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CRISPR/Cas9-mediated multiple gene editing in *Brassica oleracea* var. *capitata* using the endogenous tRNA-processing system

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Abstract

Cabbage (Brassica oleracea var. capitata) is a biennial plant with strong self-incompatibility and an obligate requirement for prolonged vernalization by exposure to low temperatures to induce flowering. These characteristics significantly increase the difficulty of exploiting novel germplasm induced by physical or chemical mutagens. In this study, we report a CRISPR/Cas9 gene-editing system based on endogenous tRNA processing to induce high efficiency and inheritable mutagenesis in cabbage. Using the phytoene desaturase gene BoPDS, the S-receptor kinase gene BoSRK, and the male-sterility-associated gene BoMS1 as the target genes, multisite and multiple gene mutations were achieved using a construct with tandemly arrayed tRNA-sqRNA architecture to express multiple sqRNAs. The BoSRK3 gene mutation suppressed self-incompatibility completely, converting the self-incompatible line into a self-compatible line. In addition, the BoMS1 gene mutation produced a completely male-sterile mutant, which was highly cross compatible with its nonmutant isoline at the flowering stage as a result of a simultaneous BoSRK3 gene mutation, enabling the economic propagation of the male-sterile line through bee-mediated cross-pollination. Interestingly, higher site mutation efficiency was detected when a guide sequence was inserted into a location in the tandemly arrayed tRNA-sgRNA architecture that was distal from the upstream Pol III promoter. In addition, mutation sites were also detected in the paralogous genes of the BoPDS and BoSRK genes that had fully consistent sequences or base mismatches but beyond the "seed" region in the spacer sequence compared with the target sgRNAs. Collectively, our results demonstrate that the CRISPR/Cas9 system, coupled with an endogenous tRNA-processing system, is an efficient tool to improve cabbage traits.

Introduction

Methods for rapidly and efficiently editing plant genomes are useful for research into gene function and crop improvement. In the past decades, chemical or physical mutagenesis methods were extensively used to produce random mutations that were subsequently screened for the phenotypes of interest¹. Compared with such traditional random mutagenesis, which is inefficient and laborious, targeted gene editing technologies can markedly improve the process of creating mutants. In recent years, methods involving sequence-specific nucleases to create targeted DNA double-stranded break (DSB) genetic tools, such as zinc-finger nucleases and transcription activator-like effector nucleases (TALENs), were used for genome editing^{2–4}. Given the substantial technical complexity and the experience required to design

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and assemble the gene constructs, these methods were not widely used ^{1,3}. Recently, a new gene-editing tool, the type II clustered regularly interspaced short palindromic repeats (CRISPR)-associated protein (Cas) system (CRISPR/Cas9) with its high efficiency, low cost, simple design, and versatility has been demonstrated to be efficient for gene disruption in many plant species, including *Arabidopsis*, *Nicotiana benthamiana*, *N. tabacum*, rice, wheat, maize, sorghum, tomato, potato, sweet orange, poplar, and liverwort ^{1–3,5}.

Many important traits are often regulated by multiple genes or paralogous genes. To improve such traits requires the editing of multiple genes. Fortunately, the Cas9 protein can be directed to specific sites by different single-guide RNAs (sgRNAs) that provide an opportunity to achieve multiplex gene editing by expressing Cas9 along with the multiple sgRNAs^{6,7}. In plants, the most common strategy is to stack multiple independent sgRNA-expressing cassettes in a single construct and to deliver the construct into plant cells^{6,8}. However, this sgRNA-expressing strategy is challenging for most organisms due to the limitations of the delivery method and the vector capacity. By using a synthetic gene with tandemly arrayed tRNA-sgRNA architecture, multiple sgRNAs were efficiently and precisely produced in vivo in rice by the endogenous RNases, RNase P and Z, for the first time⁹. In addition, the tRNA-processing sgRNA expression system was successfully demonstrated in other organisms 10-13.

The CRISPR/Cas9 system is becoming a powerful tool for genome editing in plants^{2,5}, but, in *Brassica*, only a few successful instances of genome editing have been reported^{7,14–16}. One of these cases was the *GA4* gene knockout in a doubled-haploid genotype AG DH1012 (a broccolilike Brassica) from the *Brassica oleracea* var. *alboglabra* (A12DHd) × *B. oleracea* var. *italica* (Green Duke GDDH33) mapping population⁷. Cabbage (*Brassica oleracea* var. *capitata*) is an important cruciferous leafy vegetable crop that is planted worldwide. With the availability of the cabbage genome sequence¹⁷, the CRISPR/Cas9 system may enable the exploration of gene functions and/or the improvement of the traits of cabbage varieties using specific gene modification.

Self-incompatible lines with different S-alleles are usually employed to breed cabbage F_1 hybrids. However, unstable self-incompatibility problems 18 prompt increasing numbers of breeders to breed cabbage hybrids using the male-sterility approach 19 . Seed purity can be guaranteed when producing F_1 hybrid seeds using the malesterile line, but hybrid seed production cost is increased using this route, because both the maintainer line and the male-sterile line have to be propagated manually at the bud stage due to the strong self-incompatibility. The S-receptor kinase (SRK) gene is the key stigma determinant

of cabbage self-incompatibility²⁰. Disrupting self-incompatibility by mutating the SRK gene to generate a self-compatible inbred line should be feasible to reduce the seed production cost of the cabbage F_1 hybrids using the male-sterility approach.

Plant male reproductive development involves stamen meristem specification to achieve pollen grain formation and pollination events; defects in any of these events can lead to male sterility, permitting the generation of valuable male-sterile lines for hybrid breeding²¹. A plant homeodomain-finger family of MALE STERILITY1 (MS1) transcription factors was shown to be involved in the regulation of male sterility by affecting tapetal and pollen wall development in *Arabidopsis*²². In addition, the PERSISTANT TAPETAL CELL1(PTC1) gene in rice, the HvMALE STERILITY1 (HvMS1) gene in barley, and the CA05g06780 gene in pepper, all with similar amino acid sequences to the MS1 gene in Arabidopsis, had been found to be involved in the regulation of male gametogenesis^{23–25}. The *Bol035718* gene (from the *Brassica* database) in cabbage has an amino acid sequence highly similar to that of the MS1 gene in Arabidopsis and is involved in pollen development as verified by RNAi technology in cabbage and genetic complementation analysis in Arabidopsis thaliana (data not shown). We propose that a male-sterile cabbage mutant could be generated using the CRISPR/Cas9 system to modify the BoMS1 gene.

In this study, we synthesized a tandemly arrayed tRNAsgRNA sequence to simultaneously produce numerous sgRNAs using plant endogenous tRNA processing. The phytoene desaturase gene BoPDS, the self-incompatibility determinant gene BoSRK3, and the BoMS1 gene associated with male sterility were selected as the target genes to investigate genome editing in cabbage separately or simultaneously. We showed that the CRISPR/Cas9 system successfully knocked out the target genes using the tRNAprocessing system-based methods. In addition, the cabbage male-sterile line caused by the BoMS1 gene mutation could be economically propagated using the BoSRK gene mutant. Our results demonstrate that the CRISPR/ Cas9 system with endogenous tRNA processing is a powerful tool to improve cabbage varieties using specific gene modification.

Results

Efficient mutation of the *BoPDS* gene via the tRNA-processing system

PDS (phytoene desaturase) is a key enzyme in the carotenoid biosynthesis pathway, and plants with a mutated *PDS* gene should show an albino phenotype that can be easily recognized²⁶. The cabbage phytoene desaturase gene *BoPDS* was selected as the target gene to first explore the genome modification efficiency using the tRNA-

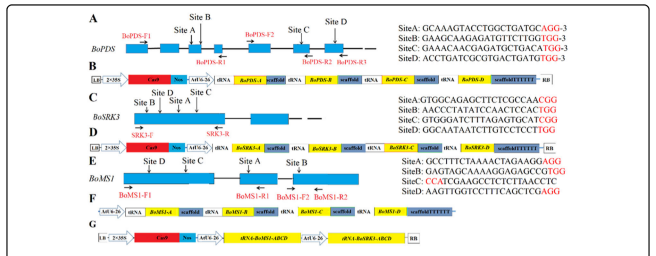


Fig. 1 Vector construct for genome editing of the *BoPDS, BoSRK3* and *BoMS1* genes. **a, c, e** Diagram of the cabbage *BoPDS, BoSRK3*, and *BoMS1* genes, respectively, with the four target sites indicated. **b, d** Diagram illustrating the engineered CRISPR/Cas9 vector with the tRNA-processing system based on multiplex sgRNAs for the *BoPDS* and *BoSRK3* genes. **f** Diagram illustrating the tRNA-sgRNA intermediate vector for the *BoMS1* gene. **g** Diagram illustrating the engineered CRISPR/Cas9 vector for the *BoMS1* and *BoSRK3* gene editing. The blue boxes indicate exons, and the black lines indicate introns. The target sequence is shown in black letters. PAM is marked in red letters. The primer sites used to evaluate the mutation types are shown by black arrows. The screening marker gene expression cassette of *Bar* was not displayed in the T-DNA region (**b, d, g**)

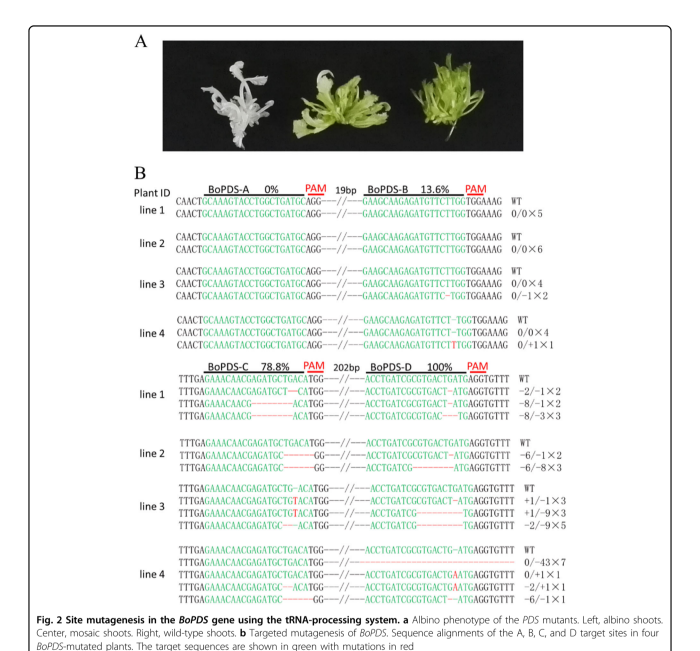
processing system. Four target sites with the protospacer adjacent motif (PAM) (NGG) on their 3'-ends in the *BoPDS* gene were manually selected. The A and B sites were both located in the third exon, while the C and D sites were located in the sixth and seventh exons, respectively (Fig. 1a). The A, B, C, and D target site sequences were inserted into the *BbsI*, *BsaI*, *BsmBI*, and *BfuAI* sites of the synthetic tRNA-sgRNA vector, respectively. The sgRNA-expressing cassette was cloned into the pCACas9 vector as a *BamHI* and *EcoRI* fragment to generate the pCACas-tRNA-sgRNA-PDS-ABCD vector (Fig. 1b).

Twenty-five T0 lines were generated for the pCACastRNA-sgRNA-PDS-ABCD vector transformation. Among them, ten lines exhibited the completely albino phenotype, while seven lines were chimeras, exhibiting a mosaic albino phenotype, resulting in 68% knockout efficiency based on the abnormal phenotypes (Fig. 2a). PCR was performed on three completely albino plants and one chimeric albino plant using the primers BoPDS-F1 and BoPDS-R1 flanking the A and B target sites and BoPDS-F2 and BoPDS-R3 flanking the C and D target sites of the BoPDS gene (Fig. 1a). The PCR products were cloned and sequenced to investigate the mutation types and frequencies for the individual sites. For target site A, no mutagenesis was detected in any of the lines. Heterozygous mutations with a 1 bp deletion or insertion were only detected in lines 3 and 4 for target site B. For target site C, biallelic mutations in lines 1 and 3, mono-allelic homozygous mutations with a 6 bp deletion in line 2, and a chimeric mutation with a 2 or 6 bp deletion in line 4 were detected. For the target site D, biallelic mutations in lines 1, 2, and 3, and a chimeric mutation in line 4 were detected (Fig. 2b). Although four sites in the *BoPDS* gene were simultaneously targeted, no large DNA fragment deletions were detected by further PCR analysis in any of the 17 transgenic plants (Supplementary Fig. 1A). In addition, the mutation efficiencies at the A, B, C, and D sites increased in order from A to D, a phenomenon that was positively correlated with the distance from the upstream *U6* promoter in the guiding sequence position of the tandemly arrayed tRNA-sgRNA architecture (Figs. 1b and 2b).

The BoSRK gene mutation suppressed self-incompatibility completely

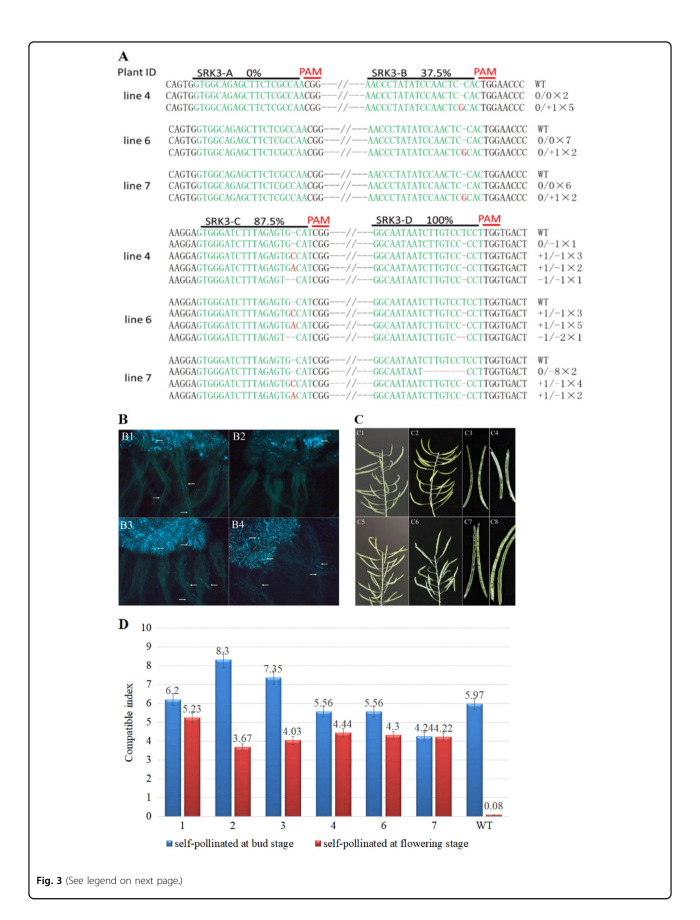
Based on the *BoSRK3* gene sequence of the "F416" inbred line (*SRK3* S-haplotype), four target sites sequence located in the first exon were inserted into the *BbsI*, *BsaII*, *BsmBI*, and *BfuAI* sites of the tRNA-sgRNA vector (Fig. 1c). A pCACas-tRNA-sgRNA-SRK-ABCD vector was generated by inserting the sgRNA-expressing cassette into the pCACas9 vector (Fig. 1d).

A total of 29 Cas9-positive T0 generation pCACastRNA-sgRNA-SRK/ABCD lines were obtained using *Agrobacterium*-mediated transformation. As with the results of the *BoPDS* gene editing, no large DNA fragment deletions occurred among the 29 lines (Supplementary Fig. 1C), but all the lines showed successful gene editing estimated by the direct target region sequencing chromatogram. As with the results from the *BoPDS* gene, we did not detect any mutations at site A of the *BoSRK3* gene



in any of the lines using clone sequencing, but a heterozygous mutation with a G nucleotide insertion was detected at site B in all the lines. Only chimeric mutations were detected at site C in all the lines, but mono-allelic homozygous mutations with a 1 bp deletion in line 4 and biallelic mutations in lines 6 and 7 were discovered at site D (Fig. 3a). In addition, the mutation efficiency at the A, B, C, and D sites in the *BoSRK3* gene was positively associated with the distance between the position of the specific guiding sequence located in the tandemly arrayed tRNA-gRNA architecture and the upstream *U6-26* promoter (Figs. 1d and 3a), the result was similar to that from the *BoPDS* gene modification.

The degree of self-compatibility of the *BoSRK3* gene mutants was estimated by the observation of pollen grain germination on the stigma surface using fluorescence microscopy and a self-compatibility index investigation. A large number of pollen grains germinated on the stigma and formed pollen tubes penetrating through the style of "F416" after self-pollination at the bud stage (Fig. 4b1), but no evident elongated pollen tube growing through the style was observed after self-pollination at the flowering stage (Fig. 4b2). Extended seed siliques and developed seeds of "F416" were observed after self-pollination at the bud stage (Fig. 4c1, c3) but not at the flowering stage (Fig. 4c2, c4) demonstrating that "F416" is a strongly self-



(see figure on previous page)

Fig. 3 Targeted mutagenesis of the BoSRK3 gene (X79432.1) by tRNA-processing system-based editing. a Summary of the mutation types and frequencies in sites A–D. The target sequences are shown in green, and the mutations are shown in red. b In situ fluorescence microscopy of pollen germination on the stigma of the mutant plants and wild-type plants. b1 Bud stage self-pollination of the wild-type plants. b2 Flowering stage self-pollination of the wild-type plants. b3 Bud stage self-pollination of the mutant plants. b4 Flowering stage self-pollination of the mutant plants. White arrows indicate pollen grain germination on the stigma and pollen tube growth down the style. c Silique growth and seed setting in the mutant and wild-type plants. c1 and c3 Silique growth and seed setting of the wild-type plants after the bud stage self-pollination. c2, c4 Silique growth and seed setting of mutant plants after the bud stage self-pollination. c6, c8 Silique growth and seed setting of mutant plants after the flowering stage self-pollination. d Investigation of the self-compatibility index of the mutant and wild-type plants at the bud or flowering stages; self-compatibility index = number of seeds/number of flowers. Bars represent the mean ± standard deviation (SD)

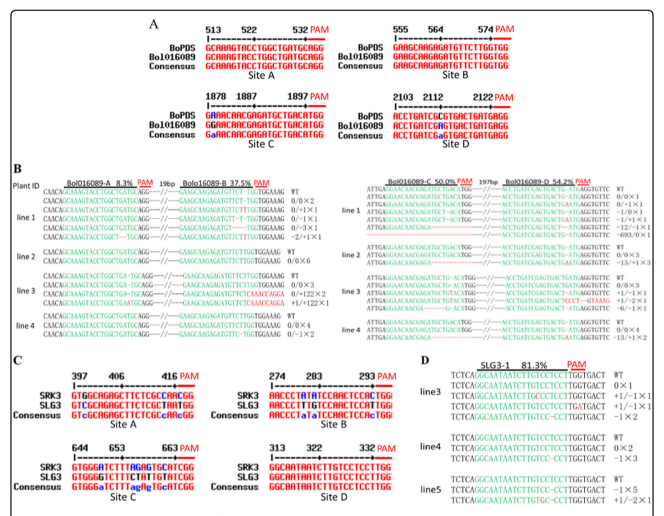


Fig. 4 Paralogous gene mutations mediated by the CRISPR/Cas9 system. a Site A, B, C, and D sequence alignment between the *Bol016089* and *BoPDS* genes. **b** Summary of the mutation types and frequencies at sites A–D in the *Bol016089* gene. **c** Sites A, B, C, and D sequence alignment between the *BoSLG3* gene (X79431.1) and the *BoSLG3* gene. **d** Summary of the mutation types and frequencies at site D in the *BoSLG3* gene. The number of each mutation type was used to estimate the genome editing efficiency. The target sequences are shown in green and the mutations in red

incompatible line. As expected, there were obvious signs of pollen grain germination on the stigma surface and pollen tube growth through the style in the *BoSRK3* gene

mutant at both the bud and flowering stages (Fig. 4b3–b4), while extended seed siliques with full seeds were observed in the *BoSRK3* gene mutant after self-

pollination at the bud and flowering stages (Fig. 4c5–c8). The self-compatibility index of the mutants was similar to or slightly lower than that of "F416" self-pollinated at the bud stage, but the index exhibited by all mutant plants at the flowering stage was significantly higher than that of "F416" self-pollinated at the same stage (Fig. 4d). The results demonstrated that the *BoSRK* gene mutation in "F416" completely suppressed the self-incompatible phenotype.

Mutations were detected in paralogous genes of the target genes

It has been shown that the CRISPR/Cassystem cleaves genomic DNA sequences containing mismatches to the single-guide (sgRNA) strand. Another gene, Bol016089, in the cabbage genome (from Brassica genome database) had a sequence highly similar to that of the BoPDS gene; the nucleotide sequences of the A and B sites in the Bol016089 gene were completely paired with the A and B sites in the BoPDS gene, but there was a base mismatch for both the C and D sites in the Bol016089 gene (Fig. 4a). Although no mutation was detected in the BoPDS gene at site A in any of the lines (Fig. 2b), we detected one cloned mutation at site A of the *Bol016089* gene in lines 1 and 3, resulting in 8.3% editing efficiency (Fig. 4b). In addition, an editing efficiency of 37.5% was found at site B and is higher than the editing efficiency at site B in the BoPDS gene (13.6%) (Figs. 3b and 4b). Because of one base mismatch with the leader sequence in the sgRNA, lower editing efficiencies were detected at sites C (50%) and D (54.2%) in the Bol016089 gene compared with the equivalent editing efficiencies at sites C (78.8%) and D (100%) in the *BoPDS* gene (Figs. 4b and 3b). Although a 693 bp fragment was deleted at the site C region in line 1, and a 122 bp fragment was inserted into the site B region in line 3 (Fig. 4b), no obvious fragment deletions in line 1 or fragment insertions in line 3 were detected using PCR (Supplementary Fig. 1B, lanes 1 and 3).

The nucleotide sequence of the receptor domain on the *SRK* gene showed a high sequence similarity with that of the S-locus glycoprotein (*SLG*) gene from the same haplotype, which is another gene involved in the regulation of self-incompatibility, and both had been proven to be a single copy gene²⁰. Four sites with nucleotide sequences highly similar to those of the *BoSLG3* gene from "F416" were found in the A, B, C, and D target sites of the *BoSRK3* gene. There were 2–6 bp mismatches in the A, B, and C sites in the *BoSLG3* gene compared with the site sequence in the *BoSRK3* gene; only the D site sequence was identical to that of the D target site in the *BoSRK3* gene (Fig. 4c). We did not detect any mutation at the A, B, or C target sites of the *BoSLG3* gene in any of the

transgenic plants. However, mutations were detected in 10 lines of the D site of the *BoSLG3* gene using PCR and clone sequencing (Fig. 4d).

Simultaneous mutation of the *BoMS1* and *BoSRK3* genes to produce a cabbage male-sterile line that can be propagated economically for F₁ variety seed production

The simultaneous mutation of multiple genes in the same cell or individual has many potential applications, such as an important trait that is regulated by multiple functionally related genes. To reduce the cost of seed production of the cabbage male-sterile line for F₁ hybrid seed propagation, it is necessary to mutate the self-incompatibility regulation gene in the cabbage male-sterile line. We assembled two tRNA-sgRNAs that targeted the *BoMS1* and *BoSRK* genes into the same vector to create a male-sterile cabbage mutant and to explore the economic feasibility of producing inbred seed of the cabbage male-sterile line.

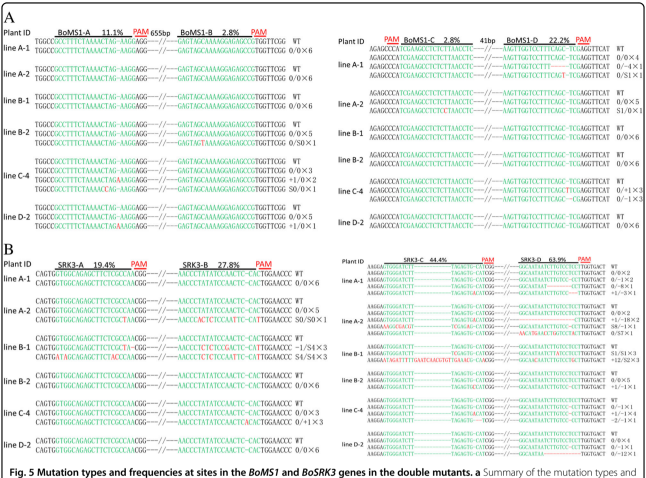
A total of 18 Cas9 gene-positive T0 lines for the PCA-Cas-tRNA-sgRNA-SRK-ABCD/MS1-ABCD vector were identified. The mutation rate was calculated by direct sequencing of the target region of the BoMS1 and BoSRK genes for each line. The BoMS1 gene in six transgenic plants had an overlap peak at the target sites. However, mutations in the BoSRK3 gene were found in 13 transgenic plants. The mutation efficiency of the BoSRK3 gene was clearly higher than that for the BoMS1 gene mutation but markedly lower than the mutation efficiency induced when only the BoSRK3 gene was targeted (100%). The frequency and type of mutation at each target site in the BoMS1 gene in the different plants were analyzed by clonal sequencing. Unexpectedly, we failed to detect any clonal mutations at any of the four target sites of BoMS1 gene in the B-1 line in which the *BoMS1* gene mutation had been detected by direct target region sequencing. We speculated that the overlapping peak of sequencing chromatograms at the target sites of the BoMS1 gene in the B-1 line should result from the purity of the PCR products and the quality of the sequence reads, because no mutation was detected by further Sanger sequencing of the BoMS1 gene in 20 T1 plants derived from the B-1 line (data not shown). Fortunately, the BoSRK3 gene mutations were detected in five other plants that harbored the BoMS1 gene mutation, so the simultaneous mutation efficiency of both genes was 27.78% (Table 1). Nucleotide deletions, insertions, and substitutions were detected at different target sites of the BoMS1 gene in the other five lines. The mutation frequency at the A site over all the plants was 11.11% (Table 1; Fig. 5a), while the lowest mutation frequency (2.78%) was detected at the B and C sites, because only one clone out of 36 was found with a

Table 1 Mutation efficiency in the TO generation lines and in different sites of the *BoMS1* and *BoSRK3* genes

Target	No. of plants	No. of plants	Mutation	No. of plants with Double gene	Double gene Site A mutation efficiency fronteneous (%)	Site A mutation	Site B mutation	Site C mutation	Site D mutation
				mutation	(%)		ובלמבור) (ימ)		
BoMS1	18	2	27.78	5	27.78	11.11	2.78	2.78	22.22
BoSRK3	18	13	72.22	5	27.78	19.44	27.78	44.44	63.89

single base substitution. However, a 22.22% mutation frequency was detected at the D site and was significantly higher than those at the A, B, and C sites (Table 1; Fig. 5a). In lines A-1, A-2, B-2, and D-2, only one mutation was detected as a heterozygous or chimeric mutation. However, mutations were detected at both the A and D sites in the C-4 line, and the D site had a biallelic mutation (Fig. 5a). For the BoSRK3 gene, the mutation frequency was 19.44% with the mutation at the A site in lines A-2 and B-1 occurring as heterozygous and biallelic mutations, respectively. A mutation frequency of 27.78% was detected at the B site in all six lines in the form of biallelic mutations (line B-1), heterozygous mutations (lines A-2 and C-4), and wild type (lines A-1, B-2, and D-2). A higher mutation frequency (44.44%) was detected at the C site than at the A and B sites in the form of biallelic mutations (line B-1 line), heterozygous mutations (lines A-1 and B-2), chimeric mutations (A-2 and C-4 lines) and wild type (line D-2). A mutation frequency of 63.89% at the D site was detected in the form of a chimeric mutation (lines A-1, A-2 and D-2), a heterozygous mutation (line B-2), a biallelic mutation (line B-1) and mono-allelic homozygous mutations (line C-4) (Table 1; Fig. 5b). In addition, more than one nucleotide substitution was detected at different sites of the A-2 and B-1 lines, which were clearly different from the single nucleotide substitution in the A-1, B-2, and C-4 lines (Fig. 5b). Among the four target sites of the BoSRK3 gene, the mutation frequency increased in order from site A to site D (Table 1; Fig. 5b), with the mutation frequencies at the different sites similar to those from the individual BoSRK3 gene modifications (Fig. 3a).

To plants with simultaneous mutations in both the BoMS1 and BoSRK genes were transferred to soil and grown to anthesis (the A-1 line died after being planted). Plants of the other five lines proved to not be phenotypically different from the wild-type plants at the vegetative stage. After flowering, we found that only the C-4 line exhibited complete male sterility; the filaments becoming shorter, and the anthers failed to produce viable pollen, but the style of line C-4 was indistinguishable from that of the wild type (Fig. 6a, b: a1-a3 show the wild type; b1-b3 show the C-4 mutant). The phenotype of pollen development was examined using light microscopy. Unlike the situation in the wild type, the microspores and the tapetum of the C-4 mutant degenerated to form a mass of undifferentiated cells; ultimately, the anther locule was completely empty with no pollen development (Figs. 6a4 and b4). The cross-compatibility at the flowering stage between the C-4 plant and the wild type plant was determined by examining different traits, including pollen germination and the cross-compatibility index. A mono-allelic homozygous mutation of the BoSRK3 gene in the C-4 plant was detected (Fig. 5b). As expected, a large number of pollen grains that germinated on the



frequencies at sites in the BoMS1 and BoSKK3 genes in the double mutants, a Summary of the mutation types and frequencies at sites A–D in the BoSRK3 gene. **b** Summary of the mutation types and frequencies of sites A–D in the BoSRK3 gene. The target sequences are shown in green and the mutations in red

stigma and bundles of the pollen tubes were observed in the style of the C-4 plant pollinated by pollen grains from the wild type plant at the flowering stage, a result that was not obviously different from the situation with pollination at the bud stage (Fig. 6c1, c2). We found that the C-4 plant had a markedly elongated silique with a normal amount of seeds following pollination by wild type pollen at the flowering stage or at the bud stage (Fig. 6c3-c8). The cross-compatibility index between the C-4 malesterile plant and the wild type maintainer line reached 5.56, indicating that the *BoMS1* gene mutation only affected the male gamete development with no negative effect on the female gamete (Fig. 6d). The results demonstrated that simultaneous BoSRK3 gene mutations could release the cross-incompatibility obstacle between the cabbage male sterile and maintainer lines at the flowering stage. The self-compatibility index at the flowering stage of the other four fertile lines with the *BoSRK3* gene mutation, specifically lines A-2 (2.28), B-1 (7.64), B-2 (12.51), and D-2 (14.29), was much higher than for the wild type plant (0.07), although considerable variability was observed among the four plants (Fig. 6d). Even without the intercession of bees or artificial pollination in a space shielded with a fly net, some male-fertile plants showed very high seed-setting ability (Supplementary Fig. 2).

Transmission of Cas9-induced mutations from the T0 to the T1 and F1 generations

To determine whether the Cas9-induced mutations were transmitted to the next generation, 20 T1 plants derived from a T0 mutant plant (line B-1) with only the *BoSRK3* gene mutation and 20 F1 plants derived from the cross between C-4 line (*BoMS1* and *BoSRK3* gene mutation) and the wild type were genotyped using PCR and amplicon Sanger sequencing. Line B-1 was a biallelic mutant at four target sites by deletion, substitution, and insertion in the *BoSRK3* gene (Fig. 5b). The individual plant sequencing results confirmed that these mutations were inherited by plants of the T1 generation in

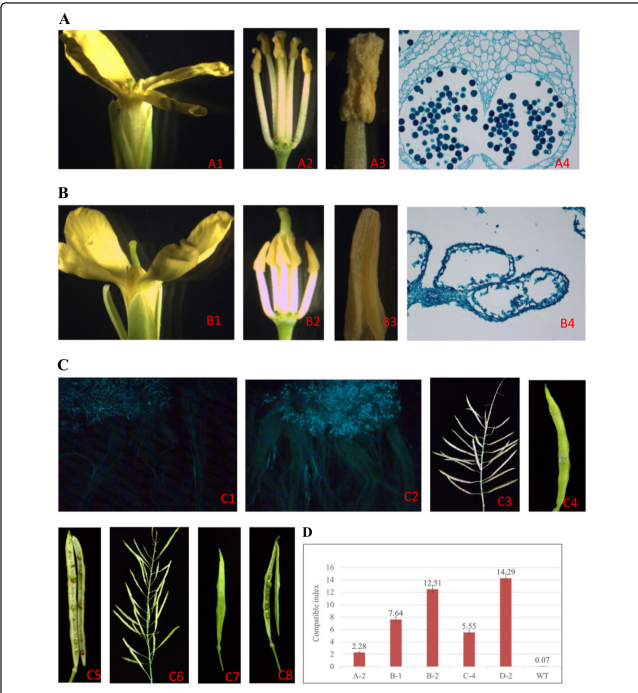


Fig. 6 Floral organ morphology, development of the pollen and cross-compatibility of the wild type and the C-4 mutant. a, b Comparison of the lower, stamen, pistil, anther, and pollen development between the wild type and the C-4 mutant. a1–a3 floral organ morphology of the wild type; b1–b3 floral organ morphology of the C-4 mutant; a4 pollen formation in the anther of the wild type (10×20); b4 empty pollen sac in the anther from the C-4 mutant (10×20). c cross-compatibility between the C-4 mutant and the wild type with the same S-allele. c1 in situ fluorescence microscopy of pollen (from the wild type) germination on the stigma of the C-4 mutant at the bud stage; c2 in situ fluorescence microscopy of pollen (from the wild type) germination on the stigma of the C-4 mutant at the flowering stage; c3–c5 silique growth and seed-set of the C-4 mutant × wild type at the bud stage; c6-c8 silique growth and seed set of the C-4 mutant plant × wild type at the flowering stage. d Self- or cross-compatibility indices of the BoMS1 and BoSRK3 gene mutant plants. Bars represent the mean \pm standard deviation (SD)

homozygotes and heterozygotes (Supplementary Fig. 3-A, A1–A2). In addition, a few new mutations were also detected in the T1 generation of the B-1 line. According to the overlapping sequencing chromatograms, new mutations at the site A were detected in 11 plants that were different from the mutations observed in the parental genome (Supplementary Fig. 3-A, A3). The similar phenomena were also found in *B. napus*¹⁶.

The T0 C-4 line could not be self-pollinated to generate T1 generation plants. Thus, it was crossed with "F416" to produce the F1 generation. The T0 C-4 line had biallelic mutations at site D due to a "T" nucleotide deletion and a "T" nucleotide insertion (Fig. 5a). The sequencing results confirmed that the two types of mutations were inherited. With the exception of one plant, the sequencing chromatograms in the 19 F1 plants had chaotic peaks at the target site (Supplementary Fig. 3-B, B1-B2), indicating that there were hemizygous mutations in the *BoMS1* gene at the target site in the 19 plants. There were 16 plants that had a "T" nucleotide deletion; 3 plants had a "T" nucleotide insertion at site D of the BoMS1 gene, and no mutation was detected at site D in one plant (Supplementary Table 2). In addition, a heterogeneous "A" nucleotide insertion was simultaneously detected at site B of the BoSRK3 gene in the T0 C-4 line (Fig. 5b), and we found that 11 plants had an "A" nucleotide insertion at site B in the BoSRK3 gene after sequencing the amplicons (Supplementary Fig. 3-B, B4; Table 2). In total, 11 F1 plants of $C-4 \times WT$ inherited both the mutations of the BoMS1 gene (site D) and the BoSRK3 gene (site B).

The presence of the transgene (T-DNA) region was also examined using Cas9 gene PCR in the T1 and F1 populations. The absence of the T-DNA region was determined to be parallel in the PCRs negative for the *Cas9*, *sgRNA*, and *Bar* genes. The results indicated that four transgene-negative T1 progeny of the line B-1 with the *BoSRK3* gene mutation had been segregated out. In addition, five F1 plants of the C-4 × WT were segregated out with the *BoMS1* gene and *BoSRK3* gene mutation but were transgene-free (Supplementary Table 2).

Discussion

Cabbage is characterized by a typical sporophytic self-incompatibility²⁰, while flowering induction also has an obligate requirement for exposure to a prolonged period at low temperatures (vernalization)²⁷. Because of the strong self-incompatibility, artificial self-pollination (at the bud stage) has to be performed to reveal the mutant phenotype after treatment with physical or chemical mutagens. This essential process is very laborious and limits the size of the population that the breeder can handle. Genome-editing tools have become well developed in recent years, providing a breakthrough in crop improvement. TALENs have been used to successfully induce site-specific mutations in the

FRIGIDA gene in cabbage in our laboratory⁴. However, as a species recalcitrant to genetic engineering transformation²⁸ until now, the gene editing of cabbage using the CRISPR/ Cas9 system has not been published, except for a report on the internet about a meal, which comprised CRISPR/Cas9edited cabbage (probably a cabbage mustard, judging from the published picture) at Umea University, Sweden. In this study, the BoPDS gene, the self-incompatibility gene BoSRK3, the BoMS1 gene associated with male sterility, and some of their paralogous genes were successfully knocked out using an array of sgRNA-tRNA units designed to express more than one sgRNA and showed obvious mutant phenotypes in the T0 generation. In addition, the CRISPR/ Cas9 system can target multiple sites or multiple genes in a single transformation event and produce homozygous knockouts, even in the T0 generation, as reported in other species^{7–10,13,14}

The Cas9 protein requires only a PAM (NGG) adjacent to the sgRNA homology region to result in efficient Cas9 binding and DSBs, but there are many possible PAM sites in a gene, and variable gene modification rates (from 0% to 100%) associated with different sites have been frequently reported 12,26,29,30. Many online tools and resources for computer-based sgRNA design for high activity site prediction have been developed³¹. However, a low relevance has frequently been found between the sgRNA efficiency predicted bioinformatically and that assessed using protoplast or leaf transient expression assays 12,32. In this study, we manually selected sgRNA target sequences based on the requirements of the Cas9 system and expressed multiple sgRNAs from a single tRNA-sgRNA transcript array to target different sites or genes. Of the T0 plants, 68% were found to contain a BoPDS gene mutation based on an abnormal phenotype. On the basis of PCR fragment direct sequencing, 100% of the T0 plants were shown to harbor a BoSRK3 gene mutation, and 27.78% of the T0 plants were shown to harbor BoMS1 and BoSRK3 gene mutations simultaneously when both genes were targeted. However, a few invalid sites (with no mutation detected) were found in the BoPDS and BoSRK3 genes (Figs. 2b and 3a) and are why more than one sgRNA is frequently assembled to simultaneously target multiple sites of a gene to guarantee successful gene modification^{9,10,15,26,29,30}.

Plant genomes typically exhibit polyploidy and have redundant genes and extensive gene family networks; therefore, multiple sgRNA designs are required for successful trait modification ^{13,14}. The most common way to edit multiple sites or multiple genes is to combine multiple sgRNAs, each under the control of its own promoter, into a single construct ^{6,8,29}. Recently, an endogenous tRNA-processing system to edit multiple genes was accomplished in rice, maize, wheat, cotton, *Arabidopsis*, *Dictyostelium*, and *Drosophila* ^{9-13,33,34}. In addition, the

tRNA^{gly} originating from plants can efficiently process the tRNA-sgRNA transcript in mammalian cells, suggesting that the processing of tRNAs is highly conserved in different organisms¹¹. In *Drosophila* and wheat, high site mutation efficiency could be obtained when the guiding sequence was inserted at a location that was proximal to the upstream Pol III promoter in the tandemly arrayed tRNA-sgRNA architecture^{11,12}. However, a comparable mutation frequency in rice was also detected even when the guiding sequences were inserted at different locations in the tRNA-sgRNA architecture⁹. In this study, higher site mutation efficiency was obtained when the guiding sequence was inserted at a location that was distal to the upstream Pol III promoter (Figs. 2b, 3a, 4b and 5b), a finding that was similar to that reported in cotton¹³. In addition, similar results were also found for tobacco PDS and Sulfur (Sul) gene modification (unpublished results).

Recently, a report described high-frequency off-target mutagenesis induced by CRISPR/Cas9 in Arabidopsis³⁵, a result that was in contrast to two earlier reports in Arabidopsis^{30,36}. This inconsistency could be because too few mutants sample were performed off-target analysis in the previous studies, and different promoters were used to drive Cas9 expression, since an increased dose of the Cas9/sgRNA complexes significantly enhances the frequency of off-target mutations³⁵. In this study, we did not analyze the potential off-target loci following the PAM sequences within the cabbage genome, but those sites in the paralogous genes Bol016089 and BoSLG3, which were highly homologous to the target sites in the BoPDS and BoSRK3 genes, respectively, were examined. The sgRNAs can accommodate mismatches varying by as many as five nucleotides in the 5' upstream region, but the PAMproximal "seed" region comprising 10–12 bp is crucial to the Cas9 cleavage activity, and mismatches in this region usually lead to a decrease in or even complete abolition of the target cleavage activity³⁷. In Brassica napus, no mutations were detected in the putative off-target sites as a result of one to three mismatches, which were found in the "seed sequence" of potential off-target sites 15. In this study, there were one to four mismatches in the "seed sequence" for the A, B, and C putative sites of the BoSLG3 gene, and, as expected, no mutations were detected in the putative off-target sites (Fig. 4c). However, mutations were detected at the putative C and D sites of the Bol016089 gene with one mismatch, but it was out of the "seed sequence" of the guide sequence (Fig. 4a, b). The frequency of off-target mutations induced by CRISPR/ Cas9 is typically well below that caused by chemical and physical mutagenesis¹. In addition, unlike CRISPR-based gene therapy in humans, any off-target modification in plants can be removed by segregation, if necessary.

Male sterility can guarantee high-purity F₁ hybrid cabbage seeds, which have recently become economically

significant, and male-sterile parents are used for cabbage hybrid breeding by plant breeders in some countries¹⁹. However, labor costs for F₁ hybrid seed production based on the male-sterile approach cannot be reduced or may even be increased due to the self-incompatibility in the maintainer line, the male-sterile line, and the paternal line. SRK is one member of the S-locus gene family involved in the regulation of cabbage self-incompatibility. The mutation of the SRK gene resulted in a selfcompatible line in B. oleracea and Brassica rapa^{38,39}. In this study, knockout of the BoSRK3 gene completely abolished the self-incompatibility of the cabbage selfincompatible line "F416", the self-compatibility index of the BoSRK3 gene mutants at the flowering stage were clearly higher than in the wild type "F416"; the mutants had become completely self-compatible lines (Fig. 3). Thus, direct modification of the SRK gene in the malesterile parent, the maintainer line and the paternal line using the CRISPR/Cas9 system, to reduce the costs of cabbage hybrid seed production, is obviously a more efficient way than the backcrossing strategy using rare natural self-compatible mutants.

Currently, ogura cytoplasmic male sterility (CMS) from radish (*Raphanus sativus*) accounts for the vast majority of *B. oleracea* male-sterility types^{40,41}. It is well known that the prevalence of a single cytoplasm in commercial crop cultivars has potential risks due to the associated low genetic diversity; the Texas cytoplasm, which was widely used as a source of CMS to produce maize hybrid cultivars, resulted in maize southern corn leaf blight and yellow leaf blight epidemics in 1970–1971 in the USA⁴². Therefore, to reduce the risk of a single cytoplasm in cabbage hybrid breeding programs, the male-sterile cabbage genetic resources can be expanded by mutating the nuclear genes regulating microsporogenesis, stamen development, or microgametogenesis.

In this study, a biallelic mutation at the D site of the BoMS1 gene in the C-4 plant induced complete male sterility with no pollen grains produced in the anther locules of the sterile mutant (Fig. 6b), and with the phenotype similar to the mutants of the MS1 gene in Arabidopsis, rice, and pepper^{22,23,25}. The ms1 mutant exhibited no defect with respect to the vegetative phenotype and flower organs with the exception of the anther in *Arabidopsis*, rice and pepper^{22,23,25}. As expected, the C-4 mutant of the BoMS1 gene exhibited complete male sterility without any morphological defects. However, male sterility controlled by recessive nuclear genes is difficult to use due to problems encountered in its maintenance, and 100% male sterility was not obtained. In 2002, strategies to obtain full maintainer lines for the male-sterile parents controlled by recessive nuclear genes were proposed by Perez-Prat and van Lookeren Campagne⁴³, and commercial applications of the recessive

male-sterile lines have been achieved in maize and rice based on these strategies 44,45 . The C-4 male-sterile mutant is highly cross compatible with the maintainer line (wild type with same S-allele) (Fig. 6c). The results indicate that the propagation of the cabbage male-sterile line is economically feasible by the simultaneous mutation of the BoSRK gene and the male-sterility gene, indicating that an F_1 hybrid cabbage seed production system from the BoMSI male-sterile line could be constructed in the near future using the same strategy as in maize and rice 44,45 .

Conclusions

Taken together, our results demonstrated that the tRNA-processing system is capable of targeting multiple sites of a gene or multiple genes in a single transformation event and can produce homozygous or biallelic mutations at multiple loci in the T0 generation that are inherited by the next generation. In addition, we have also produced simultaneous mutations of the allene oxide synthase gene (BoAOS) (a gene involved in jasmonic acid biosynthesis) and the BoSRK3 gene to develop a jasmonic acid-regulated cabbage male-sterile line using the tRNA-processing system (unpublished results). Therefore, we conclude that the multisgRNA-expression-based tRNA-processing system provides a powerful tool to study gene function and to achieve trait stacking in cabbage.

Materials and methods

Vector construction

According to the principle of the endogenous tRNAprocessing system reported by Xie et al., a polycistronic tRNA-sgRNA-expressing cassette sequence was synthesized by the Shanghai Xuguan Biotechnology Co., Ltd, Shanghai, China. A "G" nucleotide was inserted between the Arabidopsis U6-26 promoter and the first glycinetRNA sequence of Arabidopsis to meet the transcription initiation requirement of the U6 promoter. In addition, the sgRNA structure was modified by extending the duplex length by ~5 bp and mutating the fourth thymine of the continuous sequence of thymines to cytosine, modification since this was reported significantly improve the knockout efficiency in cells⁴⁶. Four tRNA-sgRNA units assembled into the sequence. BbsI, BsaI, BsmBI, and BfuAI sites were inserted between the glycine-tRNA and sgRNA scaffold sequences to facilitate the target sequence cloning (Supplementary Figure 1). Complementary oligos of targeted sites were ligated into the BbsI, BsaI, BsmBI, and BfuAI sites of the tRNA-sgRNA-expressing cassette as described at http://www.genome-engineering.org/ crispr/. All of the pairs of partially complementary oligos for the target site with 4 nt overhangs are listed in Supplementary Table 1.

The Cas9 coding sequence in the pYLCRISPR/ Cas9P35S-B vector⁸ as a NcoI and BamHI fragment was cloned into the NcoI and BglII sites of the pCA13Bar-35S binary vector (with a *Bar* marker gene, developed by our laboratory) to generate the pCACas9 vector. Four target sites were selected for each gene to generate U6-26:: tRNA-sgRNA-PDS-ABCD (BoPDS gene) and U6-26:: tRNA-sgRNA-SRK-ABCD (BoSRK gene)-expressing cassettes with a BamHI site at the 5' and an EcoRI site at the 3' ends. However, the U6-26::tRNA-sgRNA-MS1-ABCD (BoMS1 gene) had an XbaI site at the 5' and a BamHI site at the 3' ends. The U6-26::tRNA-sgRNA-PDS-ABCD- and U6-26::tRNA-sgRNA-SRK-ABCD-expressing were ligated into the BamHI and EcoRI sites in pCACas9 to obtain the pCACas-tRNA-sgRNA-PDS-ABCD and PCACas-tRNA-sgRNA-SRK-ABCD vectors, respectively. For simultaneous double gene editing of the BoSRK and BoMS1 genes, the sgRNA-expressing cassettes for the modified BoMS1 gene were ligated into the XbaI and BamHI sites of the PCACas-tRNA-sgRNA-SRK-ABCD vector to generate the PCACas-tRNA-sgRNA-SRK-ABCD/MS1-ABCD vector.

Cabbage transformation

The self-incompatible line "F416" (SRK3 haplotype) was used in this study (from stocks developed in our laboratory). A modified version of the transformation method described by Bhalla and Singh⁴⁷ was used. Briefly, hypocotyls from 7- to 10-day-old seedlings were precultured on callus initiation media (MS, 0.05 mg/L 1-naphthaleneacetic acid (NAA), 3 mg/L 6-benzylaminopurine (6-BAP), 30 g/L sucrose and 6 g/L agar) for 2 days before inoculation and coculture with Agrobacterium tumefaciens EHA105. The preincubated hypocotyls were soaked in Agrobacterium-infection buffer (MS and 30 g/L sucrose; pH 5.8-5.9) for 15 min and transferred to the cocultivation media (MS, 0.05 mg/L NAA, 3 mg/L 6-BA, 30 g/L sucrose, and 6 g/L agar) in the dark for 48 h at 25 °C. The explants were subsequently transferred to callus- and shoot-induction media (MS, 0.05 mg/L NAA, 3 mg/L 6-BA, 30 g/L sucrose, 6 g/L agar, +4 mg/L phosphinothricin (PPT; for bar selection), and 400 mg/L carbenicillin (Cb)). When the regenerating shoots reached 1-2 cm in height, they were transferred to rooting media (MS, 0.1 mg/L NAA, sucrose, 6 g/L agar, 4 mg/L PPT, and 400 mg/L Cb) to obtain transgenic cabbage plants.

Detection of the mutations

Genomic DNA was extracted from individual T0, T1 and F1 plants using cetyltrimethylammonium bromide as previously described⁴⁸. The CRISPR/Cas9 T-DNA-positive transgenic lines were genotyped for mutations using primers flanking the target sequence. The mutation

efficiency was counted as described by Ma et al.⁴⁹. The site mutation types were detected through target region cloning and sequencing. All the detection primers are presented in Supplementary Table 1.

Self-compatibility or cross-compatibility investigation of the BoSRK gene mutant plants

The flower buds were covered with a bag one day before anthesis to prevent uncontrolled cross-pollination and pollinated with pollen from the same plant or a wild-type plant with the same S-allele on the day of flowering, with self- or cross-pollination at the bud stage performed as the control. For fluorescence microscopy observation of the pollen germination on the stigma, the pistils were removed from the buds 5 h after the pollen was applied to the stigma and fixed in acetic alcohol (ethanol: acetic acid, 3:1 (v/v)) for 5 h at room temperature. The fixed pistils were hydrolyzed in 2 M NaOH for at least 2 h at 60 °C and then cleared by incubation with 50 mM potassium phosphate solution several times. They were subsequently stained with decolorized aniline blue solution (0.1% aniline blue in 2% K₃PO₄) for 2 h. Finally, the stained pistils were mounted on glass slides with 50% glycerol and observed using a UV-fluorescence microscope (Leica CTR5000, Wetzlar, Germany). The self-compatibility index was calculated using the following formula: selfcompatibility index = number of seeds/number of flowers. At least three replicates of each treatment were used in the experiment.

Floral organ and cytological analysis of the *BoMS1* gene mutant plants

On the day of flowering, individual flowers were cut from the inflorescence using a scalpel to observe the structures of the stamen, anther, and pistil under a stereomicroscope. Buds (approximately 3 mm long) from the *BoMS1* gene mutant and the wild type were fixed overnight in 50% ethanol, 5% acetic acid, and 3.7% formaldehyde in water, dehydrated through an ethanol series (30%, 50%, 70%, 80%, 90%, 100%), and embedded into paraffin blocks⁵⁰. Cross-sections were cut, approximately 2–3-µm-thick, stained in hematoxylin, and photographed using a Leica CTR5000 microscope.

Acknowledgements

Sincere thanks are extended to Dr. Yaoguang Liu of the South China Agricultural University for providing the pYLCRISPR/Cas9P35S-B vector. We thank the National Key Research and Development Program of China (2017YFD0101804) and the Science and Technology Innovation Program for the Social Undertakings and the People's Livelihood in Chongqing (cstc2015shms-ztzx80005; cstc2015shms-ztzx80007; cstc2015shms-ztzx80009) for their financial support.

Authors' contributions

H.S. designed the experiments. C.M. conducted the transformation and molecular characterization of the *BoPDS* and *BoSRK3* mutants. CZ. conducted the transformation and molecular characterization of the *BoMS1* and *BoSRK3*

gene double mutants. M.Z. and M.L. conducted the vector construction and DNA cloning. C.Z. and Q.L. conducted the fluorescence microscopy investigation. J.S. and X.R. performed cytological analysis and self-compatibility or cross-compatibility investigation. D.Z. and B.L. performed the mutation inheritance analysis. H.S. and C.M. wrote/edited the manuscript.

Conflict of interest

The authors declare that they have no conflict of interest.

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Supplementary Information accompanies this paper at (https://doi.org/10.1038/s41438-018-0107-1).

Received: 3 September 2018 Revised: 24 October 2018 Accepted: 31 October 2018

Published online: 01 February 2019

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