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# Evidence for enzyme catalysed intramolecular [4+2] Diels–Alder cyclization during the biosynthesis of pyrichalasin H†

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Cytochalasans are highly complex fungal metabolites which exhibit diverse biological activities. Little is known of the chemical steps involved in the construction of the tricyclic core, which consists of an octahydro-isoindole skeleton fused to a macrocyclic ring. Here, using a directed gene knockout and complementation strategy, we show that PylF is implicated as the proposed intramolecular [4+2] Diels–Alderase required for construction of the tricyclic core of pyrichalasin H 1.

Cytochalasans are a diverse group of fungal polyketide-non-ribosomal peptide secondary metabolites with varied biological properties including antibacterial, antiviral and antitumor potential.<sup>1,2</sup> All cytochalasans contain a tricyclic core, which consists of a macrocyclic ring fused to an amino acid-derived octahydro-isoindole. Differences arise from variations in the size and the substitution pattern of the macrocycle and the incorporated amino acid (Fig. 1).<sup>3</sup>

Several biosynthetic gene clusters (BGC) involved in cytochalasan biosynthesis are known, and their biosynthetic pathways have been partially elucidated, including those for pyrichalasin H 1,<sup>4</sup> cytochalasin E 2,<sup>5</sup> chaetoglobosin A 3<sup>6</sup> and compounds such as niduporthin 4<sup>7</sup> and the ACE1 shunt 5<sup>8</sup> which are genetically and chemically closely related. All cytochalasan BGC encode: a polyketide synthase/non-ribosomal peptide synthetase (PKS-NRPS); a *trans*-acting enoyl reductase (ER); an  $\alpha\beta$ -hydrolase (HYD); several tailoring enzymes such as cytochrome P450 monooxygenases (P450) and oxidoreductases (OXR); and transcriptional regulators (TF) and transporters. Recently, we identified the *pyi* BGC encoding the phytotoxin pyrichalasin H 1 in *Magnaporthe grisea* NI980 (Scheme 1) and elucidated the late-stage biosynthetic pathway of 1 using targeted gene inactivations.<sup>4</sup>

Formation of the tricyclic core of 1 and other cytochalasans is thought to occur in two steps. First, it is proposed that the

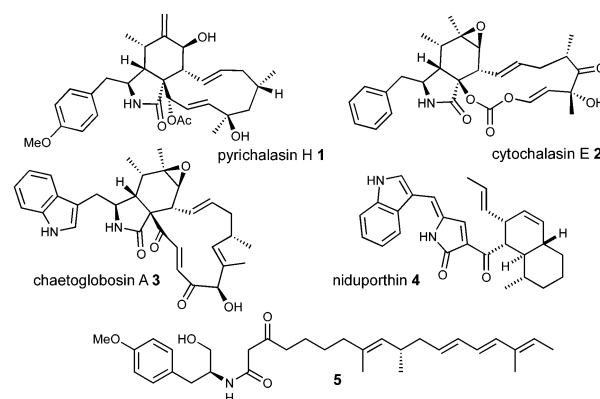


Fig. 1 Chemical structures of selected cytochalasans and related metabolites.

PKS-NRPS reductively releases an *N*-acyl  $\alpha$ -amino aldehyde 9 which is cyclised to form a 1,5-dihydropyrrone 10. This is then proposed to undergo an *endo*-selective intramolecular [4+2] Diels Alder (DA) reaction (Scheme 1).<sup>9</sup> Intermediates 9 and 10 have not been observed for pyrichalasin H 1 or related cytochalasans, although similar  $\alpha$ -aminoaldehyde and pyrrolone intermediates have been proposed during the biosynthesis of fusarin C, for example.<sup>3,10</sup> In other related systems non-reductive release (*i.e.* Dieckmann cyclisation) occurs during the biosynthesis of, *e.g.*, pretenellin A.<sup>11</sup> The heterocyclic core seems to be formed by the PKS-NRPS and *trans*-ER alone in the cases of niduporthin 4<sup>7</sup> and cytochalasans from *Parastagonospora nodorum*.<sup>12</sup>

The assumption that the DA reaction is not spontaneous is supported by observations in the total synthesis of cytochalasins.<sup>13</sup> Stork and Nakamura developed the first biomimetic macrocyclization strategy in 1982 while synthesizing cytochalasin B. To achieve the DA reaction a precursor analogous to 10 had to be heated to 180–190 °C in neat mesitylene for nearly a week to yield 35% product (4 : 1 mixture of *endo* : *exo*), strongly suggesting that the DA reaction is not favoured under physiological conditions.<sup>14</sup>

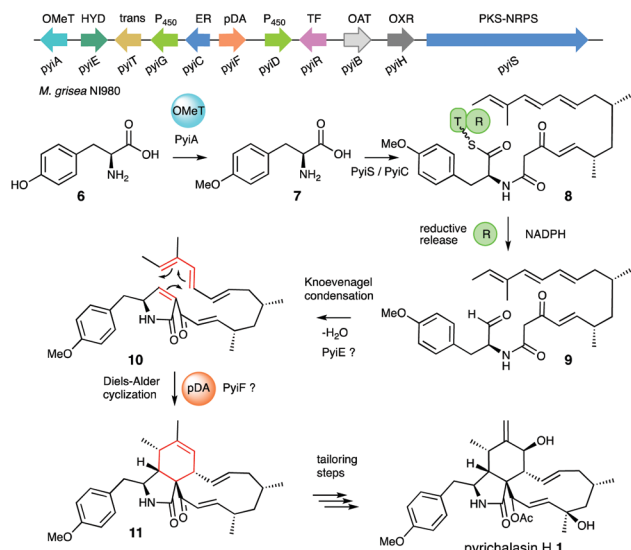
Enzymatic DA<sup>15</sup> and hetero DA (hDA)<sup>16</sup> catalysts have been discovered and validated in several biosynthetic pathways. However, while the action of a DA enzyme has been frequently

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Scheme 1 Proposed early biosynthetic steps of pyrlichalasin H 1.

proposed as the key element of cytochalasan pathways no evidence has been so far gathered for the identity of this catalyst. In the case of pyrlichalasin H 1 we have recently demonstrated the function of all the genes in its BGC except *pyiE* and *pyiF*.<sup>4</sup> Homologs of these two genes are present in all known cytochalasan BGC and they may thus be involved in the formation of the isoindole. *PyiE* is a member of the  $\alpha\beta$ -hydrolase family,<sup>17</sup> but *PyiF* and its homologs have no known function. Here we investigated the function of *PyiF* and homologs from other systems by knockout (KO) and complementation in *M. grisea* NI980.

Disruption of *pyiF* was achieved by replacing it with a hygromycin resistance cassette.<sup>18</sup> Wild-type (WT) *M. grisea* NI980 produces **1** at around 60 mg L<sup>-1</sup> (Fig. 2A), but inactivation of *pyiF* resulted in the abolition of **1** (Fig. 2B).

We expected knockout of *pyiF* to result in an accumulation of the pyrrolone **10** ( $m/z = 447$  g mol<sup>-1</sup>) but this was not observed. Instead the organic extracts of  $\Delta$ *pyiF* strains contained several new compounds which, based on their UV absorption spectra at 275 nm (see ESI<sup>†</sup>), derived from tyrosine **6** or *O*-methyl tyrosine **7** (Fig. 1B). Eight compounds (**15**, 0.4 mg; **21**, 0.3 mg; **20** and **17** (mixture, **20**:**17** ca. 2:1), 0.5 mg; **19** and **14** (mixture, **19**:**14** ca. 1:1.2), 1.5 mg; **16**, 0.5 mg; and **18**, 1.0 mg) were purified by preparative LCMS from a 1 litre  $\Delta$ *pyiF* *M. grisea* culture after 7 days.

The molecular formula for each compound was obtained (HRMS) and structure elucidation (NMR, see ESI<sup>†</sup> for full NMR and HRMS data) was then performed. In the case of the major compound **14**, HRMS indicated a molecular formula of C<sub>29</sub>H<sub>43</sub>NO<sub>6</sub> (obs. 546.3073, calc., 546.3067, ES<sup>-</sup>, [M + HCO<sub>2</sub>H-H]<sup>-</sup>) consistent with the skeleton of **1**. HMBC correlations from the methoxy group ( $\delta_{\text{H}}$  3.7 ppm) to C-27, supported by HMBC from H-25 and H-26, confirmed the presence of the *p*-substituted aromatic ring and also located benzylic C-23 ( $\delta_{\text{C}}$  39.7 ppm). COSY correlation from H-23 via H-5, H-4 and H-3 led to methine H-6, and HMBC from H-4 and H-5 located C-2 ( $\delta_{\text{C}}$  177.2 ppm) and

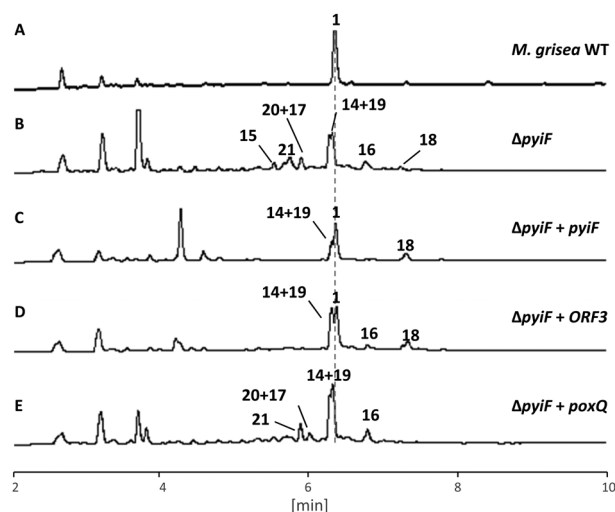


Fig. 2 LCMS traces (diode array) for key experiments: (A) *M. grisea* wild type (WT); (B) *M. grisea*  $\Delta$ *pyiF*; (C) as B, complemented with *pyiF*; (D) as B, complemented with *ORF3*; (E) as B, complemented with *poxQ*. Unnumbered peaks are unrelated.

confirmed the presence of the  $\gamma$ -lactam. Further COSY correlations then located the 7/8 olefin, and methine H-9 and methylenes H-10. HMBC correlations to methyls 21 and 22 located carbons 8–12, and further COSY correlations then revealed the diene system to C-16. Finally, HMBC from methyls-19 and 20 revealed C-17 as a tertiary alcohol and C-18 as a secondary alcohol (Scheme 2B). The related compounds were identified by comparison and close analysis of HMBC signals, particularly around the  $\gamma$ -lactam.

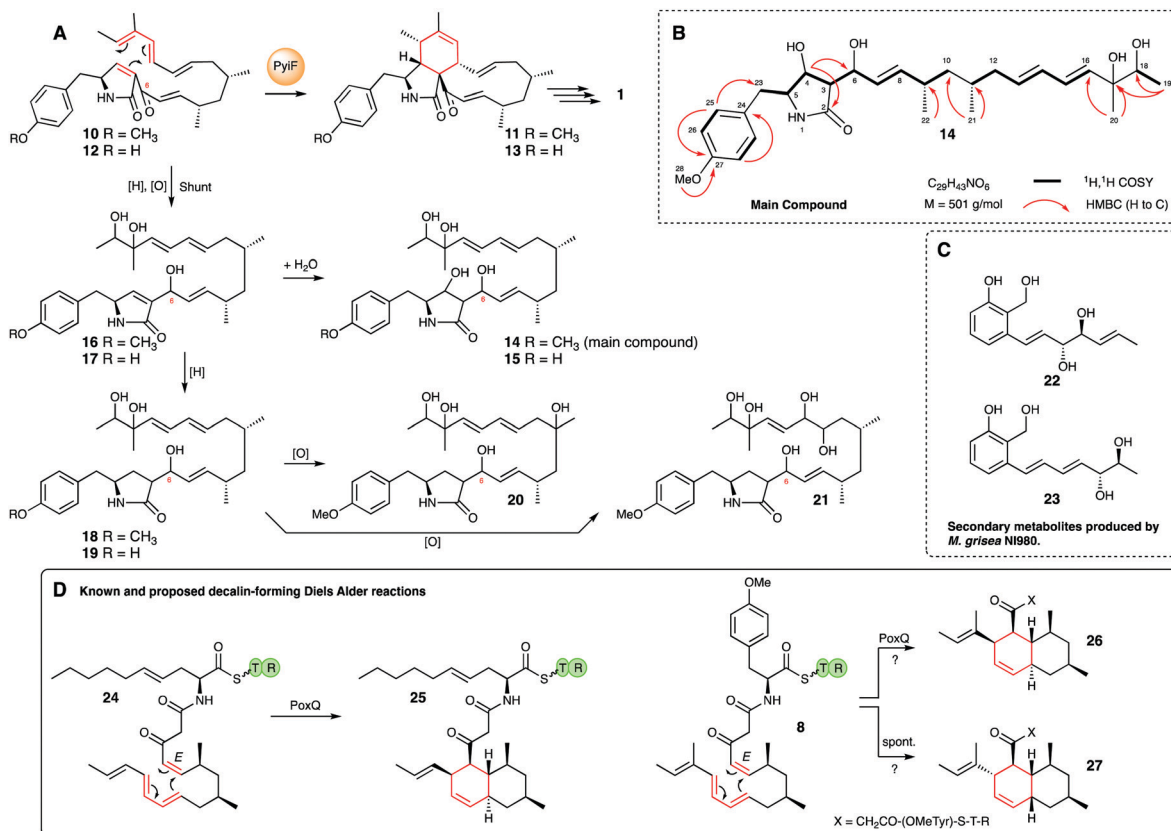
All isolated compounds displayed an underlying linear structure of the polyketide moiety. The C-17/C-18 olefin was dihydroxylated in all compounds, probably via epoxide formation and hydrolysis. Genome analysis of *M. grisea* showed it contains the dihydropyriculol/dihydropyriculariol **22/23** BGC<sup>19</sup> and dihydropyriculol **22** was purified and characterised from *M. grisea* NI980 producing strains (see ESI<sup>†</sup>). The same oxygenase responsible for the formation of the diols in **22/23** could give the diols observed in **14–21**.

Compound **21** also features C-13/C-14 olefin dihydroxylation; compounds **14** and **15** are hydroxylated at C-4; and compound **20** is hydroxylated at C-11. Another deviation from the expected pyrrolone **10** is that all compounds carry a hydroxyl at C-6 instead of a carbonyl. Furthermore, the C-3/C-4 olefin of **10** (part of the pyrrolone ring) is missing in all compounds, except **16** and **17**. Five compounds (**14**, **16**, **18**, **20**, **21**) derive from *O*-methyl tyrosine **7**; the others from tyrosine **6**. The observed compounds appeared to be shunt products of **10** (or its 27-*O*-desmethyl cogener) formed via oxidative and reductive steps.

A functional copy of *pyiF* was then introduced downstream of the inducible *amyB* promoter (*P<sub>amyB</sub>*) from *Aspergillus oryzae* and transformed into two  $\Delta$ *pyiF* *M. grisea* strains.

As expected, in all strains carrying an ectopic functional copy of *pyiF*, the production of **1** was restored. However, the obtained yields of **1** were lower compared to the wild type (ca. 10%). In addition, some of the shunt intermediates (**14–21**) observed in





Scheme 2 Putative Diels–Alder reaction catalysed by PyiF and isolated shunt products (**14–21**) purified from a  $\Delta$ *pyiF* *M. grisea* strain, and related chemistry.

the  $\Delta$ *pyiF* *M. grisea* deletion strain were also detected in most of the transformants (mainly compound **14**, Fig. 2C).

Heterologous complementation studies were carried out to investigate if the closely related homologous protein ORF3 (62% identity to PyiF, see ESI†) from the cryptic *ACE1* BGC in *P. oryzae* Guy11,<sup>20</sup> is able to restore production of **1**. The *pyi* and *ACE1* gene clusters are closely related, suggesting that ORF3 might be able to replace PyiF as a DAase. This was confirmed when  $\Delta$ *pyiF* *M. grisea* strains carrying *ORF3* driven by *P<sub>amyB</sub>* restored the production of **1**, albeit in similar titres compared to the homologous complementation with *pyiF* (ca. 10%, Fig. 2D). No new compounds were observed.

An analogous complementation experiment was also carried out with the *poxQ* gene from *Penicillium oxalicum*. PoxQ is proposed to be a decalin-forming DAase (29% identical to PyiF) during the biosynthesis of oxaleimide.<sup>21</sup> PoxQ and related enzymes<sup>22</sup> are proposed to intercept an intermediate **24** in which the polyketide moiety is nearly identical to that proposed in the case of pyrichalasin H **1** (e.g. **8**, Scheme 2D). If the PoxQ protein recognised **8** then decalins such as **26** might be expected. However, all *M. grisea*  $\Delta$ *pyiF*:*P<sub>amyB</sub>*:*PoxQ* transformants demonstrated the same secondary metabolite profile as the *M. grisea*  $\Delta$ *pyiF* control strain (Fig. 2E). Accordingly, the production of **1** could not be restored and no new decalin-containing compounds were produced.

Finally, in an attempt to demonstrate *in vitro* activity of PyiF very extensive efforts were made to obtain soluble PyiF protein *via* heterologous expression. However this proved impossible.

In conclusion, targeted gene disruption of *pyiF* in *M. grisea* resulted in complete abolition of **1**, and the accumulation of **14–21** which appear to arise from a redox shunt pathway from **10** (Scheme 2). Compound **16** probably arises by reduction of the C-6 carbonyl and likely epoxidation and hydrolysis of the C-17/18 olefin. Conjugate addition of water at C-4/5 could then give the main compound **14**. Further reduction of **16** leads to compound **18**, and compounds **20** and **22** are oxidation products of **18**. Dihydroxylation of the triene sidechain may be catalysed by enzymes from the pyriculol pathway, known to be present and active in *M. grisea*. Compounds **15**, **17** and **19** lacking the *O*-methyl likely arise from use of tyrosine by the PyiS NRPS.<sup>18</sup>

All compounds isolated from the  $\Delta$ *pyiF* mutant lack the cyclohexenyl moiety indicating that the DA reaction has not occurred. This, therefore, shows that PyiF is the likely DA enzyme required for the formation of the core cytochalasan isoindole. Complementation studies showed that the closely related ORF3 (62% identical) from the cryptic *ACE1* pathway is also a DA enzyme, and since all known cytochalasan BGCs contain a homolog of *pyiF*/*ORF3*, the results suggest this is the likely function of these genes. The results of this experiment



also add to the growing evidence that the cryptic ACE1 metabolite is a cytochalasan.<sup>18</sup>

It is notable that the titres of the DA products from the complementation experiments are lower than the WT titres of **1**. This most likely arises because the *A. oryzae* *P<sub>amyB</sub>* promoter, used to drive the expression in these cases, is not strong when used in *M. grisea*.<sup>23</sup> Lebrun and coworkers showed in a previous study that ORF3 is the most highly expressed gene in the ACE1 BGC, perhaps indicating a requirement for relatively high protein concentrations for rapid DA reaction.<sup>24</sup>

Heterologous complementation of the  $\Delta$ *pyiF* mutant with *poxQ* from *Penicillium oxalicum* did not restore the production of **1**, or produce a decalin metabolite. PoxQ is proposed to catalyse a decalin forming DA reaction of a near-identical polyketide intermediate (e.g. **8**) during the biosynthesis of **1** (Scheme 2D).<sup>19</sup> This may indicate that PoxQ and related decalin-forming catalysts<sup>20</sup> recognise more than just the polyketide intermediates, or that the triene of **8** was rapidly destroyed by olefin oxidation before PoxQ could act. Intermediates such as **8** are also known to undergo spontaneous decalin formation (albeit with different stereoselectivity)<sup>20</sup> and rapid conversion of **8** to **14** in *M. grisea* could also explain the lack of spontaneous decalin formation.

Collectively, our results strongly suggest that PyiF and ORF3 catalyse the long-proposed intramolecular DA reaction forming the tricyclic core structure of cytochalasans. For the first time the involvement of a DA enzyme in the biosynthesis of cytochalasans has been experimentally confirmed. DA enzymes involved in the biosynthesis of cytochalasans belong to the stand-alone class of pericyclases with no additional function besides catalysing the [4+2] cyclisation. Whether this reaction requires conformational or chemical catalysis is unknown and will require *in vitro* studies of soluble protein which is not yet available. To gain more insights into the biosynthetic role of the hydrolase PyiE, *in vitro* studies are underway and results of these experiments will be reported in due course.<sup>25</sup>

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## Conflicts of interest

There are no conflicts to declare.

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- 25 Full experimental details are supplied in the ESI†.

