Characterization of Chlorophenol 4-Monooxygenase (TftD) and NADH:FAD Oxidoreductase (TftC) of Burkholderia cepacia AC1100

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Burkholderia cepacia AC1100 completely degrades 2,4,5-trichlorophenol, in which an FADH2-dependent monooxygenase (TftD) and an NADH:FAD oxidoreductase (TftC) catalyze the initial steps. TftD oxidizes 2,4,5-trichlorophenol (2,4,5-TCP) to 2,5-dichloro-p-benzoquinone, which is chemically reduced to 2,5-dichloro-p-hydroquinone (2,5-DiCHQ). Then, TftD oxidizes the latter to 5-chloro-2-hydroxy-p-benzoquinone. In those processes, TftC provides all the required FADH2. We have determined the crystal structures of dimeric TftC and tetrameric TftD at 2.0 and 2.5 Å resolution, respectively. The structure of TftC was similar to those of related flavin reductases. The stacked nicotinamide:isoalloxazine rings in TftC and sequential reaction kinetics suggest that the reduced FAD leaves TftC after NADH oxidation. The structure of TftD was also similar to the known structures of FADH2-dependent monooxygenases. Its His-289 residue in the re-side of the isoalloxazine ring is within hydrogen bonding distance with a 2,5-DiCHQ. An H289A mutation resulted in the complete loss of activity toward 2,5-DiCHQ and a significant decrease in catalytic efficiency toward 2,4,5-TCP. Thus, His-289 plays different roles in the catalysis of 2,4,5-TCP and 2,5-DiCHQ. The results support that free FADH2 is generated by TftC, and TftD uses FADH2 to separately transform 2,4,5-TCP and 2,5-DiCHQ. Additional experimental data also support the diffusion of FADH2 between TftC and TftD without direct physical interaction between the two enzymes.

2,4,5-Trichlorophenol (2,4,5-TCP) and 2,4,6-trichlorophenol (2,4,6-TCP) belong to an environmentally persistent class of contaminants known as polychlorinated phenols (1). They have been used extensively as pesticides and preservatives, particularly for lumber and leather preservation, and as general herbicides and biocides. Unfortunately, human exposure likely causes various adverse health effects (2). The concern has prompted governments worldwide to classify several polychlorinated phenols as priority pollutants.

Recently, two microorganisms that degrade trichlorophenols have been characterized. Burkholderia cepacia AC1100 mineralizes 2,4,5-TCP (3, 4), and Cupriavidus necator JMP134 completely degrades 2,4,6-TCP (5). 2,4,5-TCP 4-monooxygenase (TftD) of B. cepacia AC1100, an FADH2-dependent monooxygenase, oxidizes 2,4,5-TCP to 2,5-dichloro-p-benzoquinone (2,5-DiCBQ), which is reduced to 2,5-dichloro-p-hydroquinone (2,5-DiHQ) by NADH (Fig. 1A). TftD further oxidizes 2,5-DiHQ to 5-chloro-2-hydroxy-p-benzoquinone, which is then reduced to 5-chloro-2-hydroxy-p-hydroquinone (5-Chl-2H-HQ) (6, 7). TftD can also oxidize 2,4,6-TCP to 2,6-DiCBQ but cannot further transform it or its reduced form 2,6-DiHQ. In these reactions, TftD uses molecular oxygen and FADH2 as co-substrates (8), and FADH2 is supplied by TftC, a flavin reductant.

Several FADH2-dependent monooxygenases, such as TftD, have been discovered in the biodegradation pathways of various aromatic compounds (9–12). The formation of a flavin-peroxide intermediate during catalysis has been proposed, and thus protection or stabilization of the FAD-peroxide intermediate from rapid hydrolysis seems crucial for those flavin-dependent monooxygenases (8, 13–16). All these monooxygenases have a small component as a partner, from which reduced flavins are provided. Those smaller reductases have a flavin molecule either as a tightly bound substrate or as a prosthetic group (10, 17–19). TftC and TftD belong to this two-component flavin-diffusible monooxygenase (TC-FDM) family and share se-
quency similarity with other flavin reductases and FADH$_2$-dependent monoxygenases that are involved in the degradation of various phenolic compounds.

So far, little is known about the interaction between TftC and TftD for FADH$_2$ transfer. Particularly reduced flavins can be readily oxidized by O$_2$ (20), and thus flavin dynamics is one of the critical features in the catalytic activity of monoxygenases. So far, several plausible mechanisms have been reported, e.g. (i) simple flavin diffusion between two components as in the cases of 4-hydroxyphenylacetate 3-monoxygenase of Escherichia coli and Acinetobacter baumannii (HpaB and HpaC) and phenol monoxygenase of Bacillus thermoglucosidasius (Phe-I(A) and Phe-2(A)) (21, 22); (ii) protein-protein interaction for direct substrate channeling as in the cases of bacterial luciferase (luciferase and flavin reductase P), alkane sulfonate monoxygenase (SsuE and SsuD), and arylamine oxygenase (PrnF and PrnD) (18, 23–26); and (iii) a probable combination of diffusion and complex formation for styrene monoxygenase (SMOA) and arylamine oxygenase (PrnF and PrnD) (27).

In addition, the dechlorination mechanism catalyzed by TftD is not well understood. An in-depth understanding of both TftC and TftD is important for elucidating the catalytic mechanisms, especially the mechanisms of sequential hydroxylations of 2,4,5-TCP. Here, we present a mechanistic analysis, including thermodynamic properties and crystal structures of TftC and TftD and propose a model for TCP oxidation. In addition, the plausible mechanism of flavin transfer between TftC and TftD was investigated through dynamic light scattering. This collective information could lead to the improvement of existing technologies for polychlorinated phenol bioremediation (28).

**EXPERIMENTAL PROCEDURES**

**Plasmid Construction and Cloning**—Both tftC and tftD were individually cloned into the overexpression plasmid pET30 LIC (Novagen) as described previously (8).

**Expression and Purification**—TftC or TftD expression was carried out by inoculating 100 ml of LB supplemented with 30 µg/ml kanamycin from a freezer stock of pET30TftC or pET30TftD in BL21(DE3) cells. This was allowed to grow overnight at 37 °C with constant shaking, after which this culture was used to inoculate 1.5 liters of LB medium. In addition, expression of selenomethionine-incorporated TftD in *E. coli* B834(DE3) (Novagen) was carried out using minimal media containing 30 mg/liter seleniummethionine. Protein expression was induced by addition of isopropyl β-d-thiogalactopyranoside to a final concentration of 0.5 mM at mid-log phase growth (A$_{600}$ = ~0.6). Following induction, the cells were incubated at 20 °C for 12 h with shaking at 250 rpm. Cells were harvested by centrifugation at 3,000 × g and pellets were frozen to protect the cell pellet and then thawed at room temperature, resuspended in a minimal volume of lysis buffer (50 mM Tris (pH 8.0), 300 mM NaCl, and 20 mM imidazole for TftC, 20 mM Tris (pH 8.5), and 3 mM dithiothreitol in the case of TftD), sonicated 10 times for 10 s each using a model 450 Sonifier® (Branson Ultrasonicos), and the resulting lysate cleared by centrifugation (20,000 × g for 40 min).

For the TftC purification, lysate was applied to a nickel-nitriotrilactate column and washed with several column volumes of lysis buffer. Elution took place with the lysis buffer containing 300 mM imidazole. Eluted fractions containing TftC were combined, concentrated, and buffer-exchanged into 20 mM Tris (pH 8.5) containing 50 mM NaCl by ultrafiltration in an Amicon 8050 cell with a 30-kDa cutoff membrane (Millipore). TftC was then applied to an ion-exchange column (Bio-Rad Uno Q12), and the protein of interest was collected in the flow-through. Concentrated TftC was applied to a CHT-II hydroxyapatite column and washed with several column volumes of 5 mM sodium phosphate (pH 6.8). Precipitates formed during this step were removed by centrifugation, after which the remaining solution, containing only TftC, was concentrated and exchanged into 20 mM Tris (pH 7.5).

For the TftD purification, lysate was loaded onto an anion-exchange column (Toyopearl® DEAE 650 M) equilibrated with buffer A (flow rate 6.0 ml min$^{-1}$). TftD was eluted from the column using a linear NaCl gradient between 0 and 200 mM NaCl. TftD-containing fractions were desalted and concentrated into buffer B (5 mM sodium phosphate (pH 7.0)). Concentrated TftD was applied to a CHT-II hydroxyapatite column (Bio-Rad), equilibrated with buffer B (flow rate 2.0 ml min$^{-1}$), and eluted in the column flow-through. Fractions were then concentrated into buffer C (20 mM Tris (pH 7.0), 3 mM dithiothreitol), loaded onto a Mono Q™ GL10/100 anion-exchange column (GE Healthcare), and eluted with a linear NaCl gradient at ~300 mM NaCl. Fractions containing TftD were pooled, concentrated, and exchanged into buffer D (20 mM Tris (pH 7.0)), after which the protein was loaded onto a Sephacryl S-200 column (GE Healthcare) and separated from remaining proteins.
Crystal Structures of TftC and TftD

With the exception of the nickel-nitrioltriacetate column, all purification steps were carried out using an Amersham Biosciences/BioCad 700E preparative HPLC (Applied Biosystems). All purification steps were monitored, and final homogeneity (>99%) was estimated using SDS-PAGE and Coomassie Blue staining.

Site-directed Mutagenesis and Enzyme Assay—Site-directed mutagenesis was performed using the QuikChange kit from Stratagene (La Jolla, CA). The primers used for the conversion of H289A of TftD are H289AF (5’-CGTATCTTCGACTGGTGGCTTACCACATTTTGATCCG-3’) and H289AR (5’-CGATCAAAATGTGGTAAGCCACCCAGTCGAGATACG-3’). The mutations were confirmed by sequencing, and the correct clones were transformed into E. coli BL21(DE3) for protein production. The mutant protein (TftD H289A) was purified analogous to the wild-type protein for characterization. The enzyme activity was assayed as described previously (5). Briefly, 40 μl of assay mixture contained 20 μM potassium phosphate buffer (pH 7.0), 100 μM 2,4,6-trichlorophenol, 10 μM FAD, 300 μM NADH, 1 mM ascorbic acid, 0.1 μg/ml of E. coli flavin reductase (Frec), and various amounts of wild-type TftD or H289A mutant. The reaction was initiated by adding NADH to the reaction mixture and terminated by adding 40 μl of acetonitrile/acetic acid mixture (9:1 (v/v)). The samples were centrifuged at 13,000 × g for 2 min, and the supernatants were analyzed by HPLC (8). TftC enzyme kinetics were performed by adding FAD and NADH to the enzyme at the concentrations described and then monitoring the change in absorbance of NADH at 340 nm. Km and Vmax values were determined by fitting the data to the Michaelis-Menten equation using Origin 5.0.

Molecular Mass Determination—The weight-average molecular masses of TftC and TftD were measured by combined size exclusion chromatography and multiangle light scattering as described previously (29). Briefly, 100 μg of TftC (or TftD) was loaded onto a BioSep-SEC-S 2000 column (Phenomenex) and eluted isocratically with a flow rate of 0.5 ml min⁻¹. The eluate was passed through tandem UV detector (Gilson), Optilab DSP interferometric refractometer (Wyatt Technology), and a Dawn EOS laser light scattering detector (Wyatt Technology). Scattering data were analyzed using the Zimm fitting method with software (ASTRA) provided by the instrument manufacturer.

Dynamic Light Scattering—To test for the formation of a complex between TftC and TftD, we used a DynaPro Titan (Wyatt Technology Corp.). A scan was taken of each protein individually as well as a scan containing a mixture of TftC, TftD, FAD, and NADH. NADH was added just before each experiment, and FAD reduction was confirmed by a change in color of the mixture. Measurements consisted of 10 consecutive 5-s scans, and average molecular radius was measured using a Rayleigh sphere approximation. Data were analyzed using the manufacturer-supplied software, Dynamics 6.7.3.

Isothermal Titration Calorimetry—The heats of binding of TftC and TftD with various cofactors were measured with a VP-ITC Microcalorimeter (Microcal, Northampton, MA). For calorimetric measurements, TftC or TftD in 20 mM Mops (pH 7.2), 100 mM NaCl, 1 mM dithiothreitol was titrated with one of the substrates dissolved in the same buffer. The apo-form TftC, TftD, and substrate solutions were degassed prior to titration. The experiment consisted of 29 injections of 10 μl each of ligand into the protein solution at 25 °C with constant stirring at 300 rpm employing a 300-s equilibration interval between injections. Heats of dilution of each ligand were determined by titration of the ligand into buffer without protein and were used to correct the protein titration data. The Origin software package (OriginLab Corp, Northampton, MA) was used to fit the data to an n-sites equivalent binding model using nonlinear least squares regression. Fitting the data provides the affinity, enthalpy, and entropy change for the binding reaction.

Titrations of TftC with FAD, FMN, and NADH and titrations of TftD with FAD and FMN were performed under aerobic conditions. For TftD titration with FADH₂ and FMNH₂, the titration experiment was performed in a glovebox under anaerobic conditions. For anaerobic titrations, protein was diluted into buffer followed by addition of 1 mM NADH, 1 mM glucose, and 1 μg/ml catalase. The titrant was prepared under the same conditions but without TftD and with the addition of 100 μM flavin (FAD or FMN). The samples were degassed under vacuum and then placed in the glovebox. Glucose oxidase was added to 1 μg/ml to scavenge oxygen. The flavins were reduced by incubation with 1 μg/ml of E. coli flavin reductase (Frec), and the progress of the reaction was followed visually by the color change of the flavin.

Circular Dichroism—Experimental conditions for all CD measurements were 0.013 mg/ml protein in 20 mM sodium phosphate buffer (pH 6.8) at 25 °C. CD spectra were recorded from 200 to 300 nm using an AVIV 202SF spectrophotometer (AVIV Biomedical, Inc.).

Crystallization and Data Collection—Apo-form crystals of TftC were grown at 4 °C using the hanging drop vapor diffusion method. 1.5 μl of TftC in 20 mM Tris (pH 7.5) were mixed with 1.5 μl of reservoir solution (20% (w/v) polyethylene glycol 4000, 0.2 mM potassium formate) and equilibrated against the reservoir. Crystals grew to a maximum size in 5 days and were of diffraction quality. For complex crystals of TftC with FAD and FAD-NADH, apo-form crystals of TftC were soaked for 6 and 3 h, respectively, by adding substrate directly to the drop containing the crystals.

Crystals of TftD were made by mixing 1.5 μl of protein solution (19 mg ml⁻¹) with an equal volume of crystallization solution (20% (w/v) polyethylene glycol 4000, 0.2 mM LiNO₃, (pH 7.1)). The purified TftD protein lost its FAD during purification, and attempts to make a complex crystal were unsuccessful. Addition of the reducing agent prevented growth of TftD crystals and therefore was eliminated from the final (Sephacryl) purification step. Large single crystals grew slowly (over 1 month). Both crystals of TftC and TftD were flash-frozen by transferring to the cryoprotection solutions containing the corresponding reservoir solutions plus 20% glycerol. The intensity data were collected at the Advanced Light Source (ALS, BL 8.2.1) at 100 K and reduced and scaled using HKL2000 (30) and CrystalClear 1.3.6 (Rigaku/MSC).

Structure Determination and Refinement—Initial phasing of TftC data were done by the program AMoRe (31) through the coordinates of CobR (Protein Data Bank code 3CB0). Amino
acid mutations were then performed using the graphics program O. Iterative model building and refinement took place using the programs O, X-PLOR, and PHENIX.

For TftD, location of the selenium heavy atom sites, phase calculation, and parameter refinement were performed using SOLVE (32) using a resolution range of 25 to 2.5 Å. This was followed by maximum-likelihood density modification using RESOLVE (32). After density modifications, well defined helical structures could be seen in the density, and the clear electron densities corresponding to most of the side chains allowed model building to proceed. The model was built into the electron density using the graphics program O (33) and refined using X-PLOR (34) and PHENIX (35). During the refinement processes for both TftD and TftC, the noncrystallographic symmetry restraint has not been applied. The final coordinates have been deposited in the Protein Data Bank as follows: apo-form TftC (3K87), FAD-TftC complex (3K87), FAD-NADH-TftC complex (3K88), and TftD (3HWC).

RESULTS

The apo-form of TftC was crystallized in the space group R3 with unit cell dimensions of \( a = b = 111.35, c = 103.00 \) Å, and there were two TftC molecules in the asymmetric unit. Cell dimensions for the complex crystals of TftC were \( a = b = 113.09, c = 101.68 \) Å, and \( a = b = 111.53, c = 102.52 \) Å for FAD-NADH and FAD, respectively. TftD was crystallized in the space group C222₁, with cell dimensions of \( a = 148.20 \) Å, \( b = 149.87 \) Å, \( c = 212.01 \) Å, and four molecules in the asymmetric unit. A summary of the crystallographic data for both TftC and TftD is given in Table 1.

| TABLE 1 Summary of crystallographic data |
|------------------------------------------|
| Space group                              |
| TftD                                     |
| TftC                                     |
| TftC-FAD                                 |
| TftC-FAD-NADH                            |
| Unit cell parameters                     |
| \( a \)                                  |
| 113.46 Å                                 |
| 111.35 Å                                 |
| 111.53 Å                                 |
| 113.09 Å                                 |
| \( b \)                                  |
| 149.87 Å                                 |
| 149.87 Å                                 |
| 148.20 Å                                 |
| 148.20 Å                                 |
| \( c \)                                  |
| 212.01 Å                                 |
| 212.01 Å                                 |
| 102.52 Å                                 |
| 101.68 Å                                 |
| Data                                     |
| Energy (Å)                               |
| Resolution (Å)                           |
| Completeness (%)                         |
| \( R_{merge} \)                           |
| \( R_{free} \)                           |
| No. of atoms                             |
| Nonsolvent                               |
| 33.6                                     |
| 43.8                                     |
| 40.7                                     |
| Solvent                                  |
| 2.0                                      |
| 2.0                                      |
| 2.0                                      |
| Wilson B-factor                         |
| 0.17                                     |
| 0.17                                     |
| 0.22                                     |
| Average B-value (Å²)                     |
| 0.17                                     |
| 0.17                                     |
| 0.22                                     |
| Bond lengths                             |
| 0.004                                    |
| 0.018                                    |
| 0.016                                    |
| Bond angles                              |
| 1.76                                    |
| 1.73                                    |
| 1.73                                    |
| Ramachandran plot                        |
| Favored                                  |
| 0.001                                    |
| 0.001                                    |
| 0.001                                    |
| Allowed                                  |
| 0.001                                    |
| 0.001                                    |
| 0.001                                    |

* Ramachandran plots were created using the program MolProbity (51).

Crystal Structures of TftC and TftD
Crystal Structures of TftC and TftD

A.

B.

C.

FIGURE 2. Ribbon diagram representing the crystal structures of TftC and its binding site. A, stereo view of the TftC dimer with associated FAD molecule with one subunit shown in green and the other in red. Secondary structural elements are labeled for one subunit and the noncrystallographic 2-fold axis is indicated by a half-arrow. B, difference map \( F_o - F_c \) map contoured at \( \sigma = 1.0 \) showed FAD molecule bound to TftC. Residues mentioned in the text as binding to the substrate are shown in yellow. C, difference map \( F_o - F_c \) map contoured at \( \sigma = 1.0 \) ternary complex of TftC showed both FAD and NADH. Residues important to substrate binding are shown in yellow. Several secondary structural elements near the FAD and NADH are labeled. Features marked with a prime are from the symmetry-related subunit. These figures were generated using Open-Source PyMOL™ (version 1.2).

follows: (i) between \( \beta_5 \) and \( \alpha_2 \), (ii) between \( \alpha_3 \) and \( \alpha_4 \), and (iii) \( \beta_{10} \) and \( \beta_{11} \). As mentioned in detail later, the first and second loops among those three participate in coordinating an FAD molecule. Accordingly, the corresponding temperature factors for Asn-67, Tyr-71, Ala-99, and Val-104, which are located in those two loops (Fig. 3), were reduced significantly upon complex formation with FAD.

Substrate-binding Site of TftC—The \( F_o - F_c \) maps of the FAD-soaked crystal data of TftC showed the corresponding electron density for the bound FAD molecule (Fig. 2B), with its isoalloxazine ring in a wide groove located at the dimer interface. The boundary of the binding pocket is made of \( \alpha_1 \), \( \beta_1 \), \( \beta_3 \), and \( \beta_5 \) from one subunit and \( \beta_4 \) of the other subunit. The isoalloxazine ring sits on top of antiparallel sheet made of \( \beta_2 \), \( \beta_3 \), and \( \beta_6 \), facing its re-side toward the bulk solvent. The two FAD molecules in the homodimer have somewhat different conformations of their ribityl moieties, one adopting an extended conformation and the other a folded-back conformation. The corresponding quality and coverage of electron density for the extended conformation was similar to that of the folded-back site, but the extended confirmation can be superimposed well with that of FAD in the FAD-NADH complex structure. In addition, there is crystal packing potentially hindering a proper diffusion for the folded-back site. Therefore, only the extended form of FAD will be discussed. The O2 and O4 atoms of the isoalloxazine ring are hydrogen-bonded to backbone nitrogen atoms of Ala-51 and Asn-67. The N5 atom of the isoalloxazine ring is also hydrogen-bonded to the hydroxyl side chain of Ser-50. Among those, only the alanine residue is relatively conserved among seven related flavin reductases (Fig. 3). The dimethyl benzene moiety of the isoalloxazine ring is located in a pocket formed by residues Tyr-171, Tyr-166, and Val-33, all of which are highly conserved in all flavin reductases (Fig. 3). The pyrophosphate group of FAD is located between \( \alpha_2 \) and \( \alpha_3 \), which are in a perpendicular orientation with respect to each other. Its phosphate and ribose moieties form hydrogen bonds with Thr-48, Tyr-71, Ala-99, and Arg-109, among which only Arg-109 provides a side chain interaction.

The \( F_o - F_c \) maps generated with the data collected from the crystal soaked with both FAD and NADH also clearly showed the corresponding electron density for both FAD and NAD(H) molecules stacked via the isoalloxazine and nicotinamide rings (Fig. 2C). In our FAD-NADH ternary complex of TftC, it is likely that the FAD molecules were in their reduced state judged by the pale yellow color of the soaked crystal in contrast to the intense yellow color of the FAD-containing crystal. The stacked isoalloxazine and nicotinamide rings were properly oriented at a distance of 3.5 Å between the C4 of NAD(H) and the N5 of FAD(H2). The position of the FAD molecule in this FAD-NADH complex was superimposable with the FAD molecule in the FAD complex structure. The side chain of His-145 was within hydrogen bond distance with the amide group of nicotinamide, whose re-face (A-side) was facing the re-face of the isoalloxazine ring. The pyrophosphate group of NAD(H) was hydrogen-bonded through the backbone of Tyr-166 as well as the backbone and side chain of Arg-169. The N3A atom of the adenine ring is within hydrogen bonding dis-
tance from the hydroxyl group of Ser-54 from the other subunit. Significantly, all residues in the NAD(H)-binding site are highly conserved among all compared flavin reductases. Consequently, the coordination mechanism observed in TftC is almost identical to those of previously reported flavin reductases, including the folded NAD(H) confirmation, first reported for Phe-2(A), with a distance between the adenine C6 and nicotinamide C2 atoms of 4.4 Å.

**Structure of TftD**—Four tightly associated subunits in the crystal asymmetric unit are arranged in a noncrystallographic 2-fold symmetric assembly (Fig. 4B). The tetrameric nature of TftD in solution was confirmed by the elution profile and molecular weight estimates from a MALLS experiment (supplemental Fig. S3). The individual TftD molecules consist of 14 β-strands and 17 α-helices (Fig. 4A) and can be divided into three sequential segments based on their constituent secondary structural elements. The first area from residues 1 to 146 is composed of seven α-helices (a1–a7) and four very short β-strands (β1–β4). The second area, which is composed of residues 147–273, has eight β-strands (β5–β12). The third area, C-terminal segment is composed of 10 (-helices (a8–a17) and 2 short β-strands (β13 and β14). Those three sequential segments are tightly interconnected in the three-dimensional structure. Noticeably, all of the 10 helices in the C-terminal segment, α8–α17, are involved in an inter-subunit interaction as discussed in detail later. All the eight β-strands in the second segment, β5–β12, are arranged in such a way as to appear almost like a β-barrel. At one side of this barrel, the β1 and β2 strands from the first segment fold back and together form a continuous β-sheet in the order of β2-β1-β10-β11-β7-β6 (Fig. 4A). The remaining four strands in the segment cover the other side of the same barrel in the order of β5-β8-β9-β12. The amino acid side chains of those 10 β-strands and the connecting loops between them form a hydrophobic core inside of the barrel.

Another noticeable structural feature is that three long α-helices, α9, α10, and α11, and three short helices, α14, α15, and α16, which are all from the C-terminal segment, form a sizeable helix bundle together with two short α-helices from the first segment, a4 and a5. The surfaces of most of those α-helices are mostly hydrophobic, establishing a hydrophobic interaction among them.

The corresponding electron densities for the C-terminal residues 442–445 and 483–515 are not visible, probably due to being disordered. In addition to the immediate neighbors of these two regions, the temperature factors for residues 158–167, which form an exposed loop connecting β5 and β6, showed higher temperature factors than the rest of the molecule.

The observed tetramer interface had an extensive network of inter-subunit interaction among the C-terminal α-helices. In particular, α16 and α17 extend out of the individual subunit to embed themselves into the hydrophobic surface made by α9, α10, and α14 of the adjacent subunit. In addition, a large portion of two anti-parallel C-terminal helices, α10 and α11, from two neighboring subunits contact closely in a perpendicular orientation generating a substantial hydrophobic interface between subunits. These hydrophobic interfaces involve residues such as Val-326, Leu-334, Leu-360, Phe-364, Leu-450, Phe-454, and Met-458. The remaining three α-helices at the C-terminal segment also form another type of subunit interaction together with the above-mentioned β-barrel-like motif, i.e. α12, α13, α15, their connecting loops, and the surface-exposed residues in one side of the barrel form a tightly associated non-crystallographic 2-fold related interface. This interface also possesses tightly packed hydrophobic residues as follows: Val-187, Val-190, Phe-230, Val-257, Leu-389, Met-398, Trp-405, and Phe-406. However, contrary to the above-mentioned hydrophobic interface involving α16 and α17, a substantial amount of polar inter-subunit interaction was observed, which includes Asp-159, Glu-233, Lys-237, Asp-250, Asp-259, Arg-386, Gln-396, Gln-401, Asn-410, Arg-420, Arg-428, Asp-434, and Arg-438.
Substrate-binding Site of TftD—In the apo-structure of TftD, a unique entry site and binding pocket for FADH$_2$ was clearly distinguishable (Fig. 4C). The perimeter of this widely open entry site is made with three flexible loops as follows: (i) between H9252 and H9253, (ii) between H9254 and H9255, and (iii) between H9215 and H9216. Among those three, the first and third loops have high temperature factors and show the most heterogeneous conformations among the four subunits, reflecting their high level of flexibility. The residues lining this entry site are predominantly hydrophilic, and a few ordered solvent molecules were located in the entry site of the determined apo-form structure of TftD.

The position of the riboflavin moiety of the FADH$_2$ was readily obtained through the combination of superposition with the 4-hydroxyphenylacetate 3-monoxygenase (HpaB) from *Thermus thermophilus* HB8 (2YYJ) (36) and the solid docking module on Quanta (BioSYM/MSI, Inc.), which is based on conformational space, followed by a quick energy minimization by CNS version 1.1 (37). The wall of this binding pocket is made with small portions from H9252, H9253, H9254, H9211, H929, and H9215 of one subunit and portions of H929, H910, and H911 of another subunit.

Isothermal Titration Calorimetry (ITC) for TftC and TftD—ITC was used to characterize the binding of FAD, FMN, and NADH to TftC. As shown in Fig. 5A, a significant amount of heat is released when TftC is titrated with FAD. However, a negligible amount of heat was released when TftC was titrated with FMN and NADH, indicating a lack of binding. A calculated $K_d$ of FAD is $1.8 \pm 0.1$ $\mu$M (mean $\pm$ S.D.), with a large enthalpic contribution ($\Delta H = -11.5$ kcal/mol). The data also indicated an unfavorable entropic contribution, with a calculated $\Delta S$ of $-12.29$ cal/mol/degree, possibly indicating that the TftC structure was slightly stabilized upon binding of FAD, and very few solvent molecules were freed from the pocket. These thermodynamic data were consistent with structural data, as indicated by the significant reduction of the $B$-values for two loops constituting the binding pocket upon formation of the FAD binary complex. There were few solvent molecules in the FAD binding pocket.
As discussed under “Discussion,” the structure of TftD indicated that one of the residues in the active site, His-289, might play a critical role in its catalysis, and thus TftD H289A mutant enzyme was generated, and its structural integrity was compared with that of the wild type by measuring their CD spectra. As shown in the supplemental Fig. S4A, the spectral pattern of H289A mutant in a buffer containing 20 mM sodium phosphate (pH 6.8) was superimposable to that of the wild type. Both have two negative bands around 208 and 222 nm, which are the characteristics of CD spectra of typical α-rich or αβ proteins. The tetrameric nature of the H289A mutant in solution was also confirmed by a MALLS experiment (supplemental Fig. S4B). The enzyme activities for the mutant were assayed using 2,4,6-TCP, 2,4,5-TCP, or 2,5-DiCHQ as the substrate. TftD H289A mutant still used 2,4,5-TCP and 2,4,6-TCP as substrates with about 28- and 53-fold decreases in K_m, respectively, and the k_cat values were not significantly altered (Table 2). Most significantly, the mutant enzyme completely lost activity toward 2,5-DiCHQ.

Dynamic Light Scattering—To probe the dynamics of flavin transfer between the small component flavin reductase and the large component monoxygenase, chemical cross-linking and dynamic light scattering were employed. The average Rayleigh sphere radius of TftC and TftD was measured independently and in combination with TftC substrates. When a solution of TftC and TftD (also containing FAD and NADH) was measured, FAD was reduced to FADH_2. As was confirmed by the color change and the FADHOOH spectrum (8), the bound FADH_2 in TftD was immediately oxidized to FADHOOH. Thus, most added FAD is in the form of the FADHOOH-TftD complex. If a higher order complex were formed between the two enzymes to transfer FADH_2, the height of the TftC peak would either decrease or disappear, whereas the TftD peak would appear to shift to a larger radius. As Fig. 7 indicates, addition of substrates (FAD and NADH) causes no significant shift in intensity or any shift in average particle size, indicating a lack of interaction between the two enzymes. Similar experiments were carried out using chemical cross-linking, and the results indicated the same outcome (data not shown). This supports the growing body of evidence that flavin transfer in TC-FDM systems occurs via a purely diffusive mechanism (21, 38, 39).

### Table 2: Kinetic properties of TftD and TftD H289A

| Substrate | K_m (µM) | k_cat (s⁻¹) | k_cat/K_m (M⁻¹ s⁻¹) |
|-----------|----------|------------|----------------------|
| 2,4,5-TCP | 35.8 ± 3.9 | 0.67 ± 0.03 | 1.9 × 10⁴ |
| TftD H289A | 36.6 ± 3.4 | 0.024 ± 0.003 | 6.6 × 10² |
| 2,4,6-TCP | 39.9 ± 7.6 | 0.41 ± 0.03 | 1.0 × 10⁴ |
| TftD H289A | 18.5 ± 2.4 | 0.0078 ± 0.0006 | 4.2 × 10² |
| 2,5-DiCHQ | 4.3 ± 1.1 | 0.10 ± 0.01 | 2.3 × 10⁴ |
| TftD H289A | 0 | No activity |

* Experiments were done in 40 mM KPi buffer (pH 7.0) at 24 °C. Values are means of triplicate experiments with standard deviations. The kinetic properties of TftD were reported previously (8).

As in the case of TftC, ITC was used to confirm the differential binding affinities of FAD and FMN for TftD and their corresponding reduced forms. As shown in Fig. 5B, FAD, FMN, and FMNH_2 molecules did not show any significant binding to TftD. However, a significant amount of heat was released when FADH_2 associated with TftD, indicating that the binding interactions had significant enthalpic contributions (ΔH = −8.33 kcal/mol). The data also showed a slightly unfavorable entropic contribution (ΔS = −0.78 cal/mol/degree); however, it is less significant than ΔS of TftC. ITC data analysis yielded a K_d for FADH_2 binding to TftD of 1.2 ± 0.2 µM.

**Enzyme Kinetics and Site-directed Mutagenesis**—Steady-state kinetics experiments were performed by holding the concentration of TftC constant while varying the FAD concentration at 1, 2, 4, 6, and 8 µM. The NADH concentrations in the corresponding four experiments were varied at 25, 50, 100, and 200 µM, and each point was performed in triplicate. As shown in Fig. 6, the double-reciprocal plot of the inverse rate versus inverse substrate concentration yielded lines intersecting to the left of the y axis, indicating a sequential reaction mechanism. The calculated K_m values for FAD and NADH were 2.0 (±0.3) and 45.3 (±4.5) µM, respectively. The V_max value was 120.3 (±7.7) µmol min⁻¹ mg⁻¹.
Thermophilic Sulfolobus tokodaii Geobacillus thermoglucosidasius Psychrobacter psychrophilus somnus interaction between the two proteins, indicating the reduced flavin could transfer via a diffusive mechanism.

A mixture of the two proteins with FAD and NADH is shown as a solid line. The plot indicates a lack of physical interaction between the two proteins, indicating the reduced flavin could transfer via a diffusive mechanism.

**DISCUSSION**

We have determined the crystal structures of both TftC and TftD to shed light on their reaction mechanism and structural relationship to other flavin-dependent monooxygenases and flavin reductases in TC-FDM. In particular, TftC and TftD together perform two sequential oxidative dechlorination steps in the degradation process of 2,4,5-TCP (Fig. 1) (8), even though most monooxygenases hydroxylate their substrate only once (40, 41). For TftC, in addition to its apo-form, structures of the binary complex with FAD and the ternary complex with FAD and NADH were also determined. However, we were unable to produce a complex crystal of TftD with FAD under aerobic environments, likely due to the negligible affinity of TftD for FAD as shown by our ITC data (Fig. 5B). Consequently, the position of flavin was determined through superimposition with structural homologues and a substrate docking approach.

**Structural Classification for TftC**

The primary and tertiary structural comparison of TftC revealed its high similarity with other flavin reductases. A Dali search showed that the most similar structure was corrin reductase (CobR) from *Brucella melitensis* (3CB0) with a high Z-score of 33.5, followed by putative flavin reductase from *Haemophilus somnus* (2ROX) with a Z-score of 26.3, Phe-2(A) from *Geobacillus thermodenitrificans* (1RZ1) with 23.3, HpaC from *Sulfobolus tokodaii* str. 7 (2D37) with 21.8, and HpaC from *T. thermophilus* (2ECR) with 21.0. In general, the β-barrel-like core structure and dimeric nature of TftC are similar to the structures of high similarity scores, which include other small component members of the TC-FDM family, such as Phe-2(A), HpaCS, and HpaCT. However, a detailed visual inspection revealed a structural heterogeneity in terms of the number and size of helices in the peripheral regions. Additionally, the C terminus of each TftC subunit forms interlocking anti-parallel β-strands with each other, which has not been observed in other flavin reductases. Noticeably, the structure of TftC is quite different from that of EmoB, which is also a small component of TC-FDM (19). A search for similar amino acid sequences in the Protein Data Bank using BLAST (42) revealed that putative flavin reductase from *H. somnus* (2ROX) and CobR (3CB0) showed the highest score (119 bits) with 40 and 38% identity among matched amino acids to TftC, respectively, followed by Phe-2(A) (1RZ0, 70.5 bits, 31% identity), HpaCSt (2D36; 57.4 bits, 25% identity), and HpaCSt (2ECR; 56.6 bits, 26% identity). The conserved residues were distributed sporadically through the entire secondary structural elements. The α2 and its neighboring loops show the least similarity among the above reductases (Fig. 3). In addition, there are insertions or deletions between α3 and β7 producing heterogeneity in the secondary structural elements among those compared flavin reductases. Significantly, this area of heterogeneity happens to be involved in binding of the adenine moiety of flavin in those reductases. In Phe-2(A) and HpaC, binding for the adenine moiety is through multiple hydrogen bonds with this region referred to as Loop 7. However, as noted previously in the structure of CobR (43), Loop 7 is replaced by an additional α-helix, α4, in TftC, which alters the binding interactions with AMP. It should also be noted that HpaC of *S. tokodaii*, which prefers FMN to FAD, contains a 310 helix in the same position as α4 in TftC.

The residues constituting the hydrophobic binding pocket for the dimethyl benzene moiety of the isoalloxazine ring are highly conserved among those enzymes. However, the corresponding sites facing the 2,4-pyridine side and the ribose group of the flavin molecule are not well conserved among the structurally related proteins (Fig. 3). Despite this poor sequence similarity, the backbones of the corresponding regions are similarly engaged in coordination in all the compared reductases whose complex structures are available.

**Structural Classification for TftD**

A Dali search was also used for identifying structural homologues for TftD (45) and showed that the highest match was to the 4-hydroxyphenylacetate 3-monooxygenase (HpaB) of *T. thermophilus* HB8 (2YYL) (36) with a Z-score of 43.7. It was followed by 4-hydroxybutyryl-CoA dehydratase (4-BUDH) of *Clostridium aminobutyricum* (1U8V) (46) with a Z-score of 36.5 and a couple of other structural homologues with much high similarity, the backbones of the corresponding regions are similarly engaged in coordination in all the compared reductases whose complex structures are available.
Substrate-binding Site and Reaction Mechanism

TftC—The observed conformation of the AMP moiety of the bound FAD in TftC is quite different from that in the complex crystal of the Phe-2(A) and HpaC; likewise, the local conformation of TftC shows difference from those, i.e. the exposed adenine ring of the FAD stacks with the phenyl side chain of tyrosine 71 in the TftC structure, in contrast to the definite binding pocket offering multiple hydrogen bonds to the adenine ring observed in both Phe-2(A) and HpaC. This apparent difference in binding of the AMP moiety of FAD might result in a lower affinity for FAD ($K_d$) and loss of FAD during the purification. In addition, there was no flavin occupancy in the apo-form crystal structure. This is in contrast with what has seen in other flavin reductases, such as Phe-2(A), which has a nanomolar affinity for flavin.

The NADH:FAD oxidoreductase activity of TftC was through oxidation of an NADH molecule by a bound FAD molecule, which caused reduction of the flavin. The ternary complex structure of TftC showed that the stacked nicotinamide:isoalloxazine rings were at a proper distance for hydride transfer. Compared with the FAD molecule, the NAD(H) molecule in the ternary complex of TftC had less interaction with the enzyme despite its interactions with highly conserved residues and many water-mediated interactions, which are very similar to that of Phe-2(A). The NAD(H) molecule had a negligible affinity with TftC that had no bound FAD in it, as indicated by the ITC results (Fig. 5). Therefore, the apparent affinity for NADH existed only after the FAD occupies its site in TftC. In addition, in the crystal structure for the FAD:NADH ternary complex form, the adenine ring is folded back in a stacking position on top of the nicotinamide ring, probably shielding the lower Z-scores such as medium chain-specific acyl-CoA dehydrogenase (MCAD) from pig mitochondria (3MDE) (47). Careful manual comparisons of these crystal structures show that they share similarity with TfdD in terms of the location of the secondary structural elements and topological connectivity (Fig. 8). Consistent with a structural similarity among TfdD, MACD, HpaB, and 4-BUDH, there is significant sequence similarity among these structural homologues. For example, TfdD and many water-mediated interactions, which are very similar to that of Phe-2(A). The NAD(H) molecule had a negligible affinity with TftC that had no bound FAD in it, as indicated by the ITC results (Fig. 5A). Therefore, the apparent affinity for NADH existed only after the FAD occupies its site in TftC. In addition, in the crystal structure for the FAD-NADH ternary complex form, the adenine ring is folded back in a stacking position on top of the nicotinamide ring, probably shielding the
C4 atom of nicotinamide from bulk solvent (Fig. 2C). This is also observed in both HpaC and Phe-2(A) structures (48), allowing the efficient hydride transfer to FAD. Once FAD in TftC is reduced by NADH, the FAD molecule will leave as FADH2. Based on the results from the double-reciprocal plot, it is obvious that the production of FADH2 by TftC is through a sequential reaction mechanism (Fig. 6). In the catalytic cycle NADH reduces the associated FAD and NADH leaves TftC. The diffusible FADH2 is then used by TftD for TCP metabolism. In this process, the NADH molecule delivers its hydride through transiently stacked interactions of its nicotinamide ring with the isoalloxazine ring of FAD as similarly observed in the complex structures of EmoB, Phe-2(A), and HpaC (19, 44, 48, 49).

**TftD**—The secondary structural elements of TftD around the putative substrate-binding pocket were superimposable with those in available holo-form structures of the above-mentioned enzymes, MCAD (3MDE), HpaB (2YYJ), and 4-BUDH (1U8V). Therefore, a reliable position for the riboflavin moiety of the FADH2 could be established. However, the residues for coordinating the pyrophosphate or adenosine group of the FAD(H2) could not be pinpointed due to the conformational differences among the four subunits of apo-form TftD. The two conserved residues (Fig. 8), Gln-157 and Arg-160, which reside in the flexible loop between β5 and β6 and coordinate the phosphate group of FADH2 in HpaB (36), are not in the coordination position in TftD. Therefore, upon FADH2 binding, a conformational change might follow in that local area to accommodate binding of the pyrophosphate and adenosine moieties. However, considering a noticeably weaker affinity of TftD (micromolar range) for FADH2 compared with that of HpaB (nanomolar range) (5) and negligible affinity for both FAD and FMN, TftD might have intrinsically poor coordination for the ADP portion of the FAD.

The geometry and several polar residues in the si-face of the modeled isoalloxazine ring are well conserved among TftD, MCAD (3MDE), HpaB (2YYJ), and 4-BUDH (1U8V). Thr-192 is completely conserved among all four FADH2-dependent enzymes, and their corresponding hydroxyl group side chains are in ~3.0 Å distance from the N5 atom of the isoalloxazine ring (Fig. 4C). In addition, three other nearby residues, Arg-100, Asp-253, and Arg-438, are conserved among TftD, TcpA, and HpaB (Fig. 8). The side chain of Arg-100 has been proposed to form a hydrogen bond with the peroxy moiety of the C4a-hydroperoxy flavin intermediate and to contribute to the formation and/or stabilization of that transient intermediate (36, 50). As observed in the HpaB-FAD complex structure, the Nε atom of Arg-100 of TftD is located 4.8 Å from the C4a atom of the modeled isoalloxazine ring.

The affinity of HpaB for FADH2 is in a nanomolar range compared with the micromolar range for FAD binding (21). This substantially higher affinity of HpaB for FADH2 over FAD has been attributed to the immediate peroxidation of FADH2 and its subsequent stabilization through the guanidinium side chain of Arg-100 (36). In contrast to micromolar range affinity of HpaB for FAD, our ITC data show that TftD does not have significant affinity to FAD (Fig. 5B). However, in a completely anaerobic setting, which does not allow the formation of FAD or FAD-peroxide, TftD shows a Kd of 1.2 μM for FADH2. Therefore, there is a clear difference in flavin affinity between TftD and HpaB.

In the si-side of TftD, there exists a significant difference among apolar residues in the surroundings of the bound isoalloxazine ring compared with the same side of the above-mentioned flavoenzymes. Noticeably, Val-154 is unique for TftD; the amino acid at the equivalent position is a threonine in HpaB, MCAD, and 4-BUDH. In particular, the side chain of that threonine residue in MCAD and 4-BUDH is in a position to hydrogen bond with the N1 and O2 atoms of the isoalloxazine ring. Therefore, the substitution of this residue to an apolar...
Crystal Structures of TftC and TftD

amino acid in TftD can be another reason for its negligible affinity for FAD in addition to the above-mentioned poor coordination of its AMP moiety. In addition, the corresponding residue for Val-190 is threonine, histidine, and tryptophan in HpaB, 4-BUDH, and MACD, respectively. Both histidine and tryptophan are located in proximity to the riboflavin part of the attached FAD, which could reflect its tighter affinity as a cofactor in 4-BUDH and MACD. In both TftD and HpaB, FADH$_2$ is a substrate instead of a cofactor.

The residues facing the re-side of the isoalloxazine ring, which is presumably a 2,5-DiCHQ or TCP-binding site, are quite different among the compared enzymes, as expected from their substrate specificities. In comparison with the corresponding residues of HpaB, the residues of TftD in the re-side of the isoalloxazine ring are much more apolar. For example, the corresponding residues for Leu-151, Ile-205, and Val-288 are polar in HpaB. In addition there are several other noticeable differences between these two closely related flavin-dependent monooxygenases. Previously, Tyr-104 and His-142 residues in HpaB have been noticed to coordinate the hydroxyl group of its substrate, 4-hydroxyphenyl acetate, and to abstract the proton from it, respectively. The corresponding amino acids in TftD are Ala-104 and Val-151; thus, neither of these TftD residues is able to coordinate the hydroxyl group of 2,5-DiCHQ or TCP. His-289 in the putative substrate-binding site, which is located on the re-side of the isoalloxazine ring, is the only candidate that could coordinate the hydroxyl group or proton abstraction from the bound substrate. Establishing a hydrogen bond between the imidazole side chain of His-289 and one of the hydroxyl groups of the modeled 2,5-DiCHQ resulted in the other hydroxyl group facing toward the solvent-exposed side of the pocket (Fig. 4C). Our results from site-directed mutagenesis, MALLS and CD spectra confirmed that the His-289 residue is involved in catalysis. Despite the completely abolished activity toward 2,5-DiCHQ, the H289A mutant still maintains the specificity toward 2,5-DiCHQ, the H289A mutant still maintains the activity toward 2,5-DiCHQ. Despite the completely abolished activity toward 2,5-DiCHQ, the H289A mutant still maintains the activity toward 2,5-DiCHQ. Therefore, we propose that the His-289 residue of TftD plays critical bifunctional roles in its unique sequential catalysis of 2,4,5-TCP and 2,5-DiCHQ.

Closing Remark

Characterizing the participating key enzymes is the prerequisite for informed and successful bioremediation of major pollutants, which will improve our understanding of the reaction mechanisms and increase our ability to remove these pollutants from the environment. Our data offer the mechanistic understanding of TftC and TftD activities. The results support that free FADH$_2$ is generated by TftC and that TftD uses this diffused FADH$_2$ to transform 2,4,5-TCP and 2,5-DiCHQ. Furthermore, the different roles of His-289 of TftD in the catalysis of 2,4,5-TCP and 2,5-DiCHQ are proposed, explaining how a monooxygenase attacks two substrates. Besides contributions to basic biochemistry, the results may facilitate bioremediation of polychlorinated phenols.

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