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## New insights into the disulfide bond formation enzymes in epidithiodiketopiperazine alkaloids†

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Epidithiodiketopiperazines (ETPs) are a group of bioactive fungal natural products and structurally feature unique transannular disulfide bridges between  $\alpha$ ,  $\alpha$  or  $\alpha$ ,  $\beta$  carbons. However, no enzyme has yet been demonstrated to catalyse  $\alpha$ ,  $\beta$ -disulfide bond formation in these molecules. Through genome mining and gene deletion approaches in *Trichoderma hypoxylon*, we identified a putative biosynthetic gene cluster of pretrichodermamide A (1), which requires a FAD-dependent oxidoreductase, TdaR, for the irregular  $\alpha$ ,  $\beta$ -disulfide formation in 1 biosynthesis. *In vitro* assays of TdaR, together with AcIT involved in aspirochlorine and GliT involved in gliotoxin biosynthesis, proved that all three enzymes catalyse not only the conversion of red-pretrichodermamide A (4) to  $\alpha$ ,  $\beta$ -disulfide-containing 1 but also that of red-gliotoxin (5) to  $\alpha$ ,  $\alpha$ -disulfide-containing gliotoxin (6). These results provide new insights into the thiol-disulfide oxidases responsible for the disulfide bond formation in natural products with significant substrate and catalytic promiscuities.

### Introduction

Natural products containing sulfur display remarkable structural diversity and have a wide variety of biological activities. 1,2 A found fungi includes special subgroup in dithiodiketopiperazines (ETPs) featuring a cyclodipeptide (CDP) scaffold, which is decorated by a transannular disulfide bridge. They exhibit a broad range of bioactivities including anticancer, antibacterial, antifungal, antiviral, immunosuppressive and cytotoxic activities. 1,3,4 Typically, structural diversity of ETPs possessing different core CDPs is generated by nonribosomal peptide synthetases (NRPSs) and multiple postassembly modifications.5 Commonly derived from phenylalanine-, tyrosine- and tryptophan-containing CDPs, two subtypes of bioactive ETPs carrying disulfide bridges across  $\alpha$ ,  $\alpha$ - or  $\alpha$ ,  $\beta$ positions of the diketopiperazine (DKP) skeleton have been identified in diverse fungal species (Fig. 1). Representatively,

gliotoxin, <sup>6-9</sup> sirodesmin PL, <sup>10-14</sup> acetylaranotin, <sup>15</sup> dithiosilvatin, <sup>16</sup> sporidesmin<sup>7,17</sup> and epicorazine <sup>18</sup> bear a disulfide bridge across the  $\alpha$  and  $\alpha$  positions of the DKP skeleton. In contrast, pretrichodermamide A (1), <sup>19</sup> gliovirin, <sup>20,21</sup> adametizine A, <sup>22,23</sup> aspirochlorine as well as its biosynthetic intermediate <sup>24,25</sup> and lasiodipline D<sup>26</sup> carry an irregular disulfide bridge across

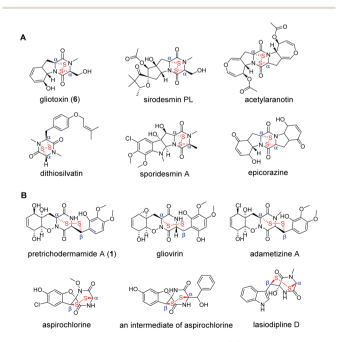


Fig. 1 Representative fungal ETPs with an  $\alpha$ ,  $\alpha$ - (A) or  $\alpha$ ,  $\beta$ - (B) disulfide bond.

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corresponding  $\alpha$  and  $\beta$  positions (Fig. 1). To date, these unique structures have captured the attention of scientists for their total synthesis<sup>27,28</sup> and biosynthesis.<sup>1,9,12</sup>

Previous studies have elucidated many aspects of the molecular mechanism involved in the biosynthesis of gliotoxin from Aspergillus fumigatus, including cyclo-L-Phe-L-Ser generation by an NRPS enzyme GliP, hydroxylation by cytochrome P450, sulfur moiety incorporation by the addition of glutathione (GSH) and its sequential degradation to the dithiol precursor. 5,29,30 A flavin adenine dinucleotide (FAD)-dependent oxidoreductase GliT is responsible for the oxidation of the dithiol group yielding the  $\alpha$ ,  $\alpha$  disulfide-containing gliotoxin.<sup>31,32</sup> In addition, biosynthesis of several ETPs including sirodesmin PL from Leptosphaeria maculans and acetylaranotin from Aspergillus terreus was extensively studied by multiple genetic and biochemical approaches. 12-15 The halogenated aspirochlorine was also investigated with respect to its crucial roles in the formation of a cyclo-L-Phe-Gly containing ETP from cyclo-L-Phe-L-Phe.<sup>24</sup> Although the interesting α, β-disulfide bridge formation was speculated to be catalysed by a GliT homolog, AclT, no experimental evidence was provided.

To identify the enzyme involved in  $\alpha$ ,  $\beta$ -disulfide bridge formation, we focused on **1** reported in *Trichoderma* sp. BCC5926, which exhibits antibacterial activity towards *Mycobacterium tuberculosis* H37Ra. <sup>19</sup> The typical  $\alpha$ ,  $\beta$ -disulfide bridge

and 1,2-oxazadecaline moiety of 1 as well as its derivatives have shown to be essential for the observed biological activities. 21,22 A biosynthetic route was proposed for gliovirin, a derivative of 1, in Trichoderma virens, based on genome mining and core gene deletion experiments.21 It was speculated that the FADdependent oxidoreductase Glv4 or Glv16 was possibly involved in the  $\alpha$ ,  $\beta$ -disulfide bridge formation. However, data for neither its detailed biosynthesis nor for the  $\alpha$ ,  $\beta$ -disulfide formation was reported. In this study, we identified 1 from a fungicolous fungus Trichoderma hypoxylon and proposed a possible biosynthetic pathway by genome mining and deletion of two targeted genes. Biochemical investigations proved that a FADdependent oxidoreductase, TdaR, was responsible for the  $\alpha$ ,  $\beta$ disulfide formation in the biosynthesis of 1. Further in vitro assays demonstrated that three oxidases TdaR, AclT and GliT from different pathways catalysed both  $\alpha$ ,  $\alpha$ - and  $\alpha$ ,  $\beta$ -disulfide bond formation in the biosynthesis of fungal ETP alkaloids.

#### Results

### Identification of the biosynthetic gene cluster for pretrichodermamide A

To identify the ETP **1**, we examined the secondary metabolites of previously reported fungus *T. hypoxylon* CGMCC 3.17906, because several ETP compounds including trichodermamide A

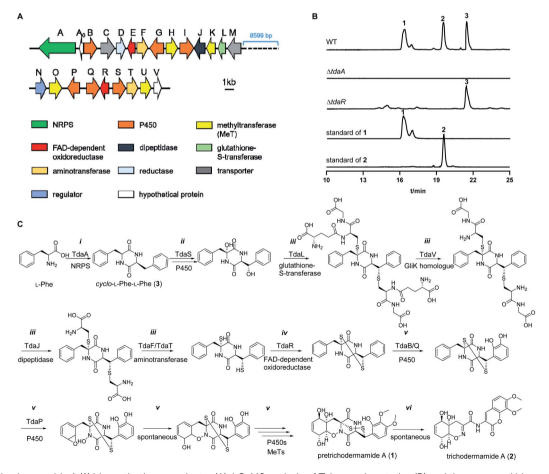


Fig. 2 Pretrichodermamide A (1) biosynthetic gene cluster (A), LC-MS analysis of *T. hypoxylon* strains (B) and the proposed biosynthetic pathway for 1 (C).

(2), aspergillazine A and aspergillazine C have been identified by our group.<sup>33</sup> Spontaneous conversion of **1** to **2** was demonstrated by Seephonkai *et al.* due to the instability of **1** under alkaline conditions or even in methanol.<sup>19</sup> Therefore, we believe that **1** should exist in this strain under controlled culture and extraction conditions. Herein, *T. hypoxylon* was cultivated in rice medium and the ethyl acetate extracts were analysed on HPLC (Fig. S1 in the ESI†). As expected, one predominant peak with an  $[M+H]^+$  ion at m/z 499.0845 was detected in a 14 day-old rice culture, which was subsequently identified as **1** after isolation and structure elucidation by comparing to reported data (Fig. S2, S3 and Table S3†).<sup>19</sup> The spontaneous conversion of **1** to **2** was observed, even under cultivation conditions and during isolation (data not shown). This result indicated that **1** was the natural ETP metabolite in *T. hypoxylon*.

To determine the biosynthetic pathway of 1, the genome of T. hypoxylon was sequenced and the draft genome sequence was used for the prediction of putative gene clusters by using Anti-SMASH.34 Based on its ETP structural character, 1 should be derived from cyclo-L-Phe-L-Phe (3), which is expected to be assembled by an NRPS.35 One of the eleven NRPS genes T\_hypo\_11188, termed tdaA, within a large 49.6 kbp cluster (Fig. 2A and Table S4†), contains a T-C-A-T-C domain structure, being similar to the core enzymes AtaP15 and AclP24 in phenylalanine-containing ETP biosynthesis (abbreviations for NRPS domains are as given before<sup>35</sup>). TdaA shares a sequence similarity of 31% with GliP9 from A. fumigatus and 88% with Glv21 21 from T. virens on the amino acid level (Table S4†). To prove its function, tdaA was deleted from the genome by PEGmediated protoplast transformation using hygromycin B as a selection marker.36 LC-MS analysis revealed complete disappearance of ETP products 1 and 2, as well as the NRPS product 3, indicating that TdaA acts as the core enzyme for the biosynthesis of 1 and 2 (Fig. 2B). To find the candidate gene for the  $\alpha$ , β-disulfide bridge formation in 1, further blast analysis was performed using the disulfide bond formation enzyme GliT in the gliotoxin biosynthetic pathway as a probe.32 Two putative FAD-dependent oxidoreductases TdaE (T\_hypo\_11193) and TdaR (T\_hypo\_11206) are located in the pretrichodermamide A (tda) cluster. Alignments with known oxidases for thioldisulfide formation showed the existence of a conserved active binding site CLFC (CXXC) box in TdaR but not in TdaE (Fig. S4†). Therefore, we performed tdaR deletion according to the aforementioned method for tdaA, leading to complete disappearance of 1 and 2, but accumulation of 3 as indicated by the analysis of extracted ion chromatograms (EICs) (Fig. 2B). This proved the involvement of TdaR in the conversion of 3 to 1 and 2. Inspection of the EICs of the culture extracts and structure analysis implied more conversion steps between 3 and 1. Thus, a biosynthetic pathway of 1 and 2 was proposed as shown in Fig. 2C. Briefly, the DKP core is assembled by NRPS TdaA (i), followed by hydroxylations at  $\alpha$  and  $\beta$  positions by a putative cytochrome P450 TdaS (ii). After sulfur incorporation and sequential degradation (iii), TdaR catalyses the  $\alpha$ ,  $\beta$ -disulfide bond formation (iv). Subsequently, further hydroxylations catalysed by multiple cytochrome P450s and methylations catalysed by different methyltransferases (MeTs) lead to the

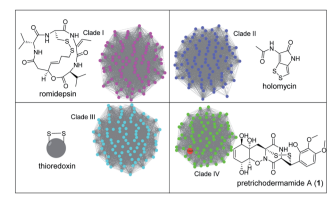


Fig. 3 Sequence clustering and phylogenetic analysis of FAD-dependent oxidoreductases involved in thiol-disulfide formation. Four distinct clades are shown in different colors with their representative sulfur-containing products. Romidepsin and holomycin were formed by DepH (clade I) and Hlml (clade II), respectively. TrxR (clade III) is a thioredoxin reductase responsible for the reduction of disulfide. TdaR, together with GliT (involved in gliotoxin biosynthesis) and AclT (involved in aspirochlorine biosynthesis), was located in clade IV with other potential fungal thiol-disulfide oxidases. Amino acid sequences were obtained from the GenBank database.

formation of  $\mathbf{1}$  with a 1,2-oxazadecaline moiety and methoxy groups (v). Lastly, spontaneous degradation results in the transformation of  $\mathbf{1}$  to  $\mathbf{2}$  (vi).

#### TdaR responsible for $\alpha$ , $\beta$ -disulfide bond formation

To address whether TdaR is involved in the  $\alpha$ ,  $\beta$ -disulfide bond formation, we performed *in vitro* biochemical characterisation. Previous studies reported that disulfide bond formation was catalysed by FAD-dependent oxidoreductases from different bacterial and fungal sources.<sup>37</sup> Hence, a sequence similarity network of TdaR homologs including 370 FAD-dependent oxidoreductases was generated which led to the identification

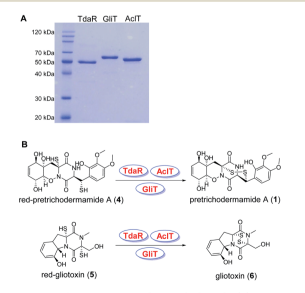


Fig. 4 Analysis of TdaR, AcIT and GliT on SDS-PAGE (A) and enzyme-mediated disulfide bond formation in the biosynthesis of 1 and 6 (B).

of four distinct clades at an *e*-value threshold of 80 (Fig. 3 and S5†).<sup>38</sup> TdaR belongs to clade IV which includes GliT and 81 other not yet fully identified fungal enzymes possibly from ETP biosynthetic pathways. Prototypes of the other three clades are DepH (clade I),<sup>39</sup> HlmI (clade II)<sup>40</sup> and TrxR (clade III)<sup>41</sup> from *Chromobacterium violaceum*, *Streptomyces clavuligerus* and *Escherichia coli*, respectively. Interestingly, proteins from clade I, II and IV presumably oxidize small molecules yielding disulfide bonds, while clade III probably acts as reductases for intracellular redox regulation.

To investigate its function, the 933 bp long coding sequence of tdaR comprising three exons was cloned from cDNA of T. hypoxylon and overexpressed in E. coli BL21(DE3) cells. The recombinant His6-tagged protein was purified with the aid of Ni-NTA agarose resin to near homogeneity which was confirmed on SDS-PAGE (Fig. 4A), yielding 5.1 mg of purified TdaR per liter of bacterial culture. Due to the instability of the unidentified substrate of TdaR for the  $\alpha$ ,  $\beta$ -disulfide bond formation, we proposed that this enzyme could also catalyse the conversion of reduced pretrichodermamide A (red-pretrichodermamide A, 4) to 1. Therefore, 1 was chemically reduced to 4 using dithiothreitol (DTT) to provide an appropriate substrate for the enzyme assays (Fig. S6†). HPLC analysis revealed a complete conversion of 0.5 mM 1 to 4 by 1 mM DTT at 30 °C for 30 min (Fig. 5A and S6†). Subsequently, this mixture was directly incubated with 0.1 µM purified TdaR at 37 °C for 30 min. As expected, the enzyme assay led to the formation of 1 which was verified by comparing the retention time with that of an authentic standard (Fig. 5A(i, ii and viiii)). This proved that the FAD-dependent oxidoreductase TdaR was responsible for the  $\alpha$ , β-disulfide bond formation in the biosynthesis of 1. Furthermore, the time course of this reaction was examined and monitored by HPLC, indicating that 17% of 0.5 mM 4 was consumed by 0.1 µM TdaR in 5 min (Fig. S7†). However, no conversion of 4 could be observed with heat-denatured TdaR or without O2 in the glove box (Fig. 5A(i and iii)). This indicated a mechanism of  $\alpha$ ,  $\beta$ -disulfide bond formation similar to GliTcatalysed  $\alpha$ ,  $\alpha$ -disulfide bond formation, which required  $O_2$ . Accordingly, we proposed that initial transformation occurred in TdaR from the oxidized form (disulfide) to reduced form (dithiol) by the putative CLFC redox system. Subsequently, electron pairs were transferred to FADox from the active site of TdaR. FAD<sub>red</sub> is then reoxidized by O<sub>2</sub> to form H<sub>2</sub>O<sub>2</sub>, leading to the concomitant production of the disulfide-containing product (Fig. 4B and 6).

To directly confirm the oxidation of TdaR on 4, 4 was purified from the mixture of 1 with tris(2-carboxyethyl)phosphine hydrochloride (TCEP) as a reducing agent on a semi-preparative HPLC (acetonitrile/ $H_2O$ , 35:65) (Fig. S9†). Incubation of TdaR with the purified 4 revealed a complete conversion of 4 to 1 after incubation at 37 °C for 30 min (Fig. S10A†). However, the formation of 1 was also observed in the negative control containing heat-denatured TdaR and 4, demonstrating the spontaneous disulfide formation. In comparison, disulfide formation in 1 catalysed by TdaR was more efficient than the spontaneous oxidation.

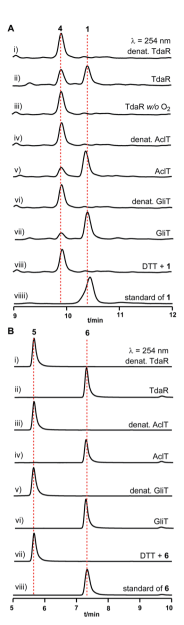


Fig. 5 HPLC analysis of enzyme assays of TdaR, AclT and GliT with 4 (A) or 5 (B). UV absorption at 254 nm was illustrated.

### Fungal thiol-disulfide oxidases catalysed both $\alpha$ , $\alpha$ - and $\alpha$ , $\beta$ -disulfide bond formation

In addition to TdaR, AclT was also predicted to catalyse an  $\alpha$ ,  $\beta$ -disulfide bond formation in the biosynthesis of aspirochlorine in *Aspergillus oryzae*. However, no detailed investigation on AclT has been reported until now. We presumed that AclT could also catalyse the  $\alpha$ ,  $\beta$ -disulfide bond formation in the biosynthesis of **1**. Therefore, the *aclT* ORF (945 bp) was amplified from *A. oryzae* cDNA and cloned into an *E. coli* expression vector pET28a(+)SUMO. The resulted plasmid pYHL87 was introduced into BL21(DE3) cells for protein overproduction, leading to 11.0 mg of purified AclT per liter of bacterial culture. It showed a yellow color and was confirmed on SDS-PAGE with the predicted size of 50.4 kDa (Fig. 4A). Subsequently, incubation of 4

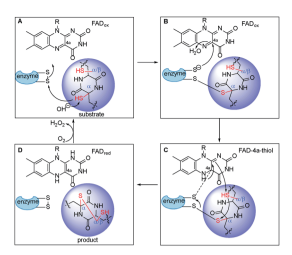


Fig. 6 Proposed mechanism of disulfide bond formation in ETP alkaloids by FAD-dependent thiol-disulfide oxidases. A transient mixed-disulfide bond was generated between substrate and enzyme (A and B) leading to a FAD-4a-thiol adduct (C). Subsequently, the disulfide in the ETP alkaloid was formed with concomitant reduction of FAD (D).

with 0.1  $\mu$ M AclT was carried out in a similar way to TdaR. HPLC analysis revealed 1 as the enzymatic product, proving AclT-catalysed  $\alpha$ ,  $\beta$ -disulfide bond formation (Fig. 5A(iv and v)). Further time course of AclT-catalysed reaction revealed that increasing amounts (18–72%) of compound 4 were consumed by 0.1  $\mu$ M AclT from 5 min to 30 min (Fig. S7†). In addition, incubation of heat-denatured AclT with the purified 4 also revealed that a trace amount of 1 is formed spontaneously (Fig. S10A†). This was consistent with the enzyme assay of TdaR with 4, indicating that AclT and TdaR performed the same function in the  $\alpha$ ,  $\beta$ -disulfide bond formation.

Notably, bioinformatics study showed that TdaR and AclT involved in the  $\alpha$ ,  $\beta$ -disulfide bond formation were located in clade IV with several known  $\alpha$ ,  $\alpha$ -disulfide bond-forming enzymes including GliT, 9,32 AtaTC15 and SirT12 (Fig. 3 and S5†). The close phylogenetic relationship triggered our interest to test if TdaR and AclT could also catalyse α, α-disulfide bond formation. Thus, reduced gliotoxin (red-gliotoxin, 5) was acquired by co-incubation of gliotoxin (6) with DTT at 30 °C for 30 min or purified from a mixture containing 6 and TCEP (Fig. S6 and S9†). Interestingly, we observed that 5 (0.5 mM) was readily converted into 6 in the presence of TdaR (0.1  $\mu$ M) or AclT (0.1 µM) after 5 min (Fig. S8†). 5 was completely transformed to 6 in 30 min by both TdaR and AclT (Fig. 5B, S8 and S10B†). These results supported our hypothesis that TdaR and AclT catalyse not only  $\alpha$ ,  $\beta$ - but also  $\alpha$ ,  $\alpha$ -disulfide bond formation in the biosynthesis of ETP alkaloids. In addition, TdaR and AclTcatalysed  $\alpha$ ,  $\alpha$ -disulfide bond formation with 5 as the substrate was more efficient than the  $\alpha$ ,  $\beta$ -disulfide bond formation.

In analogy, the coding sequence of GliT, which was proved to catalyse the  $\alpha$ ,  $\alpha$ -disulfide bond formation in the biosynthesis of **6**, was cloned into pET28a(+)SUMO and overexpressed in *E. coli* BL21(DE3). 11.2 mg purified GliT was obtained in a yellow color

from 1 liter of bacterial culture and subsequently incubated with 4 and 5 in a similar way to TdaR and AclT. In accordance with the previous study, GliT acts as a FAD-dependent oxidase for the  $\alpha,$   $\alpha$ -disulfide bond formation to generate 6 from 5 after 5 min incubation (Fig. 5B and S8†).  $^{32}$  Unprecedentedly, incubation of 4 with GliT also led to the formation of 1, as demonstrated by HPLC analysis (Fig. 5A(vi and vii)). This confirmed that the known  $\alpha,$   $\alpha$ -disulfide bond related enzyme GliT also catalysed the  $\alpha,$   $\beta$ -disulfide bond formation. The time course of GliT-catalysed oxidation of 4 was monitored by HPLC, revealing that an increasing amount of 4 ranging from 28–82% was consumed by 0.1  $\mu$ M GliT from 5 min to 30 min (Fig. S7†).

### Discussion

As a ubiquitously distributed element, sulfur is essential for mediating amino acid, peptide, and RNA synthesis in primary metabolites and secondary metabolites. 42-44 There are various examples of natural products containing sulfur atoms as key units for biological functions. In the past decades, ETPs have garnered considerable attention regarding their biosynthesis, chemical synthesis and biological application, because the disulfide group contributes promising activities probably by promoting redox cycling, formation of reactive oxygen species and protein conjugation. 45,46 Mostly, the sulfur bridge functionality of ETPs is attached to the  $\alpha$ ,  $\alpha$ -positions or  $\alpha$ ,  $\beta$ -positions of the DKP core in fungi. Significant progress in understanding the biosynthetic pathway of  $\alpha$ ,  $\alpha$ -disulfidecontaining 6 has been achieved through genetic and biochemical approaches. 5,29,30,32,37 However, fungal  $\alpha$ ,  $\beta$ -disulfidecontaining metabolites have been rarely investigated with respect to their biosynthetic mechanism. Although gene clusters coding for the biosynthesis of  $\alpha$ ,  $\beta$ -disulfide-containing aspirochlorine and gliovirin have been predicted, the required biochemical functions have remained elusive. 21,24

In this study, we identified an  $\alpha$ ,  $\beta$ -disulfide-containing metabolite 1, together with its spontaneous desulfurized product 2, in T. hypoxylon. Genome mining led to the identification of a tda cluster encoding analogues of all aforementioned gliotoxin (gli) cluster enzymes and five additional cytochrome P450s as well as three methyltransferases (Fig. 2A and Table S4†).5 Deletion of the NRPS coding gene tdaA and the FADdependent oxidase gene tdaR proved their involvement in the biosynthesis of 1 and 2 from the CDP 3 (Fig. 2B and C). Plausibly, dihydroxylation of 3,  $\alpha$ ,  $\beta$ -disulfide formation and subsequent hydroxylations as well as methylations will yield the ETP alkaloid 1. Further biochemical investigation of TdaR provided evidence for its involvement in the metabolism of 3 to 1 and proved its crucial role in the formation of the  $\alpha$ ,  $\beta$ -disulfide bridge (Fig. 4B). Thus, our results first represent an example for understanding the biosynthesis of  $\alpha$ ,  $\beta$ -disulfide-containing ETP alkaloids, especially the  $\alpha$ ,  $\beta$ -disulfide formation. Other pretrichodermamide A-like compounds derived from cyclo-L-Phe-L-Phe, including pretrichodermamides,<sup>22</sup> FA2097,47 aspergillazines,48 peniciadametizines49 and aspirochlorine,24 are very likely also formed in a similar way regarding the installation of the typical  $\alpha$ ,  $\beta$ -disulfide and a 1,2Edge Article Chemical Science

oxazadecaline moiety. Therefore, further attempts to mine  $\alpha$ ,  $\beta$ -disulfide related enzymes would provide evidence for our hypothesis.

Previous study on aspirochlorine biosynthesis has already implied a putative  $\alpha$ ,  $\beta$ -disulfide related oxidase AclT.<sup>24</sup> Herein, we proved that AclT is an analogous oxidase of TdaR catalysing the  $\alpha$ ,  $\beta$ -disulfide formation. Further biochemical investigations on TdaR and AclT revealed, interestingly, their dual function of catalysing not only the  $\alpha$ ,  $\beta$ -disulfide formation in **1** but also the  $\alpha$ ,  $\alpha$ -disulfide formation in **6** (Fig. 4B and 5). These results indicated that  $\alpha$ ,  $\beta$ -disulfide related enzymes also catalysed  $\alpha$ ,  $\alpha$ -disulfide formation, and *vice versa*. Indeed, enzyme assays confirmed that the  $\alpha$ ,  $\alpha$ -disulfide related GliT was also able to catalyse  $\alpha$ ,  $\beta$ -disulfide formation in **1**. It can be deduced that hydroxylations on the CDP core catalysed by P450 enzymes determine the positions of the two hydroxyl groups and finally the type of the disulfide bond ( $\alpha$ ,  $\alpha$  or  $\alpha$ ,  $\beta$ ).

Taken together, TdaR, AclT and GliT are dual oxidases independently forming the  $\alpha$ ,  $\alpha$ - or  $\alpha$ ,  $\beta$ -disulfide bridge in fungal ETP biosynthesis. It might well explain why most fungal related FAD-dependent oxidases were located in the same clade IV (Fig. 3), no matter they naturally catalyse  $\alpha$ ,  $\alpha$ - or  $\alpha$ ,  $\beta$ -disulfide formation. In the phylogenetic analysis of TdaR homologs, the unique clade III, exemplified by TrxR from E. coli, catalysed the reduction of a disulfide bond in thioredoxin.41 In contrast to DepH (clade I) from C. violaceum and HlmI (clade II) from S. clavuligerus involved in intra- or interchenar disulfide bonds, TdaR, AclT and GliT (clade IV) catalysed transannular disulfide bond formation.39,40 They shared a catalytically active CLFC box for redox activation, but different FAD binding sites, i.e. His139 of TdaR and AclT, but Asp139 in GliT (Fig. S4†). Plausibly in analogy to GliT-catalysed disulfide bond formation, fungal oxidases utilized a similar mechanism to install disulfide bonds across  $\alpha$ ,  $\alpha$ - or  $\alpha$ ,  $\beta$ -positions (Fig. 6). <sup>32</sup> A sulfhydryl group in the DKP-containing substrate attacks a disulfide bond in the thioredoxin motif (CXXC) within the oxidoreductase, creating a transient mixed-disulfide bond between the substrate and oxidoreductase (Fig. 6A and B). In this step, FADox receives the electrons at the C<sub>4a</sub> position, leading to a FAD-4a-thiol adduct. Subsequently, it undergoes attack at the other sulfhydryl group with disulfide formation in the ETP alkaloid and concomitant reduction to FAD<sub>red</sub> (Fig. 6C and D). Eventually, FAD<sub>red</sub> is reoxidized by molecular oxygen instead of NADP with the elimination of H<sub>2</sub>O<sub>2</sub>. We speculated that the large pocket across the active site and FAD-binding motif of GliT as well as TdaR and AclT<sup>50</sup> might result in the broad substrate specificity, making it attractive for further enzyme structure analysis and protein engineering.37

### Conclusions

In conclusion, a biosynthetic pathway of pretrichodermamide A (1) was proposed by genetic manipulation in *T. hypoxylon* and further chemical extract analysis. *In vitro* biochemical investigations first demonstrated that TdaR and AclT, which are responsible for the  $\alpha$ ,  $\beta$ -disulfide bond formation in 1 biosynthesis, also catalyse the  $\alpha$ ,  $\alpha$ -disulfide bond formation in 6.

Surprisingly, GliT involved in the  $\alpha$ ,  $\alpha$ -disulfide bond formation was proved to catalyse  $\alpha$ ,  $\beta$ -disulfide bond formation as well. Despite their unrelated fungal sources and different natural metabolite pathways, they share obviously common catalytic ability to produce  $\alpha$ ,  $\alpha$ - or  $\alpha$ ,  $\beta$ -disulfide-containing products. Our results provided representative examples of fungus-originated thiol-disulfide oxidases catalysing different types of disulfide formation in the biosynthesis of ETP alkaloids. It can be speculated that in the near future more fungus-originated thiol-disulfide oxidases would be discovered which catalyse both  $\alpha$ ,  $\alpha$ - and  $\alpha$ ,  $\beta$ -disulfide formation, even exhibiting wider specificities towards more substrates.

### Conflicts of interest

There are no conflicts to declare.

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