Reverse Structural Genomics

AN UNUSUAL FLAVIN-BINDING SITE IN A PUTATIVE PROTEASE FROM BACTEROIDES THERAIOTAOMICRON**

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Background: The structure of a putative protease from Bacteroides thetaiotaomicron has a unique binding site for a flavin.
Results: The protein tightly binds to lumichrome, riboflavin, FMN, FAD, and flavin derivatives.
Conclusion: The putative protease is a scavenger of flavin and may function as a storage protein in gut bacteria.
Significance: The wealth of information available for cofactors is invaluable to assess protein structures.

The structure of a putative protease from Bacteroides thetaiotaomicron features an unprecedented binding site for flavin mononucleotide. The flavin isoalloxazine ring is sandwiched between two tryptophan residues in the interface of the dimeric protein. We characterized the recombinant protein with regard to its affinity for naturally occurring flavin derivatives and several chemically modified flavin analogs. Dissociation constants were determined by isothermal titration calorimetry. The protein has high affinity to naturally occurring flavin derivatives, such as riboflavin, FMN, and FAD, as well as lumichrome, a photodegradation product of flavins. Similarly, chemically modified flavin analogs showed high affinity to the protein in the nanomolar range. Replacement of the tryptophan by phenylalanine gave rise to much weaker binding, whereas in the tryptophan to alanine variant, flavin binding was abolished. We propose that the protein is an unspecific scavenger of flavin compounds and may serve as a storage protein in vivo.

Structural genomics efforts have led to an increasing number of structures of proteins that are not characterized on a functional biochemical level. In the case of homologous structures, biochemical function may be inferred to some extent and thus provide an initial lead for further functional investigations. In a recent survey of flavin-dependent proteins, we identified a putative protease from Bacteroides thetaiotaomicron, a microbe inhabiting the human gut (1, 2) that presumably binds compounds and may serve as a storage protein in vivo.

In contrast to zinc, the flavin cofactor occurs predominantly in oxidoreductases (~90% of flavin-dependent enzymes) and not hydrolases. Although the flavin cofactor is also present in other enzyme classes, like transferases, lyases, and isomerases, it has never been reported in a hydrolase (3). The large distance of ~16 Å between the edge of the flavin cofactor and the zinc argues against a direct cooperation between these two cofactors (Fig. 1D). This intriguing combination of unusual cofactors sparked our interest in the protein. Here, we report the biochemical characterization of the flavin-binding site of the protein and demonstrate that the recombinant protein is capable of binding not only naturally occurring flavin derivatives, such as lumichrome, riboflavin, FMN, and FAD (supplemental Fig. S1A) but also a variety of chemically modified “artificial” flavin analogs, such as iso-riboflavin (6.7-dimethyl-8-nor-riboflavin) and iso-FMN as well as 8-amino- and 1- and 5-deaza-riboflavin (12–14) (supplemental Fig. S1B). Thus it appears that this putative protease from B. thetaiotaomicron (ppBat)2 unspecifically

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This work is dedicated to Prof. Sandro Ghisla, a pioneer in the field of chemically modified flavins, on the occasion of his 70th birthday.
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2 The abbreviations used are: ppBat, putative protease from B. thetaiotaomicron; ITC, isothermal titration calorimetry; Ni-NTA, nickel-nitrilotriacetic acid.
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binds tricyclic aromatic systems such as the alloxazine and isoalloxazine ring systems. Furthermore, we showed that replacement of Trp^{164} (Fig. 1C) by alanine completely abolishes the ability to bind the flavin ring system, whereas the W164F variant retained some affinity. Sequence alignment of ppBat with other related putative proteases from the genus Bacteroides and related bacteria revealed that Trp^{164} is conserved in the majority of homologs. Therefore, we propose that ppBat may serve as a flavin storage protein in gut bacteria.

EXPERIMENTAL PROCEDURES

Reagents—All of the chemicals were of the highest grade commercially available from Sigma-Aldrich, Fluka (Buchs, Switzerland), or Merck. Ni-NTA-agarose was from GE Healthcare, and Sephadex resin was from Pharmacia Biotech. Flavin analogs were a generous gift from Prof. Sandro Ghisla and the late Prof. Peter Hemmerich (University of Konstanz, Konstanz, Germany).

Expression of Recombinant Protein in E. coli Host Cells—The gene *gatI* of *B. thetaiotaomicron* was cloned into a derivative of pET-28 for isopropyl β-D-thiogalactopyranoside-inducible expression in *E. coli* BL21 (DE3) host cells (kanamycin resistance). The N terminus of the expressed protein possesses a hexahistidine tag to facilitate purification. Precipitated cells were resuspended in lysis buffer (50 mM NaH₂PO₄, 300 mM NaCl, 10 mM imidazole, pH 8.0), using 2 ml of buffer/g of wet cells. At this stage lumichrome, riboflavin, FMN, FAD, or modified flavin analogs were added in excess to saturate the protein with the respective ligand. To obtain the apoprotein (protein preparation “as isolated”), no ligands were added. Resuspended cells were disrupted by 0.5-s sonication pulses for 10 min while cooling on ice. The cell debris was removed by centrifugation at 18,000 × g for 30 min at 4 °C. The hexahistidine-tagged protein was purified by Ni-NTA affinity chromatography, loading the supernatant onto a Ni-NTA HisTrap column (GE Healthcare), previously equilibrated with lysis buffer. After loading of the filtrated lysate, the column was washed with 10 column volumes of wash buffer (50 mM NaH₂PO₄, 300 mM NaCl, 20 mM imidazole, pH 8.0), and bound protein was recovered with elution buffer (50 mM NaH₂PO₄, 300 mM NaCl, 150 mM imidazole, pH 8.0). The purity of the eluted fractions was determined by SDS/PAGE. Fractions containing ppBat were pooled, dialyzed against 20 mM Tris/HCl buffer containing 100 mM NaCl (pH 8.0) overnight, and concentrated using Centripreps (Millipore). The final concentration was determined at 280 nm using an ε₂₈₀ of 13,075 M⁻¹ cm⁻¹. The protein was flash-frozen and stored at −20 °C.

Site-directed Mutagenesis—The W164F and W164A variants were prepared using a QuickChange XL-site directed mutagenesis kit (Stratagene) with a slight modification of the protocol. A two-step PCR was carried out, where separate reactions for forward and reverse primer were performed for four cycles prior to another 18 cycles where an equal amount of the forward and reverse reaction were mixed. Following primers were used for the insertion of the mutations: 5’-CTGACAAGATGAGAATACGATCGCGACAATGTTGCCGAAAGTCA-3’ and 5’-CTATGACTTTCCGGCAACATTGTGAAAGATCGTATTCTCATCTTGTGCA-3’ for W164F and 5’-CTGCAAGATGAGAATACGATCGCGACAATGTTGCCGAAAGATCGTATTCTCATCTTGTGCA-3’ and 5’-CTATGACTTTCCGGCAACATTGTGAAAGATCGTATTCTCATCTTGTGCA-3’ for W164A, respectively. After verification of positive transformants by sequencing, expression and purification was achieved as described for the wild-type protein.

UV-visible Absorbance and Fluorescence Spectroscopy—UV-visible absorbance spectra were recorded with a Specord 210 spectrophotometer (Analytik Jena, Jena, Germany). Fluores-
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cence emission spectra were recorded with a Shimadzu RF301 PC spectrofluorophotometer. Difference titrations were carried out at 25 °C in tandem cuvettes by the addition of ppBat to the flavin in the measurement cell and to buffer (20 mM Tris/HCl, 100 mM NaCl, pH 8.0) in the reference cell at 2-min intervals. Fluorescence titrations were performed at 25 °C in 20 mM Tris/HCl, 100 mM NaCl, pH 8.0 buffer. For quenching of the FMN fluorescence, 1.8 ml of 1 μM flavin solution were titrated with 2-μl aliquots of 100 μM protein solution every 2 min under stirring. Emission at 524 nm was measured at an excitation of 467 nm until an end point was reached. The spectra were corrected for dilution as well as for fluorescence of enzyme-bound flavin. In case of tryptophan fluorescence, 2 ml of 15 mM protein solution were titrated with 2-μl aliquots of 1.5 mM FMN solution every 2 min (emission at 343 nm and excitation at 282 nm).

Isothermal Titration Calorimetry (ITC)—Dissociation constants for the binding of flavins to ppBat were determined using a VP-ITC system (MicroCal). All of the experiments were performed at 25 °C in 20 mM Tris/HCl, 100 mM NaCl, pH 8.0 buffer, and solutions were degassed immediately before measurements. Titration experiments for FMN, FAD, riboflavin, and lumichrome consisted of 21 injections (7 μl; duration time, 14 s; spacing time, 300 s) of a flavin solution (530 μM) to 1.495 ml ppBat (40 μM). 40 μM enzyme were titrated with iso-FMN (370 μM; 21 × 10 μl; duration time, 10 s; spacing time, 300 s), 5-deaza-riboflavin (200 μM; 25 × 10 μl; duration time, 20 s; spacing time, 300 s), and 25 μM protein were titrated with 8-amino-riboflavin and iso-riboflavin (300 μM; 25 × 7 μl; duration time, 14 s; spacing time, 300 s), respectively. 40 μM of the W164F variant were titrated with 7-μl aliquots of 300 μM riboflavin for 25 times (duration time, 14 s; spacing time, 300 s). One set of sites fitting with Origin version 7.0 (MicroCal) for ITC data analysis was used to obtain dissociation constants.

Determination of Oligomeric State by Size Exclusion Chromatography—Determination of the oligomerization state for the wild-type protein and variants was carried out with a Superdex 200 analytical 10/300 GL column (GE Healthcare) using a ÄKTAPro explorer system. The calibration curve was prepared according to the manufacturer’s protocol. Approximately 1 mg of protein samples as isolated and saturated with FMN, respectively, were dissolved in 20 mM Tris/HCl, 100 mM NaCl, pH 8.0 buffer prior to size exclusion chromatography (flow rate, 0.5 ml/min).

Photoreduction—Photoreduction was carried out as outlined in a previous study (15). The protein was diluted in 20 mM Tris/HCl, pH 8.0, containing 100 mM NaCl, 1 mM EDTA, and 2 μM benzylioviolane. Quartz cuvettes were rendered anaerobic by repeated cycles of evacuation and flushing with nitrogen. Reduction of the flavin cofactor was achieved by irradiation of the sample with a 250-W halogen lamp that was cooled to 15 °C during the time of irradiation. Spectra from 300 to 700 nm were recorded at the same temperature until no further changes were observed. Then the cuvette was opened to air, and the absorbance spectrum was recorded.

Rapid Reaction Studies—Stopped flow measurements were carried out with a Hi-Tech (SF-61DX2) stopped flow device (TgK Scientific Limited, Bradford-on-Avon, UK) positioned in a glove box from Belle Technology (Weymouth, UK) at 25 °C. Reoxidation of FMN free in solution and bound to ppBat was measured by monitoring changes at A451 (ppBat) or A449 (FMN), respectively, with a KinetaScanT diode array detector (MG-6560; TgK Scientific Limited). Reduction of FMN was achieved by the addition of solid sodium dithionite. Reoxidation was then monitored by rapidly mixing the reduced solution with buffer previously equilibrated with air.

HPLC Determination of Flavins and Lumichrome—Cofactors were released by thermal purification of ppBat at 95 °C for 10 min, cooling on ice for 5 min, and centrifugation for 5 min. The flavin extracts were loaded onto an Atlantis dC18 8-μm 4.6 × 250-mm column (Waters) and eluted with a water/acetoni-trile multi-step gradient (0–1.5 min, 0–5% acetonitrile; 1.5–20 min, 5–60% acetonitrile; 20–22 min, 95% acetonitrile) at a flow rate of 1 ml/min. The elution was monitored by UV absorption at 370 nm, and the cofactors were identified according to their elution time and UV-visible absorbance spectra.

Zinc Determination—The amount of zinc in ppBat samples was determined by means of inductively coupled plasma optical emission spectrometry.

RESULTS

Expression and Purification of ppBat—Cells transformed with the plasmid carrying the gene of ppBat were cultured, and protein production was induced by isopropyl-β-D-thiogalactopyranoside. The majority of the target protein was soluble and purified by a single Ni-NTA affinity chromatography step (the progress of purification is shown in supplemental Fig. S2). The purified protein was obtained with an average yield of 30 mg/g of wet cells (~140 mg/l bacterial culture).

Characterization of the Isolated Protein—The purified protein was pale yellow and had absorbance maxima at 372 and 449 nm indicative for bound flavin. The absorbance ratio A280/A449 was very high (≈45) for a small flavoprotein and suggested that the protein was predominantly present in a flavin-free apo-form (Fig. 2, dotted line). When FMN was added to the extraction buffer, the protein was isolated with a reproducibly lower A280/A449 ratio of 5, indicating that the apoprotein present in the crude extract was reconstituted to the holo-protein (Fig. 2).
The extinction coefficient of bound flavin was determined by denaturation of the protein with subsequent determination of the UV-visible absorbance spectrum of the released FMN ($\epsilon_{\text{free}} = 11,500 \text{ M}^{-1} \text{ cm}^{-1}$) to $\epsilon_{\text{bound}} = 8,960 \text{ M}^{-1} \text{ cm}^{-1}$. Using this extinction coefficient, it was estimated that only 8–10% of the isolated protein is present in the holo-form when no flavin was added to the crude extract. The hypochromic effect at both absorbance maxima is accompanied by a long wavelength absorbance extending beyond the typical flavin absorbance (Fig. 2, compare dashed and solid lines). In contrast to the low flavin content of the purified protein, zinc was found in stoichiometric amounts, i.e., one zinc atom/protomer.

FMN bound to ppBat showed fluorescence excitation and emission maxima similar to that of free flavin; however, fluorescence intensity was $\sim 47\%$ lower as compared with free FMN. Similarly, FMN binding to ppBat quenched the tryptophan fluorescence emission at 343 nm to $\sim 28\%$.

Because the deposited structure of the protein has a truncated N-10 side chain (only four carbon atoms of the side chain were modeled; Fig. 1B), it is not clear which flavin derivative is bound to the protein. Therefore, we also tested riboflavin and FAD as potential binding partners of the recombinant protein. We found that riboflavin and FAD also bind to the protein with spectral changes similar to those seen with FMN. Further analysis by difference UV-visible absorbance spectroscopy revealed that all three flavin compounds bind tightly to the apoprotein with similar spectral perturbations (Fig. 3) and molar extinction coefficients (Table 1). All three flavin derivatives produced a sharp end point at equimolar concentrations of flavin and dimeric protein in agreement with a 1:1 stoichiometry as suggested by the three-dimensional structure.

To identify the ligand(s) bound to ppBat as isolated from E. coli cells, the protein was denatured, and the released compounds were subjected to HPLC analysis. As shown in the inset of Fig. 2, four peaks were observed. Based on the retention times and UV-visible absorbance properties these compounds were identified as FAD (peak 1), FMN (peak 2), riboflavin (peak 3), and lumichrome (peak 4).

**Determination of Dissociation Constants**—Because of the apparently tight binding of riboflavin, FMN, and FAD to apo-ppBat (Fig. 3), we employed ITC to determine their dissociation constants. As shown in Fig. 4, binding of these compounds to apoprotein is exothermic at 25 °C. The experimental data were fitted with a single site binding model and yielded dissociation constants in the range of 74.3 and 800.6 nM for riboflavin and FAD, respectively (Table 2). Obviously, flavin binding in the dimer interface of ppBat is adversely influenced by the size of the side chain attached to N-10.

**Binding of Chemically Modified Flavins**—In addition to the naturally occurring flavin compounds, ppBat also hosted a variety of derivatized riboflavin and FMN compounds, such as iso-riboflavin and iso-FMN (6.7-dimethylisoalloxazine), 2-thio-, 8-amino-, 1-, and 5-deaza-riboflavin, as well as 5-deaza-lumichrome.

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**TABLE 1**

| Flavin          | $\epsilon$ (free/bound) | $\lambda_{\text{max}}$ (nm) | Reference |
|-----------------|-------------------------|-----------------------------|-----------|
| Lumichrome      | 8,200/6,400             | 347/348                     | 28        |
| Riboflavin      | 12,500/9,800            | 444/449                     | 28        |
| iso-Riboflavin  | 7,300/5,650             | 448/454                     | 29        |
| 1-Deaza-riboflavin | 6,800/6110          | 536/558                     | 30        |
| 5-Deaza-riboflavin | 11,500/9,620         | 399/398                     | 14        |
| 8-Amino-riboflavin | 42,000/24,570       | 479/487                     | 31        |
| 2'-Deoxy-riboflavin | 12,500/9950        | 443/452                     | 32        |
| FMN             | 12,500/8,960            | 444/449                     | 28 and 32 |
| FAD             | 11,300/8,500            | 448/449                     | 28, 29, and 32 |
| iso-FMN         | 7,200/5,600             | 449/454                     | 33        |

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**FIGURE 3.** UV-visible absorbance difference titration of riboflavin (80 M, top panel), FMN (61 M, central panel), and FAD (60 M, bottom panel) with ppBat. The data points shown in the inset were manually fitted by straight lines to indicate the titration end points.

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FIGURE 4. Representative example for the determination of a dissociation constant by isothermal titration microcalorimetry. The measurement shows the titration of 40 μM ppBat with riboflavin in 20 mM Tris/HCl, 100 mM NaCl, pH 8.0 at 25 °C.

TABLE 2
Dissociation constants for binding of lumichrome, riboflavin, FMN and FAD as well as modified flavin analogs to ppBat as determined by ITC

| Flavin ligand      | $K_d$ (mM) |
|--------------------|------------|
| Lumichrome         | 272 ± 61   |
| Riboflavin         | 74 ± 5     |
| 5-Deaza-riboflavin | 84 ± 5     |
| iso-Riboflavin     | 96 ± 2     |
| 8-Amino-riboflavin | 227 ± 26   |
| 2'-Deoxy-riboflavin| 117 ± 3    |
| FMN                | 487 ± 19   |
| iso-FMN            | 295 ± 21   |
| FAD                | 801 ± 82   |

Properties of Variants W164F and W164A—To investigate the importance of the indole side chain of Trp$^{164}$ for dimerization of the protein and for flavin binding, two single replacement variants were generated by site-directed mutagenesis: W164F and W164A. Similar to wild-type ppBat, both variants were found to form dimers in solution as judged by size exclusion chromatography (supplemental Fig. S6). In the presence of FMN in the crude extract, the W164F variant retained the ability to bind FMN, whereas the W164A variant was essentially devoid of FMN (Fig. 7). In the absence of FMN in the crude extract, only trace amounts of flavin could be detected in both variants. Detailed HPLC analysis of ligands released upon denaturation of the protein variants revealed that small amounts of FAD, FMN, riboflavin, and lumichrome were bound (Fig. 7, insets). For the W164F variant, the dissociation constant of riboflavin was determined by ITC to 1.19 ± 0.15 mM, i.e., 16 times higher than for wild-type ppBat.

Recombinant ppBat loaded with FMN was found to be present as a dimer in solution using analytical size exclusion chromatography. Both tryptophan variants yielded a virtually identical peak, indicating that neither of the two amino acid substitutions affected dimerization of ppBat (supplemental Fig. S6).

Photoreduction and Reoxidation—Wild-type ppBat reconstituted with FMN was subjected to photoreduction under anoxic conditions. The course of the reduction is shown in Fig. 8. The spectral changes upon photoreduction featured a loss of absorption at 449 nm and an isosbestic point at 345 nm. The final spectrum is characteristic for the fully reduced dihydroquinone form of the flavin. During reduction, spectral hallmarks of a flavin radical species, like an increased absorbance at 370 nm or absorbance above 550 nm, were absent, indicating that no radical species is thermodynamically stabilized during reduction. The fully reduced dihydroflavin could be readily reoxidized by allowing access of oxygen to the cuvette (Fig. 8).

The rate of reoxidation was determined in a stopped flow apparatus by rapid mixing of sodium dithionite-reduced ppBat with air-equilibrated buffer (Fig. 8, inset). This yielded a $k_{obs} = 1.1 ± 0.02$ s$^{-1}$ (average of five independent measurements). Using the same experimental set-up, the reoxidation of free FMN yielded $k_{obs} = 6.7 ± 0.09$ s$^{-1}$, i.e., 6 times faster than for FMN bound to ppBat (all measurements were performed at 25 °C).

DISCUSSION

Flavins are widely used coenzymes for reduction-oxidation processes, with well over 300 different enzymatic reactions described in the literature (3, 16–18). In addition to their role as redox catalysts, flavins are also found in some none redox enzymes, such as isomerases, lyases, and transferases (3, 19). However, the occurrence of a flavin (purportedly FMN) in a ppBat was a novelty because hydrolases have never been associated with this cofactor. The discovery that ppBat features a flavin prompted us to investigate the properties of the flavin-binding site in this unusual family of proteases/chaperones (Fig. 1). At first, the low flavin content of isolated ppBat was surprising in view of the fact that the deposited crystal structure clearly showed the presence of flavin bound between two tryptophans in the dimer interface of the protein (Fig. 1, A and B). The

flavin and lumichrome. Lumazine did not bind to ppBat. UV-visible absorbance spectra of four selected examples are shown in Fig. 5 (see also supplemental Figs. S3 and S4). To determine the stoichiometry and affinity of these flavin analogs, difference titrations were performed with an UV-visible spectrophotometer. As shown in Fig. 6, we found tight binding with all of the tested flavin analogs and a stoichiometry of 1:1 (flavin: dimer) for complex formation with ppBat. The binding of five flavin analogs was further examined by ITC. The three riboflavin analogs, 5-deaza-riboflavin, iso-riboflavin, and 8-amino-riboflavin showed dissociation constants similar to riboflavin and iso-FMN bound even tighter than FMN (Table 2). Interestingly, 2’-deoxyriboflavin lacking the hydroxyl group at the 2’-position of the ribityl side chain had only a marginally higher dissociation constant (117.4 versus 74.3 nM; see also supplemental Fig. S5), suggesting that the carbonyl oxygen of Asn$^{161}$ forms only a weak hydrogen bond to the C2’-OH group (Table 2).
addition of a flavin, for example FMN, increased the flavin content close to one flavin per protein dimer, as could be demonstrated by titration experiments (see Figs. 3 and 6). According to the information given by the authors of the crystal structure, no flavin (neither riboflavin, FMN, nor FAD) was added in the crystallization trials, and hence it follows that the population of ppBat with flavin bound (10%) preferentially crystallized, i.e., was selected by the applied conditions. Because our analysis of ppBat isolated without flavin addition has revealed the presence of several chromophores (lumichrome, riboflavin, FMN, and FAD), the nature of the bound species obviously does not matter with regard to the propensity to generate crystals. The deposited crystal structure is also lacking information on the exact nature of the side chain because only the four carbon atoms proximal to N-10 were modeled. Therefore the structure cannot distinguish between riboflavin, FMN, and FAD. In the present study, we have shown that lumichrome as well as riboflavin, FMN, and FAD all bind to ppBat with dissociation constants in the nanomolar range (Table 2).

The UV-visible absorbance spectra of all flavins bound to ppBat showed pronounced hypochromicity and long wavelength absorbance indicative of a charge-transfer complex (Table 1). This is in contrast to the rather modest quenching of both the flavin and the tryptophan fluorescence emission.
Interestingly, virtually complete fluorescence quenching was observed for both fluorophores in model compounds, where stacking interactions between the isoalloxazine and indole ring were assumed (20). Apparently, aromatic stacking interactions do not necessarily lead to complete fluorescence quenching.

A salient feature of flavoproteins is the stabilization of either the red (anionic) or blue (neutral) flavin radical (semiquinone) (12). In the case of ppBat, reduction of bound flavin to the hydroquinone proceeded without the observable occurrence of a radical species (Fig. 7). Similarly, reoxygenation of the reduced flavin yielded the oxidized form without any observable radical species (Fig. 7). Similarly, reoxygenation of the reduced flavin yielded the oxidized form without any observable radical species (Fig. 7).

Our studies have clearly shown that Trp164 is crucial for binding of flavins to the dimer interface of ppBat. In contrast to the aromatic stacking interaction, the assumed contact between the C2′-hydroxyl group of the ribityl side chain and the main chain carbonyl of Asn161 contributes very little (1.14 kJ/mol) to the binding energy (Fig. 9). On the other hand, lumichrome, which features an alloxazine rather than an isoalloxazine π-electron system and has no side chain at N-10, binds weaker than riboflavin (~6 kJ/mol) and 2′-deoxy-riboflavin (~4.86 kJ/mol), suggesting that the side chain favorably affects binding to ppBat. Alignment of 29 sequences with at least 50% overall sequence identity showed that Trp164 and Asn161 are conserved in 18 and 22 proteins, respectively (Fig. 10). On the other hand, the two cysteines that complex the zinc ion are present in all of the 29 proteins. This suggests that the possibly catalytic function of the protein is related to the presence of the zinc center, and flavin binding may have evolved as an additional asset in a subset of homologous proteins.

The binding mode in ppBat also rationalizes our findings that the protein is rather insensitive to the N-10 side chain carried by the isoalloxazine ring. Most flavoenzymes exhibit pronounced preference for either FMN or FAD and riboflavin is not used at all except in one case (3). In addition, flavins with modifications in the isoalloxazine ring have lower affinity because of adverse effects on specific interactions with amino acids in the flavin-binding pocket. As demonstrated by our affinity measurements, binding of modified flavins is in the same range as for the naturally occurring flavins and mainly governed by the side chain (i.e., iso-riboflavin and 5-deaza-riboflavin have similar affinities than riboflavin, and iso-FMN is similar to FMN). Therefore it appears that ppBat does not utilize the bound flavin for catalysis. Likewise, because flavin binding is also not required to maintain the dimeric structure of ppBat, we conclude that flavin binding to ppBat does not concern the potential catalytic function of the protein. On the other hand, the binding of flavins, in particular of riboflavin, suggests that ppBat may play a role as a storage protein for gut bacteria.
A related flavin binding mode was recently discovered in a family of archael and bacterial proteins. These so-called dodecins consist of 12 protomers of 68–71 amino acids with six flavin-binding sites (23). As in ppBat, two tryptophans sandwich the isoalloxazine or alloxazine (lumichrome) ring system, although each binding site accommodates two ligands instead of one. In addition, other amino acids engage in hydrogen bond interactions to the flavin ligand, for example an arginine side chain forms hydrogen bonds to C-4\textsuperscript{O} of the isoalloxazine ring (23–25). In the case of ppBat, the only contacts are between the tryptophans of the two protomers and the isoalloxazine ring and a (weak) hydrogen bond between the carbonyl group of Asn161 of one protomer and the C\textsuperscript{2-hydroxyl group of the ribityl side chain. Contacts between the pyrimidine ring and ppBat are not observed in the structure, and hence the affinity of flavins to ppBat is mainly governed by the interaction with the indole side chains of the two tryptophans. Therefore it is not surprising that replacement of the tryptophans by alanine and phenylalanine has marked effects on the binding affinity. In the case of alanine, binding was virtually abolished, whereas the W164F exchange has retained some affinity (Fig. 7). In contrast, replacement of Trp38 in dodecin from \textit{Thermophilus} has apparently no consequence on the propensity to bind flavins because several other interactions still ensure stable complex formation (see Protein Data Bank entry 2vyx (23)). This protein also binds coenzyme A, and it appears that the dodacamer binds a mixture of cofactors \textit{in vivo} (23, 25). Therefore dodecins appear to serve as cofactor storage proteins and may also be important as scavengers of potentially harmful cofactor degradation products, such as lumichrome, which is produced in the course of photodegradation of riboflavin \textit{in vivo} (23, 25, 26). The dissociation constants determined for dodecin from \textit{Thermophilus} are similar to those found for ppBat and support the notion that ppBat may also serve as a storage protein in gut bacteria \textit{in vivo}. A BLASTp search using the dodecin sequence of \textit{Thermus thermophilus} against the \textit{B. thetaiotaomicron} genome failed to retrieve a dodecin homolog.

The distance between the plane of the indole ring of the two tryptophans and the isoalloxazine in ppBat is 3.5–3.7 Å (Fig. 9). A similar distance for \textit{H}-stacking interaction of a tryptophan and an isoalloxazine ring was seen in the dodecins (3.4 to 3.8 Å) and was also found in the case of FMN adenylyltransferase (27). Apparently, this distance is optimal for \textit{H}-stacking interactions, although the relative orientation of the interacting aromatic ring systems differs in the above mentioned proteins. Because the main contribution for flavin binding stems from the \textit{H}-stacking interaction with the two tryptophan residues, we assume that the asymmetric isoalloxazine binds in two possible orientations in the symmetric binding site (2-fold axis) provided by the two protomers of the dimeric protein. The electron density of Protein Data Bank entry 3cne is consistent with 50\% occupancy of the two possible orientations of the tricyclic ring system.

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