**Brief Communication**

Robust CRISPR/Cas9 mediated genome editing and its application in manipulating plant height in the first generation of hexaploid Ma bamboo (*Dendrocalamus latiflorus Munro*)

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Bamboo is a special grass to human due to its great economic and ecological values. Around 2.5 billion people are directly producing and consuming bamboo, and its international trade reached 68.8 billion US dollars in 2018 (Data from International Bamboo and Rattan Organization). One major bamboo species in Asia is Ma bamboo (*Dendrocalamus latiflorus Munro*), which is a hexaploid species with three subgenomes 

$2n = 72$, AABBCC (Guo et al., 2019). Despite its agronomic importance, it is nearly impossible to modify bamboo traits by traditional breeding as it takes over 70 years to flower. Bamboo research largely lagged behind due to the lack of efficient genetic manipulation tools. 

The CRISPR (Clustered Regularly Interspaced Short Palindromic Repeat)/Cas9 provides straightforward ways for genome editing in many plants (Yin et al., 2017), but has never been applied in bamboo. Here, we reported the generation of bamboo mutants with CRISPR/Cas9 technology by targeting one specific copy or all homoeologous genes. 

Since our recently established genetic transformation protocol is time-consuming (~1.5 years; Ye et al., 2017), we optimized the CRISPR/Cas9 system in bamboo protoplast. We first improved the protoplast preparation methods and could isolate $3.0 \times 10^6$ protoplasts/g fresh leaves. Next, we improved the PEG-mediated transformation method and reached efficiencies of 53.3% for a single plasmid and 29.8% for two cotransformed plasmids (Figure 1B). Around 1.8% of the protoplasts transformed with the *UBI-Cas9/OsU6b-sgRNA* construct effectively works in bamboo protoplast and was used for the following endogenous gene editing in Ma bamboo. 

The putative phytoene synthase (*PSY1*) in bamboo, whose homolog in maize functions in carotenoid biosynthesis (Zhu et al., 2016), was selected for the initial test. Three bamboo *PSY1* alleles (*DlmPSY1-A, DlmPSY1-B* and *DlmPSY1-C*) were identified and cloned by a homology cloning strategy (Figure 1D). To mutate all copies of *DlmPSY1*, sgRNA1 targeting a conserved site among all *DlmPSY1* loci was designed (Figure 1D). In addition, the sgRNA2 target site containing 2–3 single-nucleotide polymorphisms (SNPs) in the spacer region among three *DlmPSY1* homoeoalleles was selected to test the tolerance of sgRNA mismatches (Figure 1D).

A total of 1600 bamboo calluses induced from stem were transformed as described previously (Ye et al., 2017). In total, 34 independent transgenic lines were confirmed positive (2.1%) by PCR. Based on Sanger sequencing results, 22 (100%) and 10 (83.3%) independent T0 lines were edited in the sgRNA1 and sgRNA2 regions, respectively (Figure 1E), indicating that both constructs effectively induce endogenous gene editing. 

The editing profiles were further analysed by sequencing. Eighteen lines (81.8%) contained putative homozygote/biallelic mutations in all subgenomes at the sgRNA1 target site. In some lines, putative homozygote/biallelic mutations exist in one subgenome, while heterozygote or chimeric mutations appear in other subgenomes (T0-10 and T0-26; Figure 1F). Eight mutation types were identified from 590 independent clones (Figure 1G). The most frequent mutation type was deletion (75%), of which 59.1% are small deletions (<2 bp). The ratios of large fragment deletions ($\geq 14$ bp), insertions and combined indels were 15.9%, 2.21% and 7.82%, respectively (Figure 1G). Since bamboo propagates through asexual budding, these homozygote/biallelic mutations will remain in the genome of their offspring clones during breeding.

sgRNA2 that perfectly targets *DlmPSY1-A1*, but not *DlmPSY1-B1* or *DlmPSY1-C1* was designed to study the recognition specificity (Figure 1D). Sequencing results confirmed that 10
Figure 1 Genome editing in Ma bamboo using CRISPR/Cas9 technology (A) Bamboo protoplast isolation and transformation. a. Microscopic image of isolated bamboo protoplast transformed with 3SS:tdTomato. b-d. Images of bamboo protoplasts co-expressing the fluorescence proteins mGFP and driven by the 35S promoter, and their overlay (d). (B) CRISPR/Cas9 plasmids for bamboo protoplast. Top: CRISPR/Cas9 constructs expressing the isolated bamboo protoplast transformed with

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OsU6a/b/c sgRNA expression construct, mGFP contains one additional guanine (lower-green case) downstream of the translational start site (red); and bottom: GFP expression construct. The sgRNA was designed to produce the

mutation types (right) at the sgRNA1 site of

DlmPSY1. (C) Representative bamboo protoplasts cotransfected with mGFP and OsU6b-sgRNA1/UBI-Cas9 reproducibly emitting fluorescence signals (red arrows). (D) DlmPSY1 gene structure and sequences of the target sites. Grey boxes: exons; black lines: introns; number in brackets: positions of start codon, stop codon and sgRNA target sites (red and orange rectangles). The PAM

osU6b-sgRNA

and

DlmPSY1

mutants at the sgRNA2 site. The represents homozygote (T0-12), biallelic (T0-14) and

mutants. a-c, Wild-type; d-f, dlmpsy1

mutants. (E) Frequencies of CRISPR/Cas9-induced mutations in two target sites of the

DlmPSY1. (F) Representative DlmPSY1 mutations at the sgRNA1 site. TO-1, TO-2, TO-3, TO-12 and TO-13 represent loss of function mutants. TO-10 and TO-26 lines contain heterozygote mutations in the C1 subgenome and chimeric mutations in the A1 subgenome, respectively; middle: mGFP expression construct, mGFP contains one additional guanine (lower-green case) downstream of the translational start site (red); and bottom: GFP expression construct. The sgRNA was designed to produce the presumptive cleavage site at the third nucleotide upstream of the PAM sequence (blue). (C) Representative bamboo protoplasts cotransfected with mGFP and OsU6b-sgRNA1/UBI-Cas9 reproducibly emitting fluorescence signals (red arrows). (D) DlmPSY1 gene structure and sequences of the target sites. Grey boxes: exons; black lines: introns; number in brackets: positions of start codon, stop codon and sgRNA target sites (red and orange rectangles). The PAM regions (blue), SNPs (red) and nucleotide sequences of the

sgRNA target regions (red), PAM regions (blue), nucleotide insertions (green) and their length (right side) are shown.

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transgenic lines contain mutations in DlmPSY1-A1, but none in DlmPSY1-B1 and DlmPSY1-C1 (Figure 1E). Two lines (20%) were putative homozygous or biallelic mutations (T0-12 and T0-14), while 7 lines (70%) were heterozygous/chimeric (T0-30 to T0-32 as representative examples, Figure 1H). The ratios of deletions, insertions and combined mutations were 86%, 9% and 5%, respectively (Figure 1I). The mutations were predominantly short nucleotide changes (1–26 bp), and 22.7% were 1bp nucleotide deletions (Figure 1I). Those data demonstrated the successful application of the CRISPR/Cas9 system in mutating a specific DlmPSY1 allele.

Eighteen lines (81.8%) with homozygote/biallelic mutations in all subgenomes at the sgRNA1 site exhibited albino phenotypes (Figure 1J), which appeared at an early stage during tissue culture and persisted at the plantlets stage (Figure 1J). Those results suggest that genome editing takes place at an early stage in embryonic cells and led to the loss of function of all DlmPSY1 alleles. Similar results were reported in rice, wheat or cotton (Wang et al., 2014; Wang et al., 2018; Zhang et al., 2014). In case of sgRNA2, although DlmPSY1-A was mutated, no visible phenotypic change was observed due to the existence of the wild-type DlmPSY1-B and DlmPSY1-C alleles.

Next, we applied this technology in bamboo molecular research. Bamboo is the tallest grass in the world, while the underlying mechanism is unknown. Previously, we identified several Gibberellin-responsive genes including GRG1 (GA-responsive gene 1, PH01004823G0070) that potentially acts in controlling bamboo height (Zhang et al., 2018). Here, two homozygote grg1 mutants (efficiency 40%) in Ma bamboo were produced using our optimized CRISPR/Cas9 technology. Mutation in GRG1 increased plant height (Figure 1K), mostly due to elongated internodes (Figure 1L-N). Sequencing results confirmed that the grg1 mutant has the putative homozygous mutation in A1 subgenome, biallelic mutation in B1 subgenome and homozygous mutation in C1 subgenome (Figure 1O), indicating the loss of function of GRG1 in transgenic bamboo. To our knowledge, this is the first example on controlling bamboo height through gene manipulation, which will contribute to subsequent studies on the molecular mechanisms behind the fast growth of bamboo.

In summary, for the first time we engineered the hexaploid Ma bamboo through CRISPR/Cas9 technology. The homozygote mutations were obtained in the first generation of transgenic lines, which are extremely important for bamboo species due to its long vegetative growth periods. We also confirmed the albino phenotype of dlmpsy1 mutant in bamboo and generated a bamboo mutant with altered plant height. This demonstrates the applicability of CRISPR/Cas9 in bamboo and thereby boosts future bamboo research and breeding.

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Conflict of interests

The authors declare that they have no conflict of interests.

Authors’ contributions

Q.Z. conceived this project. L.F.G., Y.S.Z., X.Q.M., C.T.L. and Q.Z. designed experiments and interpreted the results. S.W.Y., G.C. and M.V.K. performed the experiments and analysed the data. W.J.W., C.Y.C., C.W. and D.W.S. helped to perform the experiments and collect the data. All authors read and approved the submission of this manuscript.

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