Structures of Ternary Complexes of BphK, a Bacterial Glutathione S-Transferase That Reductively Dechlorinates Polychlorinated Biphenyl Metabolites*

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Prokaryotic glutathione S-transferases are as diverse as their eukaryotic counterparts but are much less well characterized. BphK from Burkholderia xenovorans LB400 consumes two GSH molecules to reductively dehalogenate chlorinated 2-hydroxy-6-oxo-6-phenyl-2,4-dienoates (HOPDAs), inhibitory polychlorinated biphenyl metabolites. Crystallographic structures of two ternary complexes of BphK were solved to a resolution of 2.1 Å. In the BphK-GSH-HOPDA complex, GSH and HOPDA molecules occupy the G- and H-subsites, respectively. The thiol nucleophile of the GSH molecule is positioned for S₂ attack at carbon 3 of the bound HOPDA. The respective sulfur atoms of conserved Cys-10 and the bound GSH are within 3.0 Å, consistent with product formation of a mixed disulfide intermediate. In the BphK-(GSH)₂ complex, a GSH molecule occupies each of the two subsites. The three sulfur atoms of the two GSH molecules and Cys-10 are aligned suitably for a disulfide exchange reaction that would regenerate the resting enzyme and yield disulfide-linked GSH molecules. A second conserved residue, His-106, is adjacent to the thiol of Cys-10 and the GSH bound to the G-subsite and thus may stabilize a transition state in the disulfide exchange reaction. Overall, the structures support and elaborate a proposed dehalogenation mechanism for BphK and provide insight into the plasticity of the H-subsite.

Glutathione S-transferases (GSTs)* transform a wide variety of electrophilic compounds in a reaction typically involving

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The atomic coordinates and structure factors (code 2D5A and 2GD5) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

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The abbreviations used are: GST, glutathione S-transferase; PCB, polychlorinated biphenyls; HOPDA, 3-chloro-2-hydroxy-6-oxo-6-phenyl-2,4-dienoate; TCHQ, tetrachlorohydroquinone; MES, 2-morpholinoethanesulfonic acid; CDNB, 1-chloro-2,4-dinitrobenzene; GS, deprotonated GSH.

GSH (γ-L-Glu-L-Cys-Gly) conjugation (1). Best known for their roles in detoxification in eukaryotes, GSTs are also broadly distributed in prokaryotes as homo- or heterodimers with ~25-kDa subunits (2). To date, GSTs have been grouped into at least 12 distinct classes on the basis of substrate specificity and primary structure (3). The nomenclature is in flux due to the continued identification of new GSTs, particularly in prokaryotes (3, 4). Despite low sequence identity, all GSTs share a conserved two-domain structure (5) in which the N-terminal domain consists of a central core of a mixed four- or five-stranded β sheet buttressed by a pair of α helices, and the C-terminal domain is entirely β helical (1, 6, 7). The active site architecture is formed in part from two binding subsites located at the interface of two adjacent monomers. The G-subsite binds GSH, and the H-subsite binds hydrophobic electrophiles (8).

Most eukaryotic GSTs consume a single GSH that is converted to an electrophilic substrate during catalytic turnover (1). Bacterial GSTs function in a wide range of detoxification and catabolic pathways (9–11). Moreover, they display considerable diversity with respect to the utilization of GSH. For example, enzymes such as maleylacetocetate and maleylpyruvate isomerases utilize but do not consume GSH (12). Interestingly, some bacterial GSTs, such as tetrachlorohydroquinone (TCHQ) dehalogenase from Sphingomonas chlorophenolica, consume two GSH equivalents (13). The first GSH is used in the dehalogenation step leading to the formation of a mixed disulfide with the enzyme. The second equivalent of GSH is used to regenerate the free enzyme, yielding a disulfide-bridged product, GSSG. In a variation of this theme, the catabolism of lignin derivatives by Sphingomonas paucimobilis SYK-6 also consumes two GSH molecules but requires two enzymes to do so; LigF and LigG catalyze glutathione conjugations similar to eukaryotic enzymes and GSSG formation, respectively (14, 15). All of these GSTs are thought to involve GSH conjugation. However, their structural basis is poorly understood.

Insights obtained from the reported crystal structures of bacterial GSTs (8, 16) are limited as the second physiological substrates for these enzymes are unknown. The structure of PmGST B1-1 from Proteus mirabilis reveals the binding of GSH to the G-subsite with the thiol group covalently attached to Cys-10 (8). PmGST B1-1 binds antibiotics such as tetracyclines and rifamycin but at a site remote from the H-subsite (17). Similarly, the structure of an Escherichia coli GST was determined in a complex with the inhibitor glutathione sulfonate (N-(N-γ-L-glutamyl-3-sulfo-L-alanyl)-glycine) (16), a GSH ana-
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logue. In this structure, Cys-10 and His-106 form hydrogen bonds to the glutathione sulfonate inhibitor; however, the bulky sulfonate group is unlikely to mimic a transition state (16). In many eukaryotic GSTs, Tyr-5 and Ser-11 are catalytically essential residues that are located in proximity to the sulfur atom of the bound GSH (8, 18). In the bacterial GSTs, these residues are not in contact with the bound GSH. Instead, two conserved residues, Cys-10 and His-106, interact directly with the GSH substrate and are ascribed catalytic roles (8). Site-directed mutagenesis studies using 1-chloro-2,4-dinitrobenzene (CDNB) as a reporter substrate failed to identify active site residues that activate the glutathione (19, 20).

The bphK gene of Burkholderia xenovorans LB400 encodes a GST that is part of the bph (biphenyl catabolic) pathway (21). The latter is responsible for the aerobic catabolism of biphenyl and the transformation of some polychlorinated biphenyls (PCBs). Biphenyl catabolism is typical of the aerobic catabolism of aromatic compounds in that it proceeds via a catechol, in this case 2,3-dihydroxybiphenyl (22). The extradiol cleavage of the latter produces 2-hydroxy-6-oxo-6-phenyl-2,4-dieneoate (HOPDA), which is hydrolyzed to benzoate and 2-hydroxy-penta-2,4-dienoate in a reaction catalyzed by BphD. BphD is competitively inhibited by 3-Cl and 4-Cl HOPDAs (23). BphK is not essential for growth of B. xenovorans LB400 on biphenyl (24); however, BphK was shown recently to dechlorinate 3-Cl and 5-Cl HOPDAs (25). This activity would alleviate inhibition of the bph pathway by these PCB metabolites and likely contributes to the potent PCB-degrading ability of B. xenovorans LB400 (25). The reductive dehalogenation catalyzed by BphK is similar to that of TCHQ dehalogenase from S. chlorophenolica (26) in that it consumes two GSH equivalents (13). The BphK-catalyzed dehalogenation of 3-Cl HOPDA is proposed to proceed via either a simple addition of GSH followed by elimination of chloride or an S_{c2}-mediated displacement (25). The second equivalent of GSH is used to regenerate the free enzyme.

To investigate the unusual catalytic mechanism of BphK and to probe the structural basis for catalysis in bacterial GSTs, we have obtained structural data for two ternary complexes of this enzyme. Each structure provides unique insights into the catalytic cycle of GSTs. In one structure, a HOPDA molecule (the dechlorinated product of 3-Cl HOPDA) and a GSH molecule are bound at the active site (BphK-GSH-HOPDA). In the other structure, two GSH molecules are bound at the active site of the enzyme (BphK-(GSH)_2). These structures support and extend the recently proposed mechanism (25). Furthermore, the structures reveal surprising flexibility of the active site to accommodate both HOPDA and GSH molecules at the H-subsite.

### EXPERIMENTAL PROCEDURES

BphK was heterologously expressed in E. coli GJ1158 and purified using glutathione-Sepharose 4B resin as described previously (25). The stock BphK solution used in crystallization experiments contained 65 mg/ml of protein, 50 mM Tris, pH 8, and 10 mM GSH. The reservoir solution was 0.8–1.2 M K^+/Na^+ tartrate, 0.1 M MES, pH 6. BphK was crystallized by hanging drop vapor diffusion using a 1:1 ratio of reservoir to protein, and crystals grew within a week.

#### TABLE 1

**Data collection and refinement statistics**

| Data                      | BphK-GSH-HOPDA | BphK-(GSH)_2 |
|---------------------------|----------------|--------------|
| Resolution (Å)            | 50-2.1 (2.1–2.18) | 50-2.1 (2.1–2.18) | |
| Space group               | R3             | R6           | |
| Unit cell dimensions (Å)  | a = b = 112.2, c = 221.9 | a = b = 103.4, c = 262.3 |
| Completeness (%)          | 94.6 (88.8)    | 97.0 (95.4)  | |
| Average I/σ               | 21.4 (1.6)     | 24.5 (6.2)   | |
| Redundancy                | 4.5 (2.9)      | 4.6 (3.4)    | |
| Rmerge                    | 0.058 (0.54)   | 0.061 (0.58) | |

- Numbers in parentheses correspond to those in the highest resolution shell.
- Rmerge = Σ[I(–I)]/Σ[I], where for each unique reflection, I is the observed intensity for each reflection merged, and <I> is the mean intensity.
- r.m.s. deviations from ideality
- Bond lengths (Å)
- Bond angles (degrees)

Two co-crystallization experiments were performed: one with HOPDA and GSH and one with GSH alone. In the first experiment, the protein was diluted 1:1 with a buffer containing 2 mM HOPDA, 10 mM glutathione, and 50 mM Tris, pH 7. Before mounting, the crystals were soaked for an additional 30 min in 10 mM HOPDA, produced as described previously (23). In the second co-crystallization experiment, the protein was diluted 1:1 with a buffer containing 10 mM GSH and 50 mM Tris, pH 7. All crystals were transferred into mother liquor supplemented to 30% glycerol as a cryo-protectant.

The BphK-GSH-HOPDA co-crystals grew in space group R3 with unit cell dimensions a = b = 112.2 Å, c = 221.9 Å, and two dimers in the asymmetric unit (A-B, C-D). The BphK-(GSH)_2 co-crystals grew in space group P6_5 with unit cell dimensions a = b = 103.4 Å, c = 282.3 Å and three dimers (A-B, C-D, E-F) in the asymmetric unit. Data sets were collected at the Stanford Synchrotron Radiation Laboratory (SSRL) BL 9-1 to 2.1 Å resolution. The data were processed with DENZO and scaled with SCALPACK (27). The overall Wilson R-factors were 42 and 30 Å^2 for the BphK-GSH-HOPDA and BphK-(GSH)_2 structures, respectively. The structure of a bacterial GST from S. paucimobilis (Protein Data Bank entry code 1F2E) with all non-conserved residues set to alanines was used as a starting model for molecular replacement using the program AMoRe with data to 2.8 Å resolution (28). The two proteins share 54% amino acid sequence identity, and the correlation coefficient and R-factor of the top solution were 0.59 and 0.53, respectively. Data collection and refinement statistics are presented in Table 1.

The BphK-GSH-HOPDA structure was refined using Refmac5 (29). Five percent of the data were set aside to calculate the free R-factor (30). Models were built and visualized using the program O (31). Electron density was observed at each of the four G-subsites, and GSH molecules were modeled at full occupancy. Electron density for the HOPDA molecule was present in all monomers and when modeled at half-
FIGURE 1. **Crystal structures of BphK ternary complexes.** 

**A**, stereo image of the bound ligands at the active site of the BphK-GSH-HOPDA complex. Residue Cys-10, GSH, and HOPDA molecules are shown as ball-and-stick models. 2Fo − Fc (gray) and omit Fo − Fc (green) electron density maps are contoured at 0.9 and 2.9 σ, respectively. 

**B**, in the active site of the BphK-(GSH)₂ complex, the 2Fo − Fc electron density map (gray) is contoured at 0.9 σ. An omit Fo − Fc electron density map of the GSH molecule at the H-subsite is shown in green and contoured at 2.5 σ. In panels **A** and **B**, carbon atoms are colored orange, nitrogen atoms are colored blue, oxygen atoms are colored red, and sulfur atoms are colored yellow. Ribbon diagrams of the BphK-GSH-HOPDA and BphK-(GSH)₂ homodimers are in panels **C** and **D**, respectively. The two monomers forming a functional dimer along (left) and perpendicular to (right) the local 2-fold axis are colored in slate and cyan. GSH and HOPDA molecules are shown as ball-and-stick models. Carbon atoms of the molecule bound to the G-subsite and the H-subsite are colored yellow and orange, respectively. Oxygen, nitrogen and sulfur atoms are colored red, blue, and green, respectively. Figs. 1 and 2 were generated using the programs Bobscript (40), Molscript (41), Raster3D (42), and PyMOL (43).
occupancy, refined with an average B-factor of 42 Å². The final model of the BphK-GSH-HOPDA complex contains four monomers, four GSH molecules, four HOPDA molecules, and 498 water molecules. The BphK-(GSH)₂ structure was refined using CNS (32). The crystal was almost perfectly twinned (33); the twin fraction for refinement and map calculations was refined to 0.45. Each of the six monomers contains one GSH substrate at the G-subsite. In three out of the six monomers, additional electron density was observed at the H-subsite and was modeled as a second GSH molecule. The occupancy of these second GSH molecules in monomers B, D, and F was set to 0.5, and the average B-factors were refined to 35 Å². The final model of the BphK-(GSH)₂ structure consists of six peptide chains, nine GSH molecules, and 216 water molecules. Ramachandran plots of the main chain torsional angles in each structure show that 95.0% of the residues are in the most favored regions, and none are observed in the disallowed regions as defined by PROCHECK (34).

RESULTS

The Overall Structure—The crystal structures of BphK-GSH-HOPDA and BphK-(GSH)₂ have two and three dimers in the asymmetric unit, respectively. The overall fold of BphK in the two structures is essentially identical (Fig. 1, C and D). The root mean square deviation of the CA positions on superposition between each of the respective dimer pairs is less than 0.2 Å. Moreover, the main protein fold and subunit arrangement in the dimer are similar to those of other GSTs (1, 6, 7). Briefly, the monomer is comprised of two domains. The N-terminal domain (residues 1–79) contains a /H9252/H9251/H9252/H9252 module (residues 1–38) connected to a /H9252/H9252/H9251 module (residues 54–77) via a long surface-exposed loop. A 12-residue linker connects the N-terminal domain to the C-terminal domain (residues 89–202). The latter domain contains five /H9251 helices. Each homodimer contains two equivalent active sites, which are found at clefts formed at the interface between the monomers (Fig. 1, C and D). A small portion of the active site cleft is formed by residues derived from the other monomer in the dimer (Fig. 2, A and B). The active site is comprised of two subsites, a common GSH binding site (G-subsite) and an H-subsite that typically binds the electrophilic second substrate of the reaction.

BphK-GSH-HOPDA Complex—Within the enzyme-GSH-product ternary complex, the GSH molecule is tightly held in an extended conformation at the G-subsite by 10 polar interactions (Fig. 2A, Table 2) and a single hydrophobic interaction with Tyr-51. The average B-factor of the atoms of the six GSH molecules is 41 Å², indicating that these sites are fully Occupancy, refined with an average B-factor of 42 Å². The final model of the BphK-GSH-HOPDA complex contains four monomers, four GSH molecules, four HOPDA molecules, and 498 water molecules. The BphK-(GSH)₂ structure was refined using CNS (32). The crystal was almost perfectly twinned (33); the twin fraction for refinement and map calculations was refined to 0.45. Each of the six monomers contains one GSH substrate at the G-subsite. In three out of the six monomers, additional electron density was observed at the H-subsite and was modeled as a second GSH molecule. The occupancy of these second GSH molecules in monomers B, D, and F was set to 0.5, and the average B-factors were refined to 35 Å². The final model of the BphK-(GSH)₂ structure consists of six peptide chains, nine GSH molecules, and 216 water molecules. Ramachandran plots of the main chain torsional angles in each structure show that 95.0% of the residues are in the most favored regions, and none are observed in the disallowed regions as defined by PROCHECK (34).

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BphK-GSH-HOPDA Complex—Within the enzyme-GSH-product ternary complex, the GSH molecule is tightly held in an extended conformation at the G-subsite by 10 polar interactions (Fig. 2A, Table 2) and a single hydrophobic interaction with Tyr-51. The average B-factor of the atoms of the six GSH molecules is 41 Å², indicating that these sites are fully occupied
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Table 2

| Interactions | Distances^a | BphK-GSH-HOPDA | BphK-(GSH)_2 |
|--------------|-------------|----------------|--------------|
| BphK-BphK    |             | 4.17–4.30      | 3.83–3.93    |
| His-106 NE2–Cys-10 SG | 2.73–2.83 | 2.57–2.65      |
| His-106 ND1–Tyr-157 OH |  |

BphK-G-subsite ligand

| His-106 NE2–GSH SG2 | 3.60–3.79 | 3.72–3.85 |
| His-106 ND2–GSH O12 | 3.40–3.57 | 3.34–3.51 |
| Asn-99 NZ–GSH O21 | 3.09–3.19 | 3.08–3.20 |
| Glu-65 OG–GSH O11 | 2.70–2.79 | 2.68–2.83 |
| Ser-103 OE2–GSH N1 | 2.97–3.07 | 3.00–3.05 |
| Gly-66 N–GSH O11 | 3.00–3.05 | 3.00–3.20 |
| Glu-104 O1–GSH N1 | 2.59–2.71 | 2.77–2.94 |
| Val-52 G–GSH N2 | 2.77–2.90 | 2.68–2.93 |
| Lys-110 O–GSH O2 | 2.78–2.87 | 2.89–3.19 |
| His-106 NE2–HPD OA1 | 3.28–3.43 | 3.27–3.44 |
| Val-52 G–GSH N2 | 2.77–2.90 | 2.68–2.93 |
| Lys-110 O–GSH O2 | 2.78–2.87 | 2.89–3.19 |
| His-106 NE2–GSH O31 | 2.96–4.80 | 2.64–2.98 |

BphK-H-subsite ligand

| His-106 NE2–GSH O31 | 3.28–3.44 | 3.27–3.44 |
| Ser-103 OG–GSH O31 | 2.38–3.48 | 2.38–3.84 |
| His-106 NE2–HPD OA2 | 3.28–3.43 | 3.27–3.44 |
| Ser-103 OG–GSH O32 | 2.66–3.19 | 2.66–3.19 |
| Ser-103 OG–HPD OA2 | 2.43–3.64 | 2.43–3.64 |

Ligand-Ligand

| GSH SG2(G-subsite)–GSH SG2(H-subsite) | 4.07–4.66 |
| GSH SG2(G-subsite)–HPD CA3 | 3.16–3.41 |

^a Range of distances over all the monomers in the asymmetric unit with both ligands bound.

Discussion

Structural Plasticity in the H-Subsite—BphK is typical of GSTs in that it binds a GSH molecule at its G-subsite and a second substrate at its H-subsite. Our crystal structures demonstrate the ability of the H-subsite of the enzyme to accommodate both a product and subsequently a second GSH molecule to accomplish the reductive dechlorination reaction. This flexibility seems to be linked to two properties of the H-subsite: a predominantly hydrophobic character and strategically positioned Ser and His residues. Thus, both ligands make hydro-

and that the bound molecules are well ordered. The glycy1 moiety is orientated toward the surface of the protein, whereas the γ-glutamyl moiety is directed toward the core of the protein. The main interactions of the glycy1 moiety are three hydrogen bonds involving Lys-35 and Val-52. The thiol of GSH forms a hydrogen bond with Cys-10 SG (3.0–3.1 Å) and is near to His-106 NE2 (3.6–3.79 Å). Three hydrogen bonds from the bound GSH are formed with residues from the adjacent monomer: Asn-99, Ser-103, and Glu-104. In addition, the γ-glutamyl moiety interacts with Glu-65 (a hydrogen bond) and Tyr-51 (hydrophobic interactions).

After refinement with GSH molecules bound at each of the four G-subsites and computing a difference map, strong electron density was observed in all of the monomers at the H-subsite adjacent to the modeled GSH. This density was fit and refined with a HOPDA molecule (Fig. 1A) in the trans transoid configuration of the enol tautomer, the predominant form in solution (23, 35). A HOPDA restraint library was created using the Monomer Library Sketcher within Refmac5. The enol tautomer is defined by CA2=CA3 and CA4=CA5 double bonds, whereas the keto tautomer is defined by a CA3=CA4 double bond (Fig. S2). When the keto form of the HOPDA molecule was modeled, the fit of the dioneoyl moiety was poor due to the disallowed rotation of the CA3=CA4 double bond.

When compared with the G-subsite, the H-subsite is more solvent-accessible, and BphK makes fewer direct interactions with the HOPDA product. Nonetheless, the HOPDA molecule refines with B-factors similar to those of the BphK atoms that form the H-subsite. The H-subsite is lined mainly with aromatic residues: Tyr-167, Phe-113, and Trp-164 (Fig. 2A). The base of this subsite is constructed from the peptide chain of Gly-8 and Ala-9. Only three hydrogen bonds are formed with the HOPDA molecule, and all involve the carboxylate oxygen atoms. Two of these hydrogen bonds are with Ser-110 OG, and one is with His-106 NE2 (Table 2).

The single direct contact involving the two bound molecules in each subsite is a van der Waals interaction (~3.3 Å) between the GSH thiol and carbon 3 of the HOPDA molecule (Fig. 2A). Importantly, this carbon is chlorinated in the physiological substrate.

BphK(GSH)_2 Complex—As in the BphK-GSH-HOPDA structure, the BphK-(GSH)_2 structure shows well defined electron density for a GSH molecule found in the G-subsite (Fig. 1B). The GSH molecule is in an extended conformation similar to the binding mode of the GSH molecule in the BphK-GSH-HOPDA structure. A shift of the glycy1 moiety (corresponding to ~3.0 Å for the carboxylate oxygen atoms) is observed between the two structures (Fig. 2C). All the G-subsite protein-GSH interactions observed in the BphK-GSH-HOPDA structure are conserved in the BphK-(GSH)_2 structure (Table 2). An additional hydrogen bond is formed between Lys-107 NZ and OE1 of the GSH molecule bound at the G-subsite. More importantly, the position of the GSH sulfur atom with respect to Cys-10 SG and His-106 NE2 is similar in the two structures (distances of 3.1–3.4 and 3.7–3.9 Å, respectively). When modeled in at full occupancy, the average B-factor of the GSH molecules was refined to 31 Å^2.

A second GSH molecule is observed at the H-subsite in three out of the six monomers (Figs. 1B and 2B). Although the second GSH was modeled at partial occupancy, clear density extends over the molecule, and the sulfur atom occupies the highest peak in an omit difference map (Fig. 1B). Similar to the binding of the HOPDA molecule, this second GSH molecule is sandwiched between the first GSH in the G-subsite and a hydrophobic pocket created by Tyr-167, Trp-164, and Phe-113. Only two hydrogen bonds are observed between the second GSH molecule and the protein. These interactions are between the glycy1 carboxylate oxygen O31 of the GSH molecule and the residues His-106 and Ser-110 (Fig. 2B, Table 2).

This second GSH molecule is also bound in an extended conformation; however, the glycy1 moiety is orientated toward Cys-10, and the γ-glutamyl moiety lies at the surface of the protein. As a result, the second GSH molecule is approximately antiparallel to the GSH molecule in the G-subsite such that the GSH sulfur atoms are 4.1–4.7 Å apart. Notably, His-106 NE2 serves as a bridge between the two GSH molecules by forming hydrogen bonds to the SG2 atom of GSH in G-subsite and O31 of GSH bound at the H-subsite (Table 2).

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The HOPDA complex, the side chain of Tyr-167 rotates 80° about To accommodate the HOPDA phenyl group in the BphK-GSH-configuration of the H-subsite residues involve Tyr-167 and Phe-113. Differences in the configuration between these enzymes.

The relative paucity of specific interactions between the shallow and open H-subsite and each of the HOPDA and GSH molecules suggests that these ligands bind weakly. Indeed, these ligands were modeled at half-occupancy in the crystal structures despite being added in excess in the co-crystallization experiments. Consistent with these observations, BphK has a moderate specificity ($K_m$) for 3-Cl HOPDA, suggesting that this enzyme is not highly adapted to its role in degrading PCBs (25). Weak binding is consistent with the broad substrate specificity typical of GSTs. For example, TCHQ dehalogenase binds the second GSH with a $K_D$ value of 19 mM (36). In addition, the TCHQ substrate inhibited the formation of GSSG, suggesting that the aromatic substrate has higher affinity for the H-site (36). Thus, the observation of HOPDA binding to BphK in the presence of excess GSH is not surprising based on analogy between these enzymes.

Dehalogenation Reaction—Fortin et al. (25) proposed a mechanism for the BphK-catalyzed dehalogenation of 3-Cl and 5-Cl HOPDAs. This mechanism is similar to the better supported mechanism of TCHQ dehalogenase (26, 36) and can be divided into two reaction demarcated by the formation of a mixed disulfide between the enzyme and GSH. The first half-reaction was proposed to proceed via one of two routes: the addition of GS$^-$ at carbon 3 of HOPDA followed by elimination of Cl$^-$ or the tautomerization of the enol(ate) followed by an $S_{N}$2 nucleophilic attack by GS$^-$ at carbon 3 to release Cl$^-$. Either mechanism gives rise to a GS-HOPDA adduct from which the dehalogenated HOPDA molecule is released by nucleophilic attack of Cys-10 SG on the sulfur of the GS$^-$ moiety, yielding a mixed disulfide linkage. In the second half-reaction, regeneration of the resting enzyme is initiated by the binding of a second GSH at the H-subsite for a final disulfide exchange. The result is a reduced sulfhydryl at Cys-10 and the formation of GSSG that leaves the active site (Fig. 3).

The binding mode and conformation of the GSH and HOPDA molecules at the H-subsite provides additional insights into the proposed mechanism. Although the dechlorination product is bound in the BphK-GSH-HOPDA complex, this structure is likely analogous to that for the binding of the 3-Cl substrate. Accordingly, a 3-Cl substituent would be orientated toward the indole ring of Trp-164. The nearest atom of the HOPDA molecule to the GSH thiol is carbon 3 (3.2–3.4 Å). Furthermore, the Burgi-Dunitz angle (~95°) is reasonable but not optimal (109°) for a nucleophilic substitution to proceed on an sp$^2$ carbon 3 (37). Carbon 5 is more distant from the nucleophile (~4.6 Å), but the enzyme readily dechlorinates 5-Cl HOPDA (25). Variation in the proposed mechanism or an alternate substrate binding mode may provide an explanation for the difference in catalytic activity on substrates chlorinated at different sites.

Of the two mechanisms proposed for the first half of the reductive dehalogenation reaction (25), the structural data are most consistent with those involving tautomerization and nucleophilic substitution (Fig. 3). In particular, the elimination mechanism involves attack of GS$^-$ on the sp$^2$ hybridized carbon 3 to form a tetrahedral intermediate resulting in the generation of a second negative charge on the carboxyl end of the HOPDA molecule. However, the structural data indicate that only Ser-110 coordinates to the HOPDA carboxylate and that this residue would poorly stabilize two negative charges on the carboxylate (Figs. 2A and 3). By contrast, the tautomerization-substitution mechanism eliminates the need for compensating groups to stabilize the additional negative charge on the carboxylate. Inspection of the H-subsite reveals no residue that could easily serve as a proton donor to catalyze the initial tautomerization. However, an intriguing possibility is that the HOPDA molecule is bound as the enol and that the 2-hydroxyl is the source of the proton at carbon 3. The pK$\alpha$ of 3-Cl HOPDA in solution is 6.1 (24) but may be perturbed by binding to the enzyme. Finally, the proposed mechanism for the first half-reaction of BphK differs from that of TCHQ dehalo-
genase, which involves chloride elimination from the aromatic ring followed by GSH attack (36).

The BphK-(GSH)₂ structure gives insight into the second half-reaction to regenerate the resting enzyme. In this structure, the distances between the sulphydryls of Cys-10 and the two bound GSH molecules (≥3.3 Å) indicate that all three sulfurs are reduced (Fig. 2B). Nonetheless, the sulfur atoms are aligned such that a disulfide could be formed easily either between Cys-10 and the GSH molecule (at the G-subsite) or between the two GSH molecules by small rotations of the χ₁ angles. His-106 is positioned to stabilize the anticipated development of a negative charge on Cys-10 that would result from a disulfide exchange reaction.

Comparison with Other β-GSTs—The overall fold of BphK is similar (CA root mean square deviation of <1.5 Å) to those of E. coli GST (16), PmgST B1-1 from P. mirabilis (8), and GST from S. paucimobilis (reported in Protein Data Bank entry 1F2E). Recently, site-directed mutagenesis studies on conserved structural motifs between residues Phe-151 and Ala-156 have identified their importance in protein folding and stability (38, 39). Conservation of structure is greater in the G-subsite, and the binding mode of a GSH molecule in this subsite of BphK is similar to that of PmgST B1-1 (8) and to the glutathione sulfonate analogue bound to the E. coli GST structure (16). A multiple sequence alignment (Fig. S1) reveals that BphK residues involved in the formation of the G-subsite are highly conserved among bacterial GSTs sharing at least 40% amino acid sequence identity. Residues Cys-10, Val-52, Glu-65, His-106, and Tyr-157 are absolutely conserved. Conservative substitutions are observed for Asn-99, Ser-103, and Glu-104. Position 51 has significant variability in the alignment and results in the greatest difference in the G-subsites between the BphK and the PmgST B1-1 structures. In PmgST B1-1, Gln-51 forms a weak hydrogen bond (3.6 Å) to the GSH, whereas in BphK, a Tyr at the equivalent position forms a hydrophobic interaction. A subtle difference between the two G-subsites is the distance between NZ of Lys-107 and OE₁ of GSH. In the BphK-GSH-HOPDA complex, this distance is 3.7–3.9 Å, whereas in the PmgST B1-1 structure, that same interaction is a hydrogen bond (3.3 Å). Interestingly, this hydrogen bond is conserved in the BphK-(GSH)₂ structure (3.4 Å).

The hydrophobic character of the H-subsite in BphK is also conserved among bacterial GST families. Three conserved small residues (Pro-7, Gly-8, Ala-9) form one side of this subsite, and two conserved large hydrophobic residues (Phe-113, Trp-164) form the other side. Tyr-167 varies substantially and is substituted by histidine, arginine, or alanine in other bacterial GSTs. The identity of the residue at position 110 does not vary quite as much, being serine, glycine, valine, or threonine. The ternary complexes reported here suggest that Ser-110 and His-106 are important in defining H-subsite substrate specificity. The naturally observed substitutions at position 110 allow for variation in substrate specificity without sterically precluding the binding of a second GSH molecule.

In PmgST B1-1 and the GST from S. paucimobilis, a disulfide bond is formed between the Cys-10 sulfur atom and SG₂ of the GSH (8). The formation of this mixed disulfide provides a precedent for the intermediate formed at the end of the first half-reaction in the BphK dehalogenation mechanism (Fig. 3). Cys-10 SG and His-106 have been hypothesized to activate GSH for the initial nucleophilic attack in related systems (8, 16); however, in our structures with reduced GSH in the active site, the hydrogen bonds to the GSH sulfur atom are weak, suggesting that little activation is necessary. Mutation of Cys-10 to Ser or Ala of PmgST B1-1 did not significantly reduce activity toward CDNB, a synthetic reporter substrate (19). Furthermore, the conservative mutation of His-106 to Asn had a greater impact on substrate binding than on catalysis using the same reporter substrate (20). Finally, mutational analysis of Ser, Tyr, and Thr residues in the active site failed to identify catalytically essential groups for GSH activation toward CDNB (19).

The reaction of some bacterial β-GSTs with CDNB is likely analogous to that observed in the larger GST superfamily, whereby a single GSH molecule is consumed and transferred giving a conjugated product (5). In contrast, the mechanism for BphK is a reductive dehalogenation requiring two GSH equivalents; the transferase reaction constitutes only a part of the first half-reaction. Thus, use of CDNB as a reporter substrate provides information only for the first two steps of a more complex mechanism (Fig. 3). Interestingly, mutation of an active site Cys in TCHQ dehalogenase does not prevent the initial GSH nucleophilic attack on substrate but does reduce overall activity 10-fold (36). The proposed mechanism for BphK and the crystal structures presented here suggest that conserved active site residues are not critical for the initial GSH transferase reaction but facilitate the subsequent catalytic steps.

Mutational studies on BphK directed at Cys-10 and His-106 are not yet available; however, a conserved region from Ser-152 to Leu-158 was targeted in a recent study (39). In the crystal structure of BphK, the phenolic oxygen of Tyr-157 within this region forms a hydrogen bond to ND₁ of His-106, which likely determines the orientation of the imidazole ring. An analysis of activity toward CDNB and 4-chlorobenzoate, a proposed alternative physiological substrate for BphK, showed that this region is not directly involved in substrate recognition and catalysis (39). The chlorinated HOPDAs are 1000-fold better substrates than 4-chlorobenzoate (25), and the reaction for the latter is likely to be analogous to that of CDNB. Further evaluation of the role of Cys-10 and His-106 awaits studies using chlorinated HOPDA substrates or appropriate analogues. Conservation of these residues in the β-GSTs suggests that a reductive mechanism involving the consumption of two GSH molecules may be a general feature of this bacterial class.

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