# Simple cloning of large natural product biosynthetic gene clusters from Streptomyces by an engineered CRISPR/Cas12a system

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Abstract: Directly cloning of biosynthetic gene clusters (BGCs) from microbial genomes has been revolutionizing the natural product-based drug discovery. However, it is still very challenging to efficiently clone, for example, large (> 80kb) and GC-rich (> 70%), streptomycete originating BGCs. In this study, we developed a simple, fast yet efficient and low-cost *in vitro* platform for direct cloning large BGCs from streptomycete genomic DNA, named as CAT-FISHING (<u>CRISPR/Cas12a- and Agarose plug-based sysTem for Fast bIoSyntHetIc geNe cluster cloninG</u>), by combining the advantages of CRISPR/Cas12a cleavage and bacterial artificial chromosome (BAC) library construction. CAT-FISHING was demonstrated by directly cloning large DNA fragments ranging from 47 to 139 kb with GC content of > 70% from the *S. albus* J1074 genome in a relatively efficient manner. Moreover, surugamides, encoded by a captured 87-kb BGC with GC content of 76%, was heterologously expressed in a *Streptomyces* chassis. These results indicate that CAT-FISHING is a powerful platform for BGCs batch cloning, which would be greatly beneficial to the natural products-based drug discovery. We believe that this system will lead a renaissance of interest in microorganisms as a source for drug development.

#### 1. Introduction

Microorganisms, especially actinobacteria and fungi, remain unrivalled in their ability to produce bioactive small molecules (BSMs), some of which reached the market without any chemical modifications required, a testimony to the remarkable potential of *Streptomyces* to produce novel drugs. The expedition of *Streptomyces* genomes deciphered a large unexploited pool of novel biosynthetic gene clusters (BGCs), responsible for new but silent BSMs <sup>[1]</sup>. However, the cloning of BGCs in *Streptomyces* is often very difficult because of the high GC and large size of those BGCs. It was found that 92% (1760/1910) of the characterized BGCs are smaller than 85kb, and 40% (756/1910) with > 70% GC content. In *Streptomyces*, 84% (534/634) BGCs have a GC content over 70% (Figure S1). To date, various processes have been developed for BGC cloning, such as genomic library (*i.e.*, cosmid and fosmid) construction, recombination-based or RecET/Redaβ-based cloning, and Gibson assembly *etc* (Table 1). Additionally, the emergence of the CRISPR/Cas9 technique has enabled several new DNA cloning methods such as ExoCET, and CATCH *etc* <sup>[2-4]</sup>. However, a simple, fast and efficient strategy for large BGCs, especially with high GC content, is still in urgent need to make natural BSMs accessible and affordable.

CRISPR/Cas12a is a single RNA-guided (crRNA) endonuclease of a Class II CRISPR/Cas system <sup>[16]</sup>. Unlike Cas9 proteins, Cas12a recognizes a T-rich protospacer-adjacent motif (PAM) instead of a Grich PAM and generates dsDNA breaks with staggered ends instead of blunt ends. Besides the genome editing applications <sup>[16, 17]</sup>, CRISPR/Cas12a has been widely used in nucleic acid-based diagnostic applications <sup>[18]</sup>, small molecule detection <sup>[19, 20]</sup> etc. Moreover, it is worth noting that CRISPR/Cas12a possesses obvious superiority in DNA assembly with regard to its programable endonuclease activity and the DNA sticky ends of 4- or 5-nt overhangs <sup>[16]</sup>. Based on these features, Li et al. developed a CRISPR/Cas12a-based DNA assembly standard, namely C-Brick<sup>[21]</sup>. It was further reported that the accuracy of CRISPR/Cas12a cleavage could be significantly improved by shortening the length of crRNA, based on this, a DNA assembly method CCTL (Cpf1-assisted Cutting and Taq DNA ligaseassisted Ligation) was developed for efficient replacement of promoter region (< 1kb) in a 36-kb antibiotic BGC<sup>[22]</sup>. And we foresee the potential of CRISPR/Cas12a reported by Lei et al might help to achieve the cloning of large DNA fragment with high GC content from genomic DNA sample. However, the susceptibility of large DNA fragments (> 80kb) to shearing in solution makes in vitro cloning of large BGC technically difficult, and how to achieve such a challenging objective by CRISPR/Cas12a was not clear.

As a classical method, by using agarose plug, bacterial artificial chromosome (BAC) library construction has been widely used for large DNA fragments cloning, but it is often time-consuming, labour intensive, expensive and technical demanding <sup>[5, 23]</sup>. Though some new approaches of DNA cloning have been developed, for example, PCR-based cloning, Gibson assembly and recombination-based cloning, they often require additional homologous sequences among different DNA parts for assembly, which is relatively inefficient when meeting complicated DNA sequences (e.g. large fragment, high GC and/or highly repetitive) (Table 1). On the contrary, the BAC library is also

indiscriminate toward insertion in a DNA sequence, this makes BAC library construction suitable for application to high GC content DNA samples.

In this study, by combining the DNA cleavage activity of CRISPR/Cas12a with the unique advantages of a BAC library construction, we have developed a CRISPR/Cas12a-and agarose plug-based method, designated CAT-FISHING (<u>CRISPR/Cas12a- and Agarose plug-based sysTem for Fast bloSyntHetIc geNe cluster cloninG</u>), which uses agarose plug-based *in situ* DNA isolation, digestion and liagtaion. As a proof of concept, large DNA fragments (or BGCs) of > 80kb have been fast cloned from *Streptomyces* genomic DNA (73% GC) by CAT-FISHING. Furthermore, the captured 87-kb surugamides BGC has also been successfully expressed in the *S. albus* J1074-derived cluster-free chassis strain <sup>[24]</sup>.

#### 2. Results

#### 2.1. Design and workflow of CAT-FISHING for genome mining-based drug discovery

The flow chart of CAT-FISHING for capturing genome mining-based BGCs for drug discovery is presented in Figure 1. Mining microbial resources (e.g., Streptomyces) with advanced genome sequencing and bioinformatics provides new opportunities for natural product-based drug discovery. By using CAT-FISHING, target BGCs can be captured only with three steps. The first step is the capture plasmid construction and CRISPR/Cas12a- based plasmid digestion. In this step, two homolog arms (each arm containing at least one PAM site) that flank the target BGC were selected as adapter sequence and amplified by PCR. Then the BAC plasmid backbone containing the two adapter sequences and selection marker (e.g., antibiotic resistance gene, counter selection gene or lacZ), and the designated capture plasmid was constructed via the DNA assembly method. Under the guidance of crRNAs, the two selected PAM motif regions on the left and right adapter sequence were simultaneously digested by CRISPR/Cas12a, resulting in the linear capture plasmid. The second step is agarose plug-based genomic DNA isolation and in situ CRISPR/Cas12a digestion. According to the BAC library construction protocol<sup>[7]</sup>, genomic DNA plugs from the target strain were prepared. And the genomic DNA was digested in situ by the CRISPR/Cas12a system guided by the two designed crRNAs that were previously used in step one. The last step is ligation, desalting and transformation. The resulting linear capture plasmid and the digested genomic DNA from steps I and II, respectively, were gently mixed and ligated by T4 DNA ligase. Then desalted ligation products were introduced into E. coli by electroporation. The successfully captured BGCs can then be identified by a PCR screening approach. Lastly, the encoded compound of the captured target cryptic BGC can be obtained by heterologous expressing it in a suitable host with further chemical identification.

#### 2.2. Evaluation of CRISPR/Cas12a-based DNA cloning efficiency

As CCTL, the principle underlying CAT-FISHING is the cohesive end ligation of two linearized DNA fragments by T4 DNA ligase. However, different from the widely used restriction endonuclease-based DNA cloning methods, here the cohesive ends were generated by paired crRNA-guided CRISPR/Cas12a cleavage. Though our previous study demonstrated that CRISPR/Cas12a cleaves

more specifically on target DNA with a shorter spacer (*e.g.*, 18nt) <sup>[22]</sup>, the Cas12a cleavage is still not as accurate as NEB restriction endonucleases do. It is well known that the successful rate of direct cloning of the targeted large DNA fragment is often determined by the cloning efficiency. This study therefore carefully evaluated and compared the cloning efficiencies achieved by applying two different kinds of cohesive ends that were individually generated by NEB restriction endonuclease and CRSIPR/Cas12a.

As shown in Figure 2A, plasmid pGY2020 derived from the pCC2-FOS Fosmid vector (Epicentre) was constructed, and this plasmid contains two PAM sites (PAM1 and PAM2) as well as two NEB restriction endonucleases (EcoRI and HindIII). And the specific DNA fragment (ampicillin resistant gene  $Amp^R$ , < 1kb) was cloned into pGY2020 by the CRISPR/Cas12a or NEB restriction enzymesbased method. There was no significant difference (P > 0.05) in the true positive rate between these two cloning methods (Figure 2B; Figure S2). In addition, three randomly selected clones containing pGY2020/P1P2 also confirmed by junction sequencing and found no mutations (Figure 2C-D). These results indicate that CRISPR/Cas12a-based cloning strategy is able to clone short DNA fragments with a relatively high efficiency. Next, we would like to assess the ability of CRISPR/Cas12a-based method in direct cloning of large DNA fragments.

# 2.3. Cloning of a target DNA fragment from a BAC plasmid

In order to further demonstrate CRISPR/Cas12a-based cloning strategy in a simplified system, an 137kb BAC plasmid pBAC-ZL (68% GC) was used to mimic and evaluate its cloning performance on a large DNA fragment. As shown in Figure 3A, a 50-kb fragment and an 80-kb fragment could be obtained by using the corresponding crRNAs-guided CRISPR/Cas12a cleavage. Under the guidance of the corresponding crRNA pairs, the BAC plasmid pBAC-ZL was digested by CRISPR/Cas12a, and 50-kb and 80-kb target bands were observed on the agarose gel after PFGE (pulsed field gel electrophoresis) (Figure 3B). By using the corresponding capture plasmid, two target DNA fragments were also successfully cloned from the BAC plasmid pBAC-ZL, as shown in Figure 3C-3E and Figure S3-S4. For 50-kb DNA fragment, of more than 100 transformants about 95% were the right clones. For the 80-kb DNA fragment, the number of transformants and the true positive rate were both lower, and about 50% of the transformants were the right clones. These results indicated that in a simplified system, by applying CRSIPR/Cas12a and corresponding capture plasmids, it could achieve high cloning rates of the 50-kb and 80-kb DNA fragments from the purified BAC plasmid DNA sample. It also needs to be noted that, for the 80-kb DNA fragment, the cloning difficulty was obviously greater.

# 2.4. Direct cloning of target BGCs from Streptomyces genomic DNA by CAT-FISHING

In order to setup CAT-FISHING for directly cloning large BGCs from *Streptomyces* genomic DNA, CRISPR/Cas12a has been introduced into the initial process of BAC library construction by replacing the type II restriction enzymes (e.g. *Hind*III, *Eco*RI, *Bam*HI), which are involved in partial digestion of the genomic DNA (Figure 5). The optimized workflow of the target BGC capturing by CAT-FISHING is shown in Figure 4A. Based on our extensive investigation and test, we found that after

genomic DNA isolation by agarose plugs and subsequent CRISPR/Cas12a *in situ* digestion, the resulting sample containing a mixture of genomic DNA could be directly used for subsequent ligation and transformation without prior DNA fragment isolation by PFGE and purification. This would allow CAT-FISHING to easily clone the large target DNA fragment.

In this study, a 49-kb paulomycin gene cluster <sup>[25]</sup>, an 87-kb surugamides gene cluster <sup>[26]</sup> as well as a 139-kb candicidin gene cluster <sup>[27]</sup> were selected from the chromosome of *S. albus* J1074 to demonstrate our method. As shown in Table 2, the 49-kb paulomycin gene cluster (GC content 71%) and the 87-kb surugamides gene cluster (GC content 76%) were successfully cloned by CAT-FISHING, as confirmed by PCR and restriction mapping (Figure 4B-4C). And it was found the ratio of right clones that contain a 49-kb target BGC was  $4\sim5\%$ , and that for an 87-kb target BGC was  $2\sim4\%$ . Additionally, the 139-kb candicidin gene cluster (GC content 75%) was also cloned with CAT-FISHING, albeit with a much lower efficiency.

# 2.5. Expression of the target BGC in a cluster-free Streptomyces chassis strain

To thoroughly check the sequence and functional integrity of these BGCs that were captured by CAT-FISHING, as well as to prove access to genome mining through this route, a captured 87-kb surugamides gene cluster was expressed in a cluster-free *Streptomyces* chassis strain. The *aac(3)IVoriT-attP(\PhiC31)-int(\PhiC31)* cassette was introduced into the target plasmid by Red/ET recombination. By applying ET12567/pUC307-mediated triparental conjugation, the resulting plasmid was integrated into the chromosome of *S. albus* Del14 (Figure 6A). During the subsequent fermentation study, surugamide A was produced in *S. albus* Del14-87kb by compared with *S. albus* Del14 and *S. albus* J1074 (Figure 6B). LC–MS/MS analysis further confirmed the production of surugamide A ([M+H]<sup>+</sup> = 912.6252 Da, RT = 6.97 min) (Figure 6C).

# 2.6. The advantages of CAT-FISHING over other selected large DNA fragment cloning methods

The comparison between CAT-FISHING and BAC library construction as well as CAT-FISHING and other differents protocols of large DNA fragment cloning were presented in Figure 5, Table 3 and Table S1. From the comparison, it was concluded that CAT-FISHING is very simple, fast and efficient in cloning of large DNA fragment with high GC content. And the target large BGC could be obtained from *Streptomyces* samples in several days. In additon, for high GC *Streptomyces* genomic DNA sample, the upper cloning limit of CAT-FISHING reached to as large as139 kb. Most importantly, by current cost accounting of each cloning steps, it was found that compared to BAC library construction, CAT-FISHING dramatically decreased the cost of *Streptomyces* originating target BGC cloning (Table 3; Table S2-S3). It makes that the CAT-FISHING is a totally affordable method for efficient and batch cloning large BGCs form *Streptomyces* or other microbial genomes.

# 3. Discussion

Cloning and manipulation of large DNA fragments is a fundamental and important platform technology for mining results BSMs from rich microbial genome sources. In order to avoid shearing

of large DNA fragment during *in vitro* manipulation, in this study, CAT-FISHING combined the advantages of CRISPR/Cas12a cleavage and BAC library construction (agarose plug-based *in situ* DNA isolation and digestion). Instead of using restriction enzymes (*e.g.* HindIII, EcoRI, or BamHI) for the random fragmentation of the genomic DNA <sup>[7]</sup>, CAT-FISHING utilizes CRISPR/Cas12a together with specific crRNA pairs to precisely *in situ* digestion of the genomic DNA in agarose gel (Figure 1; Figure 4A). This theoretically makes CAT-FISHING very suitable for cloning of large BGCs with high GC contents.

There are two concerns about CRISPR/Cas12a-based DNA cleavage, off-target (non-specific cleavage) and inaccuracy of the cleavage sites <sup>[21, 28]</sup>. Both could reduce the cloning efficiency of CAT-FISHING. Lei *et al.* reported that the cleavage specificity of CRISPR/Cas12a would enhance with a shorter crRNA spacer (*i.e.*,  $17 \sim 19$  nt) <sup>[22]</sup>. In this study, we used crRNA with an 18-nt spacer to evaluate the cloning efficiency of CRISPR/Cas12a-based cloning strategy. As shown in Figure 2, the number of transformants obtained by CRISPR/Cas12a were fewer than the control, while no significant difference (P > 0.05) was observed. On the other hand, non-specific cleavage by CRISPR/Cas12a could be minimized by decreasing the Cas12a concentration, shortening the cleavage time and slection of more suitable crRNA. As a result, when a purified 137-kb BAC plasmid was used to test CRISPR/Cas12a-based cloning strategy, the plasmid was almost completely digested and no non-specific cleavage products appeared on the agarose gel (Figure 3A-3B). To some extent, these results verify the relatively higher efficiency of cloning 50-kb and 80-kb DNA fragments from BAC plasmid (Figure 3C-3E).

PFGE is a powerful and essential tool for isolating large DNA fragments. During BAC library construction, however, PFGE is time consuming, as it often takes  $16 \sim 24$  hours to separate specific size DNA fragments <sup>[7]</sup>. Moreover, the following operational steps, such as DNA elution and purification (Figure 5), could drastically decrease the DNA integrity/amount as well as the subsequent ligation or transformation efficiency. In this study, after many preliminary tests, we found that, following agarose digestion, the resulting mixture from a CRISPR/Cas12a-treated genomic DNA plug could be directly used for ligation with the vector and subsequent electro-transformation (Figure 4A; Figure 5). Without the need for PFGE or the preparation of purified high molecular weight genomic DNA fragments, and compared to previously reported large DNA fragment cloning approaches (*e.g.* ExoCET, CATCH, TAR *etc.*), the cloning process in CAT-FISHING has been greatly simplified <sup>[2, 4, 8, 15]</sup>.

In this study, by applying CAT-FISHING, we achieved a cloning efficiency of  $2\sim4\%$  when cloning an 87-kb target BGC (Figure 4B-4C; Table 2). It is close to Wang's result that 106 kb salinomycin BGC from *S. albus* obtained by ExoCET at screen-positive rate of 4% <sup>[4]</sup>. To further determine the upper cloning limit of CAT-FISHING, an 139-kb candicidin BGC was selected for demonstration. Although this study successfully cloned candicidin BGC from *S. albus* genome, the screen-positive rate of target clone was less than 1%. Probably due to the complexity of the un-purified DNA mixture sample,

slightly inaccurate of Cas12a cleavage site<sup>[22]</sup> and high activity of T4 DNA ligase, many short DNA fragments or incomplete pieces of target BGCs were inserted into capture plasmids. Therefore, it is reasonable to predict that, if necessary, through DNA fragment isolation and purification, the cloning performance toward a 139-kb BGC should also be dramatically improved. Anyway, compared to other methods of large DNA fragment cloning, including the BAC library, in which often only a few right strains could be screened out of thousands of clones (i.e.,  $1/1000 \sim 1/2000$ ) <sup>[5, 29]</sup>, CAT-FISHING is a simpler method with a greater efficiency for cloning target BGCs with high GC content.

*Streptomyces* are the source of a majority of antibiotic classes in current clinical and agricultural use <sup>[30]</sup>. *S. albus* J 1074 is one of the most widely used *Streptomyces* chassis for genome mining <sup>[31]</sup>. In this study, *S. albus* Del14, which is a *S. albus* J 1074-derived cluster-free chassis strain <sup>[24]</sup>, has been used to demonstrate the sequence and functional integrity of the BGCs obtained by CAT-FISHING. As shown in Figure 5, the 87-kb target BGC was successfully expressed, and the corresponding NP surugamides (inhibitors of cathepsin B) could be confirmed by LC-MS. During genome mining, BGC cloning and expression is the most important starting point for the next step of bioactivity analysis and structure identification of target BSMs. The current results present a case study for NP production in *Streptomyces* by applying CAT-FISHING. Lastly, as an *in vitro* manipulation platform, not limited to actinomyces, CAT-FISHING could easily be extended to fungi and other microbial resources <sup>[32]</sup>.

As a results, we successfully combining the accurate DNA cleavage activity of CRISPR/Cas12a with the unique advantages of BAC library construction platform that is believed one of the most efficient strategies of cloning large DNA fragment with high G+C content. This combining gave the birth of CAT-FISHING, which possesses some unique features comparing to previous methods: 1) CAT-FISHING is a very simple and low-cost cloning method that can directly capture the target large DNA fragment (>80kb) with high GC content (70%) from genomic DNA; 2) CAT-FISHING is very fast, the target large BGC could be obtained from streptomycete genomic DNA samples in a few days (Table 3); 3) CAT-FISHING does not require PFGE or the preparation of purified large DNA fragments. Moreover, in addition to genome editing, DNA assembly, nucleic acid and small molecule detection *etc.*, this study also expanded the application of CRISPR/Cas12a to direct cloning of large DNA fragments with complicated DNA sequence (e.g. high GC or sequences with repeat) *in vitro*. This innovation of a fundamental platform technology for use in genome mining through application of the CRISPR/Cas12a system would facilitate the discovery of novel BSMs from microbial sources.

#### 4. Experimental Section

*Strains, plasmids and media:* The strains and plasmids used in this work are present in Table S4. *Escherichia coli* and its derivatives were cultivated on Luria-Bertani (LB) agar plates (tryptone 10 g/L, yeast extract 5 g/L and NaCl 10 g/L, pH = 7.2). Streptomyces and its derivatives were cultivated on soybean flour-mannitol (SFM) agar plates (soybean flour 20 g/L, mannitol 20 g/L and agar 20 g/L, pH = 7.2) or ISP4 (International Streptomyces Project Medium 4, BD Bioscience, San Jose, CA, USA). In the fermentation experiments, seeds were grown in TSB (trypticase soy broth, Oxoid Ltd), and R4 medium was used for subsequent fermentations (in a 250-mL Erlenmeyer flask, 30°C and 200 rpm).

*Capture plasmid construction:* The primers for capture plasmid construction are listed in Table S5. The capture plasmid was constructed by introducing the *lacZ* gene as well as two PCR-amplified homology arms (each arm containing at least one PAM site) corresponding to the flanking regions of the target DNA fragment (or BGC) in to pBAC2015 <sup>[10]</sup>. Assembly of multiple DNA fragments was carried out by Gibson assembly or with the EZmax one-step seamless cloning kit (Tolo Biotechnology, Shanghai). Plasmid DNA was isolated from *E. coli* by using alkaline lysis <sup>[33]</sup>.

*Genomic DNA isolation:* For genomic DNA isolation, *S. albus* J1074 was cultured in Oxoid TSB (30 g/L) supplemented with glycine (5 g/L). According to *Practical Streptomyces Genetics* <sup>[33]</sup>, after cultivation at 200 rpm and 30°C for 24 ~ 36 h, mycelium was collected by centrifugation (4°C, 4000 g, 5 min). Mycelium was resuspended in TE25S (25 mM Tris-HCl pH 8, 25 mM EDTA pH 8, 0.3 M sucrose) and then the supernatant was removed (4°C, 4000 g, 5 min). The mycelium density was adjusted with TE25S and it was mixed with an equal volume of 1.0% LMP agarose (1.0% molten solution of low melting point agarose) at 50°C, and then poured into holes in a plug mould (100-µl holes). The blocks were removed from the mould and incubated at 37°C for 1 h in lysozyme solution (2 mg/mL in TE25S). The lysozyme solution was removed and the blocks were incubated at 50°C for 2 h in proteinase K solution (1 mg/mL proteinase K in NDS. NDS: 0.5 M EDTA pH 8, 10 mM Tris-HCl pH 8, 1% SDS). The proteinase K solution was removed and the blocks were washed once for 30 min in TE (10 mM Tris-HCl pH 8, 1 mM EDTA pH 8) supplemented with 0.1 mM PMSF (phenylmethanesulfonyl fluoride, serine proteinase inhibitor), then three times with TE for 30 min. After removing all of the liquid, the agarose plugs could be used for CRISPR/Cas12a digestion, but could also be stored for up to 1 month at 4°C in 70% ethanol.

*crRNA preparation:* The oligonucleotides used as templates for crRNA transcription are given in Table S6. According to our previous study <sup>[19]</sup>, crRNA was prepared via *in vitro* transcription. Templates for crRNA synthesis were synthesized and annealed by using Taq DNA Polymerase PCR Buffer (Thermo Fisher Scientific). A HiScribe<sup>™</sup> T7 Quick High Yield RNA Synthesis Kit (NEB) was used for crRNA *in vitro* transcription. The resulting crRNA was purified using RNA Clean & ConcentratorTM-5 kit (Zymo Research), and subsequently quantified using a NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific). RNase-free materials (Axygen Scientific, Union City, CA, USA) and conditions were applied during the entire experimental process.

*CRISPR/Cas12a-based DNA restriction and ligation:* The Cas12a (LbCas12a) protein used in this study was overexpressed in pET28a and then purified by fast protein liquid chromatography (FPLC; AKTA Explorer 100, GE Healthcare)<sup>[19]</sup>. In the CRISPR/Cas12a cutting system, NEBuffer<sup>TM</sup> 3.1 (100 mM NaCl, 50 mM Tris-HCl, 10 mM MgCl<sub>2</sub>, 100  $\mu$ g/mL BSA, pH 7.9) was adopted as the reaction buffer. For pBAC-ZL or capture plasmid cleavage, plasmid DNA was incubated with Cas12a protein and the corresponding crRNA pairs at 37°C for 1 h. After the reaction, the linearized capture plasmids or DNA fragments of pBAC-ZL were prepared using isopropanol and ethanol <sup>[10]</sup>. The resulting linear

capture plasmid or DNA fragments of pBAC-ZL could be used for the following ligation. If necessary, the large DNA fragments could be analysed by pulsed field gel electrophoresis (PFGE) with the CHEF-DR III apparatus (Bio-Rad, Richmond, CA). PFGE was performed in 0.5% agarose at 6 V/cm with a  $1 \sim 25$  sec switching pulse time for  $16 \sim 18$  h in  $0.5 \times$  TBE buffer. For genomic DNA cleavage, plugs were initially equilibrated in  $1 \times$  NEBuffer<sup>TM</sup> 3.1, then transferred into a cleavage system that contained the Cas12a protein and the corresponding crRNA pair, and finally incubated at 37°C for  $1 \sim 2$  h. After the reaction, following heat treatment at 65°C for 10 min to inactive Cas12a protein, the LMP agarose gel was hydrolysed using  $\beta$ -Agarase I (NEB) for 30 min at 42°C. Afterward, the resulting DNA mixture could be directly used for following the ligation with the corresponding linear capture vectors by T4 DNA ligase (NEB).

*Electro-transformation of E. coli:* Following the transfer of ligation samples into 0.1 M glucose/1% agarose gel to desalt for  $1 \sim 2$  h on ice, these samples could be used for electro-transformation. The high efficiency electro-transformation of *E. coli* cells was accomplished according to a previous study <sup>[34]</sup>. The following electro-transformations were performed in 2-mm cuvettes using the Bio-Rad GenePulser Xcell<sup>TM</sup> system (electroporator conditions: 2500 V, 200  $\Omega$  and 25  $\mu$ F). Then, to the *E. coli* cells in the cuvette, 1 mL of SOC medium (tryptone 20 g/L, yeast extract 5 g/L, NaCl 0.5 g/L, KCl 2.5 mM, MgCl<sub>2</sub> 10 mM, glucose 20 mM) was added and the mixture was transferred into a 15-mL Falcom<sup>TM</sup> tube. After shaking at 200 rpm for 1 h at 37°C, the strains were collected and spread on selective LB agar plates. The plates were incubated overnight at 37°C, and the transformants were screened and verified by PCR using the primers listed in Table S7.

Expression of BGCs in Streptomyces and LC-MS analysis: The aac(3)IV-oriT-attP( $\Phi C31$ )-int( $\Phi C31$ ) cassette amplified from pSET152 has been introduced into BAC plasmids by Red/ET recombination <sup>[35]</sup>. The resulting plasmid was introduced into *S. albus* Del14 by triparental conjugation according to a previous study <sup>[5]</sup>. The transconjugants were screened and verified by PCR using the primers listed in Table S7. After fermentation, the production of target natural product was qualitatively analysed using a high-resolution Q-Exactive Hybrid Quadrupole-Orbitrap mass spectrometer (Thermo Scientific, Waltham, MA).

# **Supporting Information**

Supporting Information is available from the Wiley Online Library or from the author.

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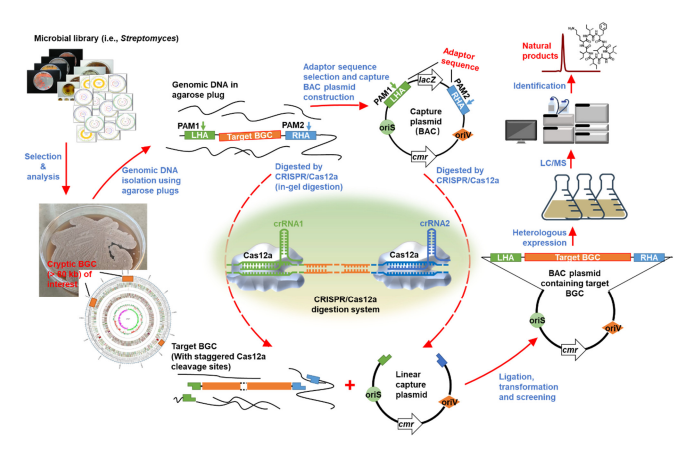
#### **Conflict of Interest**

The authors declare no conflict of interest.

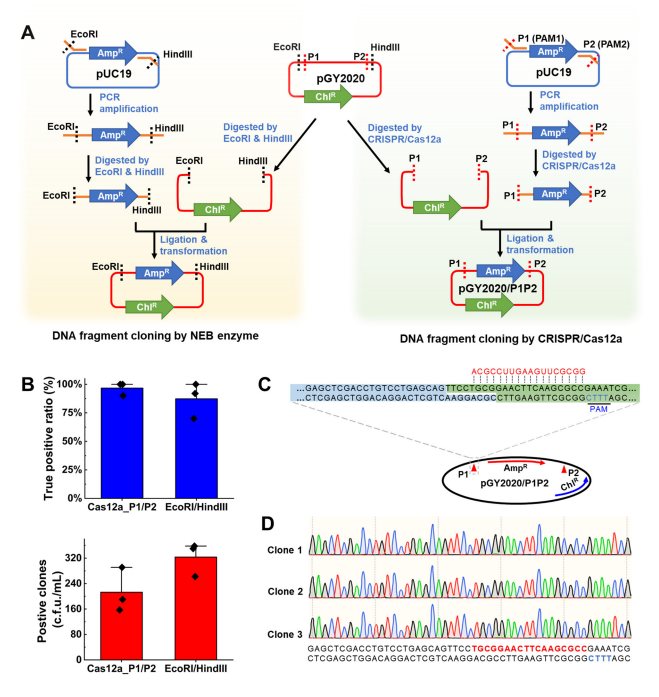
#### References

- a) V. M. D'Costa, C. E. King, L. Kalan, M. Morar, W. W. L. Sung, C. Schwarz, D. Froese, G. Zazula, F. Calmels, R. Debruyne, G. B. Golding, H. N. Poinar, G. D. Wright, *Nature* 2011, 477, 457; b) E. J. Culp, G. Yim, N. Waglechner, W. Wang, A. C. Pawlowski, G. D. Wright, *Nat. Biotechnol.* 2019, 37, 1149.
- [2] W. Jiang, T. F. Zhu, Nat. Protoc. 2016, 11, 960.
- [3] W. Tao, L. Chen, C. Zhao, J. Wu, D. Yan, Z. Deng, Y. Sun, ACS Synth. Biol. 2019, 8, 1991.
- [4] H. Wang, Z. Li, R. Jia, J. Yin, A. Li, L. Xia, Y. Yin, R. Müller, J. Fu, A. F. Stewart, Y. Zhang, Nucleic Acids Res. 2017, 46, e28.
- [5] G. Y. Tan, K. Deng, X. Liu, H. Tao, Y. Chang, J. Chen, K. Chen, Z. Sheng, Z. Deng, T. Liu, ACS Synth. Biol. 2017, 6, 995.
- [6] a) M. Xu, Y. Wang, Z. Zhao, G. Gao, S. X. Huang, Q. Kang, X. He, S. Lin, X. Pang, Z. Deng, Appl. Environ. Microbiol. 2016, 82, 5795; b) Y. Yu, L. Bai, K. Minagawa, X. Jian, L. Li, J. Li, S. Chen, E. Cao, T. Mahmud, H. G. Floss, X. Zhou, Z. Deng, Appl. Environ. Microbiol. 2005, 71, 5066.
- [7] M. Luo, R. A. Wing, in *Plant Functional Genomics*, (Ed: E. Grotewold), Humana Press, Totowa, NJ 2003, 3.
- [8] Y. Kazuya, K. A. Reynolds, R. D. Kersten, K. S. Ryan, D. J. Gonzalez, N. Victor, P. C. Dorrestein, B. S. Moore, Proc. Natl. Acad. Sci. U.S.A. 2014, 111, 1957.
- [9] V. N. Noskov, B. J. Karas, L. Young, R.-Y. Chuang, D. G. Gibson, Y.-C. Lin, J. Stam, I. T. Yonemoto, Y. Suzuki, C. Andrews-Pfannkoch, J. I. Glass, H. O. Smith, C. A. Hutchison, J. C. Venter, P. D. Weyman, ACS Synth. Biol. 2012, 1, 267.
- [10] H. Wang, Z. Li, R. Jia, Y. Hou, J. Yin, X. Bian, A. Li, R. Müller, A. F. Stewart, J. Fu, Y. Zhang, Nat. Protoc. 2016, 11, 1175.
- [11] X. Bian, F. Huang, H. Wang, T. Klefisch, R. Müller, Y. Zhang, ChemBioChem 2014, 15, 2221.
- [12] L. Li, Y. Zhao, L. Ruan, S. Yang, M. Ge, W. Jiang, Y. J. Lu, Metab. Eng. 2015, 29, 12.
- [13] C. Song, J. Luan, Q. Cui, Q. Duan, Z. Li, Y. Gao, R. Li, A. Li, Y. Shen, Y. Li, A. F. Stewart, Y. Zhang, J. Fu, H. Wang, ACS Synth. Biol. 2019, 8, 137.
- [14] a) L. Li, W. Jiang, Y. Lu, in *Synthetic Metabolic Pathways: Methods and Protocols*, (Eds: M. K. Jensen, J. D. Keasling), Springer New York, New York, NY **2018**, 203; b) A. Casini, J. T. MacDonald, J. D. Jonghe, G. Christodoulou, P. S. Freemont, G. S. Baldwin, T. Ellis, *Nucleic Acids Res.* **2013**, 42, e7.
- [15] W. Jiang, X. Zhao, T. Gabrieli, C. Lou, Y. Ebenstein, T. F. Zhu, Nat. Commu. 2015, 6, 8101.
- [16] Z. Bernd, J. S. Gootenberg, O. O. Abudayyeh, I. M. Slaymaker, K. S. Makarova, E. Patrick, S. E. Volz, J. Julia, V. D. O. John, R. J. C. Aviv, *Cell* 2015, 163, 759.
- [17] a) Y. Jiang, F. Qian, J. Yang, Y. Liu, F. Dong, C. Xu, B. Sun, B. Chen, X. Xu, Y. Li, R. Wang, S. Yang, *Nat. Commu.* 2017, 8, 15179; b) H. Manghwar, B. Li, X. Ding, A. Hussain, K. Lindsey, X. Zhang, S. Jin, *Adv. Sci.* 2020, 7, 1902312.

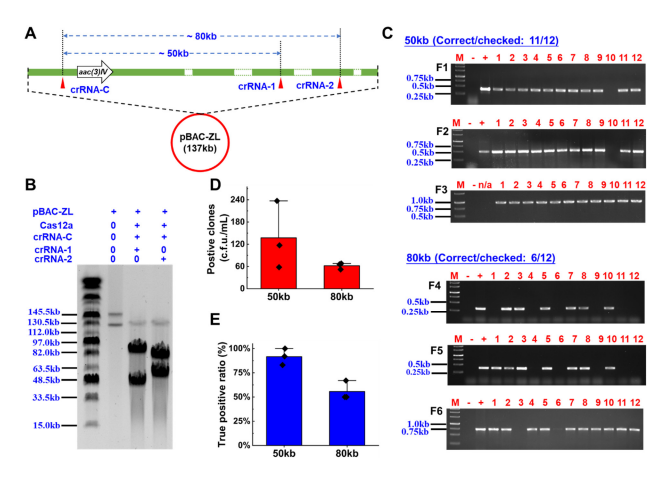
- [18] a) J. S. Chen, E. Ma, L. B. Harrington, M. Da Costa, X. Tian, J. M. Palefsky, J. A. Doudna, *Science* 2018, 360, 436; b) S.-Y. Li, Q.-X. Cheng, J.-M. Wang, X.-Y. Li, Z.-L. Zhang, S. Gao, R.-B. Cao, G.-P. Zhao, J. Wang, *Cell Discov.* 2018, 4, 20; c) S.-Y. Li, Q.-X. Cheng, J.-K. Liu, X.-Q. Nie, G.-P. Zhao, J. Wang, *Cell Res.* 2018, 28, 491.
- [19] a) M. Liang, Z. Li, W. Wang, J. Liu, L. Liu, G. Zhu, L. Karthik, M. Wang, K.-F. Wang, Z. Wang, J. Yu, Y. Shuai, J. Yu, L. Zhang, Z. Yang, C. Li, Q. Zhang, T. Shi, L. Zhou, F. Xie, H. Dai, X. Liu, J. Zhang, G. Liu, Y. Zhuo, B. Zhang, C. Liu, S. Li, X. Xia, Y. Tong, Y. Liu, G. Alterovitz, G.-Y. Tan, L.-X. Zhang, *Nat. Commu.* 2019, 10, 3672; b) X. Zhao, S. Li, G. Liu, Z. Wang, Z. Yang, Q. Zhang, M. Liang, J. Liu, Z. Li, Y. Tong, G. Zhu, X. Wang, L. Jiang, W. Wang, G-Y. Tan, L. Zhang, *Sci. Bull.* 2020, https://doi.org/10.1016/j.scib.2020.09.004.
- [20] Y. Xiong, J. Zhang, Z. Yang, Q. Mou, Y. Ma, Y. Xiong, Y. Lu, J. Am. Chem. Soc. 2020, 142, 207.
- [21] S.-Y. Li, G.-P. Zhao, J. Wang, ACS Synth. Biol. 2016, 5, 1383.
- [22] C. Lei, S.-Y. Li, J.-K. Liu, X. Zheng, G.-P. Zhao, J. Wang, Nucleic Acids Res. 2017, 45, e74.
- [23] K. Farrar, I. S. Donnison, Nat. Protoc. 2007, 2, 1661.
- [24] M. Myronovskyi, B. Rosenkränzer, S. Nadmid, P. Pujic, P. Normand, A. Luzhetskyy, *Metab. Eng.* 2018, 49, 316.
- [25] A. González, M. Rodríguez, A. F. Braña, C. Méndez, J. A. Salas, C. Olano, *Microb. Cell Fact.* 2016, 15, 56.
- [26] F. Xu, B. Nazari, K. Moon, L. B. Bushin, M. R. Seyedsayamdost, J. Am. Chem. Soc. 2017, 139, 9203.
- [27] S. Chen, X. Huang, X. Zhou, L. Bai, J. He, K. J. Jeong, S. Y. Lee, Z. Deng, Chem. Biol. 2003, 10, 1065.
- [28] Y. Fu, J. A. Foden, C. Khayter, M. L. Maeder, D. Reyon, J. K. Joung, J. D. Sander, *Nat. Biotechnol.* 2013, 31, 822.
- [29] S.-S. Woo, J. Jiang, B. S. Gill, A. H. Paterson, R. A. Wing, Nucleic Acids Res. 1994, 22, 4922.
- [30] a) G. Y. Tan, T. Liu, *Metab. Eng.* 2017, 39, 228; b) Q. Gao, G.-Y. Tan, X. Xia, L. Zhang, *Curr. Opin. Biotechnol.* 2017, 48, 251; c) G.-Y. Tan, Y. Peng, C. Lu, L. Bai, J.-J. Zhong, *Metab. Eng.* 2015, 28, 74; d) W. Wang, S. Li, Z. Li, J. Zhang, K. Fan, G. Tan, G. Ai, S. M. Lam, G. Shui, Z. Yang, H. Lu, P. Jin, Y. Li, X. Chen, X. Xia, X. Liu, H. K. Dannelly, C. Yang, Y. Yang, S. Zhang, G. Alterovitz, W. Xiang, L. Zhang, *Nat. Biotechnol.* 2020, 38, 76.
- [31] N. Zaburannyi, M. Rabyk, B. Ostash, V. Fedorenko, A. Luzhetskyy, BMC Genomics 2014, 15, 97.
- [32] N. Ziemert, M. Alanjary, T. Weber, Nat. Prod. Rep. 2016, 33, 988.
- [33] T. Kieser, M. J. Bibb, M. J. Buttner, K. F. Chater, D. A. Hopwood, *Practical streptomyces genetics*, Vol. 291, John Innes Foundation Norwich, 2000.
- [34] W. J. Dower, J. F. Miller, C. W. Ragsdale, Nucleic Acids Res. 1988, 16, 6127.
- [35] Y. Zhang, J. P. P. Muyrers, G. Testa, A. F. Stewart, Nat. Biotechnol. 2000, 18, 1314.



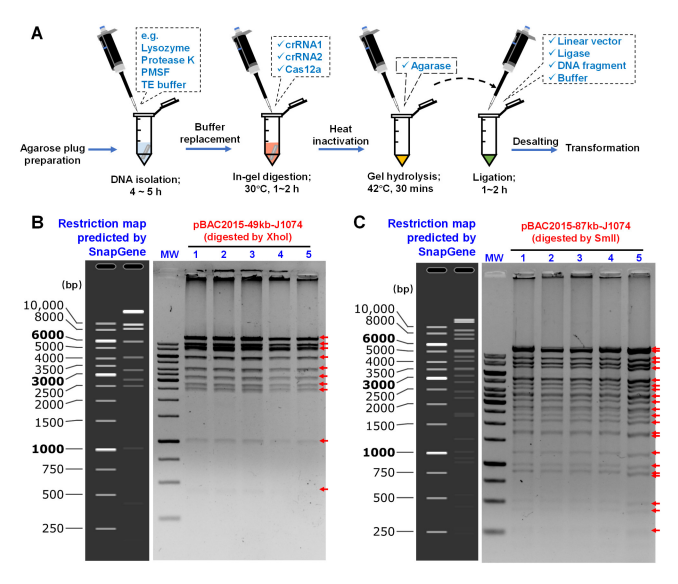
**Figure 1.** Schematic diagram of CAT-FISHING application. LHA: left homology arm, RHA: right homology arm, BGC: biosynthetic gene cluster; PAM: protospacer adjacent motif.



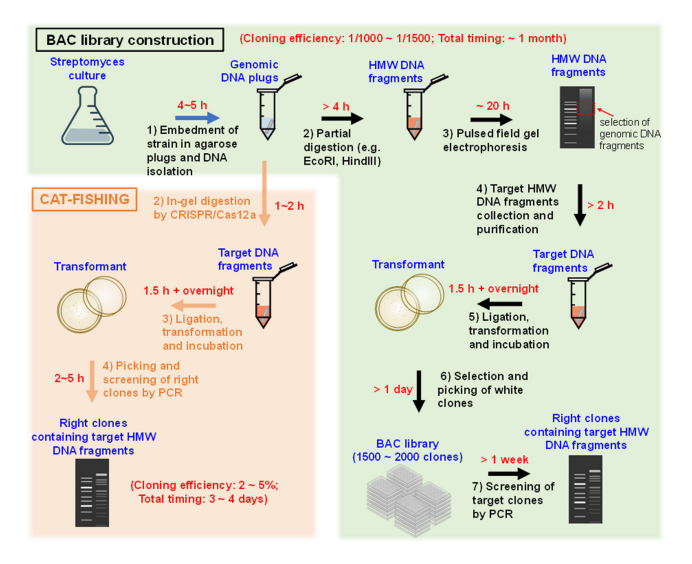
**Figure 2.** Evaluation and comparison of CRISPR/Cas12a-based and NEB restriction enzymes-based DNA cloning efficiency. A) Workflow of DNA fragment cloning by the CRISPR/Cas12a or NEB restriction enzymes-based method. B) Comparison of the clone numbers and positive rates of the CRISPR/Cas12a-based and NEB restriction enzymes-based methods. The results form three repeats are shown. C) Schematic representation of crRNA design and cohesive end ligation. 18-nt spacer crRNAs were employed, Cas12a mainly cleavage after the 14th base, generating 8-nt cohesive ends <sup>[22]</sup>. D) Conformation of Cas12a-mediated cohesive end ligation by junction sequencing. Three clones were randomly selected for sequencing.



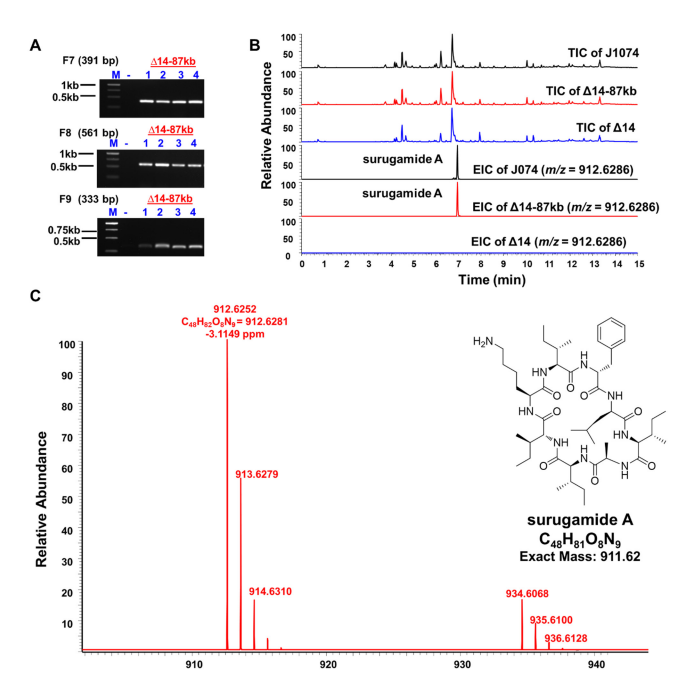
**Figure 3.** Direct cloning of 50kb and 80 kb of DNA fragments from pBAC-ZL by CRISPR/Cas12a-based cloning strategy. A) Two different target segments with different lengths (50 kb and 80 kb) in the BAC plasmid. B) Analysis of CRISPR/Cas12a-digested BAC plasmid (pBAC-ZL) by PFGE. PFGE was performed in 0.5% agarose at 6 V/cm with a  $1 \sim 25$  sec switching pulse time for  $16 \sim 18$  h in  $0.5 \times$  TBE buffer. C) PCR screening of right clones containing target DNA fragments (50kb or 80kb). Twelve clones were randamly selected from each agar plate. Three different primer pairs for regions in the middle and in the left/right boundary of target DNA fragments were used in the assay. F1, F2 and F3 are the PCR products that were amplified using 50-BAC-scr-up-F/R, 50-BAC-scr-middle-F/R and BAC-scr-down-F/R, respectively. F4, F5 and F6 are the PCR products that were amplified using 80-BAC-scr-up-F/R, 80-BAC-scr-middle-F/R and 80-scr-down-F/R, respectively. "-" represented blank control, genomic DNA of *E. coli* DH10B was used as PCR template. "+" represented positive control, pBAC-ZL plasmid DNA was used as PCR template. D-E) Determination of the clone numbers and positive rates for the DNA fragments of different lengths in three independent experiments/plates.



**Figure 4.** Direct cloning of large BGCs from genomic DNA by CAT-FISHING. A) Workflow of the target BGC capturing from genomic DNA by CAT-FISHING. B-C) Validation of five randomly selected positive clones containing a paulomycin or surugamides gene cluster by restriction enzymes digestion. XhoI and SmII have, respectively, been used for paulomycin and surugamides gene cluster restriction. Bands are indicated by arrows. pBAC2015-49kb-J1074 digested by XhoI: 19776bp, 9843bp, 8402bp, 5856bp, 4108bp, 3270bp, 2749(2499)bp, 1011bp, 435bp; pBAC2015-87kb-J1074 digested by XhoI:13832(12760)bp, 9053bp, 7701bp, 6978bp, 5176(5040)bp, 4530bp, 4091bp, 3597(3497)bp, 3135bp, 2816bp, 2412bp, 2181bp, 1731(1681) bp, 1206bp, 1016bp, 930bp, 524bp, 433bp, 277(262)bp.



**Figure 5.** Methods of cloning large DNA fragments from genomic DNA by construction of BAC library (green box) or CAT-FISHING (orange box). HMW, High Molecular Weight.



**Figure 6.** Expression of pBAC2015-87kb-J1074-int in *S. albus* Del14. A) PCR verfication of S. albus Del14-87kb. Four transformants were randomly slected. F7, F8 and F9 are the PCR products that were amplified using 87-scr-up-F/R, 87-scr-middle-F/R and 87-scr-down-F/R,respectively. "-" represented blank control, genomic DNA of *S. albus* Del14 was used as PCR template. B) Detection of surugamide components by LC-MS in S. albus Del-87kb. C) High-resolution mass spectrum of surugamide A ([M+H]+ = 912.6252 Da, RT = 6.97 min; [M+H]+ = 934.6068 Da, RT = 6.97 min).

~ 40kb								
~ 40kb		Library construction						
	Time consuming;	Validamycin,	[5, 6, 7]					
$\sim 40 \text{kb}$	Labour intensive.	Spinosad,						
> 90kb		Borrelidin, etc.						
67kb	Mis-priming or mis-annealing for high GC	Taromycin, etc.	[8]					
	content or repeated sequence (e.g. Polyketide							
	synthase)							
> 200kb	Mis-priming or mis-annealing for high GC	N/A	[9]					
	content or repeated sequence; Mutation							
	caused by PCR							
$\sim 50 \text{kb}$	Restriction enzymes limitation;	Glidobactins, etc.						
			[10-12]					
106kb		Salinomycin,						
		Spinosad.	[4, 13]					
	synthase)							
	1							
72kb		Pristinamycin	[12, 14]					
	pairings for high GC content sequence							
CRISPR/Cas9								
~ 150kb (50%	Mismatched linker pairings for high GC	Jadomycin,						
GC) or	content sequence	Chlortetracycline	[2, 15]					
~ 36kb (72% GC)								
$\sim 40 kb$	Difficult for large gene clusters cloning, e.g.	Sisomicin	[3]					
	> 50kb.							
	> 90kb 67kb > 200kb ~ 50kb 106kb 72kb ~ 150kb (50% GC) or ~ 36kb (72% GC)	> 90kb         67kb       Mis-priming or mis-annealing for high GC content or repeated sequence (e.g. Polyketide synthase)         > 200kb       Mis-priming or mis-annealing for high GC content or repeated sequence; Mutation caused by PCR         ~ 50kb       Restriction enzymes limitation; Mis-priming or mis-annealing for high GC content or repeated sequence (e.g. Polyketide synthase)         106kb       Mis-priming or mis-annealing for high GC content or repeated sequence (e.g. Polyketide synthase)         106kb       Mis-priming or mis-annealing for high GC content or repeated sequence (e.g. Polyketide synthase)         72kb       Mutation caused by PCR; Mismatched linker pairings for high GC content sequence         ~ 150kb (50% GC) or ~ 36kb (72% GC)       Mismatched linker pairings for high GC content sequence         ~ 40kb       Difficult for large gene clusters cloning, e.g. > 50kb.	> 90kb       Borrelidin, etc.         67kb       Mis-priming or mis-annealing for high GC content or repeated sequence (e.g. Polyketide synthase)       Taromycin, etc.         > 200kb       Mis-priming or mis-annealing for high GC content or repeated sequence; Mutation caused by PCR       N/A         ~ 50kb       Restriction enzymes limitation; Mis-priming or mis-annealing for high GC content or repeated sequence (e.g. Polyketide synthase)       Glidobactins, etc.         106kb       Mis-priming or mis-annealing for high GC content or repeated sequence (e.g. Polyketide synthase)       Salinomycin, Spinosad.         106kb       Mis-priming or mis-annealing for high GC content or repeated sequence (e.g. Polyketide synthase)       Salinomycin, Spinosad.         72kb       Mutation caused by PCR; Mismatched linker pairings for high GC content sequence       Pristinamycin         ~ 150kb (50% GC)       Mismatched linker pairings for high GC content sequence       Chlortetracycline         ~ 36kb (72% GC)       Difficult for large gene clusters cloning, e.g. > 50kb.       Sisomicin					

Table 1. Previous approaches that have been used for BGC cloning in vitro.

BAC: bacterial artificial chromosome; ULCC: upper limit of cloning capacity; BGC: biosynthetic gene cluster; NP: natural product; CATCH: Cas9-assisted targeting of chromosome segments; TAR: transformation-associated recombination; YA: yeast assembly; GA: Gibson assembly; ExoCET: exonuclease 466 combined with RecET recombination.

Items		Target BGCs		
		Paulomycin	Surugamides	Candicidin
Source strain		S. albus J1074	S. albus J1074	S. albus J1074
<b>Cloning vector</b>		pBAC2015	pBAC2015	pBAC2015
Genome size of source	e strain	6.8 Mb	6.8 Mb	6.8 Mb
GC content of genome		73%	73%	73%
Type of target NP		Quinone glycoside	Polyketide	Polyketide
Size of target BGC		49kb	87kb	139kb
GC content of target	BGCs	71%	76%	75%
White clone, c.f.u (/m	L)	659 ± 28 *	$508 \pm 56$ *	$330\pm47\texttt{*}$
	Repeat 1	4/96	3/96	1/192
Corrected/checked	Repeat 2	4/96	2/96	0/96
(by PCR)	Repeat 3	5/96	4/96	0/96
Percentage of right clone **		100%	100%	N/A

Table 2. The cloning efficiencies of high-GC target BGCs from genomic DNA.

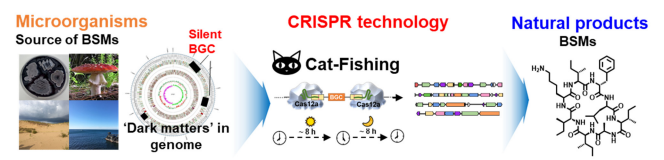
\* (Optional) Commercially available ElectroMAXTM DH10BTM cells (Thermo Fisher Scientific, Invitrogen<sup>TM</sup>) could be used to increase the number of transformants. All experiments were performed in triplicate. \*\* Verified by restriction enzyme digestion from 5 PCR-verified clones.

Methods	ULCC (high GC)*	CE	ТС	Cost	CD	Ref.
BAC library	> 120 kb	$\sim 1/1000$	> one month	> 5000 \$**	Hard	[7]
YA	NA	NA	NA	NA	Medium	[9]
САТСН	36 kb	> 90%	$3 \sim 4 \ days$	NA	Easy	[2,15]
ExoCET	106 kb	1/24	> one week	NA	Medium	[3,13]
CAT-	139 kb	1/192	2 4 4	10 ¢	<b>F</b>	This
FISHING	87 kb	~ 4/96	$3 \sim 4$ days	~ 40 \$	Easy	study

Table 3. Comparison of selected methods of large DNA fragment (> 80kb) cloning.

CE: Cloning Efficientcy; TC: Time Consuming; CD: Cloning Difficulty. \* Upper limit of cloning capacity for DNA sample with high GC content (> 70%). \*\* Price information for BAC library construction was provided by Wuhan Eightstars Bio-Technology Co. Ltd.

# **ToC figure**

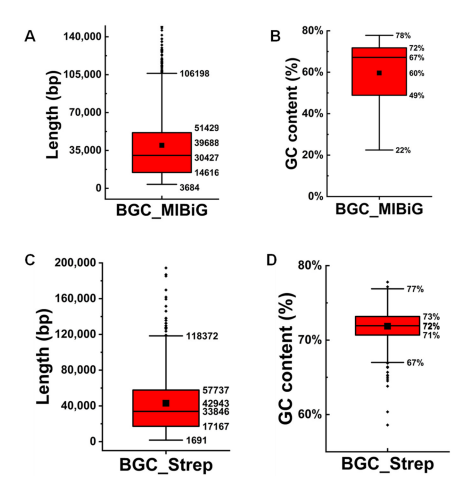


Directly cloning of large biosynthetic gene cluster (BGC) from even unculturable microbial genomes revolutionized nature products-based drug discovery. 47-139kb of target DNA fragments were simply and rapidly direct cloned from high GC genomic DNA sample by CAT-FISHING. Surugamides, encoded by a captured 87-kb gene cluster, was expressed and identified in a cluster-free Streptomyces chassis, validating the great potential of CAT-FISHING.

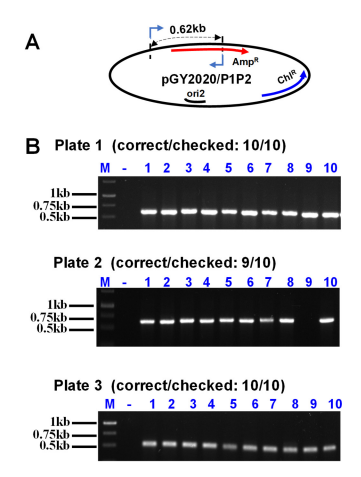
# Supporting Information

# Simple cloning of large natural product biosynthetic gene clusters from Streptomyces by an engineered CRISPR/Cas12a system

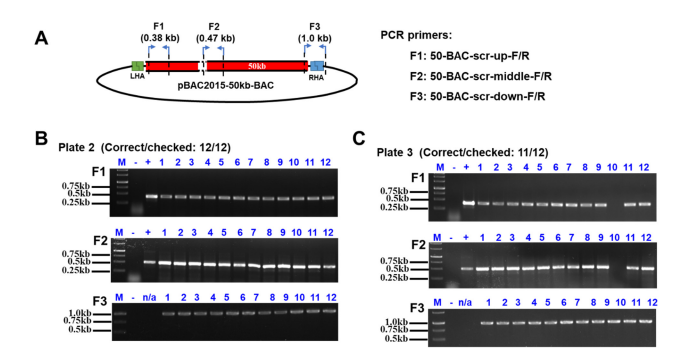
Mindong Liang, et al.



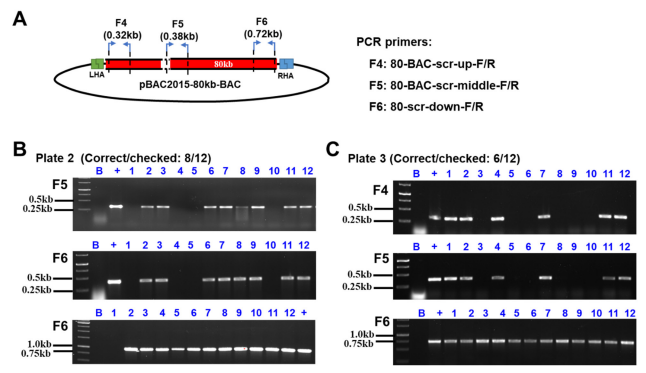
**Figure S1.** Statistical analysis of GC content and length of characterized BGCs in the MIBiG database. A-B. Distribution of BGCs length and GC content in characterized gene clusters identified in the MIBiG database. C-D. Distribution of BGCs length and GC content in characterized gene clusters identified in Streptomyces. 1910 characterized BGCs were download from MIBiG database (https://mibig.secondarymetabolites.org/stats)



**Figure S2.** Screening of right clones by PCR amplification. A. Schematic diagram of PCR screening of right clones with primers Amp-Cas12a-scr-F/R. B. PCR amplification of ten randomly selected clones. All experiments were performed in triplicate. "-" represented blank control, genomic DNA of *E. coli* DH10B was used as PCR template.



**Figure S3.** Screening of right clones that containing pBAC2015-50kb-BAC by PCR amplification. A) A. Schematic representation of crRNA design and cohesive end ligation. F1, F2 and F3 are the PCR products that were amplified using 50-BAC-scr-up-F/R, 50-BAC-scr-middle-F/R and 50-BAC-scr-down-F/R, respectively. B-C) PCR screening of 12 randomly selected clones containing pBAC2015-50kb-BAC in independent experiments/plates.



**Figure S4.** Screening of right clones that containing pBAC2015-80kb-BAC by PCR amplification. A) Schematic representation of crRNA design and cohesive end ligation. F4, F5 and F6 are the PCR products that were amplified using 80-BAC-scr-up-F/R, 80-BAC-scr-middle-F/R and 80-scr-down-F/R, respectively. Primer sequences are given in Table S7. B, C and D. PCR screening of 12 randomly selected clones containing pBAC2015-80kb-BAC in independent experiments/plates.

Method	Procedure	Timing	Cloning performance	References
	sgRNA template preparation and in vitro transcription	6 h	Strain: Bacillus subtilis, etc	
	Agarose plug preparation and in-gel cell lysis	2 d	GC% of genome: 45%	
	In-gel Ca9 digestion	3 h	Target fragment: unknown PKS	[1, 2]
CATCH*	Preparation of cloning vectors	4 h	DNA fragment length: 78 kb	[-, -]
CAICH	Ligation and electrotransformation	2 h	GC% of fragment: 46%	
	Validation of positive clones	1 d	Cloning efficiency: 12%	
		In total: $3 \sim 4 d$	ULCC: 150 kb (GC content: 51%);	
			36kb (GC content: 72%)	
	Preparation of the E. coli strain	2 d		
	Preparation of linear vectors for direct cloning	$1 \sim 7 d$	Strain: S. albus DSM4139	
	Preparation of cassettes and vectors for engineering	1 d	GC% of genome: 72.6%	
	Restriction digestion of genomic to release the target gene cluster	5 – 6 h	Target BGC: salinomycin	
RecET*	Preparation of the overnight E. coli culture	12 h (overnight)	BGC length: 106 kb	[3, 4]
(ExoCET)	Preparation of electro-competent E. coli cells	$3-3.5 \ h$	GC% of BGC: 74%	[2, 1]
	Electroporation of genomic and the linear vector into E. coli cells	1.5 h	Cloning efficiency: $4 \sim 8 \%$	
	Setting up cell cultures for restriction analysis	12 h (overnight)	ULCC: 106 kb (GC content: 74%)	
	Screening of correct recombinants by restriction analysis	7–9 h		
		In total: > one week		
	crRNA template preparation and in vitro transcription	~ 6 h	Strain: S. albus J1074	
	Genomic DNA isolation and in-gel Cas12a digestion	~ 1 d	GC% of genome: 73.3%	
CAT	Plasmid construction and Cas12a digestion	~ 1 d	Target BGC: surugamide	
CAT-	Ligation and transformation	2 h	BGC length: 87 kb	This study
FISHING	Validation of positive clones	1 d	GC% of BGC: 76%	
		In total: 3 ~ 4 d	Cloning efficiency: ~ 4%	
			ULCC: 139 kb (GC content: 75%)	

#### Table S1. Comparison of different protocols of large DNA fragment cloning.

ULCC: upper limit of cloning capacity; BGC: biosynthetic gene cluster. \* The description of procedures adapted from Nature Protocol<sup>[2, 3]</sup>.

Items	Consumption and cost (per 50 µL reaction)	Price & Sources
Cas12a	40 pmol; ~ 9\$	2000 pmol, 450\$, NEB
crRNA1	20 pmol; ~12.5\$	(See Table S3)
crRNA2	20 pmol; ~12.5\$	(See Table S3)
T4 DNA ligase	200 U; ~ 2.2\$	10,000 U, 110\$, NEB
Low-melt agarose	0.1 g; ~ 1\$	5 g, 30\$, Sangon Biotech
β-Agarase I	2 U; ~2.5\$	100 U, 125\$, NEB
		In total: ~40\$

 Table S2. Cost accounting of CAT-FISHING reaction system.

#### Table S3. Cost accounting of crRNA preparation.

Itmes	Consumption and cost (per 50 µL reaction)	Price & Sources
RNA Clean & Concentrator Kits	1 prep; 3\$	50 preps, 150\$, Zymo Research
HiScribe T7 High Yield RNA Synthesis Kit	1 reaction; ~9\$	50 reactions, 455\$, NEB
RNase Inhibitor	40 U; ~0.5\$	5000 U, 60\$, Takara
		In total: ~12.5\$

Plasmid or strains	Features	Source/Ref.
Plasmid		
pUC19	High copy number cloning vector that conveys the Amp resistance	NEB
pCC2FOS	Fosmid vector, used for fosmid library construction.	EpicentreBio
pET28a	IPTG-inducible expression vector, used for recombination protein expression	Novagen
pSET152	Site-specific integrating vector; Containing the $\Phi C31$ attachment site and <i>oriT</i> , and can be transferred by conjugation into <i>Streptomycetes</i> from <i>E. coli</i>	[5]
pBAC2015	Copy-control BAC vector without repetitive sequences.	[3]
pUB307	RPI-derived self-transmissible plasmid, providing <i>in trans</i> the function for the mobilization of the <i>oriT</i> -containing BAC plasmid	[6]
pSC101-BAD-αβγA-tet	Red $\alpha\beta\gamma$ expression plasmid for linear plus circular homologous recombination	[7]
pBAC-ZL	BAC plasmid pIndigoBAC536-S containing 137kb DNA fragment	This study
pGY2020	Amp resistance gene in pCC2FOS	This study
pBAC2015-C50	pBAC2015 derived plasmid, used for 50kb DNA fragment (from pBAC-ZL) cloning	This study
pBAC2015-C80	pBAC2015 derived plasmid, used for 80kb DNA fragment (from pBAC- ZL) cloning	
pBAC2015-CS49	pBAC2015 derived plasmid, used for 49kb paulomycin biosynthetic gene cluster cloning	
pBAC2015-CS87	pBAC2015 derived plasmid, used for 87kb paulomycin biosynthetic gene cluster cloning	
pBAC2015-50kb-BAC	pBAC2015-C50 containing the 50kb DNA fragment (from pBAC-ZL)	This study
pBAC2015-80kb-BAC	pBAC2015-C80 containing the 80kb DNA fragment (from pBAC-ZL)	This study
pBAC2015-49kb-J1074	pBAC2015-CS49 containing the whole paulomycin biosynthetic gene cluster (49kb)	This study
pBAC2015-87kb-J1074	pBAC2015-CS87 containing the whole surugamides biosynthetic gene cluster (87kb)	This study
pBAC2015-49kb-J1074-int	The $aac(3)IV$ -oriT-attP( $\Phi C31$ )-int( $\Phi C31$ ) cassette from pSET152 was introduced into the pBAC2015-49kb-J1074 by replacement of chloramphenicol-resistant (cmr) gene	This study
pBAC2015-87kb-J1074-int	The $aac(3)IV$ -oriT-attP( $\Phi C31$ )-int( $\Phi C31$ ) cassette from pSET152 was introduced into the pBAC2015-87kb-J1074 by replacement of chloramphenicol-resistant (cmr) gene	This study

#### Table S4. The strains and plasmids used in this study.

Plasmid or strains	Features	Source/Ref.
E. coli		
EPI300 <sup>™</sup> -T1R	$F-mcrA \Delta(mrr-hsdRMS-mcrBC) (StrR) \varphi 80dlacZ \Delta M15 \Delta lacX74$	EpicentreBio
	recA1 endA1 araD139 $\Delta$ (ara, leu)7697 galU galK $\lambda$ - rpsL nupG trfA tonA dhfr	
DH10B	F–mcrA Δ(mrr-hsdRMS-mcrBC) φ80dlacZ ΔM15 ΔlacX74 deoR	Gibco BRL
	recA1 endA1 araD139 $\Delta$ (ara, leu)7697 galU galK $\lambda$ – rpsL nupG	
BL21(DE3)	F-ompT hsdSB(rB-mB-) gal dcm (DE3)	TIANGEN,
		Shangai
ET12567	F–dam13::Tn9 dcm6 hsdM hsdR recF143 zjj201::Tn10 galK2	[8]
	galT22ara14 lacY1 xyl5 leuB6 thi1 tonA31 rpsL136 hisG4 tsx78 mtli	
	glnV44	
ET12567(pUB307)	ET12567 containing plasmid pUB307	This study
Streptomyces		
S. albus J1074	Wild-type strain, heterologous host	[9]
S. albus Del14	Derivative of J1074 with deletion of 15 gene cluster	[10]
S. albus Del14-49kb	BAC plasmid pBAC2015-49kb-M integrated into the genome of	This study
	Del14	
S. albus Del14-87kb	BAC plasmid pBAC2015-87kb-M integrated into the genome of	This study
	Del14	

Table S4. The strains and plasmids used in this study (continued).

Table S5. The sequence of PCR primers for plasmid construction.

Name	Sequence $(5' \rightarrow 3')$
pGY2020 construction	
pCC2-hom-arm-up-F	TACAACGACACCTAGACCACGGTGGAGCTGCGCAACCGGC
pCC2-hom-arm-up-R	ACAGACAAGCTGTGACCGTCTCCTGGGGCGTGACACCACC
pCC2-hom-arm-dn-F	GCGCGGGGAGAGGCGGTTTGGACATTGCACTCCACCGCTG
pCC2-hom-arm-dn-R	AGGAAACAGCCTAGGAACACGGAAGTTTTTTCAGGCATCG
pBAC2015-C50 construction	
50kb-BAC-arm-up-F	TTATCTATGCTCGGGGGATGCCGCGTGGTACC
50kb-BAC-arm-up-R	ACAGACAAGCTGTGACCGTCGGCGGCATCCCGATCAGCGC
BAC-arm-dn-F	GCGCGGGGAGAGGCGGTTTGGACATTGCACTCCACCGCTG
BAC-arm-dn-R	TAGAGAGGATACCGGAGATCCTTTGATCTTTTC
BAC-lacZ-F	GACGGTCACAGCTTGTCTGTAAGC
BAC-lacZ-R	CAAACCGCCTCTCCCCGCGCGTTGG
pBAC2015-C80 construction	
80kb-BAC-arm-up-F	TTATCTATGCTCGGGGGTGGAGCTGCGCAACCG
80kb-BAC-arm-up-R	ACAGACAAGCTGTGACCGTCTCCTGGGGCGTGACACCACC
BAC-lacZ-F	GACGGTCACAGCTTGTCTGTAAGC
BAC-lacZ-R	CAAACCGCCTCTCCCCGCGCGTTGG
BAC-arm-dn-F	GCGCGGGGAGAGGCGGTTTGGACATTGCACTCCACCGCTG
BAC-arm-dn-R	TAGAGAGGATACCGGAGATCCTTTGATCTTTC
pBAC2015-CS49 construction	
50kb-up-hom-arm-F	CACTCACTCACCCCGGTCACATCGTTATCTATGCTCGGCGATCGCGCTGGAGTCCTTCG
50kb-up-hom-arm-R	TTTGCGTATTGGGCAATTCTCATGTTTGACCGCTTATCTGGATCTGCCCTTTCCACTC
50kb-lacZ-bla-F	ATAAGCGGTCAAACATGAGA
50kb-lacZ-bla-R	AATGAAGTTTTAAATCAATC
50kb-dn-hom-arm-F	TTTACTCATATATACTTTAGATTGATTTAAAACTTCATTTAATCCTGTCGACCGTCAAG
50kb-dn-hom-arm-R	AGGAAATTTATCTTGATCATATAAATAGAGAGGATACCGTGACCAAGCGGTACGACTTC
pBAC2015-CS87 construction	
80kb-up-hom-arm-F	CATCGTTATCTATGCTCGGGGATGTCGACGTCCAGGGTG
80kb-up-hom-arm-R	CGGTTTGCGTATTGGGCAATTGATTCGCCGGCGTTCTG
80kb-lacZ-F	ATTGCCCAATACGCAAAC
80kb-lacZ-R	CTAAGAAACCATTATTATC
80kb-dn-hom-arm-F	TGATAATAATGGTTTCTTAGGGAAGCGGTCTCCTGAAGC
80kb-dn-hom-arm-R	ATAAATAGAGAGGATACCGGTTTTCCCCCGTTGATGAGTGG

Table S6. The sequence of crRNA for CRISPR/Cas12a cleavage.

Name	Sequence $(5' \rightarrow 3')$	Cleavage
T7-promoter-F	GAAATTAATACGACTCACTATAGGG	
50kb-BAC-up-crRNA-R	GTGCAGACCCGGATTCCGATCTACAACAGTAGAAATTCCCTATAGTGAGTCGTATTAAT	50kb BAC
80kb-BAC-dn-rRNA-R	CGATCAACGGCACTGTTGATCTACAACAGTAGAAATTCCCTATAGTGAGTCGTATTAAT	50kb BAC
80kb-BAC-up-crRNA-R	TGCGGAACTTCAAGCGCCATCTACAACAGTAGAAATTCCCTATAGTGAGTCGTATTAAT	80kb BAC
80kb-BAC-dn-crRNA-R	CGATCAACGGCACTGTTGATCTACAACAGTAGAAATTCCCTATAGTGAGTCGTATTAAT	80kb BAC
50kb-up-crRNA-R	GTACGCGGGCAGCGTGAGATCTACAACAGTAGAAATTCCCTATAGTGAGTCGTATTAAT	49kb BGC
50kb-dn-crRNA-R	GGACAGAGATTTCGCCAAATCTACAACAGTAGAAATTCCCTATAGTGAGTCGTATTAAT	49kb BGC
80-up-crRNA-R	CCTGGCCGCCCGCCCGATCTACAACAGTAGAAATTCCCTATAGTGAGTCGTATTAAT	87kb BGC
80kb-dn-crRNA-R	CACAGTTTCGGTCCTCGGATCTACAACAGTAGAAATTCCCTATAGTGAGTCGTATTAAT	87kb BGC

Name	Sequence $(5' \rightarrow 3')$	Note
Amp-Cas12a-scr-F	GGGAACCGGAGCTGAATGAA	pGY2020-P1_P2
	CTGACGCTCAGTGGAACGAA	
Amp-Cas12a-scr-R	CIGACOCICAOIOGAACGAA	pGY2020-P1_P2
50-BAC-scr-up-F	GGCAGTTTCATCGTGGCGTA	50kb BAC-Ch-U
50-BAC-scr-up-R	GCGGGACTCACATGGGTTTT	50kb BAC-Ch-U
50-BAC-scr-mid-F	CAATCCGATGACACGGCACA	50kb BAC-Ch-M
50-BAC-scr-mid-R	CCGTGGTTGTCGCTGTACTC	50kb BAC-Ch-M
BAC-scr-down-F	TGCCATCAACTCGGCAAGAT	50kb BAC-Ch-D
BAC-scr-down-R	TCTTTCGCGAAGGCTTGAGT	50kb BAC-Ch-D
80-BAC-scr-up-F	TGGTGTGTTGTCGTTGTCGG	80kb BAC-Ch-U
80-BAC-scr-up-R	GGAGATCTGGGCGAACTCCT	80kb BAC-Ch-U
80-BAC-scr-mid-F	GGCAGTTTCATCGTGGCGTA	80kb BAC-Ch-M
80-BAC-scr-mid-R	GCGGGACTCACATGGGTTTT	80kb BAC-Ch-M
BAC-scr-down-F	TGCCATCAACTCGGCAAGAT	80kb BAC-Ch-D
BAC-scr-down-R	TCTTTCGCGAAGGCTTGAGT	80kb BAC-Ch-D
50-scr-up-F	TCCATTCCGTGCCATGCG	49kb BGC-Ch-U
50-scr-up-R	GCGACGAGAGAGGATGTG	49kb BGC-Ch-U
50-scr-middle-F	ACAAGGCTCCTGACAGG	49kb BGC-Ch-M
50-scr-middle-R	GAGGTGGTGCACCTGG	49kb BGC-Ch-M
50-scr-down-F	GTGTATCGCGCCGCTG	49kb BGC-Ch-D
50-scr-down-R	CTATGCTCCAGACATC	49kb BGC-Ch-D
87-scr-up-F	GCAGCGAACTGCCTGGT	87kb BGC-Ch-U
87-scr-up-R	CCGATGAGGTCGTTCAC	87kb BGC-Ch-U
87-scr-middle-F	ATGCCCGTCAACTGCTCCTG	87kb BGC-Ch-M
87-scr-middle-R	TGATTTCCCGACCGTTT	87kb BGC-Ch-M
87-scr-down-F	GCCGGTTCAGGCGCGCT	87kb BGC-Ch-D
87-scr-down-R	AGAAGGGCAAGTTGTGC	87kb BGC-Ch-D
Redαβ-scr-F	ACGTATGTGGTGTGACCGGA	49kb BGC-Red
Redaβ-scr-R	CATTGCACTCCACCGCTGAT	49kb BGC-Red
Redαβ-scr-F	ACGTATGTGGTGTGACCGGA	87kb BGC- Red
Redαβ-scr-R	CATTGCACTCCACCGCTGAT	87kb BGC- Red

Table S7. The sequence of PCR primers for screening, verification and modification.

#### **SI references**

- [1] W. Jiang, X. Zhao, T. Gabrieli, C. Lou, Y. Ebenstein, T. F. Zhu, Nat. Commu. 2015, 6, 8101.
- [2] W. Jiang, T. F. Zhu, Nat. Protoc. 2016, 11, 960.
- [3] H. Wang, Z. Li, R. Jia, Y. Hou, J. Yin, X. Bian, A. Li, R. Müller, A. F. Stewart, J. Fu, Y. Zhang, Nat. Protoc. 2016, 11, 1175.
- [4] H. Wang, Z. Li, R. Jia, J. Yin, A. Li, L. Xia, Y. Yin, R. Müller, J. Fu, A. F. Stewart, Y. Zhang, Nucleic Acids Res. 2017, 46, 2697.
- [5] M. Bierman, R. Logan, K. O'Brien, E. T. Seno, R. Nagaraja Rao, B. E. Schoner, Gene 1992, 116, 43.
- [6] F. Flett, V. Mersinias, C. P. Smith, FEMS Microbiol. Lett. 1997, 155, 223.
- [7] Y. Zhang, J. P. P. Muyrers, G. Testa, A. F. Stewart, Nat. Biotechnol. 2000, 18, 1314.
- [8] D. J. MacNeil, K. M. Gewain, C. L. Ruby, G. Dezeny, P. H. Gibbons, T. MacNeil, Gene 1992, 111, 61.
- [9] K. F. Chater, L. C. Wilde, J Gen. Microbiol. 1980, 116, 323.
- [10] M. Myronovskyi, B. Rosenkränzer, S. Nadmid, P. Pujic, P. Normand, A. Luzhetskyy, *Metab. Eng.* 2018, 49, 316.