Functional dissection and module swapping of fungal cyclooligomer depsipeptide synthetases†

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BbBSLS and BbBEAS were dissected and reconstituted in Saccharomyces cerevisiae. The intermodular linker is essential for the reconstitution of the separate modules. Module 1 can be swapped between BbBEAS and BbBSLS, while modules 2 and 3 control the product profiles. BbBSLS is a flexible enzyme that also synthesizes beaucrivins.

Beaucrivin, beaucrivins A-C and bassianolide (Fig. 1A) are a group of promising anticancer cyclooligomer depsipeptides (CODs) from several fungal species.1 The genes encoding the synthetases for these compounds have been reported.2,3 The beaucrivin synthetase (BbBEAS, 352 kDa) and bassianolide synthetase (BbBSLS, 348 kDa) from Beauveria bassiana ATCC 7159 have the same domain organization of C1-A1-T1-C2-A2-MT-T2a-T2b-C3 (Fig. 1B) and share 66% identity and 79% similarity overall. Most reported fungal COD synthetases (CODSs) have the same structural organization except a bassianolide synthetase from Xylaria sp. BCC1067 that has an additional C-terminal reductase (R) domain.3

The assembly of fungal CODs has been previously proposed.4 CODSs take a unit of α-hydroxycarboxylic acid and l-amino acid to form the corresponding dipeptides. These intermediates will then undergo catalytic cycloligimerization to form the final CODs. BbBEAS synthesizes cyclohexadepsipeptides including beaucrivin and its congeners beaucrivins A-C, while BbBSLS uses α-hydroxyisovaleric acid (α-Hiv) and l-leucine (l-Leu) to generate the cyclohexadepsipeptide bassianolide. We have recently successfully reconstituted BbBEAS and BbBSLS in the intact form in Saccharomyces cerevisiae B15464-NpgA and achieved high-titer production of beaucrivins and bassianolide.5 In this paper, we report the functional dissection and reconstitution of these modular enzymes in S. cerevisiae, characterization of BbBSLS as a flexible enzyme that synthesizes both bassianolide and beaucrivins, module swapping between BbBEAS and BbBSLS, and construction of hybrid CODSs.

The boundary of BbBEAS and BbBSLS between M1 and M2 was analyzed by multiple sequence alignment of nonribosomal peptide synthetases (NRPSs) (Figs. S1 and S2) and a web-based software tool.6 The intermodular linker between M1 and M2 (M1-M2 linker, abbreviated as “linker” hereafter) is shown in Figs. 1B and 1C. We used two Escherichia coli/S. cerevisiae shuttle vectors, pXW55 and pXW06, to express the dissected enzymes in S. cerevisiae BJ5464-NpgA.7 The primers used and plasmids constructed in this study are shown in Tables S1 and S2, respectively (ESI†). The first two expression plasmids for the dissection of BbBEAS were pDY45 and pDY58, which harbor the linker-containing BbBEAS fragments bbBeasM1– and bbBeasM23–, respectively (Fig. 2). Expression of either of these plasmids in the yeast yielded no products. Co-expression of these two plasmids in S. cerevisiae BJ5464-NpgA led to the production of beaucrivins, including beaucrivin and beaucrivins A-C (ESI-MS spectra are shown in Fig. S3), at 10.67±1.44 mg L–1 (Fig. 2). Although the titer is lower than that of the intact enzyme (33.82±1.41 mg L–1), this result for the first time demonstrated that BbBEAS can be dissected into modules and functionally reconstituted in the yeast. To examine whether the intermodular linker influences the interaction and function of the modules, we also cloned the linker-lacking bbBeasM1 and bbBeasM23 to construct two additional expression plasmids pDY99 and pDY98. Co-expression of these two plasmids in the yeast yielded no products (Fig. 2), suggesting that the linker is essential for the reconstitution of beaucrivin biosynthesis by the dissected fragments. We further co-expressed pDY45 with pDY98 and pDY58 with pDY99. Interestingly, the former did not yield any products, while the latter generated beaucrivins in a much lower titer (Fig. 2). These results suggested that the N-terminal linker of BbBEASM23 is required for either the stability of the fragment.
In contrast, the yeast strain EA (Fig. 4, any co-expression). BbBEASM1 and BbBSLS were dissected into two fragments, M1 and M23, with or without the linker. The fragments were co-expressed in S. cerevisiae BJ5464-NpgA and the production of beauvericins and bassianolide was analyzed by LC-MS.

of modules 2 and 3 (M23) or the interaction between M1 and M23. Inclusion of the same linker in the C-terminus of BbBEASM1 significantly improved the collaborative work of M1 and M23, as reflected by the increased titer of beauvericins.

Using a similar approach, we dissected BbBSLS into M1 and M23 (with or without the linker) by constructing four plasmids pDY73, pDY78, pDY97 and pDY102 (Table S2, ESI†). Co-expression of pDY73 and pDY78, which harbor BbBSLSM1– and BbBSLSM23–, respectively, reconstituted the production of bassianolide (ESI-MS spectrum is shown in Fig. S3) in the yeast at 8.87±0.53 mg L⁻¹ (Fig. 2). Unexpectedly, beauvericin and trace amounts of beauvericins A-C were also produced by this strain. To find out whether the production of beauvericins were due to the dissection of BbBSLS, which may have influenced the overall structure of the modular enzyme, we re-examined the product profile of S. cerevisiae BJ5464-NpgA/pDY42. LC-MS analysis revealed that the intact BbBSLS also produced beauvericins (Fig. 3) in addition to bassianolide. In contrast, the yeast strain expressing the intact BbBEAS only produced beauvericins (Fig. 3). This indicated that BbBSLS is a flexible CODS which can incorporate either L-Phe or L-Leu and catalyzes trimerization or tetramerization of the monomers. We also searched for the potential hybrid products containing both N-methyl-L-Leu and N-methyl-L-Phe by extracting the target ion peaks from the ESI-MS spectra. However, no hybrid codons were found. We propose that bassianolide as the major product perfectly fits the active site of BbBSLS. The trimeric beauvericins containing N-methyl-L-Phe which is larger than N-methyl-L-Leu may occupy similar space as the tetramer bassianolide does. The possible hybrid products may not fit the active site well and thus cannot be synthesized efficiently to be detected. In addition, the condensation of the monomers in COD biosynthesis is not well understood, and it is also likely that missing of the hybrid CODSs is due to insufficient capacity of the domains involved in the oligomerization to incorporate monomers containing different amino acid precursors into the products. When pDY102 and pDY97, which respectively harbor the linker-lacking M1 and M23 of BbBSLS, were co-expressed in the yeast, no products were detected. This is consistent with the result for BbBEAS. Co-expression of pDY73 with pDY97 and pDY102 with pDY78 yielded similar results to those for BbBEAS as well (Fig. 2).

Functional dissection of BbBEAS and BbBSLS provides a useful tool for further investigation of these giant enzymes. Expression of truncated fragments of the emniatin synthetase from Fusarium scirpi has been previously attempted in E. coli. However, the proteins were mainly produced as inclusion body and required renaturation for testing their adenylation activity. No CODS have ever been synthesized using these fragments. In this work, we dissected two CODSs into functional modules and reconstituted the production of the corresponding products by co-expression of the modules with the important intermodular linker. This opens a new opportunity to investigate and engineer the functional modules instead of the giant enzymes for combinatorial biosynthesis of new bioactive molecules.

The primary function of BbBEASM1 and BbBSLSM1 is supposed to activate and load t-Hiv onto the CODSs, although we have recently found that BbBEASM1 can also utilize t-Hmv to synthesize beauvericins A-C. We hypothesize that the M1 from BbBEAS and BbBSLS can be exchanged because of the similar function. To test this hypothesis, we made eight co-transformations using the eight expression plasmids constructed above. As shown in Fig. 4, when BbBSLSM1– and BbBEASM23– were co-expressed in the yeast, beauvericins were synthesized. Similarly, co-expression of BbBEASM1– and BbBSLSM23– led to the production of bassianolide and beauvericins. These results clearly indicated that M1 of BbBEAS and BbBSLS can be exchanged without influencing the product profiles. The final products are determined by M23, which controls the selection of the L-amino acid, forms the corresponding dipeptidol, and catalyzes the subsequent cycloligomerization. Additional module swapping experiments that contained different combinations of heterologous M1 and M23, with or without the linker, were conducted to further test the role of the M1-M2 linker in the collaborative work between heterologous modules. As shown in Fig. 4, any co-transformants
containing the linker-lacking BbBEASM23 or BbBSLSM23 yielded no products. Combinations of a linker-lacking M1 with a heterologous linker-containing M23 afforded products corresponding to the M23 (Fig. 4) in relatively low titers. These results indicated that the N-terminal linker of M23 of BbBEAS and BbBSLS is required for the collaborative work of heterologous M1 and M23. It has been previously reported that short communication-mediating (COM) domains play decisive role in protein–protein recognition in bacterial modular NRPSs such as those in tyrocidine biosynthesis. Similarly, intermodular linkers in polyketide synthases are involved in the assembly of functional modules and the interpolyketide chain transfer. Our results revealed that a similar role of the intermodular linker in fungal CODSs. We propose that the C2 domain has specific binding sites for the donor and acceptor that will allow these two thiolated precursors, which are respectively linked to T1 and T2, to recognize and bind to C2, as seen in other NRPSs such as the gramicidin synthetase. In addition to testing module swapping of CODSs using dissected enzymes, we also made two hybrid enzymes BbBEASM1–BbBSLSM23 and BbBSLSM1–BbBEASM23. The linkers between the heterologous M1 and M23 were made up of the C-terminal portion of the linker of M1 and N-terminal portion of the linker of M23 in the original enzymes. As shown in Fig. 3, LC-MS analysis revealed the biosynthesis of beauvericins in S. cerevisiae BJ5464-NpgApDY71 and the production of bassianolide and beauvericins in S. cerevisiae BJ5464-NpgApDY70. These results were consistent with those from the co-expression of the separate modules. These same results from both intact enzymes and co-expressed dissected enzymes further confirmed that BbBEASM1 and BbBSLSM1 can be swapped.

Although module/domain swapping represents an attractive tool to create new molecules, it is often hampered by the heavily impaired biosynthetic capacity of the chimeric NRPSs. Successful construction of hybrid CODSs in the intact form indicated that these fungal enzymes are friendly to genetic manipulations. There were several reports on the construction of bacterial hybrid NRPSs. For example, hybrid NRPSs were made by fusions of modules from TycA, TycB and TycC involved in tyrocidine biosynthesis. Another paper has reported the construction of a new bi-modal NRPS by replacing an activation domain with one with N-methylation activity using the actinomycin NRPS. While these enzymes were made of modules from the same biosynthetic systems, other studies on intermodular linkers/COM domains have allowed the crosstalk between modules from different NRPSs. However, no module-swapping work on fungal NRPSs has been reported. This work thus represents the first example of construction of functional hybrid fungal CODSs, which provides a comprehensive platform for future engineering of fungal CODSs in both intact and dissected forms for novel "unnatural" natural products.

In summary, this work revealed that modular CODSs from fungi can be functionally dissected into modules and expressed in S. cerevisiae. Co-expression of the dissected M1 and M23 of these CODSs reconstituted the production of their products. The linker between M1 and M2 and plays a pivotal role in the reconstitution of the dissected CODSs. BbBEASM1 and BbBSLSM1 are exchangeable and the final products are determined by M23. BbBSLS is a flexible CODS that synthesizes both bassianolide and beauvericins.

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<tr>
<th>Plasmids</th>
<th>Module co-expression</th>
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<tr>
<td>pDY73+pDY58</td>
<td>M1+M23</td>
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<td>Beauvericins 7.2±0.34</td>
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<td>Beauvericins 5.5±0.54</td>
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<td>pDY5+pDY78</td>
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<td>Beauvericins 0.4±0.15 Bassianolide 2.1±0.12</td>
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Fig. 4 Module swapping between BbBEAS and BbBSLS. Heterologous modules from BbBEAS and BbBSLS were co-expressed in S. cerevisiae BJ5464-NpgA and the products were analyzed by LC-MS.

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Notes and references