

REVIEW

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Unusual cysteine modifications in natural product biosynthesis

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L-Cysteine is a highly reactive amino acid that is modified into a variety of chemical structures, including cysteine sulfinic acid in human metabolic pathways, and sulfur-containing scaffolds of amino acids, alkaloids, and peptides in natural product biosynthesis. Among the modification enzymes responsible for these cysteine-derived compounds, metalloenzymes constitute an important family of enzymes that catalyze a wide variety of reactions. Therefore, understanding their reaction mechanisms is important for the biosynthetic production of cysteine-derived natural products. This review mainly summarizes recent mechanistic investigations of metalloenzymes, with a particular focus on recently discovered mononuclear non-heme iron (NHI) enzymes, dinuclear NHI enzymes, and radical-SAM enzymes involved in unusual cysteine modifications in natural product biosynthesis.

Introduction

L-Cysteine (L-Cys, **1**) is a highly reactive amino acid that can be transformed into a variety of chemical structures because it contains sulfur, which can adopt a wide range of oxidation states, ranging from -2 to $+6$, with -2 (H_2S , thiol, sulfide, and sulfonium ion), -1 (disulfide), 0 (elemental, sulfoxide, and sulfenic acid), $+2$ (sulfone and sulfinic acid), $+4$ (SO_2 , sulfonic

acid, and sulfite ester), and $+6$ (SO_3 and sulfate ester).¹ Protein redox signaling studies revealed that the oxidation of L-Cys in proteins results in the generation of a range of sulfur-containing scaffolds, including thiyl radical, sulfenic acid, sulfonic acid, sulfenyl amide, sulfinamide, sulfonamide, S-nitrosothiol, disulfide, S-glutathione adduct, S-sulfhydryl persulfide, thiosulfinate, and thiosulfate.² In the biosynthesis of natural products, including nonproteinogenic amino acids, alkaloids, and peptides, the L-Cys modifications also lead to various unique chemical structures, such as 2-sulfamoylacetic aldehyde in the sulfonamide alkaloid altemicidin/SB-203208 biosynthesis,³ thiohydroximate in the copper chelating agent fluopsin C biosynthesis,⁴ nor-Cys in the nonproteinogenic amino acid 3-thiaglutamate (3-thiaGlu) biosynthesis,⁵ and

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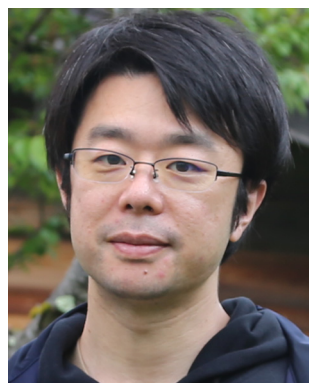
enethiol in lantibiotic peptide biosynthesis.^{6,7} Therefore, understanding the catalytic mechanisms of such Cys-modifying enzymes is a prerequisite for expanding the structural complexity of natural products. In particular, metal-dependent enzymes are versatile catalysts that facilitate dynamic scaffold transformations, which would be challenging with purely chemical catalysts.⁸ These enzymes utilize metallocofactors that mediate electron transfers, enabling reactions with oxygen and the formation of organic radicals or anions which can react with **1** or **1**-derived scaffolds, leading to various oxidative transformations.

This review mainly focuses on the recent mechanistic investigations of three groups of Cys-modifying metalloenzymes in natural product biosynthesis. The first group comprises mononuclear nonheme iron (NHI) enzymes, which belong to the cupin enzyme superfamily and utilize a mononuclear metal ion to activate and transfer electrons to molecular oxygen, as exemplified by isopenicillin N synthase (IPNS)⁹ and cysteine dioxygenase (CDO).^{10,11} Recent studies of CDO and its novel homologs, including SbzM in alkaloid altemicidin/SB-203208 biosynthesis³ and OvoA/EgtB in mercaptohistidine ovothiol biosynthesis,^{12,13} will be described. The second group comprises dinuclear NHI enzymes, which utilize two iron atoms to activate molecular oxygen and generate diiron-peroxo

intermediates.¹⁴ This group includes heme oxygenase diiron-like (HOD) enzymes such as FlcD and FlcE,¹² and DUF692-type enzymes such as MbnBC in CuMbn biosynthesis,¹⁵ TglHI in 3-thiaGlu biosynthesis,⁵ TmoHI in 3-thia- α -amino acid biosynthesis,¹⁶ and ChrHI in the ribosomally-synthesized and post-translationally modified peptide (RiPP) ChrA* biosynthesis.¹⁷ The third group consists of radical-SAM enzymes that utilize a [4Fe-4S] cluster and SAM to generate a reactive 5'-deoxyadenosyl radical (5'-dA[•]) that abstracts a hydrogen atom from substrates,¹⁸ including TmoD in 3-thia- α -amino acid biosynthesis¹⁶ and SycC in AviCamCys biosynthesis.⁷ This review summarizes recent mechanistic investigations of the striking variety of metalloenzyme-catalyzed reactions that enhance the complexity of Cys-derived natural products, highlighting potential applications for the creation of new molecules and supranatural artificial biocatalysts.

1. Cysteine dioxygenase (CDO)

CDO is a well-studied mononuclear NHI enzyme that plays an important role in human L-Cys (**1**) metabolism. The product of CDO, L-cysteine sulfinic acid (**2**), is catabolized by an



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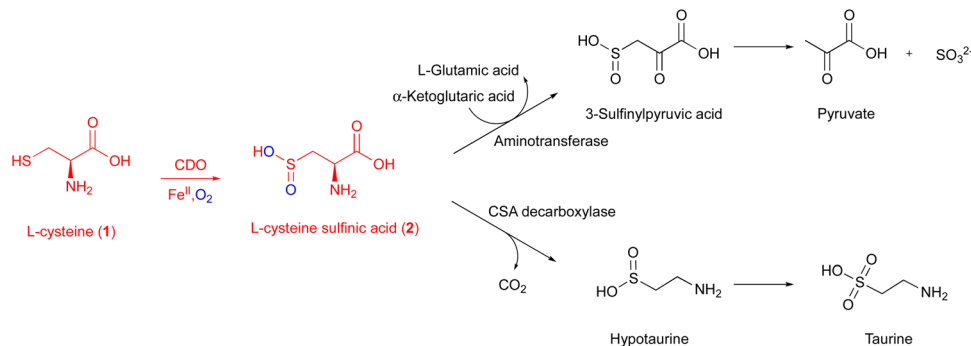


Fig. 1 Metabolic pathway of L-cysteine sulfinic acid (2) generated from L-Cys (1) by CDO.

aminotransferase or cysteine sulfinate decarboxylase in different pathways and transformed into taurine or pyruvate (Fig. 1).¹⁹

The first CDO (from mouse) crystal structure was determined at a resolution of 1.75 Å in 2006.²⁰ The protein core consists of a conserved overall β-barrel-like structure, which is characteristic of cupin superfamily proteins.¹⁰ In the crystal structure of rat CDO, the ferrous iron is coordinated by the three His facial triad motif (His86, His88, and His140), and additionally coordinated by one water molecule (Fig. 2(A)).^{21,22} In the crystal structure of human CDO solved in 2007, 1 was found in the substrate binding site.²³ In addition, 1 was coordinated to the ferrous iron *via* an amino nitrogen and a thiol sulfur, and supported by several hydrogen bonds with surrounding residues (Fig. 2(B)).

A covalent linkage between Cys93 and Tyr157 near the metal binding site was found as a post-translational modification in human CDO (Fig. 2(B)).^{21–23} The covalent linkage between the corresponding Cys and Tyr residues is conserved in eukaryotic CDOs, whereas in bacteria, Cys93 is substituted by glycine, which precludes the formation of the cross-link in BsCDO (CDO from *B. subtilis*, Fig. 2(C)).^{24,25} The subsequent mutation study suggested that the covalent linkage enhances the activity by alleviating the inhibitory effect of Cys93 in human CDO. In addition to the Tyr–Cys linkage, other examples of naturally cross-linked Cys residues in enzymes have been reported. Wensien and his co-workers recently discovered a special lysine–cysteine redox switch with an N–O–S bridge in the transaldolase from *Neisseria gonorrhoeae*. The N–O–S bridge

regulates the enzyme activity through a loaded-spring mechanism, changing the conformations of several key residues.²⁶ A more recent study showed that Orf1, which is involved in the biosynthesis of BD-12, contains an N–O–S bridge that plays an important role in catalysis.²⁷ Orf1 is a noncanonical FAD-dependent enzyme responsible for appending a glycine-derived *N*-formimidoyl group to glycinothricin, producing the antibiotic BD-12. The X-ray crystallographic analysis revealed that Orf1 forms an oximinoglycine with an N–O–S-Cys281 substrate–enzyme bridge, which mediates the nucleophilic addition of glycinothricin and decarboxylation, leading to the cleavage of the N–O bond and the production of a formimidoylated final product.²⁷

Several mechanisms for the CDO reaction have been proposed, based on its X-ray crystal structures solved in early studies.^{20,21,23} Recently, to obtain experimental evidence for the catalytic mechanism, researchers have conducted experiments to mimic or capture potential intermediates during the CDO reaction. Nitric oxide (NO) was utilized as an O₂ analog to investigate the binding order of substrates 1 and O₂ through electron paramagnetic resonance (EPR) spectroscopy. In the absence of 1, CDO does not interact with NO, but in the presence of 1, CDO interacts with NO to generate a low-spin [FeNO]⁷ (*S* = 1/2) signal,²⁸ suggesting that 1 binds to CDO prior to molecular oxygen. Substitutions of 1 with analogs, including cysteamine, 3-mercaptopropionic acid, and propane thiol, abolished this signal, indicating the strict substrate specificity of the CDO reaction. This [FeNO]⁷ (*S* = 1/2) signal was assumed to

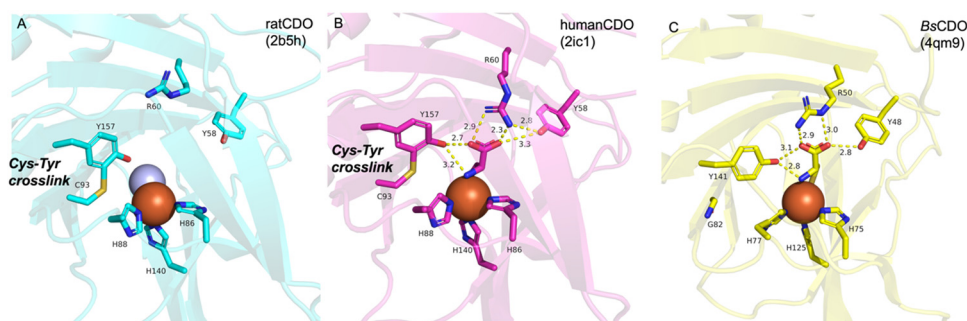


Fig. 2 Crystal structures of CDOs. (A) Metal binding site in rat CDO. (B) Substrate and metal binding site and the Tyr–Cys cross-link in human CDO. (C) Substrate and metal binding site in BsCDO. Fe(II) is colored brown, and water is colored silver. Hydrogen bonds are shown with yellow dotted lines.

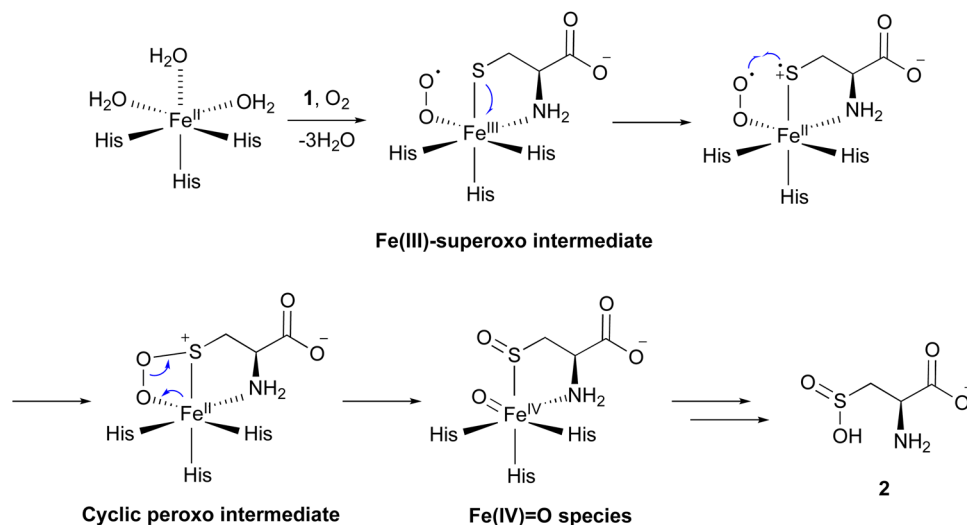


Fig. 3 Proposed reaction mechanisms of CDO.

originate from the bidentate thiol/amine coordination of **1** to the iron-nitrosyl complex,²⁸ suggesting the presence of an Fe(III)-superoxo intermediate (Fig. 3). In addition, azide was employed to mimic superoxide. As suggested by EPR and DFT calculations, azide directly binds to Fe(III), but not to Fe(II). In the DFT-optimized structure of this complex, the central nitrogen atom of the azide moiety is 3.12 Å away from the sulfur of **1**, supporting the attack from the superoxide group of the putative Fe(III)-superoxo intermediate.²⁹

Fe(III)-bound CDO was mixed with superoxide anions generated by xanthine oxidase, in an attempt to detect the characteristic signals of the Fe(III)-superoxo complex. The addition of superoxide anions generated a transient species observed at 565 nm by UV-vis spectroscopy. The decay of this 565 nm absorption kinetically matched the formation of **2**. Further analyses by EPR spectroscopy suggested that this transient species is an Fe(III)-superoxo intermediate.³⁰ These data provided additional evidence for the presence of the Fe(III)-superoxo intermediate.

By mixing CDO:Fe(II):**1** with oxygenated buffer at a low temperature, characteristic signals for intermediates were captured through UV-vis and ⁵⁷Fe-Mössbauer spectrum analyses. Transient absorption peaks at 500–640 nm formed and disappeared within 20 ms after mixing. This transient intermediate was further captured by a characteristic signal in the ⁵⁷Fe-Mössbauer spectrum,³¹ and the DFT calculation suggested that this signal corresponds to the cyclic peroxo intermediate in Fig. 3.

Iron(II)-thiolate models were adopted for understanding the reaction mechanism of CDO. In the CDO mimic models, oxygen can only be activated by specific thiolate ligands, suggesting that they play a critical role. Further calculation results suggested that the thiolate ligand must transfer a charge to Fe(II), resulting in a lower redox potential, which is crucial for the reaction of oxygen with iron.³² Sulfur oxidation was also investigated in an [Fe(III)(LN₃S)]⁺ model, in which LN₃S represents a tetradentate ligand with a bis(imino)pyridyl

scaffold and a pendant arylthiolate group. Furthermore, additional calculations suggested that oxygen preferentially binds to the iron center, instead of attacking the sulfur group directly. Together, these data suggested that the formation of the Fe(III)-superoxo complex initiates the catalysis, followed by the step-wise oxidation of the sulfur group.³³

Consequently, the mechanism of the CDO reaction has been proposed as described below (Fig. 3). The molecular oxygen binds to Fe(II), leading to the formation of the Fe(III)-superoxo intermediate. The distal oxygen of the superoxo complex reacts with activated sulfur, generating a cyclic peroxo intermediate. Heterolytic O–O bond cleavage generates a sulfoxide and Fe(IV)=O species. The Fe(IV)=O species reacts with the sulfoxide, and **2** is released from the enzyme in the final step of the catalysis. Another mechanism was also considered, involving a persulfenate intermediate detected in the X-ray crystal structure analysis.³⁴ However, this pathway exhibited a high energy barrier in the DFT calculations, suggesting that the formation of such an intermediate is unlikely in the catalytic reaction of CDO.

1.1. Unusual CDO homologs

1.1.1 SbzM. Altemicidin (**3**), isolated from *Streptomyces sioyaensis* SA-1758, is a sulfonamide antibiotic with promising antitumor activity.^{35,36} The *sbz* gene cluster was shown to be responsible for the biosynthesis of altemicidin and its analog SB-2032308 (**4**).^{3,37} An isotope feeding study and *in vitro* assay revealed that 2-sulfamoylactic acid (**5**) was generated from **1** by SbzM and SbzJ.³ First, **1** is oxidized by the CDO homolog SbzM to 2-sulfamoylactic aldehyde (**6**), which is subsequently transformed into **5** by the aldehyde dehydrogenase SbzJ. **5** is further modified to 2-sulfamoylacyl-AMP by the AMP-ligase SbzL. A 2-sulfamoylacyl-group, loaded on the acyl carrier protein SbzM (7), is subsequently installed onto the 6-azatetrahydroindane intermediate, generating **8** by the GCN5-related *N*-acetyltransferase SbzM. Eventually, **8** is transformed into **3** and **4** (Fig. 4(A)).



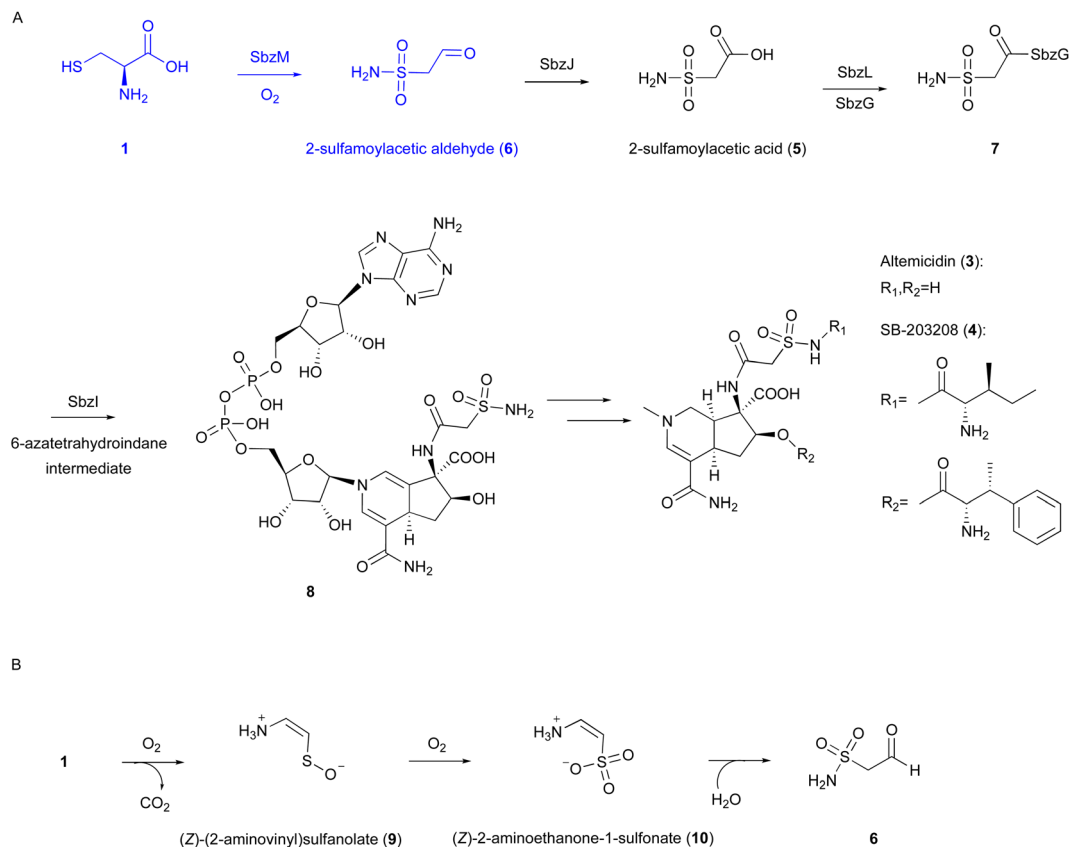


Fig. 4 Reaction and proposed mechanism of SbzM. (A) Biosynthetic pathway of altamycin and analogs. (B) Proposed reaction of SbzM.

Our recent investigations on the metal dependency of the CDO homolog enzyme SbzM suggest that nickel and iron are potential active metals.³ A plausible mechanism was proposed for the SbzM reaction (Fig. 4(B)),³ in which **1** is transformed into (Z)-(2-aminovinyl)sulfanolate (**9**) through sulfur monooxygenation followed by decarboxylation, and then the sulfanolate is further oxidized to a second intermediate, (Z)-2-aminoethanone-1-sulfonate (**10**). The subsequent intramolecular rearrangement of the amino group on the sulfur atom, accompanied by the attack of water, results in the formation of the N-S bond in the final product **6**. An alternative mechanism involving an Fe(IV)=O complex,³⁸ as in the cases of CDO and IPNS,^{9,39,40} is also possible, in which a mercapto group is oxidized by the superoxo complex. However, since the activity can be restored with the addition of Ni(II), and considering that nickel cannot readily form the Ni(IV)=O intermediate, this hypothesis would be excluded. Detailed structural and mechanistic investigations are still in progress.

1.1.2 OvoA and EgtB. Ovothiol (**11**) is a 4-mercapto-1-methyl-L-histidine that was first isolated from marine invertebrates, and is biosynthesized from **1** and L-histidine (**12**) (Fig. 5(A)).^{41–43} The mercapto group attached to the imidazole ring of **12** is highly acidic (pK_a = 1.4) and easily oxidised,⁴⁴ imparting **11** with robust reducing ability to effectively scavenge free radicals and superoxide ions.^{45,46}

OvoA catalyzes the oxidative coupling between **1** and **12** to generate the 5-histidylcysteine sulfoxide conjugate (**13**), which

is the precursor of the final product **11** (Fig. 5(A)).¹² OvoA is an NHI enzyme functionally similar to CDO, with the iron coordinated by the 2xHis–1xGlu triad motif within a conserved HX₃HXE sequence in the N-terminal domain. Besides the main reaction, OvoA also catalyzes a CDO-type reaction to produce **2**, as a minor reaction (Fig. 5(B)).⁴⁷ OvoA generates L-cystine (**14**) when the substrate is only **1**, and produces **13** as the main product along with **2** as a co-product when the substrates are both **1** and **12** (Fig. 5(C)).⁴⁷ A study using an OvoA homolog from *Methyloversatilis thermotolerans* (OvoA_{met}) showed that only **1**, but not **12**, is required for CDO activity.⁴⁸

Structure–function analyses of the homology model of OvoA and the functionally similar enzyme EgtB showed that Tyr417 of OvoA corresponds to Tyr377 of EgtB. By substituting Tyr417 with an unnatural amino acid, 2-amino-3-(4-hydroxy-3-(methylthio)phenyl) propanoic acid (MtTyr), the CDO-type activity was considerably enhanced.⁴⁹ These data suggested that Tyr417 plays an important role in bifurcating the two pathways, oxidative C–S bond formation and CDO-type oxidation. The crystal structure of OvoA_{Th2} (OvoA homolog from *H. thermophila*) in complex with two substrates, **12** and **1**, was recently reported (PDB ID: 8KHQ). This structure provided critical information to explain the regioselectivity of C–S bond formation.⁵⁰ In the substrate binding pocket, Co(II) is coordinated by three His residues (His71, His162, and His166), with **12** coordinating to Co(II) from the opposite side of His71 and **1**

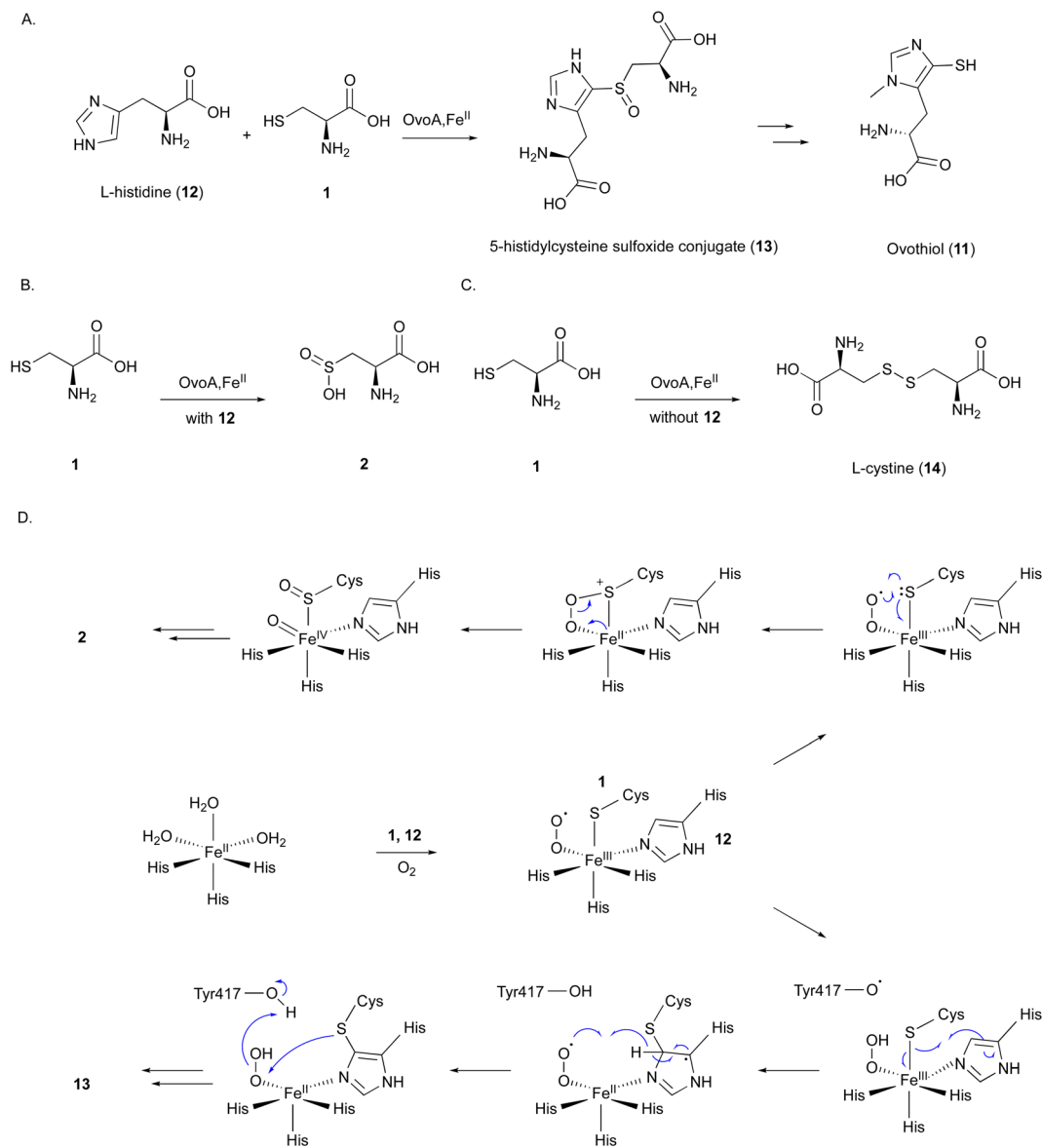


Fig. 5 Reaction and mechanism of OvoA. (A) C-S bond formation between **1** and L-histidine (**12**) catalyzed by OvoA. (B) Formation of **2** catalyzed by OvoA in the presence of **1** and **12**. (C) L-Cystine (**14**) formation by OvoA in the presence of **1**. (D) Proposed catalytic mechanism of OvoA to produce **2** or **13**.

coordinating from the opposite side of His162 (Fig. 6(A)). The possible molecular oxygen binding site, situated opposite to His166, is occupied by a water molecule, and the conserved Tyr406 forms a hydrogen bond with it. A hydrogen bond exists between the amino group of **1** and the carboxyl group of **12**, bringing them closer together.

A catalytic mechanism for OvoA has been proposed that explains the generation of both **13** and **2** with Tyr417 (Fig. 5(D)).⁴⁹ A ternary Fe(III)-superoxo radical complex is formed, and a hydrogen is transferred from Tyr417 to the superoxo species, resulting in a peroxo species. The radical on the sulfur, formed by single electron transfer to Fe(III), reacts with the imidazole to form a C-S bond between **1** and **12**. Dehydrogenation from the peroxo species by Tyr417 and

oxidation by the generated superoxo species lead to the production of the product **13**. In the Y417MtTyr variant, the CDO-type reaction occurs, producing **2**. An Fe(IV)=O intermediate in the proposed mechanism was observed in OvoA_{met} through stopped-flow, ⁵⁷Fe-Mössbauer spectra, and X-ray absorption near edge structure (XANES) analyses.⁵¹

In ergothioneine biosynthesis, the NHI enzyme EgtB, which is functionally related to OvoA, catalyzes the O₂-dependent C-S bond formation between γ -glutamyl cysteine (γ -GC, **15**) and *N*- α -trimethyl histidine (TMH, **16**), producing **17** (Fig. 7(A)).¹³ To investigate the structural basis of EgtB catalysis, the crystal structure complexed with **15** and **16** was solved.⁵² Tyr377, located close to the binding site of **16**, is assumed to be the key residue for oxygen binding (Fig. 6(B)). Interestingly, EgtB

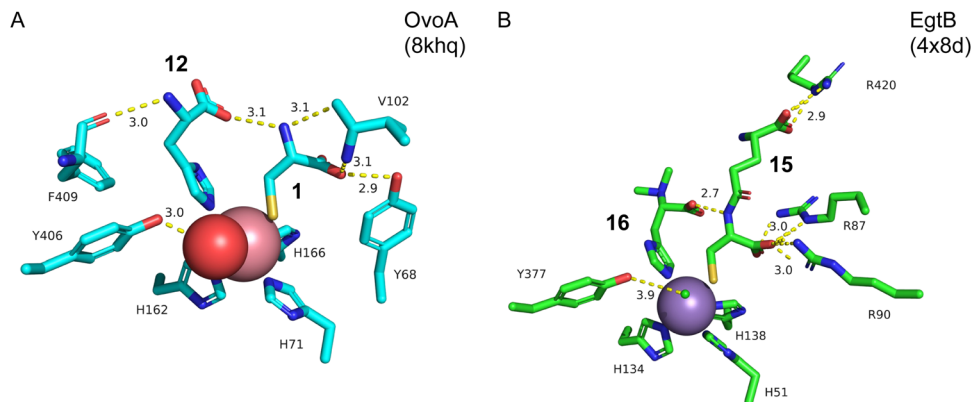


Fig. 6 Crystal structures of OvoA_{Th2} (homolog of OvoA) and EgtB. (A) The active site complexed with substrates **1** and **12** in OvoA_{Th2}. Tyr406 corresponds to Tyr417 in OvoA. (B) The active site complexed with substrates **15** and **16** in EgtB. Hydrogen bonds are shown as yellow dotted lines, the manganese ion is colored silver, and the chloride ion is shown as a small green ball.

Y377F oxidized the thiol of **15**, similar to CDO (Fig. 7(A)), indicating that the hydroxyl group of Tyr377 determines the fate of the reaction.⁵³

Based on the crystal structure and the mutation experiments, the reaction mechanism of EgtB has been elucidated (Fig. 7(B)). The Fe(III)–superoxo species is transformed to Fe(III)–hydroperoxo species through a hydrogen radical transfer from Tyr377. The C–S bond is formed by the attack of the sulfur of **15** on C-2 of the imidazole of **16**, followed by the removal of H-2 of the imidazole by Tyr377. Finally, the sulfur group undergoes sulfoxidation to generate the final product **17**. As observed in the case of Y377F, without the hydrogen radical transfer from the hydroxyl group of Tyr377, the Fe(III)–superoxo species will directly attack the thiol group of **15**, generating a thiol-dioxygenated product *via* a CDO-like mechanism.⁵³

2. FlcD and FlcE in fluopsin C biosynthesis

Fluopsin C (**18**) (Fig. 8) was first isolated in 1970 from environmental *Pseudomonas aeruginosa* and *P. fluorescens*, and also in 1972 from a *Streptomyces* strain,^{54,55} as a copper-containing, dark green complex with specific d–d transition UV-vis absorption for copper(II) in the range of 550–600 nm. **18** exhibits antibiotic activity against a broad spectrum of bacteria and fungi, and even inhibits multidrug-resistant bacterial pathogens, such as *Acinetobacter baumannii* and *Staphylococcus aureus*. However, the producer, *P. aeruginosa*, is resistant to **18**, consistent with the ability of *P. aeruginosa* to tolerate high concentrations of copper.⁵⁶

The author focused on the gene cluster *flc* (PA3515–PA3523), which is upregulated under copper stress, as a putative biosynthetic cluster for **18**.^{57,73} The *flc* operon encodes five proteins, including the methyltransferase FlcA, two adenylosuccinate lyases (ASLs)⁵⁸ FlcB and FlcC, and two HOD enzymes FlcD and FlcE, together with the copper chaperone CopZ1 and the efflux pumps PA3521–PA3523. Feeding experiments, hetero-

logous expression in *E. coli*, and *in vitro* assays were employed to characterize each of the Flc biosynthetic reactions (Fig. 8). As the initial step, the ASL FlcB catalyzes the conjugation of **1** and fumarate to produce **19**.

The HOD enzyme FlcE catalyzes both the oxidative decarboxylation and *N*-hydroxylation of **19** to produce **20** (Fig. 8). FlcE requires only Fe(II) for its activity, and does not require reductants, similar to other HOD enzymes, including *Chlamydia* protein associating with death domains (CADD) in folate biosynthesis and SznF in streptozotocin biosynthesis.^{59,60} Next, **20** is oxidized to **21** by the other HOD enzyme FlcD through a four-electron oxidation process, with one carbon being released as formic acid. An assay using L-[3-¹³C]-**1** identified the lost carbon position in the reaction as C-2. In addition, like FlcE, FlcD does not require a reducing agent for the reaction.

The proposed carbon excision reaction by FlcD is given below (Fig. 9). FlcD's diiron center first binds to molecular oxygen, forming μ -peroxo-Fe₂(III/III) (Fig. 9(i)). This μ -peroxo-Fe₂(III/III) is then converted into the Fe₂(IV/IV) complex (Fig. 9(ii)). Subsequently, Fe₂(IV/IV) abstracts a hydrogen radical from the *N*-hydroxyl group (Fig. 9(iii)). The generated *N*-hydroxyl radical facilitates isomerization and hydroxylation at C- α (Fig. 9(iv)). After hydrogen atom transfer from CH₂- β , transient C–N cleavage forms the NO radical (Fig. 9(v)), and further a 1,2-shift rearrangement constructs the C–N bond at C- β (Fig. 9(vi)), as proposed in the UndA reaction in undecene biosynthesis.^{61,62} A second hydroxylation at C- α regenerates the Fe₂(II/II) center (Fig. 9(vii)). Finally, the generated *gem*-diol is cleaved to form formate and **21** (Fig. 9(viii)). The proposed sequential radical reaction mechanism, leading to carbon excision without any reducing agent, is intriguing. In the structural study of other di-iron oxygenases, such as myo-inositol oxygenase (MIOX),⁶³ its di-iron center was shown to be coordinated by four His and two Asp. The substrate myo-inositol is bound to one of the iron atoms in bidentate coordination *via* OH-1 and OH-6. Structural analysis of FlcD will provide a structural basis for this interesting reaction, as exemplified by the study for MIOX.

For the remaining two enzymes, the lyase FlcC converts **21** to the thiohydroxamate **22**, and the SAM-dependent methyltransferase



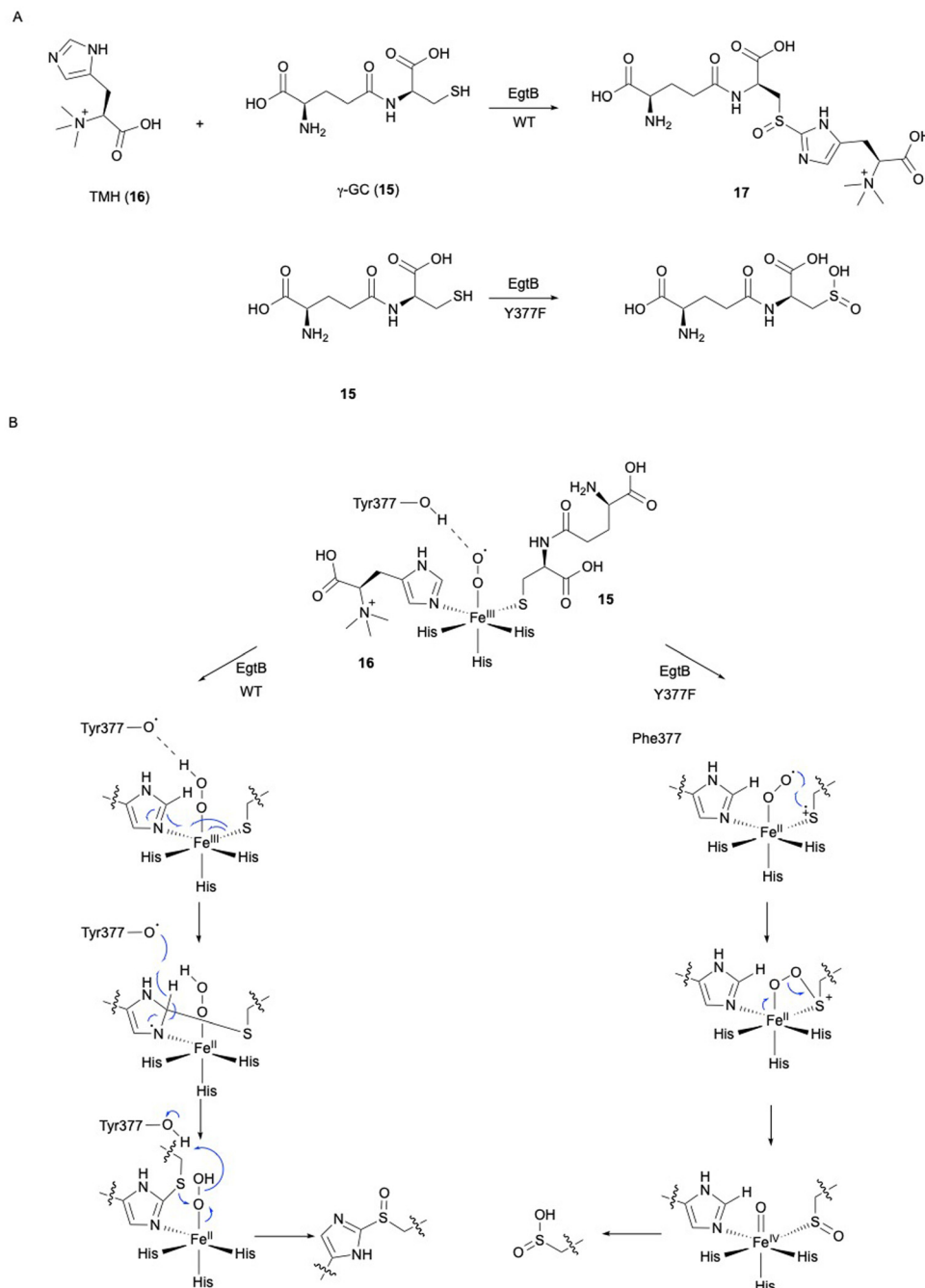


Fig. 7 Reaction and mechanism of EgtB. (A) C–S bond formation between **15** and **16** by EgtB WT, and thiol oxidation of **15** by the EgtB Y377F variant. (B) Proposed mechanism of EgtB WT and the Y377F variant.

FlcA methylates **22** to form Cu-free fluopsin. Fluopsin chelates a Cu(II) ion to yield the final product, fluopsin C (**18**) (Fig. 8).

3. MbnBC in CuMbn biosynthesis

The copper-binding peptidic methanobactins (Mbns) chelate Cu(I) to form Cu(I)Mbn (**23**) as a siderophore,^{80,81} which was originally found in obligate methanotrophs⁶⁴ under copper-limiting conditions (Fig. 10). Mbns are synthesized by

ribosomes, and classified as RiPPs.^{70–72} During the biosynthesis of RiPPs, a cysteine residue in the peptide is typically cyclized to a thiazoline and further oxidized to a thiazole.⁶⁵ In addition to the well-known thiazoline oxidation, L-Cys-related biosynthesis in RiPPs involves several interesting oxidation reactions, as described below.

The *mbn* operon was first identified as regulated in response to copper in the methanotroph *Methylosinus trichosporium* OB3b.⁶⁶ The uncharacterized proteins MbnBC are encoded in *mbn* operons and are responsible for the post-translational



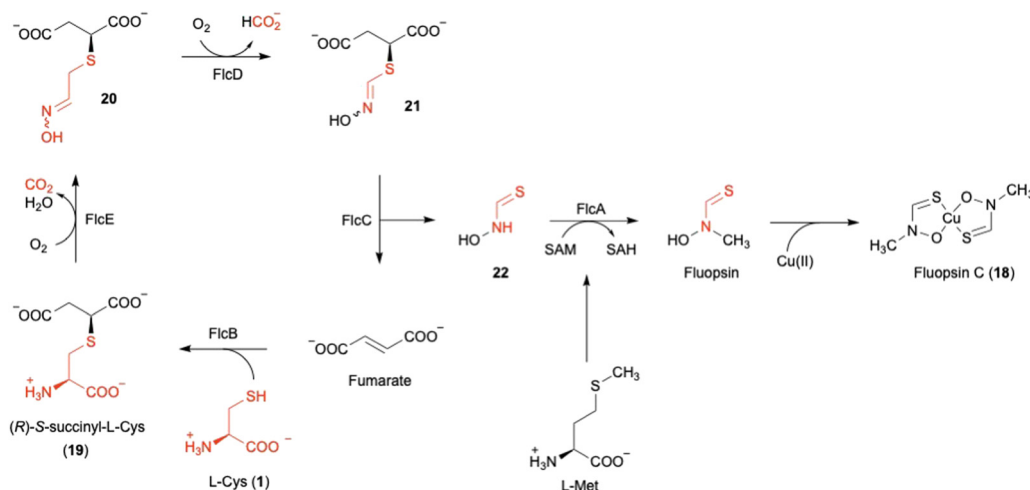


Fig. 8 Biosynthetic pathway of fluopsin C (**18**).

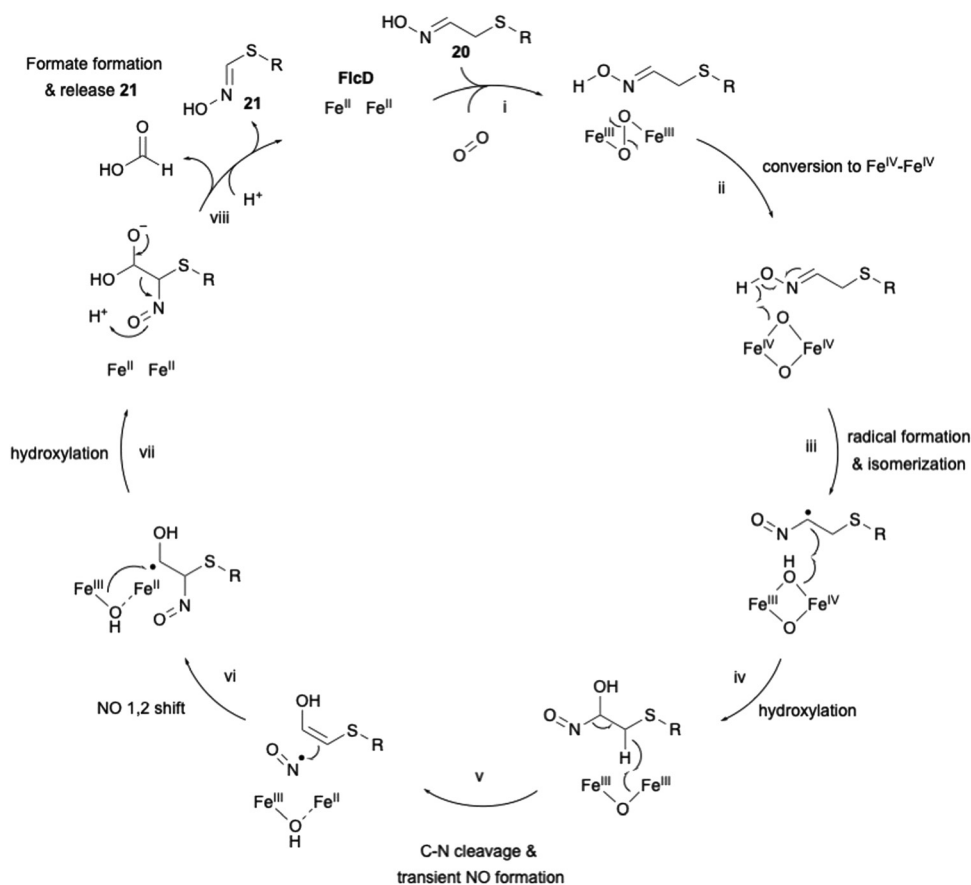


Fig. 9 Proposed mechanism of carbon excision catalyzed by FlcD.

modification of the precursor peptide MbnA. The oxazolone ring and thioamide group in mature Mbns are thought to originate from a Cys residue (Fig. 10),^{67,68} and are transformed into an oxazolone-thioamide moiety through a net four-electron oxidation catalyzed by MbnBC.

MbnB is a member of the DUF692 family and widely distributed in bacteria, but none of the DUF692 family enzymes

has been functionally characterized, although the crystal structure of one (*HsMbnB* from *Haemophilus somnus*, PDB ID: 3BWW) was solved as a TIM barrel fold protein, which has metal-binding residues. MbnC is present exclusively in the *mbn* operons and *Pseudomonas* species, but its function is unknown. Co-expression of MbnBC resulted in soluble proteins, which were purified as a heterodimer.⁵⁷ Fe-Mössbauer spectral analysis



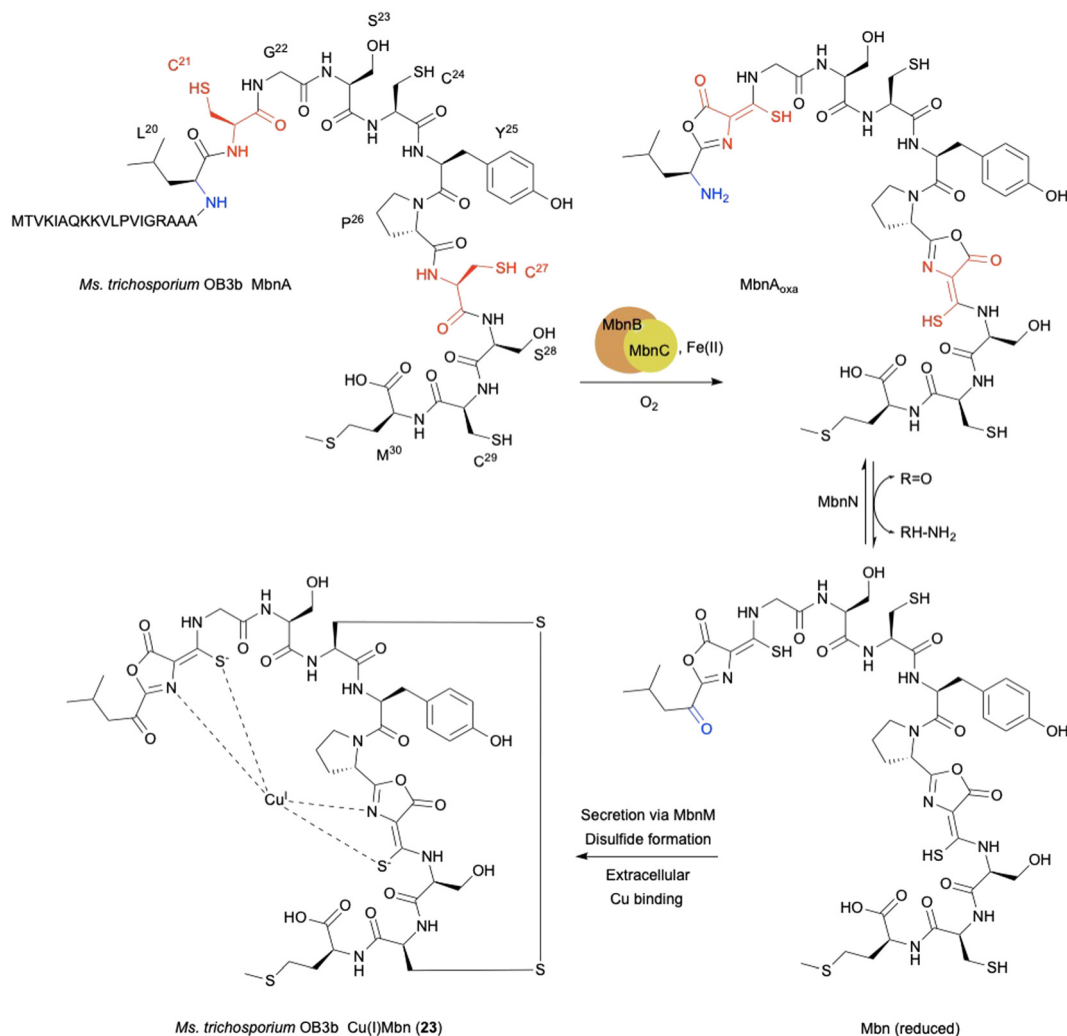


Fig. 10 Biosynthetic pathway of Cu(I)Mbn (23). The scheme includes oxidation catalyzed by MbnBC, transamination at leucine catalyzed by the aminotransferase MbnN, and secretion by the multidrug exporter MbnM.

and native top-down mass spectrometry indicated that MbnBC has three Fe(II) in each subunit.⁶⁹ The LC-MS/MS analysis of the products showed that MbnBC oxidizes Cys21 of MbnA to produce the oxazolone-thioamide moiety of MbnA_{oxa} in the presence of Fe(II) and O₂, and does not require any reducing agents.

The oxidation of MbnA by MbnBC was proposed as described below (Fig. 11). A hydrogen atom is abstracted from C-β of Cys21 by an Fe(III)-superoxo intermediate, forming a radical (i). A thioaldehyde is then formed *via* radical recombination (ii). The thioaldehyde is attacked by an amide nitrogen, aided by the deprotonation by the peroxide coordinated to Fe(III), forming a β-lactam (iii), as seen in the ring formation by IPNS.⁷⁰ Base-mediated deprotonation from the other amide nitrogen forms an amidate (iv). The generated amidate reacts with the β-lactam to produce an oxazolone ring bound to a thioaminal (v).⁷¹ Abstraction of another hydrogen atom by Fe(IV)=O lead to the formation of a C-β radical (vi). A hydroxide bound to Fe(III) then abstracts H-α to construct the thioamide group of MbnA_{oxa} and reduce Fe(III) to Fe(II) (vii). This is the first example of an

oxazolone and thioamide synthase, and an oxidation reaction catalyzed by a DUF692-type NHI enzyme.

A quite recent study presents the X-ray crystal structures of MbnABC complexes, revealing that the leader peptide of MbnA binds to MbnC and facilitating the recruitment of the MbnBC holoenzyme. Meanwhile, the core peptide of MbnA resides in the catalytic cavity formed by the interaction between MbnB and MbnC, which contains a unique tri-iron cluster. The thiol group of the substrate is ligated to this tri-iron center, leading to a dioxygen-dependent reaction for oxazolone-thioamide installation. This study also identified the conserved residue D240 as the base required for reaction step iv in Fig. 11.⁷²

4. TglH in 3-thiaGlu biosynthesis

In the biosynthesis of 3-thiaGlu 24, the second example of a DUF692-type NHI enzyme family member was reported. A complex of the DUF692-type NHI enzyme (TglH) and a leader



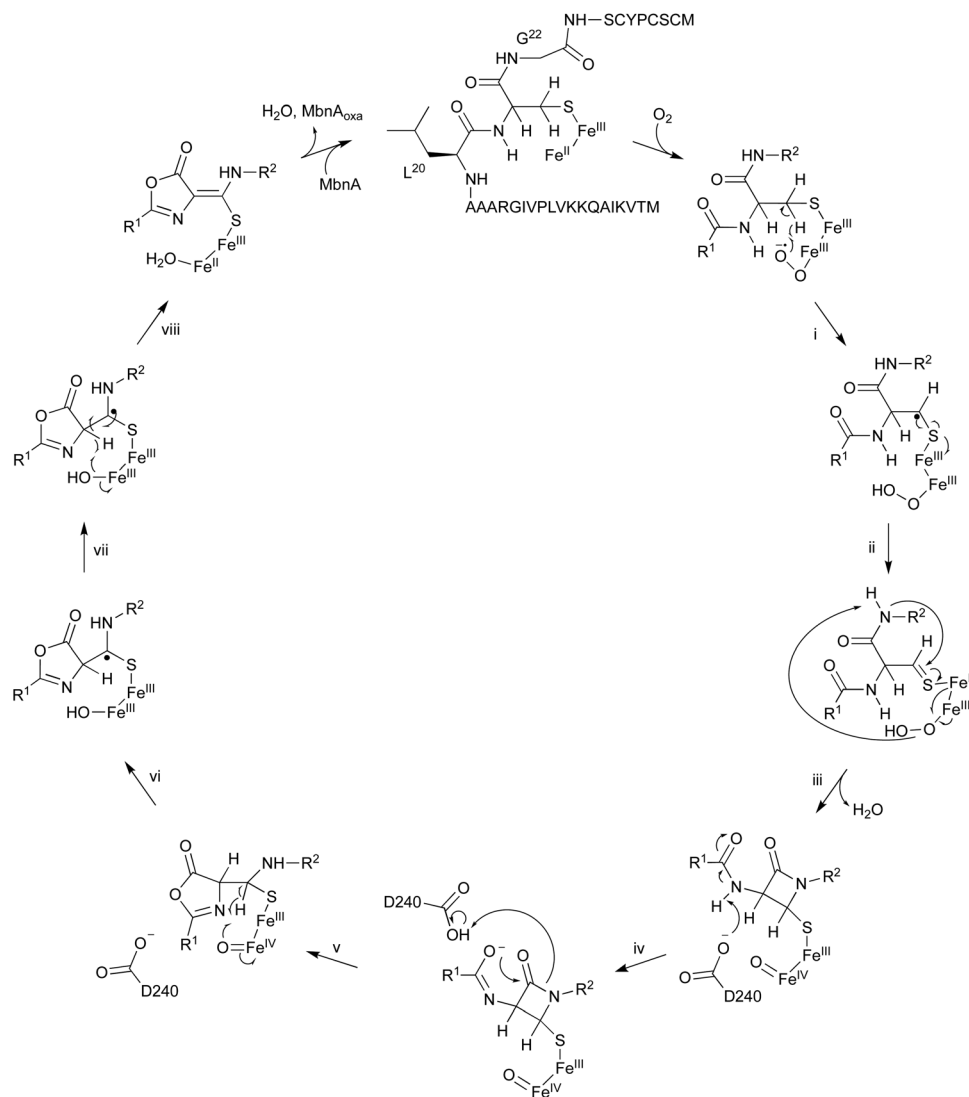


Fig. 11 Proposed mechanism of oxazolone/thioamide formation catalyzed by MbnBC.

peptide-binding protein (TgII) was shown to catalyze the unique C- β excision reaction of a Cys residue in the substrate peptide, producing the nor-cysteine scaffold of TgIA-norCys (25), which is a precursor of 24 (Fig. 12A). TgIHI is encoded in the RiPPs biosynthetic gene cluster (*tgl* cluster) in the plant pathogen *P. syringae* pv. *maculicola* ES432. The *tgl* cluster encodes a leader peptide (TgIA), a peptide-aminoacyl tRNA ligase (PEARL,⁵ TgIB), TgIH, TgII, a carboxy-S-adenosyl-L-methionine (Cx-SAM⁷³) synthase (TgIE), and a Cx-SAM-dependent transferase (TgIF). Each of the biosynthetic reactions catalyzed by Tgl enzymes was characterized through heterologous expression in *E. coli* and *in vitro* assays, as shown in Fig. 12A.⁷³

TgIA-Cys is generated in the reaction catalyzed by the PEARL TgIB, from TgIA, Cys-tRNA, and ATP (Fig. 12B), and oxidized to 25 by TgIHI without any reducing agent. In the TgIHI assay with TgIA-3-¹³C-Cys, ¹³C-labeled formate was detected as a by-product, suggesting that the β -carbon of Cys was removed from the substrate.

The proposed reaction mechanism of the β -carbon scission by TgIHI is described below (Fig. 13). First, the activation of the Fe(II) complex by O_2 lead to the formation of the Fe(III)-superoxo species (Fig. 13(i)). This Fe(III)-superoxo species abstracts the C- β hydrogen to form a radical intermediate (Fig. 13(ii)). The Fe(III)-hydroperoxo species reacts with the C- β radical to generate a thioacetal Fe(IV)=O complex (Fig. 13(iii)), which abstracts a hydrogen from OH- β to form an oxygen radical (Fig. 13(iv)). This leads to the homolysis of C α -C β , generating a C- α radical (Fig. 13(v)), through a mechanism similar to that reported for 2-hydroxyethylphosphonate dioxygenase (HEPD).⁷⁴ The following radical recombination generates a new C-S bond in the *S* configuration as 25, with the reduction of the Fe(III) complex (Fig. 13(vi)), similar to thioether formation by IPNS.³⁹ Finally, the thioformyl moiety is hydrolyzed by a hydroxide bound to Fe(II), producing both formate and 25 (Fig. 13(vii)). The TgIHI-catalyzed methylene excision reaction is a novel type of post-translational modification in a natural product.

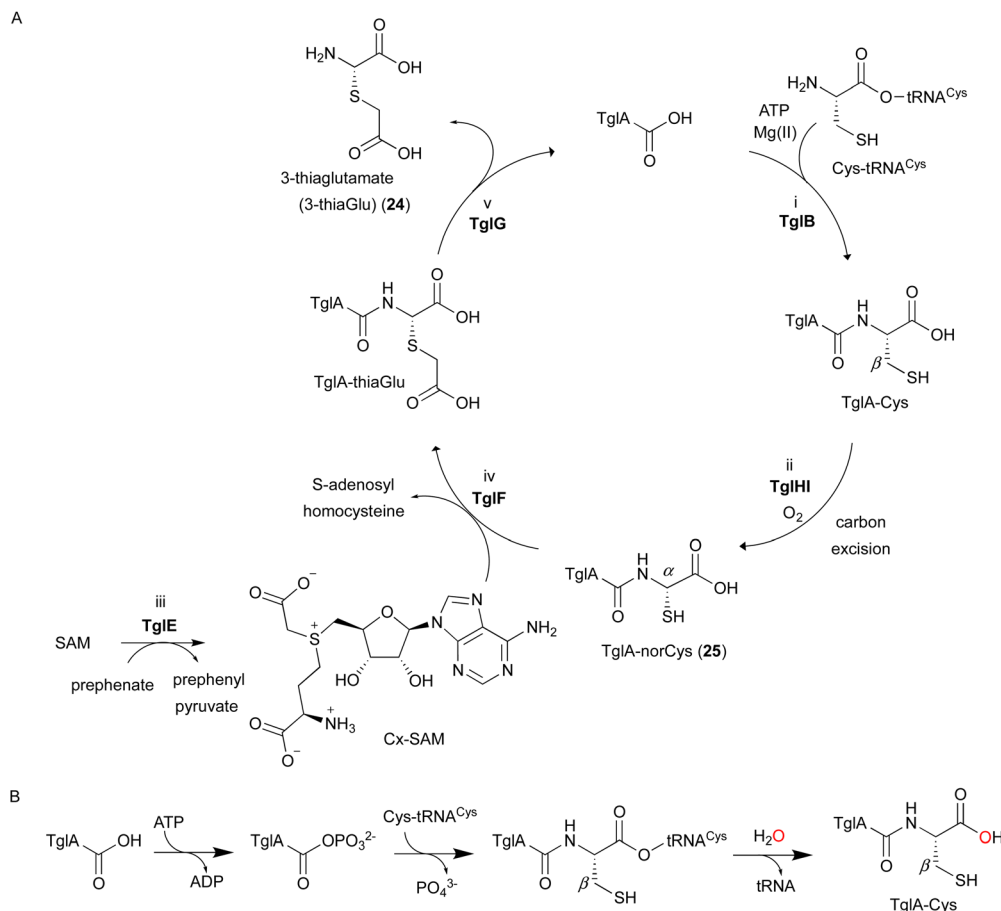


Fig. 12 Biosynthetic pathway of 3-thiaglutarate (**24**). (A) 3-thiaglutarate (**24**) formation catalyzed by Tgl enzymes. (B) Cysteine addition by the PEARL TglB to the leader peptide TglA. The reaction starts with phosphorylation of the C-terminal carboxylate of TglA, which then forms an amide bond with the amino group of Cys-tRNA^{Cys}, followed by the hydrolysis of the tRNA to generate TglA-Cys.⁵

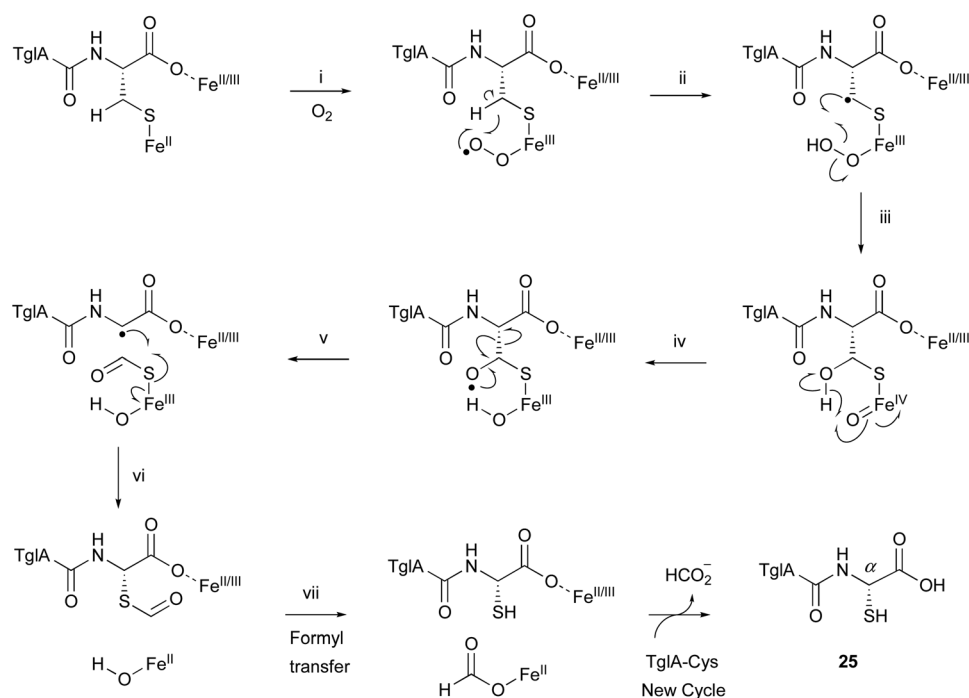
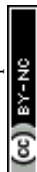


Fig. 13 Proposed mechanism of the TglH reaction.



Very recently, the X-ray crystal structures of TglHI complexed with and without iron ions were reported.⁷⁵ TglH adopts a TIM barrel structure with β -sheets wrapped around helices, forming a stable complex with TglI, in a similar manner to MbnBC.^{72,76} These structures revealed the formation of a catalytic pocket between TglH and TglI, and two iron-iron binding sites in the pocket and a third auxiliary iron binding site were identified. The complex structure model of TglHI and TglA-Cys generated by AlphaFold2⁷⁷ identified the key amino acid Asn73, which could contribute to the H- β abstraction by orienting the thiol of TglA-Cys *via* hydrogen bonding.

After the TglHI reaction, TglF reacts with **25** and Cx-SAM, generating TglA-thiaGlu (L-3-thiaglutamate). Its absolute configuration was confirmed by microcrystal electron diffraction (MicroED) analyses with synthesized diastereoisomers (Fig. 12(A)). TglG, a membrane-bound protease, cleaves the amide bond of TglA-thiaGlu, releasing 3-thiaGlu.

5. TmoHI and TmoD in 3-thia- α -amino acid biosynthesis

In the biosynthesis of 3-thiahomoleucine (**26**), the DUF692 protein (TmoH, 67% sequence identity with TglH) and a vitamin B₁₂-dependent radical-SAM (rSAM) enzyme (TmoD), were characterized as new L-Cys oxidases in *Tistrella mobile*. The biosynthetic cluster (*tmo* cluster) encodes a precursor peptide (TmoA), a PEARL (TmoB), TmoH, a SAM-dependent methyltransferase (TmoS), a RiPPs recognition element-containing protein (TmoI), TmoD, a metalloprotease (TmoG), and a transporter (TmoT). Each of the reactions by Tmo enzymes was identified *in vitro* and in an *E. coli* expression system (Fig. 14).¹⁶

TmoHI cleaves the β -carbon of TmoA-Cys (**27**) to generate nor-Cys (**28**), in a similar manner to TglHI. TmoS catalyzes the S-methylation of **28** to produce **29**. TmoD is classified as a class B vitamin B₁₂-dependent rSAM enzyme. This rSAM enzyme family utilizes SAM, methylcobalamin, and a [4Fe-4S]²⁺ cluster to initiate methyl transfer onto a non-nucleophilic phosphorus or carbon atom. TmoD was anaerobically reconstituted with Na₂S, (NH₄)₂Fe(SO₄)₂, and hydroxocobalamin (HOCbl), enabling the catalysis of the tandem trimethylation of the C-terminal residue of **29** to yield **30** (Fig. 15).

The hypothetical reaction mechanism of TmoD is described below (Fig. 15). HOCbl is first reduced to cobalamin Co(i) (i) and then reacts with SAM to form methylcobalamin Me-Co(iii) (ii). The [4Fe-4S]²⁺ cluster is reduced to [4Fe-4S]⁺ (iii), and generates 5'-dA $\cdot$ from another SAM (iv).⁷⁸ The 5'-dA $\cdot$ abstracts a hydrogen from **29** to produce **29 $\cdot$ (v), which then accepts a methyl group from Me-Co(iii), forming a monomethylated compound **31** (vi). After repeating another two rounds of methylation, the final compound **30** is produced. CysS,⁷⁹ ThnK,⁸⁰ and TokK,⁸¹ which belong to the same group as TmoD in the similarity network of vitamin B₁₂-dependent rSAM enzymes, have similar functions, adding methyl groups iteratively. For example, TokK catalyzes three tandem methylation reactions of the intermediate to generate the isopropyl group of the carbapenem asparenomicin.⁸¹ The trimethylation by TmoD, which construct an isopropyl group on the S-methyl group, expands the reaction diversity of the vitamin B₁₂-dependent rSAM enzymes responsible for peptide modifications.**

Finally, the membrane-bound TmoG cleaves the C-terminal **26** from **30** to regenerate TmoA. The substrate specificity of TmoG was tested with TmoA-Cys, TmoA-Leu, TmoA-Gln, and TmoA-Met, and only TmoA-Met was accepted. This preference is consistent with the recognition of the extended hydrophobic side chain of **30**.

6. ChrHI in RiPP ChrA* biosynthesis

To discover novel DUF692 proteins, 13 108 DUF692 sequences were retrieved from the Sequence Similarity Network (SSN) and the Genomic Neighborhood Network (GNN).¹⁷ As a result, the DUF692 protein ChrH encoded in the gene cluster in *Chryseobacterium* sp. (*chr* cluster) was analyzed, and its product was identified as the peptide ChrA*. Heterologous expression in *E. coli* and *in vitro* assays revealed the details of the ChrHI reaction that converts ChrA to ChrA*, which has one additional sulfide and one additional methyl group, with the loss of a single thiol group compared to its substrate (Fig. 16).

Extensive 1D and 2D NMR analyses, including HSQC, dqCOSY, NOESY, and HMBC, coupled with an HR-MS/MS analysis showed the structural changes on C63 and C66 in ChrA*, produced from *E. coli* expressing ChrA and ChrHI. The reaction of ChrHI with ChrA as the substrate was reconstituted

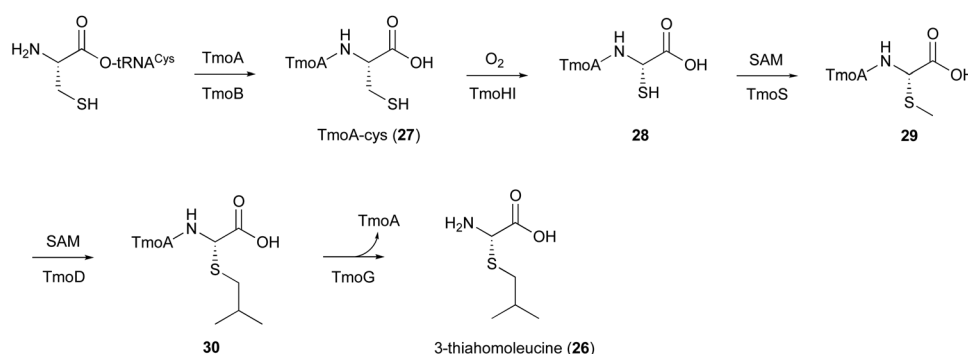


Fig. 14 Biosynthetic pathway of 3-thiahomoleucine (**26**).

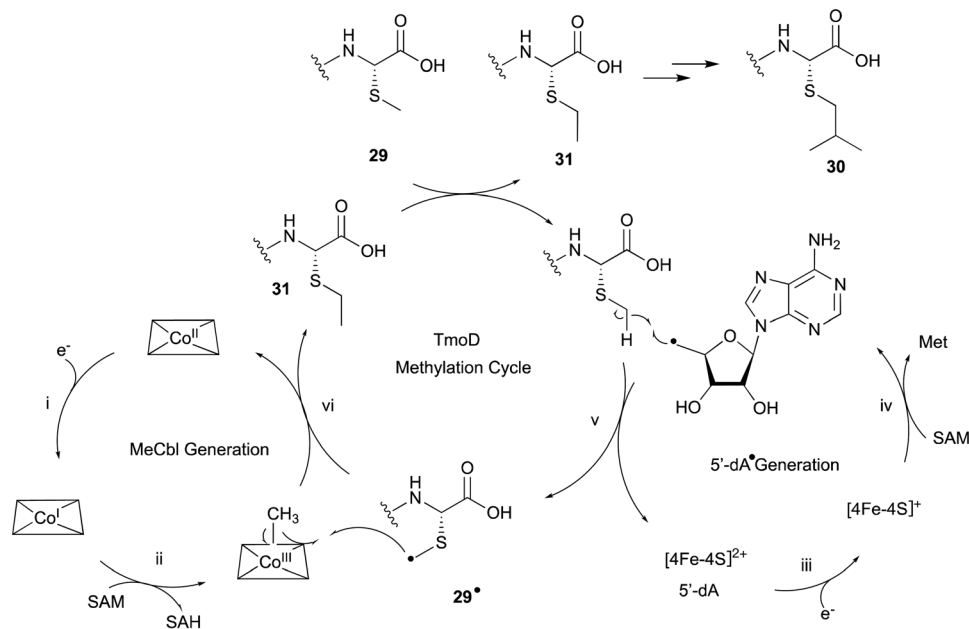


Fig. 15 Proposed mechanism of the methylation catalyzed by TmoD.

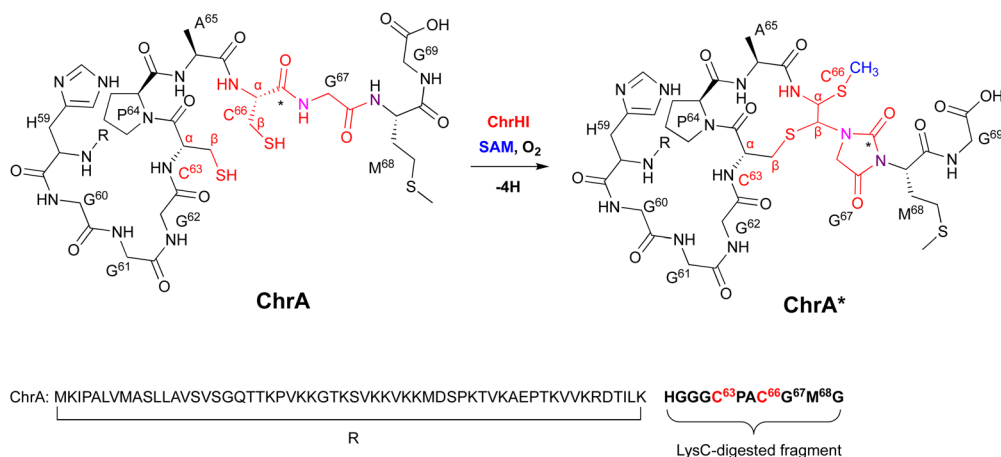


Fig. 16 Modification of ChrA by ChrHI. Purposed macrocyclization catalyzed by ChrHI. R: the remaining sequence of ChrA after digestion of the C-terminal fragment with the endoproteinase LysC.

in vitro. The reaction with SAM yielded the same product ChrA*, while the reaction without SAM generated a new product whose m/z is 36 Da lower than ChrA (32, Fig. 17).

Based on these data, the proposed reaction mechanism of ChrHI is described below (Fig. 17). First, the Fe(II) in the active site of ChrH reacts with molecular oxygen to form the Fe(III)–superoxo intermediate. Next, in a similar manner to TglH and MbnB, the Fe(III)–superoxo species abstracts H- β of Cys66 to produce a thioketyl radical (i), which forms a thioaldehyde and transfers one electron to Fe(III) (ii). The amide nitrogen of Gly67 then attacks the thioaldehyde to form a β -lactam (iii), similar to MbnBC and the mononuclear NHI enzyme, IPNS.³⁹ A second nucleophilic attack by the amide nitrogen of Met68 to the β -lactam forms a bicyclic intermediate (iv). The generation of Fe(IV)=O is expected to occur after one electron is transferred

from Fe(II) to the Fe(III)–hydroperoxy species, although the exact timing has not been determined. The Fe(IV)=O abstracts a hydrogen atom from the hemiaminal in the bicyclic intermediate, generating an oxygen radical (v). Subsequent ring opening leads to the C- α radical (vi), which can follow either pathway A [generation of the product *via* episulfide formation, and episulfide opening *via* nucleophilic attack by C₆₃ thiol and methylation by SAM, (vii)] or pathway B [first oxidized to a cation (viii), followed by *S*-methylation of the hemiaminal by SAM (ix), then addition of the thiol of Cys63 to Cys66 (x), and MeS migration to Cys66 (xi)], to generate ChrA* (Fig. 17). In absence of SAM, the radical intermediate is reduced (pathway C). Subsequent imine formation (xii) and the addition of the thiol of Cys63 to Cys66 (xiii) lead to the generation of 32. ChrHI is the first DUF692-type protein that catalyzes reactions by using SAM as a cofactor,

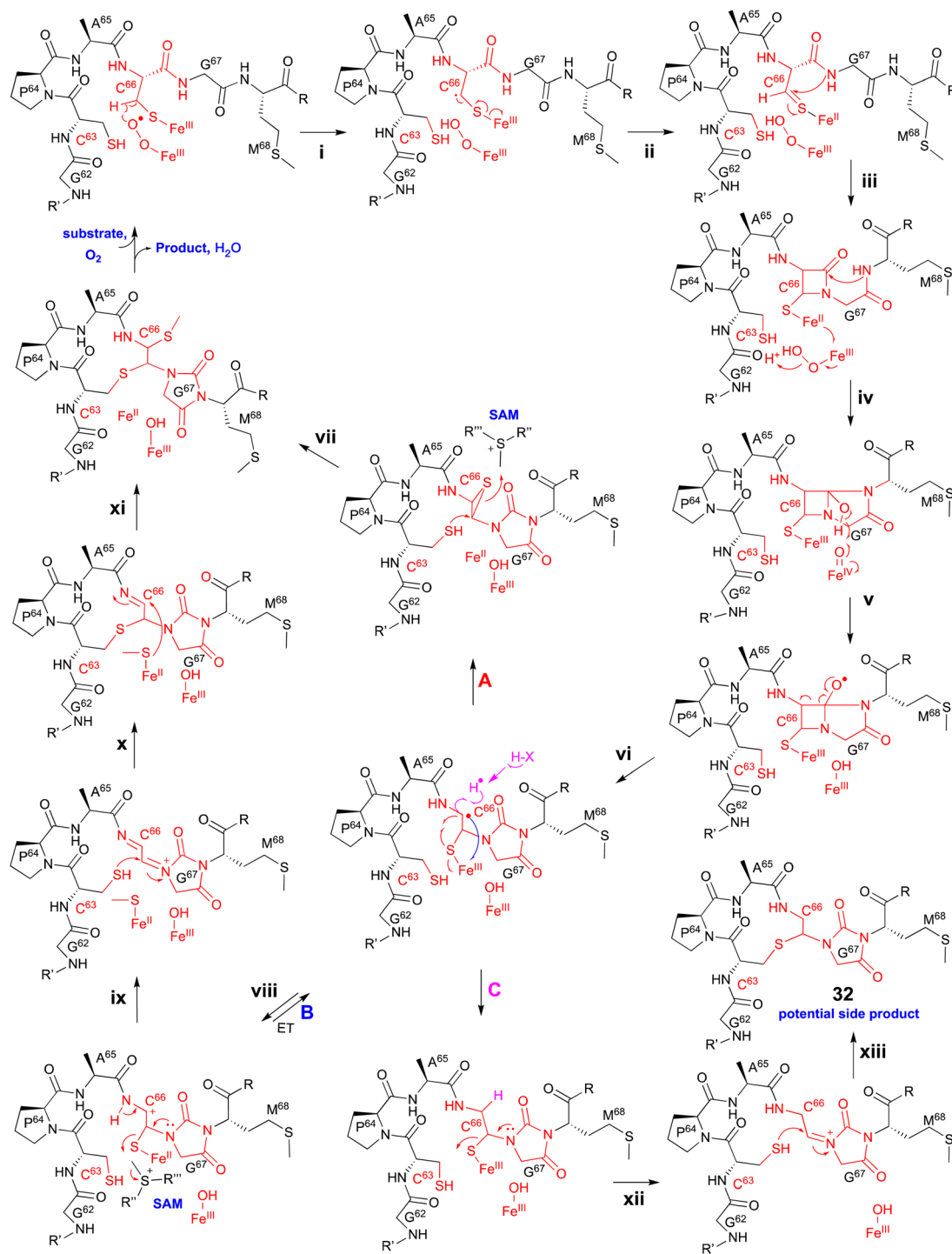


Fig. 17 Proposed mechanism of the macrocyclization catalyzed by ChrHI.

in addition to iron. The reaction bifurcation with or without SAM is intriguing.

7. SycC in AviCamCys biosynthesis

The LanD-like flavoproteins, belonging to the family of homooligomeric flavin-containing cysteine decarboxylases, catalyze

the oxidative decarboxylation of **1** at the C-terminus of the peptide to form an enethiol (Fig. 18).^{82,83} This Cys-derived enethiol is highly nucleophilic and easily reacts with Dha or Dhb to form *S*-[2-aminovinyl]-cysteine (AviCys) and *S*-[2-aminovinyl]-3-methylcysteine (AviMeCys), as part of a macrocyclic ring containing 4–6 amino acids.^{84,85}

Genome mining, using LanD as a query, was performed for the biosynthesis, including the generation of the Cys-derived



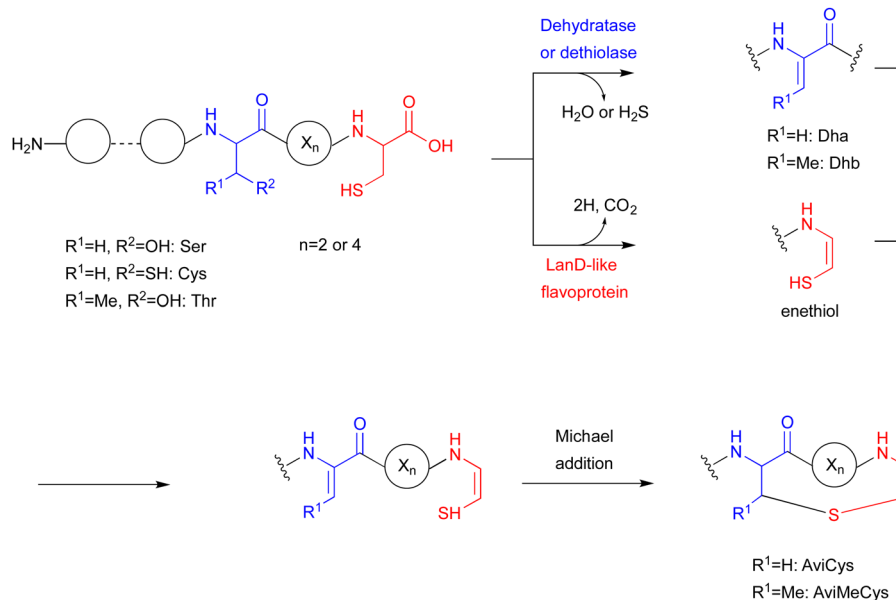


Fig. 18 Reaction catalyzed by the LanD-like protein, and the formation of AviCys and AviMeCys.

enethiol. As a result, a novel radical SAM enzyme (SycC), which generates a radical to react with an enethiol, forming the sulfide ring of AviCamCys (33) (Fig. 19), was identified in *S. ureilyticus* YC419. SycC is encoded in the *syc* cluster with a

22-aa precursor peptide (SycA), a major facilitator superfamily transporter (SycB), and a LanD-like protein (SycD). Heterologous expression in *E. coli* and *in vitro* assays showed that SycD reacts with SycA to produce the C-terminal enethiol, and SycC

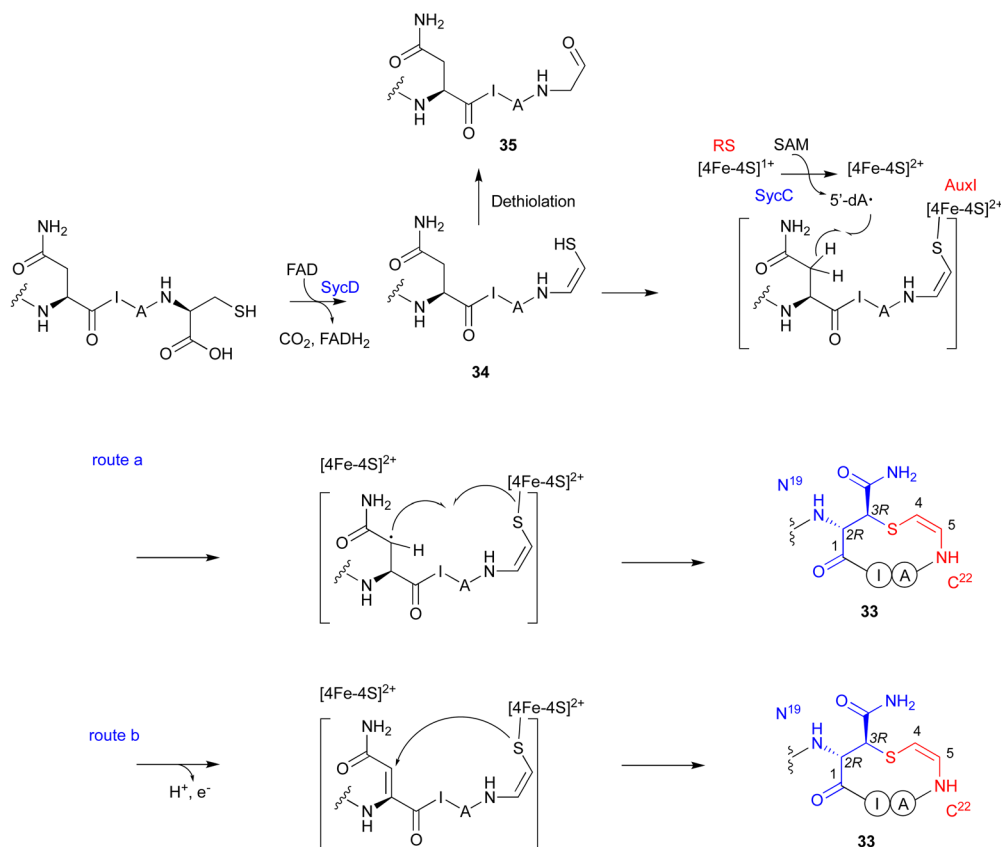


Fig. 19 Proposed mechanism of the macrocyclization catalyzed by SycD and SycC.



oxidizes the enethiol intermediate to produce **33**. Advanced Marfey's method⁸⁶ and NOESY analyses identified the absolute configuration of C-3 as *R*.

SycC belongs to the radical SAM protein family, including NxxcB⁸⁷ and CteB (PDB ID: 5WHY, contains three iron-sulfur cluster),⁸⁸ which is known for forming saturated thioether bonds. Similar to NxxcB and CteB, SycC contains both the CxxxCxxC motif for binding to the canonical [4Fe-4S]⁺ cluster and the SPASM domain for binding to the auxiliary [4Fe-4S]⁺ cluster, which enhances the activity.⁸⁹ Similar to the NxxcB-catalyzed process, the reaction of SycD was proposed as described below (Fig. 19). SycC first utilizes the canonical [4Fe-4S]⁺ cluster to provide an electron to SAM, generating the 5'-dA radical. The 5'-dA radical then abstracts the C-β hydrogen of Asn19, forming the C-β radical. The auxiliary [4Fe-4S]⁺ cluster either activates/conjugates the enethiol of the C-terminal Cys22 with the Asn19Cβ radical (route a) or deprotonates H-α to form an α,β-unsaturated diamide for the following enethiol addition (route b). In the absence of SycC, the intermediate **34** undergoes hydration-coupled dethiolation to form the shunt aldehyde **35**.

Previously, the knowledge about enethiol modification was limited to the generation of Avi(Me)Cys by dehydration or dethiolation.⁸³ SycC is a novel enzyme that forms a radical at the upstream Asn residue and promotes sulfide formation between the generated radical and enethiol to produce the unsaturated thioether, AviCamCys. This finding enriches our knowledge about the enzymatic transformation of enethiol-containing peptides. Several radical SAM enzymes involved in the modification of Cys-containing peptides have also been reported, such as Alba for thioether formation in subtilisin A biosynthesis,⁹⁰ but this review presents the more recently discovered example involving the modification of a Cys-derived enethiol.

8. Conclusion

Several new Cys-modifying enzymes, including CDO homologs, dinuclear NHI-dependent enzymes, and radical SAM enzymes, have been identified in natural product biosynthesis, revealing unusual chemical reactions in L-Cys metabolism. These advances provide detailed insight into the novel secondary metabolism of L-Cys-related natural products, expand the repertoire of L-Cys modifying enzymes, and allow more efficient access to unique molecules. With growing interest in the biosynthesis of natural products related to amino acids, the biosynthetic study of peptides or alkaloids from nature has emerged as an important research direction. With significant progress in the study of L-Cys metabolism, such mechanistic elucidation based on structural, biochemical, and computational analyses will not only provide useful insights into the Cys-derived biosynthetic pathways, but also pave the way for enzyme engineering toward future drug discovery.

Conflicts of interest

There are no conflicts to declare.

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