

Research review paper

New strategies and approaches for engineering biosynthetic gene clusters of microbial natural products

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ABSTRACT

With the rapidly growing number of sequenced microbial (meta)genomes, enormous cryptic natural product (NP) biosynthetic gene clusters (BGCs) have been identified, which are regarded as a rich reservoir for novel drug discovery. A series of powerful tools for engineering BGCs has accelerated the discovery and development of pharmaceutically active NPs. Here, we describe recent advances in the strategies for BGCs manipulation, which are driven by emerging technologies, including efficient DNA recombination systems, versatile CRISPR/Cas9 genome editing tools and diverse DNA assembly methods. We further discuss how these approaches could be used for genome mining studies and industrial strain improvement.

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1. Introduction

Microbial natural products (NPs, also referred to as secondary metabolites) and their semisynthetic derivatives continue to be important sources of pharmaceutically active compounds for health care and plant crop protection (Butler et al., 2014; Katz and Baltz, 2016; Newman and Cragg, 2016). These valuable NPs exhibit diverse chemical structures with a broad spectrum of bioactivities, including antibiotic (penicillin), antitumor (bleomycin) and insecticide (avermectins) properties (Fig. 1). In microbial genomes, all of the genes responsible for the biosynthesis, regulation and resistance of NPs are generally compacted into biosynthetic gene clusters (BGCs) that vary in size from a few to >100 kb (Smanski et al., 2016) (Fig. 1). With the rapid advances in microbial (meta)genome sequencing, a large number of cryptic BGCs has been found in bacteria and fungi, which are regarded as an untapped treasure trove for the discovery of new medicines (Baltz, 2016a; Cimermancic et al., 2014; Doroghazi et al., 2014). For example, the most gifted actinomycete strains are capable of producing 30–50 chemical skeletons, approximately ten-fold more than previously reported (Baltz, 2016a). Over the past decade, the development of a variety of powerful computational approaches, such as antiSMASH and Pep2Path, has facilitated the rapid and high-throughput prediction of cryptic BGCs (Medema and Fischbach, 2015; Medema et al., 2014; Weber et al., 2015; Ziemert et al., 2016). More recently, a community standard for the annotation of BGCs (MiBiG) was proposed, providing a generally applicable reference dataset to promote the discovery and identification of novel NPs (Medema et al., 2015).

With the continuing accumulation of uncharacterized BGCs in public databases, the next key step is to rapidly express these cryptic BGCs in suitable hosts. Therefore, it is of great importance to develop a series of powerful tools for manipulating BGCs, including

using Red $\alpha\beta$ or the truncated version of RecET is inefficient at mediating homologous recombination between two linear DNA molecules, hampering its use for direct cloning of target BGCs (Fu et al., 2012).

In 2012, the Müller group surprisingly discovered that the full-length RecE and its partner RecT could mediate high-efficiency linear-linear homologous recombination (LLHR), allowing for the direct capture of large genome regions (Fu et al., 2012) (Fig. 3A). Using this approach, nine polyketide synthase (PKS) and non-ribosomal polypeptide synthetase (NRPS) BGCs (with sizes from 10 to 37 kb) from the gram-negative bacterium *Phototrhobdus luminescens* were directly cloned into linear expression vectors. Two new molecules, luminmycin A and luminmide A/B, were successfully identified in the heterologous *E. coli* host. However, they failed to clone a large *plu2670* gene cluster (52 kb). To address this limitation, a two-step cloning strategy was developed: LLHR was first used to capture target BGCs, and LCHR was then used to introduce a selection marker to reduce background. It is worth noting that, to insure a high efficiency for the LLHR-mediated cloning method, genomic DNA must be cleaved by unique restriction enzymes near the 5' and 3' ends of target BGCs. However, it is not always easy to find suitable restriction enzyme cutting sites (Wang et al., 2016). With the advent of the programmable CRISPR/Cas9 system, which is able to recognize and cut DNA sequences near target BGCs easily, this bottleneck could be overcome (Jinek et al., 2012; Lee et al., 2015).

2.1.2. TAR cloning

Transformation-associated recombination (TAR) cloning, which is based on the native recombination capacity of *Saccharomyces cerevisiae*, was originally developed to isolate large pieces of mammalian genomic

DNA in the 1990s (Kouprina and Larionov, 2016; Larionov et al., 1996; Larionov et al., 1997). In 2010, the Brady group extended TAR cloning to the capture of microbial natural product BGCs (Kim et al., 2010) (Fig. 3A). To facilitate the rapid expression of target BGCs in *Streptomyces*, the yeast-*E. coli*-*Streptomyces* shuttle vector pTARa was constructed containing three elements: a CEN6/ARS4 sequence and a selection marker allowing for single-copy maintenance in yeast, bacterial artificial chromosome (BAC) elements for plasmid enrichment and verification in *E. coli* and shuttle/integration components for expression in *Streptomyces*. Using pTARa, multiple BGCs (>50 kb in size) were directly cloned or reassembled from environmental DNA (eDNA) libraries (Chan et al., 2016; Katz et al., 2016). Afterwards, the Moore group created a similar cloning vector pCAP01, which is derived from the SuperCos1 cosmid equipped with a pUC *ori*. A 67-kb silent non-ribosomal peptide (NRP) gene cluster from the marine actinomycete *Saccharomonospora* sp. *CNQ-490* was captured and successfully activated in the heterologous host *Streptomyces coelicolor* M1146 by remodeling the cluster-situated regulatory genes (Yamanaka et al., 2014).

Although TAR cloning can be used to directly clone BGCs of interest, the method exhibits a very low cloning efficiency (0.5–2%) due to vector recircularization via non-homologous end joining (NHEJ), which leads to time-consuming screening of hundreds of clones. Recently, two different strategies were introduced to increase positive rates in the TAR-mediated cloning method (Lee et al., 2015; Tang et al., 2015). The first strategy is to use a counter-selection marker for colony selection. The *URA3* gene, which encodes orotidine 5'-phosphate decarboxylase and is under control of the strong promoter pADH1, is introduced to pCAP01, generating pCAP03. Because pADH1 can tolerate an insertion

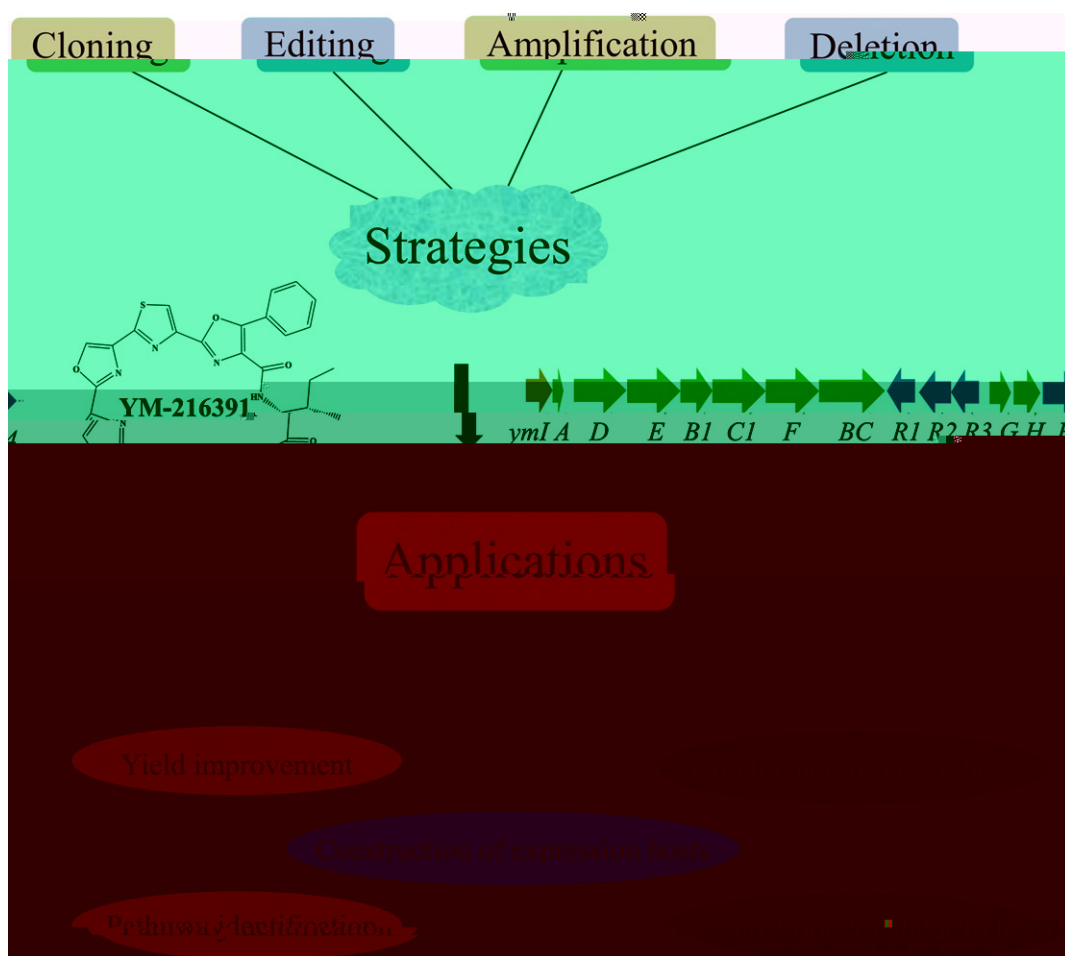


Fig. 2. Overview of the strategies for engineering biosynthetic gene clusters (BGCs) and their applications to NP discovery and development. As an example, an anti-tumor compound YM-216391 and its gene cluster are shown.

the TATA box, yeast transformants that contain pCAP03 with target BGCs can survive in the presence of 5-fluoroorotic acid (5-FOA). Using pCAP03, the Moore group successfully captured two thiotetronic acid BGCs ~30 kb in size with an efficiency of 20–75% (Tang et al., 2015). The second strategy is to use the RNA-guided Cas9 endonuclease to cleave chromosomal DNA instead of restriction enzymes. In the past, the Kouprina group discovered that homologous recombination is more efficient when the linearized capturing vector hooks (homology arms) are located closer to the ends of the target DNA sequences (Kouprina and Larionov, 2006). Although unique restriction enzymes can be theoretically obtained to cleave near the 5'- and 3'-ends of target DNA, it is often difficult to find suitable cutting sites. In 2015, the Kouprina group used the programmable CRISPR/Cas9 system to precisely cleave two sides of the target DNA, significantly improving TAR cloning efficiency up to 32% (Lee et al., 2015) (Fig. 3A). Currently, capturing target chromosomal regions only requires screening fewer than a dozen transformants. It is conceivable that TAR cloning, combined with a counter-selection marker and the CRISPR/Cas9 system, will further accelerate the direct cloning of microbial natural product BGCs.

2.1.3. SSR-mediated cloning

Derived from temperate *Streptomyces* bacteriophages, such as Φ C31 and Φ BT1, a variety of serine integrase-mediated site-specific recombination (SSR) systems has been developed to stably integrate exogenous DNA into the chromosomes of prokaryotes, eukaryotes and archaea (Baltz, 2012). Recently, the Φ BT1 integration system was extended for the cloning of target BGCs in native *Streptomyces* hosts (Du et al.,

2015). First, the paired Φ BT1 integration sites *attB/attP* and the replicative plasmid pKC1139 are individually introduced on either side of the target BGC via two single cross-over recombination events. Then, the Φ BT1 recombinase is expressed, which mediates the cleavage of the two paired integration sites, resulting in circularization of target BGC in the plasmid pKC1139 (Fig. 3B). Using this strategy, three different BGCs, including the actinorhodin (ACT) gene cluster (23 kb) from *S. coelicolor* and the napsamycin (45 kb) and daptomycin (157 kb) gene clusters from *Streptomyces roseosporus*, were successfully acquired with high efficiency (>80%). Then, ACT and daptomycin production were enhanced by increasing the copy numbers of their corresponding BGCs in the native hosts. However, the SSR-mediated cloning method has two major limitations. One limitation is that the procedure is time-consuming and unsuitable for difficult-to-manipulate microorganisms. The second is that the multi-copy plasmid pKC1139 carrying the target BGC is unstable in *Streptomyces*.

2.1.4. CATCH

As described above, *in vivo* cloning strategies have several limitations. LLHR in *E. coli* can only be used to clone BGCs smaller than 50 kb. The other two methods (TAR cloning in yeast and SSR-mediated cloning in *Streptomyces*) are time-consuming due to the slow growth rates of yeast and *Streptomyces*. Recently, a Cas9-associated targeting of chromosome segments (CATCH) method by combining the CRISPR/Cas9 system with Gibson assembly, was established for *in vitro* capture of bacterial genomic segments up to 100 kb in length (Jiang et al., 2015). After excision by the RNA-guided Cas9 endonuclease at two designated

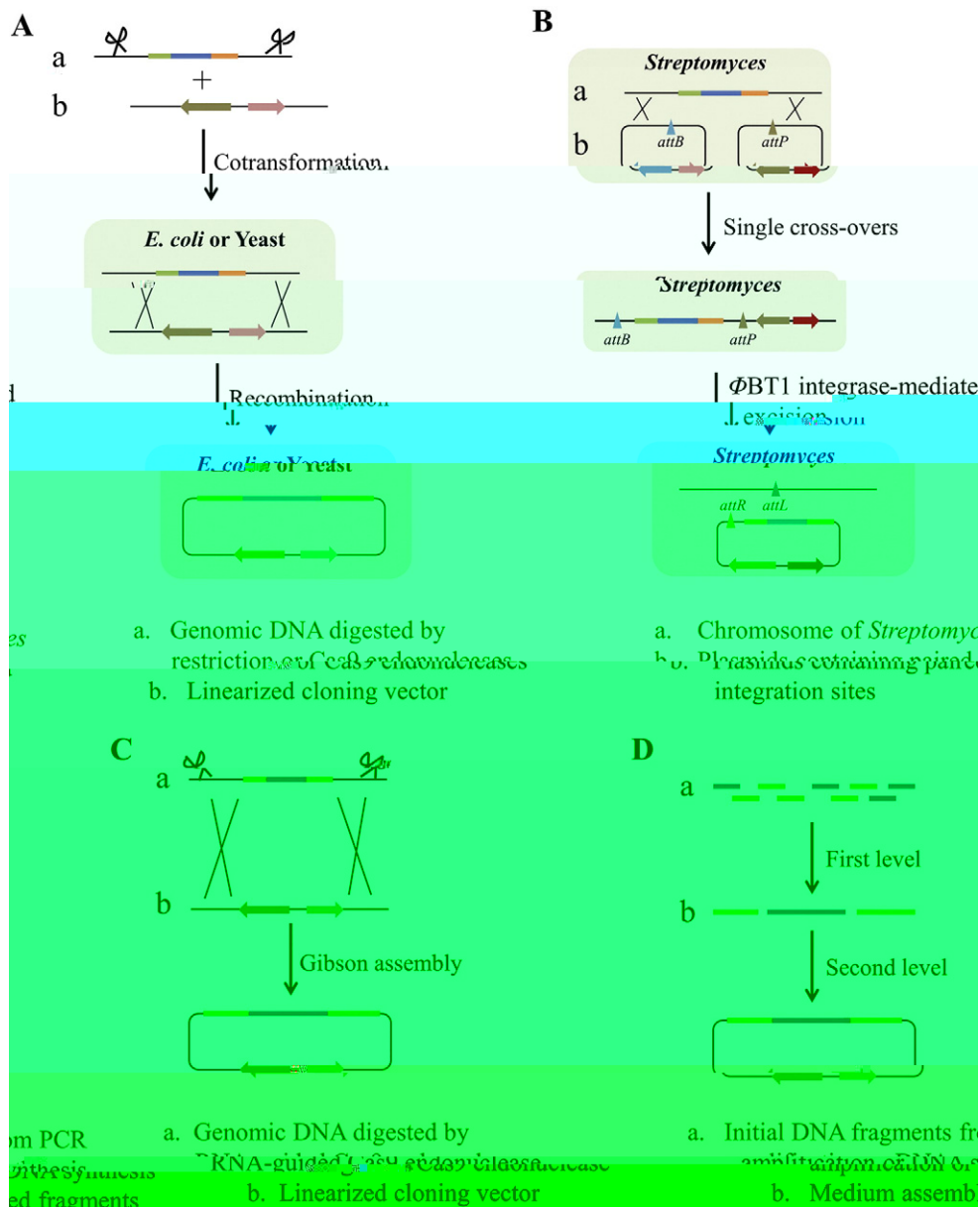


Fig. 3. Approaches for producing large DNA fragments. (A) Direct cloning of target BGCs by Cas9-facilitated transformation-associated recombination (TAR) in yeast or full-length RecET-mediated linear-linear homologous recombination (LLHR) in *E. coli*. (B) Direct cloning of antibiotic gene clusters via ϕ BT1 integrase-mediated site-specific recombination in *Streptomyces*. (C) One-step targeted cloning of large gene clusters by combining the *in vitro* CRISPR/Cas9 system with Gibson assembly (CATCH). (D) Various DNA assembly methods for constructing large DNA fragments, including DNA assembler and Gibson assembly.

chromosomal loci, target DNA fragments can be one-step ligated to the capturing vector in a Gibson assembly mix (Fig. 3C). Using this approach, a 78-kb bacillaene-producing PKS gene cluster from *Bacillus subtilis* was successfully cloned into a BAC vector with a 12% positive rate. Two ~30 kb antibiotic gene clusters from *Streptomyces* were also captured into the p15A vector with an efficiency of approximately 90%. It is noteworthy that this approach requires very careful operation as preparing gel plugs to guarantee the integrity of the genomic DNA and high assembly efficiency (Jiang and Zhu, 2016).

2.2. Bottom-up assembly

With the exception of direct cloning of BGCs, DNA assembly provides an alternative strategy for acquiring target BGCs in a bottom-up manner (Fig. 3D and Table 1). With the rapid development of synthetic biology,

numerous DNA assembly methods have been developed and are widely utilized to generate complex libraries, biosynthetic pathways and even microbial genomes (Ellis et al., 2011; Merryman and Gibson, 2012). Typically, these methods can be divided into three main types: (1) nontypical restriction enzyme-mediated methods, including Golden gate and MASTER ligation (Chen et al., 2013; Engler, 2009); (2) *in vitro* recombination-based assembly methods, including SLIC, CPEC and Gibson assembly (Gibson et al., 2009; Li and Elledge, 2007; Tian, 2009); (3) DNA assembler based on yeast homologous recombination (Shao and Zhao, 2009). Herein, four major DNA assembly methods for obtaining natural product BGCs are reviewed.

2.2.1. MASTER ligation

Type II restriction enzymes, which recognize asymmetric DNA sequences and cleave outside of their recognition sequences, hold great

Table 1
Strategies for cloning or assembling NP biosynthetic gene clusters (BGCs).

Methods	Principles	Sizes	Advantages	Disadvantages	Refs
Direct cloning strategy	Linear-linear homologous recombination (LLHR)	52 kb (PKS-NRPS BGC)	High efficiency (17–50%); user friendly	Low efficiency for cloning large BGCs	(Fu et al., 2012)
	Transformation-associated recombination (TAR)	67 kb (tarmycin BGC)	Cas9-facilitated high efficiency cloning (30%)	Time-consuming	(Lee et al., 2015)
	Site-specific recombination (SSR)-mediated cloning CATCH	157 kb (daptomycin BGC)	Suitable for capturing large and complex BGCs	Time-consuming; instability of BGCs in the vector pKC1139	(Du et al., 2015)
Bottom-up assembly strategy	CRISPR/Cas9 and Gibson assembly	78 kb (Psk BGC)	High efficiency (12–90%); suitable for cloning large genomic regions (>100 kb)	Fine operation is required to guarantee genomic DNA integrity	(Jiang et al., 2015)
	SSRTA	62 kb (epothilone BGC)	Useful for assembly of large DNA with high GC contents	Scars	(Zhang et al., 2011)
	MASTER ligation	29 kb (actinorhodin BGC)	Scarless cloning; user friendly	Low efficiency for assembling large DNA constructs	(Chen et al., 2013b)
	Modified Gibson assembly	67 kb (pristinamycin II BGC)	High efficiency (20–40%) for assembling BGCs with high GC contents	Time-consuming	(Li et al., 2015)
	DNA assembler	45 kb (PKS BGC)	Scarless cloning; high efficiency (30%) for assembling fewer fragments	Time-consuming; instability of homologous regions in yeast	(Shao et al., 2011)

potential for a wide range of applications in the construction of gene libraries and biochemical pathways. For instance, the “golden gate cloning” method that commonly uses *Bsa* I, *Bsm*B I and *Bbs* I can assemble at least nine DNA fragments in a defined linear order in a single step with high efficiency (90%) (Werner et al., 2012). Recently, a similar strategy called MASTER (methylation-assisted tailorable ends rational) ligation based on *Msp*II, a specific type IIs endonuclease, was developed for sequence-independent hierarchical DNA assembly (Chen et al., 2013). *Msp*II only recognizes methylation-specific 4-bp sites, (m)CNR (R = A or G), and can produce four-base 5'-overhangs of any sequence. Using the *Msp*II-mediated method, the ACT biosynthetic gene cluster (29 kb) from *S. coelicolor* was successfully assembled and heterologously expressed in the fast-growing strain *Streptomyces* sp. 4F. However, the assembly efficiency of the MASTER method is relatively low (~6.7%) when constructing large DNA fragments, which indicates that the ligation conditions still require further optimization (Chen et al., 2013).

2.2.2. SSRTA

In addition to the cloning of target BGCs in native hosts as described above, the site-specific Φ BT1 integration system has also been exploited for the *in vitro* assembly of multiple DNA parts, resulting in the establishment of a novel technology named ‘site-specific recombination-based tandem assembly’ (SSTRA) (Fogg et al., 2014; Zhang et al., 2011). Considering that DNA recombination cannot occur between *attB/attP* sites with different central di-nucleotides in the Φ BT1 integration system, the Ding group systematically identified 16 pairs of non-compatible *attB/attP* recombination sites (Zhang et al., 2008). Using these paired, non-compatible *attB/attP* sites, a 62.4 kb epothilone gene cluster was successfully assembled from seven DNA fragments in a single step. However, it is worth noting that this method will introduce multiple scar sequences (*attR*, 42 bp) between the assembly modules, which may be problematic for the efficient expression of target BGCs.

2.2.3. Modified Gibson assembly

The Gibson assembly method exhibits high simplicity and speed to produce large DNA constructs (up to 900 kb), and permit sequence-independent, one-pot assembly of multiple DNA fragments (Gibson et al., 2009). Three enzymes, T5 exonuclease, Phusion DNA polymerase and Taq DNA ligase, are included in the Gibson reaction mix to perform ligation reactions between DNA molecules. However, we found that this method could not be effectively used to assemble DNA fragments with high GC contents (>70%), mainly due to vector self-ligation (Li et al., 2015). It has been proposed that mismatched linker pairings are easily generated between overhangs with high GC contents due to the low reaction temperature (50 °C) (Casini et al., 2014). To address this limitation, a modified Gibson assembly method was developed in our group (Li et al., 2015). First, a pair of universal terminal single-stranded DNA overhangs with high AT contents (21 bp) was added to the ends of the BAC vector. Second, to allow for hierarchical and seamless assembly of large DNA molecules, two restriction enzyme sites were introduced to the respective sides of the designed overlaps. A 67 kb pristinamycin II (PII) gene cluster from *Streptomyces pristinaespiralis* was hierarchically assembled from 15 PCR-amplified fragments. Although the assembly efficiency is increased from 2.5% to 20–40%, the modified Gibson method requires further optimization, such as extending the length of overlaps between the DNA fragments and the vector.

2.2.4. DNA assembler

DNA assembler, an *in vivo* DNA assembly approach based on homologous recombination in *S. cerevisiae*, was recently developed to construct large biochemical pathways with high efficiency (70–100%) (Shao and Zhao, 2009). Afterwards, this robust method was used to

assemble microbial natural product BGCs. Using DNA assembler, expression vectors containing both target BGCs and genetic elements needed for DNA maintenance in different hosts can be generated in a single-step manner (Shao et al., 2011). For example, seven PCR-amplified fragments (4–5 kb) from the aureothin gene cluster (29 kb) were co-transformed with three helper fragments into *S. cerevisiae*, generating a single circular DNA molecule with high efficiency (60%). Furthermore, because the assembled fragments are derived from PCR amplification, the DNA assembler method allows different genetic modifications, such as point mutations and gene deletions, to be easily introduced into target BGCs (Fig. 4A). Several new aureothin derivatives have been generated by introducing point mutation(s) into the DH domain of AurB (Shao et al., 2011). As such, DNA assembler exhibits unprecedented versatility and flexibility for assembling and refactoring natural product BGCs.

3. Strategies for editing BGCs—optimizing NP biosynthetic pattern

After acquiring target BGCs, the next key step is to edit or refactor BGCs to achieve the following three goals: (1) increasing the expression levels of target BGCs for strain improvement; (2) generating new derivatives by combinatorial biosynthesis; (3) activating silent BGCs for novel NP discovery. Currently, the available molecular genetic/synthetic biology approaches applicable to editing BGCs mainly include: (1) Red/ET recombineering in *E. coli* (Jiang et al., 2013); (2) multiplexed CRISPR/Cas9 and TAR-mediated promoter engineering in yeast (Kang et al., 2016); (3) *in vitro* CRISPR/Cas9-mediated editing systems (Liu et al., 2015); and (4) combinatorial design and assembly (Smanski et al., 2014; Temme et al., 2012) (Fig. 4). It should be noted that, as just mentioned, because initial fragments are obtained from PCR amplification or DNA synthesis, DNA assembler can be conveniently used to refactor BGCs (Shao et al., 2011) (Fig. 4A).

3.1. Red/ET recombineering

Red/ET recombineering technology has been exploited for a wide variety of applications (Wang et al., 2016). In addition to LLHR-mediated direct cloning of microbial natural product BGCs, Red/ET recombineering has also been used to edit BGCs to generate novel antibiotic derivatives *via* combinatorial biosynthesis

3.3. mCRISTAR

Large-scale sequencing efforts of the microbial (meta)genome have revealed that most natural product BGCs are not expressed in laboratory fermentation conditions. Along with the new era of genomics-driven NP discovery, activation of these silent BGCs provides a simple and generic strategy to unearthing novel NPs. In recent years, a variety of pleiotropic and pathway-specific approaches has been developed to induce the expression of BGCs, including variations of growth conditions, transcriptional/translational machinery engineering, epigenetic perturbation and manipulation of global or pathway-specific regulators (Rutledge and Challis, 2015). Interested readers are referred to some excellent reviews on the awakening of silent BGCs using such approaches (Chiang et al., 2011; Rutledge and Challis, 2015). Here, we focus on the refactoring of BGCs of interest, especially *via* promoter exchange involving the replacement of natural promoters with constitutive promoters. This straightforward strategy has been widely used to trigger the expression of many NPs from *Streptomyces* and *Burkholderia* by Red/ET recombineering or DNA assembler (Biggins et al., 2011; Franke et al., 2012; Luo et al., 2013; Olano et al., 2014).

Recently, a more powerful tool that combines the CRISPR/Cas9 system and TAR in yeast achieved marker-free, multiplexed replacement of up to four BGC promoters in a single round (Kang et al., 2016) (Fig. 4D). In this mCRISTAR (multiplexed-CRISPR-TAR) method, the CRISPR/Cas9 system is first used to induce DNA double-strand breaks (DSBs) in BGC promoters. The resulting linearized cluster fragments are then reassembled with synthetic promoters containing BGC-specific homology sequences. To avoid inter-promoter recombination, a set of orthogonal, active promoters and RBS sequences with different expression strengths was constructed. It is worth noting that natural CRISPR arrays rather than sgRNA (single guide RNA) were used to simplify the construction of CRISPR expression modules. In addition, a mCRISTAR webapp was developed to automatically generate all of the sequences required for the modified process. With this highly efficient DNA editing method, three silent BGCs, including the tetracycline, lazarusimide and aromatic polyketide AB1210 gene clusters, were rapidly refactored and successfully expressed in the genetically tractable heterologous *Streptomyces albus* host.

3.4. Functional optimization by combinatorial design and assembly

Because of the complex and redundant host regulation of natural product biosynthesis, it is sometimes difficult to achieve expected goals only *via* genetic engineering of BGCs of interest. Rebuilding target BGCs in a bottom-up manner using orthogonal, well-characterized parts provides an alternative approach to bypass all native regulation. As a proof-of-concept, the Voigt group developed a systematic platform to completely refactor the nitrogen fixation gene cluster from *Klebsiella oxytoca* (Smanski et al., 2014; Temme et al., 2012). At first, all seven operons in the native gene cluster were organized and expressed under the control of synthetic regulatory parts, including promoters, RBS and terminators (Temme et al., 2012). To further obtain radically different cluster architectures, Smanski et al. constructed 122 variants of the complete 16-gene pathway, in which 103 genetic parts were combinatorially designed and assembled by varying gene order, gene orientation and operon occupancy. A fully refactored nitrogen fixation gene cluster was finally produced, which recovered 57% of its wild-type activity (Smanski et al., 2014). It is conceivable that this rebuilding approach will provide a useful synthetic biology tool for the functional optimization of BGCs of interest in the future (Smanski et al., 2016).

4. Strategies for amplifying BGCs—improving NP production

Today, microbial NPs remain an important source for novel drug discovery and development; however, their production titers are often very low and must be improved (Baltz, 2016b; Katz and Baltz, 2016;

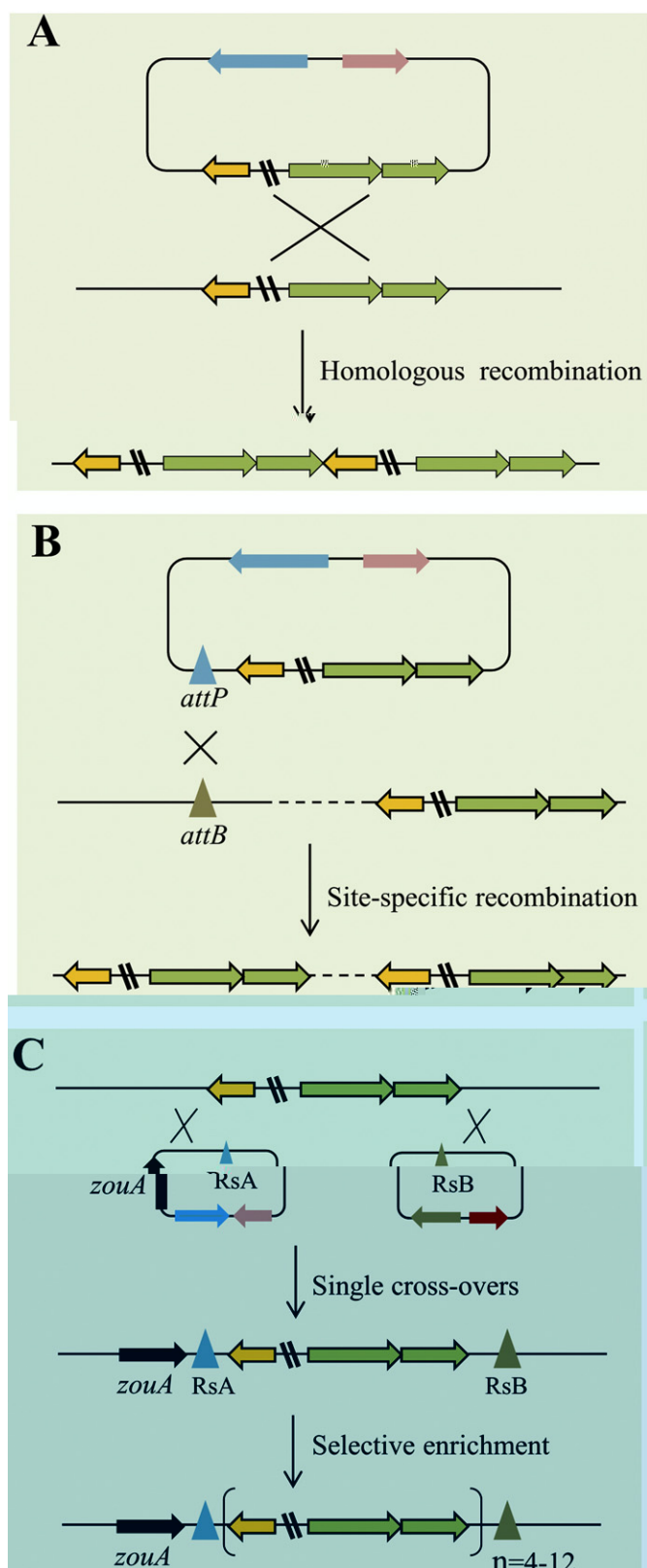


Fig. 5. Tools for the amplification of natural product BGCs in actinomycetes. (A) Homologous recombination-mediated tandem amplification of BGCs in native hosts. (B) Bacteriophage site-specific recombination-mediated chromosomal integration of BGCs in native or heterologous *Streptomyces*. (C) ZouA relaxase-mediated site-specific recombination. This method facilitates the tandem amplification of genomic regions between the *RsA* and *RsB* sites.

Lok, 2015). Over the last two decades, rational metabolic engineering strategies, including increasing precursor supply, manipulating regulatory genes and amplification of BGCs, have been successfully employed to improve the biosynthesis of important NPs in industrial strains or to activate the expression of silent BGCs in heterologous hosts (Baltz, 2016b; Olano et al., 2008; Zhang et al., 2016). As a biotechnological tool, the duplication or amplification of special regions of genomic DNA plays important roles in industrial applications, including NP biosynthesis, bioconversions and degradation of toxic compounds. Intriguingly, some overproducing strains generated *via* traditional mutagenesis and screening, such as *Penicillium chrysogenum*, *Streptomyces kanamyceticus* and *Streptomyces lincolnensis*, contain tandem amplifications of their antibiotic gene clusters (Fierro et al., 1995; Peschke et al., 1995; Yanai et al., 2006).

Till now, three different strategies have been reported for BGC amplification. In general, entire BGCs can be quickly duplicated in native hosts based on homologous recombination (Nah et al., 2015) (Fig. 5A). However, this simple approach only increases one copy of target BGCs, and the resultant strains are typically genetically unstable due to the presence of a tandem amplification. Based on site-specific recombination, two different approaches have been developed for multi-copy amplification of BGCs. The serine integrase-mediated recombination system has been exploited to produce stable recombinants by discrete insertion of target BGCs into actinomycete chromosomes (Baltz, 2012) (Fig. 5B). The relaxase ZouA-mediated DNA recombination system can facilitate the tandem amplification of more than ten BGC copies (Murakami et al., 2011) (Fig. 5C).

4.1. Bacteriophage Att/Int system

Phage-encoded large serine integrases have been widely used in the genetic engineering of actinobacteria, particularly *Streptomyces* (Baltz,

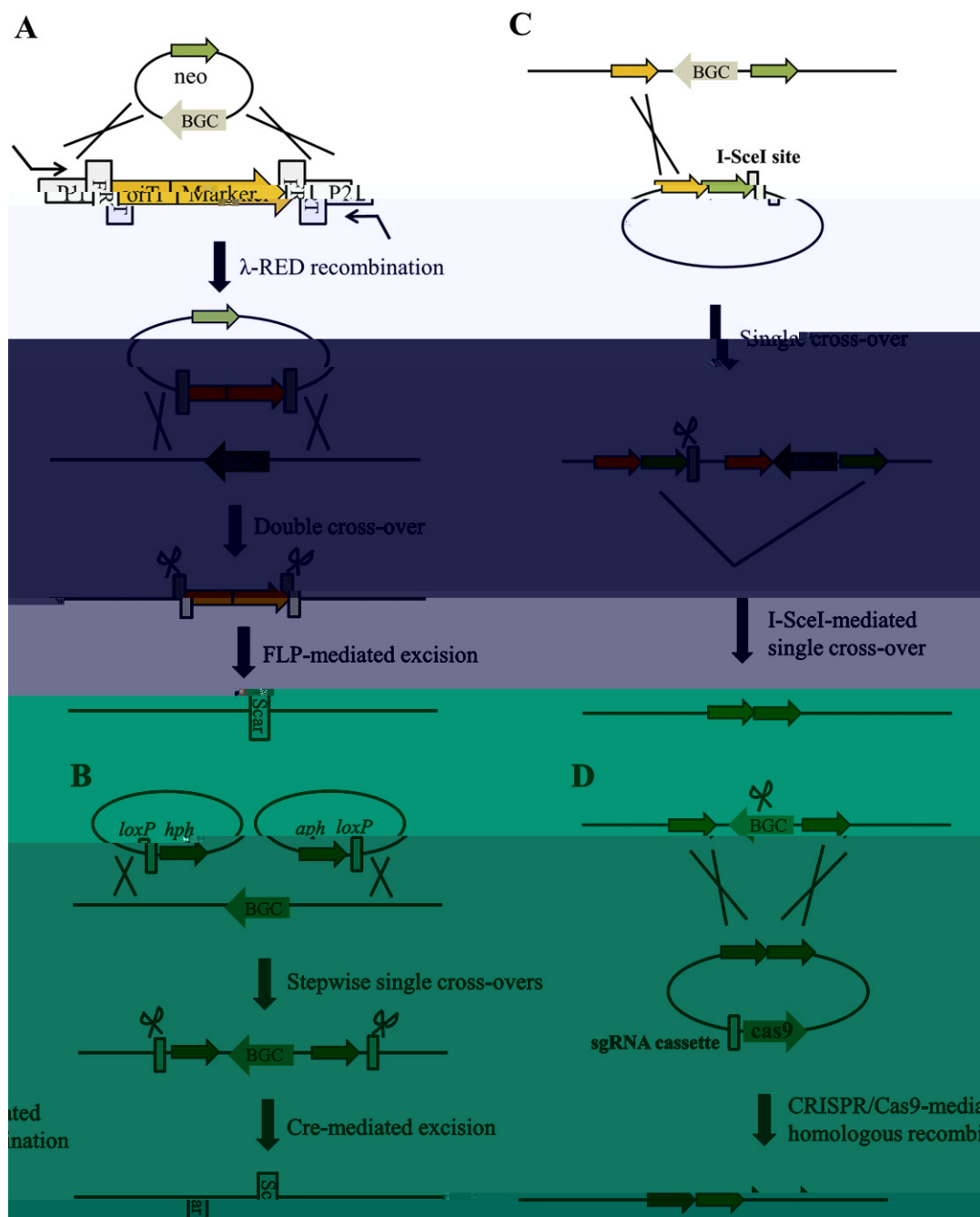


Fig. 6. Four major strategies for deleting large genomic regions in actinomycetes. (A) PCR-targeting method achieved by combining homologous recombination and site-specific recombination. (B) Cre-loxP recombination-mediated deletion of large chromosomal regions. (C) I-SceI meganuclease applied to the markerless deletion of BGCs. (D) Cas9-facilitated one-step deletion of large-size BGCs.

(Bentley et al., 2002; Gust et al., 2003). Three steps are usually required in the PCR-targeting method: (1) the target gene within the cosmid in *E. coli* is replaced with the disruption cassette (containing a selectable marker flanked by *FRT* or *loxP* sites); (2) after the mutant cosmid is transferred into *S. coelicolor*, the single gene can be efficiently deleted by a double cross-over event; (3) the disruption cassette, flanked by *FRT* or *loxP* sites, is finally excised by tyrosine recombinase FLP or Cre to generate the unmarked, nonpolar mutation (Fig. 6A). Recently, this approach was successfully used to remove all three endogenous type III PKS genes in the engineered *S. coelicolor* strain M1152, generating an excellent expression host for the discovery and identification of actinobacterial type III polyketide metabolites (Thanapipatsiri et al.,

2015). Although PCR-targeting is one of the most frequently method for single or multiple gene disruptions in streptomycetes, two major bottlenecks, including time-consuming cosmid library construction and limited size of cosmid for carrying DNA fragments (<50 kb), have hampered its application for deleting target BGCs.

5.2. Cre-loxP recombination system

In the PCR-targeting method, the Cre-loxP recombination system is used as an assistant tool to remove resistance markers. In fact, the site-specific recombination system is also alone used to delete unnecessary BGCs. First, two *loxP* sites in the same orientation are directly

inserted into flanking genomic regions by stepwise single cross-overs. Then, the Cre recombinase is expressed to excise the unnecessary genomic DNA ([Komatsua et al., 2010](#)) ([Fig. 6B](#)). In general, this strategy is faster than the tedious PCR-targeting method and can knock out large

recombination (Yin et al., 2015), respectively; (3) considering that a large number of BGCs are not expressed under standard laboratory conditions, it is still necessary and urgent to develop powerful heterologous hosts and universal refactoring strategies to rapidly activate silent BGCs (Iqbal et al., 2016; Katz et al., 2016). Nonetheless, the highlighted strategies for engineering microbial natural products BGCs presented in this review and future developments in this area are sure to play increasingly important roles in genome mining for NP discovery and development, as well as yield improvement for large-scale manufacturing.

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