh Ewald an N.Log(N) Method for Ewald Sums in Large Systems. J Chem Phys. 1993; 98:10089–10092.
- Essmann U, Perera L, Berkowitz ML, Darden T, Lee H, Pedersen LG. A Smooth Particle Mesh Ewald Method. J Chem Phys. 1995; 103:8577–8593.
- 38. Bruice TC. A view at the millennium: The efficiency of enzymatic catalysis. Acc Chem Res. 2002; 35:139–148. [PubMed: 11900517]
- 39. Kim J, Tsai PC, Chen SL, Himo F, Almo SC, Raushel FM. Structure of diethyl phosphate bound to the binuclear metal center of phosphotriesterase. Biochemistry. 2008; 47:9497–9504. [PubMed: 18702530]
- 40. Aubert SD, Li YC, Raushel FM. Mechanism for the hydrolysis of organophosphates by the bacterial phosphotriesterase. Biochemistry. 2004; 43:5707–5715. [PubMed: 15134445]
- 41. Lewis VE, Donarski WJ, Wild JR, Raushel FM. Mechanism and Stereochemical Course at Phosphorus of the Reaction Catalyzed by a Bacterial Phosphotriesterase. Biochemistry. 1988; 27:1591–1597. [PubMed: 2835095]
- 42. Benning MM, Hong SB, Raushel FM, Holden HM. The binding of substrate analogs to phosphotriesterase. J of Biol Chem. 2000; 275:30556–30560. [PubMed: 10871616]
- 43. Raushel FM. Bacterial detoxification of organophosphate nerve agents. Current Opinion in Microbiology. 2002; 5:288–295. [PubMed: 12057683]
- 44. Lum, KT. PhD dissertation. Department of Toxicology, Texas A& M University; 2004. Directed evolution of phosphotriesterase: Towards the efficient detoxification of sarin and soman.
- 45. Pettersen EF, Goddard TD, Huang CC, Couch GS, Greenblatt DM, Meng EC, Ferrin TE. UCSF Chimera--a visualization system for exploratory research and analysis. J Comput Chem. 2004; 25:1605–1612. [PubMed: 15264254]
- 46. Couch GS, Hendrix DK, Ferrin TE. Nucleic acid visualization with UCSF Chimera. Nucleic Acids Res. 2006; 34:e29. [PubMed: 16478715]



Figure 1.

The substrate binding pocket of wild-type PTE. The residues assigned to the small, large, and leaving group pockets are depicted in purple, blue, and green, respectively. The two metal ions are depicted in light blue and the bridging hydroxide is in red. The image was constructed from PDB code: 1HZY.

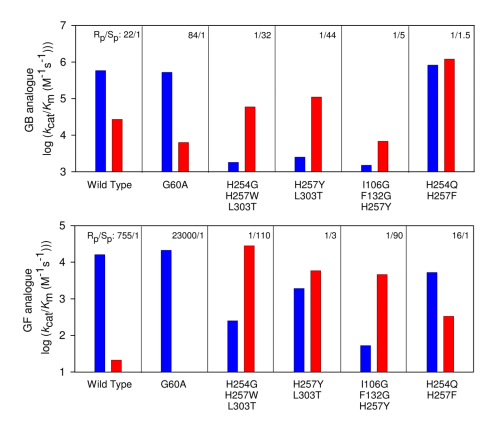


Figure 2. Bar graph illustrating the values of $k_{\text{cat}}/K_{\text{m}}$ for the R_{p} - and S_{p} -enantiomers of compounds 1 and 2 (9).

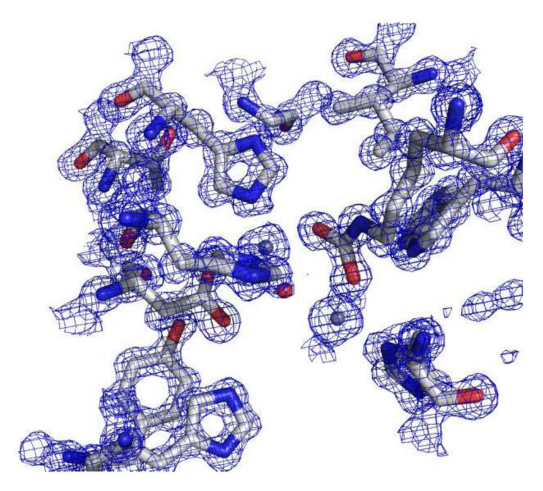


Figure 3. $2F_o - F_c$ map of the crystal structure for the YT mutant near the active site at a contour level of $\sigma = 1.5$. Carbon atoms are shown in light-grey, nitrogen in blue, oxygen in red and arsenic in purple. Grey spheres represent zinc.

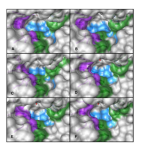


Figure 4.

The active site structure of wild-type PTE and five mutant enzymes (A) Wild-type; (B) G60A; (C) I106G/F132G/H257Y; (D) H257Y/L303T; (E) H254G/H257W/L303T; and (F) H254Q/H257F. The images were drawn with Chimera (45,46). The surfaces are colored grey; the surface for C_{α} and side chains for residues 254, 257, 317, and 271were colored purple. The surfaces for residues 60, 303, 308 and 106 were colored blue and the surface for residues 131, 132, 306, and 309 were colored green. The stick figures of the residues were colored a shade darker than their corresponding surface. The metals are represented as black spheres and the bridging water, when present, was colored as a red sphere. A transparency of 15% was placed on the surface with a "one layer" transparency.

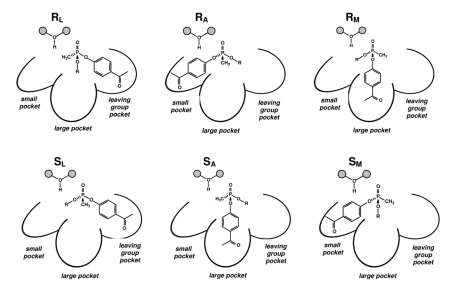
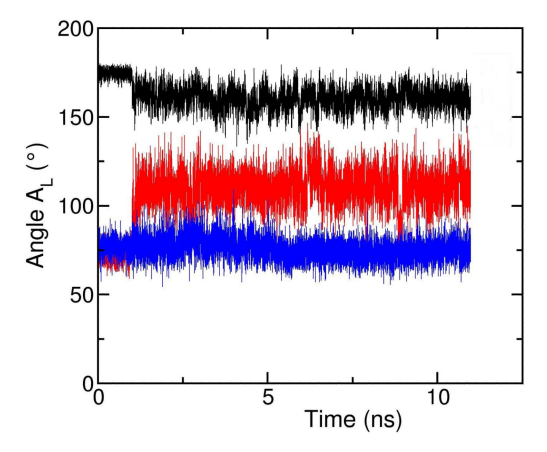


Figure 5.

Cartoon representations for the initial binding poses for the $R_p\text{-}$ and $S_p\text{-}\text{enantiomers}$ of compounds 1 and 2 in the active site of PTE. Each enzyme-substrate complex was oriented to have three different initial binding poses during the MD simulations. The binding poses that place the p-acetyl phenol group, the O-R group, and the methyl substituent towards the leaving group pocket for the $R_p\text{-}\text{substrate-enzyme}$ complexes are denoted as R_L , R_A and R_M , respectively. The binding poses that have the p-acetyl phenol group, the O-R group, and the methyl substituent toward the leaving group pocket for the $S_p\text{-}\text{substrate-enzyme}$ complex are denoted as S_L , S_A and S_M , respectively.



 $\label{eq:Figure 6.} \textbf{ Changes in the value of the } A_L \text{ angles for the } R_L \text{ (black), } R_A \text{ (red), and } R_M \text{ (blue) initial binding poses during MD simulation for } R_p\textbf{-1} \text{ with wild-type PTE.}$

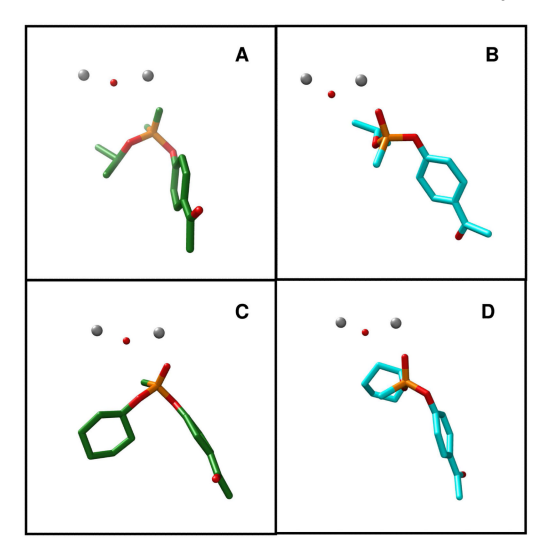


Figure 7. Orientation of the R_p - and S_p -enantiomers of compounds 1 and 2 with wild-type PTE determined from the MD simulations. Panel A: R_L for compound 1; panel B: S_L posed for compound 1; panel C: R_L posed for compound 2; panel D; S_L pose for compound 2.

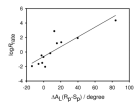
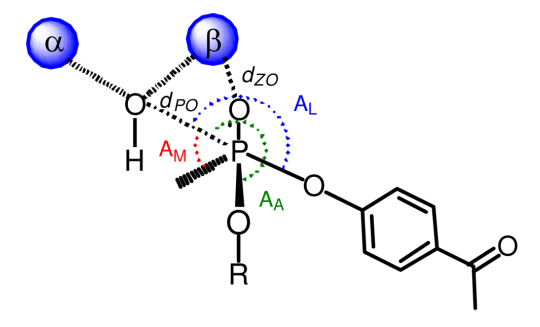


Figure 8. Correlation between the experimental logarithm of $(k_{cat}/K_m)_R/(k_{cat}/K_m)_S$ and $\Delta A_L(R_p-S_p)$.

Scheme 1. Structures of organophosphonate substrates for PTE



Scheme 2. Definition of the angles and distances: $A_L,\,A_M,\,A_N,\,d_{PO},$ and $d_{ZO}.$

 Table 1

 Data collection and refinement statistics for the QF, GGY, and YT mutants of PTE.

		I106G	
Enzyme	H254Q H257F	F132G H257Y	H257Y L303T
		Data Collection	_
Space Group	P1	P1	P1
Cell Dimension			
a, b, c (Å)	43.53, 45.37, 79.11	56.76, 68.91, 89.67	43.36, 45.37, 79.21
α , β , γ (°)	104.75, 93.13, 97.97	90.03, 100.29, 94.12	104.86, 93.27, 97.81
Resolution (Å)	50.0-1.80 (1.86-1.80)	50.0-1.58 (1.64-1.58)	50.0-1.04 (1.08-1.04)
Ι/σ	12.9 (3.0)	10.2 (1.5)	17.3 (6.3)
Completeness (%)	91.8 (70.2)	93.1 (73.5)	85.5 (41.4)
Redundancy	1.9 (1.8)	1.9 (1.7)	3.4 (2.6)
R_{merge}	0.055 (0.245)	0.075 (0.397)	0.061(0.188)
		Refinement	
Observed Reflections	49,147	155,682	235,873
Protein Nonhydrogen Atoms	5,174	10,277	5,178
Water Molecules	700	1,636	855
R_{work}	0.146	0.189	0.105
R_{free}	0.189	0.244	0.127
Average B-factor, (Å ²)	14.48	19.87	9.99
RMSD from Ideal Geometry			
Bond Length (Å)	0.013	0.016	0.015
Bond Angles (°)	1.4	1.6	1.5

Table 2

Structure parameters for the complexes between the wild-type PTE and the Rp- and Sp-enantiomers of 1 and 2.

Tsai et al.

G14.				Initia	Initial Pose		
Substrate	rarameters	\mathbf{R}_{L}	\mathbf{R}_{A}	$\mathbf{R}_{\mathbf{M}}$	$\mathbf{S}_{\mathbf{L}}$	$\mathbf{S}_{\mathbf{A}}$	$\mathbf{S}_{\mathbf{M}}$
	$A_{\rm L}$	160 ± 6	111 ± 10	74 ± 6	140 ± 6	68 ± 5	142 ± 9
	A_A	70 ± 5	137 ± 10	96 ± 4	90 ± 11	160 ± 8	104 ± 10
Compound 1	A_{M}	95 ± 7	70 ± 5	164 ± 5	98 ± 16	7 ± 06	75 ± 7
	q_{PO}	3.4 ± 0.1	3.5 ± 0.1	3.6 ± 0.1	3.8 ± 0.2	3.4 ± 0.1	3.6 ± 0.2
	d_{ZnO}	2.1 ± 0.1	2.1 ± 0.1	2.1 ± 0.1	2.1 ± 0.1	2.1 ± 0.1	2.1 ± 0.1
	$ m A_L$	156 ± 6	75 ± 7	108 ± 8	143 ± 8	64 ± 6	95 ± 6
	A_A	<i>67</i> ± <i>6</i>	159 ± 7	78 ± 4	101 ± 9	149 ± 6	94 ± 4
Compound 2	A_{M}	100 ± 7	87 ± 8	146 ± 7	80 ± 8	110 ± 7	148 ± 5
	d_{PO}	3.5 ± 0.1	3.4 ± 0.2	3.8 ± 0.1	3.8 ± 0.2	3.6 ± 0.2	4.0 ± 0.1
	$d_{ m ZnO}$	2.1 ± 0.1	2.1 ± 0.1	2.1 ± 0.1	2.1 ± 0.1	2.1 ± 0.1	2.1 ± 0.1

Page 23

Comparison of the attack angle, A_L , and the distance, d_{ZnO} , for the productive binding poses of 1 with wild-type PTE and five mutants.

Table 3

Tsai et al.

Enzyme		Wide-type	G60A	H254G H257W L303T	H257Y L303T	1106G F132G H257Y	H254Q H257F
Ratio (R _p /S _p)		22	84	1/33	1/44	1/5	1/1.5
6	${\bf R}_{\rm p}$	160 ± 6	163 ± 7	148 ± 9	146 ± 9	150 ± 5	155 ± 7
A _L (7)	\mathbf{S}_{p}	140 ± 6	123 ± 14	150 ± 8	152 ± 6	150 ± 9	147 ± 9
$\Delta A_L(R_p-S_p)$ (°)		20 ± 6	40 ± 14	-2 ± 9	-2 ± 9 -6 ± 9	6 = 0	8 + 9
4	${\bf R}_{\rm p}$	3.4 ± 0.1	3.4 ± 0.1	3.8 ± 0.2	3.9 ± 0.2	3.5 ± 0.1	3.6 ± 0.2
<i>d</i> _{PO} (A)	\mathbf{S}_{p}	3.8 ± 0.2	3.9 ± 0.1	3.6 ± 0.2	3.6 ± 0.1	3.6 ± 0.1	3.7 ± 0.2
$\Delta d_{PO}(R_p-S_p) (Å)$		-0.4 ± 0.2	-0.5 ± 0.1		0.2 ± 0.2 0.3 ± 0.2	-0.08 ± 0.1	-0.12 ± 0.2

Page 24

Table 4

Comparison of the attack angle, A_L, and the distance, d_{ZnO}, for the productive binding poses of 2 with wild-type PTE and five mutants...

Tsai et al.

Enzyme		Wide-type	G60A	H254G H257W L303T	H257Y L303T	1106G F132G H257Y	H254Q H257G
Ratio (R _p /S _p)		755	23000	1/110	1/3	1/90	16
6	${\bf R}_{\rm p}$	156 ± 6	162 ± 5	151 ± 12	150 ± 8	130 ± 9	166 ± 5
A _L ()	$^{\rm S}_{\rm p}$	143 ± 8	78 ± 12	149 ± 8	152 ± 5	144 ± 8	150 ± 5
$\Delta A_L(R_p\text{-}S_p)~(^\circ)$		13 ± 8	84 ± 12	2 ± 12	-2 ± 8	-14 ± 9	16 ± 5
(V) F	${\bf R}_{\rm p}$	3.5 ± 0.1	3.4 ± 0.1	3.5 ± 0.2	3.8 ± 0.2	4.2 ± 0.2	3.3 ± 0.1
<i>a</i> po (A)	\mathbf{S}_{p}	3.8 ± 0.2	3.6 ± 0.2	3.8 ± 0.2	3.7 ± 0.1	3.8 ± 0.2	3.8 ± 0.1
$\Delta d_{PO}(R_p\text{-}S_p)\;(\text{Å})$		-0.3 ± 0.2	-0.2 ± 0.2	-0.3 ± 0.2	0.1 ± 0.2	0.4 ± 0.2	-0.5 ± 0.1

Page 25