Efficient genome editing in filamentous fungus

Trichoderma reesei using the CRISPR/Cas9 system

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Filamentous fungi have wide applications in biotechnology. The CRISPR/Cas9 system is a powerful genome-editing method that facilitates genetic alterations of genomes in a variety of organisms. However, a genome-editing approach has not been reported in filamentous fungi. Here, we demonstrated the establishment of a CRISPR/Cas9 system in the filamentous fungus Trichoderma reesei by specific codon optimization and in vitro RNA transcription. It was shown that the CRISPR/Cas9 system was controllable and conditional through inducible Cas9 expression. This system generated site-specific mutations in target genes through efficient homologous recombination, even using short homology arms. This system also provided an applicable and promising approach to targeting multiple genes simultaneously. Our results illustrate that the CRISPR/Cas9 system is a powerful genome-manipulating tool for T. reesei and most likely for other filamentous fungal species, which may accelerate studies on functional genomics and strain improvement in these filamentous fungi.

Keywords: genome-editing; CRISPR/Cas9; filamentous fungus; Trichoderma reesei

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Introduction

Filamentous fungi, such as Trichoderma reesei, Aspergillus niger, A. oryzae, and Penicillium chrysogenum, have been used to produce diverse enzymes or metabolites for centuries [1]. To obtain hyper-producers, repeated classical mutagenesis and screening have been applied for long-term strain improvements. Genetic engineering approaches for filamentous fungi have been well developed. However, these approaches are not as efficient as those available for yeast and bacteria due to the additional complexity of filamentous fungi [2], such as multicellular morphology, cellular differentiation, thick chitinous cell walls, and the lack of suitable plasmids.

Genome-editing is a type of genetic engineering in which DNA is inserted, replaced, or removed from a genome using artificially engineered nucleases, or 'molecular scissors'. It is a useful tool to elucidate the function and effect of a gene or protein in a sequence-specific manner [3]. Recently, developed engineering nucleases, zinc-finger nucleases [4] and transcription activator-like effector nucleases [5], are believed to facilitate precise genome modifications. However, the time- and labor-intensive process of using zinc-finger nucleases and the requirements of specific enzyme engineering for different targets in transcription activator-like effector nucleases limit their applications. The type II clustered regularly interspaced short palindromic repeats (CRISPR)/CRISPR-associated gene (Cas) system is the most popular genome-editing tool at this time. A single multidomain Cas9 catalyzes a double-strand break (DSB) in the target DNA composed of a 20-bp sequence matching the protospacer of the guide RNA (gRNA) and an adjacent downstream 5′-NGG nucleotide sequence (termed as the protospacer-adjacent motif (PAM)) [6]. Over the past 2 years, many studies have demonstrated that the
CRISPR/Cas9 system is a powerful genome-editing method that facilitates genetic alterations in genomes in a variety of organisms. Until now, there have been no reports on the CRISPR/Cas9 system or other genome-editing approaches in filamentous fungi, even in the model organism *Neurospora crassa*, despite the successful application of this technique in yeast [2, 7].

The filamentous fungus *T. reesei*, which is recognized by its Generally Recognized as Safe status by the US Food and Drug Administration, is the most widely used producer of commercial lignocellulolytic enzyme preparations [1] and is expected to be a potential cell factory for different heterologous proteins with its powerful ability to synthesize and secrete proteins with high quantities. However, the genetic basis of desired domestication traits is poorly understood, because robust genetic tools do not exist for this industrial host. The establishment of a genome-editing system can be used to develop *T. reesei* as a super cell factory for lignocellulolytic enzyme preparations and other heterologous proteins, as well as to characterize the regulatory mechanisms for induction, synthesis, and secretion of proteins in *T. reesei*.

In this study, we constructed a CRISPR/Cas9 system in *T. reesei* that allowed us to induce mutagenesis or introduce a new gene by homologous recombination into a target site of the *T. reesei* genome. In addition, it can be used to generate multiple genome modifications simultaneously by co-transforming gRNAs and donor DNAs (dDNAs) for different targets. This system demonstrates great potential as a tool for genome engineering in filamentous fungi, owing to user-designated site specificity of Cas9 endonuclease activity and the simplicity of gRNA construction.

**Results**

*Expression of codon-optimized Cas9 in T. reesei*

Construction of the CRISPR/Cas9 system in an organism requires successful expression of Cas9. Unlike in *Saccharomyces cerevisiae* [7], the codon-optimized Cas9 gene for human cells does not function in *T. reesei* Qm6a or Rut-C30 under control of the constitutive promoter Ppdc (the promoter of *pdc*, the gene encoding pyruvate decarboxylase, which participates in glucose metabolism [8]) or the inducible promoter Pcbh1 (the promoter of *cbh1*, the gene encoding cellobiohydrolase I, which is repressed by glucose but induced by a series of oligosaccharides or cellulose [9]). On the basis of the codon frequency of *T. reesei* [http://www.kazusa.or.jp/codon/cgi-bin/showcodon.cgi?species=51453], we optimized the *Cas9* gene of *Streptococcus pyogenes* and the sequence of the SV40 nuclear localization signal (Supplementary Data S1). To detect the expression and localization of the optimized Cas9, the enhanced green fluorescent protein (eGFP) gene was fused to *toCas9* (optimized *Cas9* gene) and expressed in both the wild-type strain Qm6a and the mutant strain Rut-C30 under control of the Ppdc and Pcbh1 promoter, respectively (Supplementary

![Figure 1](image-url)  
**Figure 1** Expression and localization of Cas9-eGFP in *T. reesei* strains. (a) Detection of Cas9-eGFP expression in transformants 6a-pe and C30-ce by western blots. Transformants 6a-gfp and C30-gfp expressing eGFP were used as controls. Lanes C30-ce and 6a-pe were loaded with 0.05 mg protein. Lanes C30-gfp and 6a-gfp were loaded only with 0.01 mg protein to avoid overexposure. Anti-GFP antibody (Genscript, Nanjing, China) was used for detection. (b) Fluorescence microscopic assessment of the localization of Cas9-eGFP in C30-ce. The bars were 5 μm. DAPI, 4',6-diamidino-2-phenylindole; eGFP, enhanced green fluorescent protein.
Data S1). The expression of toCas9 in each of the selected transformants was detected based on western blots (Figure 1a). toCas9 was expressed in both the transformant 6a-pe (introduced Ppdc-toCas9-eGFP-Tpdc into Qm6a) and the transformant C30-ce (introduced Pcbh1-toCas9-eGFP-Tpdc into Rut-C30).

To determine whether the SV40 nuclear localization signal localizes Cas9 to the nucleus, C30-ce was assessed using fluorescence microscopy (Figure 1b). The mycelia of C30-ce glowed with punctiform green fluorescence after 2 days of growth on the inducing culture media containing 1% Avicel. After staining with 4',6-diamidino-2-phenylindole (1 μg ml⁻¹), the majority of the green fluorescence spots overlapped with nuclei (blue fluorescence). This demonstrated that Cas9 was located mostly in the nucleus of T. reesei, where Cas9 is believed to generate DSBs within the target gene.

To build Cas9-expressing chassis, the Ppdc-toCas9-Tpdc cassette and Pcbh1-toCas9-Tpdc cassette were introduced into Qm6a and Rut-C30, respectively. To choose the Cas9 expression chassis for further investigations, we examined the growth and cellulase activities of the selected transformant 6a-pc from Qm6a and C30-cc from Rut-C30 after three consecutive subcultures for genetic stability, which did not show any differences compared with those of the parent strains (Supplementary Figure S1).

**CRISPR/Cas9 system directed URA5 mutagenesis in T. reesei**

Using appropriately designed gRNAs, the genome can be cut at any desired location by the Cas9 protein. Owing to the lack of confirmed RNA polymerase III-based promoters in T. reesei, such as the U6 promoter in Arabidopsis thaliana [10] and SNR52 promoter in S. cerevisiae [7], we preferred to introduce gRNAs into cells after transcription in vitro [11]. As ura5 has undergone negative selection, its inactivation can be observed in 5-FOA-containing minimal medium (MM) plates (cells expressing wild-type ura5 convert 5-FOA into the toxic substance 5’ fluorouridine monophosphate, which severely limits cell growth). Thus, gRNA was designed to target ura5 and was delivered into Cas9-expressing T. reesei cells through protoplast transformation. To ensure that Cas9 is active in cells, we first chose transformant 6a-pc (a transformant constitutively expressing Cas9) as the Cas9-expressing chassis to examine the mutation frequency. The ura5 genes of 14 selected 5-FOA-resistant transformants were sequenced to confirm whether the mutations were directed by CRISPR/Cas9 (Table 1). Each ura5 gene of the 14 selected 6a-pc transformants showed a frame-shift mutation at the expected target site, proximally located upstream of the PAM sequence (Figure 2a). This was consistent with the fact that CRISPR/Cas9-guided DSBs can be repaired through nonhomologous end joining (NHEJ), which generates insertions and deletions in the vicinity of the cleavage site. By contrast, no colony was observed in 5-FOA-containing MM plates from the control transformation of 6a-pc without gRNA. It confirmed that gRNA might be the essential component to result in mutations.

**Controllability of the CRISPR/Cas9 system in T. reesei**

In industries, the majority of filamentous fungi produce metabolites or enzymes at a specific stage or under unique conditions. A spatiotemporally controllable genome-editing tool precisely controls the metabolic shift in industry bioprocesses. We tested the conditionality of CRISPR/Cas9-guided mutagenesis in the Cas9-expressing chassis, C30-cc (a transformant expressing Cas9 only under inducing conditions) (Supplementary Data S1). As expected, mutagenesis was performed conditionally by an inducer (such as lactose or cellulose) in C30-cc. In the inducible medium (lactose or cellulose as the sole carbon resource), all 15 selected transformants (AR1–AR15) showed mutations in the target gene (Figure 2a). However, in the repressing medium (containing glucose, a repressor for promoter Pcbh1 that controls the Cas9 gene expression), none of the colonies showed mutations. Besides, no colony was observed in 5-FOA-containing MM plates from the control transformation of 6a-pc without gRNA. The above results demonstrated that the CRISPR/Cas9 system functions in T. reesei as an efficient and controllable genome-editing tool in a sequence-specific manner. Besides, the expression of Cas9 under an inducible promoter or other conditionally activated promoters renders the CRISPR/Cas9 system a spatiotemporal-controller of genome-editing of filamentous fungi.

**CRISPR/Cas9 system stimulated homologous recombination in T. reesei**

Similarly to inducing mutagenesis in a sequence-specific manner, the CRISPR/Cas9 system also induces homologous recombination between endogenous and exogenous DNAs. We used poura5 (an exogenous ura5 gene, encoding UR5 in P. oxalicum [12]) as a selectable marker and the endogenous putative methyltransferase lael gene as a target to test the incidence rate of homologous recombination (Figure 2b) by the CRISPR/Cas9 system [13]. The uridine-
| Name             | Sequence (5ʹ to 3ʹ)                          | Experiment                                |
|------------------|----------------------------------------------|-------------------------------------------|
| ToCas9F          | AGCGCAGCTACAGCAATCATGGACAGGAATGAGCAT         | Cloning of toCas9                         |
| ToCas9R          | TTAGACCTTGCGCTTCTTCTGGGTGTCGCGCGGGAGTCG     | Cloning of toCas9                         |
| PpdcF            | ACGAGCGCCAGTGCAAGCTTAAAGACCTCAAGGCTTCTT    | Cloning of pdc promoter                   |
| PpdcR            | GATTGTGCTGATGCTGCGCTGTTTGATCGTTTTGAGTGC    | Cloning of pdc promoter                   |
| pcbh1casF        | ACGACGGCCAGTGCAAAGCTTCTTCTTCTTCTGATCCGCT   | Cloning of cbh1 promoter                  |
| pcbh1casR        | TTCAGGTGCTGCTTCTTCTGCTTCTGCTTCTGCTTCTG    | Cloning of cbh1 promoter                  |
| eGFPF            | GTTAGAGACAGCAGAGGCTGACGTCAGCGCCACCATTG     | Cloning of eGFP                           |
| eGFPR            | TTAGAGACAGCAGAGGCTGACGTCAGCGCCACCATTG     | Cloning of eGFP                           |
| TpdcF            | AGAAGAAGAGAGATTGGTGACCCCGCATGAAGTCTGACCG  | Cloning of pdc terminator                 |
| TpdcR            | TAATTGGCGGAGCTCAGATGGAGCGCCCTCGATGCTTCC   | Cloning of pdc terminator                 |
| ToCGFPF          | GCCGCGACCTCCGCCGCTACGGCTCGCCACATCATGTG     | Overlapping eGFP to toCas9               |
| GFPTcCasR        | GTCGCGGCGGGATCGCAGCC                  | Overlapping eGFP to toCas9               |
| lae1 up F        | TGCAGGTGAGCTGAGATTGAAGGTGAAGCTATG         | Cloning the 5ʹ flanking region of lae1    |
| lae1 up R        | TACTGGCTTAATACTTGCGCTTCCATGAAAAATGT       | Cloning the 5ʹ flanking region of lae1    |
| lae1-0.2 F       | TGCGAGGTGAGCATTTATGTGCTTTACAAATGT         | Cloning the 5ʹ flanking region (0.2 kb)   |
| lae1-0.4 F       | TGCAGGTGAGCTGATTTGCATTATCTGAAAATGT        | Cloning the 5ʹ flanking region (0.4 kb)   |
| lae1-0.6 F       | TGCAGGTGAGCTGATTTGCATTATCTGAAAATGT        | Cloning the 5ʹ flanking region (0.6 kb)   |
| lae1-0.8 F       | TGCAGGTGAGCTGATTTGCATTATCTGAAAATGT        | Cloning the 5ʹ flanking region (0.8 kb)   |
| THR Ura5 F       | GCGCGCATAGTTAACCGCCTAGTACCTCCGCTTACGTACG  | Cloning the 5ʹ flanking region of donor DNA |
| THR Ura5 R       | CAGGGCGCGGCGAGAATTTCATAGTGCAAGCTACGTACG   | Cloning the 5ʹ flanking region of donor DNA |
| lae1 down F      | AAATCCGCTCCAGCCAGCTGCGCACTAAGAGATGACAA    | Cloning the 3ʹ flanking region of lae1    |
| lae1 down R      | GCTCCTCTAGATGCTGCTAGATCCAGATGACAGC        | Cloning the 3ʹ flanking region of lae1    |
| lae1-0.2 R       | GGCCTCTAGATGCTGCTAGATCCAGATGACAGC        | Cloning the 3ʹ flanking region (0.2 kb)   |
| lae1-0.4 R       | GGCCTCTAGATGCTGCTAGATCCAGATGACAGC        | Cloning the 3ʹ flanking region (0.4 kb)   |
| lae1-0.6 R       | GGCCTCTAGATGCTGCTAGATCCAGATGACAGC        | Cloning the 3ʹ flanking region (0.6 kb)   |
| lae1-0.8 R       | GGCCTCTAGATGCTGCTAGATCCAGATGACAGC        | Cloning the 3ʹ flanking region (0.8 kb)   |
| Ulae1 F          | GGAACCCGCCAGCTGACCAC                       | Verifying the 5ʹ flanking region of lae1   |
| HR Ura5 R        | TGGCTCGATGATCCGAGAC                       | Cloning the 3ʹ flanking region of poura5 cassette |
| HR Ura5 F        | AATCGCCACTACTTTCTCAGA                      | Cloning the 5ʹ flanking region of poura5 cassette |
| Dlae1 R          | CAGGGCGCTCTAGTTACCAGC                      | Verifying the 3ʹ flanking region of lae1   |
| Tura5 F          | GCACGGCGCTACCTCAAGGTTGC                    | Sequencing and verifying ura5             |
| Tura5 R          | CAGGGCTCTCTCCGCTTACG                      | Sequencing and verifying ura5             |
dependent transformant AR3 (derived from chassis C30-cc, Figure 2a) was used as a host. After co-transformation of gRNA (gRNA-lae1, targeting lae1 site) with dDNA (dDNA-lae1, containing Pgpda-poura5-Ttrpc cassette and the 3' and 5' flanking regions of lae1) (Table 1), the transformants could be selected in MM plates. The sequence of the selectable marker cassette replaced lae1 in all 14 transformants (Supplementary Figure S2). All Δlae1 transformants showed weakened sporulation (Figure 3a) and cellulytic capability (Figure 3b) compared with the controls, as expected [13]. Compared with other genetic approaches reportedly used for homologous recombination in filamentous fungi [14], the efficiency and precision of the CRISPR/Cas9 system supported its significant advantages.

The effect of the length of homology arms on the recombination frequency was also investigated. The uridine-dependent Cas9 expression chassis AR3 was transformed with gRNA-lae1 and a set of dDNAs with various lengths of homology arms ranging from 0.2 to 1.0 kb at 0.2-kb intervals (dDNA-lae1-0.2k, dDNA-lae1-0.4k, dDNA-lae1-0.6k, and dDNA-lae1-0.8k) (Supplementary Data S1). Nine, 14 or 15 transformants (monoconidial cultures) from each transformation were selected and then checked using diagnostic PCR. These results are summarized in Table 2. The homologous recombination frequencies for all tested homology arms were >93%. These results demonstrated that a pair of 200-bp homology arms is sufficient to achieve efficient homologous integration stimulated by the CRISPR/Cas9 system in T. reesei.
Simultaneous homologous recombination of multigenes using the CRISPR/Cas9 system in *T. reesei*

Disruptions or homologous recombinations of multigenes are required during strain modification or to investigate regulatory systems for a unique metabolic process in filamentous fungi. A genetic approach that could target multiple genes simultaneously would greatly benefit such studies. In the uridine-dependent Cas9 expression chassis AR3, we tested the efficiency of multiple genome modifications directed by the CRISPR/Cas9 system. The selectable marker (Pgpda-poura5-Ttrpc cassette) was used to simultaneously target *lae1* and *vib1* (coding VIB1, a putative link between glucose signaling and carbon catabolite repression [15]). The gRNA and dDNA targeting *vib1* were termed gRNA-*vib1* and dDNA-*vib1*, respectively (containing Pgpda-poura5-Ttrpc cassette and the 3' and 5' flanking regions of *vib1*) (Supplementary Data S1). After co-transformation of the two sets of gRNA and dDNA at the same molar concentration (gRNA-*vib1*:dDNA-*vib1*:gRNA-*lae1*:dDNA-*lae1* = 1:1:1:1), 24 transformants were selected in the uridine-free medium. All 18 positive transformants integrated the selectable marker cassette into the *lae1* locus. However, only three showed homologous recombination in both *lae1* and *vib1* loci. The frequency for double recombination was only 16% (Supplementary Figure S3).

As all the single-deletion transformants were *Δlae1* strains, the molar concentration ratio of the gRNA and dDNA targeting *vib1* to those targeting *lae1* was adjusted to 1.5:1:1:1 (gRNA-*vib1*:dDNA-*vib1*:gRNA-*lae1*:dDNA-*lae1*) in the next co-transformation test. Eleven of the 24 randomly selected transformants were double-deletion strains. The frequency of double recombination was ~45%, and the remaining strains were all single-*lae1* deletion transformants (Supplementary Figure S3).

To test the efficiency of simultaneous homologous recombination of triple genes, *clr2* used as the third target, which is a conserved and essential transcription factor for cellulase gene expression in ascomycete fungi [16]. After co-transformation of the three sets of gRNA and dDNA at the same molar concentration (gRNA-*lae1*:dDNA-*lae1*:gRNA-*vib1*:dDNA-*vib1*:gRNA-*clr2*:dDNA-*clr2* = 1:1:1:1:1:1), 24 transformants were selected in uridine-free medium, in which there was one triple-gene deletion (4.2%), seven *Δlae1Δclr2* double deletions, 13 *Δlae1* single deletions, and two *Δclr2* single deletions (Supplementary Figure S3).

| Homology arm size (kb) | No. of analyzed transformants | No. of deletants | Homologous recombination frequency (%) |
|------------------------|------------------------------|------------------|----------------------------------------|
| 0.2                    | 15                           | 14               | 93                                     |
| 0.4                    | 15                           | 14               | 93                                     |
| 0.6                    | 15                           | 15               | 100                                    |
| 0.8                    | 9                            | 9                | 100                                    |
| 1.0                    | 14                           | 14               | 100                                    |

Table 2: Homologous recombination frequencies at the *lae1* locus using homology arms of various lengths

**Figure 3** Reduction of sporulation and cellulolytic capability in *Δlae1* strains. (a) The observed sporulation of mutants and controls in potato dextrose agar plates after 7 days of growth. Sporulation was reduced in all three randomly selected *Δlae1* strains compared with controls. (b) Assays for filter paper activity of *Δlae1* strains and controls in inducing medium with 1% lactose after 3 days of fermentation. The cellulolytic capability was weakened significantly (*P* < 0.01, *t*-test). Error bars represent the s.d. among three independent measurements.
S4). Although the frequency of triple recombination was quite low, the CRISPR/Cas9 system is an applicable and promising approach to target multiple genes simultaneously.

Discussion

High-efficiency genetic manipulation approaches facilitate fungal strain improvement, as well as the elucidation of fungal molecular mechanisms behind the industrial applications. The first reports of genetic transformation in Aspergillus were in 1983 [17, 18], which lagged behind those for Escherichia coli and S. cerevisiae. Over the past three decades, scientists have attempted to develop various tools or strategies to improve genetic manipulation approaches in filamentous fungi [19–21]. Different genetic transformation approaches have been applied for homologous recombination in many filamentous fungi; however, the low homologous recombination frequency (< 5%) has limited studies on their functional genomics through targeted gene replacement [22]. Deletion of the Ku70 or Ku80 heterodimer (Ku complex) increased the frequency of homologous recombination by up to 65% in many filamentous fungi by eliminating the NHEJ pathway [23]. However, Δku strains were more sensitive to growth environments with specific chemicals such as phleomycin, bleomycin, and methyl/ethyl methanesulfonate. Besides, the strategy to delete the Ku70 or Ku80 heterodimer is not suitable for Agrobacterium-mediated fungal transformation because of its incompatibility in NHEJ-deficient strains [24]. In this study, the CRISPR/Cas9 system showed the highest frequency of single homologous recombination in T. reesei (almost 100%) using a pair of ≥600-bp homology arms (Table 2). Even when the length of the homology arms was decreased to 200 bp, the CRISPR/Cas9 system still reached a 93% homologous recombination frequency (Table 2), higher than the frequency in NHEJ-deficient T. reesei using 500-bp homology arms (63%) [25].

Establishing the CRISPR/Cas9 system in filamentous fungi is based on basic transformation systems. High frequencies of multicopy insertion and ectopic integration have been observed in the application of these filamentous fungal transformation systems [25, 26]. Compared with PEG/CaCl2 protoplast, electroporation and biolistic transformation systems, A. tumefaciens-mediated transformation alleviates variable multicopy insertion and ectopic integration with its high efficiency (> 90%) of single-copy integration in T. reesei [27]. In this study, we introduced the Cas9-encoding gene into the genome of T. reesei using random A. tumefaciens-mediated transformation. To ensure that the introduction and expression of the Cas9-encoding gene does not disturb the expression and function of other genes, the morphology and important phenotypes such as growth and cellulase activities of the selected transformants were compared with their parent strains (Supplementary Figure S1). Although the confirmation process for Cas9 expression typically takes more than 1 month, similarly to previous transformation systems, the Cas9 expression chassis (including the uridine-dependent Cas9 expression chassis) could be used in the following studies for different gene mutations or gene knockout /knock-in by changing the 20-bp protospacer of gRNA to the target locus according to the fungal genomic sequence information. Thus, the CRISPR/Cas9 system in filamentous fungi would save time and labor in the study of fungal life activity regulatory systems, such as reproduction, development, metabolic pathways, protein synthesis, and secretion, which involve multiple unknown functional genes. For example, it took more than a decade to construct genome-wide single-gene deletion libraries for functional genomics research in N. crassa and A. nidulans using traditional genetic manipulation approaches [28, 29]. The CRISPR/Cas9 system is expected to shorten the process for establishing a genome-wide single-gene deletion library for a filamentous fungal species with available genomic sequence information.

Owing to the site specificity of the Cas9 endonuclease and gRNA, the CRISPR/Cas9 system is believed to introduce a genetic alteration or a heterologous gene into the target locus precisely and efficiently. In this study, variations in the target ura5 gene were detected in all selected mutagenesis transformants (Figure 2a), and poura5 was shown to replace the target gene lae1 in all homologous recombination transformants (Supplementary Figure S2). Even in the screened homologous recombination transformants simultaneously targeting triple loci, the heterologous gene poura5 was shown to replace target genes lae1, vib1, and clr2 or at least one or two of these genes (Supplementary Figure S4). Our results confirmed that the CRISPR/Cas9 system was successful in editing the genome of a filamentous fungus precisely and efficiently, similarly to other organisms.

In our simultaneous homologous recombination experiments targeting double genes, almost all of the selected single-deletion transformants were Δlae1 strains (Supplementary Figure S3). We did not obtain a Δvib1 strain, and DSBs were not observed in the gRNA-directed vib1 locus. Similarly, in the
simultaneous homologous recombination experiments targeting triple genes, the selected single-deletion transformants were Δlael or Δclr2 strains, and the selected double deletion was Δlael/Δclr2. No Δvib1, Δlael/Δvib1, or Δlael/Δvib1 strains were detected among the selected transformants (Supplementary Figure S4). The CRISPR/Cas9 system prefers to target the lael or clr2 locus rather than the vib1 locus. The gene vib1 may be located in a region of low transcriptional activity [30], which complicates the process of gRNA directing Cas9 to the vib1 locus. The increase in the molar concentration ratio of gRNA-vib1 (the gRNA targeting vib1) during co-transformation resulted in an increase in frequency for double recombination from 16 to 45% (Supplementary Figure S3). This suggests that the proportion of gRNA and dDNA of multiple genes requires optimization to achieve a high frequency of simultaneous multirecombination.

This study demonstrates precise editing of T. reesei genome through the repair of Cas9-induced DSBs in an NHEJ or homology-recombination method. However, unwanted insertion and deletion mutations were observed at off-target sites, which share high sequence similarity with the on-target site in other mutations were observed at off-target sites, which share high sequence similarity with organisms [31]. In this study, the primary applications of the CRISPR/Cas9 system have been demonstrated as model genome-engineering methods in filamentous fungi, which can accelerate research on the functional genomics of filamentous fungi. Numerous applications of the CRISPR/Cas9 system have been developed in other organisms within the past year. Similarly, more CRISPR/Cas9-mediated systems could be examined in filamentous fungi. For example, a CRISPR/Cas9-mediated large fragment deletion would facilitate genetic modification of filamentous fungal chassis [31]. A catalytically deactivated version of Cas9 can be used for genome-scale CRISPR-mediated control of gene repression and activation [10]. However, until now gRNAs could not be synthesized in vivo easily in T. reesei, which may limit its application fields. It is important to allow gRNA to be synthesized in vivo by identifying effective RNA polymerase-III-based promoters in T. reesei.

Materials and Methods

Strains and plasmids

E. coli DH5α served as the cloning host (Novagen, Gibbstown, NJ, USA). A. tumefaciens AGL1 was used to transform the Cas9 gene to T. reesei strains [39]. The T. reesei strain Qm6a (13631) and Rut-C30 (56765) purchased from American Type Culture Collection (ATCC, Manassas, VA, USA), were used as the hosts for Cas9 expression. A T-DNA binary vector, pDHt/sk, containing the hph coding for hygromycin B phosphotransferase (under control of the A. nidulans trpC promoter and terminator) was used to construct the transformation vectors. pMD-18T vector (Takara, Dalian, China) was used to maintain gRNA templates and dDNAs in DH5α.

Expression of Cas9 in T. reesei

The Cas9 gene with a SV40 nuclear localization signal was codon-optimized for expression in T. reesei (toCas9, Supplementary Data S1). To express toCas9 efficiently, a constitutive promoter Ppdc and an inducible promoter Peh1 were used to express toCas9 in Qm6a and Rut-C30, respectively. The eGFP gene was fused to toCas9 as a tag. All the cassettes with Tpdc as terminator fused into XbaI site of the linearized pDHt/sk using the ClonExpress MultiOne Step Cloning Kit (Vazyme, Nanjing, China) to generate four vectors, pDHt/sk-Ppdc-toCas9-Tpdc (pDHt/sk-PC), pDHt/sk-Ppdc-toCas9-eGFP-Tpdc (pDHt/sk-PE), pDHt/sk-Pebh1-toCas9-Tpdc (pDHt/sk-CC), and pDHt/sk-Pebh1-toCas9-eGFP-Tpdc (pDHt/sk-CE) (Table 1). We also constructed vectors only bearing the eGFP gene (pDHt/sk-Ppdc-eGFP-Tpdc and pDHt/sk-Pebh1-eGFP-Tpdc) as controls. All of the expression vectors were transformed into the recipient T. reesei using Agrobacterium-mediated fungal transformation [40]. Transformants were selected using hygromycin B (10 μg ml−1) and cefotaxime (300 μm) on potato dextrose agar. Corresponding to the expression vector, the transformants derived from Qm6a (with constitutive promoter) were named 6a-pc, 6a-pe, and 6a-gfp and ones from Rut-C30 (with inducible promoter) were named C30-cc, C30-ce, and C30-gfp. Each positive transformant was used to create monoconidial cultures for genetic stability test.

The positive transformants with eGFP tag under Ppdc and Peh1 were grown on MM [40] with 2% glucose and 1% Avicel as carbon resources, respectively. The mycelium was collected by filtration and used to extract intracellular proteins after grinding in liquid nitrogen. Fusion protein production was examined by SDS–polyacrylamide gel electrophoresis and western blots using an anti-GFP antibody (Genscript, Nanjing, China).

Protoplast transformation of gRNA

The gRNA cassette including synthetic gRNA sequence and target DNA of the ura5 gene (5′-GGCGAGGGCCGCAAC ATCGTcgg-3′; PAM is shown in italics) was controlled by T7
promoter (Supplementary Data S1). This cassette was ligated into pMD-18T and maintained in DH5α as template for the following transcription. Before transformation, the cassette was transcribed into RNA in vitro using the MEGAscript T7 Kit (Ambion, Austin, TX, USA). The generated RNA fragments (≥10 μg) were transformed into the Cas9-expressing chassis (6α-laе1 and C30-cc, respectively) using a modified polyethylene glycol-mediated protoplast transformation procedure [41]. To express Cas9 as efficiently as possible, different carbon resources were used in screening MM plates corresponding to the promoters as mentioned in the previous section. Finally, the nonsense mutants of URA5 were selected in these modified MM plates including the 1.5 mg ml⁻¹ 5-FOA (Sangon, Shanghai, China) and 10 mm Uridine (Sangon). The genomic DNAs of selected transformants were extracted using the FastPrep instrument (MP Biomedicals, Santa Ana, CA, USA) with a standard protocol and used for the amplification of the gene ura5. Sequence analysis of the partial ura5 from each transformant was carried out by BioSune (Shanghai, China).

Co-transformation of gRNA with donor DNA

The generated uridine-dependent Cas9 expression chassis (AR3) derived from C30-cc was used as a host to construct an efficient CRISPR-Cas-directed homologous recombination system. Similar to URA5 mutagenesis, the CRISPR-Cas target site (5'-GGATGGCGAATCATGGCGGG-3'; PAM is shown in italics) (Supplementary Data S1) was located in lae1 [13]. The dDNA (dDNA-lae1) containing the 5' and 3' flanking sequences of lae1 and the selectable marker cassette (the ura5 gene from P. oxalicum controlled by the Pgpda promoter and Trrpc terminator) was constructed similarly to dDNA-URA5 (Supplementary Data S1) was generated by overlapping PCR using Q5 (NEB, Beijing, China) and ligated into the pMD-18T vector. The generated vector was propagated in DH5α and purified using the Plasmid Midi Kit (Qiagen, Hilden, Germany). Accompanying the RNA fragments including gRNA and the target sequence of lae1, dDNA was introduced into T. reesei using the protoplast transformation procedure. The transformants were selected using MM plates with 1% lactose as an inducer of Cas9.

Construction of dDNAs with different homology arm sizes

To examine the effect of homology arm length on targeted gene replacements in T. reesei, dDNAs were constructed using homology arms at various sizes. In addition to dDNA-lae1, Pgpda-poura5-Trrpc was used as a selectable marker. Besides the 1-kb flanking sequences (dDNA-lae1), the 0.2-, 0.4-, 0.6-, and 0.8-kb flanking sequences of lae1 were fused to the selectable marker. The generated dDNAs were named dDNA-lae1-0.2k, dDNA-lae1-0.4k, dDNA-lae1-0.6k, and dDNA-lae1-0.8k (Supplementary Data S1) and ligated into the pMD-18T vector. The vector was propagated in DH5α and purified using the Plasmid Midi Kit (Qiagen). The co-transformation procedure was similar to that described above.

Co-transformation of multiple fragments

Multiple genome modification was also performed in the uridine-dependent Cas9-expressing chassis AR3 through co-transformation of two sets of gRNA and dDNA. In addition to the locus of lae1, another target site (5'-GGCTTCTGTCAA GACAACCTggg-3', PAM is shown in italics) (Supplementary Data S1) was located in the gene encoding VIB1, a putative link between glucose signaling and carbon catabolite repression [15]. The dDNA (dDNA-vib1) containing the 5' and 3' flanking sequences of vib1 and the selectable marker cassette (Pgpda-poura5-Trrpc) (Supplementary Data S1) was constructed as similarly to dDNA-lae1. In the triple-loci assay, clr2 (5'-GGA CGAGAGCGAGAGCggg-3', PAM is shown in italics) (Supplementary Data S1) was used as the third target, which was a conserved and essential transcription factor for cellulase gene expression in ascomycete fungi [16]. The dDNA (dDNA-clr2) containing the 5' and 3' flanking sequences of clr2 and the selectable marker cassette (Pgpda-poura5-Trrpc) (Supplementary Data S1) was constructed similarly to dDNA-lae1. After quantitating to the same molar concentration, two or three sets of gRNA and dDNA were introduced into T. reesei using the protoplast transformation procedure. The transformants were also selected using the MM plate with an inducer.

Conflict of Interest

The authors declare no conflict of interest.

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References

1. Punt PJ, van Biezen N, Conesa A, Albers A, Mangnus J, van den Hondo C. Filamentous fungi as cell factories for heterologous protein production. Trends Biotechnol 2002; 20: 200–206.
2. Jiang D, Zhu W, Wang Y, Sun C, Zhang KQ, Yang J. Molecular tools for functional genomics in filamentous fungi: recent advances and new strategies. Biotechnol Adv 2013; 31: 1562–1574.
3. Burgess DJ. Technology: a CRISPR genome-editing tool. Nat Rev Genet 2013; 14: 80.
4. Carroll D. Genome engineering with zinc-finger nucleases. Genetics 2011; 188: 773–782.
5. Cermak T, Doyle EL, Christian M et al. Efficient design and assembly of custom TALEN and other TAL effector-based constructs for DNA targeting. Nucleic Acids Res 2011; 39: e82.
6. Cong L, Ran FA, Cox D et al. Multiplex genome engineering using CRISPR/Cas systems. Science 2013; 339: 819–823.
7. DiCarlo JE, Norville JE, Mali P, Rios X, Aach J, Church GM. Genome engineering in Saccharomyces cerevisiae.
using CRISPR-Cas systems. *Nucleic Acids Res* 2013; 41: 4336–4343.

8 Li J, Wang J, Wang S, Xing M, Yu S, Liu G. Achieving efficient protein expression in *Trichoderma reesei* by using strong constitutive promoters. *Microb Cell Fact* 2012; 11: 84.

9 Ilmén M, Onnella ML, Klembsdal S, Keränen S, Penttilä M. Functional analysis of the cellulobiohydrolase I promoter of the filamentous fungus *Trichoderma reesei*. *Mol Genet Genom* 1996; 253: 303–314.

10 Gilbert LA, Larson MH, Morsut L et al. CRISPR-mediated modular RNA-guided regulation of transcription in eukaryotes. *Cell* 2013; 154: 442–451.

11 Wang Y, Li Z, Xu J et al. The CRISPR/Cas system mediates efficient genome engineering in *Bombyx mori*. *Cell Res* 2013; 23: 1414–1416.

12 Liu G, Zhang L, Wei X et al. Genomic and secretomic analyses reveal unique features of the lignocellulolytic enzyme system of *Penicillium decumbens*. *PLoS One* 2013; 8: e55185.

13 Seiboth B, Karimi RA, Phatale PA et al. The putative protein methyltransferase LAE1 controls cellulase gene expression in *Trichoderma reesei*. *Mol Microbiol* 2012; 84: 1150–1164.

14 Meyer V, Arentshorst M, El-Ghezal A et al. Highly efficient gene targeting in the *Aspergillus niger* kusA mutant. *J Biotechnol* 2007; 128: 770–775.

15 Xiong Y, Sun J, Glass NL. VIB1, a link between glucose signaling and carbon catabolite repression, is essential for plant cell wall degradation by *Neurospora crassa*. *PLoS Genet* 2014; 10: e1004500.

16 Coradetti ST, Craig JP, Xiong Y, Shock T, Tian CG, Glass NL. Conserved and essential transcription factors for cellulase gene expression in ascomycete fungi. *Proc Natl Acad Sci USA* 2012; 109: 7397–7402.

17 Ballance DJ, Buxton FP, Turner G. Transformation of *Aspergillus nidulans* by the orotidine-5'-phosphate decarboxylase gene of *Neurospora crassa*. *Biochem Biophys Res Commun* 1983; 112: 284–289.

18 Tilburn J, Scassacchio C, Taylor GG, Zabicky-Zisman JH, Lockington RA, Davies RW. Transformation by integration in *Aspergillus nidulans*. *Gene* 1983; 26: 205–221.

19 Penttilä M, Nevalainen H, Rättö M, Salminen E, Knowles J. A versatile transformation system for the cellulolytic filamentous fungus *Trichoderma reesei*. *Gene* 1987; 61: 155–164.

20 Herzog RW, Daniell H, Singh NK, Lemke PA. A comparative study on the transformation of *Aspergillus nidulans* by microprojectile bombardment of conidia and a more conventional procedure using protoplasts treated with polyethylene glycol. *Appl Microbiol Biot* 1996; 45: 333–337.

21 Frandsen RJN. A guide to binary vectors and strategies for targeted genome modification in fungi using *Agrobacterium tumefaciens*-mediated transformation. *J Microbiol Methods* 2011; 87: 247–262.

22 Kanaar R, Hoeijmakers JH, van Gent DC. Molecular mechanisms of DNA double strand break repair. *Trends Cell Biol* 1998; 8: 483–489.

23 Weld RJ, Plummer KM, Carpenter MA, Ridgway HJ. Approaches to functional genomics in filamentous fungi. *Cell* 2006; 16: 31–44.

24 van Attikum H, Bundock P, Hooykaas PJ. Non-homologous end-joining proteins are required for *Agrobacterium* T-DNA integration. *EMBO J* 2001; 20: 6550–6558.

25 Zhang G, Hartl L, Schuster A et al. Gene targeting in a nonhomologous end joining deficient *Hypocrea jecorina*. *J Biotechnol* 2009; 139: 146–151.

26 Krappmann S, Sasse C, Braus GH. Gene targeting in *Aspergillus fumigatus* by homologous recombination is facilitated in a nonhomologous end-joining-deficient genetic background. *Eukaryot Cell* 2006; 5: 212–215.

27 Fu K, Liu L, Fan L, Liu T, Chen J. Accumulation of copper in *Trichoderma reesei* transformants, constructed with the modified *Agrobacterium tumefaciens*-mediated transformation technique. *Biotechnol Lett* 2010; 32: 1815–1820.

28 Colot HW, Park G, Turner GE et al. A high-throughput gene knockout procedure for *Neurospora* reveals functions for multiple transcription factors. *Proc Natl Acad Sci USA* 2006; 103: 10352–10357.

29 Park G, Colot HW, Collop PD et al. High-throughput production of gene replacement mutants in *Neurospora crassa*. *Methods Mol Biol* 2011; 722: 179–189.

30 Bi X. Heterochromatin structure: Lessons from the budding yeast. *IUBMB Life* 2014; 66: 657–666.

31 Essletzbichler P, Konopka T, Santoro F et al. Megabase-scale deletion using CRISPR/Cas9 to generate a fully haploid human cell line. *Genome Res* 2014; 24: 2059–2065.

32 Hruscha A, Krawitz P, Rechenberg A et al. Efficient CRISPR/Cas9 genome editing with low off-target effects in zebrafish. *Development* 2013; 140: 4982–4987.

33 Veres A, Gosis BS, Ding QR et al. Low Incidence of off-target mutations in individual CRISPR-Cas9 and TALEN targeted human stem cell clones detected by whole-genome sequencing. *Cell Stem Cell* 2014; 15: 254–254.

34 Ran FA, Hsu PD, Lin CY et al. Double nicking by RNA-guided CRISPR Cas9 for enhanced genome editing specificity. *Cell* 2013; 155: 479–480.

35 Ansai S, Kinoshita M. Targeted mutagenesis using CRISPR/Cas system in medaka. *Biol Open* 2014; 3: 362–371.

36 Fu Y, Sander JD, Reyon D, Cascio VM, Joung JK. Improving CRISPR-Cas nuclear specificity using truncated guide RNAs. *Nat Biotechnol* 2014; 32: 279–284.

37 Miller GL. Use of dinitrosalicylic acid reagent for determination of reducing sugar. *Anal Chem* 1959; 31: 426–428.

38 Grau J, Boch J, Posch S. TALENoffer: genome-wide TALEN off-target prediction. *Bioinformatics* 2013; 29: 2931–2932.

39 Ram AFJ, Michielse CB, Hooykaas PJJ, van den Hondel CAMJJ. *Agrobacterium*-mediated transformation of the filamentous fungus *Aspergillus awamori*. *Nat Protoc* 2008; 3: 1671–1678.

40 Zou G, Shi S, Jiang Y et al. Construction of a cellulase hyper-expression system in *Trichoderma reesei* by...
promoter and enzyme engineering. *Microb Cell Fact* 2012; 11: 21.

41 Liu T, Wang T, Li X, Liu X. Improved heterologous gene expression in *Trichoderma reesei* by cellobiohydrolase I gene (*cbhI*) promoter optimization. *Acta Bioch Bioph Sin* 2008; 40: 158–165.

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