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# An overlooked cyclase plays a central role in the biosynthesis of indole diterpenes†

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Indole diterpenes (IDTs) are a large class of highly complex fungal natural products that possess a wide array of intriguing bioactivities. While IDTs are structurally diverse, the first four steps of IDT biosynthesis are highly conserved and result typically in the formation of a tetrahydropyran (THP)-ring containing structure, most commonly paspaline. The biosynthetic genes responsible for these steps are the most extensively studied of all IDT genes and collectively define the core biosynthetic pathway. Here we show that the fourth fundamental step, formation of the THP ring, is catalysed by a terpene cyclase encoded by an overlooked and uncharacterised fifth gene, *idtA*. All previously delineated biosynthetic routes have incorrectly attributed this step to the terpene cyclase *IdtB*, leading to imprecise pathway reconstructions and ignoring the fully evolved biosynthetic solution for core IDT generation. Moreover, while *IdtA* terpene cyclases are found in Eurotiomycetes fungi, in Sordariomycetes fungi this step is catalysed by the unrelated protein *IdtS*, demonstrating that two distinct solutions to this chemistry exist. All biosynthetic gene clusters known to specify production of THP-containing IDTs include an *idtA* or *idtS* gene. These findings reset the paradigm for core IDT biosynthesis and support accurate heterologous biosynthesis of these complex natural products.

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## Introduction

Indole diterpenes (IDTs) are a complex and structurally diverse class of filamentous fungal natural products that are comprised of indole connected to a variably cyclised diterpenoid moiety.<sup>1–3</sup> Additional tailoring steps on the core scaffold then massively amplify the structural diversity and associated bioactivity of IDTs. Several hundred IDTs have been identified to date, the majority of which possess a hexacyclic paspaline-type IDT core, which includes a tetrahydropyran (THP) ring derived from the diterpenoid moiety (Fig. 1).<sup>2</sup> These THP-containing IDTs include the archetypical IDT paxilline, as well as the terpenoles, lolitrems, penitrems, shearinines, janthitrems and aflatrems.<sup>4–13</sup> There are only a few known examples that lack this THP ring, such as the nodulisporic acids (Fig. 1A).<sup>14</sup> Here we demonstrate that, contrary to all previously published biosynthetic routes that proposed THP ring formation was catalysed by *IdtB*-type terpene cyclases, this chemistry is delivered by a discrete cluster-encoded cyclase.<sup>2,15–23</sup>

The first IDT biosynthetic gene cluster (BGC), which specifies the production of the paspaline-type IDT paxilline was

identified in 2001 from *Penicillium paxilli*.<sup>24</sup> Through a series of gene deletion, complementation, and heterologous reconstruction experiments four genes from this PAX BGC (*paxG*, *paxC*, *paxM* and *paxB*) were identified as being sufficient to deliver paspaline (Fig. 1B).<sup>20,22,25</sup> Paspaline biosynthesis is initiated with the formation of geranylgeranyl pyrophosphate (GGPP) catalysed by the cluster-encoded GGPP synthase, PaxG. An indole prenyl transferase (PaxC) then catalyses the first dedicated step of IDT biosynthesis, producing 3'-geranylgeranylindole (3'-GGI).

Construction of hexacyclic paspaline from this acyclic precursor then requires iterative cycles of activation and cyclisation. In the first of these, the third olefinic moiety of the diterpene tail is epoxidised by the flavin-dependent monooxygenase PaxM. A precise regio- and stereo-specific cyclisation cascade is then catalysed by the unusual integral membrane type II terpene cyclase PaxB, producing the first stable cyclised IDT, emindole SB.<sup>1,26,27</sup> Generation of the THP ring of paspaline is then primed through a second PaxM-catalysed epoxidation on the terminal terpene olefin of emindole SB. A subsequent cyclisation generates the THP ring, a reaction which was previously also attributed PaxB, or the equivalent *IdtBs* in BGCs from other fungi.<sup>15–17,19,20,22,23,25,28,29</sup> Two P450 monooxygenases, PaxP and PaxQ, then convert paspaline to paxilline.<sup>20–22</sup> Three additional genes are also present in the paxilline BGC: *paxD* and *paxO* encode enzymes that further decorate the indole moiety of paxilline—although these modifications that are not generally

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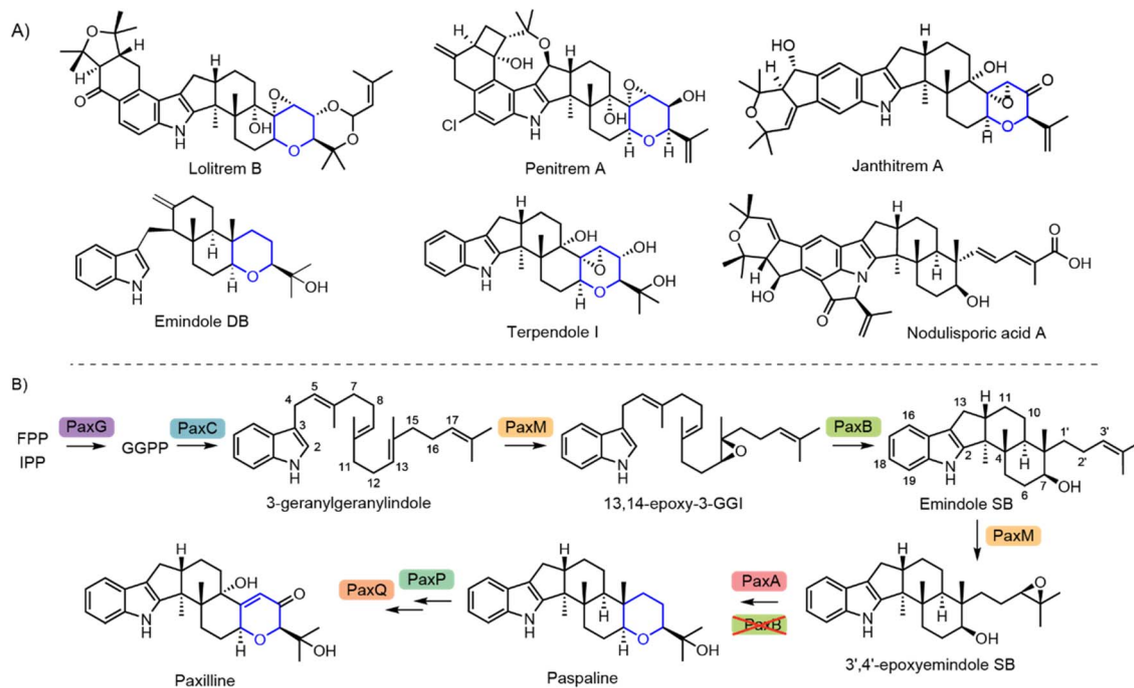


Fig. 1 (A) Representative IDTs with the THP ring shown in blue. (B) The biosynthetic pathway for paxilline.

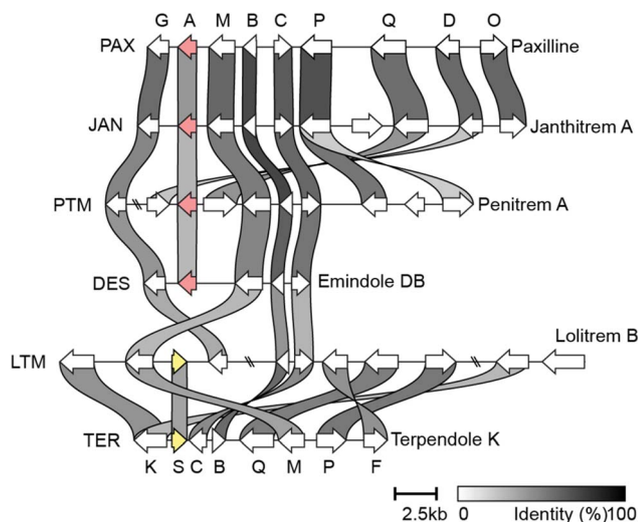


Fig. 2 IDT biosynthetic gene clusters from the Eurotiomycetes fungi *P. paxilli* (PAX), *Penicillium janthinellum* (JAN), *Penicillium crustosum* (PTM), *Aspergillus desertorum* (DES) showing *idtA* homologues in pink. The BCGs from Sordariomycetes fungi *Epichloe festucae* (LTM) and *Tolypocladium album* (TER) contain an alternative gene, *idtS* (shown in yellow). Letters above the PAX cluster correspond to the gene name and function of the encoded enzyme – as shown in Fig. 1B. This figure was generated using clinker.<sup>30</sup>

observed in *P. paxilli*—and a third gene, *paxA*, encoding a predicted helical transmembrane protein of unknown function. Intriguingly, orthologues (*idts*) of *paxA* are also found in other BGCs known to specify THP-containing IDTs (Fig. 2).

## Results and discussion

To investigate the function of PaxA we created *paxA* deletion strains ( $\Delta paxA$ ) in *P. paxilli* (Fig. S1 and ESI 1.2†). In contrast to wild type,  $\Delta paxA$  strains were attenuated for the production of paxilline (Fig. 3, S5 and ESI 1.2.3†) and accumulated an IDT not observed in wild type (Fig. 3A and S4†). This compound, with a molecular weight of 421 Da, was isolated and characterized as 3',4'-epoxyemindole SB (ESI 1.3†), the proposed substrate for THP ring formation (Fig. 1B).<sup>20</sup>

Reintroduction of *paxA* into  $\Delta paxA$  strains complemented this chemotype, attenuating the 3',4'-epoxyemindole SB peak and restoring paxilline biosynthesis (Fig. 3A, S6, S7 and ESI 1.2.4†). In a parallel study, we used a biomimetic approach to construct synthetic BGCs containing *paxG*, *paxC*, *paxM*, *paxB*, and optionally *paxA*, which were then transformed into the  $\Delta PAX$  locus of a *P. paxilli* strain in which the entire *PAX* BGC had been deleted. Consistent with our hypothesis that PaxA catalyses THP ring formation, the synthetic BGC containing *paxA* conferred the ability to synthesise paspaline without appreciable accumulation of 3',4'-epoxyemindole SB. Whereas the strains transformed with the synthetic BGC lacking *paxA* accumulated the epoxidised intermediate (Fig. S8, S9 and ESI 1.2.6†).

To interrogate directly the catalytic role of PaxA we set up a series of feeding experiments using  $\Delta PAX$  strains into which *paxM* alone, *paxM* + *paxA*, or *paxM* + *paxB* had been reintroduced. These strains were fed by the addition of purified emindole SB. Whereas strains expressing *paxM* + *paxA* efficiently converted this to paspaline, the strains expressing *paxM* alone or *paxM* + *paxB* accumulated 3',4'-epoxyemindole SB (Fig. 4, S15 and ESI 1.4†). Combined, these results demonstrate



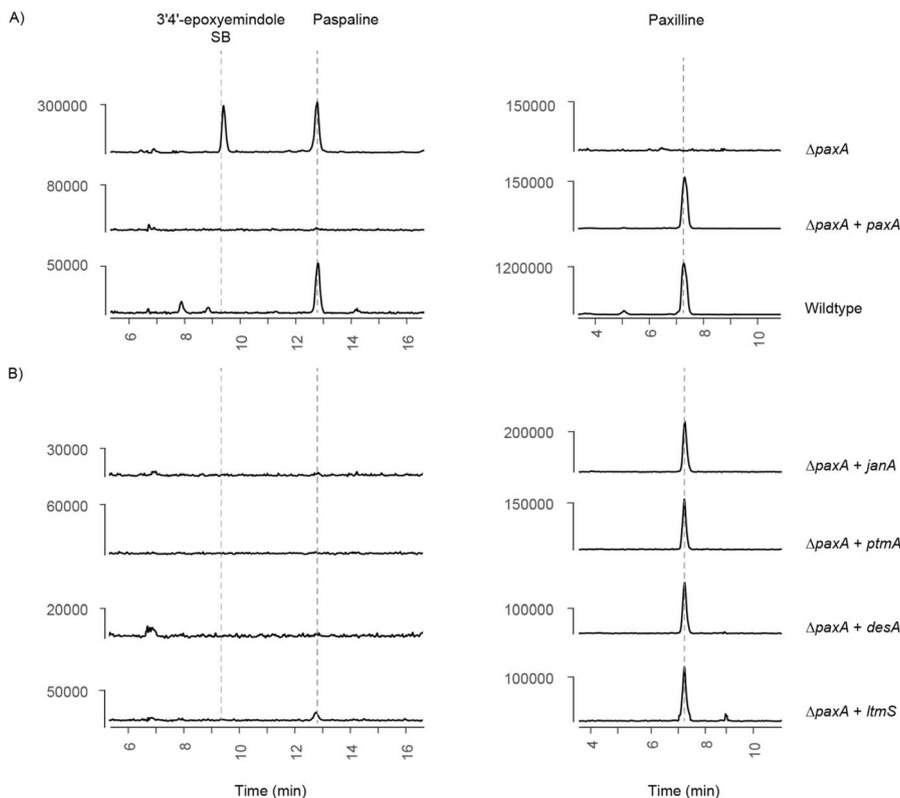


Fig. 3 Extracted ion chromatograms showing IDT production of  $m/z$  422 (LHS) and 436 (RHS) species. Maximum ion intensity counts are shown on the y-axis. (A) IDT production of wildtype *P. paxilli*,  $\Delta paxA$  and a *paxA* complemented knockout strain. (B) Complementation of *P. paxilli*  $\Delta paxA$  with alternative *idtA/S* homologues (*janA*, *ptmA*, *desA* and *ltmA*).

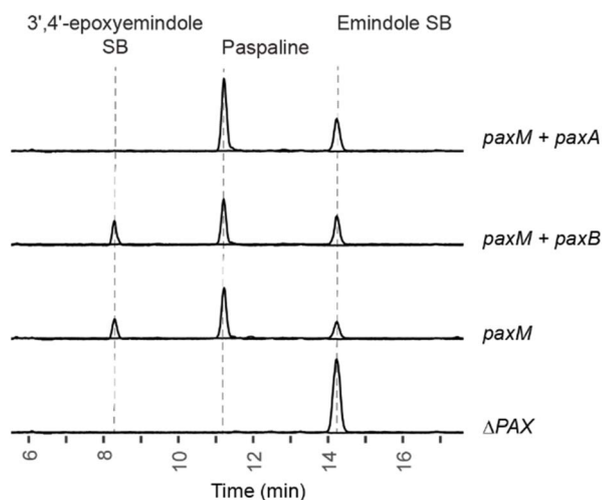


Fig. 4 Combined EIC traces showing IDT production at  $m/z$  406 and 422 in *P. paxilli* strains fed with emindole SB. Ion intensity counts are shown on the y-axis (y-axis range: 0 to  $1 \times 10^6$ ). Chromatograms are annotated with the genes present in each strain.

that PaxA catalyses the conversion of 3',4'-epoxyemindole SB to paspaline *in vivo*. The terpene cyclase PaxB, which was previously assigned this role, does not contribute to THF ring formation.<sup>20,22</sup>

The structural model for PaxA reveals that this protein is a helical integral membrane protein reminiscent of the terpene cyclase PaxB, but with no clear structural or sequence relationship (Fig. 5 and S16–S18<sup>†</sup>). Interestingly however, PaxA bears similarity to other meroterpenoid cyclases that specifically catalyse the formation of THF or tetrahydrofuran (THF) rings.<sup>31–33</sup> Notably in *Acremonium egypitiacum*, which generates meroterpenoids ascofuranone and ascochlorin, both IdtA- and IdtB-like terpene cyclases are encoded.<sup>31</sup> However, rather than operating sequentially as they do in paspaline biosynthesis, in *A. egypitiacum* they operate to bifurcate the pathway to the two alternative meroterpenoid outcomes. Our modelling of 3',4'-epoxyemindole SB into PaxA reveals a likely active site nestled between helices at the C-terminal end of the helical bundle (Fig. 5A). While the putative binding site is lined by hydrophobic residues as expected to support the required conformation for reaction, an aspartate residue is located appropriately to promote THF formation through a favoured 6-*exo*-tet cyclisation (Fig. 5B and C). Accordingly, we found that substituting Asp279 for Ala (D279A) also generated a 3',4'-epoxyemindole SB accumulating chemotype (Fig. 5D, S19 and ESI 1.5<sup>†</sup>). An equivalent Asp residue has been shown to be important for the ascofuranone pathway THF-forming biosynthetic cyclase AscI.<sup>31</sup> We note that Cys66, which is conserved across all confirmed IdtA cyclases (Fig. S18<sup>†</sup>), resides  $\sim 4$  Å from



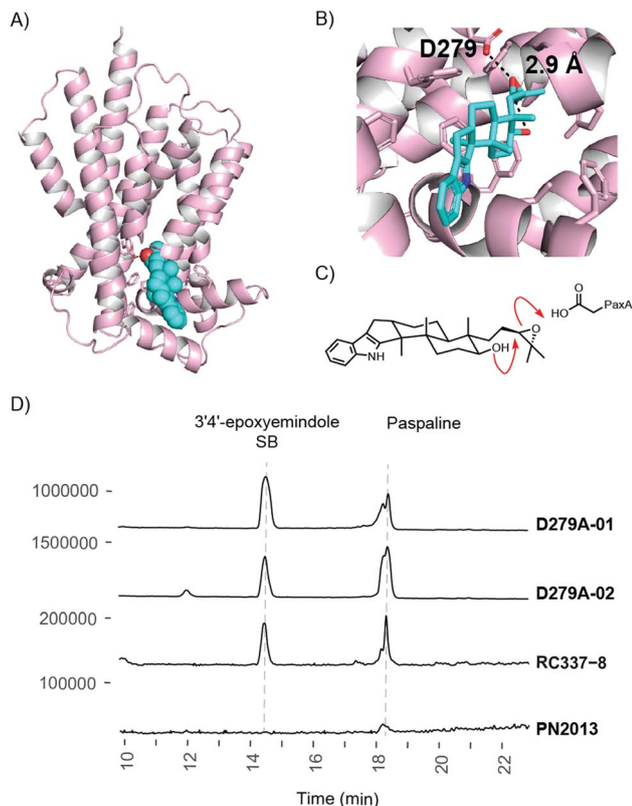


Fig. 5 Structural model of PaxA (generated by AlphaFold<sup>34</sup>), with substrate 3',4'-epoxyemindole SB (cyan). (A) PaxA is a helical integral membrane protein, 3',4'-epoxyemindole SB shown as spheres (B) Close up of predicted binding site showing Asp279. (C) Acid catalysed THP formation. (D) EIC traces showing IDT production at  $m/z$  422 for two Asp to Ala substitution mutants (D279A). Maximum ion intensity counts are shown on the y-axis.

the hydroxyl group of 3',4'-epoxyemindole SB and may play a role in proton loss in THP formation.

It was notable that some paspaline formation was detected in cultures of  $\Delta paxA$  strains (Fig. 3, S4 and ESI 1.2.3<sup>†</sup>), in the *paxGCMB* heterologous reconstruction strains that lacked *paxA* (Fig. S8 and ESI 1.2.6<sup>†</sup>), and in feeding studies where PaxM was present in the absence of PaxA (Fig. 4 and S15<sup>†</sup>). These observations indicate that spontaneous conversion of 3',4'-epoxyemindole SB to paspaline is occurring either *in vivo* or during chemical extraction. The stereochemical outcome in this reaction is set by the prior PaxM-catalysed epoxidation step, and 6-*exo*-tet THP formation is the expected reaction outcome. Indeed, we have observed this conversion directly *in vitro*, with isolated samples of 3',4'-epoxyemindole SB slowly converting to paspaline, and as expected, we show that this reaction is accelerated by the addition of weak acid (Fig. S20 and ESI 1.6<sup>†</sup>). This observed PaxA-independent formation of paspaline (and correspondingly paxilline in strains containing *paxP* and *paxQ*) explains why previous investigations have overlooked the role of PaxA.<sup>15–23</sup> Similarly, the equivalent non-enzymatic formation of the THF ring for the epoxy intermediate in ascofuranone biosynthesis was also observed.<sup>31</sup>

As noted, we identified orthologues (*idtAs*) of *paxA* in BGCs from Eurotiomycetes fungi that are known to biosynthesise similar THP-containing IDTs (Fig. 2, S16 and S17<sup>†</sup>). Accordingly, we tested the function of JanA and Ptma, encoded within the *JAN* and *PTM* clusters that specify the paspaline-derived janthitremene and penitrem IDTs, respectively. Introduction of the *janA* or *ptma* genes into one of our  $\Delta paxA$  strains eliminated 3',4'-epoxyemindole SB accumulation and restored paxilline production (Fig. 3B, S21–24 and ESI 1.7<sup>†</sup>), demonstrating that JanA and Ptma are indeed functional orthologues of PaxA. Additionally, we showed that *desA*, which is a *paxA* homologue located within the emindole DA-specifying *DES* cluster from *Aspergillus desertorum*, also complemented these chemotypes when added to a  $\Delta paxA$  strain (Fig. 3, S25, S26 and ESI 1.7<sup>†</sup>).<sup>15</sup> This is notable as while emindole DB, like paspaline, contains a THP ring (Fig. 1A), this product likely arises from the epoxidation of emindole DA and its subsequent cyclisation catalysed by DesA, implying a wider repertoire of chemistry for the IdtA cyclases solely than the formation of paspaline.<sup>15</sup> This finding implies that some substrate promiscuity is demonstrated by DesA.

Our attention then turned to the IDT BGCs from Sordariomycete fungi, which do not include *paxA* homologues (Fig. 2). For some this was to be anticipated; the nodulisporic acids do not contain a THP ring, being derived directly from emindole SB rather than *via* paspaline, so a *paxA* homologue is not required.<sup>23,35,36</sup> However, several of these BGCs are known to specify paspaline-derived IDTs, yet do not contain an *idtA* gene. Notably, these include the *LTM* cluster from *Epichloë festucae* and the *TER* cluster from *Tolypocladium album*, which are respectively responsible for delivering the lolitremes and the terpendoles.<sup>17,37</sup> We noted that these clusters include the homologous uncharacterised genes *ltsM* and *terS*, which encode proteins with a predicted helical transmembrane topology that is similar to PaxA despite lacking any significant sequence homology (Fig. S27 and ESI 1.8<sup>†</sup>). Introduction of the *ltsM* gene into a *paxA* deletion strain restored efficient conversion of 3',4'-epoxyemindole SB to paspaline and production of paxilline, demonstrating that LtsM is functionally equivalent to PaxA (Fig. 3, S28, S29 and ESI 1.8<sup>†</sup>). Consistent with this observation and supporting an important role for LtsM in lolitrem biosynthesis, *E. festucae*  $\Delta ltsM$  strains grown inside their grass hosts accumulated 3',4'-epoxyemindole SB and showed a reduction in the production of lolitrem B (Fig. S30–33 and ESI 1.8.3<sup>†</sup>).

Collectively these results indicate clearly that these unassigned *idtA* and *idtS* genes encoded within IDT BGCs specify cyclases with key roles in IDT biosynthesis. While the previously proposed biosynthetic pathways for the paspaline and related IDTs included a dual role for the terpene cyclase IdtB in constructing both emindole SB and the full paspaline skeleton, our work here shows that this long-held paradigm is incorrect; the distinct cluster-encoded IdtA and IdtS cyclases deliver the THP functionality.<sup>15–17,19,20,22,23,25,28,29</sup> Moreover, through IdtA and IdtS, Eurotiomycetes and Sordariomycetes fungi have evolved two seemingly independent solutions for delivering this chemistry,



reflecting the importance of this catalysis for efficient IDT biosynthesis.

## Data availability

The data supporting this article have been included as part of the ESI† available at [<https://doi.org/10.1039/d5sc02009c>].

## Author contributions

RCC & EJP conceived the study; RCC, EJP, DB, ATR, LJS, YL & RMM designed the experiments; NSLN provided key materials; RCC, DB, ATR, LJS, YL & KAS collected and analysed the data. RCC, ATR & EJP wrote the manuscript. EJP and RCC acquired the funding. All authors corrected the manuscript.

## Conflicts of interest

There are no conflicts to declare.

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