Biosynthesis

Chemical and Genetic Studies on the Formation of Pyrrolones During the Biosynthesis of Cytochalasans

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Abstract: A key step during the biosynthesis of cytochalasans is a proposed Knoevenagel condensation to form the pyrrolone core, enabling the subsequent 4+2 cycloaddition reaction that results in the characteristic octahydroisoindoline motif of all cytochalasans. In this work, we investigate the role of the highly conserved α,β-hydrolase enzymes PyiE and ORFZ during the biosynthesis of pyrichalasin H and the ACE1 metabolite, respectively, using gene knockout and complementation techniques. Using synthetic aldehyde models we demonstrate that the Knoevenagel condensation proceeds spontaneously but results in the 1,3-dihydro-2H-pyrrol-2-one tautomer, rather than the required 1,5-dihydro-2H-pyrrol-2-one tautomer. Taken together our results suggest that the α,β-hydrolase enzymes are essential for first ring cyclisation, but the precise nature of the intermediates remains to be determined.

Introduction

Cytochalasans are secondary metabolites that have been isolated from a wide range of fungi and that possess numerous potent bioactivities extensively reviewed elsewhere. [1, 2] As might be expected for metabolites that have been known since the mid-1960s, [3] their biosynthesis has been extensively studied, [4, 5] and their intriguing structures have been the target for numerous total synthesis campaigns. [6]

We have investigated the biosynthesis of pyrichalasin H 1 from the fungus Magnaporthe grisea (Scheme 1). [7] The results to-date are consistent with an overall pathway in which O-methyl tyrosine 2 is created by PyiA and used as the initial building-block by the polyketide synthase non-ribosomal peptide synthetase (PKS-NRPS) PyiS which acts in concert with the trans-enoyl reductase (ER) PyiC. PyiS has a reductive release system and it is hypothesised that an aldehyde intermediate

Scheme 1. Comparison of the pyi and ACE1 BGC and proposed biosynthesis of metabolites encoded by these BGC.

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such as 3 is released. Cyclisation of 3 by a Knoevenagel process is then proposed to give 4,[8] and Diels–Alder (DA) reaction could then give the observed cytochalasan skeleton of 5. The function of tailoring enzymes PyiBDGH has been proven using gene knockout (KO) and heterologous expression experiments during the formation of the mature metabolite 1.[19] Meanwhile, pyiR encodes a transcription-factor which we have shown to be responsible for increasing the titre of 1.[10] Recently we showed that PyiR is involved in the proposed DA reaction.[11,12]

These observations leave PyiE, which encodes a putative hydrolytic enzyme, to be proposed as the Knoevenagel catalyst by default since all other proteins encoded by the pyrichalasin biosynthetic gene cluster (pyi BGC) have proven roles. In addition, compounds 3 and 4 (or their analogues in other pathways) have never been directly observed, so uncertainty remains regarding the early biosynthetic steps.

We have also investigated the parallel ACE1 biosynthetic pathway[13] in the closely related fungus *Pyricularia oryzae* (Scheme 1). *P. oryzae* causes rice blast disease and the ACE1 metabolite is involved in avirulence signalling between the fungal pathogen and resistant rice plants. The ACE1 BGC is only active in single cell appressoria for a matter of hours during initial fungal penetration of plant leaves, so the identity of the ACE1 metabolite is not known. It has also been impossible, for the same reasons, to perform productive KO experiments in the ACE1 system.[14] However, the high gene-for-gene homologies between the ACE1 and *pyi* genes, and the fact that *ACE1* genes can complement knockouts in the *pyi* BGC mean that the ACE1 metabolite is very likely to be a cytochalasan derived from 2. Heterologous expression experiments[15] have also reinforced this hypothesis. Expression of *ACE1* (PKS-NRPS) and *RAP1* (trans-ER) in *Aspergillus oryzae* lead to the production of various shunt compounds such as 6a and 6b in the absence of 2,[15] and 6c when 2 is supplied. It is assumed that these are derived by rapid reduction of putative aldehyde intermediates such as 7.[16] Interestingly, putative intermediate 7 differs from the proposed pyrichalasin aldehyde intermediate 3 by being two carbons longer showing that the *ACE1* PKS is differently programmed.

Hydrolitic enzymes similar to PyiE, and its *ACE1* homolog ORFZ, are encoded in the BGC of all known cytochalasans,[17] as well as in other BGC which encode the biosynthesis of compounds such as fusarin C where the PKS-NRPS has a reductive release mechanism.[18] However, the precise function of these hydrolitic enzymes remains unknown. As part of ongoing studies into the biosynthesis of cytochalasan we therefore decided to investigate PyiE from *M. grisea* and ORFZ from *P. oryzae* in more detail.

**Results**

**Knockout of pyiE**

Disruption of *pyiE* in *M. grisea* was achieved by insertion of a hygromycin resistance cassette using the bipartite method of Neilsen and co-workers.[19] Examination of organic extracts from *M. grisea ΔpyiE* showed that, in comparison to extracts from wild-type (WT) *M. grisea*, production of 1 was significantly reduced, concomitantly with an increase in production of known pyriculols.[20] We then re-introduced a copy of either *pyiE* itself or its homolog ORFZ, under the control of the *Aspergillus oryzae* amyB promoter (*PamyB*), into the KO strain using previously reported methods.[21] In previous work we had shown that *PamyB* is functional in *M. grisea* and we therefore expected that biosynthesis of 1 should be restored to WT titres.[21] However, LCMS analysis showed that 1 was not produced in significantly increased titre for either complementation experiment (Figure 1 and Figures S6.1–S6.6).

The LCMS profiles obtained from these strains were similar to those we had observed during KO of *pyiA*.[21] Since the target gene *pyiE* is adjacent to *pyiA* (Scheme 1) we considered it possible that either *pyiA* itself, or the putative bidirectional promoter region between *pyiE* and *pyiA*, may have been damaged by the *pyiE* engineering. We tested this possibility by feeding O-methyl tyrosine 2 to cultures of *M. grisea ΔpyiE*. *M. grisea ΔpyiE* *amyB*-*PamyB*/*pyiE* and *M. grisea ΔpyiE PamyB* ORFZ. In the case of the *M. grisea ΔpyiE PamyB*/*pyiE* and *M. grisea ΔpyiE PamyB* ORFZ strains production of 1 was substantially restored (Figure 1 and Figures S6.1 to S6.6).

In the *ΔpyiE*+2 experiments a major new peak eluting at 5.1 minutes was also observed (Figure 2). Mass data corresponded to the pre-Diels–Alder intermediates or shunt metabolites. Mass spectrometric analysis indicated a mass of 519 Da, also consistent with compounds related to 1 (e.g. 10, m/z 521). This compound was detected in both WT and the *ΔpyiE* mutant, but in much lower amounts (Figure S6.9). The major isomer eluting at 5.1 min was purified by mass-directed preparative reverse-phase chromatography. HRMS indicated a formula of C$_27$H$_{30}$NO$_5$ (observed 542.3096, calculated 542.3094 for C$_27$H$_{29}$NNaO$_5$) consistent with the 29 main-chain carbons of 1. Full NMR analysis (Supporting Information, section S7) showed the compound to be 9, confirming it as a pre-cyclisation shunt compound. Reduction of the expected C-21 aldehyde to a primary alcohol is consistent with the known propensity of fungi to reduce free aldehydes.[15,16a]

![Figure 1. Production of pyrichalasin H 1 in WT M. grisea N980 and selected mutants. UV at 275 nm.](image-url)
Dihydroxylation of the skeleton of 9 at C-15/C-16 and hydroxylation at C-9 are consistent with modifications previously observed in hunt metabolites (e.g. 10) in M. grisea, for example during KO studies of the Diels–Alderase gene pyiF. The structural assignment was further confirmed by observation of a major mass fragment at m/z 364 corresponding to a facile homo-allylic fragmentation of the polyketide triol sidechain (Figure 3).

Heterologous expression of ACE1, RAP1 and ORFZ in Aspergillus oryzae

Previously we had observed that when we co-expressed ACE1 + RAP1 in A. oryzae, linear alcohol shunt products 6a–c were observed rather than expected aldehyde 7 (Scheme 1). To examine whether the inclusion of the corresponding α,β-hydrolase ORFZ prevented reduction and enabled biosynthesis of the expected aldehyde 7 and ring-closed congeners analogous to 4, we co-expressed ACE1, RAP1 and ORFZ in A. oryzae using previously reported methods. However, the same shunt intermediates 6a–c were observed in the absence of exogenous 2 (Figure S6.7), while 6c was produced in the presence of 2 (Figure S6.8). To investigate the possibility that the Diels–Alderase, ORF3 is also required, we co-expressed ACE1, RAP1, ORFZ and ORF3 in A. oryzae. However, the same shunt intermediates 6a–c were again observed (Figure S6.7). PCR and RT-PCR analyses showed that all transformed genes were being expressed in A. oryzae, and previous complementation studies had shown that ORFZ and ORF3 were correctly cloned. We therefore concluded that aldehyde reduction out-competes any ring-closing in this host. Realising that we could not investigate the function of PyiE or ORFZ in vivo, we decided to investigate the Knoevenagel reaction in vitro.

Chemical investigation of Knoevenagel step

In order to investigate the chemistry of the proposed Knoevenagel step we synthesised the model compound 11a which features a phenylalanine-derived aldehyde similar to proposed intermediate 3, but with a truncated β-keto polyketide designed for simpler synthesis and analysis. Thus, amino-alcohol 2S-12 was treated with acetone diketene adduct 13, to give 14 and oxidised under mild Pfitzner-Moffatt conditions to give the required aldehyde 11a. 11a was treated with 1M NaOH in an attempt to generate 12a (Scheme 2). However, in aqueous solution 12a rapidly equilibrates to its 1,3-dihydro tautomer 12b. This structure is supported by: chemical shift data, especially for the C-13 methyl (δH 2.15 ppm); the chemical shift of H-4 (δH 5.54 ppm); the chemical shifts of C-12 and C-3 (δC 137.3 ppm).

Scheme 2. Synthesis of model pyrrol-2-one 12. Reagents and conditions: (a) toluene, 100 °C; (b) EDCI, ClCH2CO2H, DMSO:toluene 1:1, RT, 16 h; (c) 1M aq. NaOH, 0 °C, 5 min.
172.3 and 106.9 ppm, respectively); the observation of a distinct UV absorption (329 nm) corresponding to the aromatic heterocycle; and loss of optical rotation (e.g. for 20a \( [\alpha]_D = -12.0 \) (c = 0.9, CH\(_2\)Cl\(_2\)); for 23, \( [\alpha]_D = -16.9 \) (c = 0.3, MeOH); for 12 \( [\alpha]_D = 0 \).

As pyrchalasin H 1 is derived from O-methyl tyrosine 2 we also synthesised a \( p \)-methoxy derived aldehyde 20a (Scheme 3). L-Tyrosine 15 was protected as its N-methyl carbonate (MOC) and then methylated to give 16 (Scheme 3). LiAlH\(_4\) reduction led to 17 and saponification then gave amino-alcohol 18. Treatment with acetonide diketene adduct 13 under thermolytic conditions gave 19. Modified Pfitzner–Moffatt oxidation of 19 resulted in the desired compound 20a, which once again equilibrated with its hydrate form 20b in water. Aldehyde 20a, however, was not easy to purify by flash chromatography, although mg amounts could be purified by reverse-phase HPLC. In an alternative route, Moffatt oxidation of 17 earlier in the synthesis was followed by formation of the corresponding diethyl acetal to give 21. Then basic removal of the MOC group to 22 and attachment of the N-acetoacetyl group using acetonide diketene adduct 13 gave 23 which could be prepared and purified at larger scale. This material could be deprotected under mild aqueous acid conditions, followed by neutralisation for convenient production of 20a in situ.

Once again, brief base treatment achieved cyclisation to give the 1,3-dihydro-2H-pyrrol-2-one tautomer 24b. This compound proved significantly less stable than the phenylalanine analogue 12b and it was difficult to purify and obtain full NMR data. In aqueous solution 24b rapidly reacted further to give a co-eluting mixture of two new compounds 25 with \( m/z \) 263 which were more stable and which could be purified and fully characterised. Full NMR analysis showed these compounds to be a pair of diastereomers based on the 5-hydroxy-pyrrolid-2-one skeleton, presumably derived by addition of water to 24b (Scheme 4).

In order to obtain further evidence that the obtained heterocycles exist predominantly as their 1,3-dihydro-2H-pyrrol-2-one tautomers we synthesised a model compound unable to tautomerise in this way. Briefly, \( p \)-methoxyaniline 26 was diazotized at low temperature, and the resulting diazonium salt 27 was treated with methallyl alcohol 28 in the presence of iron sulfate to give diazo compound 29 (Scheme 5). AZn/HCl reduction then gave the amino alcohol 30, and the usual sequence of acetoacetylation to 31, Moffatt oxidation to 32 and aqueous base cyclization gave the expected 1,2-dihydro-2H-pyrrol-2-one tautomer 33. This differed from the non-methylated compound 12b in lacking a distinct UV absorption for the pyrrole (e.g. \( \lambda_{\text{max}} \) shift from 329 to 275 nm); having a \( \Delta\delta \) for H-4 of 7.71 rather than 5.54 ppm; different carbon chemical shifts for C-12 and C-3 (\( \Delta\delta \) 194.3 and 135.2 ppm); and a more obvious methyl-13 ketone resonance (\( \Delta\delta \) 2.51 ppm).

With synthetic aldehydes in hand we next set out to study their behaviour under aqueous conditions, and in the presence of preparations of the putative Knoevenagel catalyst ORFZ. The native \( M.\ oryzae \) ORFZ gene was cloned into the expression vector pET28a(+), confirmed by sequencing. The plasmid was transformed into \( E.\ coli \) BL21 (DE3) and ORFZ was expressed by addition of IPTG to a final concentration of 0.1-

\[ \text{Scheme 3. Synthesis of \( p \)-methoxy model compounds. Reagents and conditions: (a) MOCI, THF, H}_2\text{O, NaHCO}_3\text{; (b) MeI, K}_2\text{CO}_3, \text{DMF;} \ (c) EDCI, Cl}_3\text{CHCO}_2\text{H, DMSO:toluene 1:1, RT, 16 h;} \ (d) (EtO)\text{Cl}_3\text{CH, EtOH, p-TsOH, 50 °C, 16 h;} \ (e) 25\% \text{KOH, MeOH, H}_2\text{O, 50 °C, 12 h;} \ (f) \ 13, \text{toluene 100 °C, 2 h.} \]

\[ \text{Scheme 4. Formation of hemiaminal diastereomers 25ab.} \]
1 mM (Figure S3.1). His$_6$-ORFZ was isolated from the soluble protein fraction using Ni-NTA chromatography and the purified protein was analysed by ESI Q-TOF to confirm the correct sequence.

### In vitro assays

Aldehyde **20a** was prepared at a concentration of ca 0.5 mM in water and incubated at 28°C. The composition of the solution was monitored by analytical LCMS. Under these conditions **20** appeared to be relatively stable as the hydrate form **20b** (Figure 4), but cyclised slowly to form the previously characterised alcohols **25ab** (ca 10% after 17 h). Aldehyde **20** was then tested in buffer solutions. Buffers such as Tris which contain a free amine are unsuitable for use with aldehydes so phosphate buffer was initially tested at pH values between 6.0 and 8.0. Under these conditions **20** behaved similarly to its reactions in water (Figure 5). However a small amount of an additional new compound **34** with m/z 522 was observed after 23 h. In phosphate at pH 7.0 a much higher proportion of alcohols **25** were observed and the new compound with m/z 522 also increased significantly (Figure 5).

At pH 8.0 in phosphate buffer aldehyde **20** appeared to be quantitatively converted to the m/z 522 compound after 24 h (Figure 6). A mass of 522 corresponds to a potential dimerisation and oxidation of **25**. The experiments were therefore repeated under an atmosphere of N$_2$. Under these conditions the m/z 522 compound was no longer observed, but at pH 8.0, for example, complete conversion to the alcohols **25** was achieved (see Supporting Information, section S8).

New compound **34** was purified. Full NMR analysis confirmed this to be an oxidised dimer of the alcohol **25**. A series of intra-unit HMBC correlations (Figure 7) confirmed the presence of two different monomers, and a series of inter-unit...
HMBC correlations confirmed the carbon-carbon linkage shown in Figure 7. Chemical shift considerations suggest that carbons L and J (Figure 7) are linked via oxygen rather than the amide nitrogen.

Further investigations into buffers indicated that HEPES buffer induced the minimum formation of 25 and dimer 34. Therefore this buffer was used in subsequent in vitro assays with enzymes. Attempts were then made to observe the effect of adding either purified ORFZ or cell-free extracts of E. coli expressing ORFZ, to 20 (e.g. Figures S8.1–S8.11). However, despite numerous attempts, no catalysis could be observed.

**Sequence analysis**

α,β-Hydrolases related to PyiE and ORFZ are commonly found encoded in fungal PKS-NRPS biosynthetic gene clusters, especially those with a reductive release mechanism.[17] For example, the cytochalasin E (CcsE),[18] fusarain (Fus2),[18a, 24] pseudoretin (PsoG),[18c] and phomacin (SnoG)[19] gene clusters all encode homologous α,β-hydrolases. No structural data for orthologs of ORFZ/PyiE has been reported, but database (ESTHER)[20] and structural prediction (Phyre-2,[21] Swiss-Model)[22] searches indicate that well-studied enzymes such as acylaminoacyl peptidase (3fnb), 2,6-dihydroxy-pseudo-oxynicotine (DHPON, 2jbw) hydrolase,[23a, 24] and AntI[23] (6hx) possess the same overall protein fold. DHPON hydrolase catalyses a retro-aldol C–C bond cleavage in *Arthrobacter nicotinovorans* (Scheme 6A), while AntI catalyses retro-Claisen reaction followed by a ring-closing cyclisation during the biosynthesis of polyketide-derived anthraquinones in *Photorhabdus luminescens* (Scheme 6B).

Sequence alignment of DHPON hydrolase, AntI, PyiE, ORFZ, CcsE, Fus2, PsoG and SnoG was performed using COBALT,[23c] and a model structure of ORFZ was constructed in Swiss-Model using 3fnb as a template (see Supporting Information) to give an adequate resulting structure (QMEAN –3.48). The ORFZ model aligns with AntI with an RMSD of 3.5 Å over 344 residues and comparison of the sequence and structural alignments gives useful information (Figures S9.2).

AntI has a nucleophilic serine (S245) located at a ‘nucleophilic elbow’ in the highly conserved motif GXSXGG.[23] This motif and the catalytic serine is conserved in DHPON-hydrolase and all the PyiE/ORFZ family members. In AntI S245 forms part of a catalytic triad with D326 and essential H355. H355 has been shown to protonate the retro-aldol nucleofuge.[23] Structural homologs of D326 and H355 are present in DHPON-hydrolase (D300 and H329 respectively) which also catalyses C–C bond cleavage, but these catalytic residues are absent from PyiE, ORFZ and the other cytochalasan hydrolases.

After the retro-Claisen reaction, AntI also catalyses a ring-closing reaction (Scheme 6B). This has been shown to involve enolisation followed by a Dieckmann cyclisation. Residues D175, S176, E181 and R24 of AntI are involved in catalysis of this step, while Y20 is thought to be involved in substrate orientation.[23] These residues are conserved in DHPON-hydrolase and in the PyiE family of hydrolases (e.g. D174, S175, E178, R17 and F13 respectively in PyiE and D165, E169, S243, F12 and R16 in ORFZ, Figure S9.2).

**Discussion**

Knock-out of *pyiE* in *M. grisea* significantly reduced the titres of 1, but initial complementation of *pyiE* did not restore 1 to WT levels. When exogenous O-methyltyrosine 2 was fed to the *pyiE* complemented strain, production of 1 was restored, and new compounds were detected in the Δ*pyiE* strain (Figure 2). This is consistent with damage to the promoter region of *pyiA* which encodes the essential tyrosine O-methyl transferase. The major new compound from the Δ*pyiE* strain chemically complemented with 2 was isolated, purified and structurally characterised as the linear O-methyltyrosine octaketide 9. This compound has not undergone Knoevenagel condensation and instead the expected aldehyde has been reduced to an alcohol. The shunt compound has also been oxidized at C-9, C-15 and C-16 (Figure 3) through addition of hydroxyl groups. These modifications are highly similar to the shunt pyrichalasin precursor 10 observed when *pyiF* was disrupted in *M. grisea*.[27] This KO result is similar to the results of KO experiments ob-
erved in the cases of fusarin (fus2)[18a] and pseurotin (psoG)[18a-c] which also demonstrated that these genes are essential for biosynthesis, although intermediates or precursors were not identified.

Heterologous complementation of ΔpyiE strains with ORFZ from the ACE1 BGC shows that PyiE and ORFZ have the same function. This reinforces previous observations in which we showed that other enzymes from the ACE1 cluster can complement the pyi pathway: for example ORF3 can complement Pyf (the Diels–Alderase); and late stage tailoring enzymes from the ACE1 pathway such as CYP1 and CYP3 are also functional in the pyrichalasin system.[7]

In vitro investigations using synthetic model compounds 11 and 20 show that they do not cyclise rapidly. In water or phosphate buffer at pH 7.0 a very slow spontaneous Knoevenagel cyclisation is observed, followed by further addition of water at the β-position to give 25, and these reactions are accelerated at higher pH. In strong base the cyclisation is fast, but further water addition does not occur. Detailed studies of the major pyrrolinone tautomer show conclusively that the 1,3-dihydro-form 24b is formed: this tautomer cannot undergo the required Diels–Alder cyclisation. Formation of the 1,3-dihydro tautomer 24b also removes the α-stereocentre which is conserved in the cytochalasans, strongly suggesting that non-enzymatic equilibration of 1,3- and 1,5 dihydro tautomers does not occur during cytochalasan biosynthesis. In phosphate buffer in the presence of atmospheric oxygen further oxidation and dimerisation of 24b takes place to give 34, emphasising the inherent reactivity of these intermediates. Our attempts to show in vitro activity of expressed and purified ORFZ with model compound 20 were not successful. It may be that ORFZ is incorrectly folded, as observed for many fungal protein targets of E. coli expression,[31a] or that 20 is not a sufficiently elaborate substrate for the protein. Alternatively, ORFZ may require interaction with other protein partners (vide infra) or the PKS-NRPS itself. No doubt the lack of soluble and active ORFZ/PyiE family members has limited progress in this area.

Conclusions

Taking the above observations into account we can make the following conclusions. First, Knoevenagel reaction of putative intermediate 3 (Scheme 1) requires catalysis by PyiE/ORFZ at physiological pH. The model studies show the uncatalysed reaction is slow at pH 7 and the pyiE KO studies and A. oryzae heterologous expression studies show that intermediate aldehydes such as 3 are quickly reduced in vivo. This agrees with observations from other heterologous expression studies in our group[13] and from Oikawa and co-workers[6b,16a] which show that expression of early genes from cytochalasin pathways in A. oryzae always leads to alcohol shunt intermediates. Corroborating observations from Lebrun and co-workers show that ORFZ (hydrolase) and ORF3 (Diels–Alderase) are the most highly expressed genes from the ACE1 BGC (up to 7-fold higher than the ACE1 PKS-NRPS itself), suggesting a high concentration of these catalysers is required to avoid the reductive shunt pathways.[18a]

Second, the model studies show that rapid tautomerisation of 24a to 24b creates a substrate which is incompetent for further Diels–Alder reaction because the alkene is no longer present at C-3,C-4. Tautomer 24b is itself reactive and undergoes other reactions leading it away from the cytochalasan pathway. Interestingly the observed hemi-amin functionality of 25 ab mirrors the same functionality found in fusarin C[18a] and talaroconvolutin B (Figure 8),[35] for example.

These conclusions show that the previously widely held view that Knoevenagel cyclisation leads to the formation of the 1,5-dydro-2-pyrrol-2-one tautomer 4 prior to Diels–Alder catalysis is an over-simplification. We have shown that in vivo, Knoevenagel catalysis must be rapid enough to out-compete aldehyde shunt reduction, and that Diels–Alder reaction must also occur very rapidly before (non-enzymatic) tautomerisation and loss of chirality.

It is intriguing that the PyiE enzyme family preserve the nucleophile active site serine at a ‘nucleophilic elbow’ position found in DHPON hydrolase and Antl, but lack the acid/base residues required for retro-Claisen reaction. However the acid/ base residues shown by Bode and co-workers to be involved in cyclisation in Antl are conserved in PyiE and ORFZ.[31] This suggests that an alternative possible mechanism for PyiE and its homologues could involve addition of the serine nucleophile of PyiE/ORFZ to the β-ketone of 20 during reaction (Scheme 6C) to give an enzyme-bound tetrahedral intermediate, without retro Claisen cleavage because of the lack of the catalytic triad H and D residues. The formation of a tetrahedral centre at the β-position would prevent incorrect tautomerisation, providing the correct (enzyme-bound) substrate for the subsequent Diels–Alder reaction. This in-turn suggests that the hydrolase (e.g. PyiE/ORFZ) and Diels–Alderase (e.g. Pyf/ORF3) catalysts may work closely together or as a hetero-dimer, or conceivably with the PKS-NRPS itself. However, further detailed in vitro experiments will be required to test this idea, and more reliable methods for the production of soluble and active PyiE/ORFZ and Pyf/ORF3 proteins will be required to probe this hypothesis further. However synthetic and analytic progress made here sets the scene for future advances in this area.

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Conflict of interest

The authors declare no conflict of interest.

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