New tools for reconstruction and heterologous expression of natural product biosynthetic gene clusters

Yunzi Luo, Behnam Enghiade and Huimin Zhao*

Covering: 2010 to 2015

Natural product scaffolds remain a major source and inspiration for human therapeutics. However, generation of a natural product in the post-genomic era often requires reconstruction of the corresponding biosynthetic gene cluster in a heterologous host. In the burgeoning fields of synthetic biology and metabolic engineering, a significant amount of efforts has been devoted to develop DNA assembly techniques with higher efficiency, fidelity, and modularity, and heterologous expression systems with higher productivity and yield. Here we describe recent advances in DNA assembly and host engineering and highlight their applications in natural product discovery and engineering.

1. Introduction

Natural products produced by bacteria, fungi, and plants have been an immense reservoir for new pharmaceuticals, therapeutic agents and industrially useful compounds for centuries. To date, natural products and their derivatives command a substantial pharmaceutical market share, comprising 61% of anticancer drugs and 49% of anti-infective drugs in the past 30 years. Thus the discovery and engineering of the related biosynthetic gene clusters have been intensively studied. Heterologous expression of these biosynthetic gene clusters represents one of the most effective strategies for natural product discovery. With the advent of fast and relatively inexpensive next-generation DNA sequencing technologies, new tools for heterologous expression have been developed to investigate the vast amount of genome sequences. The heterologous expression process is typically comprised of three steps: target biosynthetic gene cluster identification, pathway design and expression, and product detection (Fig. 1). Briefly, target natural product biosynthetic gene clusters are identified from sequenced genomes using powerful computational tools such as AntiSMASH. Various DNA assembly tools, along with engineered heterologous hosts are then used for reconstruction and expression of those large biosynthetic gene clusters. To identify the target natural products, the resulting metabolite profiles are evaluated and characterized by advanced powerful metabolomics and detection techniques.

In this highlight, we will focus only on advances in the development of new design and expression strategies in the recent five years (2010–2015), including DNA assembly tools that have been developed for reconstruction of large biosynthetic gene clusters and new technologies that have recently emerged to enable better regulation control and host engineering in diverse microbial systems. With these new synthetic biology tools, microbes can now potentially be designed/redesigned with a great speed for engineered biosynthesis of natural products.

2. Pathway reconstruction in heterologous hosts

The rapid development of genomics in the past decade has revised our view of the biosynthetic potential and metabolic capabilities of microorganisms. Many organisms have the biosynthetic capability to produce many more natural products than what was originally expected, which sparks a renaissance in natural product discovery. However, most of these putative new natural product biosynthetic gene clusters are only predicted by bioinformatics analysis of sequenced genomes, and their corresponding natural products have not been determined because they are either not expressed or expressed at very low levels under laboratory conditions. As genetic manipulation is either difficult or yet-to-be established for the majority of those organisms, heterologous expression of a single gene, a cassette of genes, or an entire biosynthetic gene cluster in a genetically
tractable host is a practical alternative route for identifying and engineering the corresponding natural products.\textsuperscript{11,12} Therefore, in order to introduce those natural product biosynthetic gene clusters into heterologous hosts, tools for rapid reconstruction of those gene clusters in microbes are essential (Table 1).

2.1 Pathway construction via DNA assembly methods

A variety of \textit{in vitro} and \textit{in vivo} DNA assembly tools have been developed for efficient and fast pathway construction\textsuperscript{7} (Fig. 2). Currently, the focus has been shifted to the development of automated or automation-compatible DNA assembly tools that can be implemented in a high-throughput manner. For example, Marillonnet and coworkers recently developed a modular cloning (MoClo) system based on type IIs restriction enzymes to assemble multiple DNA fragments in a defined linear order. This cloning system was based on Golden Gate cloning technology, which takes advantage of the ability of type IIs restriction enzymes to cleave outside of their recognition sites. Other DNA assembly methods, such as the DNA assembler method based on yeast homologous recombination mechanism, are also widely used to assemble large DNA constructs, in most cases DNA fragments are required to be free of corresponding restriction sites. Other DNA assembly methods, such as the DNA assembler method based on yeast \textit{in vivo} homologous recombination mechanism, are also widely used to assemble large DNA constructs for production of known biosynthetic pathways via synthetic biology tools. She has published 12 papers in peer-reviewed international journals and 2 book chapters.

Dr Yunzi Luo received her Ph.D. in Chemical and Biomolecular Engineering from the University of Illinois at Urbana and Champaign in 2014. Currently, she is a new PI in the State Key Laboratory of Biotherapy/Collaborative Innovation Center of Biotherapy, West China Hospital, Sichuan University, Chengdu, P. R. China. She is interested in engineering biological systems, especially natural products and allows high-throughput assembly of multiple genetic elements.\textsuperscript{14} Recently, Dueber and coworkers adapted this system to establish a versatile platform for yeast, which includes a rapid, modular assembly method and a basic set of characterized parts that can be used in yeast. Additionally, genome-editing tools for making modifications directly to the yeast chromosomes are also included.\textsuperscript{15} Another novel method called “MASTER Ligation” was developed towards high-throughput seamless DNA assembly \textit{in vitro}. By using one restriction endonuclease, MspJI, which specifically recognizes methylated 4-bp sites, mCNNR (R = A or G), and generates a 4-bp arbitrary overhang, multiple DNA sequences can be seamlessly assembled through a simple and sequence-independent hierarchical procedure. By applying this method, the 29 kb actinorhodin biosynthetic cluster of \textit{Streptomyces coelicolor} was successfully assembled.\textsuperscript{16} As it avoids cuts on corresponding type IIs restriction sites within the fragments as in the MoClo method, the MASTER method is more suitable for assembling large DNA constructs.

Although restriction enzyme-based DNA assembly methods can assemble multiple DNA fragments into relatively large constructs, in most cases DNA fragments are required to be free of the corresponding restriction sites. Other DNA assembly methods, such as the DNA assembler method based on yeast \textit{in vivo} homologous recombination mechanism, are also widely used to assemble large DNA constructs for production of known

Dr Huimin Zhao is the Centennial Endowed Chair Professor of chemical and biomolecular engineering, and professor of chemistry, biochemistry, biophysics, and bioengineering at the University of Illinois at Urbana-Champaign (UIUC). He received his B.S. degree in Biology from the University of Science and Technology of China in 1992 and his Ph.D. degree in Chemistry from the California Institute of Technology in 1998. Prior to joining UIUC in 2000, he was a project leader at the Industrial Biotechnology Laboratory of the Dow Chemical Company. Dr Zhao has authored and co-authored over 220 research articles and 20 patents. His primary research interests are in the development and applications of synthetic biology tools to address society’s most daunting challenges in human health and energy, and in the fundamental aspects of enzyme catalysis and gene regulation.
### Table 1  Comparison of the utility of the different approaches for pathway construction

<table>
<thead>
<tr>
<th>Approach</th>
<th>Principle of method</th>
<th>Efficiency</th>
<th>Size assembled (numbers of fragments)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MoClo system</td>
<td>Based on Golden Gate cloning</td>
<td>90–100% for 10 fragments</td>
<td>50 kb (68 fragments)</td>
</tr>
<tr>
<td>MASTER ligation</td>
<td>Using MspJI, which specifically recognizes methylated 4-bp sites and generates a 4-bp arbitrary overhang</td>
<td>NA</td>
<td>29 kb</td>
</tr>
<tr>
<td>DNA assembler (any-gene-any-plasmid (AGAP))</td>
<td>Based on yeast in vivo homologous recombination on mechanism</td>
<td>100% for 2 fragments</td>
<td>50 kb</td>
</tr>
<tr>
<td>Reiterative recombination</td>
<td>Endonuclease-induced homologous recombination in conjunction with recyclable markers</td>
<td>99% for less than 10 kb</td>
<td>Less than 10 kb, could be larger</td>
</tr>
<tr>
<td>Site-specific recombination-based tandem assembly (SSRTA)</td>
<td>Based on øBT1 integration system</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RecET-mediated linear-plus-linear homologous recombination (LLHR)</td>
<td>Linear-linear homologous recombination</td>
<td></td>
<td>10–52 kb</td>
</tr>
<tr>
<td>Phage øBT1 integrase-mediated site-specific recombination</td>
<td>Requiring both homologous and site-specific recombinations. The homologous recombinations were used for targeted integration of the mutated attB and attP into Streptomyces chromosome, while the øBT1 integrase-mediated site-specific recombination was employed to excise targeted region of interest from the chromosome</td>
<td>90% for 23 kb, 80% for 157 kb</td>
<td>157 kb</td>
</tr>
<tr>
<td>Transformation-associated recombination (TAR)</td>
<td><em>In vivo</em> homologous recombination of <em>Saccharomyces cerevisiae</em></td>
<td></td>
<td>21.3 kb and 67 kb</td>
</tr>
</tbody>
</table>

**Fig. 1** Overview of the heterologous expression strategy for natural product discovery. The heterologous expression process is typically comprised of three steps: target biosynthetic gene cluster identification, pathway design and expression, and product detection.
or unknown natural products (Fig. 2). Recently, the efficiency, fidelity, and modularity of the DNA assembler method were greatly improved, which opens doors to a wide variety of applications. More attention has been paid to design issues, such as whether the target gene clusters need to be refactored for better expression in heterologous hosts and which transcriptional units should be chosen for refactoring these pathways. Usually, well-characterized promoters, ribosomal binding sites (RBSs), and terminators are used for pathway refactoring. Promoter libraries are created either by generation of synthetic promoters based on one strong constitutive promoter or identification and characterization from central metabolism pathways via RNA-seq technique. Shao and coworkers developed a novel plug-and-play platform in which a set of constitutive promoters, which were proven functional in the target expression host, were assembled upstream of each pathway gene to refactor silent biosynthetic gene clusters. Using this strategy, a silent spectinabilin biosynthetic gene cluster from *Streptomyces orinoci* was successfully activated. Similarly, a cryptic polycyclic tetramate macrolactams (PTMs) biosynthetic gene cluster from *Streptomyces griseus* was also successfully activated. Reiterative recombination is another robust method for building multi-gene pathways based on homologous recombination. This method uses endonuclease-induced homologous recombination in conjunction with selectable markers. Following endonuclease cleavage of the acceptor module, the donor module’s fragment would be added into the acceptor’s module via homologous recombination, meanwhile the acceptor module’s endonuclease cleavage site and selectable marker will be replaced simultaneously. Reiterative recombination provides a highly efficient and simple strategy for sequentially assembling an indefinite number of DNA constructs at a defined locus. Another related method called site-specific recombination-based tandem assembly (SSRTA) uses the *Streptomyces* phage φBT1 integrase. In this one-step approach, multiple DNA modules are flanked by pairs of non-compatible recombination sites of the φBT1 integration system. A series of mutated attB sites are placed upstream of each module, and mutated attP sites are located downstream. After incubation of all the DNA modules with the φBT1 integrase, tandem assembled products are produced. This approach can efficiently and accurately join multiple DNA molecules in a defined order in vitro.

Meanwhile, approaches for assembling multiple DNA fragments from fungal genomes have been developed as well. One method uses fusion PCR to amplify genes from genomic DNA of a target fungus, place them under control of alcA(p), and fuse them into fragments that can be used to transform *Aspergillus nidulans*. By applying this approach, all of the non-reducing polyketide synthase (NR-PKS) genes from *Aspergillus terreus* along with additional genes required for production or release of the NR-PKS products were successfully transferred into *A. nidulans*. Similarly, overlap extension PCR and yeast homologous recombination were used to clone five desired fungal polyketide synthase and one nonribosomal peptide synthetase
(NRPS) genes (5–20 kb) from various fungal species, into a yeast expression vector and corresponding bioactive molecules were produced in an engineered <i>Saccharomyces cerevisiae</i> host strain.\(^{25}\)

2.2 Pathway construction via direct cloning methods

Many prokaryotic secondary metabolites are produced from polyketide synthase (PKS) and NRPS pathways, which are relatively large gene clusters. Apart from the above-mentioned <i>in vitro</i> and <i>in vivo</i> DNA assembly techniques, methods for directly cloning a target gene cluster into an expression vector have also been developed, which are mostly based on different site-specific recombination systems consisting of a specialized recombinase and its target site\(^{26}\) (Fig. 2). One such direct cloning method is based on the discovery that the full-length Rac prophage protein RecE and its partner RecT can mediate highly efficient linear–linear homologous recombination.\(^{27}\) Using this method, ten megsynthetase gene clusters (each 10–52 kb in length) from <i>Photorhabdus luminescens</i> were cloned into expression vectors and two of them were expressed in a heterologous host which led to the identification of luminmycin A and luminmide A/B.\(^{27}\) More recently, another strategy based on phage øBT1 integrase-mediated site-specific recombination was developed, and used for simultaneous <i>Streptomyces</i> genome engineering and cloning of antibiotic gene clusters.\(^{28}\) Using this strategy, the large actinorhodin gene cluster from <i>Streptomyces coelicolor</i> M145 and the napsamycin gene cluster and daptomycin gene cluster from <i>Streptomyces roseosporus</i> NRRL 15998 were successfully cloned with a success rate higher than 80%.\(^{28}\) Similarly, using the recombination-based cloning strategy, fungal heterologous expression vectors that encode the cryptic clusters from <i>Derma</i> tophyles were constructed in yeast and integrated into <i>Aspergillus nidulans</i>, which led to the production of neosartoricin B.\(^{29}\) It is noteworthy that similar to the DNA assembly tools, promoters and other gene regulation elements can also be inserted into a target gene cluster during the direct cloning process. For example, a simplified plug-and-play system for promoter knock-in was recently developed.\(^{30}\) Promoter cassettes harboring different selection markers were incorporated into the target gene clusters based on the transformation-associated recombination (TAR) cloning in yeast.\(^{30}\)

3. Pathway regulation and engineering in heterologous hosts

After the target natural product biosynthetic gene clusters are correctly assembled and introduced into a heterologous host, regulation of the resulting heterologous pathways should be considered.\(^{31}\) Because the host cells may grow poorly due to metabolic burdens caused by overexpression of a large number of exogenous pathway enzymes or synthesis of toxic intermediates, optimization and regulation of the introduced pathways is often desired.

To eliminate or minimize the accumulation of toxic intermediates, dynamically regulated promoters can be used to either lower or increase expression of the corresponding genes. Native promoters that respond to the toxic pathway intermediates can be identified from genome-wide transcriptional analysis, and then be engineered to dynamically regulate the related metabolic pathway. For example, farnesyl dipiphosphate (FPP) and 3β-hydroxy-3-methylglutaryl-CoA (HMG-CoA) are toxic in <i>E. coli</i>. By using two FPP responsive promoters, high levels of amorphaadiene were produced without use of any inducible promoters.\(^{32}\) Similarly, high concentration of malonyl-CoA is toxic to cells while low concentration of malonyl-CoA reduces fatty acid production. A negative feedback circuit was designed to control the production of malonyl-CoA.\(^{33}\) In this negative feedback circuit, a series of malonyl-CoA responsive promoters are combined with the <i>lacI-T7</i> regulation system for expression of the acetyl-CoA carboxylase. In presence of excess malonyl-CoA, malonyl-CoA responsive promoters turns on LacI expression and hence down-regulates expression of acetyl-CoA carboxylase, thus lowering the production of malonyl-CoA (Fig. 3).\(^{33}\) Similarly, a dynamic sensor-regulator system (DSRS) was designed and constructed to produce fatty acid-based products in <i>E. coli</i>. The DSRS uses a transcription factor that senses a key intermediate (fatty acyl-CoA in the FAEE biosynthetic pathway) and dynamically regulates the expression of genes involved in biodiesel production. The titer of biodiesel-producing strains was increased to 1.5 g L\(^{-1}\) by DSRS.\(^{34}\) Another novel approach to increase production of a specific metabolite called feedback regulation evolution of phenotype (FREP) was developed in <i>E. coli</i>. In this approach, a sensor–actuator circuit is built that dynamically controls the rate of mutation in the whole genome. When the concentration of the target metabolite is low, the mutation rate increases. As a result, an increased mutation rate generates diversity in the population that can lead to evolution of new phenotypes. When the concentration of the target metabolite increases, it causes a reduction in the mutation rate by feedback regulation and hence decreases phenotype diversity. By using this method, the titers of tyrosine and lycopene were increased by 5 and 3 fold respectively compared to the parent <i>E. coli</i> strains.\(^{35}\)

In the case of the production of an antimalarial agent artesiminin, overexpression of every gene from the mevalonate

---

**Fig. 3** Construction of a negative malonyl-CoA feedback regulatory circuit to regulate fatty acid production. The expression of the <i>accABCD</i> genes is controlled by a LacI-repressed <i>pT7</i> promoter. LacI expression is controlled by the <i>PFR1</i> that was repressed by FapR, which was further controlled by malonyl-CoA. When excess malonyl-CoA is accumulated, the biosensor turns on the expression of <i>lacI</i>, which downregulates <i>accABCD</i> expression, alleviating toxicity caused by <i>accABCD</i> overexpression.
pathway to ERG20 (FPP synthase) in *S. cerevisiae* CEN.PK2 led to a 10-fold higher amorpha-4,11-diene production. However, the production of artemisinic acid was only two-fold higher. Expression of the cytochrome p450 enzyme responsible for oxidation of amorphadiene and its cognate reductase at the same level resulted in significant decrease in host viability, which was assumed to be due to the release of reactive oxygen species generated by poor coupling between the cytochrome p450 and its reductase. By optimizing the expression ratio between these two genes, improved production of artemisinic acid (25 g L⁻¹) was achieved in the engineered *S. cerevisiae* strain. In a subsequent study, a novel multivariate-modular approach based on experimental design for systematic pathway optimization was successfully used to improve the production of amorphaadiene in *E. coli* by more than 15-fold (201 mg L⁻¹).

### 4. Host engineering: minimized genomes and engineered hosts

In principle, many microbes can be used as hosts for heterologous expression of natural product biosynthetic gene clusters. However, microbes with clean background and high level of precursors required for the biosynthesis of target natural products will greatly facilitate the discovery and characterization of the target natural products. Currently, there are still several limitations for heterologous expression of a target biosynthetic gene cluster. For example, firstly, only a handful model strains are available with full sets of genetic manipulation tools. Therefore, while introducing biosynthetic gene clusters from distant related species, researchers need to consider whether the codons are optimized to help expression in the chosen heterologous hosts, as well as whether suitable substrates are provided in the chosen strain. Secondly, the introduction of a heterologous biosynthetic gene cluster may disturb the original metabolic balance, and the products or intermediates from the heterologous gene cluster may be toxic. Thus, how to fine-tune the expression and regulation of the imported heterologous within the whole cell network is another key issue to be addressed. As a potential solution to this key issue, inducible promoters may be incorporated to control the release of toxic intermediates and well-characterized promoters or RBSs with different strengths may be used to modify the flux balance. Thirdly, the determination of the minimal sets of genes required for the production of the final products always requires several error and trial. Therefore, there is a need to engineer the hosts for better heterologous expression of the target natural product biosynthetic gene clusters.

*E. coli* has been used for heterologous expression of natural product biosynthetic gene clusters. For instance, the cDNAs of 11 sesquiterpene synthetases from the Jack O’Lantern fungus *Omphalotus olearius* were expressed in *E. coli* and the resulting metabolites were analyzed to identify a series of sesquiterpene compounds. However, its role in heterologous expression of natural product biosynthetic gene clusters from Gram-positive bacteria, especially actinomycetes, is rather limited. Therefore, several *Streptomyces* hosts have been engineered to serve as heterologous expression hosts. Since *Streptomyces avermitilis* is used for industrial production of avermectins, the microorganism is already optimized for efficient supply of primary metabolic precursors and biochemical energy to support multistep biosynthesis, which makes it an excellent host for heterologous expression of natural product biosynthetic gene clusters. To obtain a cleaner background, 1.5 Mb of the sub-telomeric region of *S. avermitilis* which does not contain essential genes for host growth was deleted by a Cre–loxP site-specific recombination system (Fig. 4a). The engineered strains were further improved by removing other endogenous pathways responsible for production of known secondary metabolites. The deletions in total resulted in elimination of 78% of the putative transposase genes, which was expected to further improve the stability of the engineered mutants. Introduction of...
intact natural product biosynthetic gene clusters from other strains resulted in higher production in the engineered S. avermitilis strain. In a subsequent study, more than 20 biosynthetic pathways derived from sugar pathway (e.g. streptomycin), polyketide pathway (bacilomycin B1), amino acid pathway (holomycin), shikimate pathway (chloramphenicol) and MVA or MEP pathway (amorpha-1,4-diene) were evaluated in the engineered host. 18 out of 25 intact biosynthetic gene clusters were successfully expressed. Generally speaking, due to less competitive endogenous pathways for precursors in the genome-minimized strain, production levels of exogenous biosynthetic pathways were higher than the parent strains. Similar studies were performed using genomic mutations and synthetic antisense RNA for overproduction of various other natural products. For example, a 114% increase in L-tyrosine titer was achieved by overproduction of the 1-tyrosine titer was achieved by screening two separate libraries of mutant global transcription factors, RpoA and RpoD. Subsequent transcriptional analysis and whole genome sequencing allowed complete phenotype reconstruction from well-defined mutations and point to important roles for both the acid stress resistance pathway and the stringent response of E. coli in imparting this phenotype. As such, cell-wide measurements could be applied to elucidate the genetic and biochemical underpinnings of an engineered cellular property, leading to the total restoration of metabolite overproduction from specific chromosomal mutations.

5. Conclusion and future prospects

Thanks to exponentially accumulating data from genome sequencing projects and continued development of more sophisticated and powerful genetic manipulation techniques, there is a rapidly growing interest in natural product discovery. Reconstruction and heterologous expression of natural product biosynthetic gene clusters represents one of the most effective strategies for discovery, characterization and engineering of novel natural products. Further exploration in natural product research will not only focus on typical natural product “workhorses”, such as actinomycete bacteria and Aspergillus fungi, but also a wider array of underexplored or neglected organisms. As a result, more computational genome-mining strategies need to be developed in order to fully exploit this untapped biosynthetic potential. Meanwhile, continued innovation in design/redesign of the regulation network in heterologous hosts needs to be explored to promote the discovery of novel natural products or provide new insights into natural product biosynthesis. Furthermore, the generation of diverse engineered heterologous hosts is also of great importance as it facilitates the discovery and engineering of natural products.

Even broader applications of the heterologous expression strategy will be enabled by the development of new synthetic biology tools and an enhanced design, build and test cycle. It is expected that the cost of DNA synthesis will continue to decrease rapidly and de novo synthesis of large DNA molecules such as a natural product biosynthetic gene clusters will become increasingly more affordable. Development of a high throughput and fully automated pipeline for DNA assembly, DNA transformation, cell culture, and product isolation and identification may become a game-changing paradigm for natural product discovery. Moreover, the complete redesign and refactoring of a target natural product biosynthetic gene cluster will lead to improved production of the target natural product.
and generation of new analogs of the target natural product with better biological properties.

6. Acknowledgements

We gratefully acknowledge financial support from the National Institutes of Health (GM077596) and National Academies Keck Futures Initiative on Synthetic Biology.

7. References

Highlight