

The structure of ergothioneine biosynthetic sulfoxide synthase EgtB

Kristina V. Goncharenko^{[a]†}, Allegra Vit^{[b]†}, Wulf Blankenfeldt^{[b]*} and Florian P. Seebeck^{[a]*}

Abstract: The non-heme iron enzyme EgtB catalyzes O₂-dependent carbon-sulfur bond formation between γ -glutamyl cysteine and N- α -trimethyl histidine as the central step in ergothioneine biosynthesis. Both, catalytic activity and architecture of EgtB are distinct from known sulfur transferases or thiol dioxygenases. The crystal structure of EgtB from *Mycobacterium thermoresistibile* in complex with γ -glutamyl cysteine and N- α -trimethyl histidine reveals the two substrates and three histidine residues as ligands of an octahedral iron binding site. This active site geometry is consistent with a catalytic mechanism in which C-S bond formation is initiated by an iron (III)-complexed thiyl radical attacking the imidazole ring of N- α -trimethyl histidine.

Ergothioneine (**1**, Figure 1) occurs in a broad range of prokaryotic and eukaryotic organisms, including humans or human pathogens such as *Mycobacterium tuberculosis*.^[1] Higher eukaryotes absorb ergothioneine as a micronutrient, which stems from microbial production. The precise cellular function of ergothioneine is not known, but recent observations on animal, fungal and bacterial cells suggest that this sulfur compound may be a protectant against oxidative stress.^[2] *Mycobacteria* biosynthesize ergothioneine from glutamate, cysteine and histidine.^[1d, 3] The central step in this pathway is catalyzed by the non-heme iron enzyme EgtB which forms and sulfoxidizes a C-S bond between N- α -trimethyl histidine (**2**, TMH) and γ -glutamyl cysteine (γ GC) using O₂ as oxidant. Together with the ovothiol (**4**) biosynthetic enzyme OvoA (Figure 1),^[4] EgtB represents a distinct enzyme class (sulfoxide synthases) with no relation to sulfur oxidizing or C-S bond forming iron enzymes such as cysteine dioxygenase or isopenicillin synthase.^[5] Instead, sulfoxide synthases present a new entry to a rich collection of C-S bond forming catalysts.^[6]

To elucidate the structural basis for sulfoxide synthase activity, we determined the crystal structure of EgtB from *Mycobacterium thermoresistibile* (EgtB_{thermo}) in complex with iron and TMH, and as a quaternary complex with manganese, N- α -dimethyl histidine (DMH) and γ GC. We produced this enzyme in *Escherichia coli* following the protocols established for the EgtB homolog from *Mycobacterium smegmatis* (EgtB_{smegmatis}).^[1d] The

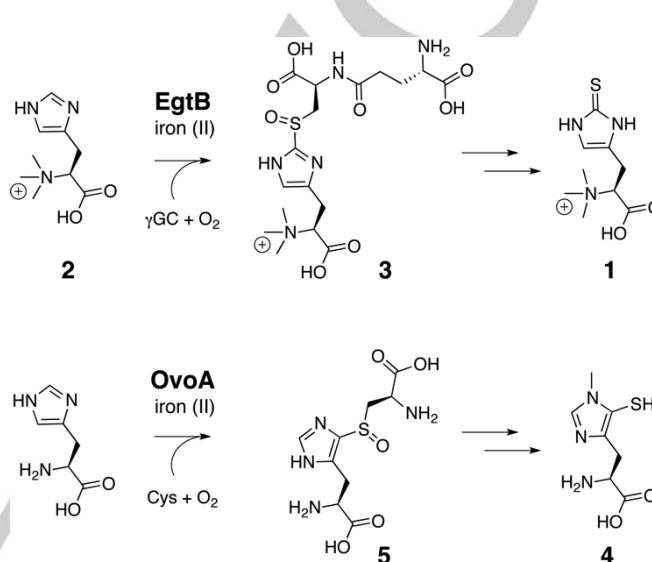


Figure 1. EgtB and OvoA catalyzed C-S bond formation and sulfoxidation between γ -glutamyl cysteine and N- α -trimethyl histidine (TMH) or between cysteine and histidine as the central step in ergothioneine (**1**) and ovothiol (**4**) biosynthesis.

two enzymes share 81 % sequence homology, both are monomeric (Figure S1) and display similar *in vitro* activities (Table 1). Both recombinant enzymes purified to a significant extent as iron-bound holoenzymes as inferred by a ferrozine-based colorimetric assay (EgtB_{thermo} > 95 %, EgtB_{smegmatis} > 50 %, Table S1) and by titration of EgtB activity with FeSO₄ (Figure S2). *In vitro* activity was assayed in HEPES buffered solutions in the presence of TMH, γ GC, 4 μ M FeSO₄, 2 mM ascorbate, 100 mM NaCl and 2 mM TCEP at 25°C. Formation of the sulfoxide product (**3**, Figure 1, Figure S3) was monitored by cation-exchange HPLC.^[1d] Under these conditions EgtB_{smegmatis} and EgtB_{thermo} catalyzed up to one turnover per second and remained active for hundreds of turnovers (Table 1, Figure S3). In absence of ascorbate the activity of EgtB_{thermo} ceased after 120 \pm 20 turnovers, but fully recovered after addition of 2 mM ascorbate (Figure S4). Incubation with γ GC alone also inactivated EgtB_{thermo} (Figure S4) and induced an absorption band at 565 nm (ϵ_{565} = 450 M⁻¹cm⁻¹, Figure S5) which also vanished after addition of ascorbate. A similar absorption feature has been observed in cysteine-bound ferric cysteine dioxygenase where it was assigned as a sulfur to iron (III) charge transfer transition.^[7] By contrast, no change in activity or absorbance was induced by incubation of EgtB_{thermo} with TMH or buffer alone. We suspect that a γ GC-dependent side reaction oxidizes EgtB into the inactive ferric state. The absorption at 565 nm of

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Table 1. Kinetic parameters of EgtB variants.^[a]

| enzyme | donor | acceptor | $k_{\text{cat,donor}}$ (s^{-1}) ^[b] | $k_{\text{cat,donor}}/K_{\text{M}}$ ($\text{M}^{-1}\text{s}^{-1}$) | $k_{\text{cat,acceptor}}$ (s^{-1}) | $k_{\text{cat,acceptor}}/K_{\text{M}}$ ($\text{M}^{-1}\text{s}^{-1}$) |
|---------------------------|-------------|----------|---|--|---|---|
| EgtB _{smegmatis} | γ GC | TMH | 1.1 | 1.4×10^4 | 1.2 | 2.8×10^4 |
| EgtB _{thermo} | γ GC | TMH | 8.6×10^{-1} | 2.0×10^4 | 8.7×10^{-1} | 2.2×10^4 |
| EgtB _{thermo} | NGC | TMH | 2.5×10^{-1} | 2.3×10^2 | n.a. | n.a. |
| EgtB _{D416N} | γ GC | TMH | 2.7×10^{-1} | 1.3×10^2 | 1.1×10^{-1} | 1.9×10^4 |
| EgtB _{D416N} | NGC | TMH | 1.0×10^{-1} | 1.1×10^3 | n.a. | n.a. |

[a] γ GC = gamma-glutamyl cysteine; TMH = N- α -trimethyl histidine; NGC = N-glutaryl cysteine; displayed values are averages from three independent measurements, errors correspond to less than 15 % of averaged value. [b] k_{cat} and $k_{\text{cat}}/K_{\text{M}}$ were determined in the presence of co-substrate at a concentration at least 3-fold higher than the corresponding K_{M} and in air saturated buffers. Michaelis-Menten plots are displayed in the supporting information.

inactive EgtB_{thermo} in complex with γ GC provides a first indication that the sulfur atom of this ligand may directly interact with the catalytic iron center. The following structural analysis will provide strong support for this notion.

We crystallized EgtB_{thermo} as apo protein (mthEgtB_apo), in complex with iron and TMH (mthEgtB_TMh) and in complex with manganese, DMH and γ GC (mthEgtB_DMh_ γ GC). The crystals belong to space group P₄₃2₁2 with cell constants a,b=135 and c=141 Å. The apo structure was phased using the anomalous signal of the intrinsic sulfur atoms and the coordinated iron. The phases were later combined with native data to yield the structure of EgtB_{thermo} at resolutions of 1.7 Å (apo). Ligand complexes of EgtB_{thermo} were obtained to a resolution of 1.6 Å (TMH) and 1.98 Å (γ GC and DMH). In all structures, the asymmetric unit includes two protein chains sharing only a relatively small and asymmetric interface.

The electron densities of the three EgtB_{thermo} structures reveal a continuous polypeptide chain from Pro7 to Asp434 (Figure 2). Residues 7 – 150 fold to a DinB-like four-helix bundle with long linkers between helices 1 – 2 (18 residues), 2 – 3 (34 residues) and 3 – 4 (7 residues). The fourth helix is followed by an extended two-stranded β -sheet (residues 151 to 210) wrapped around the C-terminal domain, which adopts a C-type lectin fold (CLec).^[8] This fold contains few secondary structure elements but instead is stabilized by a dense array of buried ionic interactions such as salt bridges between Arg to Glu residues (Arg409:Glu196; Arg413:Glu296; Arg:397:Glu300; Arg428:Glu360). In addition, a calcium cation in the center of the C-terminal domain immobilizes seven oxygen ligands from side chains and backbone amides (Met354, 2.8 Å; Gly399, 2.9 Å; Val358, 2.6 Å; Gly356, 2.7 Å; and Gln353, 3.6 Å, Glu360 3.3 Å). This unusual, loop-rich structure is conserved in at least two single domain proteins with entirely different functions, with less than 30 % sequence homology and without dependence on transition metals.^[8a, 8c] One of these structural homologs (r.m.s.d. of 0.741, Figure S6) is the human formylglycine-generating enzyme (FGE) which catalyzes O₂-dependent posttranslational maturation of sulfatases.^[8c] The second homolog is the diversity-generating retroelement variable protein TvpA from *Treponema denticola*.^[9] Iron dependent sulfoxide synthases such as EgtB and OvoA must have emerged from these FGE-like proteins by fusion with an N-

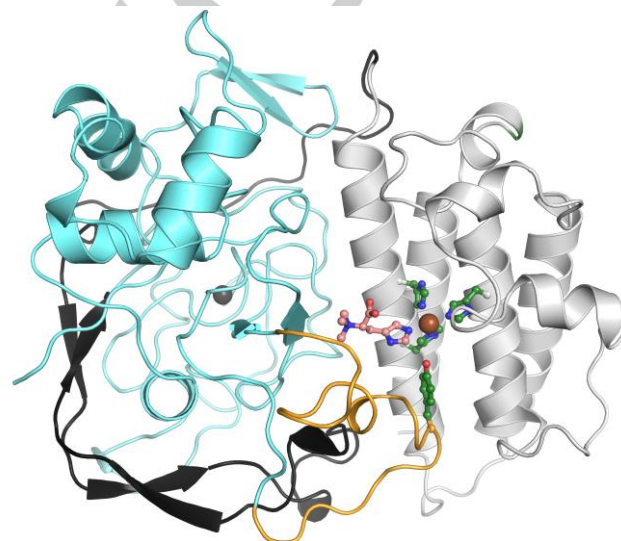


Figure 2. Cartoon diagram of EgtB_{thermo} in complex with iron (brown) and TMH (salmon). The protein consists of an N-terminal DinB domain (grey, residues 1 – 150), two-stranded β -sheet region (black, residues 151 – 210), and a C-terminal C-type lectin domain (cyan/orange, residues 211 – 434). The active site containing a 3-histidine facial triade (green, His51, His134, His138) is formed between the DinB domain and residues 370-425 (orange)

terminal DinB domain. It is not surprising that the EgtB active site maps to the interface between the two domains.

The active site of EgtB_{thermo} is housed in a 15 Å deep and 10 Å wide tunnel lined by residues 375 through 425 from the CLec domain and residues from helices 2 and 4 and the loops between helices 1, 2 and 3. At the bottom of this tunnel three histidines form the DinB domain (residues 51, 134 and 138) coordinate the catalytic iron cation (Fe-N: 2.1 Å, 2.1 Å and 2.3 Å). This structure revises our earlier prediction that Glu140 rather than His51 is a metal ligand.^[1d, 4a]

In hindsight, we note that this facial triade is fairly conserved among proteins from the DinB protein superfamily.^[10] Several single domain DinB proteins have been characterized as zinc-dependent thiol S-transferase or maleylpyruvate

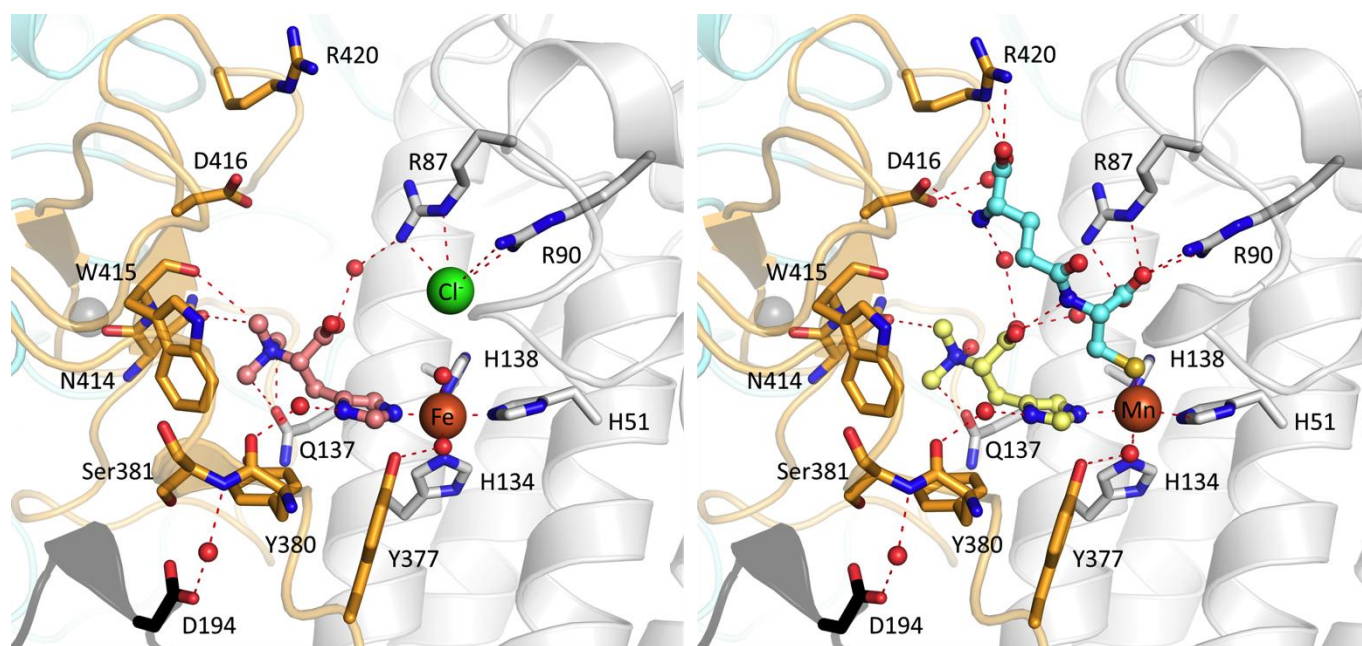


Figure 3. Active site of EgtB_{thermo} in the ternary complex with TMH and iron (left), or with DMH, γ GC and manganese (right). Localized waters are in red. A chloride ion (green) in the ternary complex occupies the cationic docking site of γ GC. Unbiased difference electron densities of bound ligands are shown in Figure S7.

isomerases,^[11] indicating that metal-catalyzed C-S bond formation may be a common activity among DinB-like proteins.^[11a] However, the mechanisms of iron and O₂ dependent oxidative C-S bond formation as catalyzed by EgtB and OvoA are most likely very different from that of zinc-promoted alkyl transfers.^[6a, 12] With this hypothesis in mind we proceeded to study the substrate binding mode of EgtB_{thermo}.

In the ternary complex containing EgtB_{thermo}, TMH and iron (ternary complex) the substrate imidazole ring (Fe-N τ : 2.2 Å, Figure 3) and two water molecules (Fe-O: 2.1 and 2.2 Å) join the 3-histidine facial triad in an octahedral coordination sphere around the iron center. The substrate imidazole ring also interacts with Tyr380 through a water-mediated hydrogen bond (5.4 Å), and the 1-carboxylate group is loosely connected to Arg87 (4.9 Å), again via a bridging water molecule. Two N- α -methyl groups of TMH pack against the indole side chain of Trp415. The third N- α -methyl group appears to make dipolar contacts to the amide side chains of Gln137 (3.2 Å) and Asn414 (3.5 Å).

To identify the binding mode of the second substrate γ GC we examined the structure of the quaternary complex between EgtB_{thermo}, DMH, γ GC and manganese (II). Initial attempts to soak the iron containing binary complex with γ GC resulted in disintegration of the crystals. As a solution to this problem we used manganese-reconstituted EgtB_{thermo}. This enzyme complex is inactive^[1d] but the corresponding crystals were tolerant to soaking with DMH and γ GC and diffracted to a resolution of 1.98 Å (Figure 3, Table S2 & S3). In this

complex the active site residues and DMH adopt superimposable positions with respect to the ternary complex (r.m.s.d. = 0.041). γ GC coordinates as the fifth ligand to the metal center (Fe-S: 2.6 Å). This direct metal-thiolate contact is consistent with the observed sulfur to iron charge transfer transition absorption band in the γ GC complex with ferric EgtB_{thermo} (Figure S5). The α -amino group and the two carboxylates of γ GC form salt bridges to Asp416 (2.8 Å), Arg420 (2.7 Å), Arg87 (3.0 Å) and Arg90 (2.6 Å). The amid function of γ GC hydrogen bonds with the 1-carboxylate of DMH (2.8 Å).

To test whether the γ GC-binding mode in the manganese complex properly reflects substrate binding by the iron containing and therefore active form of EgtB_{thermo} we probed γ GC binding by site directed mutagenesis. Specifically, we produced an EgtB_{thermo} variant with an Asp416 to Asn mutation (EgtB_{D416N}). This mutation increases $K_{M, \gamma GC}$ by 200-fold but does not significantly change $K_{M, TMH}$ or k_{cat} . In a complementary experiment we assayed wild type EgtB_{thermo} and EgtB_{D416N} with a γ GC derivative which lacks the α -amino function (N-glutaryl cysteine, NGC). NGC is a 100-fold less efficient sulfur donor for wild type but a 10-fold better substrate for EgtB_{D416N} than γ GC. Evidently, the salt bridge between the α -amino group of γ GC and Asp416 found in the manganese containing quaternary complex of EgtB_{thermo} is also important for substrate recognition during catalysis. The sixth metal ligand in the quaternary complex is a water molecule (Fe-O: 2.5 Å) which also hydrogen bonds to the phenolic side chain of tyrosine 377 (2.8 Å). In addition, this

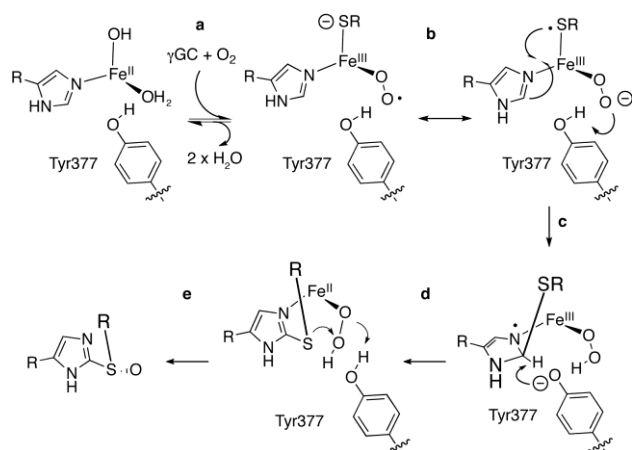


Figure 4. Proposed mechanism for EgtB catalyzed C-S bond formation and sulfoxidation.

coordination site maps to the mouth of a narrow tunnel connecting the active site with the protein exterior (Figure S8). It is tempting to view this tunnel as the path through which O_2 accesses the active site.

Based on the observation that γ GC, TMH and possibly O_2 are direct ligands to the catalytic iron center we propose the following mechanism for EgtB catalyzed sulfoxide synthesis (Figure 4): Binding of γ GC and O_2 to the EgtB complex with TMH results in an iron (III) superoxo species (a). This species may have partial character of iron (III) in complex with a peroxide anion (O_2^{2-}) and a thyl radical (b). The peroxide anion is stabilized by proton transfer from Tyr377 and the thyl radical attacks the imidazole ring to form an iminyl radical (c) which re-aromatizes by deprotonation and ligand-to-metal electron transfer (d). Sulfoxidation of the new thioether reduces the iron (II) peroxo species to return EgtB to its resting state.

Conclusions. EgtB represents a novel type of non-heme iron enzyme which catalyzes oxidative C-S bond formation and sulfoxidation. The crystal structure of EgtB provides the basis for testing mechanistic proposals by kinetic, spectroscopic and computational methods,^[13] and also opens the door for protein engineering to generate tailor made sulfur transferases. Our demonstration that a single mutation (Asp416) suffices to change the sulfur donor specificity of EgtB^{thermo} from γ GC to NGC by almost three orders of magnitude bodes well for this objective. We anticipate that the structure of EgtB will also stimulate mechanistic investigations on the ovothiol biosynthetic sulfoxide synthase OvoA.^[4]

Acknowledgements

The authors would like to thank the Swiss Light Source (PSI, Villigen) and BESSY II Synchrotron (Berlin, Germany) for beamline access; the European research council and the HZI

Graduate School for Infection Research for financial support. F.P.S. is supported by the "Professur für Molekulare Bionik" and by an ERC starting grant.

Keywords: ergothioneine • ovothiol • non-heme iron enzyme • catalysis • sulfur transfer

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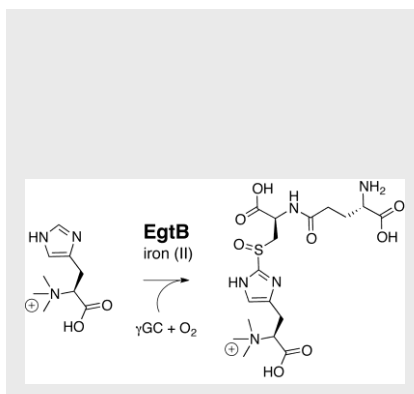
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