Structural, spectroscopic and catalytic activity studies on glutathione reductase reconstituted with FAD analogues

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FAD-modified human glutathione reductases were reconstituted from apoenzyme using the FAD analogues 6-SH-FAD, 6-SCN-FAD, 6-OH-FAD, 6-NH2-FAD and 8-OH-FAD. The catalytic activities of the modified enzymes were substantially lower than for the native enzyme. All five species could be crystallized, but only those containing 6-SH-FAD, 6-OH-FAD and 6-NH2-FAD yielded crystals that could be analyzed. X-ray analyses and structural refinements were performed at 0.27 nm and 0.30 nm resolution resulting in R factors around 13.5%. The crystal structures showed the additional non-hydrogen atoms and small conformational changes of the polypeptide that were obviously induced by the substituents of the FAD analogues. The observed changes together with spectroscopic and activity data permit some conclusions about the chemical nature of the substituents.

The flavoenzyme glutathione reductase catalyzes the reduction of oxidized glutathione at the expense of NADPH. The human enzyme is a homodimer of two 52.4-kDa subunits with one FAD/subunit. It appears structurally and functionally well known [1-3; its crystal structure has been established at 0.154 nm resolution [4]. Numerous substrate- and ligand-binding studies were performed in order to elucidate enzyme catalysis by correlating structural with kinetic data [5-7]. Recently, these experiments were supplemented by site-directed mutations of the glutathione reductase species from Escherichia coli [8].

For studying flavin-polypeptide interactions, FAD analogues carrying modifications at the isoalloxazine ring have been used as substitutes for native FAD [9]. Human glutathione reductase is a particularly good model system for such experiments, because its structure is precisely known and its apoenzyme is reasonably stable and binds various FAD analogues. Some FAD-modified glutathione reductases have already been studied with respect to their spectroscopic and catalytic properties [10-12]. Here, we analyze FAD-modified glutathione reductases by X-ray diffraction and attempt to correlate structural with spectroscopic and catalytic data.

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**Abbreviations.** GR(6-SH-FAD), the FAD-modified enzyme glutathione reductase from human erythrocytes as reconstituted from the apoenzyme and 6-SH-FAD (the same nomenclature is used for the other FAD analogues); Fmod,obs, observed structure factor amplitudes for crystalline glutathione reductase reconstituted with FAD analogues; F60H,obs, and F6NH2,obs indicate particular modifications; Fnat,obs, observed structure factor amplitudes for the native enzyme; Fcalc, calculated structure factor phases for the native enzyme.

**Enzyme.** NADPH:glutathione oxidoreductase (EC 1.6.4.2).

**MATERIALS AND METHODS**

In our experiments we used the FAD analogues 6-SH-FAD [13], 6-SCN-FAD [13], 6-OH-FAD [14], 6-NH2-FAD [15] and 8-OH-FAD [16] all of which are specified in Fig. 1. Glutathione reductase was isolated from human erythrocytes as described by Krohne-Ehrich et al. [17]. The apoenzyme was obtained by saturating a holoenzyme solution with (NH4)2SO4 at 4°C and pH 3.0 [18]. This leads to precipitation and release of FAD. The precipitate was washed several times. Because of the extreme pH conditions this procedure was finished as fast as possible, usually within 90 min. The FAD-free precipitate was dissolved and stored in 100 mM Tris/ HCl, 10 mM EDTA, pH 8.7, at concentrations of 4-7 mg apoenzyme/ml. The protein concentrations were determined photometrically using A280 = 1.0 as the absorption coefficient for 1 mg apoenzyme/ml [19].

The enzyme was reconstituted by adding a slight excess of the respective FAD analogue to the apoenzyme. After incubating for 1 h at room temperature to allow complete binding of the FAD analogue, the solution of the reconstituted enzyme was centrifuged using an Amicon Centricon 10 ultrafilter to remove excess FAD analogue and low-molecular-

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**Fig. 1. Isoalloxazine moiety of FAD showing the modified positions.** For all FAD analogues R is ribityl-5'-ADP, as in native FAD.
mass contaminants. In the same manner, the buffer was changed and the enzyme was concentrated for crystallization. The spectral studies were performed in storage buffer consisting of 100 mM potassium phosphate, 200 mM KCl, 1 mM EDTA, 14 mM mercaptoethanol, pH 7.0. For 6-SCN-FAD we used the same procedures, but avoided mercaptoethanol throughout. The enzyme activity was measured under standard conditions [17] in 100 mM potassium phosphate, 200 mM KCl, 1 mM EDTA, 0.5 mg/ml bovine serum albumin, pH 6.9, at 25°C.

The reconstituted FAD-modified enzymes were crystallized at 20°C using the hanging-drop method under the same conditions employed previously for the native enzyme [20]. The drop consisted of 0.8 M \((\text{NH}_4)_2\text{SO}_4, 0.1 M\) potassium phosphate, pH 7.0, with a protein concentration of about 16 mg/ml. The reservoir was filled with 1 ml 1.2 M \((\text{NH}_4)_2\text{SO}_4, 0.1 M\) potassium phosphate, pH 7.0. Crystallization was also achieved (often with better results) without the buffer. The crystals could be stored for months in 1.5 M \((\text{NH}_4)_2\text{SO}_4, 0.1 M\) potassium phosphate, pH 7.0. The same buffer was used for the X-ray analyses.

X-ray data were collected on a modified four-circle diffractometer [21] and processed to obtain the structure factor amplitudes \(F_{\text{mod,obs}}\). Taking the observed structure factor amplitudes \(F_{\text{nat,obs}}\) and the calculated phases \(x_{\text{nat,calc}}\) of the native enzyme from [4], a \((F_{\text{mod,obs}} - F_{\text{nat,obs}}) \cdot \exp(i\cdot x_{\text{nat,calc}})\) difference-Fourier map was calculated that showed the structural changes caused by the modifications of FAD. Electron densities and models were visualized on a graphics system (model PS330, Evans & Sutherland, USA) using the program FRODO [22]. For structural refinement of the 6-SH-FAD-modified enzyme \((\text{GR}(6-\text{SH-FAD}))\) we used program TNT [23]. When the more convenient program XPLOR [24] became available, it was applied for refining the 6-OH-FAD- and 6-NH2-FAD-modified glutathione reductases, \(\text{GR}(6-\text{OH-FAD})\) and \(\text{GR}(6-\text{NH}_2\text{-FAD})\), respectively.

RESULTS

Spectral and catalytic properties

The FAD-modified glutathione reductases can be characterized by their absorption spectra [12, 14−16, 25, 26]. The measured \(\lambda_{\text{max}}\) values of the main absorption bands of the bound FAD analogues are given in Table 1. On binding to the apo form of the enzyme, the \(\lambda_{\text{max}}\) values of the FAD analogues increased, usually in the order of 10 nm. However, no increase in \(\lambda_{\text{max}}\) was observed with 6-OH-FAD.

The analogue 6-SH-FAD can be deprotonated at the substituent S6x (i.e. sulfur bound to C6, see Fig. 1). On dissociation, an electron pair of S6x conjugates with the isoalloxazine nucleus giving rise to an absorption band around 650 nm. Using this absorption band, the \(pK\) value of 6-SH-FAD has been determined as 5.9 in the free state [13], and as \(< 5.0\) when bound to glutathione reductase [11]. The electronic and spectral properties of 6-OH-FAD are similar to those of 6-SH-FAD; it has a similar band around 650 nm. The \(pK\) value of 6-OH-FAD has been determined as 7.1 in the free state [10, 14]. We have now measured a \(pK\) of 5.6 ± 0.5 for the bound analogue in \(\text{GR}(6\text{-OH-FAD})\). Given these \(pK\) values, the enzyme-bound 6-OH-FAD and 6-SH-FAD analogues are predominantly in their anionic state at the pH 7.0 of the X-ray analyses.

The increased acidic nature of enzyme-bound 6-SH-FAD and 6-OH-FAD as compared to the free state must result from a stabilization of the anionic charge within the protein. In 6-SH-FAD and 6-OH-FAD the negative charge is partially located at the substituent atoms S6x and O6x and partially in the isoalloxazine nucleus, where it is enhanced in the region around N1 and O2x. A partial negative charge at O6x and S6x is stabilized by the contacting positively charged amino group of Lys66. A partial negative charge around N1 and O2x is stabilized by the favorably oriented amide dipoles of the \(\alpha\)-helix of amino acid residues 339−354 that contacts O2x [27].

For 6-NH2-FAD in the free state, no spectral changes, and thus no ionization, occur in the physiological pH range. There are two measurable \(pK\) values, the first one around 1 where N6x accepts a proton, and the second one around 10 where N3 loses a proton [28]. The \(pK\) for the deprotonation of the 6-NH2 group is greater than 13 [28]. As a consequence, the 6-NH2 substituent has no formal charge at pH 7.0 of the X-ray analysis. Since the lone electron pair of N6x conjugates with the isoalloxazine, nucleus, however, N6x carries a partial positive charge.

The catalytic activities of the modified glutathione reductase analogues are also given in Table 1. All of them are appreciably lower than the activity of the native enzyme. Due to the error margins, the activities of \(\text{GR}(6\text{-SCN-FAD})\) and \(\text{GR}(6\text{-OH-FAD})\) could be as high as 1% of that of the native enzyme and the activity of \(\text{GR}(8\text{-OH-FAD})\) could be essentially zero.

Structural studies

Crystallography. Crystals could be obtained from human glutathione reductase reconstituted with all five FAD analogues. Crystals suitable for X-ray diffraction, however, grew only from \(\text{GR}(6\text{-SH-FAD})\), \(\text{GR}(6\text{-OH-FAD})\) and \(\text{GR}(6\text{-NH}_2\text{-FAD})\). The enzymes \(\text{GR}(6\text{-SCN-FAD})\) and \(\text{GR}(8\text{-OH-FAD})\) could only be crystallized up to sizes of 250 × 150 × 70 \(\mu\)m\(^2\) and 200 × 100 × 100 \(\mu\)m\(^2\), respectively, which are too small for reasonable data collection on a diffractometer. The crystallization of \(\text{GR}(8\text{-OH-FAD})\) was complicated by the weak binding of this analogue to the apoenzyme, which may have given rise to inhomogeneous protein ensemles. The relatively low binding constant of 8-OH-FAD as compared to FAD was discovered when 8-OH-FAD was slowly washed out on concentrating \(\text{GR}(8\text{-OH-FAD})\) in the Centricon tube.

<table>
<thead>
<tr>
<th>Enzyme-bound FAD</th>
<th>(\lambda_{\text{max}}) of main absorption band</th>
<th>Relative catalytic activity</th>
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<tr>
<td>FAD</td>
<td>463</td>
<td>100</td>
</tr>
<tr>
<td>6-SH-FAD</td>
<td>449</td>
<td>3</td>
</tr>
<tr>
<td>6-SCN-FAD</td>
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<td>5</td>
</tr>
<tr>
<td>8-OH-FAD</td>
<td>484</td>
<td>2</td>
</tr>
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Table 1. Spectral and catalytic properties of glutathione reductase reconstituted with FAD analogues

All data were obtained at pH 7.0 in the buffers given in the text. The catalytic activity is defined as \([\text{specific activity of FAD-modified enzyme}] / [\text{specific activity of apoenzyme}]/[\text{specific activity of native enzyme}]\). The error for the 6-substituted FAD analogues is ±1% and that for 8-OH-FAD is ±2% relative catalytic activity, indicating that \(\text{GR}(8\text{-OH-FAD})\) could be inactive. The larger error for 8-OH-FAD relates to the lower binding affinity of this analogue
All three analyzed FAD-modified human glutathione reductases crystallized in space group B2 and are isomorphous with the native enzyme crystals (Table 2). Data were collected up to 0.27 nm resolution. The structural refinements converged at $R$ factors around 13.5% with good geometries (Table 2). The quality of the structure analyses is also demonstrated in the maxima of the difference-Fourier maps of GR(6-SH-FAD) and GR(6-OH-FAD). These maxima are at the S6a and O6a positions, and their heights (Table 2) correlate well with the number of electrons of sulfur and oxygen. At the given resolution and the achieved good $R$ factors, the general accuracy of the atomic coordinates can be expected to be 0.03 nm.

In the native enzyme there is a cavity around the C6 atom of isoaalloxazine [4]. This hole extends over a distance of about 0.5 nm in the direction of the C6-S6a, C6-O6a and C6-N6a bonds of the modified flavins. Given this free space, the incorporation of the additional 6-SH, 6-OH and 6-NH$_2$ groups caused only small conformational changes of the polypeptide. For the most part, these changes occurred at residues Gly62, Cys66 and Glu201, which surround the substituents at C6. It should be noted that the environment of the C6 atom is polar, which in conjunction with the available space explains the high binding constants of FAD analogues that carry a small polar group at C6. Since the obtained resolutions of the X-ray analyses are merely 0.27 nm and 0.30 nm, the quantification of these shifts is limited by the residual error.

The structure of GR(6-SH-FAD). The highest peak in the difference-Fourier map between GR(6-SH-FAD) and native glutathione reductase exceeds the second highest peak by a factor of four. As shown in Fig. 2, it corresponds to the sulfur of 6-SH-FAD. The 'observed' C6-S6a bond length was taken as the distance between the density maximum of the difference-Fourier map and the position of the C6 atom after the refinement. In this way, the observed bond length becomes largely independent of the respective bond length used in the refinement. The observed C6-S6a bond length was 0.16 nm, which is slightly, but not significantly, shorter than a double bond (0.17 nm) and a single bond (0.18 nm) [29].

Since the pK value of 6-SH-FAD is lower than 5.0 when bound to glutathione reductase [11], the FAD analogue should be present in its anionic state with deprotonated S6a. Presumably, the negative charge at S6a is partially delocalized into the flavin-ring system, where it is most likely concentrated around N1 and O2a and stabilized by the helix of amino acid residues 339–354. This is consistent with the observed C6-S6a bond being on the shorter side.

The structure of GR(6-OH-FAD). The highest peak in the difference-Fourier map between GR(6-OH-FAD) and native glutathione reductase exceeds the second highest peak by a factor of two. As shown in Fig. 2, the differences-Fourier map exceeds the next highest peak by a factor of two. It is caused...
Fig. 2. *Difference-Fourier map of glutathione reductase containing 6-SH-FAD calculated from\( (F_{\text{obs}} - F_{\text{cal}}) \cdot \exp(i\pi n)\). For clarity, only positive density is given. The contours are drawn at 15% of the electron-density maximum of this map. The model is centered at the S6a atom. Part of the FAD analogue together with chain segments around Cys58, Cys63, Lys66 and Glu201 (all labelled) is depicted, as well as parts of Tyr197, Ile198 and Arg291 (not labelled); (+) water molecules.

Fig. 3. *Difference-Fourier map of glutathione reductase containing 6-OH-FAD calculated from\( (F_{\text{obs}} - F_{\text{cal}}) \cdot \exp(i\pi n)\). For clarity, only positive density is given. The contours are drawn at 28% of the electron-density maximum of this map. The model is centered at the O6a atom. Part of the FAD analogue together with chain segments around Cys58, Cys63, Lys66 and Glu201 (all labelled) is depicted, as well as parts of Tyr197, Ile198 and Arg291 (not labelled); (+) water molecules.

Fig. 4. *Structural differences between glutathione reductase containing 6-OH-FAD (thick lines) and the native enzyme (thin lines). Atom O6a is marked by a dot. Depicted is a part of FAD together with residues Cys63, Lys66 and Glu201.

by the additional oxygen O6x (Fig. 3). In the same manner as with 6-SH-FAD, the C6-O6x bond length was determined as 0.12 nm, which corresponds to a double bond and is smaller than a single bond [29]. The short C6-O6x bond is consistent with a deprotonated O6x, the negative charge of which is delocalized towards the N1-O2x region of the isoalloxazine, where it is stabilized by the helix of amino acid residues 339–354 [27]. The deprotonation of O6x at pH 7.0 of the X-ray analysis is in agreement with the measured pK value of 5.6 ± 0.5 for 6-OH-FAD bound to glutathione reductase.

Other remarkable difference-Fourier peaks are close to O4x and C4a of isoalloxazine (Fig. 3). In the refinement one of these peaks corresponded to an appreciable shift of O4x caused by a tilt of the isoalloxazine by about 5° (Fig. 4). The other peak relates to the disulfide bridge which moves together with the adjacent isoalloxazine atoms by about 0.03 nm, keeping the distance constant (Fig. 4).

The refined coordinates indicate further that the tilt of the isoalloxazine has moved the C6 atom by about 0.04 nm toward the salt bridge Lys66-NZ–Glu201-OE2. The resulting distances O6x…NZ, O6x…OE2 and NZ…OE2 are 0.29 nm, 0.35 nm and 0.29 nm, respectively, making a partial negative charge at O6x rather likely. Obviously, O6x forms such a strong hydrogen bond to Lys66-NZ that the whole
Fig. 5. Difference-Fourier map of glutathione reductase containing 6-NH$_2$-FAD calculated from $(F_{6 \text{NH}_2, \text{obs}} - F_{\text{nat, obs}}) \cdot \exp^{2\pi i x}$, For clarity, only positive density is given. The contours are drawn at 30% of the electron-density maximum of this map. The model is centered at the N6 atom. Part of the FAD analogue together with chain segments around Cys58, Cys63, Lys66 and Glu201 (all labelled) is depicted, as well as parts of Tyr197, Ile198 and Arg291 (not labelled); (+) water molecules.

Fig. 6. Structural differences between glutathione reductase containing 6-NH$_2$-FAD (thick lines) and the native enzyme (thin lines). Atom N6 is marked by a dot. Part of the FAD analogue together with residues Cys63, Lys66 and Glu201 is depicted.

isoalloxazine follows the movement of O6 and tilts. The isoalloxazine tilt must involve an appreciable amount of energy because it concerns the most rigid region of the native enzyme [4].

The structure of GR(6-NH$_2$-FAD). The highest peak of the difference-Fourier map corresponds to the additional nitrogen N6 in (Fig. 5). Using the method described for 6-SH-FAD, the C6-N6 bond length was determined as 0.15 nm, corresponding to a single rather than a double bond [29].

As with GR(6-OH-FAD), other important peaks of the difference-Fourier map occur close to O4 and C4 of isoalloxazine (Fig. 5). The refinement revealed that these peaks are caused by a tilt of the isoalloxazine by almost 10°, similar to the tilt observed in GR(6-OH-FAD). Again, there is a substantial shift of O4 related to one of the peaks; and again the disulfide bridge follows the adjacent isoalloxazine atoms resulting in a longitudinal 0.05 nm movement (Fig. 6).

As compared with the native enzyme, the C6 atom of isoalloxazine is shifted by 0.05 nm toward the salt bridge Lys66-Glu201. Moreover, the salt bridge is rearranged. Lys66-NZ has moved away from C6, forming the closest contact to Glu201-OE1 (0.27 nm) instead of Glu201-OE2. The resulting distances N6...NZ, N6...OE2 and NZ...OE2 are 0.36 nm, 0.30 nm and 0.30 nm, respectively. The large isoalloxazine tilt in a very rigid region of the native enzyme [4] indicates that the N6...OE2 hydrogen bond formed is rather strong. N6 makes a further hydrogen bond to the water molecule Sol-89, which is shifted by 0.03 nm toward N6 as compared to the native enzyme.

Clearly, the lengths of the distances between the new substituent and the salt bridge atoms have reversed. While the substituents S6 and O6 are close to Lys66-NZ and about 0.05 nm further away from Glu201-OE2, N6 is 0.06 nm closer to OE2 than to NZ. This observation is consistent with deprotonated S6 and O6 carrying a partial negative charge and therefore attracting the positive partner NZ via a hydrogen at NZ. It also agrees with two hydrogens and presumably a partial positive charge at N6 which is repelled by the positively charged NZ amino group, but attracted via one of its own hydrogens to the carboxylate atom OE2.

DISCUSSION

The observed structural changes increase in the order GR(6-SH-FAD) < GR(6-OH-FAD) < GR(6-NH$_2$-FAD). The large sulfur of 6-SH-FAD contacts Lys66-NZ at a reasonable hydrogen bond distance of 0.3 nm and causes almost no change. Only Gly62 and the redoxactive disulfide bridge are pushed somewhat away. In GR(6-OH-FAD) and GR(6-NH$_2$-FAD) the isoalloxazine tilts and twists (Figs 4 and 6) to form hydrogen bonds between the new substituent and the adjacent salt bridge Lys66-NZ...Glu201-OE2. Most conspicuously, the O6 and N6 atoms contact different partners of the salt bridge, N6 causes a salt-bridge rearrangement in comparison to the native enzyme.

All data (i.e. the observed conformational changes, the determined bond lengths between C6 and the substituents, the protonation state at pH 7.0 of the X-ray analyses as deduced from spectroscopically derived pK values) agree with a deprotonated thiol in 6-SH-FAD, a deprotonated hydroxyl in 6-OH-FAD and an amino group in 6-NH$_2$-FAD. For 6-SH-
FAD and for 6-OH-FAD the $pK$ value drops on binding to glutathione reductase, which is consistent with the stabilization of a negatively charged substituent by Lys66-NZ. A further contribution derives most likely from a delocalization of the negative charge towards N1 and O2a of the isoalloxazine, where it is stabilized by the helix of amino acid residues 339—354.

The enzyme activities of all FAD-modified enzymes analyzed are appreciably reduced (Table 1). The highest residual activity (5%) is found for GR(6-NH$_2$-FAD), although this analogue causes the largest conformational changes. This indicates that catalysis is much more affected by changing the electronic properties of the isoalloxazine than by polypeptide rearrangements.

This notion is corroborated by the residual 3% enzymatic activity of GR(6-SH-FAD) as compared to no activity for GR(6-OH-FAD) as given in Table 1. In both cases, the flavin redox potential has dropped by the conjugation of electron pairs from thiolate or hydroxylate with the isoalloxazine nucleus. While this conjugation is limited for the large thiolate atom, it is much more pronounced for the smaller hydroxylate atom giving rise to a redox potential drop of about 100 mV [16], which renders the enzyme inactive. Using this argument, the largely reduced activity of GR(6-NH$_2$-FAD) points to a conjugation between the lone electron pair of N6a and the isoalloxazine nucleus causing some decrease of the redox potential and a partial positive charge at N6a.

In conclusion, the described analyses of FAD-modified enzymes improved our knowledge of the chemical nature of the FAD analogues and of their interactions with functional groups of the polypeptide. The internal salt bridge Lys66-Glu201 is intimately connected with protonation state and partial charge of the substituents at C6 of the bound flavin.

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