Joint Functions of Protein Residues and NADP(H) in Oxygen Activation by Flavin-containing Monoxygenase
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The reactivity of flavoenzymes with dioxygen is at the heart of a number of biochemical reactions with far reaching implications for cell physiology and pathology. Flavin-containing monooxygenases are an attractive model system to study flavin-mediated oxygenation. In these enzymes, the NADP(H) cofactor is essential for stabilizing the flavin intermediate, which activates dioxygen and makes it ready to react with the substrate undergoing oxygenation. Our studies combine site-directed mutagenesis with the usage of NADP(H) analogues to dissect the specific roles of the cofactors and stabilizing the crucial flavin-hydroperoxide. This concept surrounding protein matrix. The highlight of this “double-engineering” approach is that subtle alterations in the hydrogen bonding, electrostatics, and accessibility of the flavin will represent a challenge for the design and development of oxidases and monooxygenases for biotechnological applications.

Flavin-containing monooxygenases (FMOs) form a large family of enzymes present almost ubiquitously among living organisms (1, 2, 3). They are involved in diverse biological processes such as the biosynthesis of various natural products (4) and catabolism of xenobiotics. Humans have five different isozymes that are coded by different genes and represent a key enzymatic system in drug metabolism, whose relevance almost equals that of cytochrome P450. Among known FMO substrates, there are metabolites directly or indirectly deriving from food digestion such as trimethylamine, drugs such as tamoxifen, and toxic molecules such as nicotine. Mutations in the gene for the isozyme 3 of human FMO (FMO3), which is most abundant in the liver, are directly responsible for trimethylaminuria, also known as fish odor syndrome (TMAU). This genetic disease causes the accumulation of trimethylamine so that the body of the patients tend to have an unpleasant smell (5, 6).

Irrespective of their biological context, FMOs always catalyze the same reaction: the oxygenation of a soft nucleophile using molecular oxygen as oxygen source and NADPH as electron donor (7). At the heart of this very complex reaction is the stabilization of the flavin-hydroperoxide adduct resulting from the reaction of dioxygen with the two-electron reduced FAD (Fig. 1A). Equally essential is the role of NADP(H). This cofactor not only acts as the electron donor that reduces the flavin but it directly takes part also in formation/stabilization of the flavin-hydroperoxide. This notion is best illustrated by the fact that using alternative electron donors (e.g. dithionite) in place of NADPH does not support catalysis because the reaction of oxygen with the artificially reduced flavin generates hydrogen peroxide rather than the stable flavin-hydroperoxide needed for monooxygenation (8, 9, 10). In the cellular milieu, FMOs are thought to be mainly in the flavin-hydroperoxide state, ready to attack a suitable substrate (the so-called “cocked gun”) (11).

The peculiar catalytic role of NADP(H) in oxygenation makes FMOs a very attractive system to study the more general problem of oxygen reactivity in flavoenzymes. Because it has been established that the nicotinamide cofactor remains bound to enzyme during oxygenation, NADP(H) can be exploited as a probe to gain insight into the reaction of dioxygen with the reduced flavin and to interrogate the elements that promote the stabilization of the crucial flavin-hydroperoxide. This concept represents the starting point of our study that builds on the previous analysis of the crystal structure and basic biochemical properties of the FMO from Methylophaga sp. strain SK1 (mFMO, Fig. 1B) (12). This bacterial enzyme shares many features with the human FMO3 including a considerable degree of
sequence identity (33%), a similar substrate specificity, and the ability to form a stable flavin-hydroperoxide intermediate (13). At the same time, mFMO is better suited for detailed structural and enzymological investigations because it is a soluble protein (i.e. not membrane-associated). The aim of our study is 2-fold: (i) to gain insight into the astonishing properties of FMOs that combine in a single active center two cofactors and a complex sequence of catalytic steps that lead to the formation of an “activated-form” of dioxygen (flavin-hydroperoxide, Fig. 1A), and (ii) to translate this knowledge into a molecular rational for the effects of mutations found in patients affected by TMAU. For these purposes, we have probed the mFMO catalytic properties by altering both the active site amino acids by site directed mutagenesis and the NADP(H) substrate by using analogues of this cofactor. Such a “double-engineering” strategy has revealed that very fine details in the stereochemical and hydrogen-bonding environment around the flavin determine the efficiency of the reaction with dioxygen and the nature of the resulting products.

EXPERIMENTAL PROCEDURES

Protein Purification and Reagents—All reagents were purchased from Sigma-Aldrich. Mutagenesis, protein expression, and purification were carried out using standard protocols as previously described (13). All the experiments were carried out using the E158A/E159A mutant. This surface mutation does not affect the enzymatic properties but is essential for the growth of suitable crystals (13). Throughout the text, we shall refer to this protein as the wild type.

Steady-state Kinetics—The steady-state activity of the mFMO variants was measured spectrophotometrically by monitoring the decrease of NADPH in time at 340 nm ($\epsilon_{340\text{ nm}} = 6.22 \text{ mm}^{-1} \text{ cm}^{-1}$). The reaction mixture (1.0 ml) typically contained 50 mM Tris/HCl pH 8.0, 100 mM NaCl, 5% (v/v) glycerol, 3 mM $\beta$-mercaptoethanol, 0–200 $\mu$M NADPH, 0–2 mM trimethylamine, and 0.2–0.6 $\mu$M mFMO. Trimethylamine was chosen because it is the best substrate for mFMO (13) and is commonly used for FMO activity evaluation (7). The oxidase activity was probed in the same conditions except for the absence of trimethylamine. The corresponding $k_{\text{uncoupling}}$ (Table 1) was measured by monitoring the consumption of NADPH (spectrophotometrically) and oxygen (using an oxygen electrode, Hansatech Oxygraph). Furthermore, the production of hydrogen peroxide was verified using the well-known amplex red/peroxidase assay as described (14). Reduced NADP analogues (3-acylpyridine adenine dinucleotide and thionNADP) were generated enzymatically using glucose-6-phosphate dehydrogenase or phosphite dehydrogenase. In a typical experiment, solutions (150 $\mu$M) of reduced dinucleotides were produced by overnight incubation in 50 mM Tris-HCl pH 8.0, 150 $\mu$M glucose-6-phosphate or phosphite, and 1 unit of dehydrogenase. Dinucleotide reduction was monitored spectrophotometrically. Enzymatic activities were then carried out as with NADPH. For 3-acylpyridine adenine dinucleotide (APADP), the commercially available reduced form was used as additional check (OYC Europe). Lack of a monoxygenase activity was inferred from the absence of an increased rate of oxygen or NADPH consumption upon addition of trimethylamine. When no monoxygenase activity was detected with trimethylamine, a very sensitive colorimetric assay using indole (whose monoxygenation product spontaneously dimerises to indigo) was used as an additional check for oxygenation activity. Kinetic parameters were determined by measuring at $\geq 5$ different substrate concentrations and the observed rates were measured in duplicate or triplicate. Michaelis-Menten kinetic parameters were obtained by fitting...
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TABLE 1

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<tr>
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<th>Steady state</th>
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<th>Presteady state</th>
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a Performed using 150 $\mu M$ thioNADPH and monitoring oxygen consumption.

b ND, not determined.

 webcam performed using 150 $\mu M$ APADPH and monitoring oxygen consumption.

the data using Origin software (version 7.0 for Windows) and were within 15% standard deviation.

Presteady-state Kinetics—Reductive and oxidative half-reactions of mFMO proteins were analyzed using a stopped-flow apparatus from Applied Photophysics Ltd. (model SX17MV) equipped with a diode array detector. Spectral data (300–720 nm) were collected at time intervals of 2.5 ms. The obtained spectra were analyzed by means of numerical integration methods using Pro-K software (Applied Photophysics Ltd.), yielding individual observed rate constants and deconvoluted spectra (Table 1 and Fig. 2). All experiments described below were performed at 25 °C in the same buffer used for steady-state experiments using 8–12 $\mu M$ enzyme (after mixing). Each reaction condition was measured in triplicate. Rate constants obtained by fitting were within 20% standard deviation. Anoxic conditions were achieved by flushing the system and solutions with N2 and removing traces of oxygen upon addition of 10 mM NADP$^+$.

Steady-state and presteadate-state kinetic parameters (Table 1 and Fig. 2). All experiments described below were performed at 25 °C in the same buffer used for steady-state experiments using 8–12 $\mu M$ enzyme (after mixing). Each reaction condition was measured in triplicate. Rate constants obtained by fitting were within 20% standard deviation. Anoxic conditions were achieved by flushing the system and solutions with N2 and removing traces of oxygen upon addition of 10 mM NADP$^+$.

Reduction of mFMO by NADPH was measured anoxically at various concentrations of the reduced nicotinamide coenzyme. The observed rates increased hyperbolically with increasing NADPH concentration and could be fitted with Equation 1 using non-linear regression analysis (Origin version 7.0 for Windows).

$$k_{\text{obs}} = \frac{k_{\text{red}[\text{NADPH}]}}{[\text{NADPH}]} + K_0 \quad \text{(Eq. 1)}$$

For measuring the oxidation of reduced mFMO, the enzyme was first anoxically reduced outside the stopped-flow apparatus by adding a slight 1.2-fold excess of NADPH. Only in the case of the N78K mutant a 10-fold excess of NADPH was needed to obtain the reduced enzyme form, due to the relatively poor NADPH affinity of this mutant. The reduced enzyme was subsequently mixed with oxygenated buffer containing equimolar amounts of NADP$^+$. By varying the oxygen concentration (125, 360, and 595 $\mu M$), the oxygen dependence of the rates for the reoxidation events was probed. The affinity for NADP$^+$ ($K_{p, \text{NADP}^+}$) was determined by measuring the change in the flavin absorbance spectrum upon titration of 10 $\mu M$ of each mFMO variant in 50 mM Tris·HCl, pH 7.5 with 0–500 $\mu M$ NADP$^+$ in the same buffer.

Crystalization and Structure Determination—All crystals were obtained using microbatch technique under 100% paraffin oil at 4 °C by mixing equal volumes of 8 mg protein/ml in 25 mM Tris·HCl, pH 8.0, 250 mM NaCl, 1 mM NADP$^+$ (or analog), and 20% (w/v) PEG4000 in 100 mM Na/HEPES pH 7.5 as described (13). Data were collected at the European Synchrotron Radiation Facility (Grenoble, France) and processed with the CCP4 package (15). The structures were solved by difference Fourier methods. Phases were further improved by density 4-fold averaging with DM (16). The programs COOT (17) and Refmac5 (18) were used for model rebuilding and refinement, which was carried out using strict non-crystallographic symmetry restraints (Table 2). Figures were generated using PyMOL (DeLano, 2002 on World Wide Web).

RESULTS

Functional Properties of Wild-type mFMO—To provide a starting point for our studies, we have first investigated the enzymatic properties of the wild-type enzyme. Steady-state kinetic analysis revealed that, similar to mammalian FMOs, trimethylamine, and NADPH are efficient substrates (Table 1). Furthermore, it was found that the enzyme has considerable NADPH oxidase activity as gathered from the rate of NADPH depletion in the absence of an organic substrate ($k_{\text{uncoupling}} = 0.5 \text{ s}^{-1}$, Table 1). The ability of mFMO to act as NADPH oxidase was further confirmed by measuring the formation of hydrogen peroxide, which approximately equaled the amount of consumed NADPH.

The presteady-state analysis using the stopped-flow technique showed that mFMO is rapidly and almost fully reduced by NADPH ($k_{\text{red}}$ in Table 1). The reductive-half reaction data also indicated that the enzyme has a high affinity for the reduced nicotinamide coenzyme. Titration of the oxidized mFMO revealed a somewhat lower affinity for NADP$^+$ (Table 1). The NADPH-reduced protein was employed to monitor the oxidative half-reaction of mFMO, which can be best described by a three-step process (rate constants are listed in Table 1 and Fig. 2A). At first, the relatively stable flavin-hydroperoxide (or flavin peroxide) intermediate is rapidly formed (spectrum B in Fig. 2A) with a typical absorbance maximum at around 360 nm. Next, the intermediate is converted to a second species (spectrum C), which might represent a conformational change occurring during re-oxidation and/or protonation of the intermediate (i.e. conversion of flavin peroxide into flavin hydroperoxide). A partly similar spectral intermediate has been observed also in the oxidative half-reaction of the pig-liver FMO (10).

Finally, the enzyme is completely re-oxidized (spectrum D). It is of note that the actual reaction of the flavin with oxygen (i.e. formation of the flavin-hydroperoxide; $A \rightarrow B$ in Fig. 2A) linearly depends on the oxygen concentration as typically found in
flavin-dependent oxidases and monooxygenases (rate constant $k_{\text{ox}}$ of 6.5 x $10^3 \text{M}^{-1} \text{s}^{-1}$, Table 1). By repeating the experiments with 1 mM trimethylamine, it was found that $k_{\text{ox}}$ does not depend on the binding of substrate. This is in contrast with flavoprotein hydroxylases and P450 monooxygenases but is fully in line with previous kinetic studies on eukaryotic FMOs (10). Consistent with this mode of function, the three-dimensional structure of mFMO exhibits a two-domain organization that defines a large cleft at the domain interface (Fig. 1B). At the bottom of this cleft, a small pocket is defined by the nicotinamide moiety of NADP$^+$, the side chain of Asn-78, and the N5-C4a flavin atoms. This is the site where dioxygen reacts with the reduced flavin forming the flavin-hydroperoxide, which remains exposed to the incoming substrate (Fig. 3A).

The N78S Mutant—The position of Asn-78 in direct contact with the C4a atom of the flavin and close to the NADP$^+$ nicotinamide ring. This indicated that this side chain could play a key role in the reaction with dioxygen (Figs. 1 and 3A). Furthermore, two mutations affecting the corresponding residue in human FMO3 have been identified in TMAU patients (19, 20, 21). Thus, Asn-78 represented an obvious target for site-directed mutagenesis.

At first, we generated the N78S mutation, which is one of the mutations known to be the cause of TMAU. It is a relatively conservative amino acid replacement that does not greatly perturb the bulkiness of the side chain and maintains a group capable of H-bonding interactions. Nevertheless, we found that the mutant is inactive against various potential FMO substrates (trimethylamine, nicotine, methimazole, and indole). The mutant was further investigated for its reactivity toward NADPH and oxygen. Steady-state kinetics showed that N78S has oxidase activity with a $K_m$ for NADPH of about 4 $\mu$M and a rate constant ($k_{\text{uncoupling}}$) of 0.7 s$^{-1}$, which is very similar to that of the wild-type (Table 1). Stopped-flow kinetics confirmed that the mutant enzyme readily reacts with NADPH. It also demonstrated that the enzyme reacts with oxy-

NADP$^+$ nicotinamide ring, which is poorly defined by the electron density whereas the ADP-ribose moiety is clearly visible in the density map. This feature indicates that the nicotinamide is partly disordered. The main lesson from the N78S analysis is that an apparently small alteration in the flavin C4a environment is sufficient to convert FMO into a relatively efficient oxidase, explaining why the corresponding N61S in human FMO3 leads to enzyme deficiency in patients affected by TMAU (20, 21).

Adding Charges: N78D and N78K—Having analyzed the effect of a small hydrophilic side chains at position 78, we further probed this site by introducing charged side chains. We generated and studied the mutations N78D and N78K, the latter also being found in TMAU patients (19). Both mutants turned out to be unable to catalyze trimethylamine monooxygenation but exhibited drastically different enzymatic properties.

N78D functions as moderately efficient oxidase ($k_{\text{uncoupling}}$ of 0.19 s$^{-1}$) although it binds NADPH with a $>$20-fold worse affinity than the wild-type (Table 1). As for the N78S protein, the presteady-state kinetics showed that the N78D mutant is unable to effectively stabilize the flavin-hydroperoxide with the re-oxidation process occurring in one step without detectable intermediates ($k_{\text{ox}}$ of 2200 $M^{-1} \text{s}^{-1}$). These functional properties can be evaluated in light of the crystallographic analysis of the mutant enzyme bound to NADP$^+$. The structure displays a peculiar feature: the nicotinamide is flipped out and hosted at the rim of the active site cleft (Fig. 3B). This binding mode is brought about by a rotation of the nicotinamide ribose, which does not affect the conformation of the ADP fragment of NADP$^+$. In this position, the cofactor carboxamide is engaged in a H-bond with the Asp-78 side chain, which compensates for the nicotinamide positive charge. As a result, the flavin N5 atom remains exposed to solvent without the shielding exerted by NADP$^+$ as in the wild-type protein (Fig. 3A). Although this binding mode does not require significant conformational
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Changes in the active site residues, it is evident that such a modified NADP$^+$-binding drastically alters the structure, shape, and electrostatics of the active site which becomes unable to promote accumulation of the flavin-hydroperoxide intermediate. This fully supports the idea that proper NADP$^+$ binding is an essential element for the oxygenation reaction.

The N78K mutant exhibited strongly altered catalytic properties and remarkable structural features. In addition of being inactive as monooxygenase, the protein is very inefficient in using NADPH for flavin reduction (Table 1). Equally impaired is the reaction of the NADPH-reduced enzyme with oxygen, which occurs without any observable intermediate at a rate ($k_{ox}$ of 670 M$^{-1}$ S$^{-1}$), which is not dissimilar to that of the free flavin in water. The structural data further emphasize the large perturbation caused by the mutation (Fig. 3C). The Lys-78 side chain points toward the nicotinamide ribose possibly forming a H-bond with the ribose hydroxyl groups (the NE atom of Lys-78 has only weak electron density). The nicotinamide of NADP$^+$ is disordered, lacking any significant electron density. This feature is coupled to a conformational change in the position of Tyr-212, which swings into active site pointing toward interior of the protein (Fig. 3C). This conformation of Tyr-212 combined with the bulkiness of Lys-78 makes the flavin C4a very crowded and inaccessible, which accounts for the poor reactivity of this mutant with dioxygen.

The structure of the N78K mutant indicates that Tyr-212 may function as an active-site backdoor. In this scenario, the conformation found in the N78K protein may well correspond to that present in the NADP$^+$-free enzyme. Thus, the flavin of the unliganded enzyme is predicted to be partly shielded by Tyr-212, which, upon nicotinamide cofactor binding, swings out to position itself above the nicotinamide ring, establishing favorable stacking interactions with the nicotinamide ring (Fig. 3, A and C). Tyr-212 could also play a role in product release, displacing NADP$^+$. A fascinating observation concerning this aspect is that the function of Tyr-212 of mFMO is reminiscent of that played by an active-site tyrosine in human glutathione reductase, a flavoenzyme that uses NADPH as flavin reductant and shares the same dinucleotide-binding topology as mFMO (22).

Modifying the Substrate: NADPH Analogues—Having established how critical is the environment around the flavin C4a atom for the oxygen reaction and for attaining proper NADP$^+$ binding, we investigated the effects on catalysis exerted by alterations in the NADP(H) substrate. To this aim, we employed two NADP(H) analogues, thioNADP(H) and APADP(H) (Fig. 4A). Both compounds, in the oxidized form, bind tightly to the protein as inferred by the significant increase (5–7 °C) in the protein melting temperature measured by ThermoFAD method as reported (23).

The crystal structure of the complex with thioNADP$^+$ revealed that the dinucleotide binds exactly as NADP$^+$. The only noticeable difference concerns the pyridine ring, which is disordered as gathered from the lack of well-resolved electron density in all four crystallographically independent subunits present in the crystal asymmetric unit (Table 2). We found that thioNADPH is an efficient substrate both in the oxidase and in the monooxygenase assays (Table 1). Thus, replacement of the
and FAD (carbons in blue) probe the enzymatic properties. First of all, the reduced form of NADP (Fig. 4A) seems to significantly affect catalysis (Fig. 4A).

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FIGURE 4. Binding of NADP analogues to mFMO. A, chemical formulas of the two compounds using for this study (APADP on the left and thioNADP on the right). B, structure of the complex with APADP in approximately the same orientation and coloring scheme as in Fig. 1B. The carbons of APADP are in green, H-bonds are shown as dashed lines. C, comparison between the binding of NADP and APADP using a composite picture that shows the protein surface of the wild-type protein bound to NADP (carbons in cyan) and FAD (carbons in yellow). The pyridine ring of APADP is superimposed to highlight its different position at the rim of the active-site cleft surface. The NADP and APADP complexes are very similar (root-mean-square deviation of 0.21 Å for the Cα atoms).

oxygen atom of the nicotinamide with a bulkier sulfur does not seem to significantly affect catalysis (Fig. 4A).

APADP(H) turned out to be a more insightful compound to probe the enzymatic properties. First of all, the reduced form of this cofactor analog (both commercially obtained or generated enzymatically) is unable to sustain any monooxygenase activity. Instead, we found that it can function as substrate in the oxidase assay with a rate constant close to that measured with NADPH (kuncoupling in Table 1) implying that APADPH can reduce the flavin. The complex of mFMO with APADP produced crystals of very good quality resulting in excellent electron density maps that were even better than those obtained with the NADP+ bound proteins. The cofactor analog binds with the ADP-ribose group in the standard conformation (Fig. 4B). Conversely, the pyridine ring is positioned at the entrance of the active site cleft in a binding mode, which is almost identical to that observed for NADP+ in the structure of the N78D mutant (Figs. 3B and 4B). This conformation enables Asn-78 to form a H-bond interaction with the carbonyl group of APADP+, leaving the flavin ring unprotected. Combining these biochemical and structural data leads to conclusion that the reduced APADPH must be able to adopt the conformation required for reducing the flavin (Fig. 1A). However, after flavin reduction, the pyridine moiety moves away from the flavin. A reason for the differing binding modes between NADP+ and APADP+ might be that the acetyl group of APADP+ is unable to donate a H-bond to the flavin N5 (Figs. 1A, 3A, and 4B). This can be expected to favor the “flipped out” conformation which allows the APADP+ acetyl group to H-bond to Asn-78 (Fig. 4B). As for the N78D mutant, such a “out” conformation is unable to support oxygenation because it does not generate the proper environment for flavin-hydroperoxide stabilization and, possibly, because it may hamper the access of the organic substrate to the flavin (Fig. 4C). On the other hand, the flavin remains accessible to dioxygen and thereby competent in the oxidase reaction.

DISCUSSION

The specific objective of our study was to investigate the reaction of FMO with dioxygen using site-directed mutagenesis and NADP(H) analogues as dissecting tools. The general aim was to improve our knowledge of the molecular basis of TMAU and to advance our current understanding of the oxygen reactivity in flavoenzymes. Along these lines, we can draw several conclusions.

The analysis of the Asn-78 mutants indicates that at least some of the TMAU-causing mutations target the Achilles heel of the enzyme: the intrinsic “fragility” of the crucial flavin-hydroperoxide intermediate, which, if not appropriately stabilized, causes the conversion of the enzyme into an oxidase.

In this scenario, the fact that the devil is in the detail is best illustrated by the N78S variant (Figs. 2A and 3A). This mutation does not affect the steric hindrance and hydrophilic nature of the side chain and does not cause any detectable change in the active site conformation. Nevertheless, it fully inactivates the enzyme. Evidently, the fine details in the environment of the C4a atom (Fig. 1A) can make a tremendous difference in the flavin reactivity. In this case, it appears that a Ser side chain in position 78 is unable to provide the precisely positioned H-bond partner required to stabilize the flavin-hydroperoxide, whose terminal oxygen can accept and donate H-bonds (Figs. 1A and 3A). Therefore, the intermediate does not form at all or, more likely, it decays too rapidly to sustain monooxygenation (Fig. 2B). As a result, the mutant enzyme retains (albeit less...
efficient reactivity with oxygen but only to function as oxidase and not as monooxygenase (Table 1).

The N78D mutant illustrates the role of electrostatics. This isosteric mutation positions a negative charge close to the N5 atom which can enormously enhance the stability of the flavin-hydroperoxide to the point that the reaction of oxygen with the flavin N5 atom can be biocatalytically useful hydroperoxide derivative (26, 27). Although the N5 atom of the two-electron reduced flavin can be isosteric with a monooxygenase (Table 1). As observed for other flavoenzymes, it is likely that this mutation disfavors the more general conclusion from this analysis is that the two alternative pathways (oxidation versus monooxygenation) along the oxygen reaction can be predicted to have very similar energy barriers so that subtle alterations in the balance of interactions around the C4a-N5 atoms of the flavin can drastically alter the outcome of the reaction of the reduced enzyme with dioxygen. This will be a crucial aspect to be considered in the design and development of FMOs and related monooxygenases for biocatalytic application.

Acknowledgment—We thank Dr. Andrea Alfieri for contributions in the earlier stages of the project.

REFERENCES

TABLE 2
Crystalllographic data collection and refinement statistics

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<td>21.4</td>
<td>21.4</td>
<td>18.4</td>
<td>21.0</td>
</tr>
<tr>
<td>R free (%)</td>
<td>21.4</td>
<td>24.4</td>
<td>21.1</td>
<td>20.2</td>
<td>25.2</td>
</tr>
<tr>
<td>Rms bond length (Å)</td>
<td>0.016</td>
<td>0.024</td>
<td>0.022</td>
<td>0.014</td>
<td>0.020</td>
</tr>
<tr>
<td>Rms bond angles (°)</td>
<td>1.7</td>
<td>1.9</td>
<td>1.9</td>
<td>1.6</td>
<td>2.0</td>
</tr>
</tbody>
</table>

n° of atoms = 15,209 15,348 15,065 16,288 15,494

Values in parentheses are for reflections in the highest resolution shell.

a Rsym = Σh|Ih|−|I|/Σh, where I is the intensity of ì-th observation and <|I|> is the mean intensity of the reflection.

b Values in parentheses are for reflections in the highest resolution shell.


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