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An Enzyme-Catalyzed [4+2] Cycloaddition is a Key Step in the **Biosynthesis of Spinosyn A**

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The Diels-Alder reaction is a [4+2] cycloaddition reaction in which a cyclohexene ring is formed between a 1,3-diene and an electron-deficient alkene via a single pericyclic transition state. 1 This reaction has been proposed as a key transformation in the biosynthesis of many cyclohexene-containing secondary metabolites.2-5 However, only four purified enzymes have thus far been implicated in biotransformations that are consistent with a Diels-Alder reaction, namely solanapyrone synthase, 6 LovB, 7,8 macrophomate synthase, 9,10 and riboflavin synthease11,12. Although the stereochemical outcomes of these reactions indicate that the product formation could be enzyme-guided in each case, these enzymes typically demonstrate more than one catalytic activity leaving their specific influence on the cycloaddition step uncertain. In our studies of the biosynthesis of spinosyn A, a tetracyclic polyketide-derived insecticide from Saccharopolyspora spinosa, 13,14 we identified a cyclase, SpnF, that catalyzes a transannular [4+2] cycloaddition to form the cyclohexene ring in spinosyn A. Kinetic analysis demonstrates that SpnF specifically accelerates the ring formation reaction with an estimated 500-fold rate enhancement $(k_{\text{cat.spnF}} \text{ versus } k_{\text{non}})$. A second enzyme, SpnL, was also identified as responsible for the final cross-bridging step that completes the tetracyclic core of spinosyn A in a manner consistent with a Rauhut-Currier reaction 15. This work is significant because SpnF represents the first example characterized in vitro of a stand-alone enzyme solely committed to the catalysis of a [4+2] cycloaddition reaction. In addition, the mode of formation of the complex perhydro-as-indacene moiety in spinosyn A is now fully established.

Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

Author contributions H.-w.L. provided the scientific direction and the overall experimental design for the studies. H.J.K designed and performed most of the experiments. S.H.C participated in the chemical synthesis of the substrate for SpnJ (2) and the characterization of the structures of the enzyme reaction products. M.W.R analyzed the kinetic experiment data. Y.-n.L. carried out the mutation studies of SpnF. H.J.K, M.W.R and H.-w.L. wrote the manuscript.

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Spinosyn A (1), an active ingredient of several highly effective and environmentally benign commercial insecticides, has a complex aglycone structure comprised of a perhydro-*as*-indacene moiety fused to a 12-membered macrolactone.13,14 How this tetracyclic ring system is biosynthesized has been a subject of much speculation.16–18 Attention has largely focused on the construction of the cyclohexene ring due to the potential involvement of an enzyme that catalyzes the [4+2] cycloaddition, which if concerted would represent a so-called "Diels-Alderase." Four genes in the spinosyn A biosynthetic gene cluster of *S. spinosa – spnF, spnJ, spnL*, and *spnM* – were proposed to convert product (2) of the polyketide synthase (PKS) to the tetracyclic aglycone (4) (see Figure 1).16 The gene product of *spnJ*, which is a flavin-dependent dehydrogenase, was recently demonstrated to catalyze oxidation of the 15-OH of 2 to form the keto-intermediate 3.19 However, the functions of the enzymes encoded by the remaining three genes, *spnF*, *spnL*, and *spnM*, which show significant sequence similarity to lipases (SpnM) and *S*-adenosyl-_L-methionine (SAM)-dependent methyltransferases (SpnF and SpnL),16 remain elusive.

To investigate the functions of SpnF, SpnL, and SpnM, the corresponding genes were heterologously overexpressed in *Escherichia coli* BL21(DE3)* and their products purified as *N*-terminal His₆-tagged proteins (>95% purity). Each of the purified enzymes was tested for activity with 3. As shown in Figure 2A, neither SpnF nor SpnL processes 3. In contrast, complete conversion of 3 to a new product was observed after a 2-hr incubation with SpnM. NMR and MS analysis of this new product (8) revealed a 15,6,5-tricyclic skeleton. Further investigation of the reaction time course led to the discovery of a transient intermediate (Fig. 2D, orange peak), which was identified by spectral analysis as the monocyclic macrolactone 5.

These findings suggested that the SpnM-catalyzed conversion of **3** to **8** might be a two-step process involving 1,4-dehydration of **3** followed by a transannular [4+2] cycloaddition between the 11,12-alkene and the conjugated 4,5, 6,7-diene of intermediate **5** to form the cyclohexene moiety in **8** (Fig. 4). To investigate whether SpnM is indeed a cyclase of dual function catalyzing both the dehydration and cyclization steps, the dependence of the rate of each step on the concentration of SpnM was determined. In this experiment the formation and consumption of **5**, which respectively reflect the dehydration and cyclization steps, was monitored by HPLC as a function of time.

As shown in Figure 3A, rate enhancement of the dehydration step was observed with the increase in SpnM concentration, while the rate of cyclization was unaffected. The full time courses of the formation and decay of 5 were analyzed using numerically integrated coupled rate equations based on different kinetic models (see SI Section S5).20 The observed data was fit best to the integrated rate equations 1a and 1b, which model Michaelis-Menten kinetics for the dehydration step ($V_{\rm SpnM}$ and $K_{\rm SpnM}$) and first order kinetics ($k_{\rm non}$) for the cyclization step. Correlation of the fitted parameters versus the concentration of SpnM (Fig. 3B) reveals a significant dependence of $V_{\rm SpnM}$, but not $K_{\rm SpnM}$ or $k_{\rm non}$, on SpnM concentration. This analysis indicates that SpnM functions only as a dehydratase, and its product (5) can cyclize nonenzymatically to 8 with a first order rate constant of ~ 0.03 min⁻¹.

Having established that SpnM functions as a dehydratase, whereas the [4+2] cycloaddition can proceed nonenzymatically, we next considered the possible involvement of the two remaining gene products, SpnF and SpnL, in the final C–C bond formation between C-3 and C-14. Surprisingly, incubation of SpnF or SpnL with **8**, which was generated *in situ* from **3** in the presence of SpnM, did not produce the aglycone **4** (Fig. 2B). However, when SpnF was added to the assay mixture, a change in the product distribution was noted including rapid disappearance of **5** (Fig. 2B, trace (c)). As shown in Figure 2E, the consumption of **5** with concomitant formation of a product having a retention time consistent with **8** was completed in 20 min in the presence of 10 μ M SpnF, instead of requiring more than 2 hr in its absence (Fig. 2D). The structure of this product was established to be **8** by NMR analysis. Additional controls including examining the effects of denatured SpnF as well as mutated SpnF on the rate of cyclization were also performed. In all cases only native (His₆-tagged) SpnF lead to appreciably increased rates of cyclization (see SI Section S4.4). These results clearly indicate that SpnF is responsible for the observed rate enhancement of the cyclization of **5** to **8**.

To quantify the effect of SpnF on the rate of the cyclization step, the production and consumption of 5 was again monitored by HPLC, this time as a function of SpnF concentration while keeping that of SpnM fixed. As shown in Figure 3C, the rate of conversion of 5 to 8 is clearly enhanced as the concentration of SpnF increases. The data was fit best using the coupled rate equations 2a and 2b, where the cyclization event is modeled as the sum of a first order and a Michaelis-Menten process. In these fits all parameters including the initial concentration of 3 were allowed to float except for $k_{\rm non}$, which was fixed at the fitted value obtained in the absence of SpnF. Correlation of the fitted parameters with SpnF concentration indicates a significant dependence of $V_{\rm SpnF}$ on SpnF concentration (Fig. 3D), which is not true for the $V_{\rm SpnM}$ parameter. These results firmly establish that SpnF is a cyclase catalyzing the conversion of 5 to 8 with an apparent $k_{\rm cat}$ of $14 \pm 1.6~{\rm min}^{-1}~(\pm~{\rm s.e.})$ for an estimated rate enhancement ($k_{\rm cat,spnF}$ versus $k_{\rm non}$) of ~ 500-fold.

This result left *spnL* as the only gene without an assigned function, and it was hypothesized that SpnL plays a role in the transannular cyclization reaction between C-3 and C-14. However, the observed inability of SpnL to convert **8** to the anticipated product (**4**) (Fig. 2B, trace (b)) prompted us to reconsider the sequence of events of the proposed biosynthetic pathway in Figure 1. Previously, glycosylations had been thought to occur on the tetracyclic aglycone (**4**), because transfer of the rhamnose moiety from TDP-β-1-rhamnose (**9**) to 9-OH of **4** by the rhamnosyltransferase, SpnG, had been demonstrated.21,22 However, this observation did not necessarily rule out an alternative pathway in which rhamnosylation of **8** precedes C-3/C-14 cross-bridging, because glycosyltransferases involved in the biosynthesis of secondary metabolites are known to be promiscuous with regard to their substrate specificity.23,24

To test this possibility, the ability of SpnG to accept **8** as a substrate was investigated. As shown in Figure 2C (trace b), incubation of **8** and **9** with SpnG resulted in the disappearance of **8** with the concomitant appearance of a new peak (retention time of 17 min). This new product was identified as **10** by NMR and MS analysis (Fig. 4). Upon addition of SpnL to

this reaction mixture, the peak at 17 min disappeared and a new peak appeared at 27 min (Fig. 2C, trace c). Both transformations were highly efficient. Identification of the new product as 13 was confirmed by high-resolution MS analysis and HPLC co-elution with an authentic sample of 13. These results strongly suggest that 10 rather than 8 is the biological substrate for SpnL, which catalyzes the final cyclization step to generate the perhydro-asindacene core.

The mechanisms by which SpnF and SpnL catalyze their respective cyclization reactions are a point of interest. The SpnF-catalyzed *endo*-mode *syn*-addition of an alkenyl to a dienyl functionality appears consistent with a Diels-Alder reaction; however, confirmation of this hypothesis will require demonstrating that the reaction progresses through a single pericyclic transition state such as **6**. Therefore, a stepwise [4+2] cycloaddition mechanism, e.g., one involving a dipolar intermediate such as **7**, cannot at present be ruled out. In contrast, the C–C bond formation catalyzed by SpnL may involve a Rauhut-Currier mechanism15 consistent with the observation that **10** is susceptible toward nucleophilic addition by a thiol (see SI Section S3.10) forming a covalent adduct that may be structurally analogous to **11** or **12** (see Figure 4), though the specific site of attack remains unknown. While these mechanistic proposals are at present speculative, it is worth noting that Roush and coworkers were able to accomplish their non-enzymatic total synthesis of spinosyn A by exploiting both the transannular Diels-Alder and Rauhut-Currier reactions in an analogous fashion.25 This precedent in chemical synthesis certainly substantiates the feasibility of the mechanisms proposed for the SpnF- and SpnL-catalyzed reactions.

In summary, the biosynthetic pathway for spinosyn A is now fully established (Fig. 4), with the specific functions of SpnM as a dehydratase and SpnF as well as SpnL as the two cyclases in the cross-bridging steps biochemically determined. SpnF represents the first enzyme for which specific acceleration of a [4+2] cycloaddition reaction has been experimentally verified as its only known function. It stands in contrast to macrophomate synthase, for which evidence has been provided suggesting a tandem Michael-Aldol reaction mechanism,26,27 as well as the multifunctional solanapyrone synthase, LovB, and riboflavin synthase, which respectively participate in hydroxyl oxidation,5,6 polyketide synthesis,7,8 and hydride transfer,12 in addition to the [4+2] cycloaddition reactions, the concertedness of which have yet to be verified. For this reason, the SpnF reaction provides a unique system for detailed mechanistic investigation of enzyme-catalyzed [4+2] cycloadditions and the existence of a *bona fide* Diels-Alderase.

Method Summary

All proteins used in this work were overexpressed in *E. coli* BL21(DE3)* (Invitrogen, Carlsbad, CA) and purified by Ni-NTA (Qiagen, Valencia, CA) affinity chromatography. Specifically, SpnF was co-overexpressed with the chaperone protein pair, GroEL/ES, to improve the solubility. Because overexpression of the protein encoded by the originally assigned *spnM* gene16 failed to afford an active soluble protein, the gene sequence was reexamined and revised to include 204 additional base pairs (Supplementary Fig. 2). Overexpression of the revised *spnM* gene produced an active enzyme with significantly improved protein yield. All enzyme reaction products (5, 8, 10, and 13) were extracted with

ethyl acetate or chloroform and purified using silica gel column chromatography or HPLC. Their structures were characterized by 1D- and 2D-NMR spectroscopy and/or highresolution mass spectrometry. In particular, the stereochemistry of 8 was assigned based on its ¹H-¹H NOESY spectrum. All substrate specificity and time course assays were run in 50 mM Tris·HCl buffer (pH 8) at 30 °C. Reaction aliquots were quenched with an excess volume of ethanol after a given incubation time and centrifuged to remove protein. The supernatant was then analyzed by reverse phase HPLC with detection by UV absorbance at 254 nm (Figure 2) or 280 nm (Figure 3). Time course assays also included pmethoxyacetophenone as an internal standard to normalize the peak areas corresponding to 5. Numerical integration of equations (1) and (2) employed the fourth order Runge-Kutta algorithm (Supplementary Ref. 9) following non-dimensionalization of substrate concentration. The resulting simulated progress curves were fit using steepest descent (Supplementary Ref. 10) directly to full time courses of normalized substrate concentration obtained via the HPLC discontinuous assay to provide the kinetic parameters and a concentration normalization factor. Further detail regarding experimental procedures as well as data fitting and analysis is described in the Supplementary Information.

Supplementary Material

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Acknowledgements

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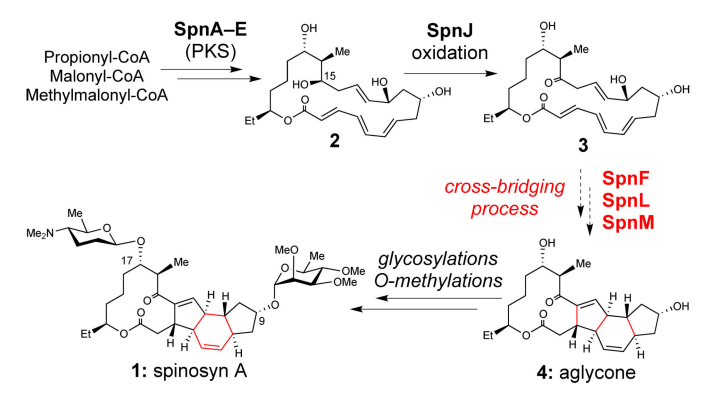


Figure 1. Proposed spinosyn biosynthetic pathway

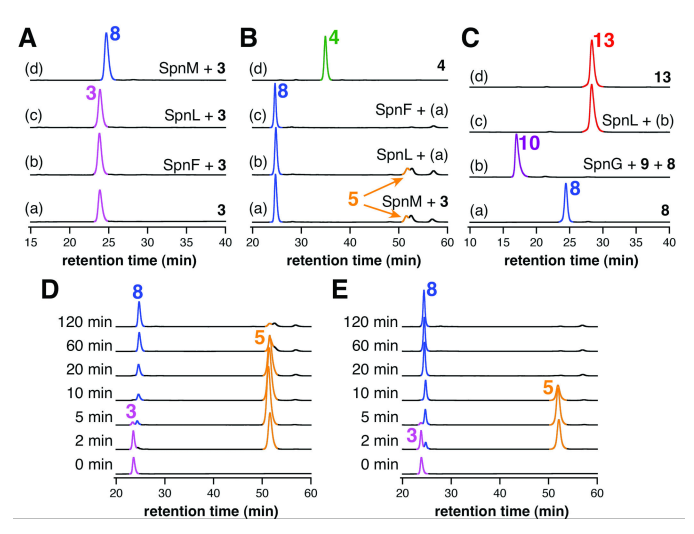


Figure 2. HPLC analysis showing the reactions catalyzed by SpnF, SpnL, and SpnM A. Incubation of 2 mM 3 for 2 hr alone (a), and with either 1.25 μ M SpnF (b), 1.25 μ M SpnL (c), or 1.25 μ M SpnM (d). B. Incubation of 2 mM 8 (generated *in situ* from 3) for 2 hr either alone (a), with 15 μ M SpnL (b), or with 15 μ M SpnF (c). Trace (d) is the authentic standard of 4. C. Incubation of 1 mM 8 for 2 hr either alone (a); with both 20 μ M SpnG and 1.5 mM 9 (b); or with 15 μ M SpnL, 20 μ M SpnG, and 1.5 mM 9 (c). Trace (d) is the authentic standard of 13. D. Time course of 2 mM 3 in the presence of 1.25 μ M SpnM. E. Time course of 2 mM 3 in the presence of both 1.25 μ M SpnM and 10 μ M SpnF.

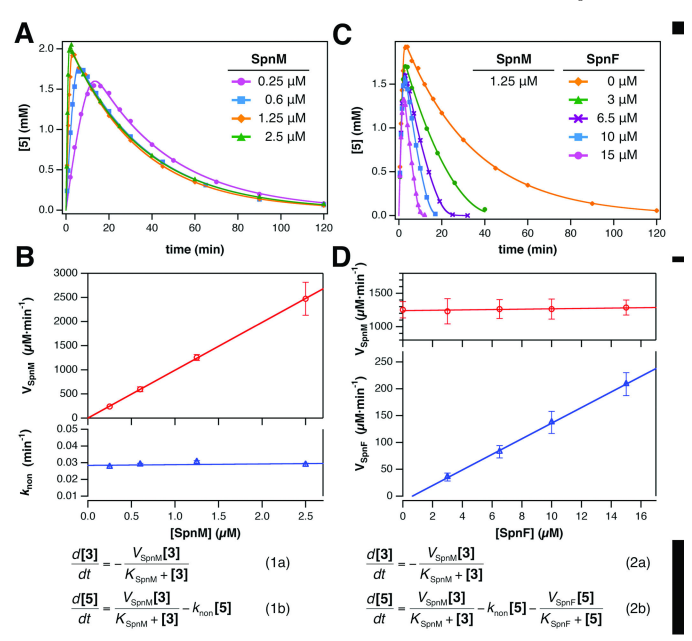


Figure 3. Kinetic analysis demonstrating that SpnM and SpnF respectively catalyze the dehydration and cyclization steps of macrolactone $\bf 3$

Formation and consumption of **5** was monitored by HPLC and the concentration ([**5**]) plotted versus time (**A** & **C**). Each reaction mixture initially contained 2 mM **3**, and the indicated amounts of SpnM and SpnF. **A.** Variable SpnM with no SpnF and fits based on the rate equations (1). **B.** Fitted parameters V_{SpnM} and k_{non} versus [SpnM]. The measured turnover number for SpnM, $k_{\text{cat},\text{SpnM}}$, is $1020 \pm 57 \, \text{min}^{-1}$ while the first order rate constant for nonenzymatic cyclization of **5**, k_{non} , is $0.0288 \pm 0.00041 \, \text{min}^{-1}$. The apparent Michaelis constant for SpnM, K_{SpnM} , is $380 \pm 51 \, \mu$ M. **C.** Variable SpnF at a fixed concentration of SpnM and fits based on rate equations (2). **D.** Correlation of V_{SpnM} and V_{SpnF} versus [SpnF]. The turnover number for SpnF, $k_{\text{cat},\text{SpnF}}$, is $14 \pm 1.6 \, \text{min}^{-1}$. The apparent Michaelis constant

for SpnF, K_{SpnF} , which is independent of SpnF concentration, is $120 \pm 46 \,\mu\text{M}$. All values and error bars are \pm s.e.

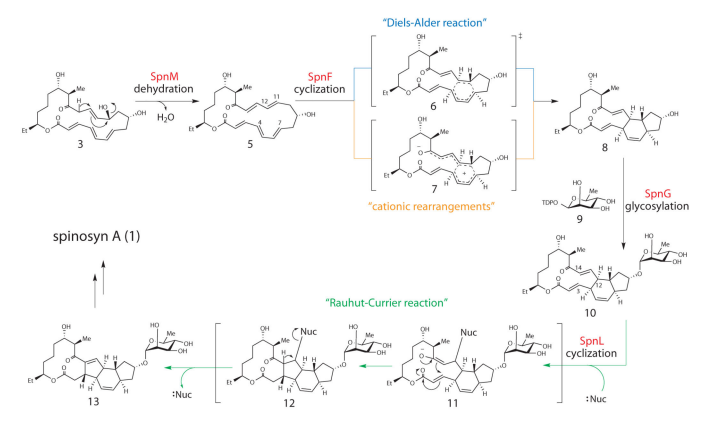


Figure 4. The spinosyn aglycone biosynthetic pathway

SpnM catalyzes a dehydration reaction to convert **3** to **5**, and SpnF subsequently catalyzes cyclization of **5** to afford **8**. The resulting tricyclic macrolactone **8** is then modified with a rhamnose moiety at the C-9 hydroxyl group by SpnG rather than being converted directly to the aglycone **4** (see Scheme 1) as previously thought. SpnL completes the cross-bridging process by interlinking the C-3 and C-14 carbon centers of **10** to produce the tetracyclic nucleus (**13**) of spinosyn A.