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Discovery of a polyketide carboxylate phytotoxin from a polyketide glycoside hybrid by β -glucosidase mediated ester bond hydrolysis†

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Fungal phytotoxins cause significant harm to agricultural production or lead to plant diseases. Discovering new phytotoxins, dissecting their formation mechanism and understanding their action mode are important for controlling the harmful effects of fungal phytopathogens. In this study, a long-term unsolved cluster (polyketide synthase 16, PKS16 cluster) from *Fusarium* species was thoroughly investigated and a series of new metabolites including both complex α -pyrone-polyketide glycosides and simple polyketide carboxylates were identified from *F. proliferatum*. The whole pathway reveals an unusual assembly and inactivation process for phytotoxin biosynthesis, with key points as follows: (1) a flavin dependent monooxygenase catalyzes Baeyer–Villiger oxidation on the linear polyketide side chain of α -pyrone-polyketide glycoside **8** to form ester bond compound **1**; (2) a β -glucosidase unexpectedly mediates the ester bond hydrolysis of **1** to generate polyketide carboxylate phytotoxin **2**; (3) oxidation occurring on the terminal inert carbons of **2** by intracellular oxidase(s) eliminates its phytotoxicity. Our work identifies the chemical basis of the PKS16 cluster in phytotoxicity, shows that polyketide carboxylate is a new structural type of phytotoxin in *Fusarium* and importantly uncovers a rare ester bond hydrolysis function of β -glucosidase family enzymes.

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Introduction

Fungal phytotoxins are a large group of specialized secondary metabolites mainly produced and secreted by fungal phytopathogens, which lead to significant losses in crop yields or plant diseases.^{1–3} They usually serve as the virulence factors that subvert the host defense system and promote the invasion of fungal phytopathogens at different growth stages of plants.^{4–7} Throughout the entire infection process, fungal phytopathogens typically produce different phytotoxins, utilizing their different chemical spaces and structural types to cope with complex interactions with plants.^{8,9} Therefore, a deep understanding of the biosynthetic pathway, regulation mechanism, maturation mechanism, and action mode of phytotoxins is of great significance for effectively controlling the harmful effects of fungal phytopathogens.¹⁰

So far, a series of phytotoxins, for instance, polyketides, terpenoids, polyketide–nonribosomal peptide (pk–nrp) hybrids and peptide alkaloids have been identified from many famous fungal phytopathogens like *Alternaria*, *Botrytis*, *Colletotrichum*, *Helminthosporium*, and *Phoma*.² In addition to the above-

mentioned phytopathogenic fungal species, we mainly focused on the *Fusarium* genus, which is one of the top ten fungal phytopathogens worldwide,¹¹ comprising more than 70 species with a wide range of plant hosts,¹² and possessing a good ability to produce various phytotoxins¹³ such as fumonisins,¹⁴ deoxynivalenol,¹⁵ fusaric acid,¹⁶ apicidin,¹⁷ enniatins¹⁸ and so on (Fig. 1a and S2†).

As a continuation of our recent research on genome mining for new enzymes and the biosynthesis of mycotoxin natural products from *Fusarium* sp.,^{19–23} an important but long-term unsolved cluster (polyketide synthase 16, PKS16 cluster, Fig. 1b) from the main phytopathogenic fungus causing the “bakanae” disease of rice, *F. fujikuroi* IMI 58289,^{24–27} has attracted our attention. Previous studies revealed the following information: (1) the PKS16 cluster is located at the left end of chromosome 11 of *F. fujikuroi* IMI 58289;²⁴ (2) similar to other phytotoxins like fumonisins and enniatins, the expression of the PKS gene in the PKS16 cluster is also co-induced by low-nitrogen conditions;²⁵ most importantly, (3) the wide distribution of the PKS16 cluster among all sequenced *F. fujikuroi* species isolates strongly imply that the products derived from this cluster may act as phytotoxins in the *F. fujikuroi* species,²⁶ although their structures and mode of actions are still unclear to date.

In this work, a complete and homologue gene cluster (namely *pro* cluster) of PKS16 was discovered from *F. proliferatum* by genome mining. A series of new products were successfully identified from the *pro* pathway, which include

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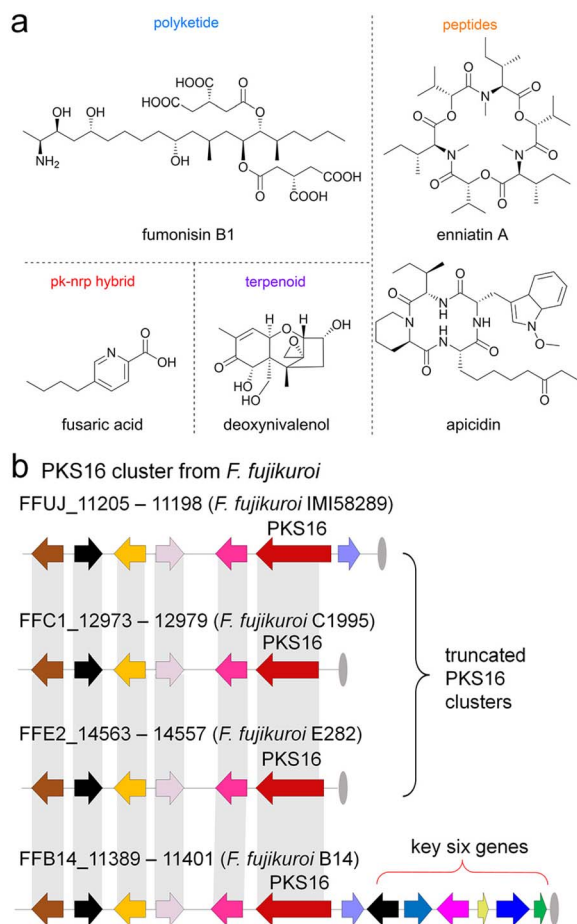


Fig. 1 Fungal phytotoxins and PKS16 clusters from *Fusarium* species. (a) The well-known phytotoxins produced by *Fusarium* sp. (b) PKS16 and its homologue clusters in *F. fujikuroi* species. The PKS16 cluster in *F. fujikuroi* IMI 58289 lacks the key tailoring-step genes, while the core hrPKS is truncated in strains C1995 and E282.

both complex α -pyrone-polyketide glycosides and simple polyketide carboxylate compounds. The whole process reveals an unusual assembly and inactivation process for phytotoxin biosynthesis, with key points including (1) a flavin monooxygenase (FMO) catalyzing Baeyer–Villiger (B–V) oxidation on the linear polyketide side chain of α -pyrone-polyketide glycoside **8** to form ester bond compound **1**; (2) a β -glucosidase unexpectedly mediating the ester bond hydrolysis of **1** to generate a new polyketide carboxylate phytotoxin **2** from *Fusarium* sp., representing a rare function of β -glucosidase family enzymes; and furthermore, (3) the oxidation occurring on the terminal inert carbons of **2** by intracellular oxidase(s), which eliminates the phytotoxicity of **2**.

Results and discussion

Genome mining discovers the complete and homologue *pro* cluster of PKS16 in *F. proliferatum*

Since the production of phytotoxins is always transient and sometimes requires interactions between fungal phytopathogens and plants to stimulate their generation,⁸ we initially

inferred that the PKS16 products may be produced in a very small amount, making them difficult to identify in *F. fujikuroi* IMI 58289. However, the position of PKS16 at the left end of chromosome 11 highlights another possibility that, the absence of some key genes in the PKS16 cluster leads to *F. fujikuroi* IMI 58289 losing its ability to produce the corresponding compounds.

To test this hypothesis, we first carefully investigated the homologue gene clusters from publicly available genome sequences of *F. fujikuroi* species. As shown in Fig. 1b, (1) compared to PKS16, the two clusters from *F. fujikuroi* C1995 and *F. fujikuroi* E282 are also incomplete; moreover, the truncated core PKS genes in these two clusters indicate that they have no function; however, (2) the cluster in *F. fujikuroi* B14 is considered to be complete, with additional six genes located downstream of the core highly reducing PKS (hrPKS) gene. We further compared this cluster with other publicly available *Fusarium* genomes and our lab databases using local BLAST with the sequence of the core hrPKS gene. Indeed, the complete and homologue gene clusters of PKS16 have also been found in many *Fusarium* strains, such as *F. denticulatum*, *F. mexicanum*, *F. subglutinans*, *F. sphaerosporum* and *F. proliferatum* (Fig. S3[†]), which indicates the wide distribution of PKS16 homologue clusters in *Fusarium* species. Interestingly, in addition to *Fusarium* species fungi, the PKS16 homologue clusters are also found in other phytopathogenic fungi like *Neonectria ditissima* and *Xylona heveae* (Fig. S3[†]).

The representative *pro* cluster (NCBI accession number PQ271635) from *F. proliferatum* CGMCC 3.4710 is shown in Fig. 2a. Aside from the core hrPKS gene (*proF*), the *pro* cluster contains other genes encoding α -galactosidase (*proA*), transporters (*proB* and *proH*), cytochrome P450 (CYP450, *proC*), DUF3129 family protein (*proD*), B–V monooxygenase (BVMO, *proE*), α , β -hydrolase (*proG*), FMO (*proI*), membrane-bound glycosyltransferase (mGT, *proJ*, Fig. S4[†]), a protein (*proK*) of unknown function and β -glucosidase (*proL*). It is worth noting that the discovery of a glycosyltransferase in the *pro* cluster indicates that the products of PKS16 and its homologue clusters are possible new polyketide glycoside hybrid compounds in *Fusarium* species;²⁸ the only previously identified polyketide glycoside from *Fusarium* is fusapyrone.^{29–31} Moreover, *proA*, *proC* and *proD* are not conserved genes in other homologue clusters (Fig. 2a and S3[†]), indicating that they may not be involved in the biosynthesis of *pro* pathway products.

The α -pyrone-polyketide glycoside **1** and polyketide carboxylate **2** are products of the *pro* cluster

We first screened various low-nitrogen incubation conditions and found that all *pro* genes were under transcription when *F. proliferatum* was incubated in glycerol medium (Fig. S5[†]), which strongly implies that the corresponding products of the *pro* cluster were produced. To probe the products of the *pro* pathway, we knocked out (KO) the whole *pro* cluster (*Fp- Δ proA–L*, Fig. S6[†]) and compared the metabolite profiles between the *F. proliferatum* wild type and the *Fp- Δ proA–L* strain, and it was found that the production of six compounds (**1–6**) was

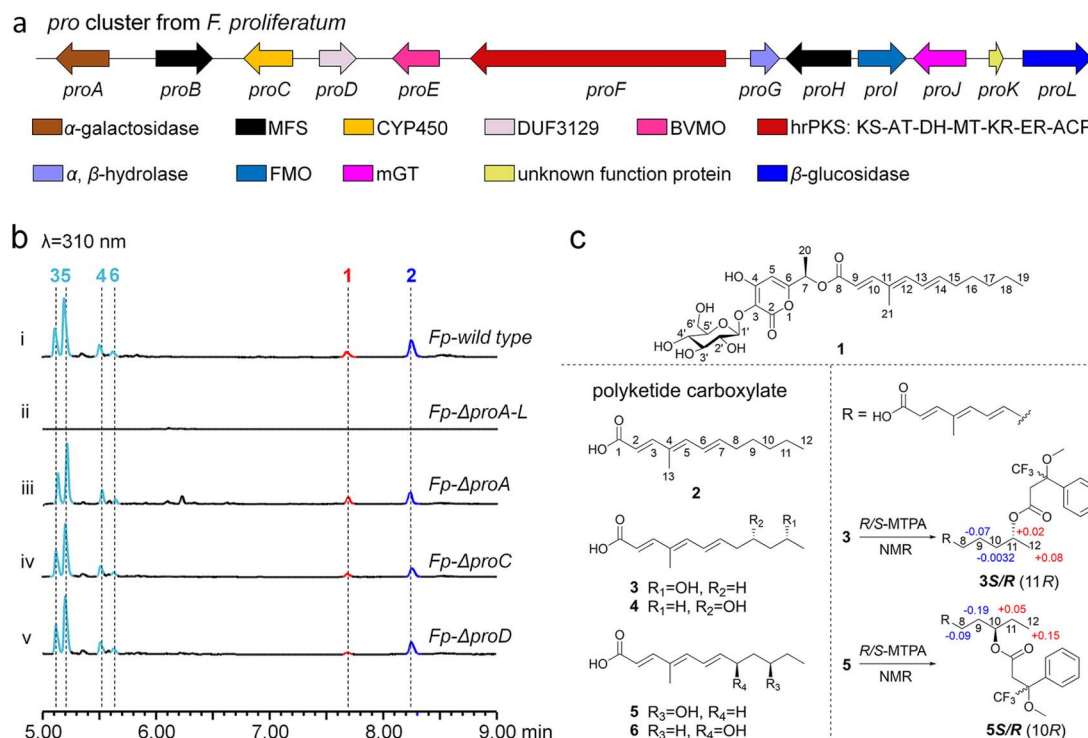


Fig. 2 Identification of the homologue *pro* cluster and confirmation of the corresponding products in *F. proliferatum*. (a) Organization and proposed gene functions of the *pro* cluster. (b) LC-MS analysis of the extracts from *F. proliferatum* and its knockout mutants. (c) Chemical structures of compounds 1–6.

eliminated in *Fp-ΔproA-L* (Fig. 2b(i and ii)). Subsequent large-scale fermentation of the *F. proliferatum* wild type, purification, and confirmation of the structures of 1–6 via HR-MS and NMR analysis showed that (1) compound 1, namely, proliferapyrone A, is an α-pyrone-polyketide glycoside (Fig. 2c, Table S6 and Fig. S36–S42†), and its linear polyketide side chain is linked to α-pyrone via an unusual ester bond, and O-glucosylation occurs at the C3–OH of the α-pyrone moiety via a β-glycosidic bond (C3–O–C1' ether bond), which was further confirmed to be D-glucose by comparison with the standard after hydrolysis with 4 M trifluoroacetic acid (Fig. S7†); (2) unlike the complex structure of 1, compounds 2–6, namely, proliferic acids A–E, are all polyketide carboxylate compounds, where 3–6 are likely hydroxyl-derivatives of 2 via oxidation of the inert carbon atoms at C8–C11 of 2, respectively (Fig. 2c, S43–S77 and Tables S7–S11†); (3) using 3 and 5 as examples, based on the Mosher derivation³² results (Fig. 2c, S78–S81, Tables S12 and S13†), the stereochemistry of the hydroxyl group in 3–6 was assigned as *R* configuration, respectively.

Next, we further investigated the relationship between 1–6 and three non-conserved genes (*proA*, *proC* and *proD*) of the *pro* cluster. Individual KO mutants of *proA*, *proC* and *proD* were first generated (Fig. S6†), and subsequent analysis of the metabolites from each KO mutant showed that these three non-conserved genes are indeed not involved in 1–6 formation; the *Fp-ΔproA*, *Fp-ΔproC* and *Fp-ΔproD* mutants retain the ability to produce compounds 1–6, respectively (Fig. 2b(iii–v)). The exclusion of CYP450 *proC* acting as the radical-type oxidase in the

hydroxylation of the inert C8–C11 of 2 to form 3–6 strongly suggests that other oxidase(s) besides the *pro* cluster is(are) responsible for these conversions. Indeed, when 2 was fed into the *Fp-ΔproA-L* strain, the formation of 3–6 was observed (Fig. S8†). These results clearly show that compounds 3–6 are over-oxidized off-pathway derivatives of 2 by *F. proliferatum*; thus, 1 and 2 are the final on-pathway products corresponding to the *pro* cluster.

hrPKS and α,β-hydrolase collaboratively produce α-pyrone-polyketide precursor 7

Structural analysis of 1 and 2 indicates that these two compounds feature significantly different skeletons, suggesting that an unusual biosynthetic process possibly occurred during their biosynthesis. We initially inferred that the generation of 2 results from the ester bond hydrolysis of 1; therefore, according to the proposed function of *pro* cluster enzymes, the preferred candidate responsible for this step should be α,β-hydrolase *proG*. However, when 1 was incubated with purified *proG* (Fig. S9†), the conversion of 1 to 2 was not observed (Fig. S10†). We also considered that the hydrolysis of 1 to 2 could be an off-pathway conversion in *F. proliferatum*; however, when 1 was fed into the *Fp-ΔproA-L* strain, only trace amounts of 2 were observed (Fig. S11†). Moreover, the chemical conversion of 1 to 2 was investigated under various pH buffer conditions, and the formation of 2 was only detected under strongly basic conditions (pH > 11, Fig. S12†). These conversion investigations indicate that the formation of 2 should be an enzyme-catalyzed

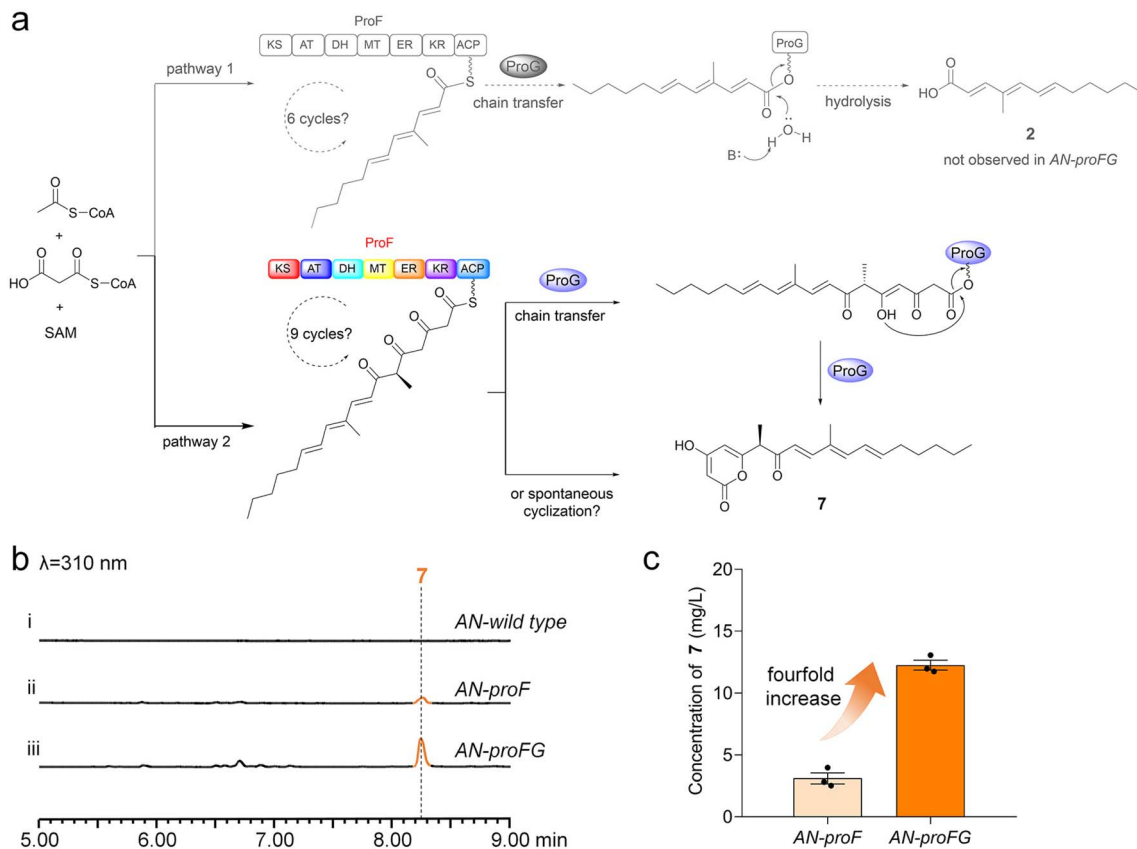


Fig. 3 The hrPKS and α,β -hydrolase collaboratively produce α -pyrone-polyketide precursor 7. (a) Two proposed pathways for the collaboration of hrPKS ProF and α,β -hydrolase ProG. (b) LC-MS analysis of the culture extracts from *AN-proF* and *AN-proFG*. (c) The yields of 7 increased nearly fourfold in *AN-proFG* indicating that ProG plays a supportive role with ProF in the biosynthesis of 7.

route, with the corresponding enzyme(s) located in the *pro* cluster.

Based on these observations, we alternatively reasoned that the skeletal differences between 1 and 2 may be due to differences in the collaboration of hrPKS ProF and α,β -hydrolase ProG. As shown in Fig. 3a, (1) during six cycles of catalysis by ProF, ProG may catalyze the hydrolysis of the ACP-bound polyketide intermediate to form 2; (2) after nine cycles of catalysis by ProF, an α -pyrone polyketide is ultimately generated through the spontaneous cyclization of the ACP-bound terminal triketone polyketide intermediate.³³ This cyclization may also be catalyzed by ProG.

To validate this hypothesis, we cloned and transferred these two genes into the typical heterologous host *A. nidulans*.^{34–37} In contrast to the *A. nidulans* wild type control, (1) a compound (7, proliferapyrone B) was produced by *AN-proF* (Fig. 3b(i and ii)), and its structure was confirmed to be α -pyrone polyketide through HR-MS and NMR analysis (Fig. 3a, Table S14 and Fig. S82–S88[†]). The C7-methyl stereochemistry of 7 has been confirmed to be the *R* configuration by comparing the experimental and calculated ECD spectra³⁸ (Fig. 3 and S13[†]); (2) the expected compound 2 was not observed in *AN-proFG*; instead, the yields of 7 increased nearly fourfold (Fig. 3b(iii) and 3c), indicating that ProG plays a supportive role to ProF in the biosynthesis of 7. Therefore, although these results do not

support the hypothesis that 2 originates from the ProF branch pathway (pathway 1, Fig. 3a), they clearly elucidate the collaborative mechanism of ProF and ProG in the production of 7, where ProF undergoes complete nine cycles to generate an on-pathway intermediate with the terminal triketone moiety; ProG then mediates its transfer and cyclization to yield the α -pyrone polyketide precursor 7 (pathway 2, Fig. 3a).³³

Confirmation of the biosynthetic genes and the catalytic order from 7 to 1

We next investigated the biosynthetic order from 7 to 1. A structural comparison between 7 and 1 indicates that some tailoring steps, such as sequential hydroxylation and *O*-glucosylation at the C3 of the α -pyrone ring and subsequent B–V oxidation at the C7–C8 of 7, could accomplish the conversion of 7 to 1.

Based on this assumption, three enzymes ProI, ProJ and ProE are ideal candidates for catalyzing these steps. Indeed, when *proI*, *proJ* and *proE* were co-expressed with *proFG* in *A. nidulans*, (1) compound 1 and one minor compound (8, proliferapyrone C) were produced by *AN-proFGIJE* (Fig. 4a(i)); (2) removal of *proE* eliminates the production of 1; however, 8 was accumulated in *AN-proFGIJ* (Fig. 4a(ii)). Subsequent purification, and HR-MS and NMR analysis confirmed 8 to be an α -

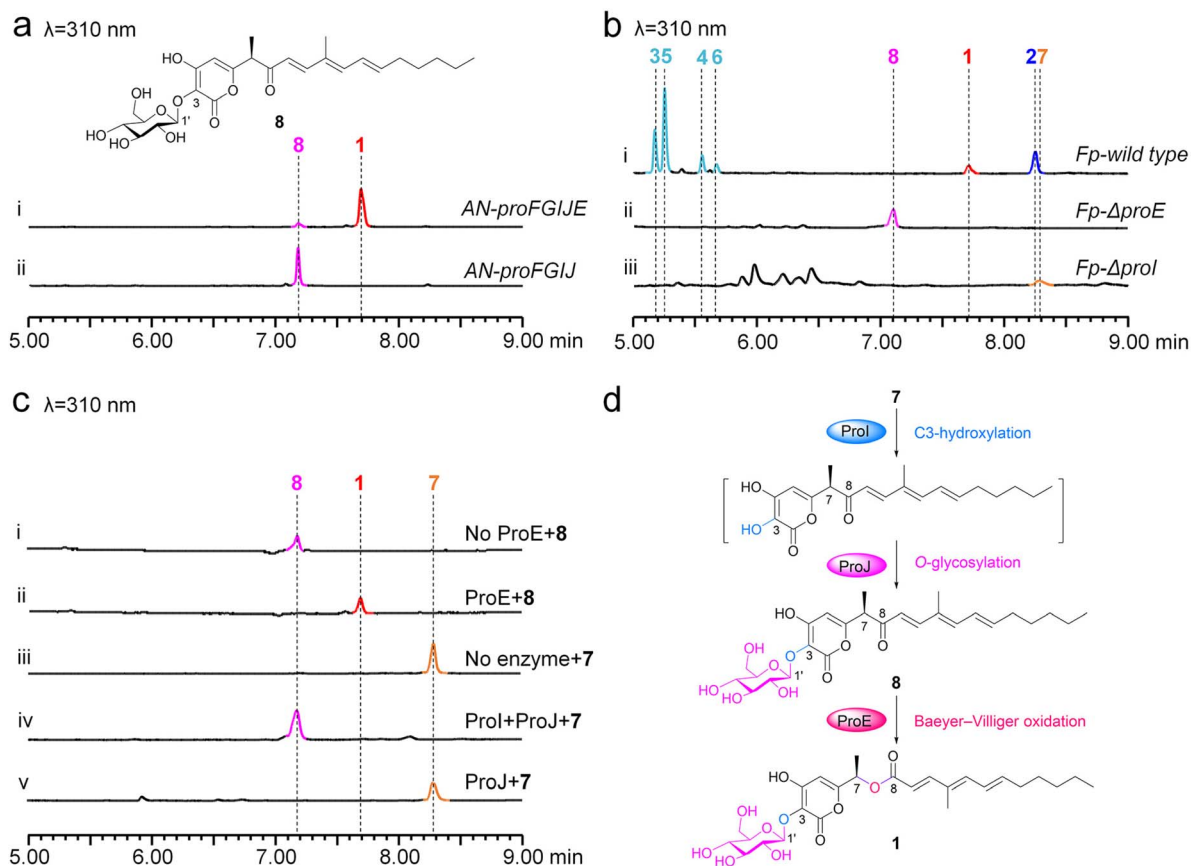


Fig. 4 Confirmation of the function of ProE, ProI and ProJ. (a) LC-MS analysis of the culture extracts from *AN-proFGIJE* and *AN-proFGIJ*. (b) LC-MS analysis of the culture extracts from *proE* and *proI* knockout mutants in *F. proliferatum*. (c) The *in vitro* biochemical assays of ProE, ProI and ProJ. (d) The proposed pathway from 7 to 1.

pyrone-polyketide glycoside lacking the C7–C8 ester bond (Fig. 4a, Table S16 and Fig. S89–S95[†]). Therefore, ProE is responsible for the conversion of 8 to 1 by catalyzing the BV oxidation between C7 and C8; this observation is further confirmed by the *in vivo* KO *proE* in *F. proliferatum* (*Fp- Δ proE*, Fig. S6[†]) to yield 8 (Fig. 4b(ii)) and the *in vitro* incubation of 8 with ProE to form 1 (Fig. 4c(i and ii)). Moreover, (3) when 7 was incubated with purified FMO ProI (Fig. S9[†]), along with microsome fractions of mGT ProJ and cofactors (FAD, NADPH and UDP-glucose), the formation of 8 was clearly observed (Fig. 4c(iii and iv)), confirming that these two enzymes are indeed responsible for converting 7 to 8; (4) when 7 was incubated with ProJ and UDP-glucose, no products were observed (Fig. 4c(v)), indicating that ProJ does not recognize C4–OH of 7 to perform O-glycosylation, showing a particular preference towards the C3–OH; (5) when 7 was incubated with ProI, a time-dependent decrease in 7 was observed; however, the generation of the proposed C3-hydroxyl derivative of 7 was not observed (Fig. S14[†]), implying that the hydroxylation occurring at the C3 position of the pyrone ring of 7 possibly makes it unstable.^{33,39} We further confirmed that 7 is the substrate of ProI, as the *in vivo* KO *proI* in *F. proliferatum* (*Fp- Δ proI*, Fig. S6[†]) yields 7 (Fig. 4b(iii)). Therefore, based on these results, the route from 7 to 1 by ProIJE was established (Fig. 4d).

β -Glucosidase is involved in the formation of 2 from 1

It is worth noting that, clarification of the biosynthesis process from 7 to 1 confirms that 2 is not a product of this conversion stage, thus it should have been originated from 1. Previous results excluded the involvement of ProG and other enzymes outside the *pro* cluster in this conversion step (Fig. S8 and S11[†]); thus ProK and ProL seem to be the last candidates for the conversion of 1 to 2.

Careful bioinformatic analysis showed that (1) ProL belongs to the glycoside hydrolase family 3 (GH3) of β -glucosidase proteins (Fig. S15[†]);^{40–44} members of this family usually employ an acid/base mechanism to hydrolyze the glycosidic bond (ether bond) of glycosides and oligosaccharides (Fig. S16[†]), and the corresponding catalytic residues⁴³ (aspartate and glutamate) are indeed conserved in ProL (Fig. S15[†]); (2) ProK is a very small protein (142 aa); unlike ProL, it is difficult to predict its function through sequence alignment or structural simulation. Therefore, from a bioinformatic analysis perspective, it seems that neither of these enzymes should catalyze the ester bond hydrolysis process from 1 to 2.

To probe the function of ProK and ProL, *in vivo* KO of these two genes was performed (Fig. S6[†]). Analysis of the metabolic profiles of *Fp- Δ proK* and *Fp- Δ proL* showed that, compared to the *F. proliferatum* wild type (Fig. 5a(i)), (1) the production of 2 and

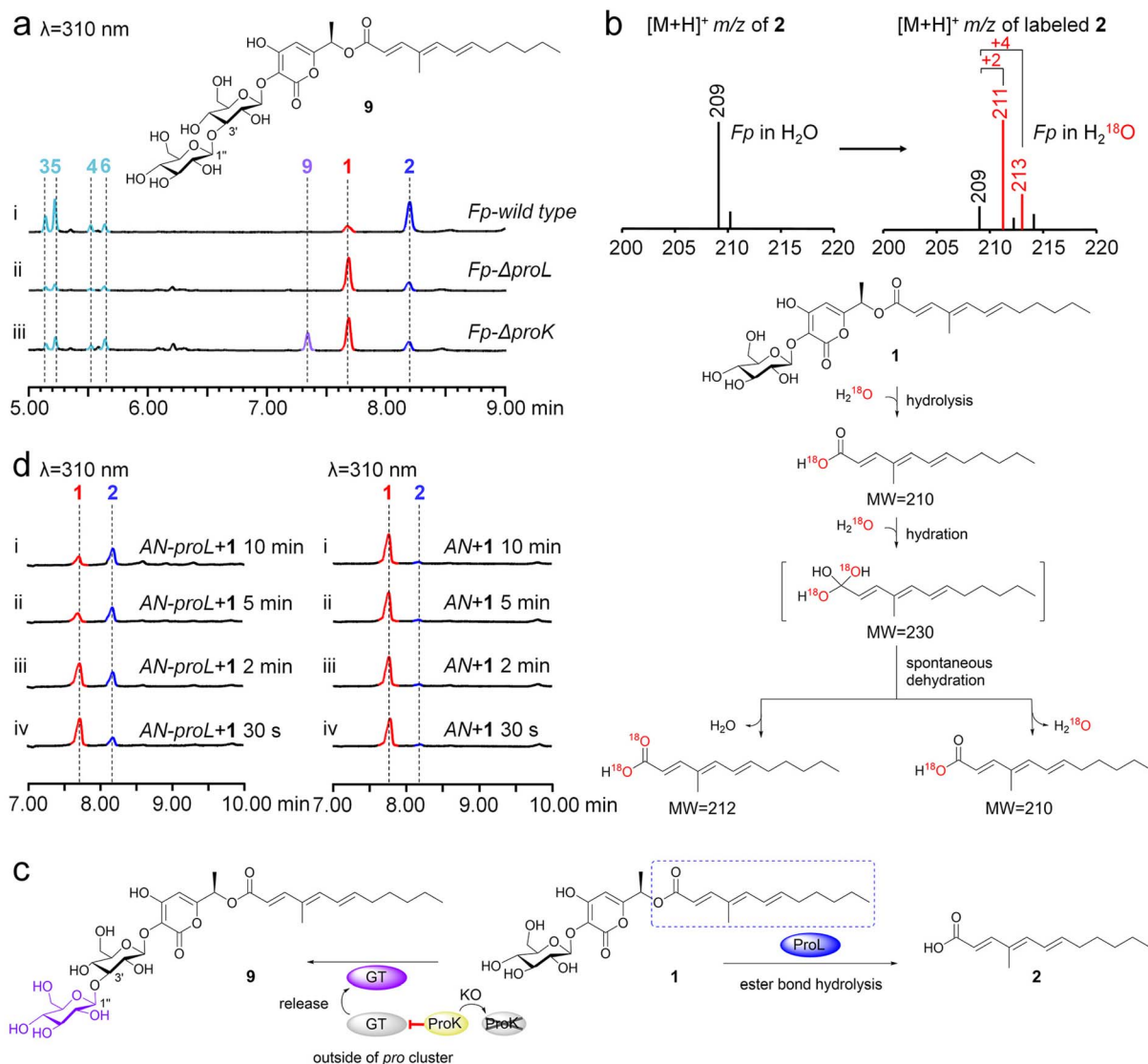


Fig. 5 Investigation of the function of ProL and ProK. (a) LC-MS analysis of the culture extracts from *proL* and *proK* knockout mutants in *F. proliferatum*. (b) LC-MS analysis of the incorporation of H_2^{18}O into 2. (c) The proposed pathway from 1 to 2 and 9, respectively. β -Glucosidase ProL is involved in the hydrolysis of 1 to form 2. (d) The bioconversion of 1 to 2 by the crude enzymes of *AN-proL*.

its oxidized derivatives (3–6) was almost eliminated, while 1 was accumulated in *Fp*- Δ proL (Fig. 5a(ii)); (2) unexpectedly, the conversion efficiency of 1 to 2 was greatly decreased; meanwhile a new compound (9, proliferapyrone D) was produced by *Fp*- Δ proK (Fig. 5a(iii)). Subsequent structural determination through HR-MS and NMR analysis confirmed that 9 features a di-glucose moiety at the C3-OH position; the key HMBC correlation between H1'' and C3' shows that the linkage of di-glucose is through an unusual β -1,3 glycosidic bond (Tables S17, S18 and Fig. S96–S109[†]).⁴⁵ Thus, the di-glucose moiety of 9 is laminaribiose,⁴⁶ which is further confirmed by comparison with the standard after hydrolysis (Fig. S17[†]); moreover, (3) when *F. proliferatum* was incubated in H_2^{18}O -labeled medium, the incorporation of ^{18}O from H_2^{18}O into 2 was successfully observed (Fig. 5b), which confirmed that the hydrolysis reaction indeed occurs during the formation of 2.

These results revealed the following important information (Fig. 5c): (1) the conversion of 1 to 2 is mediated by ProL, where the β -glucosidase-mediated hydrolysis of ester bonds is rarely reported;^{47,48} (2) the production of 9 seems to be the result of an outside *pro* cluster enzyme catalyzing the glycosylation of 1, as the formation of 9 was not observed when 1 was incubated with UDP-glucose and the in-cluster mGT Proj (Fig. S18[†]); (3) the function of ProK is likely to act as an inhibitor of glycosyltransferase(s) outside the *pro* cluster; its deletion may eliminate its inhibitory effect, thereby allowing unknown glycosyltransferase(s) to catalyze the glycosylation of 1 to form 9 *in vivo*. We further searched for homologue proteins of ProK using sequence similarity network (SSN) analysis⁴⁹ (Fig. S19[†]) and found that they are a very small family clade in fungi ($n = 47$), with most of them belonging to the *Fusarium* species, and some ProK homologs are indeed associated with glycosyltransferases

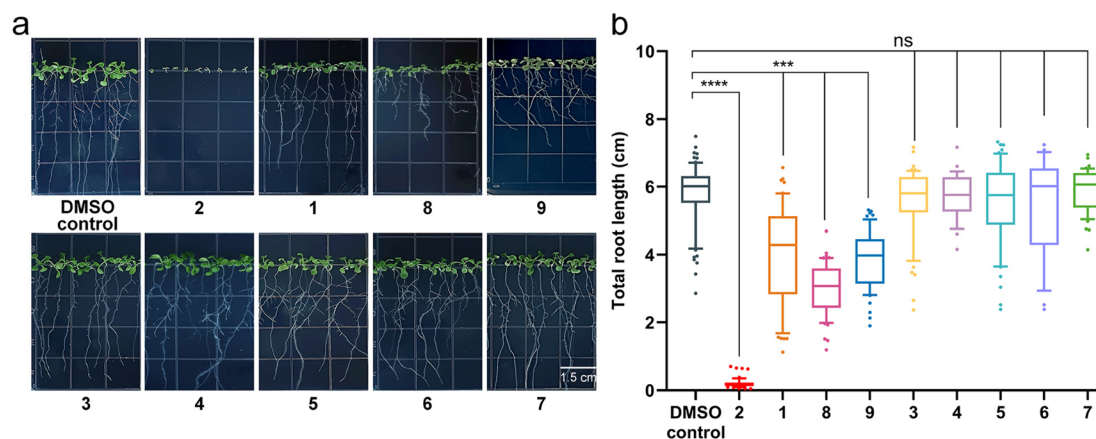


Fig. 6 The root growth inhibition activity tests against *Arabidopsis thaliana* of compounds 1–9. (a) Phenotypic characteristics of seven-day-old *Arabidopsis thaliana* grown on Murashige and Skoog basal medium containing 50 µg per mL 1–9 (DMSO: dimethyl sulfoxide). (b) Quantification of the total root length of seven-day-old *Arabidopsis thaliana* grown on medium containing 50 µg per mL 1–9. In box plots, the center line is the median, box edges delineate first and third quartiles and whiskers show the range of values ($n = 3$ independent experiments), unpaired two-tailed Student's t -test (*** $p \leq 0.001$, **** $p \leq 0.0001$, ns stands for no significant difference).

in natural product gene clusters (Fig. S19[†]). However, the actual function and the relationship between ProK family proteins and glycosyltransferases need further in-depth investigation.

We initially attempted to elucidate the unusual function of ProL in conversion of 1 to 2 through *in vitro* biochemical assays; however, it was insoluble when expressed in *E. coli* (Fig. S20[†]).

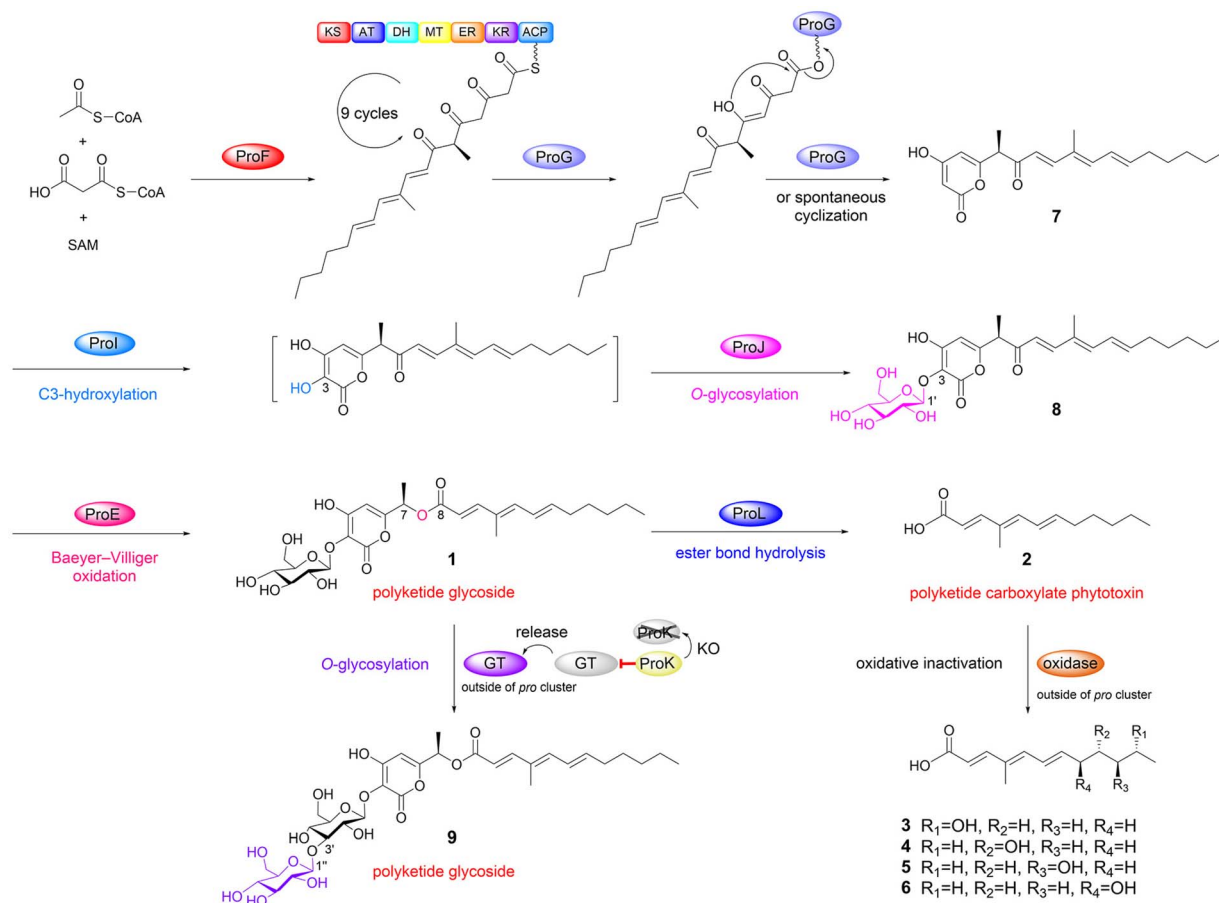


Fig. 7 Formation of polyketide carboxylate 2 from polyketide glycoside 1 occurs through β -glucosidase mediated ester bond hydrolysis.

Alternatively, we decided to confirm this conversion *via* a fungal system. As shown in Fig. 5d, compared to the *AN* wild type, when **1** was incubated with the crude enzymes from *AN-proL*, the production of **2** was observably detected within 10 min, where only trace amounts of **2** were observed in *AN*, suggesting that *A. nidulans* has endogenous enzyme(s) capable of weakly hydrolyzing **1** to **2**. Moreover, when **1** was incubated with the crude enzymes of *AN-proK*, it was indeed inactive (Fig. S21†). We further performed mutagenesis experiments on the conventional active sites D266 and E493 of β -glucosidases in ProL and found that both *AN-proL* D266A and *AN-proL* E493A lost their ability to convert **1** to **2** (Fig. S22†), respectively. Therefore, these results importantly show that the conversion of **1** to **2** is mediated by ProL; unlike traditional β -glucosidases that hydrolyze glycosidic bonds (ether bond),^{40,43} ProL harbors the classical catalytic amino acid residues (D266 and E493) of β -glucosidases; however it has an unexpected ability to mediate the hydrolysis of the ester bond of **1** to form **2**.

Polyketide carboxylate **2** is a phytotoxin

Considering that the core PKS gene of the PKS16 cluster is transcribed in the early stages of *F. fujikuroi* IMI 58289 infection,²⁷ we proposed that the compounds **1–9** identified from the *pro* cluster may be new phytotoxins in *Fusarium* species. As shown in Fig. 6, their phytotoxicity evaluation conducted through the root growth test on *Arabidopsis thaliana*⁵⁰ showed that (1) the polyketide glycosides (**1**, **8** and **9**) have moderate inhibition effects on the growth of roots; however, (2) polyketide carboxylate **2** is a phytotoxin, and it has significant inhibitory activity on the growth of roots; (3) the polyketide precursor **7** and compounds **3–6** are not phytotoxins.

The above activity tests indicate the assembly and inactivation process of polyketide carboxylate phytotoxin **2** in *Fusarium* species, where it is generated from polyketide-glycoside **1** *via* β -glucosidase-mediated hydrolysis of ester bonds and is inactivated by the intracellular oxidase-catalyzed oxidation of the terminal inert carbon atoms to form **3–6**. It should be noted that, the newly discovered compound **2** is a structural analog of the well-known mitotic progression inhibitor myrmicacin;^{51,52} thus, it or its structural derivatives are a potential and worth developing herbicide source in the future.

Conclusions

In summary, we discovered the intact PKS16 homologue clusters from various *Fusarium* strains by genome mining and identified gene functions of the representative *pro* cluster from *F. proliferatum*. A plenty of α -pyrone-polyketide glycosides and polyketide carboxylate compounds were characterized from this cluster, importantly revealing an unusual assembly and inactivation process in phytotoxin biosynthesis (Fig. 7). The newly discovered polyketide carboxylate **2**, which is generated from the α -pyrone-polyketide glycoside **1** *via* β -glucosidase-mediated hydrolysis of the ester bond, serves as a new phytotoxin that can significantly inhibit the root growth in *Arabidopsis thaliana*. Our work identifies the chemical substance basis of the long-term

unsolved PKS16 cluster in phytotoxicity, shows that polyketide carboxylate is a new structural type of phytotoxins in *Fusarium* and importantly uncovers a rare ester bond hydrolysis function of β -glucosidase family enzymes.

Data availability

Materials and methods, additional tables and figures, spectroscopic data and the sequence data of the *pro* gene cluster from *Fusarium proliferatum* CGMCC 3.4710 are available in the ESI.† The DNA sequence of the *pro* cluster has been deposited in the NCBI GenBank with the accession number PQ271635.

Author contributions

X. W. and D-K. K. performed the *in vivo* and *in vitro* experiments, as well as compound isolation. X. W. performed bioactive assays. X. W. and H-R. Z. performed compound characterization. All authors analyzed and discussed the results. Y. Z. supervised the research and wrote the manuscript.

Conflicts of interest

There are no conflicts to declare.

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