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Discovery and characterization of a terpene biosynthetic pathway featuring a norbornene-forming Diels-Alderase

Pericyclases, enzymes that catalyze pericyclic reactions, form an expanding family of enzymes that have biocatalytic utility. Despite the increasing number of pericyclases discovered, the Diels-Alder cyclization between a cyclopentadiene and an olefinic dienophile to form norbornene, which is among the best-studied cycloadditions in synthetic chemistry, has surprisingly no enzymatic counterpart to date. Here we report the discovery of a pathway featuring a norbornene synthase SdnG for the biosynthesis of sordarin-the terpene precursor of antifungal natural product sordarin. Full reconstitution of sordarin biosynthesis reveals a concise oxidative strategy used by Nature to transform an entirely hydrocarbon precursor into the highly functionalized substrate of SdnG for intramolecular Diels-Alder cycloaddition. SdnG generates the norbornene core of sordarin and accelerates this reaction to suppress host-mediated redox modifications of the activated dienophile. Findings from this work expand the scopes of pericyclease-catalyzed reactions and P450-mediated terpene maturation.
One of the best-studied pericyclic reactions is the [4 + 2] cycloaddition between a cyclopentadiene and a substituted olefinic diene to form a bridged bicyclic norbornene (Fig. 1a). This Nobel prize winning reaction, first studied by Diels and Alder in their 1928 seminal publication, has become the prototype for cycloaddition and demonstrated many important features of Diels-Alder (DA) reactions such as stereoselectivity, the concerted mechanism, and various mechanisms of rate acceleration. Surprisingly, cycloadditions involving a cyclopentadiene to form norbornene-containing compounds have not been found in biosynthesis. Instead, most reported biosynthetic DA reactions take place with a linear diene involving a cyclopentadiene to form norbornene-containing compounds. However, the IMDA reaction has not been reported to take place during the maturation of a terpene compound thal by the inhibition of fungal elongation factor. Biosynthesis of different sordarins is therefore of interest to understand how Nature biosynthesizes sordaricin. The putative biosynthetic gene cluster (BGC) of a highly decorated sordarin derivative, hypoxysordaricin, was reported by Kudo and coworkers (Fig. 1d). Notwithstanding these findings, the biosynthetic strategy to generate the reactive species for cycloaddition and the nature of the IMDA reaction (enzymatic vs. uncatalyzed) are unresolved.

In particular, it is intriguing how the hydrocarbon cycloaraneosene can be morphed to form the norbornene core in sordaricin. To the best of our knowledge, IMDA reaction has not been reported to take place during the maturation of a terpene natural product, although one example of cycloaddition between

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**Fig. 1 Biosynthesis of sordaricin is an example of the classical norbornene forming Diels-Alder reaction in Nature.** a) DA reaction between a cyclopentadiene and a substituted olefinic diene yielding norbornene. b) Well characterized IMDA reactions in biosynthesis of polyketides and polyketide-nonribosomal peptide hybrid natural products. HRPKS, highly reducing polyketide synthase. c) Retro-biosynthetic scheme of sordarins. The numbering of cycloaraneosene follows Kudo, et al. d) BGCs for hypoxysordarin (sdn) and sordarin B (putative). Genes in sdn but absent from the sordarin B cluster are colored gray. TC, terpene cyclase; GGPPS, geranylgeranyl pyrophosphate synthase; MT, methyltransferase; HP, hypothetical protein; DH, dehydratase; GT, glycosyltransferase; SDR, short chain reductase; TF, transcription factor; FMO, flavin-dependent monoxygenase.

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| A | B | C | D | E | F | G | H | I | J | K | L | M | N | O | P | Q | R | S | T |
|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|
| (TC) (P450) | (GGPPS) | (MT) (P450) | (P450) | (HP) (P450) | (DH) (GT) | (SDR) | (MT) | (TF) | (HRPKS) | (HP) | (Hydrolase) | (TF) | (P450) | (P450) | | | | | | |
| Rosellinia necatrix | Xylaria hypxon | | | | | | | | | | | | | | | | | | | |

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For Sordaricin B (putative), the numbering of cycloaraneosene follows Kudo et al. Genes in sdn but absent from the sordarin B cluster are colored gray. TC, terpene cyclase; GGPPS, geranylgeranyl pyrophosphate synthase; MT, methyltransferase; HP, hypothetical protein; DH, dehydratase; GT, glycosyltransferase; SDR, short chain reductase; TF, transcription factor; FMO, flavin-dependent monoxygenase.
a sesquiterpene dienophile and a polyketide-derived quinone methide diene has been documented. Extensive modifications of the cycloaraneosene skeleton are expected to ready the molecule for norborneone formation: (1) unlike IMDA reactions observed for polyketide chains that are conformationally flexible to position dienes and dienophiles in spatial proximity, the polycyclic terpene molecule is rigid, thereby requiring breaking one or more C-C bonds to afford rotational freedom; (2) desaturation of sp²—sp³ C-C bond(s) in the cyclized terpenes such as cycloaraneosene must take place to generate the diene moiety; and (3) the diene and especially the dienophile require activation to lower the transition state (TS) energy barrier for IMDA. Whereas dienophiles in polyketides that undergo IMDA are typically conjugated to electron withdrawing groups as a result of polyketide synthase programing, the hydrocarbon scaffold of a terpene molecule must be strategically oxidized prior to the pericyclic reaction.

Here, we report the complete reconstitution of sordaricin biosynthesis from cycloaraneosene and the chemical logic that sets up an IMDA reaction to form the norborneone structure. A pericyclic that accelerates the IMDA cycloaddition and attenuates competing shunt product formation is discovered and characterized. The findings in this work represent an example of a pericyclic reaction involved in building terpene structural complexity.

**Results**

**Identification of genes likely involved in sordaricin biosynthesis.** The previously identified hypoxysordarin BGC (sdn, GenBank accession: LC079035.1, https://www.ncbi.nlm.nih.gov/nuccore/LC079035.1) contains twenty genes (SdnA-SdnT), a majority of which are expected to encode enzymes required in the maturation of sordaricin to the final product hypoxysodarin (Fig. 1d, Supplementary Fig. 1). Given that genes encoding enzymes for sordaricin biosynthesis must be conserved in the BGCs of all sordarin analogs, we searched the sequenced fungal genome data for more sdn-like clusters for comparative analysis. Using SdnA as a query, a more compact cluster was found to be conserved in *Rotellinia necatrix* and *Xylaria hypoxylon* (Fig. 1d). This cluster contains eight genes, all of which are present in the larger sdn cluster. In addition to SdnA, the homologous sdn enzymes include four P450 oxygenases (SdnB, SdnE, SdnF and SdnH) and a hypothetical protein (SdnG). A glycosyltransferase (SdnI) and methyltransferase (SdnD) are also conserved, although these two enzymes are not expected to participate in sordaricin formation based on the putative functional annotation. We predict these more compact pathways are responsible for sordarin B biosynthesis (Fig. 1c, Supplementary Table 1), during which SdnI glycosylates the sordaricin core with rhamnose, while SdnD methylates the 4-hydroxy group of the transferred rhamnose. Therefore, we putatively assigned SdnA and the four P450s to be involved in sordaricin biosynthesis, with potential participation by the hypothetical protein SdnG. A flavin-dependent monooxygenase SdnN, which was proposed to play a central role in oxidative maturation of cycloaraneosene, is not conserved between sdn and the more compact clusters (Fig. 1d, Supplementary Fig. 1).

**SdnB is a multifunctional P450 in the sdn pathway.** To reveal the chemical logic for transforming cycloaraneosene into sordaricin, we reconstituted the four *S. araneosa* P450 enzymes SdnB/E/F/H with SdnA in the heterologous host *Aspergillus nidulans* A1145 AEMAST, SdnC, a GGPP synthase present in the sdn cluster but not in the more compact clusters, was included to increase GGPP flux in the host. Expression of SdnA and SdnC led to the production of cycloaraneosene (Supplementary Fig. 2).

We then coexpressed SdnA and SdnC with each of the four P450s (SdnB, E, F, and H) individually. While SdnE, SdnF, and SdnH did not transform cycloaraneosene to new metabolites, coexpression of SdnB yielded two new metabolites 2 (25 mg/L) and 3 (3.5 mg/L) (Fig. 2a, Supplementary Fig. 3a). Structural characterization via nuclear magnetic resonance (NMR) and high-resolution mass spectrometry (HRMS) established 2 as (8 R, 9 S)-cycloaraneosene-8,9-diol, which is derived from two successive hydroxylation of cycloaraneosene (Fig. 2b, Supplementary Notes, Supplementary Table 4, Supplementary Figs. 4, 14–19). The stereochemistries of the diol 2 were determined by NOESY based on the reported stereochemistry for cycloaraneosene. The monohydroxylated cycloaraneosene-8-ol was previously isolated from *S. araneosa* and is likely an intermediate leading to 2 (Supplementary Fig. 1).

Structure elucidation of 3 showed an unexpected shunt product and revealed an additional role of SdnB in the pathway (Fig. 2b, Supplementary Notes, Supplementary Table 5, Supplementary Figs. 4, 20–24). We proposed that 3 is likely formed via oxidative cleavage of the C-8, C-9 diol in 2 by SdnB (Fig. 2b), which was confirmed through direct feeding of 2 to *A. nidulans* expressing only SdnB (Fig. 3a). Compound 2 is stable under the same feeding conditions when an empty plasmid control was used. Since the conversion of 2 to 3 is a net redox-neutral process, the product of SdnB oxidation is likely dialdehyde 4 instead of 3. In the absence of downstream enzymes, the C-7 acrolein moiety in 4 can be reduced by endogenous reductases in *A. nidulans* to afford alcohol 3 as a cellular detoxification mechanism to remove the unsaturated aldehyde. We chemically synthesized 4 by selectively oxidizing the allylic alcohol in 3 to the corresponding unsaturated aldehyde via activated MnO₂. (Supplementary Notes, Supplementary Table 6, Supplementary Figs. 4, 25–29). Consistent with our hypothesis, 4 was readily converted to 3 when fed to *A. nidulans* expressing only empty plasmids (Fig. 3a). Overall, our results suggest that SdnB both acts as a canonical monooxygenase and also a thwarted oxygenase, an oxygenase that consumes oxygen to generate strong enzymatic oxidant but does not result in formal incorporation of oxygen atom into the product (Supplementary Fig. 5). The diol cleavage activity of SdnB enables rotation of the C-6-C-7 bond and thereby freed the C-7-C-17 double bond which is the proposed dienophile for the IMDA reaction (Fig. 2b).

**SdnH is a desaturase that generates the cyclopentadiene.** Each of the remaining P450 enzymes (SdnE, SdnF, or SdnH) was then coexpressed with SdnA-C-B to identify the next biosynthetic step. Only the coexpression of SdnH led to two new metabolites 5 (7 mg/L, Supplementary Notes, Supplementary Table 7, Supplementary Figs. 4, 30–34) and 6 (4.5 mg/L, Supplementary Notes, Supplementary Table 8, Supplementary Figs. 4, 35–39) (Fig. 2, Supplementary Fig. 3b). Both compounds are oxidatively cleaved at C-8 and C-9 as in 3, and both contain the cyclopentadiene functionality. Compound 5 is reduced at C-8 as in 3, whereas 6 is further oxidized from 5 at C-9 to a carbonyl. Similar to the formation of 3 from 4, 5 is likely a redox shunt product derived from cyclopentadiene-dialdehyde 7, which should be the product of sequential actions of SdnB and SdnH starting from cycloaraneosene (and 2). Auto-oxidation or host oxidases may subsequently convert 5 to 6. Our results suggest SdnH, another thwarted oxygenase, catalyzes the desaturation of C-12-C-13 of the cyclopentene ring present in cycloaraneosene to generate the cyclopentadiene (Supplementary Fig. 5).

Two routes can be proposed for the formation of 7 from the diol 2, either via 4 (SdnB followed by SdnH) or via 8 (SdnH followed by SdnB). Comparing the relative titers of shunt products in the heterologous strains suggests that the latter route is in play. A.
**Fig. 2 Sordaricin biosynthesis involves a concise yet well-programmed chemical logic.** a Stepwise heterologous reconstitution of genes involved in sordaricin biosynthesis. The chromatograms are extracted from mass spectra of the base peak for each compound. b Complete biosynthetic pathway for sordaricin. The main pathway is highlighted with a shaded background. All numbered compounds are structurally characterized via NMR and HRMS. The compound in bracket is proposed but not observed.

As a result, 7 is subjected to cellular redox modifications to shunt products 5 and 6. These compounds are also unable to form norbornene due to electronically mismatched substitutions on diene and dienophile pair. To examine the reactivity of 7, we chemically synthesized 7 by activated MnO₂ oxidation of the allylic alcohol in 5 (Fig. 4a, Supplementary Notes, Supplementary Table 9, Supplementary Figs. 4, 40–44). During synthesis in dichloromethane, we observed and characterized the norbornene-dialdehyde 9 as a minor product (Supplementary Notes, Supplementary Table 11, Supplementary Figs. 4, 51–56). We monitored the uncatalyzed cyclization of 7 by following the disappearance of 304 nm absorption from the cyclopentadiene moiety (Supplementary Fig. 4). In a pH 7.4 HEPES buffer, 7 cyclizes with a k_{uncat} of 0.0018 min⁻¹, which corresponds to a half-life of 390 min (Fig. 4c inset). Hence, additional modification to the diene/dienophile pair to align the HOMO/LUMO energies is necessary to form sordaricin. The presence of the C-9 carboxylate group in sordaricin hints oxidation of the C-9 aldehyde in 7 to 10 is the logical next step.

To support this proposed C-9 oxidative activation, we performed density functional theory (DFT) calculations to understand the transition state energy barrier for IMDA starting from 7 (C-8, C-9 dialdehyde, TS-1), 10 (C-8 aldehyde, C-9 carboxylic acid, TS-2), and 5 (C-8 alcohol, C-9 aldehyde, TS-3) (Fig. 4e). The calculated ΔG_{uncat} for cyclization of 10 is 20.3 kcal/mol, which is ~7 kcal/mol lower than that for dialdehyde 7.
(27.6 kcal/mol) and represents significant rate enhancement upon C-9 oxidation to the carboxylate. The sluggish cyclization of dialdehyde 7 can be rationalized since both diene and dienophile in 7 are electron deficient. Oxidation of C-9 aldehyde to carboxylate eliminates the electron withdrawing aldehyde and accelerates cyclization of 10. The calculation also confirmed a much higher activation barrier (33.9 kcal/mol) for the DA cycloaddition of 5, consistent with the removal of the C-8 electron-withdrawing group conjugated to the dienophile.

One of the two remaining P450 enzymes is a likely candidate for the oxidation of 7 to 10. Indeed, upon coexpression of SdnF with SdnA-C-B-H in A. nidulans, two new norbornene-containing metabolites 11 (1 mg/L) and 12 (1 mg/L) were formed with the concomitant disappearance of 5 and 6 (Fig. 2, Supplementary Fig. 3c). Compound 11 contains the cyclized norbene ring and is one additional C-18 hydroxylation step from 1 (Supplementary Notes, Supplementary Table 13, Supplementary Figs. 4, 62–66). Compound 12 is a shunt product derived
from 11 via reduction of C-8 aldehyde to alcohol (Supplementary Notes, Supplementary Table 14, Supplementary Figs. 4, 67–71). When chemically prepared 7 was fed to A. nidulans expressing SdnF alone, efficient conversion to 11 was observed (Fig. 5a). To assay the oxidation activities of SdnF separately from cyclization of 10, 5 was supplemented to the SdnF expression strain and a control strain expressing only empty plasmids. While the control strain is able to convert ~30% of 5 to the carboxylate 6 likely via auto-oxidation or host oxidases, the SdnF expression strain led to near complete conversion of 5 to 6 (Fig. 5b), demonstrating that SdnF is able to selectively oxidize the C-8 aldehyde to carboxylate.

To measure the rate of IMDA cyclization after C-8 oxidation, we chemically synthesized 10 by oxidizing 6 with activated MnO₂ and followed its cyclization by HPLC (Supplementary Notes, Supplementary Table 12, Supplementary Figs. 4, 57–61). In a pH 7.4 HEPES buffer, 10 cyclizes to form 11 with a \( k_{\text{cat}} = 0.0018 \pm 0.0001 \text{ min}^{-1} \) and \( k_{\text{cat}} = 0.103 \pm 0.004 \text{ min}^{-1} \) for compounds 7 and 10, respectively, using the same buffer at pH 7.4. These results suggest that the oxidation of 7 to 6 leads to an increase in the rate of IMDA cyclization.

**Fig. 4** In vitro characterization of non-enzymatic and enzymatic norbornene formation in sordaricin biosynthesis. a Overall synthetic scheme for compound 7 and 10. b Cyclization of compound 10 in the absence (left) and presence (right) of SdnG in a 50 mM HEPES buffer, pH 7.4. Decrease at A₂₈₆ nm (disappearance of the cyclopentadiene chromophore) was used to monitor the cyclization. c First order kinetics of non-enzymatic cyclization of compounds 10 and 7 (inset). The data points and error bars represent the mean and standard deviation of \( n = 3 \) independent measures respectively. Errors associated with the kinetic parameters were obtained from fitting. Source data are provided as a Source Data file. d Michaelis-Menten kinetics of SdnG with compound 10 as substrate. The data points and error bars represent the mean and standard deviation of \( n = 3 \) independent measures respectively. Errors associated with the kinetic parameters were obtained from fitting. Source data are provided as a Source Data file. e DFT calculations of IMDA transition states in water for dialdehyde 7 (TS-1), and carboxylate-aldehyde 10 (TS-2), and aldehyde-alcohol 5 (TS-3).
The molar ratio of combined shunt products to sordaricin is 10:1, suggesting that at least 20% of sordaricin is diverted to shunt pathways that outcompete the IMDA reaction. This is not unexpected since the measured k_{cat} is 0.103 min^{-1} for 10 is relative slow compared to those expected for endogenous enzyme-catalyzed redox modifications. This apparent biosynthetic inefficiency prompted us to examine if additional biosynthetic enzymes are needed to accelerate the IMDA reaction and prevent shunt product accumulation.

**Complete reconstitution of sordaricin biosynthesis.** To complete sordaricin biosynthesis, we introduced the remaining P450 SdnE in *A. nidulans* expressing SdnA-C-B-H-F. The resulting host indeed biosynthesized sordaricin 1 (20 mg/L) via hydroxylation of the C-18 methyl in 11 (Fig. 2). The 1H and 13C NMR spectra and specific rotation of 1 match with those reported for sordaricin (Supplementary Notes, Supplementary Table 3, Supplementary Figs. 4, 9–13). The pathway was also successfully reconstituted in *S. cerevisiae* RC01 with a titer of 2 mg/L (Supplementary Fig. 6). Notwithstanding the complete reconstitution of sordaricin pathway, we observed co-accumulation of previously isolated shunt product 6, as well as two new shunt metabolites 13 (2.3 mg/L) and 14 (1.5 mg/L) (Fig. 2a). The same three shunt products were also present together with 5 in the strain that coexpressed SdnA-C-B-H-F (Fig. 2a). Scaled up cultures led to isolation and characterization of 6, 13, and 14 (Supplementary Notes, Supplementary Table 15, Supplementary Figs. 4, 7–82) as unacylated shunts derived from reduction of the dienophile in 10. Furthermore, the C-8 aldehyde in 13 is oxidized to carboxylate, while reduced to alcohol in 14. The molar ratio of combined shunt products to sordaricin 6 is 1 to 4, suggesting that at least 20% of 10 is diverted to shunt pathways that outcompete the IMDA reaction. This is not unexpected since the measured k_{cat} is 0.103 min^{-1} for 10 is relative slow compared to those expected for endogenous enzyme-catalyzed redox modifications. This apparent biosynthetic inefficiency prompted us to examine if additional biosynthetic enzymes are needed to accelerate the IMDA reaction and prevent shunt product accumulation.

**SdnG is a norbornene synthase.** To test if the sdn uses a dedicated enzyme to further accelerate the IMDA reaction of 10 to 11, the remaining uncharacterized enzyme SdnG conserved between the sdn and more compact clusters shown in Fig. 1d was expressed in the strains that produced 1. The predicted open reading frame of SdnG encodes a 146 amino acid protein with no conserved domain. A BLAST search in NCBI genome database did not identify any homologs of SdnG outside sdn-like clusters (Supplementary Fig. 7). When coexpressed, a dramatic change in metabolite profile was observed in which 1 is now nearly the exclusive product, with the shunt products present only at trace levels (Fig. 2a). Similarly, when SdnG was coexpressed in strain that produced 11 (SdnA-C-B-H-F), shunt product titers were greatly attenuated (Fig. 6a). Suppression of shunt products suggests SdnG acts as a pericyclase, more specifically a Diels-Alderase, to accelerate the IMDA reaction and outpace cellular redox modifications. It is worth noting here that once the final molecule 1 is formed, we can only detect trace amount of reduction or oxidation of the C-8 and C-9 groups in *A. nidulans*, for reasons we do not fully understand.

To verify the role of SdnG, the protein was expressed and purified from *E. coli* and assayed directly in the presence of the putative substrate carboxylic acid 10 (Supplementary Fig. 8a). In the absence of SdnG, up to 90% 10 (100 μM) is cyclized within 30 min, whereas cyclization of 90% 10 was observed within 2.5 min upon addition of 1 μM SdnG (Fig. 4b). This rate acceleration requires SdnG since a generic globular protein such as bovine serum albumin (BSA) does not accelerate cyclization of 10 (Supplementary Fig. 8b). The steady-state
kinetics of SdnG was measured to give $K_M$ towards 10 of 19 $\mu$M and $k_{cat}$ of 56 min$^{-1}$ (Fig. 4d). The value of $k_{cat}$ is comparable to previously characterized pericyclases$^{9,11,13,15}$ and corresponds to a ~560-fold acceleration of the $k_{cat}$ of 9. This rate acceleration is similar to that reported for SpnF, the first Diels-Alderase characterized$^9$. SdnG is specific for the carboxylylate-aldehyde 10, as its pericyclase activity towards 10 is at least 100-fold higher than dialdehyde 7 in vitro (Supplementary Fig. 8b). This is consistent with coexpression of SdnG with SdnA-C-B-H in A. nidulans, which did not lead to detectable amount of norbornene product 9 (Fig. 6b). Together, the results demonstrate SdnG as a dedicated Diels-Alderase catalyzing norbornene formation.

Discussion

Our bioinformatics, heterologous reconstitution, and bio-transformation analysis revealed the complete biosynthetic pathway for sordaricin. This pathway features a concise chemical logic to transform the linear primary metabolite GGPP to a highly functionalized norbornene scaffold via only one terpene cyclase and a four-P450 cascade that consumes six molecules of oxygen and incorporates four oxygen atoms into the 5-8-5 core to other diterpenes such as fusicoccadiene$^{36}$, cyclooctat-9-en-7-ol$^{37}$, and ophiobolin M$^{38}$, cycloaraneosene has a distinguishing exocyclic olefin at C-7 which becomes the dienophile for the IMDA reaction (Fig. 1c).

Following SdnA, three P450s catalyze a series of intriguing reactions to overcome the intrinsic difficulty for a terpene scaffold to undergo IMDA reactions. The dihydroxylation of cycloaraneosene and the diol cleavage of 8 catalyzed by SdnB immediately rearrange the cycloaraneosene scaffold to free the C-7 dienophile from the rigid tricyclic scaffold. Our mechanism of dienophile release differs significantly from the previous proposal where the C-8-C-9 bond is to be cleaved via Baeyer-Villiger oxidation and the subsequent hydrolysis of resulting ester$^{27}$ (Supplementary Fig. 1). P450-catalyzed vicinal diol cleavage to yield fragmented aldehydes has been reported in biosynthesis of pregnenolone$^{39}$, biotin$^{40}$, and botrydial$^{41}$, but the SdnB reaction is particularly interesting since in this case the cleavage of one C-C bond readies the molecule to form two new C-C bonds via the later DA reaction. The idea of break a bond to make a bond, or molecular editing, has been increasingly employed in total synthesis of complex natural products as a strategy to reshuffle the molecular scaffold and generate new reactivity$^{42}$. SdnB functions represent a biosynthetic example of generating the diene/dienophile pair via such carbon skeleton reconstruction.

The desaturation catalyzed by SdnH generates a key conjugated diene which otherwise are absent in a typical terpene. Previous study by Kudo, et al. suggested that the cyclopentadiene was generated by C-13 hydroxylation of 2 and the subsequent dehydration of the resulting cycloaraneosene triol via an unspecified enzyme$^{27}$ (Supplementary Fig. 1). Our results show that SdnH alone is sufficient for the desaturation. While P450 desaturases can perform both direct desaturation and hydroxylation, it is generally believed that the hydroxylated compounds are premature OH rebound products rather than intermediates of the desaturation activity$^{43}$. The disruption of SdnB’s hydroxylation and diol cleavage activities by SdnH is reminiscent of the multifunctional P450 TamI in tirandamycin biosynthesis$^{44}$. Such an
unusual reaction sequence is likely programed to avoid premature generation of the reactive acrolein dienophile which is cytotoxic and leads to redox shunt products. Besides freeing the dienophile, the diol cleavage catalyzed by SdnB serves another purpose towards the formation of sordaricin: installing conjugated aldehyde groups to both the dienophile and the cyclopentadiene. While the dienophile is sufficiently activated with an electron deficient aldehyde group, the HOMO/LUMO energy gap of the diene/dienophile pair is widened by a similarly electron deficient aldehyde group on the diene. Therefore, a third P450 SdnF is necessary to further activate the diene/dienophile pair for IMDA by converting the aldehyde conjugated to the diene to a less electron withdrawing carbonyl group. Such redox activation of diene/dienophile is analogous to the role played by the flavin-dependent enzyme SoI5 catalyzing a decalin-forming IMDA in solanapyrone biosynthesis where SoI5 is required to activate the dienophile by oxidizing an electron donating hydroxymethyl group conjugated to the dienophile to an electron withdrawing aldehyde. Whereas P450 modification of terpene scaffolds are widely found in biosynthetic pathways, the collective roles played by SdnB, SdnH and SdnF to set up the IMDA is a rare catalytic strategy. We also discovered a Diels-Alderase catalyzing norbornene formation. Although the IMDA reaction of 10 to 11 can take place uncatalyzed and stereoselectivity is substrate-controlled, the sdn pathway uses a dedicated pericyclase SdnG to accelerate the reaction as a means to suppress shunt product formation. It is not clear how endogenous redox metabolism towards reactive species such as the acrolein dienophile in the sordarin producers differs from that of A. nidulans, but in the heterologous host the redox modifications drain pathway intermediates towards uncylized shunt products. SdnG contains no recognizable cofactor binding domain and does not depend on any exogenous cofactors for activity, joining a group of cofactor-free pericyclases including SpinF, CghA, PyrfA, IccD, and the recently reported TsS1. However, SdnG shares no sequence homology with any characterized pericyclases and therefore likely represents an undescribed class of pericyclases. Further structural and functional characterizations of SdnG are currently underway to reveal mechanistic insights of such a norbornene synthase. Norbornene ring formation from a cyclopentadiene and an olefinic dienophile has been a mainstay of mechanistic investigation and synthetic use of Diels-Alder chemistry for the past 80–90 years. The sordaricin biosynthetic pathway featuring SdnG represents an example of a biologically accelerated DA reaction to construct this classical bicyclic ring system. The pathway discovered here utilizes a concise yet well-programmed chemical logic that is unmatched by chemical synthesis. The small size and the cofactor-free nature of the norbornene synthase SdnG sets up further mechanistic and biocatalytic exploration of pericyclases. Together, our finding adds to the ever-expanding reservoir of biological pericyclic reactions and enriches Nature’s toolbox for building terpene complexity. Lastly, synthetic sordarin analogs derived from sordaricin have shown great promise as leads for antifungal therapy. Our results provide a direct and scalable route to sordaricin.

Methods
Bioinformatics. BGCs containing cycloaraneosene synthase SdaA were identified by a BLAST search of SdaA homologs in NCBI genomic database (https://www.ncbi.nlm.nih.gov/genome/). Genomic scaffolds containing SdaA homologs were annotated by 2ndfnd. The NCBI Conserved Domain Search (https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi) was used for conserved domain analysis. Genomic DNA extraction from S. araneosaene and cDNA synthesis. S. araneosaene NRRL 3196 was obtained from the Agricultural Research Service Culture Collection (NRRL) and maintained on potato dextrose agar (Sigma). A 10 ml liquid culture of S. araneosaene NRRL 3196 in potato dextrose broth (Sigma) was shaken for 7 days at 28 °C and 250 rpm. The cell body was then collected for genomic DNA extraction with Quick-DNA” Fungal/Bacterial Mini Prep Kit (Zymo research) following the manufacturer’s protocol.

For mRNA extraction and cDNA synthesis, a 10 ml liquid culture of S. araneosaene in production medium (10% glucose, 1.5% polypeptone, 1.0% corn steep liquor, 0.5% yeast extract, 0.2% L-tryptophan, 0.5% K2HPO4, 0.1% MgSO4·7H2O, 0.05% CaSO4·H2O, 0.01% FeSO4·7H2O, 0.05% CoCl2·6H2O) was shaken for 7 days at 28 °C and 250 rpm. The cell body was then collected for mRNA extraction with Ribopure” Yeast RNA Purification Kit (Thermofisher) following the manufacturer’s protocol. First strand synthesis was performed with SuperScript III First-Strand Synthesis System (Invitrogen).

General DNA manipulation techniques. The DNA sequence of previously identified sdnA (GenBank accession: LC079031.1) was used for all subsequent studies. Plasmids for overexpression in A. nidulans were constructed as follows. The genes involved in sordaricin biosynthesis were amplified from genomic DNA of S. araneosa via polymerase chain reaction (PCR) and inserted into plasmids pFTU, pYTP, and pYTP via homologous recombination. PCR was performed with Phusion (Thermofisher) or Q5 (NEB) high fidelity polymerases. Transformation of yeast was performed with Frozen-EZ Yeast Transformation II kit (Zymo research) following the manufacturer’s instructions. Plasmids were then transferred from yeast to ZymoPrep Yeast Plasmid Miniprep I (Zymo research) following the manufacturer’s instructions and transformed into E. coli Top10 electrocompetent cells (Invitrogen) for propagation. Plasmids extracted from Top10 (Zippy Plasmid Miniprep Kit, Zymo Research) were screened via Sanger sequencing (Laragen). Plasmids for overexpression in S. cerevisiae were constructed similarly except that the genes were PCR-amplified from genomic DNA of S. araneosa. All primers used in this study are listed in a separate file (Supplementary Table-sequences of oligonucleotides used in the study).

Heterologous reconstitution of genes involved in sordaricin biosynthesis. Fungal transformation was performed as follows. A. nidulans ΔMMST1 was grown on oatmeal agar (Sigma) supplemented with uracil (5 mM), uridine (10 mM), riboflavin (0.125 µg/mL), and pyridoxine (0.5 µg/mL) at 28 °C for three days. Spores were filtered through a 40 µm cell strainer (Fisher), transferred to 45 ml CD media (1% glucose, 5% nitrate salt mix, 0.12 ml NaNO3, 0.104 µg/ml KCl, 0.104 µg/ml MgSO4·7H2O, 0.304 µg/ml KH2PO4, 0.1% trace element mix (0.022 µg/ml ZnSO4·7H2O, 0.011 µg/ml B2O3, 0.005 µg/ml MnCl2·4H2O, 0.0016 µg/ml FeSO4·7H2O, 0.0016 µg/ml CoCl2·6H2O, 0.0916 µg/ml CaSO4·2H2O, 0.0016 µg/ml (NH4)MoO4·2H2O, 0.126 µg/ml (H2)MoO4·2H2O), 3% starch, 5% agar, 2% casamino acid) and shaken at 28 °C and 250 rpm for 12–16 h. This culture was centrifuged to collect the fungal biomass which was subsequently washed with 10 ml osmotic media (1.2 M MgSO4·7H2O, 10 mM sodium phosphate, pH 5.8). The resulting fungal body was resuspended in 10 ml osmotic media supplemented with 30 µg of lysine (Sigma), 0.4 µg of Triton X100 (Sigma), filtered through a 40 µm cell strainer and then resuspended in 10 ml CD media (1.2 M MgSO4·7H2O, 10 mM sodium phosphate, pH 5.8) at 28 °C and 80 rpm for 5 h. The suspension was then poured into a centrifuge tube and 10 ml trapping buffer (0.6 M sorbitol, 0.1 M Tris, pH 7.0) was slowly added to the suspension so that the layers of fungal suspension and trapping buffer remained undisturbed. The tube was then placed at 4200 g for 10 min. The resulting trapped protoplast (greenish white layer at the interface of the fungal suspension and trapping buffer layers) was washed with 10 ml STC buffer (1.2 M sorbitol, 10 mM CaCl2, 10 mM Tris, pH 7.5) and then resuspended in 1.2 ml STC buffer on ice.

The resulting transformants were grown on CDST-Agar medium (2% starch, 5% nitrate salt mix, 0.1% trace element mix, 2% agar) plates which were then incubated at 37 °C for 2–3 days until colonies appeared.

For reconstitution in S. cerevisiae, the markers were amplified from genomic DNA of S. araneosa via polymerase chain reaction (PCR) and inserted into plasmids pYTP, pYTP via homologous recombination. The two plasmids were subsequently transformed into S. cerevisiae RC01 with Frozen-EZ Yeast Transformation II kit (Zymo research). A 3 ml YPD culture (2% glucose, 1% yeast extract, 2% bacto-agar) was grown to mid-log phase in 250 ml baffled Erlenmeyer flasks at 28 °C and 250 rpm. The resulting cell body was then collected for genomic DNA extraction with Ribopure” Yeast RNA Purification Kit (Thermofisher) following the manufacturer’s protocol. First strand synthesis was performed with SuperScript III First-Strand Synthesis System (Invitrogen).
extract, and 2% peptone) was inoculated with a single transformant colony and incubated in a 28 °C shaker, 250 rpm for 72 h. A 1 mL aliquot of this culture was then removed, cooled centrifuged at 17,000 × g for 5 min with the media and the cells which were then extracted separately with ethyl acetate and acetone, respectively. The organics were combined and dried under vacuum. The resulting residues were reconstituted with 100 μL HPLC grade methanol and analyzed similarly with A. nidulans reconstitution.

Compound isolation and characterization. Typically, 30 mL CDST agar plates of 2L culture inoculated with A. nidulans transformants were placed in a 28 °C incubator for 3–4 days. The agar plates were cut into pieces and soaked in 2L acetone for 24 h. Agar was removed by filtration and extracted again with acetone. The two extractions were combined and evaporated to dryness by a rotary eva- porator. The residues were extracted with acetone and ethyl acetate. The crude extracts were separated by silica flash chromatography with a Combiflash® system and a gradient of hexane and ethyl acetate. The targeted compounds were further purified by an UltiMate™ 3000 Semi-Preparative HPLC (ThermoFisher) with an Eclipse XDB-C18 column (5 μm, 9.4 × 250 mm, Agilent) and a gradient of water and acetonitrile (both mobile phases contain 0.1% formic acid). The gradient started with 30% acetonitrile and then linearly increased to 95% acetonitrile in 35 min. Purified compounds were dried in vacuum and analyzed by the Agilent UHPLC-HRMS as stated above and a Bruker AV500 NMR spectrometer with a 5 mm dual cryoprobe (500 MHz, UCLA Molecular Instrumentation Center). NMR data were collected with Topspin 3.5pl4 (Bruker) and analyzed with Topspin 4.1 (Bruker). Specific rotation was measured with an Autopol III Automatic Polari- meter equipped with a 50 mm polarimeter cell (Rudolph Research Analytical). The yield and spectroscopic data of each compound are listed in the Supplementary Notes.

Heterologous biotransformation. A. nidulans transformed with the gene of interest was grown on an 8 cm2 agar plate with 200 μL substrates for 5 days at 28 °C. Agar plugs from these plates were then extracted and analyzed as stated above. A. nidulans transformed with empty plasmids served as a control.

For biotransformation of compound 7 by SdnF, A. nidulans transformed with the SdnF was grown on a 5 mL CDST-Agar plate for 3 days. Compound 7 was dissolved in CH2Cl2 (2.5 mM final) and 10 μL of this solution was dropped directly onto the fungal mycelia. The plate was incubated at 28 °C for 24 h and an agar plug containing the fed compound was analyzed as stated above.

General synthetic procedures for compounds 4, 7, and 10. Compounds 4, 7, and 10 were synthesized from their corresponding reduced shunt products 3, 5, and 6 respectively by selective oxidation of the allylic alcohol with activated MnO2.45 Briefly, 20 mM alcoholic compounds were dissolved in CH2Cl2. Twenty molar equivalents of activated MnO2 (Millipore) were added and the reaction mixture was vigorously stirred at room temperature for 2 h. MnO2 was removed by centrifugation and the supernatant containing the desired product was generally used for downstream studies without the need for further purification.

In vitro non-enzymatic IMDA cyclization of compounds 7 and 10. Compound 7 or 10 was dissolved in a buffer of 50 mM HEPES, pH 7.4 with 5% DMSO (final concentration 100 μM). The reaction mixture was maintained at 25 °C with a MyBlock Mini Dry Bath (Benchmark Scientific) throughout the experiment. Ali- quotes (10 μL) of the reaction were taken at indicated time and immediately mixed with an equal volume of ice cold acetonitrile. The mixture was then directly analyzed by the previously mentioned UHPLC-QTOF. The solvent program was an isotropic gradient of 80% acetoni trile for 2.5 min. Decrease at A260 nm (for compound 7) or A280 nm, (for compound 10) was used to monitor the IMDA cyclization. The remaining reactant was quantified by a standard curve of compound 7 or 10. Concentration of the remaining reactant versus reaction time was fitted with first order kinetics to obtain the rate constant kcat (GraphPad Prism 8).

DFT calculation. Initial conformational searches were conducted on all reported structures using xtb and CREST57,58. The output geometries were recalculated with the density function and basis set o897X-D/dde2-SVP as implemented in Gaussian 16 Rev. A.03 (see ref 59-62). This functional was chosen for its ability to reproduce CCSD geometry calculations of asynchronous Dieder-Alder reactions as well as its general applicability for describing reaction barriers64. Following Head-Gordon’s suggested basis set for energetics64, we computed single point energies at the DFT calculation level of theory since the 1930s: Woodward, Houk with Woodward, and Head-Gordon. Transition states were located by Berny optimization. The output geometry was subjected to an constrained conformational search to ensure that all the substituents were in their lowest energy conformation. Then, the structure was re-optimized at the o897X-D/dde2-QZVPP level of theory. Frequency calculations were conducted to verify whether each structure was indeed a transition state. Intrinsic reaction coordinate calculations were conducted to further verify the connectivity of the transition states. We have included one such calculation in the supplementary information for TS-2: the calculation indicates that TS-2 connects reactant 10 to product 11 (Supplementary Fig. 82). The molecular coordinates of all calculated structures are listed in a separate Supplementary file (Supplementary Data).

Cloning, expression, and purification of SdnG. SdnG was PCR amplified from cDNA of S. araneosa and subcloned into a PET28a plasmid with a N-(His)6-SUMO tag via NEBuilder™ HiFi DNA Assembly Master Mix (NEB) (Supplementary Table 2). The resulting plasmid harboring N-(His)6-SUMO-SdnG was transformed into che- mically competent E. coli Rosetta 2 (Millipore). The transformant was used to inoculate 1 L LB medium with 200 μg/mL kanamycin and 34 μg/mL chloramphenicol. The culture was shaken at 37 °C and 220 rpm until OD600 reached 0.6–1 and cooled to 16 °C. IPTG was added to the culture to a final concentration of 100 μM and the culture was shaken at 16 °C for 16–20 h. After expression was completed, cells were harvested by centrifugation, flash-frozen in liquid nitrogen, and stored at −80 °C. Cell pellet from 1 L culture (~5 g) was resuspended in cell lysis buffer (50 mM sodium phosphate, 500 mM sodium chloride, 10% glycerol, 250 mM imidazole, pH 8.0) by vortexing. All subsequent purification procedures were performed at 4 °C unless otherwise stated. Cells were lysed by sonication on ice (2 s on, 5 s off cycle for 24 min, 50% maximum amplitude) and then centrifuged for 30 min at 4 °C and 21,000 x g to pellet cell debris. The supernatant was mixed with 1 mL Ni-NTA resin (ThermoFisher, pre-equilibrated with 5 column volumes of cell lysis buffer) and agitated mildly for 1 h. The mixture was then loaded onto a gravity column. The resin was washed with at least 20 column volumes of cell lysis buffer. The N-(His)6-SUMO- SdnG was then eluted from the column with 10 column volumes of elution buffer (50 mM sodium phosphate, 500 mM sodium chloride, 10% glycerol, 250 mM imidazole, pH 8.0). Ulp1 protease (−40 μg) was then added to the eluted protein and the mixture was incubated on ice overnight to cleave the N-(His)6-SUMO tag. The cleaved tag and Upl1 protease which contains a N-(His)6 tag were separated from SdnG by running the mixture through a fresh Ni-NTA column. SdnG, now (His)6 tag free, passed through the column while the SUMO tag and Ulp1 protease remained bound. Purified SdnG was then concentrated by an Amicon spin concentrator (Millipore), aliquoted, flash-frozen in liquid nitrogen, and stored at −80 °C. The purified SdnG was analyzed by 12% SDS-PAGE and protein concentration was estimated from its predicted extinction coefficient (ε280 nm = 8940 cm−1 M−1).49

In vitro assay of SdnG. SdnG (0.18 μM for compound 10 and 10 μM for compound 7) was preincubated in 95 μL of 50 mM HEPES, pH 7.4 buffer for 5 min. This solution was maintained at 25 °C with a MyBlock Mini Dry Bath (Benchmark Scientific) throughout the experiment. Reaction was initiated by adding 5 μL DMSO solution of compounds 10 or 7. The final concentration of the substrate is 20–175 μM for compound 10 and 100 μM for 7. After 1 min, 10 μL of the reaction were taken and immediately mixed with an equal volume of ice cold acetonitrile. The mixture was then directly analyzed by UHPLC-QTOF same as the non- enzymatic cyclization reactions. The measured rate was subtracted with the rate of non-enzymatic cyclization to obtain the true enzymatic reaction rate. The concentra tion of compound 10 versus rate was fitted with Michaelis-Menten equation to deduce steady-state kinetics parameters (GraphPad Prism 8).

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. All data supporting the findings of this study are available within the article and its Supplementary Information files. All unique biological materials, such as plasmids, generated in the study are available from the authors. The DNA sequence of sdn cluster (Accession: LCC079035.1) and genomes of Rosellinia necatrix (Accession: Ab6D0108871+) and Xylaria hypoxylon (Accession: GCA_902806585.1) are available from NCBI. The protein sequence of SdnG (Accession: AA0144441) is available from Uniprot. Source data are provided with this paper.

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Author contributions
Z.S., C.S.J., M.O., K.N.H., and Y.T. developed the hypothesis and designed the study. Z.S. performed all in vivo and in vitro studies. C.S.J. performed all computational studies. Z.S., C.S.J., M.O., K.N.H., and Y.T. analyzed and discussed the results and prepared the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
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