Genome editing using preassembled CRISPR-Cas9 ribonucleoprotein complexes in *Fusarium graminearum*

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Abstract

Genome editing using the clustered regularly interspaced short palindromic repeats (CRISPR)-CRISPR-associated protein 9 (Cas9) system has greatly facilitated the genetic analysis of fungal pathogens. The head blight fungus, *Fusarium graminearum*, causes destructive losses of economically important cereal crops. The recent development of the CRISPR-Cas9 system for use with *F. graminearum* has enabled more efficient genome editing. In this study, we described a CRISPR-Cas9-based genome-editing tool for the direct delivery of preassembled Cas9 ribonucleoproteins (RNPs) into the protoplasts of *F. graminearum*. The use of RNPs significantly increased both the number of transformants and percentage of transformants in which the target gene was successfully replaced with a selectable marker. We showed that a single double-strand DNA break mediated by the Cas9 ribonucleoprotein was sufficient for gene deletion. In addition, short-homology recombination required only 50 base pair regions flanking the target gene. The high efficiency of Cas9 RNPs enables large-scale functional analysis, the identification of essential genes, and gene deletion that is difficult with conventional methods. We expect that our approach will accelerate genetic studies of *F. graminearum*.

Introduction

The clustered regularly interspaced short palindromic repeats (CRISPR)-CRISPR-associated protein 9 (Cas9) system has been developed into a powerful gene editing method for gene insertion, gene knockout, and gene replacement. The RNA-guided Cas9 endonuclease generates a double-strand break at a DNA target when the target is followed by NGG, the protospacer-adjacent motif sequence. The induced double-strand break can be repaired by either the non-homologous end-joining (NHEJ) or homology-directed recombination (HDR) mechanisms [1]. The NHEJ mechanism tends to repair the DNA incorrectly resulting in short nucleotide insertions and deletions (indels) whereas the HDR mechanism is highly accurate and
requires a repair DNA template with a region of homologous sequence [2]. Therefore, the CRISPR-Cas9 system has been widely used to induce or enhance genome mutations in many eukaryotic organisms, including fungi [3].

CRISPR-Cas9-mediated mutagenesis has been widely applied to various filamentous fungal species. In Trichoderma reesei, the codon-optimized Cas9 gene was constitutively expressed to induce random mutagenesis [4]. CRISPR expression plasmids were transformed into cells for genome editing in Neurospora crassa, Aspergillus oryzae, and Fusarium graminearum [5–7]. The acceleration of Cas9 expression using the AMA1-based plasmid increases the efficiency of genome editing in Penicillium chrysogenum [8]. Moreover, an AMA1-bearing plasmid integrated into the genome could be easily removed because of its great instability [9–11]. However, these methods have practical limitations such as labor-intensive molecular cloning, off-target effects, and prolonged effects of the integrated CRISPR construct.

The use of a Cas9-single guide RNA (sgRNA) complex ribonucleoprotein (RNP) has overcome the limitations of a plasmid-mediated CRISPR-Cas9 system. This expression-free method has several advantages compared to conventional transformation methods. The RNP complex is preassembled in vitro; thus, it does not rely on the gene expression mechanism. The RNP is rapidly degraded after genomic DNA cleavage, thereby minimizing off-target cleavage activity and associated toxicity. Recently, RNP transformation has been adapted in various organisms such as plants and animals, including humans [12–14]. With the increasing interest in gene replacement via RNP transformation, there have been several reports of applications to filamentous fungi including Aspergillus niger [15], Fusarium oxysporum [16], Penicillium chrysogenum [17], and Magnaporthe oryzae [18]. However, the potential for easy and practical transformation strategy using Cas9 RNP has not been sufficiently demonstrated.

F. graminearum is an important plant pathogen that causes Fusarium head blight in major cereal crops; it also produces mycotoxins in the infected hosts [19]. In this study, we present genomic modification of F. graminearum, using CRISPR RNP transformation along with a repair construct that contains the desired genome modification. Thus far, homologous gene replacement has been used to generate deletion mutants where transformants can be selected by antibiotic marker genes. Gardiner and Kazan first adapted the CRISPR-Cas9 system in F. graminearum [7]. However, mutation of the target site is rare in the absence of selection. Based on this previous study, we developed an easy and convenient Cas9-mediated transformation system without labor-intensive molecular cloning. Here, we showed that Cas9 RNP comprised commercially available Cas9 protein; the synthesized sgRNA markedly increased the number of transformants and the gene deletion frequency. Moreover, our CRISPR-Cas9 system enabled an efficient homologous recombination-mediated gene replacement with only 50 base pair (bp) homology arms. This method will facilitate genetic analyses of filamentous fungi including F. graminearum.

Materials and methods

Strains and culture conditions

We used the wild-type (WT) F. graminearum strain Z-3639; all mutants were derived from the WT strain in this study. The strains were stored as mycelia in 20% glycerol at -80°C. Media used in this study were described in the Fusarium laboratory manual [19], and all strains were cultured at 25°C.

Protein subcellular localization

Protoplasts and conidia were incubated with Cas9 protein fused to green fluorescent protein (GFP) (Applied Biological Materials, Canada) containing the SV40 T antigen nuclear
localization sequence (NLS). One hundred fifty microliters of protoplast \((5 \times 10^5)\) and conidium \((5 \times 10^5)\) solutions were incubated with 0.5 \(\mu\)L Cas9 protein \((50 \text{ pmol}/\mu\text{L}; \text{Abcam})\) for the translocation assay. Samples were incubated at 4°C for an hour. Microscopic observation was achieved using the DM6 B microscope (Leica Microsystems, Wetzlar, Germany), which was equipped with the Leica DMC6200 camera and used the fluorescent filter L5 (Part No. 11504166).

**sgRNA design and in vitro cleavage assay**

The sgRNAs were selected where the protospacer adjacent motif sequence \((N)_{20}\text{NGG}\) was near the 5’ or the 3’ end of the target site. These sgRNAs were checked by a web-based tool CHOP-CHOP [20], which could evaluate the off-target potential and GC percentages. Suitable sgRNAs were selected randomly among them. Each synthesized sgRNA was purchased from Macrogen (Seoul, Republic of Korea). The TrueCut™ Cas9 Protein v2 (TrueCut™ Cas9 Protein v2; Thermo-Fisher Scientific, Waltham, MA, USA) was used for the *in vitro* cleavage assay. The 1392 bp PCR fragment was amplified by primers FgPks12/5F and FgPks12 clv-R (Table 1), which contains the target site (S1) that induces cleavage into lengths of 1023 bp and 369 bp. All components were added together in a 1.5-mL microtube up to 10 \(\mu\)L \((0.1 \text{ ng Cas9 protein, 160 ng sgRNA, 1x Cas9 nuclease reaction buffer, appropriate PCR fragment concentration, and nuclease-free water})\) and incubated for 30 min at 37°C. Samples were analyzed by gel electrophoresis on a 0.8% agarose gel.

**Fungal transformation**

To generate deletion mutants, the 5’ and 3’ flanking regions of *FgPKS12* and *FGSG_04274*, as well as the geneticin resistance gene cassette (*GEN*), were amplified from the genomic DNA WT strain Z-3639 and plasmid pII99, respectively. The double-joint PCR method [21] was performed to construct the fusion PCR products; the resulting amplicons were used for transformation into the WT strain. The constructed short-homology-mediated transformation donor DNAs were amplified by the primers FGSG_04274 50bp primer R and FGSG_04274 50bp primer F (Table 1). These primers were designed to contain 20 bp of the *GEN* cassette with 50 bp of sequence upstream and downstream of the open reading frame (ORF) of the target gene. The amplified PCR constructs were directly used as donor DNAs for gene replacement with the RNP complexes required in the protocol. For fungal transformation, 150 \(\mu\)L fungal protoplasts \((5 \times 10^5)\) were mixed with 10 \(\mu\)L Cas9 RNP complexes and donor templates for gene integration. After incubation in polyethylene glycol for 15 min, 1 mL STC buffer (1.2 M sorbitol, 10 mM CaCl\(_2\), 10 mM pH 7.5 Tris-HCl) was added to the mixture. The other fungal transformation procedures were performed as previously described [22].

**DNA extraction and Southern blotting**

Genomic DNA was extracted from freeze-dried mycelial powder as described [19]. Standard protocols were used for restriction endonuclease digestion, agarose gel electrophoresis and Southern blotting [23].

**Results and discussion**

**Nuclear localization of the commercial Cas9 protein in *F. graminearum***

The subcellular localization of the commercial Cas9 protein was investigated to determine the optimal Cas9 for *F. graminearum*. Commercial Cas9 proteins generally contain an NLS [24] to enable protein translocation into the nucleus where genome editing occurs. In *F. oxysporum*, a
previous study showed that the SV40 NLS is not functional for the nuclear localization of Cas9; therefore, endogenous histone H2B NLS was utilized for the successful translocation of Cas9 proteins into nuclei [16]. In this study, we assessed whether SV40 NLS is sufficient for the nuclear localization of Cas9 protein in *F. graminearum* (Fig 1). For colocalization analysis, Cas9-eGFP containing SV40 NLS was incubated with protoplasts of the *F. graminearum* hH1-RFP strain, in which histone H1 was fused to red fluorescent protein (RFP) [25]. We found that GFP fluorescence was exclusively present in the nuclei of the *F. graminearum* protoplasts, suggesting that protoplasts can be used for RNP-mediated fungal transformation.

### Table 1. Primers used in this study.

| Name               | Sequence                          | Purpose                                      |
|--------------------|-----------------------------------|----------------------------------------------|
| FgPks12/5F         | CGTTCATGATAACCTCTCTGAGTGG         | HR donor DNA construction for PKS12          |
| pks12-sgRNA1 1kb 5N| GATAAAGGTAAGGTGATCGCTG          |                                              |
| pks12-sgRNA2 1kb 3R| AATCCGTAGCATCAGCCGGAAGG         |                                              |
| pks12-sgRNA1 1kb 3N| GCTTGAAGCTATTTGAGAAGAA          |                                              |
| pks12-sgRNA1 250bp 5F| TGTGCTCAAGCAAGTAAAAGG          |                                              |
| pks12-sgRNA2 250bp 5N| TGAATGCAGCCAGGCGACG             |                                              |
| pks12-sgRNA1 50bp 5F| TAGTTCAACATGACCCCATTCA          |                                              |
| pks12-sgRNA1 50bp 3N| GGAGGTATTCGTTTTTGGGCAC          |                                              |
| pks12-sgRNA2 50bp 3N| GGCAATCTCGTCTCTATCTG           |                                              |
| FgPks12/5R         | GCACAGGTGACACACGGAAGAAGAATTA    |                                              |
| pks12-sgRNA2 3F     | CTTCTCAATCTCAAGCTGAGTGAAGTTGTGGAGGG |                                              |
| FGSG_04274 5F      | TCAATTGAGATGCAGCCTGAAAGAA       |                                              |
| FGSG_04274 3N      | TCGTGGAAAGCTCAGCATACATCT        |                                              |
| FGSG_04274 5R      | GCAAGGTGATACCATTTTTGAGCTG       |                                              |
| FGSG_04274 3R      | AGAAGGTGAGAGGGAGCTACATC         |                                              |
| FGSG_04274 3N      | GCTCAAGTGAGAGGAGCTACAT          |                                              |
| FGSG_04274 3F      | CTCCTCAAGATTCATTTCTCAGTGG       |                                              |
| FGSG_04274 50bp primer R | CTCCTTCTCTGCTGCGGCTTGAATTTAAACACCCTTCAAAGTGG | PCR detection primer                      |
| FGSG_04274 50bp primer F | ATGAGAGGCAGAATCTCATTTCTTCTACAGG |                                              |
| FGSG_04274 with 5F | TCCCTGGCTTTTGGCTGATACATCCTTACC  |                                              |
| GEN with 5F        | GTGCAGATACAGCCACTGCCGTAG         |                                              |
| neo/G2             | GCAAATTCAGGCGTGACCCAAGG         |                                              |
| pIF99/G3           | GGGAGAGGGAGCTGCTCATTTG          |                                              |
| FgPks12/5F         | CGTTCATGATAACCTCTCTGAGTGG       |                                              |
| FgPks12 clv-R      | GCTCAAGTGAGAGGAGCTACAT          |                                              |
| FGSG_04274 5F seq-F | TTGTCGGAGAGCAAAGAAAT            |                                              |
| FGSG_04274 5F seq-R | GCCGCCTGGTGGTCAAGAGG           |                                              |
| FGSG_04274 3F seq-F | CGCTACTGCTACAAATGGGCG          |                                              |
| FGSG_04274 3F seq-R | GAAGTTAAACCTCGGAATGAAG          |                                              |

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In vitro nuclease activity of the preassembled Cas9 RNP

The purpose of our experiment was to validate the efficiency of the Cas9 RNP-mediated gene deletion system in *F. graminearum*. First, we examined the endonuclease activity of the preassembled Cas9 RNP. The *PKS12* gene was targeted because the *pks12* mutations result in a visible albino phenotype, thus simplifying their identification (Fig 2A) [26]. We designed two sgRNAs, which target both ends of the Pks12-coding sequence near the start and stop codons. To predict the possibility of off-targeting, the CHOPCHOP web-based tool for identifying CRISPR/Cas9 off-target sites [20] was used to evaluate mismatches with each genomic site (Table 2). The nuclease activity of the preassembled Cas9 RNP was tested using an *in vitro* cleavage assay. The 1392 bp linear DNA segment included a target cleavage site, and the DNA fragments were designed to be 1023 bp and 369 bp (Fig 2B). The linear DNA was incubated with the preassembled Cas9 RNP comprising Cas9 and the sgRNA S1. The preassembled Cas9 RNP efficiently cleaved the linear DNA into two expected sizes of DNA fragments.

Construction of *PKS12* deletion mutants via RNP-mediated transformation

In *F. graminearum*, transformation for genetic manipulation is generally achieved by the integration of a selectable marker gene via homologous recombination [27, 28]. Here, the deletion construct containing the *GEN* for selection was synthesized by the double-joint PCR strategy [21] with three different homology arm sizes (1 kb, 270 bp, and 50 bp) (Fig 3A). After transformation with or without RNPs, *PKS12* deletion was identified by the albino phenotype because Δ*pks12* mutants could not produce the red pigment aurofusarin (Fig 3B). When deletion constructs with shorter homology arm size were used for fungal transformation without RNPs, both the transformant numbers and deletion efficiencies were markedly...
decreased in *F. graminearum* (Table 3 and Fig 3C). In the absence of the RNP, only 5.5% of the colonies were identified as Δpks12 mutants when the deletion construct had 270 bp homology arms. No deletion mutant was produced when the deletion construct with 50 bp homology arms was transformed without the RNP. In the presence of the RNP, however, the transformant number increased by at least 10-fold, and the deletion efficiency was significantly higher. Notably, the transformant number and deletion efficiency were not dependent on the arm sizes when the RNPs were used. This suggests that the 50 bp homology arm size is sufficient for homology-directed recombination in *F. graminearum*. We examined whether a single RNP with 50 bp homology arms could induce gene deletion (Table 3). The transformation frequencies and deletion efficiencies were different when sgRNA S1 or sgRNA S2 was used for the RNP. Although sgRNA S2 showed a lower yield than did S1, it was sufficient for target gene deletion (Table 3 and Fig 3C).

**Confirmation of short-homology-mediated integration as an efficient gene deletion strategy**

To determine whether Cas9 RNP would allow gene deletion for loci other than *PKS12*, we used the system to delete the uncharacterized gene *FGSG_04274*. The deletion mutants have
an abnormal colony morphology (i.e. defective vegetative growth), thus simplifying their identi-
fication (Fig 4B). Two sgRNAs were designed to target near the start codon (sgRNA S3) and the stop codon (sgRNA S4) (Table 2 and Fig 4A). As expected, transformation without RNP resulted in poor deletion efficiencies, compared to transformation with the RNPs (Table 4).

We designed a gene deletion construct that had 50 bp arms homologous to sequences upstream or downstream from the target gene. This construct can be synthesized with chimeric primers using a simple PCR method, thus minimizing time and financial cost for researchers. The GEN selection marker was amplified with 70 bp primers, which included 50 bp

![Fig 3. Construction of deletion mutants by Cas9 RNP-mediated transformation with different homology arm sizes.](https://doi.org/10.1371/journal.pone.0268855.g003)

**Table 3.** Deletion efficiencies were lower when the homologous arm size of the construct was shorter. Because of variation in the transformation results, this table shows a representative experiment.

| Target gene | Arm size | sgRNA | Total no. of transformants | Expected phenotype frequency | Deletion efficiency (%) |
|-------------|----------|-------|---------------------------|----------------------------|-------------------------|
| PKS12       | 1 kb     | 1 + 2 | 276                       | $5.5 \times 10^{-4}$          | 91.7                    |
| PKS12       | 1 kb     | -     | 28                        | $5.6 \times 10^{-5}$          | 22.9                    |
| PKS12       | 270 bp   | 1 + 2 | 296                       | $5.9 \times 10^{-4}$          | 95.8                    |
| PKS12       | 270 bp   | -     | 18                        | $3.5 \times 10^{-5}$          | 5.5                     |
| PKS12       | 50 bp    | 1 + 2 | 316                       | $6.4 \times 10^{-4}$          | 95.8                    |
| PKS12       | 50 bp    | -     | 4                         | $8.0 \times 10^{-4}$          | 0                       |
| PKS12       | 50 bp    | S1    | 222                       | $4.4 \times 10^{0}$           | 95.8                    |
| PKS12       | 50 bp    | S2    | 44                        | $8.8 \times 10^{0}$           | 63.6                    |

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homology arms to the target gene FGSG_04274 (Table 1). In the absence of the RNP, the transformation produced few colonies but did not show abnormal colony morphology (Table 4). However, short-homology-mediated integration of the donor gene coupled with the RNP markedly increased transformant frequency and deletion efficiency. We also observed that PCR genotyping could successfully detect randomly chosen transformants, which were presumably deletion mutants (S1 Fig). Moreover, deleted regions were confirmed by Southern blot and sequence analysis (S2 Fig).

In this study, we established an efficient gene deletion approach using the Cas9 RNP and a deletion construct with various homology arm sizes. An important benefit of this approach is the ability to use commercially available Cas9 protein and custom-synthesized RNAs. Therefore, no additional laboratory equipment or techniques are required for CRISPR expression; only the deletion construct must be synthesized using PCR. Through our RNP transformation

Table 4. High efficiency of gene deletion via RNP transformation was tested by FGSG_04274 for verification. Deletion mutants were identified by the dark red color and abnormal shapes of the colonies. *50 bp indicates that GEN is amplified by 70 bp primers, which contain 50 bp arms homologous to sequences upstream or downstream from FGSG_04274.

| Target gene | Arm size | sgRNA       | Total no. of transformants | Deletion efficiency (%) |
|-------------|----------|-------------|----------------------------|-------------------------|
| FGSG_04274  | 1 kb     | S3 + S4     | 652                        | 97.2                    |
| FGSG_04274  | 1 kb     | S3          | 56                         | 72.2                    |
| FGSG_04274  | 1 kb     | S4          | 579                        | 97.2                    |
| FGSG_04274  | 1 kb     | -           | 12                         | 58.3                    |
| FGSG_04274  | * 50 bp  | S3 + S4     | 435                        | 93.8                    |
| FGSG_04274  | * 50 bp  | S3          | 74                         | 87.5                    |
| FGSG_04274  | * 50 bp  | S4          | 262                        | 91.7                    |
| FGSG_04274  | * 50 bp  | -           | 7                          | 0                       |

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approach, we expect that challenging genetic modifications can be more effectively applied to filamentous fungi, including *F. graminearum*. The ability to transform protein and RNA into *F. graminearum* implies the potential for transforming other proteins such as enzymes, biosensors, or inhibitory proteins.

**Supporting information**

**S1** Raw images. (PDF)

**S1** Fig. PCR detection strategy for FGSG_04274 and ΔFGSG_04274::GEN. Five abnormal phenotype transformants and a wild-type-like transformant were randomly chosen. The left electrophoresis band indicates the amplified ORF by internal primers and the right band is for detecting deletion mutants that are amplified by primer included in GEN. Each transformants was detected linearly. (TIF)

**S2** Fig. Confirmation of gene replacement. (a) Three ΔFGSG_04274 transformants and a wild-type-like transformant were chosen for Southern blotting. (b) Sequence analysis was performed by Bioneer (Seoul, Republic of Korea) and the marker gene fragments were integrated as designed. (TIF)

**Author Contributions**

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