Structural insights into a flavin-dependent dehalogenase HadA explain catalysis and substrate inhibition via quadruple π-stacking

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HadA is a flavin-dependent monooxygenase catalyzing hydroxylation plus dehalogenation/denitration, which is useful for biodecontamination and biodetection. In this study, the X-ray structure of wild-type HadA (HadAWT) co-complexed with reduced FAD (FADH+) and 4-nitrophenol (4NP) (HadAWT−FADH+−4NP) was solved at 2.3-Å resolution, providing the first full package (with flavin and substrate bound) structure of a monooxygenase of this type. Residues Arg101, Gln158, Arg161, Thr193, Asp254, Arg233, and Arg439 constitute a flavin-binding pocket, whereas the 4NP-binding pocket contains the aromatic side chain of Phe206, which provides π–π stacking and also is a part of the hydrophobic pocket formed by Phe155, Phe286, Thr449, and Leu457. Based on site-directed mutagenesis and stopped-flow experiments, Thr193, Asp254, and His290 are important for C4a-hydroperoxylavin formation with His290, also serving as a catalytic base for hydroxylation. We also identified a novel structural motif of quadruple π-stacking (π-π-π-π) provided by two 4NP and two Phe441 from two subunits. This motif promotes 4NP binding in a nonproductive dead-end complex, which prevents C4a-hydroperoxy-FAD formation when HadA is premixed with aromatic substrates. We also solved the structure of the HadA Phe441Val−FADH+−4NP complex at 2.3-Å resolution. Although 4NP can still bind to this variant, the quadruple π-stacking motif was disrupted. All HadA Phe441 variants lack substrate inhibition behavior, confirming that quadruple π-stacking is a main cause of dead-end complex formation. Moreover, the activities of these HadA Phe441 variants were improved by 20%, suggesting that insights gained from the flavin-dependent monooxygenases illustrated here should be useful for future improvement of HadA’s biocatalytic applications.

Halogenated phenols (HPs) and nitrophenol (NP) are persistent pollutants resulting from anthropogenic activities including the use of agro- and household chemicals such as pesticides, herbicides, and flame retardants. Their accumulation in the environment can pose a hazard to human health, with toxic effects ranging from causing chronic diseases to acute death (1). Through natural evolution, microbes have developed enzymes and metabolic pathways to combat against these chemicals by degrading them into common metabolites, which can then be used as cellular energy sources (2, 3). The had pathway from Ralstonia pickettii is one of the most well-known pathways for biodegrading HPs and NP. It contains several enzymatic reactions catabolizing pesticides such as 2,4-dichlorophenol, 2,4,5-trichlorophenol, and 2,4,6-trichlorophenol to generate benzoquinone (BQ) derivatives, which can be assimilated into the tricarboxylic acid cycle (4–8).

The initial and committing step of the had pathway catalyzed by HadA monooxygenase is hydroxylation with removal of either halide substituents (F, Cl, Br, or I) from HPs or a nitro group (-NO2) from NP (6–8). HadA can detoxify a wide range of toxicants, and its ability to convert monosubstituted HPs and NP to a valuable compound such as D-luciferin has recently been shown (9). The ability of HadA to synthesize D-luciferin highlights its potential value for use in toxicant waste refineries. Rather than allowing HPs or NP to be dissipated into the open environment, thereby creating toxic ecological effects, a proper waste collection and detoxification procedure can be performed using HadA to detoxify and convert them to valuable compounds.

HadA belongs to the class D two-component flavin-dependent monooxygenases, which use reduced FAD (FADH+) as a substrate (7, 10–14). Kinetic mechanisms of wild-type HadA (HadAWT) using 4-chlorophenol (4CP), 4-bromophenol, 4-iodophenol, 4-fluorophenol, 4-nitrophenol (4NP), and phenol as substrates have shown that a binary complex of HadA-FADH+ is the first species to form before reacting with O2 to form a C4a-hydroperoxy-FAD intermediate (Fig. 1) (7, 8). A substrate such as 4CP then binds and receives a terminal –OH group (electrophile) from the C4a-hydroperoxy-FAD, which is incorporated into the C4 position of 4CP (nucleophile) via an electrophilic aromatic substitution mechanism, resulting in a C4a-hydroxy-FAD intermediate and the hydroxylated product. Next, a halide or nitro group is eliminated to form benzoquinone as a final

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product while C4a-hydroxy-FAD is dehydrated to form oxidized FAD before the product is released from HadA to complete the catalytic cycle. The quantitative structure–activity relationship analysis revealed that the overall reaction of HadA is controlled by the ability of the substrate to be deprotonated because the rate constants of HadA reactions with substrates having different substituents at the 4-position depend directly on the pKₐ values of the compounds (8).

Despite its potential use in biodetoxification and known reaction mechanisms, the X-ray crystallographic structures of HadA or related homologs in complex with flavin and aromatic substrates have not been reported. Only structures of apo-HadA_WT and its homologs including apo-ReTcpA from Ralstonia eutropha JMP134 and apo-TftD from Burkholderia cepacia AC1100 without bound ligand were reported (15–17). Thus, insights into the substrate recognition and catalytic mechanism of HadA are still not understood.

The reaction of HadA also has a side pathway that impedes the application of this enzyme in real and scalable usage. HadA_WT displays strong substrate inhibition by the preformed HadA_WT–4CP complex or “dead-end complex,” which prevents formation of C4a-hydroperoxy-FAD and thus inactivates the overall reaction (Fig. 1) (7). We note that substrate inhibition by formation of a dead-end complex is also found in other two-component flavin-dependent monooxygenases such as bacterial luciferase (class C) (18–20). However, no structural features that can promote substrate inhibition in these enzymes are known. This phenomenon limits the enzyme usage in biocatalysis because the flavin and substrate need to bind to the enzyme in a strict sequential order to avoid inhibition. This can be achieved under low substrate concentrations because the rate of flavin binding is generally faster. However, under high substrate concentrations, the rate of substrate binding (bimolecular reaction), which is dependent on the concentration, can be faster than the flavin binding. This can cause the turnovers to cease (7). Therefore, molecular insights into the mode of substrate inhibition gained by structural analysis of HadA would be useful for shedding light on the mechanisms underlying catalysis and inhibition and would allow improvement in applications of HadA and other two-component flavin-dependent monooxygenases in general.

In this work, we present the crystal structures of a ternary complex of HadA_WT and FADH⁻ and 4-nitrophenol (HadA_WT–FADH⁻–4NP), representing the first comprehensive ligand-bound structure of dehalogenating/denitrating flavin-dependent monooxygenases. The binding pocket of substrates 4NP and FADH⁻ was identified, and the interactions of the substrates with the surrounding and key catalytic residues that enable HadA catalysis were determined by site-directed mutagenesis, stopped-flow experiments, and product analysis. Moreover, we found rigid quadruple π-stacking (π-π-π-π) interactions between the aromatic moieties of the two bound 4NP molecules and the two aromatic side chains of Phe441 from two subunits at the dimer interface. This unusual π-π-π-π interaction explains the root cause of substrate inhibition by forming a dead-end complex. We verified this hypothesis by disrupting the quadruple π-stacking interaction by site-directed mutagenesis of Phe441. The results indeed showed that the substrate inhibition was alleviated in the HadA_Phe441Val and HadA_Phe441Leu variants. This is the first time that the substrate inhibition mechanism in flavin-dependent monooxygenases can be explained.
structurally. The outcome of this work should help improve future applications of flavin-dependent monooxygenases in general.

Results

The overall structure of the HadA<sub>WT</sub>−FAD<sup>+</sup>−4NP ternary complex

In order to gain structural and mechanistic insights into HadA<sub>WT</sub>, we cocrystallized the enzyme with FAD<sup>+</sup> and 4NP to obtain the first ternary structure of HadA<sub>WT</sub>−FAD<sup>+</sup>−4NP at a 2.3-Å resolution. Data and refinement statistics are listed in Table 1. The HadA<sub>WT</sub>−FAD<sup>+</sup>−4NP ternary complex (Protein Data Bank [PDB] code: 7E8P) adopts a common structural fold of the acyl-CoA dehydrogenase flavoenzyme family with two dimers associated to form a tetramer (Fig. 2A), similar to that of apo-HadA<sub>WT</sub> (PDB code: 6HM) (RMSD of 0.439) previously reported (Fig. S1) (16). Each protomer is composed of three domains: the N-terminal domain (residues 1−146), the β-sheet domain (residues 147−275), and the C-terminal domain (residues 276−517) (Fig. 2B). The co-complex with FAD<sup>+</sup> and 4NP allowed identification of four binding sites for ligands in which strong electron densities could be observed for all four FAD<sup>+</sup> molecules, whereas only one 4NP molecule could be clearly seen in the expected pocket of subunit A (Fig. 2B, Inset). Of interest, two 4NP molecules were found to locate to the dimer interface between subunits C and D. The key difference between the HadA<sub>WT</sub>−FAD<sup>+</sup>−4NP structure reported here and the previously reported apo-HadA<sub>WT</sub> structure is the loop closure upon FAD<sup>+</sup> binding (22). In contrast, this loop was not observed in the structure of p-hydroxyphenylacetate (4HPA) (23), in which C<sub>2</sub> can bind well to all types of flavins including FMNH<sup>+</sup>, FAD<sup>+</sup>, and reduced riboflavin (24−26). Therefore, this structural feature may be a universal motif specific for selectively binding FAD<sup>+</sup> in two-component flavin-dependent monooxygenases.

The ribose moiety of adenosine is hydrogen bonded to Arg<sub>326</sub> and Asp<sub>383</sub> in the C-terminal domain of the neighboring subunit of the dimer, whereas the adenine ring resides within the van der Waals contacts of Ile<sub>315</sub>, Thr<sub>317</sub>, Ile<sub>320</sub>, Val<sub>323</sub>, Leu<sub>384</sub> of the neighboring C-terminal domain and in a water tunnel of the tetrameric inner-interface. The 2D scheme for HadA−FAD<sup>+</sup> interactions in the presence of 4NP within its active site is shown in Figure 3B.

Table 1

| Complex          | HadA<sub>WT</sub>−FAD<sup>+</sup>−4NP | HadA<sub>Pha</sub><sub>441Val</sub>−FAD<sup>+</sup>−4NP |
|------------------|----------------------------------|---------------------------------|
| Wavelength (Å)   | 1.54                             | 1.54                            |
| Space group      | P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub> | P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub> |
| Unit cell parameters | a = 97.8, b = 97.8, c = 97.8, α = 97.8, β = 97.8, γ = 90° | a = 97.8, b = 97.8, c = 97.8, α = 97.8, β = 90° |
| Molecules in asymmetric unit | 4 | 4 |
| Resolution (Å)  | 20.74−2.30                      | 21.10−2.30                       |
| No. of measured reflections | 583,953                      | 583,953                          |
| No. of unique reflections | 118,305                       | 118,305                          |
| Redundancy      | 4.9 (3.04)                       | 4.9 (3.04)                       |
| Completeness (%)| 99.1 (98.4)                      | 99.6 (98.8)                      |
| R<sub>merge</sub> | 10.2 (2.8)                      | 10.3 (2.9)                       |
| Wilson B factor (Å<sup>2</sup>) | 17.5                        | 17.3                            |
| R<sub>factor/R<sub>free</sub></sub> | 20.71/26.25                  | 21.10/26.25                     |
| Average B-factor (Å<sup>2</sup>) | 20.9                        | 15.1                            |
| R.m.s. bond deviation (Å) | 0.01                        | 0.01                             |
| R.m.s. angle deviation (°) | 1.67                       | 1.65                             |
| Favored, Allowed, Outliers | 90.6, 9.1, 0.2 | 91.4, 8.4, 0.2 |
| PDB code        | 7E8P                            | 7E8Q                            |

The overall FAD<sup>+</sup>-binding site of HadA<sub>WT</sub>

FAD<sup>+</sup> binds to each subunit in a tabular-shaped pocket, created by the N-terminal and β-sheet domains, and extends to the C-terminal domain. The adenosine moiety protrudes out toward the dimer interface and is stabilized by polar interactions and hydrophobic residues of the dimer (Fig. 3A). The loop of residues 157 to 170 (Fig. 2B) forms a lid to confine the bound FAD<sup>+</sup> via interactions with an adenosine diphosphate moiety. This loop is designated as the “flavin-binding loop” that was previously identified in other two-component flavin-dependent enzymes (21, 22). Gln158 and Arg161, which are parts of the flavin-binding loop, and Arg387 (from a neighboring subunit) anchor the diphosphate backbone via salt-bridge and hydrogen-bonding interactions. The geometry of these arginine side chains is strengthened by salt-bridge interactions with Asp156. Similar to the structure of p-hydroxyphenylacetate (4HPA) (23), in which C<sub>2</sub> can bind well to all types of flavins including FMNH<sup>+</sup>, FAD<sup>+</sup>, and reduced riboflavin (24−26). Therefore, this structural feature may be a universal motif specific for selectively binding FAD<sup>+</sup> in two-component flavin-dependent monooxygenases.

The ribose moiety of adenosine is hydrogen bonded to Arg<sub>326</sub> and Asp<sub>383</sub> in the C-terminal domain of the neighboring subunit of the dimer, whereas the adenine ring resides within the van der Waals contacts of Ile<sub>315</sub>, Thr<sub>317</sub>, Ile<sub>320</sub>, Val<sub>323</sub>, Leu<sub>384</sub> of the neighboring C-terminal domain and in a water tunnel of the tetrameric inner-interface. The 2D scheme for HadA−FAD<sup>+</sup> interactions in the presence of 4NP within its active site is shown in Figure 3B.

Interactions around the flavin isoalloxazine ring in the structure of HadA<sub>WT</sub>−FAD<sup>+</sup>−4NP

The isoalloxazine ring of FAD<sup>+</sup> binds deeply at the base of the tabular pocket and interacts with residues of both subunits of the dimer (Fig. 3A). The aromatic ring of the isoalloxazine moiety makes van der Waals contacts with Ile<sub>191</sub>, Val<sub>442</sub>, Phe<sub>443</sub>, Phe<sub>446</sub>, and Arg<sub>387</sub> and Ile<sub>391</sub> of the neighboring subunit, whereas the ribityl side chain forms hydrogen bonds to the backbones of Phe<sub>446</sub> and Asn<sub>447</sub>. The 2-carbonyl (O2) and 3-NH(H(N3)) groups of the isoalloxazine are hydrogen bonded to the amino and carboxyl backbones of Phe<sub>446</sub> and Pro<sub>152</sub> with distances of 2.8 and 3.1 Å, respectively, whereas the 4-carbonyl (O4) and 5-NH(H(N5)) are fixed by hydrogen bonding from side chains of Arg<sub>101</sub> (2.7 Å) and Thr<sub>193</sub> (3.1 Å), respectively. The position of a guanidine side chain of Arg<sub>101</sub> is controlled by a nearby Asn<sub>447</sub> side chain (3.0 Å).

The hydroxyl group of Thr<sub>193</sub> is 2.7 Å from the carboxyl ion of Asp<sub>254</sub>, which is salt bridged to the guanidine side chains of Asp<sub>254</sub> and 3-NH(H(N3)) groups of the isoalloxazine are hydrogen bonded to the amino and carboxyl backbones of Phe<sub>446</sub> and Pro<sub>152</sub> with distances of 2.8 and 3.1 Å, respectively, whereas the 4-carbonyl (O4) and 5-NH(H(N5)) are fixed by hydrogen bonding from side chains of Arg<sub>101</sub> (2.7 Å) and Thr<sub>193</sub> (3.1 Å), respectively. The position of a guanidine side chain of Arg<sub>101</sub> is controlled by a nearby Asn<sub>447</sub> side chain (3.0 Å).

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Arg233 and Arg439. This FADH$^-\text{con}$figuration suggests that a molecular O$_2$ would diffuse to reach the C4a atom on the re-face to form C4a-hydroperoxy-FAD; this geometric mode of oxygen reaction is also commonly found in other two-component flavin-dependent monooxygenases (27).

The interactions contributed by Thr193 together with Asp254, Arg233, and Arg439 should be important for the flavin chemistry of HadA, as they are directly involved in the alteration of the hybridization of the isoalloxazine N5 during the reaction. To identify the functions of these residues, four HadAThr193 and two HadAAsp254 variants were constructed by site-directed mutagenesis (listed in Table. S1). Three HadAThr193 variants, including HadAThr193Ala, HadAThr193Val, and HadAThr193Ser, could be expressed in soluble form. In contrast, HadAThr193Cys could only be expressed in inclusion bodies. Both HadAAsp254Ala and HadAAsp254Asn could also be expressed in soluble form. Thus, the catalytic properties of the soluble enzymes were further characterized by monitoring 4NP consumption and by performing transient kinetics studies (Experimental procedures).

Results showed that HadAThr193Ala and HadAThr193Val could not utilize 4NP as a substrate (Fig. 4). Kinetic data of HadAThr193Ala and HadAThr193Val revealed that only free FADH$^-\text{oxidation}$ (Fig. S3) was observed when mixing FADH$^-$ with these enzymes, indicating impairment in their FADH$^-\text{binding}$. These two variants did not show C4a-hydroperoxy-FAD formation as in the HadAWT (Fig. 5, A–C). For the HadAThr193Ser variant, its 4NP consumption activity was 72% relative to HadAWT (Fig. 4). Kinetic traces of the HadAThr193Ser reaction with aerobic 4CP (Fig. 5D) revealed two observable phases (0.01 – 1 s and 1 s – 100 s) when detected at wavelength 450 nm ($\lambda_{450}$). These kinetic phases indicate oxidations of free FADH$^-\text{adduct formation}$ (84%) and the HadA-bound FADH$^-\text{adduct formation}$ (16%) species. A small amount of C4a-adduct formation could be observed at wavelength 380 nm ($\lambda_{380}$). Therefore, the HadAThr193Ser variant could maintain partial ability to bind FADH$^-$ but with less efficiency than HadAWT. This indicates the importance of the hydroxyl group of residue 193, which is required for interacting with H(NS) of FADH$^-$. Therefore, we propose that the -OH group of Thr193

Figure 2. Crystal structure of the HadAWT–FADH$^-\text{4NP complex}$. A, tetrameric quaternary structure of HadA monooxygenase co-complexed with FADH$^-$ and 4NP. B, three domains of the HadA subunit include the N-terminal domain (green part), the β-sheet domain (orange part), and C-terminal domain (purple part). The flavin-binding loop (residues 157–170) is labeled in red. Inset in (B) is an electron density map of FADH$^-$ and 4NP bound in the active site. 4NP, 4-nitrophenol.
directly interacts with H(N5) of FADH\(^-\) for binding and stabilization of FADH\(^-\) and for formation of its intermediates. Apart from Thr193, Asp254 is also important for FADH\(^-\) binding. Disruption of hydrogen bonding networks in the variants HadAAsp254Ala and HadAAsp254Asn also resulted in complete loss of 4NP consumption activity (Fig. 4). Based on kinetic data, these variants also lose FADH\(^-\) binding ability because only free FADH\(^-\) oxidation was observed in their reactions (Fig. 5, E and F). We propose that hydrogen bonding and salt-bridge interactions between the Thr193 side chain and neighboring residues such as Asp254, Arg233, and Arg439 may render greater negative charge on the hydroxyl group of Thr193, leading to stronger interactions with H(N5) on the surface of the isoalloxazine ring.

A unifying FADH\(^-\) binding feature found in HadA and other two-component flavin-dependent monooxygenases

To analyze whether the nature of the amino acids found in the FADH\(^-\)–binding pocket of the ternary complex structure of HadA is also common in other group D monooxygenases, amino acid sequences and structures of these enzymes were analyzed. Sequences of (1) flavin-dependent monooxygenases catalyzing hydroxylation plus dehalogenation and denitration activities (HadA monooxygenase from \(R.\) \(p\)ickettii [this study]; (2) flavin-dependent monooxygenases catalyzing hydroxylation plus dehalogenation (\(R\). \(e\)utropha JMP134 (15, 28), \(C\). \(t\)cpA from \(C\). \(i\)nangolens \(X\)1\(^{T}\) (29), \(Tf\) from \(B.\) \(c\)epacia \(A\)C1\(100\) (17, 30), \(C\). \(p\)h-p from \(A\). \(c\)lorophenolicus \(A\)6 (31), \(D\). \(c\)mB1 from \(R\). \(p\)hodococcus \(s\). \(p\)J\(-\)3 (32), and \(H\). \(n\)pA from \(C\). \(i\)nangolens sp. \(C\)NP-8 (33)); (3) flavin-dependent monooxygenases catalyzing hydroxylation plus denitration (NpcA from \(R\). \(o\)pacus \(S\)AO101 (34), NpdA2 from \(A\). \(r\)throbacter sp. \(S\)train \(J\)443 (35), and NpsA1 from \(R\). \(p\)hodococcus sp. strain \(P\)1 (36)); and (4) flavin-dependent monooxygenases catalyzing merely hydroxylation.
of phenolic compounds such as phenol and 4-hydroxybenzoylacetate (THpaB from *T. thermophilus* HB8 (22), EcHpaB from *Escherichia coli* (37), PheA1 from *Bacillus thermoglucosidasius* A7 (38), and C2 from *A. baumannii* (24, 26)) were analyzed by the Clustal omega software (EMBL-EBI) (Fig. S4). Sequences that are similar to, or different from, HadA were identified. Their available enzyme X-ray structures were also compared with that of HadA to identify structural features associated with specific functions. Particularly, the C2 structure is important for identifying features specific for FADH\(^-\) binding because unlike other enzymes, C2 can also bind to FMNH\(^-\). Therefore, we propose that these residues are important for discriminating between FADH\(^-\) and FMNH\(^-\) binding and that they are an important feature responsible for the enzyme selectivity for only binding FADH\(^-\).

For residues surrounding the isoalloxazine ring, all FADH\(^-\)-utilizing monooxygenases in class D have residues equivalent to Arg101, Thr193, Arg233, Asp254, and Arg439 of HadA and their structural configurations are also conserved in the same manner (15–17, 21, 22) (Fig. S5, A–E). This is different from C2 in which Ser171 (in C2) instead of threonine was identified as being important for FMNH\(^-\) binding and C4a-hydroperoxy intermediate stabilization by H-bond interactions with both

**Figure 5. Transient kinetics of HadA variants.** Rapid kinetics of reactions of HadA\(^\text{WT}\), FADH\(^-\) (75 and 25 μM, respectively) mixing with an aerobic solution of 4-chlorophenol (0.5 mM 4-chlorophenol and 0.13 mM O\(_2\)) using the single mixing mode of a stopped-flow apparatus. Absorption changes at wavelengths 380 nm (solid line) and 450 nm (dashed line) were monitored to observe the C4a-hydroperoxy-FAD intermediate and oxidized FAD, respectively. C4a-hydroperoxy-FAD absorbs mainly at A\(_{380}\), whereas oxidized FAD absorbs at both wavelengths. HadA variants are (A) HadA\(^\text{WT}\), (B) HadA\(^{\text{Thr193Ala}}\), (C) HadA\(^{\text{Thr193Val}}\), (D) HadA\(^{\text{Thr193Ser}}\), (E) HadA\(^{\text{Asp254Ala}}\), (F) HadA\(^{\text{Asp254Asn}}\), (G) HadA\(^{\text{His290Ala}}\), (H) HadA\(^{\text{His290Cys}}\), and (I) HadA\(^{\text{His290Asn}}\). Inset in (A) and (I) are spectra of the C4a-hydroperoxy-FAD intermediate detected at 10 s of reaction time.
H(N5) and O4 of FMNH− (39). The hydroxyl side chain of Ser171 is stabilized by hydrogen bonding with two adjacent serine residues (23) instead of the salt bridge found in HadA (Fig. S5F). Although reactions of HadA homologs have not been investigated by transient kinetics, we can infer that these conserved residues likely function similarly to those of HadA in that they are important for facilitating the reaction of FADH− and oxygen. Residues of HadA important for its catalytic functions are highlighted in the figure generated by WebLogo (40) (Fig. 6).

**Identification of hydrophobic interactions for 4NP binding**

In the active site of HadA, the 4NP molecule binds in a hydrophobic pocket (consisting of Val155, Phe206, Phe286, Thr449, and Leu457) located between the β-sheet and C-terminal domains of each subunit. The aromatic ring of 4NP forms contacts with Phe206 face-to-face with π-π stacking and sits in van der Waals contact of Phe286, which forms π-edge interactions with Phe206 (Fig. 3). The hydrophobic environment found in the HadA active site is similar to the environment of an empty pocket speculated to be the phenol-binding pocket in apoenzyme structures of dehalogenating flavin-dependent enzymes previously crystallized (Fig. S6) (15–17).

We tested the importance of hydrophobic residues surrounding the 4NP-binding pocket using site-directed mutagenesis to decrease the hydrophobicity of the pocket via replacing the aromatic side chain of Phe206 and Phe286 with aliphatic side chains of Val, Leu, Ile, or Ala. The variants HadA Phe206 Val, HadA Phe206 Leu, HadA Phe286 Ile, HadA Phe286 Ala, HadA Phe286 Val, HadA Phe286 Leu, and HadA Phe286 Ile showed a drastic decrease in 4NP conversion activities (Fig. 4). Molecular docking of 4NP into the active sites of HadA WT and Phe206 and Phe286 variants of HadA performed by AutoDock Vina to determine the binding energy (Experimental Procedures) showed that the affinities of 4NP in HadA Phe206 variants decreased because the π-π stacking interactions between 4NP and Phe206 were lost. Although the binding affinities of 4NP in the HadA Phe286 variants were not significantly perturbed (Fig. S7), the denitration activities of these HadA Phe286 variants were impaired. The data suggest that, in addition to the ability to bind, proper geometric arrangement of 4NP in the active site is also important for hydroxylation and denitrination of 4NP by HadA. Altogether, we demonstrated that the aromatic side chains of Phe206 and Phe286 are important for promoting the correct configuration of 4NP binding in HadA catalysis.

When apoenzyme structures of enzymes (Re TcpA and TtfD) with similar activities to HadA were analyzed, the residues Phe206 and Phe286, along with other hydrophobic residues such as Val155 and Leu457, could be identified. Based on this information and the HadA structure, we propose the substrate-binding regions of these enzymes as shown in Figures 6 and S4.

**His290 is a key catalytic residue for electrophilic aromatic substitution**

In the 4NP-binding site, a polar hydroxyl group of 4NP interacts closely with His290 (2.5 Å) on helix α9, whereas the nitro (−NO2) moiety of 4NP points toward a guanidine side chain of Arg208 (3.4 Å). Based on apoenzyme structures, His290 in HadA and His289 in TtfD were previously proposed as a general base to abstract a proton from a phenolic substrate, which triggers delocalization of a lone pair of electrons from O1 to the C4 position of the substrate, facilitating monoxygenation by C4a-hydroperoxy-FAD (shown in Fig. 1) (8, 17, 41). To elucidate the role of His290 in catalysis, we constructed and overexpressed nine His290 variants (listed in Table. S1). Only HadA His290 Ala, HadA His290 Cys, and HadA His290 Asn could be overexpressed as soluble enzymes, and these enzymes were purified to homogeneity. HadA His290 Ala and HadA His290 Cys exhibited a complete loss of 4NP conversion activity, whereas HadA His290 Asn showed a very low 4NP consumption activity (Fig. 4).

We carried out stopped-flow investigation of these three variants and found that transient kinetics of HadA His290 Ala and HadA His290 Cys showed mixed kinetics of free and enzyme-bound FADH− oxidation (Fig. 5, G and H). However, formation of C4a-hydroperoxy-FAD could not be detected in these variants, explaining why these enzymes are completely inactive. Of interest, the kinetic traces of the HadA His290 Asn reaction showed three observable phases (Fig. 5I). The first phase (0.001–0.4 s) showed simultaneous increase in A380 and A450 with an observed rate constant of ~6 s−1 similar to kinetics of
Structures of a HadA flavin-dependent dehalogenase

free FADH\(^{-}\) oxidation (Fig. S3). This phase was interpreted as oxidation of unbound FADH\(^{-}\). The second phase (0.4–20 s) exhibited only a large increase in \(A_{430}\) with almost no change in \(A_{450}\). This phase likely corresponded to formation of a C4a-hydroperoxy-FAD intermediate with an observed rate constant of \(~\sim 2\) s\(^{-1}\). The absorption characteristics of the intermediate formed at this phase fits well with that of C4a-hydroperoxy-FAD (Fig. 5I, inset). The last phase was an increase of \(A_{450}\) and decrease of \(A_{380}\), indicating that this phase was likely a decay of the C4a-hydroperoxy-FAD intermediate with generation of oxidized FAD with an observed rate constant of \(~\sim 0.002\) s\(^{-1}\). The decay rate of C4a-hydroperoxy-FAD in HadAHis290Asn (0.002 s\(^{-1}\)) is in the same range as that of C4a-hydroperoxy-FAD decay in HadAWT (0.007 s\(^{-1}\)) (7). We could not detect product from a single turnover reaction of HadAHis290Asn:FADH\(^{-}\) carried out in an aerobic solution of 4CP (Experimental Procedures) (data not shown). This suggests that, although C4a-hydroperoxy-FAD can be formed in HadAHis290Asn, the enzyme mostly eliminates H\(_2\)O\(_2\) via an unproductive pathway without catalyzing dehalogenation/denitration, possibly due to the lack of His290. Therefore, we conclude that His290 is important for stabilization of C4a-hydroperoxy-FAD as well as for facilitating product formation in dehalogenation/denitration reactions as previously proposed (8, 17, 41).

Of interest, His290 is only conserved among mono-oxygenases with additional dehalogenation and denitration activities but not in monoxygenases catalyzing only hydroxylation (Fig. 6 and Fig. S4). Therefore, this histidine residue should be key for enabling dehalogenation and denitration reactions. Its position is also different from the catalytic base found in enzymes catalyzing ortho-hydroxylation of 4HPA, such as Tyr104 and His142 for THpaB (22) and His120 for C\(_2\) (42).

Proposed mechanisms of hydroxylation and group elimination by HadA

The combined results from structural analysis and site-directed mutagenesis experiments suggest a possible model for HadA reaction mechanisms as described in Figure 7. In the first step, FADH\(^{-}\) binds to HadA such that the flavin H(N5) forms a hydrogen bond to the Thr193 hydroxyl side chain in which its hydroxy dipole is polarized by the hydrogen bond network of the neighboring residues Asp254, Arg233, and Arg439. The next step is diffusion of O\(_2\) toward the re-side of FADH\(^{-}\) to form a C4a-hydroperoxy-FAD intermediate. The substrate then binds to the aromatic cage formed by Phe206 and Phe286 with its C4 position (para-position) pointing toward the flavin C4a position.

We noted that, in the HadAWT complex structure (Fig. 3), the substrate C4 position is 9 Å away from the C4a position of FADH\(^{-}\), which is significantly farther than the distance between the substrate hydroxylation site and the C4a-position in C\(_2\) and THpaB structures (5.0 and 4.9 Å, respectively) (22, 23). We propose that significant conformational changes in the HadA structure possibly occur after C4a-hydroperoxy-FAD is formed to allow the terminal –OOH moiety of C4a-hydroperoxy-FAD to move closer to the C4 position of the substrate. Based on our previous kinetic investigation (7, 8), we found that, for active hydroxylation, the substrate should bind after C4a-hydroperoxy-FAD is formed. Therefore, the structure obtained likely does not represent the near attack conformation. Subsequently, the flavin, 4NP, and residues around the active site need to be rearranged in order to accommodate an oxygen molecule used for the formation of C4a-hydroperoxy-FAD. Furthermore, the 4NP substrate position needs to be reoriented in order to be ready for reacting with the C4a-hydroperoxy FAD intermediate.

For the hydroxylation step, His290 abstracts a proton from the hydroxyl group of a phenolic substrate to facilitate hydroxylation and group elimination to form BQ and C4a-hydroxy-FAD as products. Our previous investigation on the mechanism of HadAWT has shown that the deprotonation of aromatic substrates is important for controlling HadA catalysis (8). As His290 is the only residue surrounding 4NP that has the potential to act as a catalytic base at the working pH (pH 7.5), it is the most probable candidate responsible for substrate deprotonation. Dehydration of C4a-hydroxy-FAD occurs prior to release of both BQ and oxidized FAD from the HadA active site to complete the catalytic cycle.

Unusual 4NP-binding site at the subunit interface of HadAWT

Apart from electron density of 4NP in the active site, we observed additional electron density of two 4NP molecules bound at the dimer interface between subunits C and D (Fig. 8, A and Fig. S8A). The area around this region consists of many hydrophobic side chains forming a "hydrophobic cage" that captures two 4NP molecules. The aromatic moieties of the 4NP molecules are sandwiched with the side chains of Phe441 from subunits C and D to perfectly form face-to-face \(\pi\)-\(\pi\) or quadruple \(\pi\)-stacking (Phe441\(_C\) = 4NP \(\approx\) Phe441\(_D\)) interactions (Fig. 8A). This intermolecular hydrophobic interaction is not crucial for assembly of the four subunits because apo-HadAWT can efficiently form a tetramer in the absence of 4NP (16). As Phe441 and the quadruple \(\pi\)-stacking interaction are located at the \(\alpha\)15 helix near the FADH\(^{-}\)-binding site, we hypothesize that this interaction may be the cause of the dead-end complex formation seen for HadA, which hinders procession to the productive pathway when the substrate first binds to the enzyme prior to flavin binding.

Disruption of quadruple \(\pi\)-stacking in the HadAPhe441Val variant

To further investigate the role of Phe441 and 4NP binding in substrate inhibition, the quadruple \(\pi\)-stacking of 4NP and Phe441 were disrupted by mutating phenylalanine to aliphatic residues including HadAPhe441Val, HadAPhe441Leu, and HadAPhe441Ile. The three-dimensional structure of HadAPhe441Val–FADH\(^{-}\)–4NP was also solved at a 2.3-Å resolution (PDB code: 7E8Q) using a similar approach as used for the structure of HadAWT–FADH\(^{-}\)–4NP (Fig. S9). Data and refinement statistics of this variant three-dimensional
structure are listed in Table 1. The overall structure of the ternary complex of HadA_{Phe441Val} is very similar to that of HadA_{WT} (RMSD of 0.143) without any significant changes in the active site area (Fig. S10). Electron density of two 4NP molecules are found in the same hydrophobic tunnel at the interface between the two subunits of the HadA_{Phe441Val} dimer; however, the orientation of the two 4NPs is altered by \(-\pi/2\) from those observed in HadA_{WT} (Fig. S8B). Thus, \(\pi-\pi\) interaction cannot occur in the variant. The two 4NPs bind \(\text{via}\) van der Waals interactions with nearby residues on the \(\alpha_{15}\) helix with a nitro group (-NO\(_2\)) of 4NP hydrogen bonded with a carboxamide side chain of Asn445 (Fig. 8B).

### No dead-end complex formation in HadA_{Phe441} variants

We then further tested whether substrate inhibition by the formation of a dead-end complex still exists in these variants by preincubating the enzymes (HadA_{Phe441Val} and HadA_{Phe441Leu}) with various concentrations of 4CP (0.1–6.4 mM) for 5 min to ensure complete formation of the dead-end complex (\(k_{on}\) of 25 M\(^{-1}\) s\(^{-1}\) for HadA_{WT}) (7). The solutions were then rapidly mixed with oxygen to monitor their ability to form C4a-hydroperoxy-FAD using stopped-flow experiments as previously explored in the HadA_{WT} (7). For the reaction of HadA_{WT} (Fig. 9, A), increasing the 4CP concentration significantly altered the kinetics of the reaction, as monitored at both \(A_{380}\) and \(A_{450}\). Incubation with high concentrations of 4CP showed that the amplitude of \(A_{380}\) at 1 s was decreased concomitantly with an increase in amplitude of \(A_{450}\) observed with \(t_{1/2}\) \(\sim 0.2\) s. At the highest concentration of 4CP, the kinetics of the reaction monitored at both \(A_{380}\) and \(A_{450}\) were nearly identical, indicating that these changes were due to the formation of oxidized FAD directly without formation of C4a-hydroperoxy-FAD.

In contrast, the signals and amplitude of both \(A_{380}\) and \(A_{450}\) in both HadA_{Phe441Val} and HadA_{Phe441Leu} were independent of 4CP concentrations under the same conditions as HadA_{WT} (Fig. 9, B and C), indicating that the binding of 4CP does not prevent C4a-hydroperoxy-FAD formation. In both variants, the rates of C4a-hydroperoxy-FAD formation were slightly slower at a 4CP concentration of 6.4 mM (blue bold line), whereas the rate of FAD oxidation at the final phase was almost constant (red dashed line). Kinetic traces of the HadA_{Phe441Leu} reactions also showed a similar trend (Fig. S11).
All data suggest that no substrate inhibition via dead-end complex formation was observed in these HadAPhe441 variants, which are void of the quadruple π-stacking complex at the subunit interface.

Hydroquinone (HQ) product from the reactions of HadA-Phe441Val and HadAPhe441Leu was analyzed and found to be constant over a range of 4CP concentrations of 0.1 to 6.4 mM 4CP without any signs of substrate inhibition (Fig. 9D). These data were significantly different from the reaction of HadA WT. These results agreed well with the stopped-flow experiments discussed above in which no substrate inhibition was observed from the behavior of C4a-hydroperoxy-FAD formation. Altogether, the data indicate that for HadAPhe441 variants in which the π-π-π-π complex at the subunit interface is interrupted, no dead-end complex or substrate inhibition could be observed. These enzymes should be able to perform superior catalysis than the HadAWT because they can avoid being trapped in the dead-end enzyme–substrate complex.

Figure 8. Intermolecular binding of 4-nitrophenol at the interface of HadA dimer. A, a quadruple π-stacking, an unusual π-π-π-π interaction formed by two 4-nitrophenol (4NP) substrates sandwiched with two Phe441 residues of the HadAWT dimer observed in the HadAWT–FADH–4NP structure. B, disruption of a quadruple π-stacking in the HadAPhe441Val–FADH–4NP structure. The lower panel shows side chains of residues from subunit C (yellow) and subunit D (blue) residing within 4 Å distances from the two 4NP molecules bound in the cavity of the subunit interface.

Protein rigidity in HadAWT caused by binding of 4NP

To understand how HadAWT is affected by the binding of 4NP at the subunit interface, we performed thermofluor assays to determine the melting temperatures ($T_m$) of HadAWT and HadAPhe441 variants in the presence and absence of 4NP. The $T_m$ values are summarized in Table 2 with melting curves of all enzymes shown in Figure S12. The results showed that the $T_m$ values of the apo forms of HadAWT and the three HadAPhe441 variants were similar (~48 °C). The data indicate that mutation of Phe441 did not alter the structural stability of the apoenzyme. However, the $T_m$ value of HadAWT–4NP (in which a quadruple π-stacking exists) was 52 °C which is 4.2 °C higher than that of apo-HadAWT. Of interest, the complexes of HadAPhe441 variants and 4NP (in which 4NP can bind but the quadruple π-stacking is absent) showed increased $T_m$ values ($ΔT_m$) of only around 1.5 to 2.1 °C. Therefore, the $T_m$ results imply that, upon 4NP binding at the subunit interface, especially with the quadruple π-stacking found in the HadAWT, the quaternary structures of HadA are likely more rigid than the apoenzyme forms. This is indicated by the increase in $T_m$ values. The rigidity of the HadAWT–4NP complex likely obstructs conformational flexibility particularly at helix α15, leading to the substrate inhibition phenomenon.

Substrate inhibition mechanism in HadA

The data in Figure 9 indicate that HadAPhe441 variants are productive over a wide range of 4CP concentrations without showing significant substrate inhibition. The difference between the structures of HadAWT and HadAPhe441Val around the subunit interface areas (Fig. 8) and $T_m$ values (Table 2) clearly explain how the dead-end complex is stabilized in
HadAWT but not HadA_{Phe441Val}. The unusual face-to-face quadruple π-stacking interactions of four aromatic moieties is the site of “dead-end” complex stabilization, which prevents the formation of C4a-hydroperoxy-FAD, causing inactivation of HadAWT. As the distance between the C1 position of 4NP in the productive and the dead-end binding sites is ~20.8 Å apart (Fig. S13), the inhibition is not caused by direct blocking of 4NP binding to the active site. We propose that the binding of 4NP at the subunit interface may trigger a conformational change in the protein, especially at the α15 helix of the HadA structure, obstructing the proper catalytic site or rigidifying the protein such that it loses its flexibility to properly assume the dynamic changes required for progression of the reaction.

We then searched among enzymes in the same family as HadA to identify whether this feature is prevalent in other systems. The analysis showed that the presence of phenylalanine at the subunit interface is conserved in HadA, ReTcpA, CcTcpA, TrD, and HnpA dehalogenases, whereas for other enzymes including CphC-I, DcmB1, NpsA1, NpcA, NpdA2, PheA1, TtHpaB, and EcHpaB, a glutamate is conserved at this position (Fig. 6 and Fig. S4). At the subunit interface of TtD’s structure (PDB: 3HWC) (17), two phenylalanines identified have the potential to form π-π-π-π interactions with a phenolic substrate similar to HadAWT. In contrast, the negatively charged glutamate in EcHpaB interface (PDB: 6EBO) does not allow the dead-end complex formation by the quadruple π-stacking (21, 37) (Fig. S14). We propose that this unusual quadruple π-stacking protein–ligand interaction is a distinct property specifically found in HadAWT and closely related enzymes.

The improvement of HadA catalysis by HadA_{Phe441} variants

We further explored whether this understanding of the structural features causing substrate inhibition and dead-end complex formation in HadA can be used to improve HadA biocatalysis and its applications. Reactions of HadA_{Phe441} variants and HadA WT were carried out to compare their abilities to detoxify 4NP. The molar ratio between 4NP and HadA was set as 1000:1 to attain conditions in which 4NP significantly inhibits the reaction of HadAWT. The results

**Table 2**

Melting temperature (Tm) of HadAWT and HadA_{Phe441} variants

| HadA variants | Melting temperature, Tm (°C) | ΔTm (°C) |
|---------------|-----------------------------|----------|
|               | Apoenzyme                   | With 4NP | Apoenzyme |
| HadAWT        | 48.1 ± 0.9                  | 52.3 ± 0.6 | 4.2 ± 0.5 |
| HadA_{Phe441Val} | 47.3 ± 0.7                  | 49.3 ± 0.8 | 2.1 ± 0.4 |
| HadA_{Phe441Leu} | 48.2 ± 0.7                  | 49.6 ± 1.1 | 1.5 ± 0.5 |
| HadA_{Phe441Ile} | 48.9 ± 0.5                  | 50.5 ± 2.2 | 1.7 ± 0.9 |

**Figure 9. Regulation of dead-end complex formation by Phe441.** An anaerobic solution of FADH⁺ (25 μM) was rapidly mixed with an air saturated (A) HadAWT, (B) HadA_{Phe441Val}, or (C) HadA_{Phe441Leu} (75 μM) that was preincubated with 4CP (0.1–6.4 mM) in 20 mM Hepes pH 7.5. Absorption changes at wavelengths 380 (blue solid line) and 450 nm (red dash line) were monitored to detect formation of the C4a-hydroperoxy-FAD intermediate and oxidized FAD, respectively. C4a-hydroperoxy-FAD absorbs mainly at 380 nm, whereas oxidized FAD absorbs at both wavelengths. D, HQ product formation at various concentrations of 4-chlorophenol (4CP) catalyzed by HadA_{WT} (black circle), HadA_{Phe441Val} (blue square), and HadA_{Phe441Leu} (red diamond).
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showed that multiple turnover reactions of HadAPhe441 variants were more superior than that of HadAWT. Within 1 h, the HadAPhe441 variants could convert 4NP into BQ at a conversion rate of 100%, whereas only 90% of the 4NP was depleted in the HadAWT reaction (Fig. 10). The rates of 4NP conversion by HadAPhe441Val, HadAPhe441Leu, and HadAPhe441Ile were 21 ± 8%, 24 ± 13%, and 17 ± 10% greater than that of HadAWT, respectively (p-value < 0.05) (Fig. 10, inset). The data indicate that alteration of Phe441, which removes the quadruple π-stacking interactions and, thus, dead-end complex formation ability, indeed improved biocatalysis by HadA.

Discussion

The work herein reports the first full-package structure of the group D flavin-dependent monooxygenase (HadA) that catalyzes dehalogenation/denitration in addition to hydroxylating. The ternary complex obtained clearly identified important interactions, which were also confirmed by site-directed mutagenesis and stopped-flow experiments. The structural analysis also revealed another binding site of 4NP, which prevents C4a-hydroperoxy-FAD formation and results in the formation of a dead-end complex. Construction of variants in which 4NP inhibition site was perturbed resulted in variants with improved catalytic activities because the substrate inhibition was abolished.

The tetrameric quaternary structure of HadA monooxygenase co-complexed with FADH− and 4NP is similar to that of apo-HadA, apo-ReTtcpA, and apo-TtFl (15–17), which share an acyl-CoA dehydrogenase fold like other enzymes in the group D two-component flavin-dependent monooxygenases, including C2-FMNH−–HPA, TtHpaB-FAD-HPA, and EcHpaB-HPA, previously reported (10, 11, 14, 21–23) (Fig. S15). These data indicate that the binding of ligands does not affect oligomerization of HadAWT because all forms of the apo or ligand-bound enzymes are tetramers. The FADH−-binding region in the HadA active site is also similar to those of group D FADH−-utilizing monooxygenases. Binding of FADH− is involved in positioning of the flavin-binding loop (Fig. 2B), particularly at residues 157 to 170 because the loop interacts with the adenosine diphosphate moiety of the bound FADH−. The flavin-binding loop does not interact closely with the nearby subunit since the distances between the closest regions of the flavin loop (residues 157–160) and the nearby subunit residues (residues 410–415) are longer than 3.5 Å. Specific functions of amino acid residues surrounding FADH− investigated by site-directed mutagenesis, transient kinetics, and product analysis indicate that the bound isoalloxazine ring is stabilized by hydrogen bonding interactions with conserved amino acids such as threonine (Thr193), aspartate (Asp254), and three arginine (Arg101, Arg233, and Arg439) residues. These structural features for FADH− binding are also universal among other two-component monooxygenases in the same family.

On the other hand, the binding pockets of phenol compounds are varied in these enzymes depending on the type of substrates they employ. Since most of their substrates are aromatic compounds, these enzymes including HadA have hydrophobic pockets lined with different hydrophobic amino acid residues. For a single-component flavin-dependent pentachlorophenol hydroxylase (PcpB) from *Sphingobium chlorophenolicum* (member of class A flavin-dependent monooxygenases), which can degrade highly hydrophobic substrates such as pentachlorophenol (43), its binding pocket is rather different from those of the HadA family. The modeled active site of PcpB consists of four phenylalanine residues to maintain hydrophobic interactions with substrate (44, 45). For class D two-component monooxygenases that catalyze hydroxylation, only two X-ray structures are available for the enzyme co-complexed with the flavin and aromatic substrate, i.e., TtHpaB-FAD−HPA from *T. thermophilus* HB8 (PDB code: 2YYI) (22) and C2-FMNH−–4HPA from *A. baumannii* (PDB code: 2BT) (23) with RMS of Cα alignment of 1.773 and 10.105, respectively, are available. Although the overall foldings of these enzymes are similar to that of HadA, the architecture inside their active sites is quite different (Fig. S15) because HadA catalyzes hydroxylation with dehalogenation/denitration at position C4 of phenol substrates, whereas TtHpaB and C2 catalyze only hydroxylation at the ortho-position of p-hydroxyphenylacetate.

The binding of reduced flavins seems to be crucial for phenolic substrate binding. We could not crystalize a binary complex of HadA−4NP or HadA−4CP in the absence of FADH−. X-ray structures of similar binary complexes also could not be obtained for C2 and TtHpaB (22, 23). Kinetic investigation of p-hydroxyphenylacetate hydroxylases from *A. baumannii* and *Pseudomonas aeruginosa* indicates that the substrate 4HPA only binds after the flavin binding (24, 46, 47). These imply that the bound flavin may create a recognition site
for a phenolic substrate. For HadA<sub>WT</sub>, although a phenolic substrate can bind, it leads to the formation of the dead-end complex. Under physiological conditions, as the rate of FADH<sup>−</sup> binding to HadA<sub>WT</sub> is faster than 4CP binding by four orders of magnitude (7), these properties would help prevent the enzyme from getting trapped in the dead-end complex.

The structure reported here also explains the root cause of substrate inhibition, which is a common obstacle in biocatalytic applications. The rigidity of a perfect face-to-face quadruple π-stacking complex formed from four aromatic moieties (two aromatics from phenylalanine side chains and two aromatics from substrates) may interfere with the FADH<sup>−</sup> binding site. This conclusion was confirmed by single point mutations at Phe441 that disrupted this quadruple π-stacking complex, resulting in enzyme variants with improved biocatalytic activities. Usually, the π–π interaction is observed in protein–protein or protein–ligand interactions to facilitate protein function. For example, binding of aromatic substrate in the active site of PcpB from <i>S. chlorophenolicum</i> (44, 45), nicotine oxidoreductase (NicA2) from <i>Pseudomonas putida</i> (48), and also HadA<sub>WT</sub> itself is mediated by π–π interactions. The π–π interactions could also be applied in various biological applications such as in the development of molecular receptors, in the design of controlled drug release systems, and in the fabrication of biosensors (49, 50).

The mechanism of dead-end complex formation in HadA shares some common features with those of other flavin-dependent monoxygenases such as bacterial luciferase and halogenase. Bacterial luciferase from <i>Vibrio harveyi</i> belongs to class C flavin-dependent monoxygenases. Binding of an aldehyde substrate to apo-luciferase blocks the binding of FMNH<sup>−</sup> resulting in a decrease in the light emission reaction (18–20). The equilibrium binding experiments revealed two aldehyde-binding sites at each α and β2 subunit. The stronger-binding aldehyde binds to the active site and acts as a substrate, whereas the weaker-binding aldehyde binds to an inhibitory pocket (51). For flavin-dependent halogenases, RebH from <i>Lechevalieria aerocolonigenes</i> and Thal from <i>Streptomyces alborescens</i>, which are class F monooxygenases, stopped-flow investigation showed that preformation of the RebH–FADH<sup>−</sup> or Thal–FADH<sup>−</sup> complexes prior to binding of a trypto-phan substrate results in abolishment of C4a-hydroperoxylavin formation (52, 53). Molecular dynamics simulations of Thal–FADH<sup>−</sup> showed rearrangement of water molecules upon prolonged incubation of Thal and FADH<sup>−</sup>, which may affect the protonation status of FADH<sup>−</sup> and stabilization of C4a-hydroperoxy-FAD (52). For class A NicA2 from <i>P. putida</i>, a pocket ~12 Å away from the active site can accommodate nicotine binding, which may cause substrate inhibition (48).

As flavin-dependent monoxygenases are attractive for many applications (3, 10–12, 14, 54), identification of residues important for catalysis provides basic knowledge for future rational design in enzyme engineering. As HadA is useful for many applications including detoxification and biodetection (9), the structural basis reported here should be useful for improving HadA for industrial applications. In the past, engineering of C2 resulted in variants useful for biocatalysis. The variant Tyr398Ser of C2 catalyzes formation of trihydroxyphenolic acids with higher efficiency than the wild-type enzyme (55), whereas single mutations at either Ser146 or Arg263 resulted in C2 variants, which can use p-amino-phenylacetate and p-hydroxyphenylethylamine better than the native substrate 4HPA, respectively (56, 57). Mutation of group C flavin-dependent luciferases can also change their properties, such as alteration of emission color or prolonging light intensity of the product, which is useful for biodetection applications (58).

In conclusion, these in-depth structural insights into the catalysis and inhibition mechanisms of HadA flavin-dependent monoxygenase with additional dehalogenating/denitrating activities provide an important basis for better understanding flavin-dependent monoxygenases and for future enzyme engineering to more effectively implement these enzymes in applications including biodetoxification and biodetection of toxic compounds as well as in biocatalysis.

**Experimental Procedures**

**Protein crystallization**

HadA was purified for crystallization according to a reported protocol (7). The purified enzyme was stored in 20 mM Heps pH 7.5 containing 20% (v/v) glycerol and kept in a −80 °C freezer until use. A stock of FAD solution (high concentration) was dissolved in filtered H<sub>2</sub>O. The concentration of FAD was calculated based on the absorption at a wavelength 450 nm using an extinction coefficient ε<sub>450</sub> of 11.3 mM<sup>−1</sup> cm<sup>−1</sup>. Reduced FAD (FADH<sup>−</sup>) was prepared by stoichiometric reduction of FAD using sodium dithionite inside an anaerobic glove box. A stock of 4NP at high concentration was prepared by dissolving in dimethyl sulfoxide (Sigma). All components of crystallizing agents were filtered by passage through syringe filters with 0.22-μm pore size before use. All equipment for crystallization were made anaerobic before being used. All experiments using FADH<sup>−</sup> were performed inside an anaerobic glove box (Belle Technology Ltd) to avoid reoxidation by molecular oxygen.

For crystallization of the HadA–FADH<sup>−</sup>–4NP complex, HadA (0.3 mM), FADH<sup>−</sup> (12 mM), and 4NP (12 mM) were mixed inside the anaerobic glove box. Concentrations indicated were the final concentrations in the protein complex that was incubated for 2 h to ensure complete complex formation prior to performing crystallization. A solution of the protein complexes (1 μL) was mixed with crystallizing agents (1 μL) using the microbatch technique and a 60-well minitray (1 mm diameter at bottom of each well) covered with 6 ml of baby oil (Johnson; a mixture of mineral oil, olive oil, and vitamin E, PZ Johnson). The protein complex was crystallized at 25 °C in small grids of 0.18 to 0.22 M sodium citrate dehydrate (Sigma), 15% to 17% (w/v) polyethylene glycol 3350 or PEG3350 (Hampton research), 5% (v/v) 2,2,2-trifluoroethanol (Fluka) as an additive, and 0.1 M Bis-Tris propane (Sigma) pH 6.5 as the buffer. Crystals of HadA complexes were harvested by quick dipping in the crystallizing agent containing substrate and 15%...
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(v/v) glycerol as cryoprotectants prior to storage in liquid nitrogen.

**Data collection and structure solving**

X-ray diffraction data of HadA complexes were collected at 100 K at a wavelength of 1.54 Å using a D8 Venture single crystal X-ray diffractometer coupled with a PHOTON 100 detector (Bruker). The structure of HadA was determined by molecular replacement using Phaser in the CCP4 suite (59) with a chain A of apo-HadA (PDB code: 6JHM) as a template. Model building and structure refinement were performed using Coot (60) and REFMAC5 (61). The ligand structure was prepared using Hyperchem.

**Site-directed mutagenesis and preparation of HadA variants**

Site-directed mutagenesis was performed using PCR and the hadAWT-pET-11a plasmid as a template. In brief, the solutions (50 μl) of ~50 ng of hadAWT-pET-11a plasmid, 0.4 μM of both forward and reverse primers listed in Table S1 (Bio Basic Inc), 0.2 mM of dNTPs, and 0.05 U/μl of Pfu DNA polymerase (Thermo Scientific) or 0.02 U/μl Q5 High-fidelity DNA polymerase (New England Biolabs Inc) in 1× buffer were mixed. The PCR condition was hot started at 95 °C for 5 min followed by 16 cycles of denaturation at 95 °C for 45 s, annealing at 60 °C for 1 min, and extension at 72 °C for 15 min. The reaction was kept at 72 °C for 18 min in the final cycle. PCR products were treated with DpnI (New England Biolabs Inc) to remove the hadAWT-pET-11a template. Mutated plasmids were transformed into E. coli XL1-Blue and grown on LB agar plate containing ampicillin (50 μg/ml) at 37 °C. Colonies were selected, grown, and purified to obtain plasmid DNA. Selected plasmids were sequenced to confirm accuracy of the mutation (Macrogen Inc or U2Bio). HadA variants were purified using the same approach used for HadAWT (7).

**Activity assay of HadA**

Activity of HadA variants was analyzed using a coupled enzymatic system consisting of HadA and reduced flavin regenerating enzymes including glucose-6-phosphate dehydrogenase (Glc-6-PD) and HadX flavin reductase (6). The reaction consisting of 4NP (50 μM), glucose-6-phosphate (Glc-6-P, 1 mM), Glc-6-PD (0.5 unit/ml), NAD+ (5 μM), HadX (1 μM), HadA variants (10 μM) was carried out in 20 mM Hepes pH 7.5. The absorption change at a wavelength of 400 nm was monitored to measure the loss of 4NP using a UV-visible spectrophotometer Cary 60 (Agilent Technology).

**Transient kinetics experiments of HadA variants**

Rapid kinetics experiments of HadA variants were performed using similar methods previously used for HadAWT (7). A binary complex of HadAvariants:FADH+ (75 μM:25 μM) was prepared by reduction of HadAvariants:FAD with a slight excess amount of sodium dithionite inside the anaerobic glove box before placing them in a tightly capped anaerobic syringe to avoid auto-oxidation by oxygen. A solution of anaerobic HadAvariants:FADH+ was mixed with an aerobic solution 4CP (0.5 mM with 0.13 mM oxygen) on a single-mixing stopped-flow spectrophotometer model SF-61SX (TgK Scientific, Bradford-on-Avon) in which the flow parts were made anaerobic by flushing with an oxygen scrubbing solution overnight (25). Reagent concentrations are expressed as the final concentrations after mixing. Absorption changes at wavelengths 380 nm (A380) and 450 nm (A450) were monitored to indicate the formation of C4a-hydroperoxy-FAD and oxidized FAD, respectively. To monitor dead-end complex formation, the aerobic enzyme solution was mixed with a substrate and incubated for certain periods before mixing with FADH+ to initiate the formation of C4a-hydroperoxy-FAD similar to the approach used in the HadAWT reaction previously reported (7). 4NP was not used in this experiment because the compound has high absorbance at 400 nm, interfering with observation of the kinetics of flavin oxidation.

**HQ production analysis**

Samples from single turnover reactions were prepared and collected for analysis using the same methods described (7). The BQ product was reduced to HQ, which is more stable by adding ascorbic acid (0.5 mM). Production of HQ product was analyzed on an HPLC 1100 series (Agilent Technologies) equipped with a UV-visible diode array detector. A Nova-Pak (Waters) C18 reverse phase column with a 4-μm particle size and a 3.9 × 150-mm column size was used as the stationary phase. A gradient of H2O/methanol containing 0.1% formic acid (10%-70% methanol) was used as a mobile phase. HQ was detected at a wavelength of 289 nm at a retention time of 3.9 min.

**Molecular docking**

Binding of 4NP to the active sites of HadA variants was investigated by AutoDock Vina (62). Structures of HadA variants were prepared by mutagenesis tools using the PyMol software with appropriate rotamers. The configurations of nonpolar moieties were assigned using AutoDock Tools 1.5.7 with the cubic set at 8 to 10 Å and the coordinate of C1 of 4NP in the HadAWT structure as a center with an energy range of four and exhaustiveness of eight. The affinity energy was collected from the lowest-energy conformation aligned in the correct orientation.

**Thermofluor assays**

A solution of HadAWT or HadAPhe41 variants (10 μM) was mixed with 5× SYPRO orange dye (Sigma) in 20 mM Hepes pH 7.5 in the absence or presence of 4NP (2 mM). In the presence of 4NP, enzymes and substrate were preincubated for 1 h prior to mixing with SYPRO orange dye in 20 mM Hepes pH 7.5. A real-time PCR instrument model Rotor Gene Q (Qiagen) was used to monitor fluorescence changes due to protein unfolding to determine the melting temperature (Tm) of HadA (16). The program’s temperature was gradually increased from 35 °C to 95 °C with 0.5 °C/min increment.
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Statistical analysis
Data from at least three replications were used for calculating means ± standard deviations (SD). Statistical significance analysis was performed using the independent t test from Statistical Package for the Social Sciences Program (IBM).

Data availability
All data are included in the article and supporting information. Data of X-ray structures are available at Protein Data Bank under PDB codes indicated.

Supporting information—This article contains supporting information.

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Abbreviations—The abbreviations used are: BQ, benzoquinone; C2, p-hydroxyphenylacetate hydroxylase from Acinetobacter baumannii; CnTcpA, 2,4,6-trichlorophenol monoxygenase from C. nantongensis sp. JT-3; CphC-1, a two-component flavin monoxygenase from Arthrobacter chlorophenolicus A6; 4CP, 4-chlorophenol; DcmB1, two-component flavin monoxygenase from Rhodococcus sp. JT-3; EcHpaB, p-hydroxyphenylacetate hydroxylase from Escherichia coli; FAD, oxidized form of flavin adenine dinucleotide; FADH1, reduced form of flavin adenine dinucleotide; Glc-6-P, glucose-6-phosphate; Glc-6-PD, glucose-6-phosphate dehydrogenase; HadA, a flavin monoxygenase component from Ralstonia pickettii; HadX, a flavin reductase component from Ralstonia pickettii; HPs, halo-generate phenols; HpaA, FAD-dependent two-component monoxygenase from Cupriavidus sp. CNP-8; HQ, hydroquinone; NP, nitrophenol; 4NP, 4-nitrophenol monoxygenase from Rhodococcus opacus SAO101; NpdA2, p-nitrophenol monoxygenase from Arthrobacter sp. Strain JS443; NpsA1, p-nitrophenol monoxygenase from Rhodococcus sp. strain PN1; PcpB, pentachlorophenol hydroxylase from Sphingobium chlorophenolicum; PheA1, phenol hydroxylase from Bacillus thermoglucosidasius A7; RetCPA, 2,4,6-trichlorophenol 4-monoxygenase from Ralstonia eutropho JMP134; TtD, 2,4,5-trichlorophenol 4-monoxygenase from Burkholderia cepacia AC1100; ThHpaB, p-hydroxyphenylacetate hydroxylase from Thermus thermophilus HB8.

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