The Second Enzyme in Pyrrolnitrin Biosynthetic Pathway Is Related to the Heme-Dependent Dioxygenase Superfamily†


Centre for Biomolecular Sciences, EastChem, The University, St Andrews, Scotland, KY16 9ST, U.K., Department of Chemistry, EastChem, The University of Edinburgh, Scotland EH9 3JJ, Biochemie, Technische Universität Dresden, D-01062 Dresden, Germany, and Department of Chemistry, Biochemistry and Molecular Biology, University of Georgia, Athens, Georgia 30602

Received June 20, 2007; Revised Manuscript Received August 6, 2007

ABSTRACT: Pyrrolnitrin is a commonly used and clinically effective treatment for fungal infections and provides the structural basis for the more widely used fludioxonil. The pyrrolnitrin biosynthetic pathway consists of four chemical steps, the second of which is the rearrangement of 7-chloro-tryptophan by the enzyme PrnB, a reaction that is so far unprecedented in biochemistry. When expressed in Pseudomonas fluorescens, PrnB is red in color due to the fact that it contains 1 mol of heme b per mole of protein. The crystal structure unexpectedly establishes PrnB as a member of the heme-dependent dioxygenase superfamily with significant structural but not sequence homology to the two-domain indoleamine 2,3-dioxygenase enzyme (IDO). The heme-binding domain is also structurally similar to that of tryptophan dioxygenase (TDO). Here we report the binary complex structures of PrnB with D- and L-tryptophan and D- and L-7-chloro-tryptophan. The structures identify a common hydrophobic pocket for the indole ring but exhibit unusual heme ligation and substrate binding when compared with that observed in the TDO crystal structures. Our solution studies support the heme ligation observed in the crystal structures. Purification of the hexahistidine-tagged PrnB yields homogeneous protein that only displays in vitro activity with 7-chloro-L-tryptophan after reactivation with crude extract from the host strain, suggesting that an as yet unknown cofactor is required for activity. Mutation of the proximal heme ligand results, not surprisingly, in inactive enzyme. Redox titrations show that PrnB displays a significantly different reduction potential to that of IDO or TDO, indicating possible differences in the PrnB catalytic cycle. This is confirmed by the absence of tryptophan dioxygenase activity in PrnB, although a stable oxyferrous adduct (which is the first intermediate in the TDO/IDO catalytic cycle) can be generated. We propose that PrnB shares a key catalytic step with TDO and IDO, generation of a tryptophan hydroperoxide intermediate, although this species suffers a different fate in PrnB, leading to the eventual formation of the product, monodechloroaminopyrrolnitrin.

Biosynthetic pathways in bacteria remain a major source of interest primarily due to the value of the final natural product but often for the novel chemistry contained within them. The pyrrolnitrin biosynthetic pathway (Scheme 1) is an example of such a pathway in which the final product, a potent antifungal compound used to treat skin infections, is produced by a succession of elegant transformations whose mechanisms are not immediately obvious. This pathway is common to many organisms such as Enterobacter agglomerans (1), Myxococcus fulvus (2), Cystobacter ferrugineus (2), and Serratia spp (3) and was first isolated and studied in Pseudomonas species. Through both genetic engineering of plants to include the pyrrolnitrin biosynthetic gene cluster and direct addition of Pseudomonas spp to plants, it has been demonstrated that fungal molds can be effectively controlled (4–6). Pyrrolnitrin is believed to exert its antifungal activity through inhibition of the respiratory electron transport system (7).

Previous knockout studies in Pseudomonas fluorescens have identified the intermediate products from each of the biosynthetic proteins in the pyrrolnitrin pathway, and some

Scheme 1: Pyrrolnitrin Pathway

1 This work was supported by the Biotechnology and Biological Sciences Research Council (U.K.) through a grant to J.H.N.
2 To whom correspondence should be addressed: Centre for Biomolecular Sciences, The University, St Andrews, KY16 9ST U.K.
3 University of St Andrews.
4These authors made an equal contribution.
5University of Edinburgh.
6University of Dresden.
7University of Georgia.
8University of Georgia.
9University of Georgia.
10.1021/bi7012189 CCC: $37.00 © 2007 American Chemical Society Published on Web 10/09/2007
of the transformations have been well characterized (8). The first step is the regioselective incorporation of chlorine into D- and L-tryptophan by PrnA. The crystal structure and subsequent solution studies indicate that HOCl is formed at the FAD cofactor and passed down a channel toward the substrate (9) where an N-chlorolysine species is formed prior to the chlorination of tryptophan (10). A similar mechanism is likely to operate in the third step, as PrnC, which is known to have sequence homology to PrnA, chlorinates mono-dechloroamino-tryptophan. The final step, in which the amine of the substrate is oxidized to a nitro group, is carried out by the non-heme iron and iron–sulfur cluster-containing PrnD and has also been recently examined (11).

Although the product and substrate of PrnB are known, the enzyme has to date received little attention. Interestingly, both D- and L-tryptophan can be incorporated into the final product of the complete pathway, pyrrolnitrin, and it has been shown that PrnA can utilize either of these enantiomers as its substrate. This indicates that PrnB utilizes both enantiomers of 7-chlorotryptophan alternatively an as yet unidentified racemase interconverts the D- and L-isomers, with PrnB only acting on one. Only 7-chloro-L-tryptophan is isolated from the Pseudomonas fluorescens mutant strain lacking PrnB, suggesting that this is the preferred natural enantiomer (8).

Interest in tryptophan biochemistry is dominated by the study of its degradation in the kynurenine pathway by the enzymes tryptophan 2,3-dioxygenase (EC 1.13.11.11, TDO) and indoleamine 2,3-dioxygenase (EC 1.13.11.52, IDO). Both are b-type heme dependent dioxygenases and catalyze the conversion of L-tryptophan to L-formylkynurenine by incorporating two atoms of molecular oxygen into the indole ring of the substrate (Scheme 2). TDO is a tetrameric enzyme containing four molecules of heme b per tetramer (12, 13), and IDO is a monomeric, two-domain protein containing one molecule of heme b per protein molecule (14). The heme group is found enclosed and in the presence of substrate free from water (12). Given the similarity in the heme-binding domains of the two proteins and in their active sites, it seems likely that both use a similar mechanism, although IDO has broader substrate specificity and has been shown to perform a more versatile role in vivo. Two mechanisms have been proposed: the Criegee mechanism involves the formation of a tryptophan hydroperoxide intermediate that undergoes a Criegee-type rearrangement (12, 13); the other mechanism involves the formation of a four-membered dioxetane ring (14). In comparison to the mechanism operating in PrnB, there must be significant differences as the products are dissimilar, although both result in the opening of the indole ring of tryptophan. Most significantly for PrnB, there is no incorporation of oxygen into the product. There is no overall significant sequence similarity of PrnB to either IDO or TDO, hinting at a novel mechanism for the formation of mono-dechloroamino-tryptophan. Recently, another member of the tryptophan dioxygenase family was isolated from Shewanella oneidensis which shares the same fold as IDO, contains heme b but is not active with tryptophan, and contains a highly divergent sequence from other members of the family (PDB 2NW).

In this paper we report the initial structural and spectroscopic characterization of the 361 residue, 43 kDa PrnB protein from Pseudomonas fluorescens BL915. We have determined the X-ray crystal structure using multiple wavelength anomalous scattering in the presence of its putative substrates (7-chloro-D-tryptophan and 7-chloro-L-tryptophan) and its substrate analogues D- and L-tryptophan. These structural data show striking similarities between PrnB and both IDO (14) and TDO (12, 13). Our spectroscopic studies reveal significant differences between PrnB and IDO/TDO, suggesting that PrnB does indeed utilize a novel mechanism.

MATERIALS AND METHODS

Purification of PrnB. The cloning, purification, and crystallization of PrnB have already been reported in detail (15). We were unable to obtain folded soluble protein from any Escherichia coli system but instead relied upon the Pseudomonas fluorescens expression system. Native protein had a tendency to aggregate, and a triple mutant (C21S, C60S, and C185S) was also used. The protein had a noncleavable His tag at the N-terminus. Briefly, P. fluorescens cells with a plasmid containing the his tagged prnB gene were grown. After centrifugation, the cells were washed with 0.1 M potassium phosphate buffer (pH 7.0), and then one part of cells was resuspended in two parts of potassium phosphate buffer (pH 7.0) and the cells were disrupted by sonication for 3 min (Branson Sonifier, with 5 s pulse, 10 s break, 0 °C, 65% amplitude). After removal of the cell debris by centrifugation, the crude extract was either stored at −20 °C or immediately used for purification of the enzyme. In the latter case 3 mL aliquots of the crude extract were
PrnB Is a Heme-Dependent Enzyme

Table 1: Crystallography Statistics for the Structures

<table>
<thead>
<tr>
<th></th>
<th>native</th>
<th>L-tryptophan</th>
<th>D-tryptophan</th>
<th>7-Cl-L-tryptophan</th>
<th>7-Cl-D-tryptophan</th>
</tr>
</thead>
<tbody>
<tr>
<td>wavelength (Å)</td>
<td>1.488</td>
<td>1.739</td>
<td>0.976</td>
<td>1.54</td>
<td>1.283</td>
</tr>
<tr>
<td>space group</td>
<td>C222</td>
<td>C2</td>
<td>C2</td>
<td>C2</td>
<td>C2</td>
</tr>
<tr>
<td>cell dimensions</td>
<td>a = 67.7 Å</td>
<td>a = 68.6 Å</td>
<td>a = 68.2 Å</td>
<td>a = 68.4 Å</td>
<td>a = 68.5 Å</td>
</tr>
<tr>
<td></td>
<td>b = 80.1 Å</td>
<td>b = 79.6 Å</td>
<td>b = 79.5 Å</td>
<td>b = 78.9 Å</td>
<td>b = 79.5 Å</td>
</tr>
<tr>
<td></td>
<td>c = 129.5 Å</td>
<td>c = 92.9 Å</td>
<td>c = 92.3 Å</td>
<td>c = 93.4 Å</td>
<td>c = 93.1 Å</td>
</tr>
<tr>
<td></td>
<td>α = β = γ = 90°</td>
<td>α = γ = 90°</td>
<td>α = γ = 90°</td>
<td>α = γ = 90°</td>
<td>α = γ = 90°</td>
</tr>
<tr>
<td>resolution (Å)</td>
<td>38–1.75</td>
<td>51–2.0</td>
<td>37–1.75</td>
<td>39–2.4</td>
<td>51–2.0</td>
</tr>
<tr>
<td>(highest shell (Å))</td>
<td>1.84–1.75</td>
<td>2.11–2.0</td>
<td>1.84–1.75</td>
<td>2.5–2.4</td>
<td>2.1–2.0</td>
</tr>
<tr>
<td>Rmerge</td>
<td>0.082 (0.193)</td>
<td>0.066 (0.298)</td>
<td>0.073 (0.666)</td>
<td>0.049 (0.102)</td>
<td>0.065 (0.437)</td>
</tr>
<tr>
<td>completeness (%)</td>
<td>16.0 (4.2)</td>
<td>21.5 (6.3)</td>
<td>12.7 (2.0)</td>
<td>14.3 (6)</td>
<td>13.0 (2.0)</td>
</tr>
<tr>
<td>redundancy</td>
<td>98.1 (94.3)</td>
<td>97.1 (94.8)</td>
<td>97.5 (96.3)</td>
<td>98.3 (94.9)</td>
<td>95.8 (83.0)</td>
</tr>
<tr>
<td></td>
<td>4.4 (4.0)</td>
<td>7.1 (7.3)</td>
<td>3.7 (3.8)</td>
<td>2.1 (2.0)</td>
<td>2.6 (1.8)</td>
</tr>
<tr>
<td>Data Collection</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>no. of reflections</td>
<td>33422</td>
<td>30217</td>
<td>48852</td>
<td>17632</td>
<td>30563</td>
</tr>
<tr>
<td>Rwork/Rfree</td>
<td>17.9/20.7</td>
<td>17.7/20.6</td>
<td>18.3/20.5</td>
<td>18.2/21.3</td>
<td>17.1/19.5</td>
</tr>
<tr>
<td>(highest shell)</td>
<td>(22.7/26.4)</td>
<td>(21.8/27.7)</td>
<td>(32.3/38.7)</td>
<td>(19.4/25.8)</td>
<td>(23.1/31.6)</td>
</tr>
<tr>
<td>no. of atoms</td>
<td>2748</td>
<td>2690</td>
<td>2714</td>
<td>2714</td>
<td>2703</td>
</tr>
<tr>
<td>ligand/ion</td>
<td>52</td>
<td>67</td>
<td>67</td>
<td>57</td>
<td>67</td>
</tr>
<tr>
<td>water</td>
<td>370</td>
<td>120</td>
<td>220</td>
<td>156</td>
<td>196</td>
</tr>
<tr>
<td>rmsd</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>bond lengths (Å)</td>
<td>0.01</td>
<td>0.01</td>
<td>0.009</td>
<td>0.009</td>
<td>0.01</td>
</tr>
<tr>
<td>bond angles (deg)</td>
<td>1.21</td>
<td>1.14</td>
<td>1.10</td>
<td>1.12</td>
<td>1.14</td>
</tr>
<tr>
<td>PDB code</td>
<td>2V71</td>
<td>2V7J</td>
<td>2V7K</td>
<td>2V7L</td>
<td>2V7M</td>
</tr>
</tbody>
</table>

loaded onto a Ni-chelating Sepharose fast flow column (1 x 7 cm, 0.2 mL/min) equilibrated with 50 mM Tris phosphate buffer (pH 8.0) containing 150 mM NaCl and 20 mM imidazole. After removing all nonbond protein by washing the column with 40 mL of the same buffer, proteins were eluted with a linear gradient over a period of 500 min (0.2 mL/min) from 150 mM NaCl, 20 mM imidazole in 50 mM Tris phosphate buffer (pH 8.0) to 300 mM NaCl, 500 mM imidazole in 50 mM Tris phosphate buffer (pH 8.0) with the collection of 2 mL fractions. Fractions were analyzed by SDS-PAGE, and those containing PrnB were pooled and dialyzed against 10 mM Tris phosphate buffer (pH 8.0) overnight. The purified protein (mutant and native) was a deep red color characteristic of heme-containing enzymes.

Structural Biology. The protein was characterized by mass spectrometry including acetone extraction of lyophilized sample. Mass spectrometric analysis of the acetone extraction is consistent with the presence of heme b (peak at 615.95 nm; calculated 615.6 Å). Protein crystals were only obtained from the triple mutant protein, and the details of the crystallization have been reported (15). The best results were obtained with 1.5 µL of protein solution (16 mg/mL, 50 mM Tris pH 7.5) plus 1.5 µL of precipitant (0.22 M magnesium sulfate, 16% w/v PEG 3350) irrespective of whether protein was incubated with tryptophan (as a saturated solution) or not. We first recorded a data set to 1.75 Å resolution of protein crystallized in the presence of L-tryptophan. We attempted to solve the structure with molecular replacement using apo TDO from Xanthomonas campestris (PDB 1ZEE) but were unable to find a solution. This was unsurprising given the less than 20% identity, and we resorted to experimental phasing. We collected data on a second crystal from the same crystallization conditions at three wavelengths to a resolution of 2.0 Å chosen to optimize the anomalous (1.739 Å) and dispersive differences (1.741 Å and 1.722 Å) resulting from the iron. This crystal, although grown in the same conditions, had a different unit cell and spacegroup. SOLVE located the single iron, and density modification allowed us to quickly trace a monomer. We then extended the phases to 1.75 Å with data collected with a wavelength of 1.49 Å. This structure was used to solve the structure of the protein crystallized with D-tryptophan. Protein crystallized with a racemic mixture of D- and L-7-chlorotryptophan under the same conditions gave a 2.0 Å 7-Cl-D-tryptophan complex structure. Protein crystallized with pure 7-Cl-L-tryptophan, synthesized as previously reported (16), gave a 2.4 Å 7-Cl-L-tryptophan complex structure. The structures were refined using REFMAC5 (17). Ligands were added only when density was clear. The ligand dictionaries were taken from the REFMAC defaults. TLS parameters were employed in refinement, and we investigated the use of anisotropic thermal factor refinement, but for the two higher resolution data sets, the improvement in Rfree was less than 0.5%, and so isotropic thermal factors were used. Full crystallographic details are given in Table 1.

Activity Assay. Cultures of Pseudomonas fluorescens BL915 Δorf1-4 which had been engineered to lack the pyrroline-5-thiol synthetase pathway but harbored a plasmid containing prnB or mutants of prnB were grown overnight in HNB medium. The cultures were diluted 1 to 20 in fresh medium, and cells were grown for a further 24 h at 30 °C and 200 rpm on orbital shaker incubators. To some of the cultures sterile filtered 7-Cl-L-tryptophan (r Racemic) was added to a final concentration of 1 mM. Cells were grown for an additional 2 days. 20 mL of the cultures with or without 7-Cl-tryptophan were extracted twice with one volume of ethyl acetate. The extracted mixtures were dried in vacuo and redissolved in 130 µL of methanol. The extracts were analyzed by thin layer chromatography on silica plates using toluene/dioxane (9:1) as the eluent and by isocratic HPLC with methanol/H2O (65:35) as the eluent, using a Varian OmniSpher or Lichrosphere...
Magnetic field as field. MCD intensities are linear with respect to the magnetic room-temperature bore was used to generate a 6 T magnetic Instruments SM-1 superconducting solenoid with a 25 mm for the UV spectra were recorded using Jasco J810 and J730 models analyzed by HPLC as described above.

**FIGURE 1:** A stereoribbon representation of the structure of PrnB. The core helical domain is colored cyan, and the N-terminal capping domain is colored blue. The cyan colored helices are found in TDO, and the N-terminal domain is not. The disordered loop is marked by an asterisk (*) at each end. The side chains of three cysteine residues (20, 60, and 175) which were mutated to serine are shown in sticks. The heme group and the proximal H313 ligand are shown in sticks with carbon yellow, nitrogen blue, and oxygen red.

100, RP-18, 5 µm, 250 × 4.6 mm column. The flow rate was 1 mL/min, and compounds were detected at 220 and 303 nm. In *vitro* activity was analyzed by incubating 225 µL of PrnB-containing protein solution with 125 µL of 1 mM 7-chlorotryptophan (racemic or pure enantiomers) and 150 µL of 0.1 M potassium phosphate buffer (pH 7.0) at 30 °C for 16 h. The assay mixtures were extracted twice with 500 µL of ethyl acetate, the combined organic extracts were dried in *vacuo* and redissolved in 130 µL of methanol, and 20 µL of the resulting solution was used for HPLC analysis as described above. For reactivation of PrnB after dialysis or purification by immobilized nickel-chelating affinity chromatography, 225 µL of PrnB-containing protein solution was incubated with 125 µL of 1 mM 7-chlorotryptophan, 150 µL of 0.1 M potassium phosphate buffer (pH 7.0), and 150 µL of cell-free crude extract from the host strain *P. fluorescens* BL915 AorF1-4 at 30 °C for 16 h and analyzed by HPLC as described above.

**Magnetic Circular Dichroism (MCD) Spectroscopy.** MCD spectra were recorded using Jasco J810 and J730 models for the UV–vis and near-IR regions, respectively. An Oxford Instruments SM-1 superconducting solenoid with a 25 mm room-temperature bore was used to generate a 6 T magnetic field. MCD intensities are linear with respect to the magnetic field at room temperature and are plotted normalized to magnetic field as Δε/ε (M⁻¹ cm⁻¹ T⁻¹). Samples were prepared in deuterated Hepes buffer (50 mM) at pH 7.5.

**Electron Paramagnetic Resonance (EPR) Spectroscopy.** X-band EPR spectra (10 K) were recorded on a Bruker ER-300D series electromagnet and microwave source interfaced to a Bruker EMX control unit and fitted with an ESR-9 liquid helium flow cryostat from Oxford Instruments and a dual mode microwave cavity from Bruker (ER-4116DM). Quantification of $S = 1/2$ species was carried out by comparison with the spin standard Cu²⁺(EDTA) using the method of Aasa and Vannagard (18). Magnetic field magnitude is detected by a Hall probe within the cavity of the instrument. Additionally, the g values of the Cu²⁺EDTA standard (g = 2.32 and 2.075) were used to ensure the correct field values for the detected frequency (observed values identical to those given above). Frequency is monitored within the EMX unit and also with an additional frequency counter attached to the microwave source.

**OTTLE Potentiometry.** Anaerobic potentiometric titrations were carried out as described previously (19), at 25 °C using a modified quartz EPR OTTLE cell. A range of mediators were added (flavin mononucleotide (−200 mV vs SHE), 2-hydroxy-1,4-naphthoquinone (−145 mV), 5-hydroxy-1,4-naphthoquinone (−3 mV), phenazine ethosulfate (+55 mV), phenazine methosulfate (+82 mV), 1,2-naphthoquinone (+135 mV)) to ensure efficient reduction and reoxidation. Titrations were performed both in the absence and in the presence of L-tryptophan and D-tryptophan (15 mM), and the heme reduction potentials were determined by fitting the data to the Nernst equation for a single electron process using the Microcal Origin software. Reduction potentials are quoted versus the standard hydrogen electrode (SHE).

**Electronic Spectroscopy and Dissociation Constant Measurements.** Electronic absorption spectra were recorded using a Cary 50-Probe UV–visible spectrophotometer at 25 °C. Dissociation constants ($K_a$) were determined by established spectrophotometric procedures. The spectrum of the oxyferrous species was recorded using an Applied-Photophysics stopped-flow spectrophotometer (SX.17MV) in conjunction with a diode array detector, housed in an anaerobic glove box (Belle Technology, [O₂] < 5 ppm). A solution of prereduced PrnB (40 µM) was mixed with oxygenated buffer at 15 °C, and spectra were recorded using the diode array detector.

**RESULTS**

**Structure of PrnB.** The asymmetric unit of all crystals contains a monomer of the protein. We investigated crystal contacts for any dimerization using the Protein Interaction server (PISA) (20) at the European Bioinformatics Institute (EBI), and this indicates that the protein is a monomer, consistent with gel filtration results. The final refined structure consists of residues 5 to 361 (true C-terminus); one loop of 9 residues which connects two α helices (323 to 331) is disordered. The N-terminal tag (40 residues) is not seen in any structure. The structure contains 17 α helices connected together by a series of loops; there are two short
β-strands in the structure (Figure 1). The experimental electron density map showed clear density for a heme group in each of the five structures. The molecule can be split into two domains, a small N-terminal domain of 120 amino acids (α1–7) and the core helical domain of 240 amino acids. The helical core domain is related to myoglobin. The structure is very similar to that seen for IDO (PDB 2D0T) (root-mean-square deviation of 2.4 Å for 268 Cα atoms), and the core helical domain is topologically identical to TDO (PDB 2NW8) (root-mean-square deviation of 3.2 Å for 168 Cα atoms). There are differences in detail regarding the loops and positioning of α helices that explain why molecular replacement failed to work. The three cysteine residues that were mutated are remote from the heme. However, despite being grown in the presence of L-tryptophan, only a water molecule is found at the heme in the 1.75 Å structure. The heme is bound in a solvent exposed cleft with 7-propionate group accessible and distal face exposed. The 6-propionate group makes a salt contact with R310. The heme is also anchored to the protein by a number of van der Waal interactions with residues on helices α8, α9, α10, and α16 which pack around the distal face and hydrophobic edge of the porphyrin ring. The precise degree of solvent accessibility of the distal face of the heme must be treated with caution as the disordered loop is adjacent to heme cleft and could fold over sealing the cleft. The proximal ligand to the heme iron is His 313, which is located on α16. The distal face of the porphyrin has a large hydrophobic cavity above it. Calculations using CASTP (27) estimate the surface area of the cavity as 480 Å² with a volume of 580 Å³ in the substrate free structure. The distal ligand to the iron in this structure is a water molecule which is hydrogen bonded to a second water molecule which is in turn bound to the main chain of protein. Strangely, a crystal grown under essentially the same conditions and used in MAD phasing does unambiguously have L-tryptophan bound at the active heme. We have no explanation for this, nor have we been able to obtain good quality crystals of the substrate free form again. Crystals grown in the presence of D-tryptophan clearly show D-tryptophan bound as do crystals grown with 7-Cl-L-tryptophan. The presence or absence of a ligand at the heme does not significantly alter the PrnB structure. The changes are displacement of water molecules and movement of a few flexible side chains.

In all four complex structures, the indole is bound in the hydrophobic pocket above the distal face and the main chain is located at the iron. However, we see two modes of binding of the indole ring and two modes of binding of the main chain: in total, three distinct complexes (the D-tryptophan and 7-Cl-D-tryptophan complexes are essentially identical, the only difference being the additional chlorine atom on the ligand). In the 7-Cl-L-tryptophan and 7-Cl-D-tryptophan (and D-tryptophan) complexes the indole rings essentially superimpose. The indole ring is sandwiched between the side chains of L140 and Phe 201 on one side and the main chain between G223 to V225 on the other. The five membered ring of the indole system is capped by Y209, and the chlorine atom is found in a pocket bounded by V117, P222 and is 3.8 Å from a water molecule. In the L-tryptophan complex, the indole ring is rotated relative to other three complexes, in such a way that the 6 membered ring overlap but the five membered rings are on opposite sides (Figure 2). In the L-tryptophan complex, the indole nitrogen points toward the side chain of S143 (> 4.0 Å).

Despite the difference in the indole ring location between the L-tryptophan and 7-Cl-L-tryptophan complexes, the main chain of the tryptophans does superimpose. The amino nitrogen atom of the tryptophan is ligated to the iron, and the carboxylic acid points into an oxyanion hole G223–V225: one oxygen atom makes a hydrogen bond to the backbone amide of V225, and the other makes hydrogen bonds to water molecules. In the 7-Cl-D-tryptophan complex, one of the oxygen atoms of the carboxyl group is bound to the iron, with the other oxygen making the same hydrogen bond to V225 as seen for the L-enantiomers, and the amino nitrogen makes hydrogen bonds to water molecules.

Activity of PrnB. The native enzyme and the triple cysteine his-tagged mutant used for the biophysical characterization behave identically. The cell-free extracts containing PrnB showed a spot on silica thin layer plates with an Rf-value of 0.24 that reacted with van Urk’s reagent (reacts with indoles and their derivatives) to give a pink color immediately. This turns olive with time and is thought to correspond to 2-(2′aminophenyl)pyrrole. This is the product one would expect from the reaction of PrnB with tryptophan. Van Urk’s reagent is made by dissolving 1 g of p-dimethylaminobenzaldehyde in 50 mL of 36% HCl followed by the addition of 50 mL of ethanol; the solution is stored at 4 °C (8). HPLC analysis of cell free extracts shows that a peak at 6 min is eluted that exhibits a UV-spectrum with a maximum at 303 nm. Mass spectrometry of the peak reveals a mass of 159 (M + H+), consistent with 2-(2′aminophenyl)pyrrole. Extracts of culture medium of cells fed with 7-Cl-tryptophan show a spot on thin layer plates with an Rf-value of 0.5 that stains as described above when sprayed with van Urk’s reagent. HPLC analysis of cell free extracts of the 7-Cl-tryptophan fed cells shows a peak at 10.5 min. The sample corresponding to this peak, when analyzed by mass spectrometry, showed a parent ion peak of 193/195 amu with a ratio of 3:1, expected for monodechloroaminopyrrolnitrin (PrnB product, Scheme 1). The same peak was obtained when crude cell-free extract was incubated in the presence of racemic 7-chlorotryptophan (Figure 3) as well as the pure D- and L-enantiomers (Figure 3). The yield of monodechloroaminopyrrolnitrin with 7-chloro-L-tryptophan is about 1.5-fold higher than with the D-enantiomer. When the crude extract was dialyzed overnight or purified by Ni-chelating chromatography, no PrnB activity was detected. However, when crude extract of the host strain which did not contain PrnB (and has no PrnB activity) was added to the dialyzed PrnB, full activity was restored (Figure 3). In vitro analysis of the activity of PrnB with L- and D-tryptophan could not be performed since both cell-free extracts (the PrnB-containing one and the one from the host strain) already contain L-tryptophan which cannot be removed without inactivation of the enzyme. The H313A mutant had no activity. Attempts to purify the mutant resulted in a colorless (presumably heme free) aggregated protein. We assayed for TDO activity by looking for the formation of N-formylkynurenine at 321 nm. The oxyferrous form of the enzyme was generated and titrated against L-tryptophan in a multimixing experiment in the stopped flow. There was no evidence of reaction either by monitoring for N-formylkynurenine formation or spectral changes at the heme (versus control experiments in the absence of L-Trp).
**Figure 2:** (a) The complexes of PrnB with substrates 7-Cl-L-tryptophan, 7-Cl-D-tryptophan, and L-tryptophan (D-tryptophan overlaps with 7-Cl-D-tryptophan and is not shown). The protein is colored as in Figure 1, non carbon atoms are colored as in Figure 1, chlorine atoms are purple, for 7-Cl-L-tryptophan carbon atoms are colored yellow, carbons in 7-Cl-D-tryptophan are colored orange, and carbons in L-tryptophan are colored salmon. Although the chloro-indole rings of 7-Cl-L-tryptophan and 7-Cl-D-tryptophan overlap, the main chain atoms do not. L-Tryptophan binds in a different manner to the other complexes, although the part of the indole rings overlaps. The proximal ligand to the heme (H313) is shown. In 7-Cl-L-tryptophan and L-tryptophan, the amino nitrogen atom ligates to the iron, whereas in 7-Cl-D-tryptophan one of the carboxyl oxygen atoms ligates to the iron. (b) The active site of PrnB. The residues that surround the 7-Cl-L-tryptophan ligand are shown. The oxygen hole at A224 is labeled, and the hydrogen bonds are shown. Atoms are colored as in Figure 2a. (c) The subtle differences in the active sites of PrnB (colored as in Figure 2a) and TDO (carbon atoms green, other atoms colored as in Figure 2a). The superposition is based on the heme groups of each protein. The different orientation of the loop adjacent to heme, which contains the oxygen hole, can be seen. (d) Superposition of the L-tryptophan complexes of PrnB and TDO (PDB code 2NW8); atoms colored as in Figure 2c. The binding sites are mutually exclusive; F201 in PrnB prevents L-tryptophan binding as seen in TDO, while H55 of TDO prevents any of the binding modes for tryptophan seen in PrnB. (e) The substrate complexes of PrnB and TDO show that despite quite different orientations the CD2 atom (marked with an asterisk (*)) where the peroxide is predicted to be inserted is similarly positioned relative to the heme.

**Spectroscopy of PrnB.** UV-visible spectroscopy shows the characteristic signature of heme-binding proteins with the Fe$^{3+}$ peak at 405 nm and the Fe$^{2+}$ peak at 417 nm. The ligand titration experiments give a $K_D$ of 60 $\mu$M for L-tryptophan, 39 $\mu$M for D-tryptophan, and 4.9 $\mu$M for 7-Cl-L-tryptophan binding to Fe$^{3+}$ PrnB. We do not have sufficient quantities of pure 7-Cl-D-tryptophan for spectroscopic study. Due to poor solubility at higher concentrations, it was impossible to directly measure $K_D$ values for tryptophan with PrnB in the reduced, Fe$^{2+}$ form. However, using our electrochemical data we can estimate these values to be as follows: 11 mM $\pm$ 2 mM for L-Trp, 40 mM $\pm$ 7 mM for D-Trp, and 140 mM $\pm$ 20 mM for 7-Cl-L-tryptophan. The $K_D$ values of L-tryptophan and 7-Cl-L-tryptophan with the Fe$^{3+}$ form are very sensitive to pH and titrate with a $pK_a$ of around 6.5. The midpoint reduction potential of PrnB in the absence of ligand is $-109$ mV. The presence of ligand significantly alters the redox potential of PrnB, $-287$ mV (L-tryptophan), $-243$ mV (D-tryptophan), and $-195$ mV (7-Cl-L-tryptophan), respectively (Figure 4). In the Fe$^{2+}$ state, in the absence of substrate, PrnB readily forms an oxyferrous complex. The decay of this complex by autooxidation has a rate constant of 0.0015 s$^{-1}$ $\pm$ 0.0001 at 15 °C. The rate of decay is independent of the presence of tryptophan (as pure D or L) (Figure 4).

The MCD/EPR data (Figure 5) show that the substrate free enzyme is a mixture of low-spin high-spin six-coordinate species, consistent with the water molecule in the crystal structure. When L-tryptophan is added to PrnB, the spectrum shows formation of a low-spin species. MCD/EPR indicates that this is six-coordinate with a nitrogenous ligand in the vacant coordination site. The addition of D-tryptophan shows oxygen atom coordination similar to but distinct from native (Figure 5).

**DISCUSSION**

In crude extract, PrnB reacts with both 7-Cl-D- and 7-Cl-L-tryptophan almost equally as well. They could be equivalent substrates, or it could be that the crude extracts contain a racemase which rapidly interconverts the D-enantiomer into the L-enantiomer. This would be inconsistent with the finding that from a mutant lacking PrnB only 7-Cl-L-tryptophan (no 7-Cl-D-tryptophan) could be isolated (22). The fact that PrnB is inactivated by dialysis and purification by Ni-chelating chromatography, but can be reactivated by the addition of cell-free crude extract from the host strain *P. fluorescens* BL915 Δorf1-4, indicates that a yet unknown cofactor or chemical/enzymatic transformation of substrate is required for PrnB activity. Simply adding reducing agents such as ascorbic acid has not led to activity recovery.

The topology of the core helical domain of PrnB is very similar to that of both IDO (and TDO), and the heme binding site is located in approximately the same position in all structures. Both PrnB and IDO are monomers and contain the small N-terminal domain which helps to form the heme binding pocket. This degree of similarity would point to a functional relationship, at least for the key catalytic step, between IDO/TDO and PrnB. IDO and PrnB structures are clearly more similar, and the “disordered loop” at the heme site of PrnB is also disordered in IDO. TDO lacks the small N-terminal domain, but rather another monomer contributes an $\alpha$ helix to the heme binding pocket, requiring at least a dimeric unit for activity; in fact, TDO is a tetramer. The “disordered loop” in PrnB is ordered in TDO and partly occludes the heme binding site. In our PrnB structures, the sixth site of the heme is ligated by the substrate, either by the amino nitrogen (L-tryptophan and 7-Cl-L-tryptophan) or the carboxyl group (D-tryptophan and 7-Cl-D-tryptophan).
This is clearly inconsistent with the PrnB functioning as tryptophan 2,3-dioxygenase as this requires that O$_2$ binds at Fe$^{2+}$, not tryptophan. Of course a static crystal structure may be misleading. However, the pH dependence of L-tryptophan and 7-Cl-L-tryptophan binding are consistent with their complex crystal structures (Figure S5, Supporting Information). Further MCD and EPR studies of PrnB in solution indicate nitrogen ligation to the Fe$^{3+}$ when L-tryptophan is added. The same experiments show oxygen ligation to the Fe$^{3+}$ in D-tryptophan, thus the crystal structures are supported by solution measurements. In three complexes (D-tryptophan, 7-Cl-D-tryptophan, and 7-Cl-L-tryptophan) the indole rings superimpose. We predict that this is an indole binding site in a catalytically competent complex. In the absence of a dramatic conformational change, there is no other pocket which could accommodate a chlorinated indole ring. The binding mode observed in L-tryptophan would require a significant conformation change to avoid clashes for any chlorinated indole, and we believe that the indole ring

**FIGURE 3:** (a) HPLC-analysis of the in vitro activity assay of PrnB incubated with racemic 7-chlorotryptophan. The bottom line shows monodechloroaminopyrrolnitrin (MCAP) the product formed by PrnB from 7-chlorotryptophan as a standard. The other lines show from bottom to top increasing volume of crude extracts. The numbers on the right-hand side give the amount of MCAP corresponding to the peak. (b) HPLC analysis of the in vitro activity assay of PrnB incubated with racemic 7-chlorotryptophan and the pure D- and L-enantiomers, respectively. The lines from bottom to top show MCAP standard; incubation with racemic 7-chlorotryptophan (0.5 mM and 0.25 mM final concentration, respectively); incubation with 0.5 mM 7-chloro-D-tryptophan; incubation with 0.5 mM 7-Cl-L-tryptophan. (c) HPLC analysis of in vitro activity assays of PrnB. The lines from bottom to top show MCAP as the standard; crude extract of the host strain P. fluorescens BL915 Δorf1-4; dialyzed PrnB-containing extract; cell-free crude extract containing PrnB; restoration of PrnB activity by incubation of dialyzed crude extract together with nondialyzed crude extract of the host strain.

**FIGURE 4:** (a) OTTLE potentiometry of PrnB in the absence of L-tryptophan (circles), + 20 mM L-tryptophan (triangles pointing up), + 20 mM D-tryptophan (triangles pointing down), and 10 mM 7-chloro-L-tryptophan. In each case the filled and empty shapes correspond to the reductive and oxidative traces, respectively. Each data set is fit using the Nernst equation, and the fit is colored according to the corresponding label. (b) Decay of oxyferrous PrnB. Upper panel: Spectral changes associated with the decay of the oxyferrous PrnB species. Lower panel: Exponential decrease in absorbance at 543 nm used to determine the rate of oxyferrous PrnB decay. Red line indicates single exponential
location in the L-tryptophan complex is a noncatalytic binding mode for L-tryptophan only.

We have been unable to detect any IDO or TDO activity from isolated PrnB using assays previously employed for IDO (12). Redox potential measurements also point toward PrnB being incapable of functioning as tryptophan 2,3-dioxygenase. The redox potential of the holo PrnB is $-109$ mV, which is different than that of IDO, whose potential in the substrate free form was reported as $-267$ mV (23). Most strikingly in IDO the presence of L-tryptophan increases the redox potential to $+14$ mV, stabilizing the Fe$^{2+}$ state, whereas for PrnB, L-tryptophan significantly destabilizes the Fe$^{2+}$ state ($-243$ mV) and D-tryptophan even more so ($-280$ mV). This is consistent with both D-tryptophan and L-tryptophan providing a sixth ligand and stabilizing Fe$^{3+}$ in solution. Thus in complete contrast to IDO, tryptophan deactivates PrnB toward 2,3-dioxygenase chemistry. Comparing the TDO and PrnB L-tryptophan complexes shows that the indole rings are oriented in a similar manner with respect to the heme but shifted by over 2 Å (Figure 2). In TDO, the amino nitrogen and carboxyl group of L-tryptophan are rotated and do not bind to the Fe$^{3+}$. L-Tryptophan cannot adopt the conformation seen for TDO in PrnB, as F201 of PrnB would clash with the C$_\beta$ of L-tryptophan. Similarly H55 of IDO would prevent the orientation of any of the tryptophan rings seen in PrnB (Figure 2). Structurally as well as biochemically, PrnB does not appear to possess IDO chemistry having an altered substrate binding pocket.

The structural similarity of PrnB to IDO (and TDO) does however argue that the enzymes are related mechanistically at some level. Comparing the L-tryptophan complexes suggests that PrnB shares a similar indole binding pocket. However, the other three complexes of PrnB suggest that this pocket has been adapted to favor binding of the larger chlorinated substrates. Other changes in the active site have resulted in a significant change in redox potential at the heme.

**Figure 5:** (a) Perpendicular mode X-band EPR spectra of native ferric PrnB in the absence of tryptophan (black line), ferric PrnB $+ 20$ mM L-tryptophan (red line), and ferric PrnB $+ 20$ mM D-tryptophan (blue line). Conditions: microwave frequency, 9.67 GHz; microwave power, 2 mW; modulation amplitude, 10 G; temperature, 10 K; scan speeds and time constants are the same for all spectra. Spectra have been adjusted for differences in the enzyme concentration and receiver gain where required. Gains were typically of the order of $2 \times 10^5$ to $8 \times 10^5$. (b) Room-temperature MCD at 5 T in the UV/visible region of native ferric PrnB in the absence of tryptophan (black line), ferric PrnB $+ 20$ mM L-tryptophan (red line), and ferric PrnB $+ 20$ mM D-tryptophan (blue line). All spectra were recorded at pH 7.0. (c) Room-temperature MCD at 5 T in the near-IR region of native ferric PrnB in the absence of tryptophan (black line), ferric PrnB $+ 20$ mM L-tryptophan (red line), and ferric PrnB $+ 20$ mM D-tryptophan (blue line). All spectra were recorded at pH 7.0 with 50 µM enzyme.
and in the orientation of the tryptophan at the iron center. The defining chemistry of IDO is the formation of the oxyferrous species and its insertion into the indole ring to form the iron tryptophan hydroperoxide. IDO has an oxyanion hole (centered on N125) close to the iron center which could serve to stabilize the oxyferrous complex (12). PrnB has a similar oxyanion hole, although its precise orientation with respect to the heme is subtly different (Figure 2) and may explain the stability of this complex in PrnB. We propose that the mechanism for PrnB has retained the defining chemical step (Scheme 2) of peroxide insertion at the CD2 atom of tryptophan. In support of this we note that despite the differences in orientation between 7-Cl-L-tryptophan in PrnB and L-tryptophan in IDO (12) (Figure 2), the respective CD2 atoms are within 1.5 Å of each other when the heme groups are superimposed. Our Fe3+ complexes therefore cannot be completely accurate representations of a productive complex, as the Fe2+ must be free to bind O2 and not be ligated to substrate. We propose that the amino (and carboxyl group) bound to the heme must rotate around the Cα Cβ bond of tryptophan to free the sixth site to iron while the indole ring is fixed in position. We note that, in the 7-Cl-L-tryptophan and 7-Cl-O-tryptophan complexes, the main chain atoms are rotated relative to each other while the chlorinated indole rings are in identical positions. The difference between PrnB and IDO/TDO we believe is in the fate of the iron tryptophan hydroperoxide intermediate. PrnB does have a different architecture on the distal face to TDO/IDO. This is mirrored in its different redox potential and in differences in the indole binding site. These differences, we propose, prevent it from stabilizing the Criegee intermediate and stop tryptophan dioxygenase chemistry. In looking at the chemistry catalyzed by PrnB, the most likely intermediate would possess three fused rings (Scheme 3), and in fact chemical studies on peroxy tryptophan have shown that such a tricyclic species does form (24). The three ring structure has precedence in biomimetic organic chemistry (pyrrolobenzoxazine core unit of paeciloxazine (25) Scheme 3) and most significantly in other chlorinated and non-chlorinated natural products (pyrroindomycin A/B, Scheme 3) (26). We suggest that PrnB catalyzes the formation of a three fused ring intermediate (as opposed to the Criegee intermediate) by utilizing the electron withdrawing properties of the tryptophan hydroperoxide intermediate to activate the indole ring for nucleophilic attack (Scheme 4). The three ring system can rearrange (Scheme 4) by the reversible elimination of the peroxide. An irreversible elimination step would ultimately decide the fate of the three ring intermediate. This mechanism would give rise to three possibilities: the pyrroindolynitrin like product (loss of CO2); the pyrroindomycin (26) like product (loss of H2); an arylpyrrole compound (Scheme 4). Although arylpyrrrole compounds are well described in chemistry (27), they are not well-known natural products. However, the arylpyrrrole derivative 2-carboxy-4-(2′-amino-3′-chlorophenyl)pyrrole as well as its bromo analogue (Scheme 3) have been isolated from the pyrrolnitrin producer P. aureofaciens (28, 29). We take as support for our mechanistic hypothesis the fact that all three possible chlorinated products that could result from the iron hydroperoxide three ring intermediate are found in nature.

![Scheme 3](image)

*This has analogues in the natural product pyrroindomycin (26) and biomimetic chemistry (pyrrolobenzoxazine core) (25); an arylpyrrol structure has been found in pyrrolnitrin producing bacteria (28, 29).*

![Scheme 4](image)

*The tricyclic ring structure is boxed. Routes to the pyrroindomycin core and arylpyrrole core structures are shown (24).*
We have considered an alternative mechanism (Supporting Information) involving proton abstraction from the Cα atom and N atom of 7-Cl-L-tryptophan. However, there are no appropriate residues located close to these atoms, and such a mechanism would involve an Fe23+ species.

Although enzyme activity in isolated extracts is robust, a more accurate delineation of the mechanism requires that we obtain activity under defined conditions. We can only speculate on the lack of activity of the purified enzyme. Our protocols could inactivate the protein in some way related to the redox conditions (loss of cofactor or reducing enzyme).

Alternatively, there is an additional specific essential enzyme which complexes with PrnB or modifies substrate. The conservation of the four gene cluster and its successful insertion into plants would seem to make it very unlikely that there is a specific PrnB partner enzyme or specific 7-Cl-tryptophan modifying enzyme. Despite this, PrnB is a significant addition to the IDO/TDO superfamily. The rearrangement is surprising. Our work suggests that there may be more members of this superfamily to be discovered which catalyze related but different indole chemical modifications. These putative enzymes could well have important synthetic applications. Our investigations into this fascinating class of enzyme will continue.

ACKNOWLEDGMENT

We thank Ms. M. Thorke, Biochemie, TU Dresden, for synthesis of racemic 7-chlorotryptophan and Ms. A. Ernyei, Biochemie, TU Dresden, for enzymatic preparation of enantiomerically pure D- and L-7-Cl-tryptophan. We thank Dr. R. Goss, University of East Anglia, for pure L-7-Cl-tryptophan. We are also indebted to Myles Cheesman, University of East Anglia, for MCD and EPR assistance and Sarah Thackray, University of Edinburgh, for help with OTTLE experiments.

SUPPORTING INFORMATION AVAILABLE

Figure S1, electron density for L-tryptophan bound to PrnB; Figure S2, electron density for D-tryptophan bound to PrnB; Figure S3, electron density for 7-Cl-D-tryptophan bound to PrnB; Figure S4, electron density for 7-Cl-L-tryptophan bound to PrnB; Figure S5, pH dependence of 7-Cl-L-tryptophan binding to Fe23+ PrnB; Figure S6, alternative mechanism for the formation of the fused three ring structure. This material is available free of charge via the Internet at http://pubs.acs.org.

REFERENCES


BI7012189