Molecular insights into substrate recognition and catalysis by tryptophan 2,3-dioxygenase

Citation for published version:

Digital Object Identifier (DOI):
10.1073/pnas.0610007104

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Proceedings of the National Academy of Sciences

Publisher Rights Statement:
Copyright 2007 by the National Academy of Sciences of the United States of America; all rights reserved.

General rights
Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.
Molecular insights into substrate recognition and catalysis by tryptophan 2,3-dioxygenase

Farhad Forouhar*, J. L. Ross Anderson†, Christopher G. Mowat‡, Sergey M. Vorobiév*, Arif Hussain*, Mariam Abashidze*, Chiara Bruckmann*, Sarah J. Thackray1, Jayaraman Seetharaman*, Todd Tucker*, Rong Xiao‡, Li-Chung Ma‡, Li Zhao‡, Thomas B. Acton‡, Gaetano T. Montelione‡, Stephen K. Chapman†, and Liang Tong‡

*Department of Biological Sciences, Northeast Structural Genomics Consortium, Columbia University, New York, NY 10027; †School of Chemistry, University of Edinburgh, West Mains Road, Edinburgh EH9 3JJ, United Kingdom; and ‡Center for Advanced Biotechnology and Medicine and Northeast Structural Genomics Consortium, Rutgers University, Piscataway, NJ 08854

Communicated by Wayne A. Hendrickson, Columbia University, New York, NY, November 13, 2006 (received for review September 25, 2006)

Tryptophan 2,3-dioxygenase (TDO) and indoleamine 2,3-dioxygenase (IDO) constitute an important, yet relatively poorly understood, family of heme-containing enzymes. Here, we report extensive structural and biochemical studies of the Xanthomonas campestris TDO and a related protein SO4414 from Shewanella oneidensis, including the structure at 1.6-Å resolution of the catalytically active, ferrous form of TDO in a binary complex with the substrate L-Trp. The carboxylate and ammonium moieties of tryptophan are recognized by electrostatic and hydrogen-bonding interactions with the enzyme and a propionate group of the heme, thus defining the L-stereospecificity. A second, possibly allosteric, l-Trp-binding site is present at the tetramer interface. The sixth coordination site of the heme iron is vacant, providing a dioxygen-binding site that also would involve interactions with the ammonium moiety of L-Trp and the amide nitrogen of a glycine residue. The indole ring is positioned correctly for oxygenation at the C2 and C3 atoms. The active site is fully formed only in the binary complex, and biochemical experiments confirm this induced-fit behavior of the enzyme. The active site is completely devoid of water during catalysis, which is supported by our electrochemical studies showing significant stabilization of the enzyme upon substrate binding.

Tryptophan 2,3-dioxygenase (TDO) and indoleamine 2,3-dioxygenase (IDO) catalyze the oxidative cleavage of the L-tryptophan (L-Trp) pyrrole ring, the first and rate-limiting step in L-Trp catabolism through the kynurenine pathway (1–3). In addition, IDO has been implicated in a diverse range of physiological and pathological conditions, including suppression of T cell proliferation, maternal tolerance to allogenic fetus, and immune escape of cancers (2, 9–12).

Despite catalyzing identical biochemical reactions (Fig. 1a), the sequence similarity between TDO and IDO is extremely low. An alignment of their sequences is only possible based on their structures, which suggests a sequence identity of 10% between them (Fig. 1b). In comparison, Xanthomonas campestris TDO shares 34% sequence identity with human TDO (Fig. 1b), demonstrating the remarkable evolutionary conservation of this enzyme. TDO is a homotetrameric enzyme and is highly specific for L-Trp and related derivatives such as 6-fluoro-Trp as the enzyme. TDO is a homotetrameric enzyme and is highly specific demonstrating the remarkable evolutionary conservation of this enzyme. Our structures reveal for the first time the structural basis for the stereospecificity of this important enzyme. Our structural information is confirmed by biochemical studies and offers significant molecular insight into tryptophan dioxygenation by TDO and IDO.

Results

Structure Determination. Crystals of the reduced (Fe(II)) TDO from X. campestris in a binary complex with the substrate L-Trp or 6-fluoro-Trp were obtained after extensive efforts and by using anerobic conditions, because the oxidized (Fe(III)) enzyme has much lower affinity for L-Trp (see below). The structures at up to 1.6-Å resolution of these binary complexes [Table 1; and see supporting information (SI) Table 3] as well as that of the free enzyme were determined by molecular replacement based on the structure of the apo enzyme, in the absence of heme, which we had determined by the selenomethionyl single-wavelength anomalous diffraction method (PDB entry 1YW0) (15).

The structure of the SO4414 protein from S. oneidensis (16) was determined at 2.4-Å resolution by molecular replacement based on our structure of the apo enzyme (PDB entry 1ZEE).

Structure of TDO. The structure of X. campestris TDO monomer contains 12 helices (named αA through αL) and no β-strands (Figs. 1b and 2a). TDO is an intimately associated tetramer (Fig. 2b), and ~4,500 Å2 of the surface area of each monomer is buried in the tetramer. Helices αB and αC are located in the extensive, mostly hydrophobic interface between two of the monomers. The N-terminal segments (residues 21–40, including helix αA) of the two monomers are swapped in this dimer (Fig.


The authors declare no conflict of interest.

Abbreviations: D-Trp, D-tryptophan; IDO, indoleamine 2,3-dioxygenase; L-Trp, L-tryptophan; TDO, tryptophan 2,3-dioxygenase.

Data deposition: The atomic coordinates have been deposited in the Protein Data Bank, www.pdb.org (PDB ID codes 1YW0, 2NOW, 2WNB, 2WNA, 1ZEE, and 2WNB).

§To whom correspondence should be addressed. E-mail: ltong@columbia.edu.

This article contains supporting information online at www.pnas.org/cgi/content/full/0610007104DC1.

© 2006 by The National Academy of Sciences of the USA

www.pnas.org/cgi/doi/10.1073/pnas.0610007104

PNAS | January 9, 2007 | vol. 104 | no. 2 | 473–478
which is important for the catalysis by TDO because several residues in this segment are part of the binding site for the Trp residue in the active site (see below).

**Binding Mode of the l-Trp Substrate to TDO.** Our structure of the binary complex defines the molecular mechanism for the recognition of the l-Trp substrate by TDO. Clear electron density was observed for heme and l-Trp in the active site based on the crystallographic analysis at 1.6-Å resolution (Fig. 3). The l-Trp substrate is located in a pocket over the distal face of the heme, having interactions with residues in helices /H9251B and /H9251D, and the /H9251D–/H9251E and /H9251J–/H9251K loops (Figs. 1b and 3b). The carboxylate group of Trp is recognized by bidentate ion-pair interactions with the side chain of Arg 117 (in helix /H9251D). The carboxylate group is also hydrogen-bonded to the side chain hydroxyl of Tyr 113 (helix /H9251D). The ammonium ion of l-Trp is recognized by the 7-propionate side chain of the heme group (Fig. 3b), and it is also hydrogen-bonded to the side chain hydroxyl of Thr 254 (in helix /H9251D). The indole ring is located ~3.5 Å above and perpendicular to the heme and is held in place by van der Waals interactions with the side chains of Phe 51 (in helix /H9251B) and several other hydrophobic residues, including Tyr 24, Tyr 27, and Leu 28 from the N-terminal segment of another monomer of the tetramer (Fig. 3b). In addition, the N1 nitrogen of the indole ring is hydrogen-bonded to the side chain of His 55 (helix /H9251B) (Fig. 3b). A water molecule is present in the active site of this binary complex (Fig. 3a), hydrogen-bonded to the ammonium ion of l-Trp and the main-chain amide of residue Gly 125 (Fig. 3b). The water is 3.5 Å from the ferrous atom in the heme, too far for ligating interactions. The iron atom is still 0.3 Å out of the plane of the heme, on the side of the proximal His 240 ligand (SI Fig. 5).

The crystal was exposed to a solution saturated with nitric oxide (NO) before being flash-frozen, but we did not observe the binding of this dioxygen analog in the structure. This is confirmed by our structure of the binary complex with 6-fluoro-Trp, which was not exposed to NO but contained the same density for the water molecule (SI Fig. 6). The structure of the 6-fluoro-Trp binary complex is essentially identical to that of the l-Trp binary complex (SI Fig. 6). NO probably dissociated from the heme during the cryofreezing manipulations in the anaerobic box.

**Induced-Fit Behavior of TDO.** Our structural information suggests that TDO is an induced-fit enzyme. Although the active site
this finding, the 7-propionate of heme is ion-paired with an Arg residue from the protein surface. This interaction may also be stabilized by crystal packing interactions, as the Arg side chain is hydrogen-bonded to a water molecule, located 3.5 Å from the heme iron. Both nitrogen atoms are instead hydrogen-bonded in the binary complex, thus stabilizing the active site region. Although this region is exposed to the solvent in the free enzyme, it is completely shielded from the solvent in the binary complex, and only the carboxylate group of the 6-propionate of heme is visible on the surface.

Additional evidence for the induced-fit behavior is observed in the active site of the second TDO monomer in the asymmetric unit. The binding mode of the L-Trp substrate is very different in this monomer (Fig. 3d; and see SI Fig. 7). The Trp side chain is not positioned as deeply into the pocket, and the hydrogen bond between the ring nitrogen and the side chain of His 55 is not positioned as deeply into the pocket, and the hydrogen bond between the ring nitrogen and the side chain of His 55 is lost (distance of 3.8 Å). Both nitrogen atoms are instead hydrogen-bonded to a water molecule, located 3.5 Å from the heme iron but at a position distinct from that of the water in the active site of the other monomer (Fig. 3d). This conformation may also be stabilized by crystal packing interactions, as the 7-propionate of heme is ion-paired with an Arg residue from another TDO tetramer in the crystal. The main chain atoms of the Trp substrate appear to be disordered, because no clear electron density was observed for them (Fig. 3e). Consistent with this finding, the αJ–αK loop is disordered in this molecule, similar to that in the free enzyme (Fig. 3d). This complex may represent an initial stage in the formation of the Michaelis complex of TDO. Proper positioning of the L-Trp substrate for catalysis would lead to the recognition of its main chain atoms and the ordering of the αJ–αK loop.

Implications for Substrate Binding by IDO. Our structure of the binary complex of TDO also has significant implications for substrate recognition by the IDOs. The rms distance for 206 equivalent Cα atoms, suggesting that SO4414 may also be a dioxygenase. Moreover, the structure contains an extra domain that is formed by residues at the N terminus, similar to the small domain in human IDO (SI Fig. 8). In contrast to human IDO, SO4414 is a tetramer but with a different organization compared with that of TDO (SI Fig. 10). Our biochemical efforts so far have not been able to demonstrate IDO (or TDO) activity for this protein, suggesting that SO4414 may prefer a different substrate for oxygenation.

A Model for the Michaelis Complex. To help provide further insight into the catalysis by these enzymes, we built a model for the Michaelis complex of TDO, by placing one oxygen atom (O1) of the dioxygen substrate directly over the heme iron, at a distance of 2.1 Å (Fig. 4a). The distal oxygen atom (O2) was placed such that the O1–O2 bond is parallel to the C2–C3 bond of indole ring, giving a Fe–O–O2 angle of 135°. This conformation places the O2 atom within 0.5 Å of the water molecule observed in our structure (Fig. 4a), suggesting that this water should be ejected from the active site upon dioxygen binding. The active site is therefore completely devoid of solvent molecules in this Michaelis complex.

The modeled dioxygen-binding mode reveals the activation mechanism of this substrate for the reaction. It has been established that TDO has an ordered catalytic cycle in which the protein first binds L-Trp to the ferrous form, and then binding of dioxygen is facilitated, and nucleophilic attack from the substrate C3 is initiated (3, 13, 19). In the model, the distal oxygen atom interacts with the L-Trp amine moiety and the backbone amide nitrogen of Gly 125 (Fig. 4a). The Lewis acidity of the heme iron further assists oxygen activation, making the TDO mechanism distinct from that of oxygenases. It also provides insights into the reaction mechanism of the IDOs, which have been shown to form a complex with tryptophan, which may be the reason why it cannot completely distinguish among the indoleamine substrates.

### Table 1. Summary of crystallographic information

<table>
<thead>
<tr>
<th>Protein</th>
<th>TDO (holoenzyme)</th>
<th>TDO (holoenzyme)</th>
<th>TDO (holoenzyme)</th>
<th>TDO (holoenzyme)</th>
<th>TDO (holoenzyme)</th>
<th>SO4414 (holoenzyme)</th>
<th>SO4414 (holoenzyme)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ligand</td>
<td>L-Trp</td>
<td>6-fluoro-Trp</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Maximum resolution, Å</td>
<td>1.6</td>
<td>1.8</td>
<td>2.7</td>
<td>2.7</td>
<td>2.4</td>
<td>2.3</td>
<td>2.3</td>
</tr>
<tr>
<td>R&lt;sub&gt;merge&lt;/sub&gt;, %</td>
<td>6.9 (52.7)</td>
<td>8.6 (60.9)</td>
<td>13.0 (32.3)</td>
<td>16.6 (74.0)</td>
<td>10.4 (63.0)</td>
<td>6.6 (30.6)</td>
<td>6.6 (30.6)</td>
</tr>
<tr>
<td>Beam line</td>
<td>ESRF BM14</td>
<td>ESRF BM14</td>
<td>NSLS X4A</td>
<td>APS 21BM</td>
<td>NSLS X4A</td>
<td>NSLS X4A</td>
<td>NSLS X4A</td>
</tr>
<tr>
<td>Completeness, %</td>
<td>87 (65)</td>
<td>87 (61)</td>
<td>72 (54)</td>
<td>82 (68)</td>
<td>83 (60)</td>
<td>87 (71)</td>
<td>87 (71)</td>
</tr>
<tr>
<td>R factor, %†</td>
<td>17.1 (18.9)</td>
<td>16.6 (17.9)</td>
<td>25.7 (25.7)</td>
<td>25.0 (32.1)</td>
<td>21.7 (23.5)</td>
<td>23.4 (26.7)</td>
<td>23.4 (26.7)</td>
</tr>
<tr>
<td>Free R factor, %</td>
<td>18.9 (22.0)</td>
<td>18.4 (21.1)</td>
<td>26.3 (26.2)</td>
<td>29.4 (35.1)</td>
<td>22.5 (24.1)</td>
<td>27.5 (31.5)</td>
<td>27.5 (31.5)</td>
</tr>
<tr>
<td>rms deviation in bond lengths, Å</td>
<td>0.005</td>
<td>0.005</td>
<td>0.008</td>
<td>0.011</td>
<td>0.007</td>
<td>0.006</td>
<td>0.006</td>
</tr>
<tr>
<td>rms deviation in bond angles, °</td>
<td>1.0</td>
<td>1.0</td>
<td>1.2</td>
<td>1.6</td>
<td>1.1</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Most-favored region, %</td>
<td>92</td>
<td>91</td>
<td>89</td>
<td>86</td>
<td>92</td>
<td>91</td>
<td>91</td>
</tr>
</tbody>
</table>
of the hydrogen-bonding donors, coupled with the electron-withdrawing nature of the heme, would increase the electrophilicity of the bound dioxygen and render it more susceptible to nucelophilic attack by the substrate C3 atom. The increased hydrophobicity of the active site upon the exclusion of water would also aid the stabilization of an oxyferrous species. Studies with heme oxygenase suggest that the hydrogen-bonding interactions to the dioxygen substrate may also help to prevent its heterolysis (20), and the exclusion of water probably removes a hydrogen-bond competitor to the dioxygen. After the initial attack by the C3 atom, the reaction may proceed via an allylic Criegee rearrangement or a dioxetane intermediate (SI Fig. 11). In the model, the O1–O2 atoms are in a trans configuration relative to the C2–C3 atoms of L-Trp (Fig. 4c), which may favor the Criegee rearrangement pathway (SI Fig. 11) (19). The Criegee pathway is also favored based on chemical, thermodynamic, and quantum mechanical considerations (3).

Our model for the Michaelis complex shows that the O1 atom is 2.6 Å from the N1 atom of L-Trp and therefore can act as the general base to extract the proton from the N1 atom (SI Fig. 11) (19). The N1 atom is hydrogen-bonded to His 55 in TDO. However, our biochemical studies show that the $k_{cat}$ of TDO is relatively insensitive to pH over the range examined (pH 6 to pH 8) (Fig. 4b), and the H55A mutant had only a 10-fold decrease in the $k_{cat}$ (Table 2), suggesting that this residue is not essential for catalysis, consistent with its replacement with a Ser residue in IDO. On the other hand, the $K_m$ shows a marked increase at lower pH (Fig. 4b), probably because of the protonation of this residue.

**An Allosteric Binding Site in the Tetramer Interface.** We also observed the binding of four L-Trp residues to an allosteric site in the interface of the tetramer (Fig. 2b), with well defined electron density (SI Fig. 12). The L-Trp residue appears to be recognized specifically by the enzyme in this pocket (SI Fig. 12). There have been reports of allosteric activation by the substrate L-Trp (21, 22), and our observations offer a possibility for this effector site. Unfortunately, our kinetic studies so far have not shown any allosteric effects with *X. campestris* TDO. This site is not occupied in the 6-fluoro-Trp complex, possibly because of the lower concentration of this compound in the crystallization solution.

**Biochemical Studies Confirm the Structural Observations.** *X. campestris* TDO has robust catalytic activity toward L-Trp and 6-fluoro-Trp but is inactive toward D-Trp, tryptamine or indolepropionic acid (Table 2), confirming its designation as a TDO. In fact, D-Trp is a weak, competitive inhibitor of the enzyme at high concentrations (Table 2). Our binding data show that D-Trp has much lower affinity for the enzyme than L-Trp (Table 2), consistent with our structural information and explaining why D-Trp cannot be oxygenated by TDO.

The biochemical studies also provide direct evidence for the induced-fit behavior of TDO. There is a large increase in the affinity of the enzyme for L-Trp when the heme iron is reduced (K4 [ferric Fe(III) heme] = 3.8 mM, whereas K4 [ferrous Fe(II) heme] = 4.1 μM) (Table 2). The electrochemistry data show a large, positive shift in reduction potential (+136 mV) in the presence of 15 mM L-Trp (Fig. 4c). In fact, the shift in reduction potential almost perfectly correlates with the increase in affinity for L-Trp on reduction, both giving an estimated ΔΔG of 15 kJ/mol. These data show that there is a significant stabilization of the ferrous form when substrate is bound. This stabilization could also play a physiological role to keep the protein reduced, and therefore active, when L-Trp is present.

Our structural studies have defined the binding mode of the substrate L-Trp to TDO, revealing the structural basis for the stereospecificity of this important enzyme. The induced-fit behavior of TDO, confirmed by our biophysical studies, appears crucial for the exclusion of water from the active site and for stabilizing the enzyme in the presence of the substrates. Finally, structural comparisons among these enzymes reveal the striking evolutionary conservation of the heme-dependent dioxygenases.

**Materials and Methods**

The experimental protocols are summarized here. More detailed information can be found in SI Materials and Methods.

**Protein Expression and Purification.** Full-length *X. campestris* TDO (NESG ID XcR13) and *S. oneidensis* SO4414 (NESG ID SoRS2) were cloned into a pET-21d (Novagen, San Diego, CA) derivative, with a C-terminal hexahistidine tag, and overexpressed at 17°C in *Escherichia coli* BL21(DE3) pMGK cells. Hemin (7 μM

---

**Fig. 3.** Molecular basis for substrate recognition by TDO. (a) Final 2F_o – F_e electron density at 1.6-Å resolution for heme, L-Trp, and a water in the active site. Contoured at 1σ. (b) Stereo drawing showing the active site of *X. campestris* TDO in the binary complex with L-Trp. The segment in cyan is from another monomer of the tetramer. Hydrogen-bonding interactions are indicated with dashed lines in magenta. (c) Overlay of the structures of the free enzyme (in orchid) and the binary complex (yellow and cyan) in the active-site region. Regions of conformational differences are indicated with the red arrows. (d) Overlay of the active-site region of the second monomer (in green) and that of the first monomer (in yellow). Only the side-chain atoms of Trp are shown in the second monomer (in magenta). (e) Final 2F_o – F_e electron density at 1.6-Å resolution for heme, L-Trp, and a water in the active site of the second TDO molecule in the crystal. Contoured at 1σ. Two conformations for the main chain atoms are shown, but neither fit the density well. For the stereo version of c and d, please see SI Fig. 7. Produced with Molscript (35) and rendered with Raster3D (36).
final concentration) was included in the media for preparation of the holoenzyme samples (23). The protein was purified by using nickel-affinity and gel-filtration chromatography.

Point mutations were created with the QuickChange II site-directed mutagenesis kit (Stratagene, La Jolla, CA). The mutations and their associated ORFs were verified by DNA sequence analysis.

**Protein Crystallization.** To obtain the structure of TDO in the ferrous state, the protein was reduced by the addition of sodium dithionite, and all steps were performed in an anaerobic glove box (Belle Technology, Dorset, U.K.), with the O$_2$ concentration maintained $<2$ ppm. Excess sodium dithionite was removed by gel filtration (Sephadex G25 column) before crystallization. Crystals of TDO were grown by the sitting-drop vapor diffusion method with a well solution comprising 100 mM Mes (pH 6.3), 10–12% (wt/vol) PEG 4000, 60 mM MnCl$_2$, 10 mM sodium dithionite, and 2 mM L-Trp. Before mounting in nylon loops and flash-freezing in liquid nitrogen, crystals were immersed in a cryoprotectant solution composed of mother liquor (with L-Trp concentration increased to 50 mM) supplemented with 23% (vol/vol) glycerol and bubbled with nitric oxide for 15 min before use.

**Data Collection and Processing.** X-ray diffraction data were collected at the X4A beam line of National Synchrotron Light Source (Brookhaven National Laboratory, Upton, NY), the 21BM beam line at Advanced Photon Source (Argonne National Laboratory, Argonne, IL), and the BM14 beam line at the DLS (Diamond Light Source, Oxford, U.K.).

![Fig. 4. Molecular insights into the catalysis by TDO. (a) Model of the Michaelis complex. The water molecule is shown as a small sphere in green. Produced with Molscript (35) and rendered with Raster3D (36). (b) The pH dependence of the $k_{cat}$ and $K_m$ values of X. campestris TDO. (c) The presence of L-Trp causes a large, positive shift in the reduction potential of TDO. SHE, standard hydrogen electrode.](image)

**Table 2. Summary of kinetic data on X. campestris TDO**

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Substrate</th>
<th>$k_{cat}$ ($s^{-1}$)</th>
<th>$K_m$, mM</th>
<th>$K_d$, Ferric heme, mM</th>
<th>$K_d$, Ferrous heme, mM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>L-Trp</td>
<td>19.5 ± 1.2</td>
<td>114 ± 1</td>
<td>3.84 ± 0.14</td>
<td>4.12 ± 0.24</td>
</tr>
<tr>
<td></td>
<td>L-Trp</td>
<td>0</td>
<td>16.5 mM ± 3.3*</td>
<td>&gt;50†</td>
<td>NC</td>
</tr>
<tr>
<td></td>
<td>L-Trp</td>
<td>37.3 ± 0.6</td>
<td>186 ± 12</td>
<td>2.45 ± 0.42</td>
<td>&lt;1†</td>
</tr>
<tr>
<td></td>
<td>L-Trp</td>
<td>2.40 ± 0.10</td>
<td>100 ± 6</td>
<td>1.51 ± 0.08</td>
<td>&lt;1†</td>
</tr>
<tr>
<td></td>
<td>Indolepropionic acid</td>
<td>0</td>
<td>0</td>
<td>&gt;10†</td>
<td>126 ± 11</td>
</tr>
<tr>
<td></td>
<td>O$_2$</td>
<td>35.4 ± 0.9</td>
<td>119 ± 2</td>
<td>0</td>
<td>NA</td>
</tr>
<tr>
<td></td>
<td>L-Trp</td>
<td>2.86 ± 0.10</td>
<td>133 ± 7</td>
<td>ND</td>
<td>3.7 ± 1.3</td>
</tr>
</tbody>
</table>

NA, $K_d$ for O$_2$ cannot be measured in the absence of substrate because of oxidation or in the presence of substrate because of turnover; NC, No spectral change detected; ND, not done.

*Inhibitory constant, $K_i$.

*Although a spectral change was evident, substrate solubility prevented accurate measurement of $K_d$. Values were estimated based on the maximum substrate concentration attainable.

*Binding was too tight to be measured. Values quoted represent the minimum $K_d$ that can be measured under standard assay conditions.

The peak positions of the oxyferrous complex (O$_2$-TDO) are at 420 nm, 548 nm, and 578 nm.
European Synchrotron Radiation Facility (Grenoble, France). The diffraction images were processed and scaled with the HKL package (24). The data-processing statistics are summarized in Table 1, and more complete information can be found in SI Table 3.

**Structure Determination and Refinement.** The structures of the apo enzymes of TDO and SO4414 were determined by the selenomethionyl single-wavelength anomalous diffraction method (15). The selenium sites were located with SnB (25), and the reflection phases were calculated with Solve/Resolve (26). The structures of the holoenzymes and the ternary complex were determined by the molecular-replacement method, with the programs COMO (27) and AMoRe (28). The atomic models were built with the program XtalView (29) and TURBO-FRODO (30), and the structure refinement was carried out with CNS (31).

**Electronic Spectroscopy, Steady-State Assays, and Dissociation Constant Measurements.** Electronic absorption spectra were recorded by using a Cary-50 Probe UV-Visible spectrophotometer at 25°C. Assays for the steady-state turnover (at pH 7.5) of L-Trp and derivatives were performed as described (32, 33), except that substrate concentrations of 0–15 mM were used. The kinetic data were fitted to the Michaelis–Menten equation. The pH dependence of the steady-state kinetics was determined in the same manner, by using phosphate (pH 6.0–8.0) and Tris (pH 8.0–9.0) buffers. The electronic absorption spectra of the steady state were recorded by using a stopped-flow spectrophotometer (SX.17MV; Applied-Photophysics, Surrey, U.K.) in conjunction with a diode array detector, housed in an anaerobic glove box ([O₂], <5 ppm; Belle Technology).

**OTTLE Electrochemistry.** Anodic potentiometric titrations were carried out as described (34) at 25°C by using a modified quartz EPR OTTLE cell. Titrations were performed in both the absence and presence of L-Trp (15 mM), and the heme reduction potentials were determined by fitting the data to the Nernst equation for a single-electron process by using Origin software (MicroCal, Northampton, MA). Reduction potentials are quoted versus the standard hydrogen electrode.

We thank Randy Abramowitz and John Schwanof at the National Synchrotron Light Source (Brookhaven National Laboratory, Upton, NY) and Hassan Belrhali at the European Synchrotron Radiation Facility (Grenoble, France) for setting up the beam lines and G. DeTitta of Hauptman Woodward Research Institute (Buffalo, NY) for crystalization screening. This research was supported by Grants P50 GM62413 and U54 GM074958 from the Protein Structure Initiative of the National Institutes of Health.