ClearANCE OF aromatic carbon–chlorine bonds is critical for the degradation of toxic industrial compounds. Here, we solved the X-ray crystal structure of chlorothalonil dehalogenase (Chd) from *Pseudomonas* sp. CTN-3, with 15 of its N-terminal residues truncated (Chd\(^{\text{T}}\)), using single-wavelength anomalous dispersion refined to 1.96 Å resolution. Chd has low sequence identity (<15%) compared with all other proteins whose structures are currently available, and to the best of our knowledge, we present the first structure of a Zn(II)-dependent aromatic dehalogenase that does not require a coenzyme. Chd\(^{\text{T}}\) forms a “head-to-tail” homodimer, formed between two \(\alpha\)-helices from each monomer, with three Zn(II)-binding sites, two of which occupy the active sites, whereas the third anchors a structural site at the homodimer interface. The catalytic Zn(II) ions are solvent-accessible via a large hydrophobic (8.5 \(\times\) 17.8 Å) opening to bulk solvent and two hydrophilic branched channels. Each active-site Zn(II) ion resides in a distorted trigonal bipyramid geometry with His\(^{117}\), His\(^{257}\), Asp\(^{116}\), Asn\(^{216}\), and a water/hydroxide as ligands. A conserved His residue, His\(^{114}\), is hydrogen-bonded to the Zn(II)-bound water/hydroxide and likely functions as the general acid-base. We examined substrate binding by docking chlorothalonil (2,4,5,6-tetrachloroisophtalonitrile, TPN) into the hydrophobic channel and observed that the most energetically favorable pose includes a TPN orientation that coordinates to the active-site Zn(II) ions via a CN and that maximizes a \(\pi-\pi\) interaction with Trp\(^{227}\). On the basis of these results, along with previously reported kinetics data, we propose a refined catalytic mechanism for Chd-mediated TPN dehalogenation.

Chlorinated aromatic hydrocarbons such as polychlorinated biphenyl, chlorobenzenes, and atrazine are important industrial starting materials for the manufacture of dyes, drugs, and pesticides/fungicides (1, 2). Aromatic carbon–chlorine bonds are typically very stable, and hence, these compounds persist in soil and contaminate groundwater. Their low water solubility, coupled with their toxicity, makes them particularly important targets for environmental remediation (3). Enzymatic dehalogenation of chlorinated natural organic matter by organohalide-respiring bacteria provides more soluble compounds that are less likely to bioaccumulate and more likely to be susceptible to further degradation or cellular recycling and, thus, less toxic (4–6). Therefore, cleavage of aromatic carbon–chlorine bonds is critical for the degradation of toxic industrial compounds. Although enzymatic dehalogenation offers a possible bioremediation solution for chlorinated aromatic compounds, a major impediment to understanding their biological role and bioremediation uses is the lack of a detailed understanding of their catalytic mechanisms.

The prevailing dogma is that biological dechlorination reactions are catalyzed by either oxidative, reductive, or thiolytic dehalogenation processes (7–13). A relatively unknown biological dehalogenation process involves hydrolysis of a C-Cl bond (14). The chlorothalonil dehalogenase from *Pseudomonas* sp. CTN-3 (Chd) is a Zn(II)-dependent hydrolytic dehalogenase that selectively substitutes an aromatic chlorine-carbon moiety in chlorothalonil (2,4,5,6-tetrachloroisophthalonitrile, TPN) to an aromatic alcohol (4-hydroxylchloro-isophthalonitrile, 4-OH-TPN) (Fig. 1) (15). TPN is one of the most commonly used fungicides in the United States with more than 5,000,000 kg sprayed on crops and fruits each year (16–19). TPN has low solubility in water (100 mg liter\(^{-1}\)) but is strongly absorbed in soil, particularly soil with high organic matter such as those found in aquatic environments. TPN is highly toxic to fish and aquatic species, as well as birds and invertebrates (1, 15, 20). TPN is also a human skin and eye irritant that can cause severe gastrointestinal issues. Animal studies involving mice have shown that TPN can cause kidney cancer, so it has been classified by the U.S. Environmental Protection Agency as a probable human carcinogen (18). Given the widespread use of TPN and its toxicity, its biodegradation and environmental clean-up has become a topic of significant importance (22).

Several bacterial strains harbor Chd genes that exhibit remarkable sequence identity (>95%) (23–26). Chd was suggested to contain a conserved Zn(II)-binding domain similar to enzymes in the metallo-\(\beta\)-lactamase (MBL) superfamily and was proposed to be dimeric in solution (15). Although Zn(II) is required for Chd activity (26), its catalytic role remains unknown. A major limitation in understanding the catalytic mechanism of Chd and in developing small molecule catalysts that specifically target TPN, is a lack of knowledge regarding its structure and active site architecture. The only aromatic hydrodehalogenase that has been structurally characterized to date is atrazine chlorohydrolase from *Pseudomonas* sp. (AtzA;...
PDB code 4V1X at 2.2 Å), which contains an iron ion coordinated by four histidine residues, an aspartate, and a water molecule (27). Although both AtzA and Chd hydrolytically dehalogenate aromatic compounds, they have relatively low sequence identity (>19%) and different metal ion requirements. Moreover, AtzA is hexameric, whereas Chd is dimeric (15, 27).

Herein, we report the 1.96 Å X-ray crystal structure of a 15 residue N-terminally truncated Chd enzyme that is fully active, using single-wavelength anomalous dispersion (SAD). A truncated form of Chd (ChdT), which lacked the first 15 N-terminal residues, was designed and utilized because of poor solubility and stability of the WT Chd enzyme at the concentrations required for crystallization. The X-ray structure reveals that Chd is a homodimer with a structural Zn(II) residing at the dimer interface and a mononuclear, trigonal bipyramidal (TBP) Zn(II) active site with His117, His257, Asp116, Asn216, and water/hydroxide as ligands. Based on this structure, substrate model- and previous kinetic studies (26), a refined catalytic mechanism is proposed.

Results

Overall structure of Chd

Chd\(^\text{I}\) crystals with dimensions 0.1 × 0.2 × 0.3 mm belonged to the primitive orthorhombic space group P\(_2_1\_2_1\_2_1\) with unit cell parameters a = 90 Å, b = 80 Å, and c = 90 Å. The structure was solved \textit{via} SAD using a wavelength of 1.28 Å, close to the absorption edge of zinc. The final model was refined to a resolution of 1.96 Å with \(R_{\text{free}}\) and \(R_{\text{work}}\) values of 19.80% and 16.22%, respectively. Final data processing and refinement statistics are listed in Table 1. A single homodimer was built in an asymmetric unit. The final refined model has been deposited to the PDB under code 6UXU.

The two monomers of the homodimer display a 2-fold non-crystallographic rotational axis that passes through a noncatalytic structural Zn(II) site (Fig. 2A). From a side view (Fig. 2B), the structural Zn(II) ion is close to one side of the dimer surface. For convenience purposes, we refer to this side of the dimer protein as the “top” side, \textit{i.e.} where the structural Zn(II) ion resides, and the opposite side as the “bottom.” The crystallographic packing was analyzed using PISA (28, 29) in the CCP4 suite, and the observed homodimer in the asymmetric unit was predicted by PISA to be the only stable quaternary structure in solution. The total predicted accessible surface area is 23,000 Å\(^2\), whereas the predicted buried surface area is 1,800 Å\(^2\).

Each monomer contains a catalytic Zn(II) ion that is solvent-accessible from both the top and the bottom of the protein (Fig. 3). From the bottom, a Y-shaped channel exists with two branched channels that lead to the catalytic Zn(II) ions and share a single large (8.5 × 17.8 Å) opening to bulk solvent. On the other hand, the structural Zn(II) ion is not accessible to the bulk solvent despite being located close to the top surface. From the top, the structural features surrounding the structural Zn(II) ion help to form two bent channels that provide access the catalytic Zn(II) ions.

The monomer protein fold is an αβα-sandwich

The overall protein fold of the Chd\(^\text{I}\) monomer (Fig. 4A) was searched using the DALI server (30) for comparison against all known structures. Interestingly, known structures with any similarity to Chd were reported as either β-lactamases or alkyl-sulfatases (31–34). The most similar structures from each of these protein families, CphA and Pisa1, were selected for detailed comparison and structurally aligned with Chd\(^\text{I}\) (Fig. 4, B and C). The overall fold of the Chd\(^\text{I}\) monomer aligned well with the overall structure of the β-lactamase, CphA (PDB code 3IOG), except for an extra C-terminal motif in Chd\(^\text{I}\). The N-terminal portion (residues 1–277) of Chd forms an αβα-sandwich fold that is commonly observed in the β-lactamase superfamily; however, Chd\(^\text{I}\) also possessed an additional C-terminal motif (residues 278–327) that originates in the middle of a long helix that is not found in the CphA structure. On the other hand, when Chd\(^\text{I}\) was structurally aligned with one subunit of the alkylsulfatase, Pisa1 (PDB code 2YHE), the additional C-terminal motif observed in Chd\(^\text{I}\) was present in Pisa1, which is much larger in size than either Chd or CphA.

### Table 1

| Crystalllographic parameters and statistics for Chd\(^\text{I}\) (6UXU) |
|---------------------------------------------------------------|
| **Data processing**                                             |
| Space group                                                     | P\(_2_1\_2_1\_2_1\) |
| Cell dimension                                                  | a, b, c (Å)           | 90, 90, 90 |
| processed resolution (Å)                                        | 1.96                  |
| Resolution at 1/σ(l)                                            | 17.5 (1.1)            |
| Resolution at 1/σ(l) = 2*                                       | 1.98                  |
| Rmerge (%)                                                      | 8.7 (117.6)           |
| Rfree (%)                                                       | 2.7 (35.0)            |
| CC(p) (%)                                                       | (63.1)                |
| Completeness (%)                                                | 98.0 (87.6)           |
| Multiplicity                                                    | 7.2                   |
| No. reflections                                                 | 362,906               |
| No. unique reflections                                          | 50,386                |

| **Refinement**                                                   |
|----------------------------------------------------------------|
| Rwork/Rfree (%)                                                 | 15.79/19.75          |
| No. of atoms                                                    | Protein 4,693         |
|                                                             | Ligand 3              |
|                                                             | Water 379             |
|                                                             | B factors (Å\(^2\))   | 23.7                 |
|                                                             | Protein RMSD\(^\alpha\)  |
|                                                             | Bond lengths (Å)      | 0.010                |
|                                                             | Bond angles (deg)     | 1.014                |
|                                                             | Ramachandran plot (%) | 96.14                |
|                                                             | Most favored          | 3.19                 |
|                                                             | Allowed               | 0.67                 |

\(^\alpha\) Provided resolution at 1/\(σ\) = 2 for conventional assessment of data quality.

\(^\beta\) The values for the highest-resolution bin are in parentheses.

\(^\gamma\) Precision-indicating merging \(R\).

\(^\delta\) Pearson correlation coefficient of two “half” data sets.
The coordination geometry of the active site Zn(II) ion is identical in both monomers and the simulated annealing composite omit map around each metal ion and their coordinating residues are consistent with the active-site Zn(II) ion residing in a distorted trigonal bipyramid or TBP geometry (Fig. 5A). The catalytic Zn(II) ion ligands include the $N^\varepsilon$ nitrogens of His$^{117}$ (2.0 Å) and His$^{257}$ (2.0 Å) along with a water/hydroxide oxygen atom (2.6 Å) making up the equatorial positions, whereas a carboxylate oxygen atom of Asp$^{116}$ (2.1 Å) and a carbonyl oxygen atom of Asn$^{216}$ (2.2 Å) (Table 2 and Fig. 5A) constitute the axial ligands. The coordinating water/hydroxide forms hydrogen bonds with the axial ligands Asp$^{116}$ (2.6 Å) and Asn$^{216}$ (2.9 Å) (Fig. 5A).

Another active site residue, His$^{114}$, is observed in two alternate conformations supported by simulated annealing composite omit maps (Fig. 5B). In one conformation (a), the $N^\varepsilon$ nitrogen atom of the imidazole ring is hydrogen bound to the coordinated water/hydroxide oxygen atom (2.9 Å) and the nitrogen atom (2.7 Å) of the axial ligand, Asn$^{216}$. The second conformation (b) places the imidazole ring perpendicular to the imidazole ring in the (a) conformation, resulting in the loss of hydrogen-bonding interactions with both the coordinated water/hydroxide oxygen atom (5.4 Å) and possibly the nitrogen atom (3.1 Å) of the axial ligand Asn$^{216}$. The structural Zn(II) ion is coordinated by the $N^\varepsilon$ nitrogen atoms of His$^{143}$ (2.0 Å) and the carboxylate oxygen atom of Asp$^{146}$ (1.9 Å) from each subunit (Fig. 5, C and D), providing a tetrahedral coordination geometry.

**Role of the structural Zn(II) site**

The structural Zn(II) site ligands were eliminated by mutating both His$^{143}$ and Asp$^{146}$ to Ala. Removal of the structural Zn(II) site ligands had no apparent effect on ChdT activity, because the double mutant exhibited a $k_{cat}$ of 25, which is identical to that of WT Chd (15, 26). Surprisingly, homodimer formation was also not affected based on gel-filtration results (Fig. S1). However, elimination of the structural Zn(II) site did increase the $K_m$ value 2-fold ($K_m = 0.24 \pm 0.07 \mu M$) and severely affected protein stability because the purified mutant protein precipitated within ~2 h when incubated at 23–25 °C.

**Chd biological assembly in solution**

The intact molecular masses of both the truncated and structural Zn(II) Chd mutants were obtained by MALDI-TOF MS (Fig. 6). Masses consistent with homodimer formation were observed between 66 and 70 kDa. These MS data are consistent with the hydrodynamic volume for both samples determined by size-exclusion chromatography (Fig. S1), which indicate a molecular mass consistent with a homodimer in solution.
should also be noted that a small portion of ChdT fragmented (<8%) during MALDI-TOF MS analysis.

**Substrate docking**

Attempts to co-crystallize or soak crystals of ChdT with the substrate TPN or recently identified inhibitors were unsuccessful; therefore, substrate docking was utilized to examine possible substrate-binding poses. Substrate docking generated four different poses ranked by the docking scoring function, where a lower value indicates lower total energy after docking (Fig. 7) (35–37). The top ranked model (model 1) suggested a binding interaction between the nitrogen atom of a cyan group of TPN and the active-site zinc ion (2.3 Å), along with an energetically favorable π–π interaction between the TPN aromatic ring and that of the active-site residue Trp227. Interestingly, this pose preorganizes a possible transition state of TPN hydrolysis by placing the ortho carbon of TPN within ~3.0 Å of the O atom of the bound water/hydroxide. In each of the other poses, all possible TPN ligating groups are too far from the catalytic Zn(II) ion to be considered a bonding interaction (>3.5 Å). However, model 3 revealed a hydrogen bond between the nitrogen atom of a TPN cyan group and the amide nitrogen of the Zn(II) ligands Asn216 (2.3 Å). The resulting substrate position placed the ortho carbon of TPN within 3.1 Å of the O atom of the bound water/hydroxide. In contrast to model 1, TPN is rotated in this pose, which abolished the energetically favorable π–π interaction between the TPN aromatic ring and that of Trp227.

**Discussion**

Chd is a protein with an unusual sequence: the highest sequence similarity to a structurally characterized protein, glyoxylase II from Arabidopsis thaliana (PDB code 1XM8), is 14.7%. Here, a 1.96 Å resolution structure of ChdT, a truncated construct of Chd, is reported that reveals an αββα-sandwich fold that is also observed in the MBL superfamily (38). Chd is a “head-to-tail” homodimer, formed between two α-helices from each monomer (Fig. 4). Each monomer contains a mono-Zn(II) active site, and a third Zn(II) ion anchors an intersubunit structural site. Otherwise, and uniquely for enzymes with the same functionality, no coenzyme is associated with active Chd.

The mononuclear Zn(II) active site in each monomer of Chd is a characteristic shared with the B2 metallo-β-lactamases. The third Zn(II) ion in the Chd homodimer, which is coordinated by the N\(^{\text{ε}}\) nitrogen atom of a His\(^{143}\) and a carboxylate oxygen atom of Asp\(^{146}\) from each subunit, resides in a tetrahedral coordination geometry that is buried at the dimer interface and likely functions as a structural site. A ChdT\(^{H143A, D146A}\) construct predictably lacked the ability to bind this third zinc ion but, surprisingly, exhibited full catalytic activity, albeit with a 2-fold elevated \(K_m\). Additionally, the ChdT\(^{H143A, D146A}\) variant retained the homodimeric architecture of the native protein in solution. The most striking effect of the double mutation was markedly impaired stability in solution at ambient temperatures.

The X-ray structure of Chd revealed two previously unknown solvent-accessible channels in each monomer that likely provide different routes for substrate and products to access/leave the catalytic Zn(II) site (Fig. 3). Because TPN is hydrophobic in nature, hydrolysis of an ortho-chlorine produces more hydrophilic products: 4-OH-TPN, a chloride ion and a proton. Because the products of TPN hydrolysis vary greatly in solubility and size, it would be reasonable to have two sets of solvent channels capable of shuttling different reactants and products to and from the active site. The Y-shaped channel that originates on the bottom of the Chd homodimer contains a large opening (8.5 × 17.8 Å) with a channel that is largely hydrophobic. This Y-shaped channel likely serves to recognize substrate and guide TPN to the active site. On the other hand, water and H\(^{\text{+}}\) and/or chloride ions can migrate through the second active-site channel, which is more hydrophilic and resides on the top of the dimer. It has been reported that dimerization of SidA, which is an alkylsulfatase similar to Pisa1, ensures resistance to biocidal SDS at toxic concentrations (39). Given that TPN is toxic, Chd may have evolved as a dimer with two active-site channels to elicit a similar response.

As predicted by Wang et al. (15), the first His in the B2 β-lactamase MBL (His-Xaa-His–Asp-His) is replaced by a Ser (Ser\(^{112}\)) in Chd). This minor change in the B2 β-lactamase MBL active site may explain why a dinuclear Zn(II) site does not occur in both B2 enzymes and Chd. The single active-site Zn(II) ion in Chd resides in a distorted TBP geometry with His\(^{117}\), His\(^{257}\), Asp\(^{116}\), Asn\(^{216}\), and water/hydroxide as ligands (Fig. 5). Several potentially important hydrogen-bonding interactions also exist in the active site of Chd. Perhaps most notably, there is at least one interaction between the metal bound water/hydroxide oxy-
gen atom and the $N^e$ nitrogen atom of His$^{114}$. His$^{114}$ forms a strong hydrogen bond (−2.9 Å) with the equatorial metal bound water/hydroxide in Chd and also with the nitrogen atom of the axial Asn$^{216}$ ligand (−2.7 Å). Interestingly, His$^{114}$ clearly occupies two orientations in the structure, the second of which is perpendicular to the hydrogen bound conformation, which places the $N^e$ nitrogen atom out of hydrogen-bonding range of the metal-bound water/hydroxide (−5.4 Å) but close enough to the axial ligand Asn$^{216}$ (−3.1 Å) to possibly form a hydrogen bond. The functional role of His$^{114}$ and its flexibility is currently unknown but may play a catalytic role as a general acid/base. This is consistent with both ChpA and Pisa1, because His$^{114}$ is strictly conserved in both enzymes, and in ChpA, it was also proposed to be the general acid to protonate the leaving group (40).

Other potentially important hydrogen-bonding interactions occur between the Asn$^{216}$ carboxamide nitrogen proton and the metal bound water/hydroxide (2.9 Å) and the second axial Asp$^{116}$ ligand carboxylate oxygen, also with the metal bound water/hydroxide (2.6 Å). These hydrogen-bonding interactions...
likely assist in the deprotonation of the bound water to a nucleophilic hydroxide. Finally, a potentially important hydrogen-bonding interaction occurs between the \( \text{N}^+ \) proton of the His\(^{114} \) ligand with a side chain oxygen atom of Glu\(^{81} \) (\( \sim 2.7 \) Å), forming a Glu-His-Zn triad that has been postulated to decrease the Lewis acidity of zinc ions (41) and may further assist in facilitating the coordination of substrate or water to the active site Zn(II) ion. Similar arrangements and fully conserved Asp-His-zinc triads are observed for other hydrolases (42).

The discovery of a large hydrophobic channel on the bottom of the Chd homodimer that provides access to the catalytic Zn(II) ion suggests that it may serve as the initial substrate recognition site. To properly position an aromatic substrate, such as TPN, for catalysis the substrate would need to either bind to or near the active site Zn(II) ion. Interestingly, Trp\(^{227} \) resides in the hydrophobic channel \( \sim 9.1 \) Å from the active site metal ion. Previous mutagenesis studies indicated that Trp\(^{227} \) is catalytically essential (15), suggesting that it may be involved in recognizing and positioning TPN near the active site Zn(II) ion via a \( \pi-\pi \) interaction. To test this hypothesis, TPN was docked into the hydrophobic channel with the most energetically favorable pose occurring for a TPN orientation that maximizes the \( \pi-\pi \) interaction with Trp\(^{227} \), places a CN nitrogen atom within a bonding distance to the active site Zn(II) (\( \sim 2.3 \) Å), and expands its coordination geometry to octahedral (Fig. 7; pose 1). Because the C4 and C6 chlorines in TPN are chemically equivalent (homotopic) given the symmetry arising from the positions of the two CN substitutes, it does not matter which CN group binds to the active-site Zn(II) ion. Chd catalyzes the dehalogenation of only one chlorine, and the resulting product cannot serve as a substrate for Chd. In pose 1, the C4/C6 chlorine would be positioned so that once the hydrolysis reaction occurs, the newly formed chloride ion would be oriented toward the small backend channel for expulsion to the bulk solvent. Although a second pose was reasonably energetically favorable (Fig. 7; pose 2), TPN was rotated, thus abolishing the energetically favorable \( \pi-\pi \) interaction between the TPN aromatic ring and that of Trp\(^{227} \). However, this pose does reveal a potential hydrogen bond between the nitrogen atom of a TPN cyano group and the amide nitrogen Asn\(^{226} \), which places the ortho carbon of TPN within 3.1 Å of the O atom of the bound water/hydroxide.

In conclusion, the X-ray crystal structure of Chd\(^{5} \), in combination with previously reported kinetic data, allows us to refine our proposed catalytic mechanism for Chd (Fig. 8) (26). The first step in catalysis is likely recognition of TPN by the large hydrophobic channel that leads to the Zn(II) active site. In this scenario, TPN is positioned by a \( \pi-\pi \) interaction with Trp\(^{227} \) that was shown to be important for catalytic turnover (15). Next, a CN nitrite nitrogen atom of TPN coordinates to the active-site Zn(II), although no direct evidence exists to support this assumption at this time, expanding its coordination number from five to six. Binding of the substrate nitrite nitrogen has a 2-fold impact. First, it removes electron density from the aromatic ring activating the ortho carbon for nucleophilic attack (27, 43), and second, it activates the coordinated water/hydroxide to function as a nucleophile. Deprotonation of the metal-bound water molecule by His\(^{114} \), with assistance from the Zn(II) ligands Asp\(^{116} \) and Asn\(^{227} \), forms a nucleophilic hydroxide moiety that is consistent with the postulated \( pK_a \) of the zinc-bound water molecule (44). Once the zinc-bound hydroxide is formed, it can attack the activated ortho carbon of TPN, forming an \( \eta-1-\mu \)-transition state complex (45). Solvent kinetic isotope effect studies indicate that one proton is transferred in the transition state, likely because of the breaking of a water O–H bond and the formation of a protonated His\(^{114} \) (26). The observed conformational flexibility of His\(^{114} \) may facilitate the formation of products and enable their release. Once the products are released, the Zn(II)-bound water molecule is replaced. Although a mechanism could also be proposed where a nitrite nitrogen atom of TPN does not bind to the active site Zn(II) ion, there is no residue other than the Zn(II) ligand Asn\(^{226} \) that could reasonably form a hydrogen bond to the nitrite nitrogen of TPN, which would be important for activating the substrate for hydrolysis.

### Experimental procedures

**Materials**

NEB restriction enzymes, T4 DNA ligase, Agilent QuikChange II site-directed mutagenesis kit, nickel–nitrilotriacetic acid columns, Wizard\textsuperscript{\textregistered} SV gel and PCR clean-up systems, size exclusion columns 4-(2-hydroxyethyl)-1-piperazineethane-sulfonic acid, potassium bromide, potassium sodium tartrate, and TPN
were purchased commercially (Fisher Scientific) at the highest purity available. TEV protease (EC 3.4.22.44) was obtained via previously published methods (46).

**N-terminal sequence modification of Chd**

The first 15 residues of the N terminus of WT Chd were removed by PCR (sense primer, ACTCCATGGATGCGTGAATTTAG, and antisense primer, CATAGCTTTAATGGTGGATGATG). The modified Chd gene containing NcoI and HindIII restriction sites was amplified under standard PCR conditions for gene cloning (47). The PCR products were purified by agarose gel electrophoresis and Wizard SV gel and subcloned into pET28a/H11001, creating the new expression vector pET28a/H11001–ChdT. The modified sequence was verified by Functional Biosciences (Madison, WI, USA).

**Expression and purification of ChdT**

The plasmid containing ChdT was freshly transformed into BL21(DE3) competent cells (Stratagene), and a single colony was used to inoculate 50 ml of LB-Miller culture containing 50 μg/ml kanamycin with shaking overnight at 37 °C as previously described (26). Briefly, this culture was used to inoculate a 1-liter culture, and the cells were grown at 37 °C until an $A_{600\text{ nm}}$ of 0.8–1.0 was reached. The culture was cooled on ice, induced with 0.1 mM isopropyl β-D-1-thiogalactopyranoside supplemented with 0.05 mM ZnCl$_2$, and expressed at 25 °C for 16 h. The cells were harvested by centrifugation at 6370 × g and 4 °C for 10 min in a Beckman Coulter Avanti JA-10 rotor and resuspended in 20 mM Tris-HCl buffer, pH 7.0, containing 50 mM NaCl, 10 mM tris(2-carboxyethyl)phosphate, and 25 mM imidazole at a ratio of 5 ml/g of cells. The cells were lysed by ultrasonication (Misonix Sonicator 3000) for 4 min (30 s on, 45 s off) at 21 W. The crude extract was obtained after centrifugation in a JA-20 rotor at 31,000 × g and 4 °C for 20 min.

Crude extracts of ChdT (100 mg) were loaded onto a 5-ml nickel–nitrilotriacetic acid Superflow Cartridge (Qiagen) using an ÄKTA FPLC P-960. The column was washed with 50 ml of 20 mM Tris-HCl buffer, pH 7.0, containing 50 mM NaCl and 25 mM imidazole, followed by 50 ml of 20 mM Tris-HCl buffer, pH 7.0, containing 50 mM NaCl and 75 mM imidazole. The protein was eluted using a linear imidazole gradient (75–500 mM) at a flow rate of 2 ml/min. Active protein fractions were pooled and concentrated using 50 mM Tris buffer, pH 7.0, containing 1 mM EDTA with an Amicon Ultra-15 10,000 molecular weight cutoff centrifugal filter unit (Millipore) resulting in ~12 mg/liter of soluble ChdT-His$_6$.  

![Figure 7. Four TPN docking poses for the active site of ChdT. The best pose (pose 1) indicates ligation of a CN group from TPN to the active site Zn(II) ion.](image-url)
The His₆ tag was removed by treating His₆-tagged ChdT with His₆-tagged TEV protease (EC 3.4.22.44) for 16 h at 4 °C in 50 mM Tris, pH 8.0. Cleaved protein was concentrated with a Centricon (15,000 molecular weight cutoff; Amicon) to 3 ml and loaded on immobilized metal affinity chromatography to remove the remaining cleaved His₆ tag, uncleav protein, and the His₆-tagged TEV protease, whereas the flow-through containing ChdT was collected and washed with 50 mM HEPES buffer, pH 7.0, containing 10% glycerol. Purified protein was analyzed by SDS-PAGE with a 12.5% polyacrylamide SPRINT NEXT GEL™ (Amresco). Protein concentration of crude extracts was determined using a Coomassie (Bradford) protein assay kit (Pierce) and pure protein by measuring the absorbance at 280 nm with a Shimadzu UV-2450 spectrophotometer equipped with a TCC-240A temperature-controlled cell holder. Theoretical molecular masses and protein extinction coefficients were calculated with the ExPASy compute pI/Mw tool.

**Activity assay of ChdT**

The enzymatic activity of ChdT toward TPN was measured using a Shimadzu UV-2450 spectrophotometer equipped with a TCC-240A temperature-controlled cell holder in 1-ml quartz cuvettes. A 1-ml reaction consisted of 50 mM HEPES buffer, pH 7.0, at 25 °C and various concentrations of TPN. The rate of TPN dehalogenation was determined by continuously monitoring the formation of 4-OH-TPN at 345 nm ($\Delta\varepsilon_{345} = 3.5 \text{ mM}^{-1} \text{ cm}^{-1}$). Data analysis was performed using OriginPro 9.0 (OriginLab, Northampton, MA, USA). The kinetic constants $V_{\text{max}}$ and $K_{m}$ were calculated by fitting to the Michaelis–Menten equation. One unit of enzyme activity was defined as the amount of enzyme that catalyzed the production of 1 μmol of the TPN minute at 25 °C.

**Crystallization**

Purified ChdT was exchanged into 50 mM HEPES buffer, pH 7.5, containing 10% glycerol for crystallization and concentrated to a final protein concentration of 10 mg/ml. Initial crystallization screens using a Gryphon Crystallization Robot (Art Robins Inc.) and Emerald Biosystems Wizard III/IV screen sets, produced crystals of ChdT in 10% PEG 6000 and 100 mM HEPES buffer, pH 7.0. Crystallization optimization leads revealed the best quality ChdT crystals via sitting-drop vapor diffusion occurred by mixing 6 μl of (10 mg/ml) ChdT with 6 μl of well solution (10% PEG 6000, 100 mM HEPES, pH 7.0). The crystals were allowed to grow at room temperature and reached optimal size and morphology for data collection after 7–10 days. The crystals were manually transferred to well solution supplemented with 30% glycerol (cryo-protectant) and flash frozen in liquid N₂.

**X-ray diffraction data collection**

SAD and monochromatic data were collected at the LS-CAT 21-ID Advanced Photon Source (APS) at Argonne National Laboratory. SAD data sets were collected at a wavelength of 1.28 Å, whereas another monochromatic data set was collected at 0.98 Å at 100 K using a Dectris Eiger 9M detector. The data sets were indexed and integrated using the HKL2000 suite.
Hydrolytic dehalogenase from Pseudomonas sp. CTN-3

Model building and refinement

The structure of ChdT was solved by SAD using autosol in the Phenix software suite (48). The model building was first conducted in autobuild followed by COOT (49), refined using Phenix, and analyzed in COOT and USCF Chimera (50). Final refinement statistics are reported in Table 1, and structural figures were made using USCF Chimera.

Double mutation at the structural zinc-binding site

His143 and Asp146, structural Zn(II) site ligands, were mutated to alanine (Ala143 and Ala149) by a single mutagenic primer as follows: sense primer, TTGATGGTGCCCTGGCGGCGGATAGGCGGTTCATGGT; and antisense primer: ACCATGAAAAGCTTTATCAGGCCGCAAAGGCACCA-TCAA (mutated residues are shown in bold type). The sense primer had a completely complementary sequence to the corresponding antisense primer. The first PCR was performed using sense primer and antisense primer in separate tubes (25 μl). Following denaturation at 98 °C for 1 min, five PCR cycles were carried out with each cycle comprised of incubation at 98 °C for 15 s, followed by a 30-s annealing step at 65 °C, and a 1-min extension at 71 °C. A 50-μl solution was prepared by combining two tubes for a second PCR, which was carried out under the same conditions as the first PCR but for 55 cycles. The final products were transformed directly into XL10-gold ultracompetent cells. The mutated ChdT sequence was verified by DNA sequencing by Functional Biosciences.

MALDI-TOF MS

Saturated α-CHCA thin layer solution was prepared on a MALDI target (51). A 2-μl protein sample (10 μM) was mixed with a DHB-CHCA matrix on a MALDI target by the “sandwich method” (21). Mass spectra of each sample were obtained on a Bruker microflex LRFTM MALDI-TOF MS.

Substrate docking

TPN poses were created and optimized by Molecular Operating Environment (Chemical Computing Group, Montreal, Canada). The TPN substrate was docked into the ChdT active site pocket using Autodock Vina, which is a structural analysis panel of Chimera 1.13.1rc (UCSF Chimera, San Francisco, CA, USA). The optimized TPN molecule was deemed a ligand, whereas the active site of ChdT was assigned as a receptor. The oxygen position of the active-site Zn(II)-bound water/hydroxide was fixed as were the protein residue ligands. The best pose, based on the docking scoring function versus binding energy (37), was considered the final model.

Data availability

All data described in the manuscript are contained within the manuscript or the supporting information. The final refined model was deposited at the Protein Data Bank under code 6UXU.

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Abbreviations—The abbreviations used are: Chd, chlorothalonil dehalogenase; ChdT, N-terminally truncated Chd; TPN, 2,4,5,6-tetrachloroiso-phthalonitrile; SAD, single-wavelength anomalous dispersion; 4-OH-TPN, 4-hydroxytrichloro-isophthalonitrile; MBL, metallo-β-lactamase; TBP, trigonal bipyramidal.

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Hydrolitic dehalogenase from Pseudomonas sp. CTN-3

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