Multigene disruption in undomesticated Bacillus subtilis ATCC 6051a using the CRISPR/Cas9 system

Kang Zhang1,2,*, Xuguo Duan1,2,† & Jing Wu1,2

Bacillus subtilis ATCC 6051a is an undomesticated strain used in the industrial production of enzymes. Because it is poorly transformable, genetic manipulation in this strain requires a highly efficient genome editing method. In this study, a Streptococcus pyogenes CRISPR/Cas9 system consisting of an all-in-one knockout plasmid containing a target-specific guide RNA, \textit{cas}9, and a homologous repair template was established for highly efficient gene disruption in \textit{B. subtilis} ATCC 6051a. With an efficiency of 33% to 53%, this system was used to disrupt the \textit{srfC}, \textit{spoIIAC}, \textit{nprE}, \textit{aprE} and \textit{amyE} genes of \textit{B. subtilis} ATCC 6051a, which hamper its use in industrial fermentation. Compared with \textit{B. subtilis} ATCC 6051a, the final mutant, BS5 (\textit{Δ}srfC, \textit{Δ}spoIIAC, \textit{Δ}nprE, \textit{Δ}aprE, \textit{Δ}amyE), produces much less foam during fermentation, displays greater resistant to spore formation, and secretes 2.5-fold more 3-\textit{β}-cyclodextrin glycosyltransferase into the fermentation medium. Thus, the CRISPR/Cas9 system proved to be a powerful tool for targeted genome editing in an industrially relevant, poorly transformable strain.

Bacillus subtilis, a well-characterized gram-positive bacterium, has been widely used for the production of heterologous proteins. This species and some of its close relatives have excellent protein secretory capability and are generally recognized as safe (GRAS), making them important hosts for the production of antibiotics, medicinal proteins, and industrial enzymes. \textit{B. subtilis} 168 is a model laboratory strain that carries many mutations that have occurred during its modification via irradiation and selection\cite{1}. These modifications have made the organism tryptophan-deficient and improved its transformability. The \textit{B. subtilis} strains commonly used for recombinant protein production, such as WB600 and WB800, were constructed on the basis of \textit{B. subtilis} 168\cite{2}.

Because the recombinant protein productivity of \textit{B. subtilis} ATCC 6051a is superior to that of \textit{B. subtilis} 168, it has been widely applied to the production of industrial enzymes\cite{3,4}. However, \textit{B. subtilis} ATCC 6051a has some undomesticated properties that hamper the extracellular production of recombinant proteins. In particular, it can produce large amounts of foam, highly resistant spores, multiple types of extracellular protease, and high level of amylase during fermentation, which related to the \textit{srfC}, \textit{spoIIAC}, \textit{nprE}, \textit{aprE} and \textit{amyE}, respectively. To improve the usefulness of this important strain, we sought to modify these properties by inactivating the five genes. \textit{B. subtilis} ATCC 6051a is poorly transformable, compared with laboratory strains, because it harbours an 84-kb endogenous plasmid pBS32, which encodes a single-pass trans-membrane protein ComI that inhibits the competence of DNA uptake\cite{5}. Due to its poor competence, the genetic manipulation of \textit{B. subtilis} ATCC 6051a is difficult and require a highly efficient genome editing method. The genome sequence of \textit{B. subtilis} ATCC 6051a was recently determined by Jeong \textit{et al.}\cite{6}, which facilitates genetic manipulation.

Clustered regularly interspaced short palindromic repeat (CRISPR) systems, which are composed of CRISPR RNAs (crRNA), trans-activating CRISPR RNAs (tracrRNAs) and CRISPR-associated (Cas) proteins constitute an immune system in bacteria and archaea that efficiently cleaves foreign DNA entering the cell, including phages and plasmids\cite{7}. Some CRISPR/Cas systems require multiple proteins\cite{8}, whereas the type II CRISPR/Cas system requires a single nuclease; Cas protein 9 (Cas9). In the widely used \textit{Streptococcus pyogenes} type II CRISPR/Cas system\cite{9}, the 20-bp complementary region (N20) within the crRNA guide Cas9 nuclease to its specific target, a

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1State Key Laboratory of Food Science and Technology, Jiangnan University, 1800 Lihu Avenue, Wuxi, 214122, China. 2School of Biotechnology and Key Laboratory of Industrial Biotechnology Ministry of Education, Jiangnan University, 1800 Lihu Avenue, Wuxi, 214122, China. *These authors contributed equally to this work. Correspondence and requests for materials should be addressed to J.W. (email: jingwu@jiangnan.edu.cn)
therefore 20-nt sequence known as the protospacer, which contains a specific protospacer-adjacent motif (PAM) at its 3’ end. The PAM sequence leads cas9 to create a double-strand break at protospacer (target) sequence and it was repaired through homologous recombination using a repair template that is supplied along with the CRISPR/Cas9 system. Recently, a single chimeric guide RNA (sgRNA) containing features of both crRNA and tracrRNA has been developed, which simplify the genome editing design. As an efficient genome editing technology, the type II CRISPR/Cas9 system has been proved to be feasible in point mutation, single gene deletion/insertion and large-size gene cluster deletion. And until recently, it has been widely applied in various organisms including, but not restricted to, Escherichia coli, Streptococcus pneumonia, Saccharomyces cerevisiae, Lactobacillus reuteri, Bombyx mori, Drosophila and humans cell lines.

Although there have been several successful implementations of the CRISPR/Cas9 system in microbial systems, there have not been any reports of genome editing in B. subtilis using a CRISPR/Cas9 system. This study describes the establishment and optimization of a CRISPR/Cas9 system in B. subtilis ATCC 6051a. We used this system to disrupt five genes in the B. subtilis ATCC 6051a genome (srfC, spoIAC, napE, aprE and amyE) that hamper its use during industrial fermentation. Compared with B. subtilis ATCC 6051a, this mutant strain, named BS5, produces much less foam at the level of control, exhibits great resistance to spore formation, and secretes 2.5 times more β-cyclodextrin glycosyltransferase (β-CGTase) into the fermentation medium. β-CGTase is widely used in the production of β-cyclodextrin and mainly produced from wild strains and E. coli, which exist low secretion and food safety problem, respectively. The increase of β-CGTase secretion from Bacillus subtilis can largely reduce the cost of β-cyclodextrin production, which shows the high industrial value of BS5.

**Results**

**Construction of the CRISPR/Cas9 system all in one plasmid.** We assembled a complete CRISPR/Cas9 system in a single knockout plasmid that could be used for highly efficiently genome editing in B. subtilis (Fig. 1a). The six knockout plasmids (pHYcas9dsrf1, pHYcas9dsrf2, pHYcas9dspo, pHYcas9dnp, pHYcas9dapr and pHYcas9damy), which originated from plasmid pHY300PLK-β-CGTase, consist of the cas9 gene amplified from plasmid pwtcarg-bacterial, a sgRNA and its promoter P43, the temperature-sensitive replicon PE194, and a homologous repair template. The synthesis of Cas9 protein in vivo was driven by the α-amanase promoter (PamyQ), which originates from B. amyloliquefaciens. The P43 promoter is a constitutively expressed promoter that can drive strong transcription of the sgRNA in B. subtilis. The homologous repair template was obtained through overlap extension PCR of regions upstream and downstream of the target locus. The length of the upstream and downstream regions ranged from 450 to 550 bp, and they introduced an 500 bp upstream region and 600 bp downstream region flanking the Cas9 cleavage site. Using knockout plasmids and plasmid pHY300PLK-β-CGTase encode ampicillin resistance in E. coli and tetracycline resistance in both E. coli and B. subtilis.

**Disruption of the srfC gene using the CRISPR/Cas9 system.** We used B. subtilis ATCC 6051a as the initial strain in which to perform genetic manipulations. Fermentation of B. subtilis ATCC 6051a in a 3 L fermenter in our laboratory produced a massive amount of foam that requires large quantities of antifoam agent. Accumulation of the amphiphilic molecule surfactin promotes foam formation, and the gene srfC is crucial to the regulation of surfactin production. The CRISPR/Cas9 system developed in this study was first tested using srfC as the target (Fig. 1c). We used the knockout plasmid pHYcas9dsrf1, which produces both a sgRNA specific to srfC and the Cas9 protein, to transform the initial strain, B. subtilis ATCC 6051a. Because the temperature-sensitive knockout plasmid is maintained at low copy in transformants at 37 °C, the tetracycline-resistant transformants were confirmed by colony PCR of cas9. As for the commonly same genotype of transformation colonies, the target cleavage by Cas9 and homology-directed repair are commonly performed during the liquid incubation of transformation. The resulting mutants had an Xho I site within the repair locus, and this disruption genotype passed to the next generation through passing in liquid media or LB plate. To test the effectiveness of the CRISPR/Cas9 system, the regions upstream and downstream of the repair locus were amplified by PCR, and then the PCR products were digested with Xho I (Fig. 3). There are 66 ± 14 transformants after transformation of knockout plasmid pHYcas9dsrf1. 30 transformants were screened, and 13 ± 2 colonies yielded a recombinant genotype with a disruption efficiency of 43% ± 6%. The results, combined with DNA sequencing of the homologous regions of the mutants (Supplementary Fig. S1a) demonstrated that the editing system described above worked efficiently. Curing the knockout plasmid pHYcas9dsrf1 through overnight incubation at 51 °C produced the desired mutant, named BS1. Colonies of BS1 were used as the initial strain for next gene disruption. There are 20 ± 4 transformants after transformation of initial knockout plasmid pHYcas9d, and no srfC gene mutant was found among the transformants. Though DNA nonhomologous end joining exist in B. subtilis, the low survival rate was probably due to the double-strand breaks in chromosome created by cas9 protein. To test foam production by B. subtilis ATCC 6051a and mutant BS1, cells were grown in a 3 L fermenter for 80 h, and the growth rate of B. subtilis ATCC 6051a and BS1 has no significant difference. During the whole fermentation process, B. subtilis ATCC 6051a produced much foam and required to add antifoam (490 ul) continuously; BS1 produced much less foam at a controllable level that needed 60 ul antifoam. The foam height of B. subtilis ATCC 6051a and BS1 was similar after 42 h, while the foam of BS1 shows great sensitive to antifoam (Fig. 2).

In addition to disruption described above, which deleted a relatively small section of the srfC gene. An 1100 bp homologous repair template was designed to have a 500 bp upstream region and 600 bp downstream region flanking the Cas9 cleavage site. Using knockout plasmid pHYcas9dsrf2 that containing this repair template to delete a 284 bp region, which includes 44 bp upstream and 240 bp downstream of the PAM sequence (Supplementary Fig. S2). The efficiency of the 284 bp deletion is 9.1% in B. subtilis ATCC 6051a, which is lower than the efficiency of disruption gene. The low efficiency may related to the short length of homologous repair template that result in low homologous recombination efficiency.
Disruption of spoIIAC gene using the CRISPR/Cas9 system. In order to cope with limiting nutrient sources and high cell density, *B. subtilis* can form highly resistant spores that will germinate and grow in favourable living conditions. These spores drastically hamper *B. subtilis* fermentations and limit their application in the food industry. The *SpoIIAC* gene encodes sigma factor F, which permits cells to proceed through stage II of sporulation. Using knockout plasmid pHYcas9dspo, we disrupted *spoIIAC* gene using the method described above. The resulting mutant, named BS2, was screened by amplifying the upstream and downstream regions and digesting the PCR products with *Xho* I (Fig. 3). The homologous regions of mutant BS2 were also subjected to DNA sequencing (Supplementary Fig. S1b) to confirm the result. The disruption efficiency of *SpoIIAC* gene was 36% ± 3%. After being cultured at 40 °C for 48 h in a culture medium that favours spore formation, the sporulation efficiency of BS1 was 28.44% (262 colonies/921 colonies), while the sporulation efficiency of BS2 was 0% (0 colonies/182 colonies), which shows that mutant BS2 has great resistant to spore formation.

Disruption of *nprE* and *aprE* genes using the CRISPR/Cas9 system. Strains that lack several extracellular protease genes generally show superior extracellular protein productivity. The *nprE* and *aprE* genes encode alkaline and neutral extracellular proteases, respectively, in *B. subtilis*. We constructed the knockout
plasmids pHYcas9dnpr and pHYcas9dapr and used them to sequentially disrupt \( \text{nprE} \) and \( \text{aprE} \) gene of mutant BS2 using the CRISPR/Cas9 method described above. Amplification of the upstream and downstream regions, and then digesting the PCR products with \( \text{Xho I} \) allowed us to select the appropriate mutant, named BS3 with disruption efficiency of 53\% \( \pm \) 6\% (Fig. 3). The \( \text{nprE} \) and \( \text{aprE} \) double gene mutant, named BS4 with disruption efficiency of 33\% \( \pm \) 3\%. The homologous regions of mutant BS3 and BS4 were subjected to DNA sequencing (Supplementary Fig. S1c,d) to confirm the disruption. When cultured on specific agar plates containing 5% non-fat powdered milk, strain BS3 forms a protein clearance zone smaller than the one formed by its parent, mutant BS2, demonstrating that BS3 has reduced protease activity. And strain BS4 shows substantial inhibition of protein degradation on a 5% non-fat powdered milk plate (Fig. 4a).

Disruption of the \( \text{amyE} \) gene using the CRISPR/Cas9 system. \( B. \text{subtilis} \) releases many extracellular enzymes during the post-exponential growth phase, and \( \alpha \)-amylase is one of the major proteins released\(^{9,35} \). These proteins hamper the purification of industrial products. We constructed the knockout plasmid pHYcas9damy and used it to disrupt the \( \text{amyE} \) gene of mutant BS4 as described above. Amplification of the upstream and downstream regions, and then digesting the PCR products with \( \text{Xho I} \) allowed us to select the appropriate mutant, named BS5 with disruption efficiency of 53\% \( \pm \) 6\% (Fig. 3). The \( \text{amyE} \) deletion mutant, named BS5 with disruption efficiency of 33\% \( \pm \) 3\%. The homologous regions of mutant BS5 were also subjected to DNA sequencing (Supplementary Fig. S1d) to confirm the disruption. The results of a starch-plate assay demonstrate that BS5 fails to release \( \alpha \)-amylase activity (Fig. 4b,c).
Extracellular expression of β-CGTase using *B. subtilis* ATCC 6051a and mutant BS5. To evaluate the utility of the strain BS5, in which five genes (*srfC, spoIIAC, nprE, aprE* and *amyE*) have been disrupted, as a host for extracellular recombinant protein expression, the ability of mutant BS5 to produce β-CGTase was compared with that of *B. subtilis* ATCC 6051a. Both expression plasmid pHY300PLK-β-CGTase and empty expression plasmid pHY300PLK were transferred into *B. subtilis* ATCC 6051a and mutant BS5, and the resulting strains were used in an expression study. After 48 h of cultivation in TB medium, the culture supernatants of *B. subtilis* ATCC 6051a and BS5 that harbouring empty expression vector PHY300PLK show no β-CGTase activity. After 80 h of cultivation in 3 L fermenter, the highest β-CGTase activity of *B. subtilis* ATCC 6051a that harbours expression plasmid pHY300PLK-β-CGTase was 110.8 U/ml in 56 h, and the highest dry cell weight (DCW) was 68.8 g/L in 66 h; the highest β-CGTase activity of BS5 that harbours expression plasmid pHY300PLK-β-CGTase was 277.8 U/ml in 70 h, which was 2.5 times of *B. subtilis* ATCC 6051a, and the highest DCW was 70.3 g/L in 75 h (Fig. 5).

**Discussion**

In this study, we established a CRISPR/Cas9 system that can disrupt target genes in *B. subtilis* ATCC 6051a (Fig. 1c), and then used this system to construct a mutant strain with improved fermentation characteristics. All of the elements required by the CRISPR system are present in a single plasmid that contains a constitutively expressed *cas9*, a strongly transcribed sgRNA, and a homologous repair template. The sgRNA recognizes a specific site on the *B. subtilis* genome (Supplementary Table S1) and guides the Cas9 protein to the target genome locus, where it creates a double-stranded break. This is followed by homology-directed repair that utilizes a homologous repair template provided by the knockout plasmid. Since the knockout plasmid contains the PE194 temperature-sensitive replicon, it can be easily cured after the mutation step by incubating the mutants at 51 °C overnight. The mutant colonies cured of the knockout plasmid can be used as the parent strain for additional
genetic modification. With an efficiency of 33% to 53%, the operate procedure of CRISPR system was simple and time-saving compared with the currently existing *B. subtilis* genome editing methods (Table 1).

The transformation efficiency in *B. subtilis* 168 and *B. subtilis* ATCC 6051a are 1553 ± 213 and 60 ± 13 transformants/μg of the knockout plasmid, respectively. Its poor competence makes genetic manipulation of *B. subtilis* ATCC 6051a inconvenient. To increase the transformability of *B. subtilis* ATCC 6051a, we considered disrupting the gene encoding ComI. However, we were unable to find a target-specific sgRNA target within the 93 bp *comI* gene. The original knockout plasmid (pHYcas9d) did not contain a homologous repair template; therefore, a homologous repair template was transferred into *B. subtilis* ATCC 6051a in the form of PCR fragment, along with the knockout plasmid. This attempt did not meet the efficiency required for genetic editing, perhaps because *B. subtilis* ATCC 6051a was unable to simultaneously take up the knockout plasmid and the homologous repair PCR fragment.

The target recognition of the sgRNA mainly depends on the last 12 bp of the guide sequence; thus, the existence of highly homologous regions in chromosomal DNA may result in off-target effects. These off-target effects have been reported in eukaryotic cells, whereas little attention has been paid to this problem in prokaryotes. Although off-target effects may be less common in bacteria because of their relatively small genome size, bacterial genomes contain some high-homology clusters. There are methods that can be used to reduce the off-target efficiency; for example, designing two sgRNAs to guide the Cas9 protein to cleave the genome at the adjacent sites, using a Cas9 nickase mutant, and making sure the last 12 bp of the guide sequence is highly specific.

Like many *Bacillus* strains, *B. subtilis* produces surfactin, which contains a peptide moiety and a β-hydroxy fatty acid side chain. Because surfactin is an amphiphilic molecule, its accumulation at gas-liquid interfaces can lead to foam production. The biosynthesis of surfactin is controlled by multiple RNA polymerase sigma factors. Sigma F, encoded by *SpoIIAC*, controls the forespore, sigma E controls the early stage of sporulation, and then sigma G and K control later stages. *SpoIIAC* nonsense mutations can block the processing of sigma E precursor protein P31 and prevent transcription of *spoIID*. The transcription product, SpoIID, is a membrane-anchored enzyme essential for sporulation. The observation that mutant BS2 shows great resistance to spore formation demonstrates that inactivation of sigma F can largely block the sporulation of *B. subtilis*.

*Bacillus* species produce many different types of extracellular protease to degrade heterologous extracellular proteins, and many protease-deficient strains display favorable heterologous protein production. *B. subtilis* WB600, which is deficient in six extracellular proteases due to knockouts of the *nprB, nprE, aprE, mpr, bpr* and *epr* genes. This strain displays a level of recombinant protein secretion higher than that of its parent strain. Recently, a *B. amyloliquefaciens* strain lacking six extracellular protease genes was constructed. This strain displays improved production of levan and α-amylase. *B. subtilis* releases multiple extracellular enzymes during the post-exponential period; α-amylase is among the major proteins released. Secretion of a large amount of α-amylase by *B. subtilis* makes the isolation and purification of recombinant proteins difficult on an industrial scale. In addition, the massive expression of endogenous α-amylase increases secretion stress and influences the production of recombinant proteins.

In summary, we established a CRISPR/Cas9 system in the poorly transformable strain *B. subtilis* ATCC 6051a, which is an undomesticated strain with favorable growth characteristics. To improve the usefulness of *B. subtilis* ATCC 6051a as an industrial expression host, we disrupted the *srfC, spoIIAC, nprE, aprE* and *amyE* genes with an efficiency of 33% to 53%. Compared with *B. subtilis* ATCC 6051a, the final mutant (BS5) forms less foam during fermentation, displays greater resistance to spore formation, and secretes 2.5 times more β-CGTase. Thus, the CRISPR system developed here can be used to modify industrially relevant strains with high efficiency, and mutant BS5 can be applied as a superior expression host.

| Delivery plasmid* | Counter-selectable marker* | Cre/loxP* | CRISPR/Cas9 |
|-------------------|----------------------------|-----------|-------------|
| Plasmid construction (8 days) | Plasmid construction (9 days) | Transformation (1.5 days) | Transformation (1.5 days) |
| Transformation (1.5 days) | Transformation (1.5 days) | PCR fusion of Spc/Zeo and target homology fragment (1 days) | Transformation (1.5 days) |
| Marker deletion (3 days) | Counter-selectable marker integration by single cross-over (3 days) | PCR product transformation (1.5 days) | Verification and plasmid curing (2 days) |
| Verification (1.5 days) | Counter-selectable marker eviction by double cross-over (3 days) | pTSC plasmid transformation and recombination mediated by cre recombinase (1.5 days) | Verification and plasmid curing (2 days) |
| Verification (1.5 days) | Verification (1.5 days) | Verification and plasmid curing (2 days) | Total 12.5 days |
| Total 14 days | Total 16 days | Total 14 days | Total 12.5 days |

Table 1. Time required for *Bacillus subtilis* genome editing by the current existing methods. *The time in the table was calculated in the least time required for each manipulation.*

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at 37 °C with shaking at 200 rpm. To evaluate spore formation, 

purchased from Sinopharm Chemical Reagent Co. Ltd. (Shanghai, China). Non-fat powdered milk was purchased from BBI Life Sciences (Shanghai, China). Other chemicals were 

obtained from Oxoid (Hampshire, UK).

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performed by Shanghai RuiDi Biological Technology Co. Ltd. (Shanghai, China). Tryptone and yeast extract were 

kit were purchased from Tiangen Co. Ltd (Beijing, China). DNA sequencing and DNA primer synthesis were 

Takara (Dalian, China). The plasmid mini-prep kit, PCR purification kit, and the agarose gel DNA purification

for the fermentation in 3 L fermenter (BioFlo 110, New Brunswick Scientific Co., Edison, NJ) that containing

Table 2. Strains and plasmids used in this study.

| Strain or plasmid | characteristics | reference |
|-------------------|-----------------|-----------|
| **Strains**       |                 |           |
| *E. coli* JM109   | recA1, endA1, thi, gyrA96, supE44, hsdR17 Δ (lac-proAB) F [traD36, proAB, lacIq, lacZΔM15] | Takara    |
| *B. subtilis* ATCC 6051a | Wild type | ATCC      |
| BS1               | ATCC 6051a derivative, ΔsrfC | This work |
| BS2               | ATCC 6051a derivative, ΔsrfC, ΔpsolIAC | This work |
| BS3               | ATCC 6051a derivative, ΔsrfC, ΔpsolIAC, Δ nprE | This work |
| BS4               | ATCC 6051a derivative, ΔsrfC, ΔpsolIAC, Δ nprE, ΔapnE | This work |
| BS5               | ATCC 6051a derivative, ΔsrfC, ΔpsolIAC, Δ nprE, ΔapnE, ΔamyE | This work |
| **Plasmids**      |                 |           |
| pMD18-T           | Amp′, MCS       | Takara    |
| pHY300PLK         | Amp′ (E. coli), Tet′ (R. subtilis and E. coli), R. subtilis-E. coli shuttle expression vector | This lab   |
| pHY300PLK-β-CGTase| Amp′ (E. coli), Tet′ (R. subtilis and E. coli), α-amylase promoter (P), β-CGTase gene. | Takara    |
| pwtCas9-bacteria  | Amp′, tetracycline repressor TetR, cas9 gene | Stanley Qi|
| pHYcas9d          | Amp′ (E. coli), Tet′ (R. subtilis and E. coli), R. subtilis-E. coli shuttle expression vector, PE194 temperature-sensitive replicon, sgRNA of srfC | This work |
| pHYcas9dsrf1      | Amp′ (E. coli), Tet′ (R. subtilis and E. coli), R. subtilis-E. coli shuttle expression vector, PE194 temperature-sensitive replicon, sgRNA and repair template of srfC | This work |
| pHYcas9dsrf2      | pHYcas9dsrf1 derivative with repair template for 284 bp deletion of srfC | This work |
| pHYcas9dsrho      | pHYcas9dsrf derivative with sgRNA and repair template of psolIAC | This work |
| pHYcas9dnp       | pHYcas9dsrf derivative with sgRNA and repair template of nprE | This work |
| pHYcas9dapr      | pHYcas9dsrf derivative with sgRNA and repair template of aprE | This work |
| pHYcas9damy      | pHYcas9dsrf derivative with sgRNA and repair template of amyE | This work |

Materials and Methods

Strains and plasmids. All bacterial strains and plasmids used in this study are described in Table 2. *Escherichia coli* JM109 was used for plasmid construction. Plasmid pHY300PLK-β-CGTase was previously con- structed in our laboratory by inserting the β-CGTase gene of *Bacillus circulans* 251 into the *B. subtilis-E. coli* shuttle expression vector pHY300PLK (Takara, Dalian, China)48. Plasmid pwtcas9-bacterial was purchased from Addgene (Addgene plasmid # 44250)49. *B. subtilis* ATCC 6051a was purchased from the American Type Culture Collection (ATCC).

Reagents and enzymes. PrimeSTAR polymerase, restriction enzymes, calf intestinal alkaline phosphatase, *Dpn* I, T4 DNA ligase, In-Fusion HD Cloning Plus kit and vector (pMD18-T) were purchased from Takara (Dalian, China). The plasmid mini-prep kit, PCR purification kit, and the agarose gel DNA purification kit were purchased from Tiangen Co. Ltd (Beijing, China). DNA sequencing and DNA primer synthesis were performed by Shanghai RuDi Biological Technology Co. Ltd. (Shanghai, China). Tryptone and yeast extract were obtained from Oxoid (Hampshire, UK). β-cyclodextrin was purchased from Sigma-Aldrich (Milwaukee, WI, USA). Non-fat powdered milk was purchased from BBI Life Sciences (Shanghai, China). Other chemicals were purchased from Sinopharm Chemical Reagent Co. Ltd. (Shanghai, China).

Media and growth conditions. For routine construction of plasmids and *B. subtilis* mutants, *E. coli*, *B. sub- tilis* and *B. subtilis* derivatives were cultured in LB medium (10 g/L tryptone, 5 g/L yeast extract and 10 g/L NaCl) at 37 °C with shaking at 200 rpm. To evaluate spore formation, *B. subtilis* mutants BS1 and BS2 were cultured at 40 °C with shaking at 200 rpm for 48 h in a medium containing 30 g/L dextrin, 30 g/L bean peptone, 1 g/L CaCl₂ and 0.5 g/L NaCl. Ampicillin (100 mg/L) and tetracycline (20 mg/L) were added as needed.

Fermentation Cultivate condition. Seed culture was obtained by inoculating 100 ul frozen glycerol stock (stored at −80 °C) into 100 ml LB medium and then incubating 12 h at 37 °C with shaking at 200 rpm. As for Shake-flask cultivation, a portion of 5% (v/v) seed culture (2.5 ml) was used to inoculate 50 ml of TB medium (Supplementary material and method), which was then incubated at 30 °C with shaking at 200 rpm for 48 h. While for the fermentation in 3 L fermenter (BioFlo 110, New Brunswick Scientific Co., Edison, NJ) that containing
0.86 L fermentation medium (Supplementary material and method), before a 10% (v/v) seed culture (100 ml)
was inoculated, its pH was adjusted to 7.0 with 20% (v/v) H₃PO₄ and NH₄OH, its temperature was adjusted
to 37 °C, and it was added with 30 ml 166.7 g/L glucose. After 2 h of inoculation, the inducer of 30 ml 333.3 g/L
lactose was added (Supplementary material and method). The feed solutions (Supplementary material and method)
were fed at rate 0 to 15 g glucose/h to confirm the glucose concentration was maintained 0.2 to 0.5 g/L. During
the fermentation process, the batch cultivation was carried out at 37 °C, pH 7.0, and 30% dissolved oxygen that
were maintained by automatically adjusting stirring speed (300 rpm–900 rpm) and flow rate of air (1.5–4.0 L/min).
Tetracycline (20 mg/L) was added as needed every 24 h. The foam height was measured as needed and
antifoam was added manually. For both cultivation, the samples collected at certain time intervals were cen-
trifuged at 12,000 × g for 10 min at 4 °C. For DCW determination, the pellet was resuspended with 0.9% (w/v) NaCl
and centrifuged at 12,000 × g for 10 min, and dried to a constant weight at 105 °C. As for the host strains that harb-
our expression plasmid pHY300PLK-β and centrifuged at 12,000 g for 10 min, and stained the plates with
iodine55. Under these conditions, secreted proteases form a clear ring around the secreting colony.

**Plasmids construction.** The sequences of all of the primers used in this study are listed in Table 3. The
initial CRISPR system knockout plasmid pHYcas9d was assembled from four fragments (One, Two, Three and
Four), each of which overlaps its two adjacent fragments by 15 bp, using the In-Fusion HD Cloning Plus kit50.
Fragment One, which encodes the Cas9 protein, was amplified from plasmid pwtcas9-bacterial using the primer
pair P03/P04. Fragment Two, which includes a P43 promoter for sgRNA expression, a target-specific 20-nt
guide sequence specific for srfC fused with the sgRNA sequence, and a temperature-sensitive replicon PE194 for
plasmid curing, was synthesized and ligated into the cloning vector pMD18-T. This fragment was amplified using
the primer pair P07/P08. Fragment Three includes an E. coli replication origin (p15A ori), an ampicillin-resistance
marker (ampR) and the α-amylase promoter (PamyQ) from B. amyloliquefaciens. This fragment was amplified
from plasmid pHY300PLK-β-CGTase using primers P02 and P05. Fragment Four includes a tetracycline
resistance marker (TcR) and a terminator from B. amyloliquefaciens. This fragment was amplified from plasmid
pHY300PLK-β-CGTase using primers P01 and P06. We amplified the upstream and downstream regions of the tar-
get locus using B. subtilis ATCC 6051A genomic DNA as a template and two primer pairs: P09/P10 and P11/P12.
The amplified fragments were ligated together using overlap extension PCR, forming homologous repair tem-
plate. To form the final knockout plasmid, pHYcas9dsrf1, the appropriate homologous repair template was
inserted into the Xba I site of pHYcas9d (Fig. 1a). During double-strand repair, this repair template removes 6 bp
of native sequence and inserts an Xho I restriction site and 5 bp of random sequence (Fig. 1b).

The other five knockout plasmids (pHYcas9dsr2, pHYcas9dspo, pHYcas9dnpr, pHYcas9damy and
pHYcas9damy) were constructed from pHYcas9dsrf1 by changing the 20 nt guide sequence (except pHYcas9dsr2)
and replacing the homologous repair template. The 20 nt guide sequence was changed using inverse PCR with
the four primer pairs P17/P18 (for pHYcas9dspo), P23/P24 (for pHYcas9dnpr), P29/P30 (for pHYcas9damy), and
P35/36 (for pHYcas9damy), which hanging the new modified 20 nt guide sequence at the 5’ end. The homologous
repair templates were created by overlap extension PCR of sequences upstream and downstream of the 20 nt-guide
sequence. These sequences were amplified from the B. subtilis ATCC 6051A genome using primer pairs P13/P14 and
P15/P16 (for pHYcas9dsr2), P19/P20 and P21/P22 (for pHYcas9dspo), P25/P26 and P27/P28 (for pHYcas9dnpr),
P31/P32 and P33/P34 (for pHYcas9damy), and P37/P38 and P39/P40 (for pHYcas9damy). Then the appropriate
homologous repair template was inserted into the modified pHYcas9dsrf1 by digesting the PCR product and the
modified plasmid with Xba I and ligating the appropriate fragments.

**Genome editing.** B. subtilis competent cells were made by the method of Anagnostopoulos and Spizizen51.
Tetracycline-resistant transformants were confirmed by colony PCR of the cas9 gene. Using transformant genomic
DNA as the template, verification PCR reactions were carried out using specific primers (Supplementary Table S2)
that anneal outside the homologous repair template. PCR products from disruption mutants were identified
by digestion with Xho I, and the homologous repair regions of the mutants were subsequently verified by DNA
sequencing.

**Plasmid curing.** To cure the mutants of the knockout plasmid, edited colonies harbouring knockout plasmid
were used to inoculate 10 mL of LB medium containing tetracycline (20 mg/L). The culture was incubated at
37 °C, and then streaked onto LB plates that were subsequently incubated overnight at 51 °C52. The colonies
cured of knockout plasmid were confirmed by streaking them onto LB plates containing tetracycline (20 mg/L);
colonies cured of plasmid fail to grow at 37 °C. These colonies were used in the next round of genome editing.

**Calculation of sporulation efficiency.** After cultivation in a medium conducive to spore formation at 40 °C
for 48 h, Cells of mutants BS1 and BS2 were diluted and divided into two same amount of portions, respectively,
one of which was heat at 75 °C for 20 min53, then streak them onto LB plates that were subsequently incubated
overnight at 37 °C. The cell ratio of two portions was calculated as sporulation efficiency.

**Detection of protease activity.** To detect the protease activity of mutants BS2, BS3 and BS4, strains were
dropped onto a 5% non-fat powdered milk plate at 37 °C and incubated for 36 h54. Under these conditions, secreted
proteases form a clear ring around the secreting colony.

**Detection of α-amylase activity.** The α-amylase expression of mutants BS4 and BS5 was detected by
dropping the strains onto LB plates containing 1% soluble starch at 37 °C for 24 h, then staining the plates with
iodine55. Under these conditions, secreted α-amylases create a colourless ring around the secreting colony.

**Enzyme assay of β-CGTase.** The β-cyclodextrin-forming activity was determined using the observation
that β-cyclodextrin forms a stable, colourless inclusion complex with phenolphthalein56. A small sample (0.1 ml)
of appropriately diluted culture supernatant was incubated with 2 ml of 1% (w/v) soluble starch in 25 mM phosphate buffer (pH 5.5) at 50 °C for 10 min. The amount of β-cyclodextrin formed was determined by titrating the sample with a standard phenolphthalein solution. One unit of activity was defined as the amount of enzyme that produce 1 μmol of β-cyclodextrin per min.

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Table 3. Primers used in this study. *The restriction enzyme sites are bold and underlined.
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Author Contributions

K.Z. and X.G.D. led the design and performance of the experiments, analysis of the data and writing of the paper. J.W. participated in experimental design, analysis and editing the paper.

Additional Information

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