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Heterologous production of asperipin-2a: proposal for sequential oxidative macrocyclization by a fungi-specific DUF3328 oxidase†

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Asperipin-2a is a ribosomally synthesized and post-translationally modified peptide isolated from *Asperigillus flavus*. Herein, we report the heterologous production of asperipin-2a and determination of its absolute structure. Notably, the characteristic bicyclic structure was likely constructed by a single oxidase containing the DUF3328 domain.

RiPPs (ribosomally synthesized and post-translationally modified peptides) are a diverse and rapidly expanding class of natural products with potent biological activities as well as unique structures.¹ Although RiPPs can be found in all three domains of life (archaea, bacteria, and eukaryotes), they have rarely been identified in fungi. In Basidiomycetes, α -amanitin, phalloidin, and omphalotin were demonstrated to be RiPPs.^{2–4} In contrast, in Ascomycetes, there were no such reports until the biosynthetic gene cluster for ustiloxin B was discovered (Fig. 1).^{5–7} Recently, it was also reported that phomopsis A from *Phomopsis leptostromiformis* and epichloëcyclins from the genus *Epichloë* belong to the RiPP class.^{8,9} Led by the identification and characterization of the biosynthetic genes for ustiloxin B, Nagano and co-workers extensively searched similar gene clusters in fungal genomes and classified them into more than 40 types.¹⁰ They also identified asperipin-2a (**1**) as a product derived from one of the gene clusters. The co-occurrence of genes encoding precursor peptides and enzymes with the DUF3328 domain is a unique feature of these gene clusters. By employing an *Aspergillus oryzae* expression system, we revealed that two oxidases UstYa and UstYb, both of which contain the DUF3328 domain, and tyrosinase UstQ were

involved in the oxidation and macrocyclization of the precursor peptide that furnishes the 13-membered ring of ustiloxins.¹¹ The details of the macrocyclization process as well as the function of DUF3328 proteins in the other gene clusters are yet to be elucidated.

1 is a bicyclic peptide that possesses two macrocyclic ether rings consisting of 14- and 17-membered paracyclophans. The putative biosynthetic gene cluster for **1** is composed of four genes encoding the precursor peptide (AFLA_041400, referred to *aprA* in this study), UstYa/Yb homolog (AFLA_041390, *aprY*), a transporter (AFLA_041380, *aprT*), and isoflavone reductase-related enzyme (AFLA_041370, *aprR*) (Fig. S1†). As proposed in the biosynthesis of ustiloxin B, *AprY* containing the functionally unknown domain DUF3328 likely catalyzes oxidative macrocyclization during the biosynthesis of **1**. However, in the previous study, only *aprA* and *aprY* gene deletions were investigated and the biosynthetic pathway of **1** remains unclear.¹⁰ In the present study, we report the heterologous expression of the four genes involved in the production of **1**, showing that DUF3328 oxidase is involved in the biosynthesis of this compound. The improved production yield of **1** also allowed us to

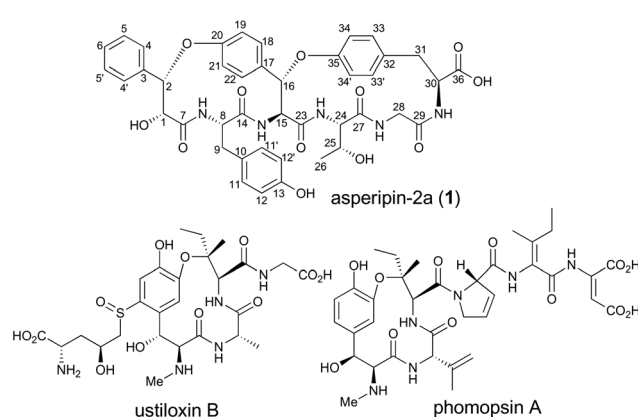


Fig. 1 Asperipin-2a (**1**) and related cyclic peptides.

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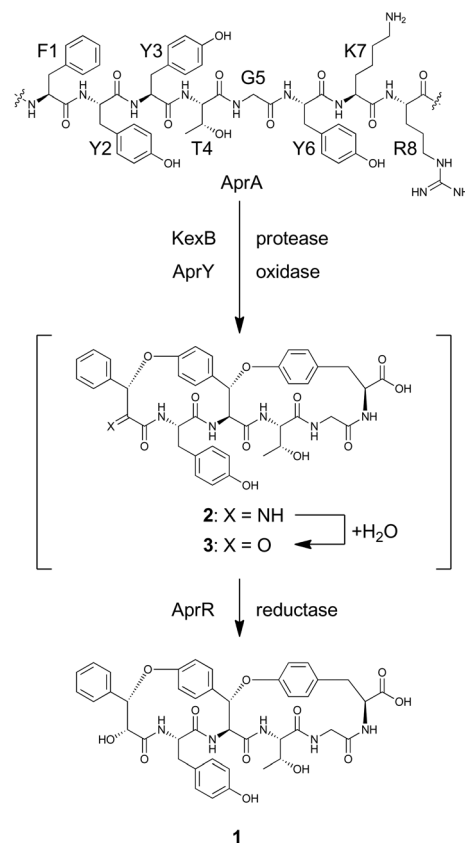
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determine its absolute configuration by chemical degradation, chiral HPLC, and NMR analyses.

To reconstitute the biosynthetic pathway, four genes, *aprA*, *aprY*, *aprR*, and *aprT*, were amplified from genomic DNA of *Aspergillus flavus*. The *aprA* gene was cloned into the plasmid pUSA2¹² to construct pUSA2-*aprA*, while the *aprY* gene was inserted into pUARA2¹² to generate pUARA2-*aprY* (Fig. S2†). The two other genes, *aprR* and *aprT*, were cloned into pAdeA2¹³ to construct pAdeA2-*aprRT* (Fig. S2†). Sequencing the *aprA* gene revealed that the precursor peptide contains eight repeats of the core sequence “FYTYGY” while the AprA sequence deposited in NCBI contains eleven repeats (see ESI, page S17†). The two plasmids, pUARA2-*aprY* and pAdeA2-*aprRT*, were introduced into *A. oryzae* NSAR1 (AO-WT) to generate AO-*aprYRT*. The resulting transformant did not produce **1** as the precursor peptide was absent in this strain. We then introduced pUSA2-*aprA* into this transformant to generate AO-*aprAYRT*. As we expected, the resulting transformant successfully produced **1** (Fig. 2). The 1D- and 2D-NMR spectra of the isolated compound as well as its HR-ESI-MS were in good agreement with previously reported spectra, confirming that the above four genes were sufficient for the biosynthesis of **1** (Table S3†). We also prepared the transformants AO-*aprAYR* and AO-*aprAYT* to test whether AprR and/or AprT were essential for biosynthesis. Although these showed significantly decreased levels of production, both transformants still produced trace amounts of **1**, suggesting that AprR and AprT were required for biosynthesis and that adventitious proteins in the host strain complement their function. Considering that deletion of *aprY* in *A. flavus* abolished the production of **1** in the previous study, this observation suggests that AprY is involved in the formation of the bicyclic structure (Scheme 1).

Using AO-*aprYRT* as a host strain, we also investigated the conversion of two AprA analogs, AprA/Y3F and AprA/Y6F. As either the third or the sixth tyrosine of the core peptide is substituted with phenylalanine, we expected that these mutants could not undergo cyclization and that monocyclic products might be obtained. We constructed the plasmids pUSA2-*aprA*/Y3F and pUSA2-*aprA*/Y6F and introduced them into AO-*aprYRT*. Neither AO-*aprAYRT*/Y3F nor AO-*aprAYRT*/Y6F



Scheme 1 Putative biosynthetic pathway for **1**.

generated the expected products, thus suggesting the importance of both the third and the sixth tyrosine for cyclization (Fig. S3†). As no related metabolites such as monocyclic compounds, were observed, the cyclization might be a sequential process. However, further experimental verification is necessary to address this hypothesis.

A BLAST search using AprA as a query retrieved orthologous precursor proteins from at least three strains including *Aspergillus oryzae*, *Aspergillus parasiticus*, and *Aspergillus arachidicola* (Table S4 and Fig. S4†). Notably, each gene was flanked by *aprY*, *aprR*, and *aprT* homologs, suggesting that they are involved in the biosynthesis of **1** or its congeners. The number of core peptide repeats of each AprA ortholog varies from one to eight. The third and the sixth tyrosine residues were strictly conserved among each repeat (Fig. S5†). The first phenylalanine was also conserved, indicating the importance of these three residues in the cyclization process. On the other hand, in some cases, the second, the fourth, and the fifth residues were substituted with phenylalanine, histidine, and asparagine, respectively (Fig. S5†). This observation indicated that these positions did not affect cyclization and could be substituted for structural diversification.

The highly-strained 14-membered paracyclophane ring of **1** has limited conformational flexibility and is rotationally restricted,¹⁴ thus allowing us to elucidate its relative stereochemistry by 1D- and 2D-NMR analyses. Both H1 and H2 were

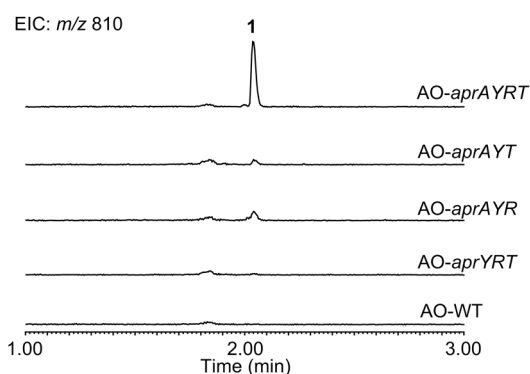


Fig. 2 LC-MS profiles of the metabolites extracted from transformants. Chromatograms were extracted at *m/z* 810.

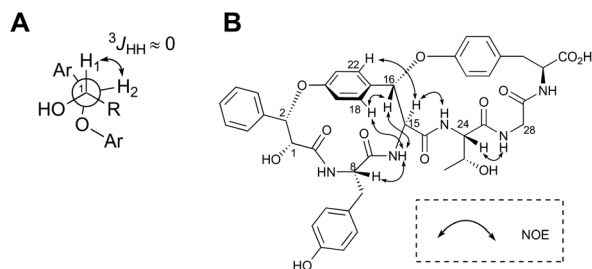
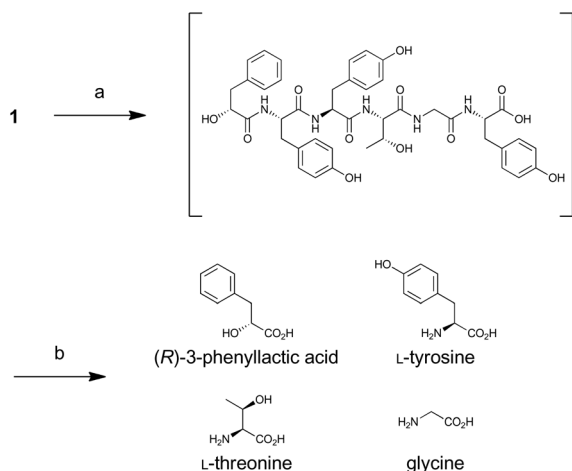


Fig. 3 Stereochemical analysis of **1**. (A) Newman projection of 3-phenyllactic acid moiety of **1**. (B) Key ROESY correlations.

observed as broad singlets, suggesting a small $^3J_{\text{HH}}$ value between these protons (Fig. 3A). On the basis of this observation, the relative stereochemistry at C1 and C2 can be deduced as (1*R*,2*S*) or (2*R*,1*S*). In addition, the NOEs between H16 and 15-NH, H16 and H18, and H15 and H22 clearly indicated that the relative configuration at C15 and C16 was either all-*S* or all-*R* (Fig. 3B).

For determination of the absolute configuration, **1** was subjected to a Pd-mediated hydrogenolysis, followed by hydrolysis with 6 M HCl to give a mixture of 3-phenyllactic acid and amino acids (Scheme 2). A mixture of amino acids were then converted with *N* $^{\alpha}$ -(5-fluoro-2,4-dinitrophenyl)-L-leucinamide (L-FDLA) to yield L-FDLA derivatives.^{15–17} LC-MS analysis revealed that the retention times of the two constituent amino acids Tyr and Thr from **1** were identical to those of the L-Tyr- and L-Thr-derivatives, respectively (Fig. 4A and B). L-FDLA-Gly was also observed (Fig. 4C). The molar ratio of L-Tyr and Gly was roughly estimated to be 3 : 1 by comparing the peak areas of FDLA derivatives (Fig. S6†). These results are consistent with the ribosomal synthesis of the precursor peptide AprA. The hydrolysate was also analyzed by chiral HPLC without derivatization. The retention time of 3-phenyllactic acid from **1** was identical to that of commercial (*R*)-3-phenyllactic acid, thus confirming the stereochemistry at C1 to be *R* (Fig. 4D). Together with the relative configuration as described above, the absolute structure of **1** was determined as shown in Fig. 1.



Scheme 2 (a) HCOONH₄, 10% Pd/C, MeOH, 70 °C; (b) 6 M HCl, 110 °C.

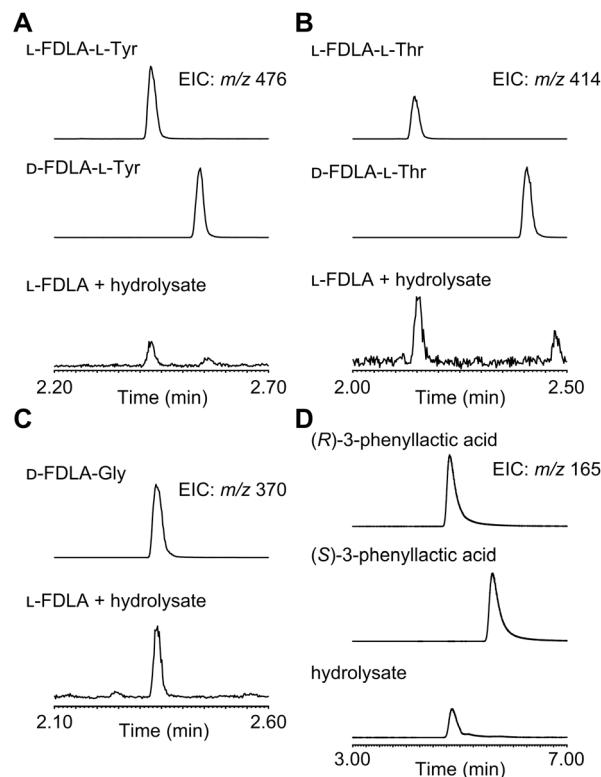


Fig. 4 LC-MS profiles of the constituents of **1**. (A) L-FDLA derivatized Tyr, (B) Thr, (C) Gly, and (D) chiral LC-MS analysis with authentic (*R*)- and (*S*)-phenyllactic acid. Chromatograms were extracted at *m/z* 476 for FDLA-Tyr, *m/z* 414 for FDLA-Thr, *m/z* 370 for FDLA-Gly, and *m/z* 165 for phenyllactic acid.

The biosynthesis of **1** starts with transcription and translation of the structural gene *aprA*. The synthesized precursor peptide AprA contains eight repeats of the core sequence FYYTGY. Each repeat is flanked by KR, which is a target site of Golgi protease, KexB, suggesting that AprA undergoes a similar proteolytic process to that known in the biosynthesis of ustiloxin B.¹⁸ AprA is also modified by AprY and AprR to furnish **1**. However, there is no information concerning the order of these post-translational modifications. As discussed above, the bicyclic structure of **1** is likely synthesized by the single enzyme AprY. Considering the relatively small size of AprY and that two C–O bonds were generated in the same stereochemical course, formation of these two bonds might be successively catalyzed in the same active site of the enzyme. In ustiloxin biosynthesis, initial hydroxylation and subsequent cyclization gave a macrocyclic system.¹¹ This transformation accompanied hydroxylation at the C $_{\beta}$ position of L-Tyr. Similarly, macrocyclization of **1** may accompany an α -hydroxylation–dehydration sequence to give imine **2**, which is readily hydrolyzed to yield putative ketone intermediate **3**. We tempted to explain that the reductase AprR may be required for the final reduction to yield **1** (Scheme 1). As AprT shows homology to major facilitator superfamily protein (MFS_1, Pfam ID: PF07690), it is likely that this protein is not involved in biosynthesis but rather exports the product.

Besides RiPPs produced by Ascomycetes such as ustiloxin B and phomopsin A, plants also produce cyclopeptides that are structurally related to **1**. Cyclopeptides, such as mauritine A,¹⁴ sanjoinin A,^{14,19,20} and ophiorrhisine A,²¹ possess 14-membered paracyclophane rings similar to that of **1** (Fig. S7†). Although DUF3328 proteins do not exist in plants, other oxidases that operate with a similar mechanism might be involved in the biosynthesis of these compounds. Future studies focusing on the mechanism of DUF3328 proteins may contribute to studies of similar cyclopeptides from other genera.

Conclusions

In this study, we achieved the heterologous production of **1** and determined its absolute structure, showing that DUF3328 protein AprY was involved in formation of the characteristic bicyclic structure. Together with our previous results on ustiloxin B, the present findings established the common strategy for macrocyclization of RiPPs produced by filamentous fungi. In both cases, oxidative cyclization catalyzed by UstY homologs accompanies a C–O bond formation between the phenol and C β of an amino acid moiety. Compared with the macrocyclization of ustiloxins, which requires three enzymes (two DUF3328s and one tyrosinase), macrocyclization of **1** would be more feasible for biochemical characterization, as it requires only one or two enzymes (AprY and/or AprR). However, preliminary attempts to reconstitute this enigmatic reaction were unsuccessful. Further studies are currently underway in our group.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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