Simultaneous induction of mutant alleles of two allergenic genes in soybean by using site-directed mutagenesis

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Abstract

Background: Soybean (Glycine max) is a major protein crop, because soybean protein has an amino acid score comparable to that of beef and egg white. However, many allergens have been identified among soybean proteins. A decrease in allergenic protein levels would be useful for expanding the market for soybean proteins and processed foods. Recently, the CRISPR/Cas9 system has been adopted as a powerful tool for the site-directed mutagenesis in higher plants. This system is expected to generate hypoallergenic soybean varieties.

Results: We used two guide RNAs (gRNAs) and Agrobacterium-mediated transformation for simultaneous site-directed mutagenesis of two genes encoding the major allergens Gly m Bd 28 K and Gly m Bd 30 K in two Japanese soybean varieties, Enrei and Kariyutaka. We obtained two independent T₀ Enrei plants and nine T₀ Kariyutaka plants. Cleaved amplified polymorphic sequence (CAPS) analysis revealed that mutations were induced in both targeted loci of both soybean varieties. Sequencing analysis showed that deletions were the predominant mutation type in the targeted loci. The Cas9-free plants carrying the mutant alleles of the targeted loci with the transgenes excluded by genetic segregation were obtained in the T₂ and T₃ generations. Variable mutational spectra were observed in the targeted loci even in T₂ and T₃ progenies of the same T₀ plant. Induction of multiple mutant alleles resulted in six haplotypes in the Cas9-free mutants derived from one T₀ plant. Immunoblot analysis revealed that no Gly m Bd 28 K or Gly m Bd 30 K protein accumulated in the seeds of the Cas9-free plants. Whole-genome sequencing confirmed that a Cas9-free mutant had also no the other foreign DNA from the binary vector. Our results demonstrate the applicability of the CRISPR/Cas9 system for the production of hypoallergenic soybean plants.

Conclusions: Simultaneous site-directed mutagenesis by the CRISPR/Cas9 system removed two major allergenic proteins from mature soybean seeds. This system enables rapid and efficient modification of seed components in soybean varieties.

Keywords: CRISPR/Cas9, Glycine max, Gly m Bd 28 K, Gly m Bd 30 K, Hypoallergenic soybean

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Background
Soybean (*Glycine max*, 2n = 2x = 40) is one of the most important protein crops used for food and forage worldwide, because its seeds contain high-quality proteins with an amino acid score comparable to that of beef and egg white [1]. Diverse soybean proteins are responsible for the physical properties of foods and other products made from soybean seeds [2, 3]. In the USA and Europe, 5 to 8% of babies and 2% of adults are allergic to soybean [4]. Several subunits of major storage proteins such as 7S and 11S globulins and 2S albumin are representative soybean allergens [5]. The vicilin-like glycoprotein Gly m Bd 28 K and the oil-body-associated protein Gly m Bd 30 K are also reported as major soybean allergens [6, 7]. Hydrophobic proteins Gly m 1A and Gly m 1B and the hull protein Gly m 2 are related to asthma outbreaks in Spain [8, 9]. Profilin Gly m 3 and the pathogenesis-related protein Gly m 4 are cross-reactive with antigens from other sources involved in sensitization and symptom induction [10, 11]. Positive response to soybean protein in allergic reaction has been reported in 14% of patients diagnosed with food allergies with atopic dermatitis [12]. Therefore, development of hypoallergenic soybean varieties or establishment of a procedure to remove allergens would be useful for expanding the market of soybean proteins and processed foods.

Protein fractionation on the basis of the differences in protein solubility at different salt concentrations and pH can be used to characterize the biochemical and physical properties of proteins [13–15]. This technique is also used for the removal of specific allergens from soy foods. Gly m Bd 30 K was efficiently removed from soy milk by acidifying it to pH 4.5 with 1 M Na₂SO₄ [16].

Genetic improvement of soybean is achieved by crossing plants carrying allergen-deficient alleles from soybean genetic resources or by mutagenesis to generate allergen-deficient mutant alleles. A number of spontaneous or induced mutants deficient in subunits of 7S or 11S globulins have been reported [17–20]. Among the germplasm of wild soybean (*G. soja*), Hajika et al. [20] found one accession lacking the α-, α’, and β-subunits of 7S globulin. The deficiency of these subunits is controlled by a single dominant gene (*Scg-1*), which is closely associated with post-transcriptional gene silencing [21]. To develop hypoallergenic soybean through crossing and subsequent back-crossing, this dominant gene has been introduced into an elite variety, Fukuyutaka [22]. The soybean variety Yumeminori lacks α- and α’-subunits of 7S globulin, and Gly m Bd 28 K, and has a decreased level of the β-subunit of 7S globulin; this variety has been developed through mutagenesis by gamma-ray irradiation [23]. Mutagenesis of the soybean variety VLSoy-2 by gamma-ray irradiation generated mutant lines lacking the A3-subunit of 11S globulin [24]. This mutagenesis also produced plants lacking α- and α’-subunits of 7S globulin [24]. Stacking of recessive mutant alleles of the genes for Kunitz trypsin inhibitor, agglutinin, and Gly m Bd 30 K was performed in the genetic background of the soybean variety Williams 82 [25]. Proteome analysis revealed that the stacking of these mutant alleles markedly decreased the accumulation of these allergens [25].

The biotechnological approach can also help to decrease the accumulation of allergens in soybean seeds. Down-regulation of the gene encoding Gly m Bd 30 K greatly suppresses the accumulation of the targeted protein in seeds of transgenic soybean [26]. The accumulation of α-, α’, and β-subunits of 7S globulin in soybean seeds can be greatly decreased through RNA interference or artificial microRNA systems [27, 28]. Recently, the transcription activator-like effector nucleases (TALENs) and clustered regularly interspaced short palindromic repeats (CRISPR)/CRISPR-associated endonuclease 9 (Cas9) systems have become the main platforms for site-directed mutagenesis in higher plants [29–32]. They enable the fine tuning of traits in soybean breeding when applied to various soybean varieties. The CRISPR/Cas9 system can be used to develop hypoallergenic soybean directly from elite varieties, because it has been optimized for various soybean varieties [33–37].

The subunits of 7S and 11S globulins are closely associated with seed characteristics important for food processing such as gel-forming and emulsifying properties [38–40]. To produce hypoallergenic soybeans without impairing the processing properties, we focused on two allergenic proteins, Gly m Bd 28 K and Gly m Bd 30 K, because no pyramiding of mutant alleles of these allergens in soybean has been reported. Here, we constructed a plasmid for simultaneous site-directed mutagenesis of these genes with the CRISPR/Cas9 system and used it for *Agrobacterium*-mediated transformation of two soybean varieties. *Cas9*-free plants carrying mutant alleles of the targeted loci, with the transgenes excluded by genetic segregation, were obtained in the T₂ or T₃ generations. Immunoblot analysis revealed that Gly m Bd 28 K and Gly m Bd 30 K proteins did not accumulate in seeds of the *Cas9*-free plants. Our results demonstrate the applicability of the CRISPR/Cas9 system for the production of hypoallergenic soybean plants.

Results
Generation of transgenic soybean plants harboring the CRISPR/Cas9 expression module.

To conduct the site-directed mutagenesis of soybean with the CRISPR/Cas9 system, we designed two guide RNAs (gRNAs) to mutagenize the Gly m Bd 28 K and Gly m Bd 30 K loci (Fig. 1a). Explants of Enrei and Kariyutaka were inoculated with *Agrobacterium* harboring the pMR284_28K_30K plasmid (Fig. 1b). Two Enrei T₀...
plants (E1 and E2) and nine Kariyutaka T0 plants (K1 to K10) were obtained. All the T0 plants set T1 seeds. In our previous study, many T0 plants produced by our soybean transformation system failed to transmit the transgenes into the T1 progeny [41]. This fact indicates that T0 plants originated from chimeric tissues which contained transformed and non-transformed cells [41]. Therefore, we did not examine transgene integration or the induction of mutagenesis in the T0 plants, and grew representative T1 plants for further analyses. The plant numbers of the T1 and T2 generations were indicated by giving the T0 individual number followed by the branch number for each generation.

**Mutations in T1 plants detected by cleaved amplified polymorphic sequence (CAPS) analysis**

Representative 20 T1 Enrei plants and 25 T1 Kariyutaka plants derived from 12 T0 plants were grown, and the induction of mutagenesis in the targeted loci was evaluated by CAPS analysis of the genomic DNA. The DNA fragments were classed into wild-type and mutant-type based on the expected size; a fragment of unexpected size was also detected and considered as mutant-type (Fig. 2). Mutations were detected in both targeted loci in plants of both varieties (Fig. 2, Additional file 2: Figure S1, Additional file 1: Tables S1, S2). Integration of the transgene was also examined by PCR analysis with Cas9-specific primers. PCR analysis revealed that 11 T1 plants (24.4% of all T1 plants examined) were Cas9-free; among these, E1–4, E1–8, and E1–9 had mutant alleles in the Gly m Bd 30 K locus, whereas the others had the wild-type alleles of both targeted loci (Additional file 1: Tables S1, S2).

**Transmission of mutations and transgenes to the T2 generation**

Because none of the Cas9-free T1 plants had mutations in both targeted loci, 13 representative T1 plants were advanced to the next generation. A total of 348 T2 seeds collected from the 13 T1 plants were evaluated for the mutations in the targeted loci (Table 1). In CAPS analysis, 227 (65%) T2 seeds showed mutant-type fragments of both targeted loci (Table 1, Additional file 2: Figs. S2, S3). No mutations were detected in 20 T2 seeds.
A Gly m Bd 28K

---AGTAATCGACGTACCTGGCAACCCAGATTCTTCTCT
Mutant-type fragment (449 bp)

Wild-type fragment (386 bp)

T₁ plant

M E1-1 E2-1 E3-1 E4-1 E5-1 E6-1 E7-1 E8-1 E9-1 E10-1 E11-1 Enrei

1,000 bp

500 bp

B Gly m Bd 30K

---GGAAAAAGAGGTGTCATCACCACAGAAGCTC
Mutant-type fragment (519 bp)

Wild-type fragment (386 bp)

T₁ plant

M E1-1 E2-1 E3-1 E4-1 E5-1 E6-1 E7-1 E8-1 E9-1 E10-1 E11-1 Enrei

1,000 bp

500 bp

Fig. 2 Confirmation of mutagenesis of targeted loci in representative Enrei-T₁ plants by CAPS analysis. The schematic diagrams above panels show Ddel or BsaII restriction sites (shaded in gray) in fragments amplified with specific primers. Red and blue nucleotide sequences have the same meaning as those in Fig. 1. Gray arrows, the sizes of expected wild-type fragments; black arrows, the sizes of expected mutant-type fragments; open triangle, a fragment of unexpected size considered as mutant type. M, molecular weight marker (100-bp ladder).

(Table 1). Thus, frequency of simultaneous site-directed mutagenesis in both targeted loci was much higher in the T₂ generation than in the T₁ generation (Table 1, Additional file 1: Tables S1, S2).

Although the Cas9 gene was detected in all 13 T₁ plants (Additional file 1: Tables S1, S2), it was removed by genetic segregation in 52 T₂ seeds (Table 2, Additional file 2: Fig. S4). Among them, 38 T₂ seeds had
mutant alleles in one or both targeted loci (Table 2). Of these 38 seeds, 14 seeds belonging to both varieties had mutant alleles in both targeted loci (double-mutants; Table 2). Representative mutational spectra of the T 2 seeds are shown in Fig. 3. The genotypes were divided into homozygous (the Gly m Bd 28 K locus in K4–1-37 and Gly m Bd 30 K locus in E1–5–17, K1–3–11, and K4–1-37 T2 seeds), heterozygous (the Gly m Bd 28 K locus in E1–2–6 and E1–5–17, and K1–3–11 T2 seeds), and biallelic mutant types (the Gly m Bd 30 K locus in E1–2–6) (Fig. 3). Deletions were predominant in the mutational spectra of these T 2 seeds (Fig. 3).

**Development of double-mutant T 3 seeds**

We used the heterozygous and biallelic mutants to develop more homozygous mutant alleles. We collected T 3 seeds and sequenced both targeted loci. In total, 4 haplotypes in Enrei and 21 haplotypes in Kariyutaka were found in the double-mutants (Table 3). Deletions (1 to 43 nucleotides) were the most common mutations (Table 3, Additional file 2: Figure S5). Predicted amino acid sequences of Gly m Bd 28 K (Figure S6) and Gly m Bd 30 K (Figure S7) are shown in Additional file 2. Three mutant alleles (d3 and d6 for the Gly m Bd 28 K locus, and d6 for the Gly m Bd 30 K locus) had in-frame mutations (Additional file 2: Figures S6, S7). In the Gly m Bd 30 K locus, the 3-nucleotide deletion generated a stop codon at the mutation site, and the 33-nucleotide deletion was not predicted as an in-frame mutation, because the deleted region contained the splicing site (Additional file 2: Figure S7).

**Analysis of Gly m Bd 28 K and Gly m Bd 30 K proteins in mature double-mutant seeds**

We selected two Enrei haplotypes (E-type1 and E-type3) and seven Kariyutaka haplotypes (K-type2, K-type4, K-type7, K-type9, K-type14, K-type15, and K-type19) from the double-mutants (Table 3), and examined the composition of crude protein fractions prepared from mature seeds. The Gly m Bd 30 K protein was visually detectable in Enrei and Kariyutaka but not mutant seeds, whereas Gly m Bd 28 K was not detectable in any seeds in the SDS-PAGE analysis (Fig. 4a). The double-mutant seeds had no signal bands that were not detected in wild-type seeds (Fig. 4a). To detect the Gly m Bd 28 K and Gly m Bd 30 K proteins specifically, immunoblot analysis was conducted in double-mutant and wild-type seeds. In the immunoblot analysis, the Gly m Bd 28 K and Gly m Bd 30 K proteins were detected only in seeds of wild-type Enrei or Kariyutaka, except that Gly m Bd 28 K was also detected in the E-type3 haplotype (Fig. 4b, c). No immunoreactive band of unexpected size was detected (Additional file 2: Figure S8).

**Expression levels of the Gly m Bd 28 K and the Gly m Bd 30 K genes**

To evaluate the expression levels of the Gly m Bd 28 K and the Gly m Bd 30 K genes, we extracted total RNA from the mature T 3 seeds of two Enrei mutants (E-type1 and E-type3), seven Kariyutaka mutants (K-type2, K-type4, K-type7, K-type9, K-type14, K-type15, and K-type19), Enrei and Kariyutaka, and conducted semi-quantitative RT-PCR analysis of the region up-stream of the mutation site (Additional file 2: Figure S9). Although

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**Table 1** Frequency of simultaneous site-directed mutagenesis in the targeted loci Gly m Bd 28 K and Gly m Bd 30 K in T 2 seeds

| T 1 plant number | Number of T 2 seeds | Mutation | Gly m Bd 28 K and Gly m Bd 30 K | Gly m Bd 28 K only | Gly m Bd 30 K only | No mutation | Total |
|------------------|----------------------|----------|---------------------------------|--------------------|--------------------|-------------|-------|
| E1–1             | 18                   | Gly m Bd 28 K and Gly m Bd 30 K | 11                 | 8                  | 2                  | 39          |
| E1–2             | 22                   | Gly m Bd 28 K only              | 0                  | 3                  | 0                  | 25          |
| E1–5             | 9                    | Gly m Bd 30 K only              | 4                  | 3                  | 1                  | 17          |
| E2–1             | 0                    | Gly m Bd 28 K and Gly m Bd 30 K | 22                 | 0                  | 10                 | 32          |
| E2–2             | 1                    | Gly m Bd 28 K only              | 20                 | 0                  | 5                  | 26          |
| K1–1             | 12                   | Gly m Bd 28 K and Gly m Bd 30 K | 1                  | 0                  | 0                  | 13          |
| K1–3             | 16                   | Gly m Bd 28 K only              | 3                  | 4                  | 2                  | 25          |
| K2–1             | 17                   | Gly m Bd 30 K only              | 3                  | 3                  | 0                  | 23          |
| K2–2             | 20                   | Gly m Bd 30 K only              | 7                  | 4                  | 0                  | 31          |
| K4–1             | 35                   | Gly m Bd 28 K only              | 0                  | 2                  | 0                  | 37          |
| K5–1             | 50                   | Gly m Bd 28 K only              | 0                  | 0                  | 0                  | 50          |
| K5–2             | 14                   | Gly m Bd 30 K only              | 1                  | 0                  | 0                  | 15          |
| K6–1             | 12                   | Gly m Bd 28 K only              | 3                  | 0                  | 0                  | 15          |

The presence of mutations was evaluated by CAPS analysis

The letter and first number correspond to the number of the parental T 0 plant

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amplified products of the 18S ribosomal RNA (18S rRNA) were detected at similar levels in all mature seeds of the mutants, Enrei, and Kariyutaka (Fig. 5), all mutants showed lower expression levels of the Gly m Bd 28 K and Gly m Bd 30 K genes than those of wild-type (Enrei and Kariyutaka) seeds (Fig. 5).

Whole-genome sequencing in T2 plants to validate the absence of foreign DNA
T2 mutant plants K2–1–16 and K4–1–37 were selected for whole-genome sequencing analysis. These plants had homozygous mutant alleles in both loci. The K2–1–16 plant had mutant alleles with a single-nucleotide insertion in the Gly m Bd 28 K and a 2-nucleotide deletion in the Gly m Bd 30 K loci. The genome of K4–1–37 contained mutant alleles with a 6-nucleotide deletion in the Gly m Bd 28 K locus and a 2-nucleotide deletion in the Gly m Bd 30 K locus. PCR analysis detected the presence of the Cas9 gene in the genome of K2–1–16, whereas K4–1–37 was Cas9 free. The whole genomes of the two T2 plants were sequenced, and the presence of foreign DNA was examined by the k-mer detection method [42]. Each 20-mer identical between the plant genome and the vector was detected (Fig. 6). The genome of K2–1–16 clearly showed significant signals in a vector-wide manner (Fig. 6a, c), whereas that of K4–1–37 had no signal of foreign DNA from the vector (Fig. 6b). A significant signal found in the G-statistic of K4–1–37 was considered as a false positive, because it had a much lower value than that of K2–1–16.

Morphological characteristics of double mutants
To assess the consequences of the site-directed mutagenesis in the targeted loci, we examined the

| T1 plant number | Cas9 integration | Number of T2 seeds | Gly m Bd 28 K and Gly m Bd 30 K | Gly m Bd 28 K only | Gly m Bd 30 K only | No mutation | Total |
|-----------------|-------------------|--------------------|-------------------------------|-------------------|-------------------|-------------|------|
| E1–1 | positive | 16 | 10 | 3 | 0 | 29 |
| free | 2 | 1 | 5 | 2 | 10 |
| E1–2 | positive | 21 | 0 | 0 | 0 | 21 |
| free | 1 | 0 | 3 | 0 | 4 |
| E1–5 | positive | 8 | 4 | 0 | 1 | 13 |
| free | 1 | 0 | 3 | 0 | 4 |
| E2–1 | positive | 0 | 20 | 0 | 5 | 25 |
| free | 0 | 2 | 0 | 5 | 7 |
| E2–2 | positive | 1 | 20 | 0 | 0 | 21 |
| free | 0 | 0 | 0 | 5 | 5 |
| K1–1 | positive | 9 | 1 | 0 | 0 | 10 |
| free | 3 | 0 | 0 | 0 | 3 |
| K1–3 | positive | 13 | 3 | 0 | 0 | 16 |
| free | 3 | 0 | 4 | 2 | 9 |
| K2–1 | positive | 16 | 3 | 1 | 0 | 20 |
| free | 1 | 0 | 2 | 0 | 3 |
| K2–2 | positive | 19 | 7 | 0 | 0 | 26 |
| free | 1 | 0 | 4 | 0 | 5 |
| K4–1 | positive | 33 | 0 | 2 | 0 | 35 |
| free | 2 | 0 | 0 | 0 | 2 |
| K5–1 | positive | 50 | 0 | 0 | 0 | 50 |
| free | 0 | 0 | 0 | 0 | 0 |
| K5–2 | positive | 14 | 1 | 0 | 0 | 15 |
| free | 0 | 0 | 0 | 0 | 0 |
| K6–1 | positive | 12 | 3 | 0 | 0 | 15 |
| free | 0 | 0 | 0 | 0 | 0 |

Table 2 Frequency of Cas9-positive and Cas9-free T2 seeds

*The letter and first number correspond to the number of the parental T0 plant.
morphology of the mutant plants. Because many mutant alleles in the targeted loci were detected in the Kariyutaka T2 generation (Fig. 3), we examined the morphological characteristics of T2 plant body and T3 seed size and shape. No difference was detected between the double-mutants and wild-type in the plant and seed morphological characteristics (Additional file 2: Figures S10, S11).

Discussion

Gly m Bd 28K and Gly m Bd 30K are the major allergenic proteins in soybean seeds [6, 43]. The mutant alleles of these loci have been identified by surveying the soybean germplasm or generated by gamma-ray irradiation mutagenesis [19, 44, 45], and stacking of these mutant alleles will enable development of hypoallergenic soybean lines. In contrast, site-directed mutagenesis mediated by the CRISPR/Cas9 system enables the induction of mutations directly in the targeted loci of the desirable donor plants such as varieties and elite breeding lines. This approach dramatically shortens breeding period and saves labor. In this study, we performed simultaneous site-directed mutagenesis of both Gly m Bd 28K and Gly m Bd 30K loci in two Japanese soybean varieties. A total of 14 T2-generation seeds possessed mutant alleles of both loci and had the Cas9 gene removed through genetic segregation (Table 3). Among all mutations, deletions were predominant and caused frame-shifts (Additional file 2: Figures S4–S6). The frame-shift mutations resulted in the deficiency in proteins recognized by the polyclonal antibodies against GlymBd28K and GlymBd30K proteins (Fig. 4). No bands of unexpected size were detected with either of these antibodies (Fig. 4). Frame-shift mutations in the targeted loci decreased the expression levels of the GlymBd28K and GlymBd30K genes (Fig. 5). These findings suggest that the frame-shift mutations produce aberrant mRNAs from the targeted locus, which induced nonsense mRNA decay (NMD), like in a site-directed mutagenesis study conducted in Brassica carinata using the hairy root transformation system [46]. The lower expression level than wild-type might result in the...
deficiency in proteins recognized by the polyclonal antibodies against Gly m Bd 28 K and Gly m Bd 30 K proteins. On the other hand, several T3 seeds had mutant alleles with putative in-frame mutations (Additional file 2: Figures S6, S7). The E-type3 haplotype with a 3-nucleotide deletion in the Gly m Bd 28 K locus showed a strong immunoreactive band with the antibody against the Gly m Bd 28 K protein, whereas the expression level of the Gly m Bd 28 K gene was lower than that in Enrei (Fig. 5). In this study, the expression level of the targeted loci was examined in only mature seeds of representative mutants and wild-type. Soybean seeds accumulate Gly m Bd 28 K and Gly m Bd 30 K proteins during seed filling [47]. Therefore, an investigation of the expression level of the targeted loci in immature seeds might lead to further understanding of accumulation mechanism of mutant proteins.

At least three immunodominant epitopes in Gly m Bd 28 K and five in Gly m Bd 30 K have been identified [48–50]. In this study, gRNAs were designed against the fourth exon of Gly m Bd 28 K and first exon of Gly m Bd 30 K (Fig. 1). Immunodetection of proteins generated by the in-frame mutations in T3 seeds would indicate the presence proteins with preserved epitopes (Additional file 2: Figures S6, S7). Analysis of sera of soybean-allergic patients may further clarify the allergenic properties of soybean seeds generated in this study.

Multiple mutant alleles were detected in the progeny of one T0 plant (Fig. 7). Three mutant alleles (i1, d2, and d5) in the Gly m Bd 28 K locus and five (d1, d2, d5, d6, and d1s1) mutant alleles in the Gly m Bd 30 K locus were ascertained in the Cas9-free T2 and T3 seeds derived from the K1 T0 plant (Fig. 7). These mutations appeared after the T2 generation, when the distribution of mutant alleles in the targeted loci was validated in the genealogy of the K1 plant and its progeny (Fig. 7). Twelve haplotypes (K-type1 to K-type12) were consequently obtained in the Cas9-free T3 seeds (Fig. 7). Previously, we showed that simultaneous site-directed mutagenesis of duplicated loci using a single gRNA resulted in heterozygous and/or chimeric mutations in the targeted loci in most of the T1 plants [36]. On the other hand, the mutant alleles of multiple targeted loci have been induced in early generations such as T0 or T1 plants in other studies on soybean site-directed mutagenesis by the CRISPR/Cas9 system [37, 51, 52]. This difference might be explained by different growth and maturity habits of the soybean varieties used. Kariyutaka has early flowering and a short period of vegetative growth [53]; the latter might decrease the chance of the occurrence of mutations in germ cells in the T0 generation, however, might produce multiple mutant alleles after the T1 generation. Therefore, the site-directed mutagenesis using Kariyutaka might be useful system for obtaining multiple mutant alleles in targeted genes efficiently in a limited number of transgenic soybean plants.

### Table 3 Mutational spectra in double-mutant T3 plants

| Donor plant | Haplotype | Gly m Bd 28 K locus | Gly m Bd 30 K locus |
|-------------|-----------|---------------------|---------------------|
| Enrei       | E-type1   | i1, i1              | d7, d7              |
|             | E-type2   | d3, d3              | d3, d3, d3          |
|             | E-type3   | d3, d3              | d7, d7              |
|             | E-type4   | d3, d3              | d4, d4              |
| Kariyutaka  | K-type1   | i1, i1              | d5, d5              |
|             | K-type2   | d5, d5              | d2, d2              |
|             | K-type3   | d5, wt              | d2, d2              |
|             | K-type4   | d5, d5              | d1, d1              |
|             | K-type5   | d5, d5              | d6, d6              |
|             | K-type6   | d5, d5              | d1, d6              |
|             | K-type7   | d5, d5              | d1s1, d1s1          |
|             | K-type8   | d5, d5              | d1s1, wt            |
|             | K-type9   | d2, d2              | d1s1, d1s1          |
|             | K-type10  | d2, d2              | d1s1, wt            |
|             | K-type11  | d2, d5              | d1s1, d1s1          |
|             | K-type12  | d2, d5              | d1s1, wt            |
|             | K-type13  | d2, d5              | d1, d1              |
|             | K-type14  | i1, i1              | d43, d43            |
|             | K-type15  | i1, i1              | d1, d1              |
|             | K-type16  | i1, i1              | d1, d43            |
|             | K-type17  | i1, wt              | d1, d43            |
|             | K-type18  | d6, d6              | d3, d3              |
|             | K-type19  | i1, i1              | d2, d2              |
|             | K-type20  | i1, i1              | d2, d1s1           |
|             | K-type21  | d6, d6              | d2, d2              |

Letters and numbers denote the alleles in the targeted loci: e.g., d7, a single-nucleotide deletion; i1, a single-nucleotide insertion; d1s1, a single-nucleotide deletion and substitution; wt, wild type

*A Heterozygous mutation in the targeted locus

*B Allelic mutations in the targeted locus

**Conclusion**

We used Agrobacterium-mediated transformation and two gRNAs for simultaneous site-directed mutagenesis of two allergenic genes, Gly m Bd 28 K and Gly m Bd 30 K, in two Japanese soybean varieties. Cas9-free plants that had mutant alleles of the targeted loci and transgenes excluded by genetic segregation were obtained in the T2 or T3 generation. Immunoblot analysis revealed that the double-mutants did not accumulate Gly m Bd 28 K or Gly m Bd 30 K protein in mature seeds. Our results showed that simultaneous site-directed mutagenesis by the CRISPR/Cas9 system removed two major allergenic proteins in mature soybean seeds.
Methods

Vector construction
We constructed a gRNA expression vector (pLeg-base) which contained two gRNA expression cassettes. The frame sequence of the gRNA scaffold was derived from the vector pEn-Chimera [54]. Promoter regions of Arabidopsis U6–26 [54] and soybean U6–16g [55] were used to control gRNA and gRNA scaffold expression (Fig. 1b). The soybean allergenic genes Gly m Bd 28 K (Glyma.08G116300.1) and Gly m Bd 30 K (Glyma.08G116300.1) were the targets for simultaneous site-directed mutagenesis (Fig. 1a). Two 20-nucleotide sequences (5′-CTCAGCGAACGGATATTGG-3′ and 5′-ACCCACAGTAAAAGTAGCAAGG-3′) identical in each gene were used to design the gRNA sequences with the web-based CRISPR-P 2.0 (http://crispr.hzau.edu.cn/CRISPR2/). The pLeg-base vector was digested with the BbsI or BsaI restriction enzyme (NEB, Ipswich, USA). Oligonucleotides designed to match the gene-specific sequence were annealed to each other to form the gRNA seed sequence, which was ligated into pLeg-base. The CRISPR/Cas9 expression plasmid (pMR284_28K_30K) was constructed by inserting the gRNA expression cassettes of pLeg-base into a Cas9-binary vector (pMR284) harboring Cas9 and glufosinate resistance gene (Bar) expression cassettes using LR Clonase (Thermo Fisher Scientific, Waltham, USA).

Fig. 4 SDS-PAGE and immunoblot analyses of the crude proteins of representative double-mutant T3 and wild-type mature seeds. a Proteins separated by SDS-PAGE and stained with Coomassie Brilliant Blue. Designations of mutations are as in Table 3. Asterisks denote signal of putative Gly m Bd 30 K protein. M, molecular weight marker. b Immunoblot analysis using polyclonal antibody against Gly m Bd 28 K protein. c Immunoblot analysis using polyclonal antibody against Gly m Bd 30 K protein. Images of full-length gel and blots are provided in additional file 2 (Figure S8)
Soybean transformation

The Japanese soybean varieties Enrei (JP 28862) and Kariyutaka (JP 86520) were obtained from Genebank, National Agriculture and Food Research Organization (https://www.gene.affrc.go.jp/index_en.php). Agrobacterium-mediated transformation was performed as described in [28], except that the concentration of glufosinate for selection of transformed cells was decreased from 6 mg/L to 4 mg/L for Enrei. *Agrobacterium tumefaciens* EHA105 harboring the plasmid pMR284_28K_30K was used. Transgenic plants were grown in commercial soil (Katakura Chikkarin Co.,
Tokyo, Japan) at 25 °C in an isolated greenhouse for transgenic plants.

Extraction of genomic DNA and detection of mutations in targeted loci by CAPS analysis

To extract leaf genomic DNA, leaf pieces (approximately 5 mm × 5 mm) were homogenized in 200 μL of extraction buffer [2% CTAB (hexadecyltrimethyl-ammonium bromide), 100 mM Tris-HCl (pH 8.0), 20 mM EDTA (pH 8.0), 1.4 M NaCl, and 0.07% 2-mercaptethanol] in a BioMasher II tube (Nippi, Tokyo, Japan). To extract genomic DNA from mature seeds, a part of cotyledon was powdered and approximately 5 mg of powder was stirred in extraction buffer [10 mM Tris-HCl (pH 8.0), 5 mM EDTA, 0.5% SDS, 0.5% NP40, 0.5% Tween 20, and 80 mg/L proteinase-K (Wako, Osaka, Japan)]. The mixture was incubated at 50 °C for 1 h. DNA extracts were deproteinized with a mixture of phenol, chloroform, and isoamyl alcohol (25:24:1). DNA was precipitated from the supernatant with 2-propanol. The targeted regions in the Gly m Bd 28 K and Gly m Bd 30 K loci were amplified by PCR with specific primers (Additional file 1: Table S3). The PCR was performed under the following conditions: 30 cycles of 94°C for 30 s, 54°C (the Gly m Bd 28 K) or 60°C (the Gly m Bd 30 K) for 30 s and 72°C for 60 s. The amplified products were digested with the Ddel and BsaII restriction enzymes (NEB), respectively, and

Fig. 6 Detection of unintended remaining foreign DNA in T2 plants. The counts of k-mer and G-statistic are shown for (a) a transgenic sample K2–1-16 and (b) a Cas9-free sample K4–1-37. c Structure of the binary expression vector pMR284_28K_30K. The horizontal axis indicates the nucleotide positions in the vector; horizontal red line is the 1% significance level by G-test. G-statistic values exceeding the 1% level of significance are in red. The k-mer counts over 500 are omitted.
separated by electrophoresis in 2.0% agarose gels. The DNA fragments of expected digested-pattern derived from the targeted region carrying mutations and those with no mutations were considered as the mutant type and wild type, respectively. DNA fragments of unexpected size were also regarded as mutant type.

DNA sequencing
The targeted and flanking regions of the Gly m Bd 28 K and Gly m Bd 30 K loci were amplified with specific primers (Additional file 1: Table S3). The amplified products were cloned into the pGEM-T-Easy vector (Promega, Madison, USA) and sequenced with the Big Dye terminator cycle method using an ABI3100 or ABI3130 Genetic Analyzer (Thermo Fisher Scientific). DNA sequencing analysis was performed by the Instrumental Analysis Division, Graduate School of Agriculture, Hokkaido University.

Selection of Cas9-free plants
To confirm the integration of the Cas9 and gRNA expression module in T1–T3 generations, PCR analysis was performed using primers specific for the Cas9 gene (Additional file 1: Table S3). PCR was also performed to simultaneously amplify endogenous Glyma.01G214600 as a positive control. The PCR was performed under the following conditions: 30 cycles of 94°C for 30 s, 54°C for 30 s and 72°C for 30 s. The existence of the Cas9 gene were identified by the existence of products amplified by the PCR.

Protein analyses in mature seeds
Soy meal was collected from mature seeds. The extraction of crude protein and protein separation were performed as described in [28]. Proteins were separated by SDS-PAGE in a precast 5–12% gradient gel (ATTO, Tokyo, Japan) and transferred onto a PVDF membrane (Hybond-P; GE Healthcare, Little Chalfont, UK). Membranes were blocked with 5% skim milk (Wako) overnight at 4°C. Recombinant Gly m Bd 30 k was prepared using the baculovirus expression system as described in [5]. Using the pET52 vector (Merck-Millipore, Burlington, USA), His10-tagged Gly m Bd 28 K was expressed in Escherichia coli BL21(DE3). After sonication and centrifugation, Gly m Bd 28 K–containing pellets were dissolved in phosphate-buffered saline containing 8 M urea, and Gly m Bd 28 K was purified using a HisTrapFF crude column (GE Healthcare). Antisera were raised in rabbits against the recombinant proteins as described in [56]. Immunoreactive bands were detected with the antisera and the ECL Plus Western Blotting system (GE Healthcare).

![Fig. 7](image_url)
Expression analysis by semi-quantitative RT-PCR
Total RNA was extracted from mature seeds of mutants, Enrei, and Kariyutaka by the LiCl precipitation procedure [28]. Semi-quantitative RT-PCR was conducted in a 20-μL volume using 30 or 38 cycles of 94 °C for 30 s, 57 °C for 30 s, and 72 °C for 10 s. The transcript level of the Gly m Bd 28 K and the Gly m Bd 30 K gene was evaluated relative to that of the 18S rRNA gene (XR_003264275).

Genome sequencing
Total DNA was isolated from fresh leaves (1.0 g) of wild-type and T2 plants as described in [41]. Genomic DNA libraries were constructed using a TruSeq DNA PCR-Free Library Preparation Kit (Illumina, San Diego, USA). Whole-genome sequencing was conducted on an Illumina HiSeq X platform to obtain 151-bp paired-end reads. Approximately 50× coverage data were obtained for each sample. Unintended remaining foreign DNA was detected as described in [42].

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12870-020-02708-6.

Additional file 1: Table S1. Induction of mutations in the targeted loci and the integration of the Cas9 gene in representative T1 plants from the transformation of Enrei. Table S2. Induction of mutations in the targeted loci and the integration of the Cas9 gene in representative T1 plants from the transformation of Kariyutaka. Table S3. Primer sequences used for vector construction, confirmation of transgenes, and CAPS, semi-quantitative RT-PCR, and sequencing analyses.

Additional file 2: Figure S1. Confirmation of mutagenesis of targeted loci in representative Kariyutaka-T1 plants by CAPS analysis. Figure S2. Detection of mutations in the Gly m Bd 28K and Gly m Bd 30K loci in representative Enrei-T1 seeds by CAPS analysis. Figure S3. Detection of mutations in the Gly m Bd 28K and Gly m Bd 30K loci in representative Kariyutaka-T1 seeds by CAPS analysis. Figure S4. Detection of the integration of the Cas9 gene in representative T1 seeds by PCR analysis. Figure S5. Mutational spectra of the targeted loci in double-mutant T1 seeds. Figure S6. Alignment of predicted amino acid sequences of the Gly m Bd 28K locus in double mutants. Figure S7. Alignment of predicted amino acid sequences of the Gly m Bd 30K locus in double mutants. Figure S8. Full-length gel electrophoresis and immunoblot of the crude protein of representative double-mutant T1 and wild-type mature seeds. Figure S9. Primer sites used for semi-quantitative RT-PCR analysis of the Gly m Bd 30K and the Gly m Bd 30K loci. Figure S10. Morphological characteristics of representative double-mutant (T2) and control Kariyutaka plants. Figure S11. Morphological characteristics of representative double-mutant (T2) and control Kariyutaka seeds.

Abbreviations
PVD: Polyvinylidene difluoride; SDS-PAGE: Sodium dodecyl sulfate-polyacrylamide gel electrophoresis

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Authors’ contributions
SS, JA, and TY: conceived and designed the experiments; AH, YK, MM, ME, and SH: constructed vectors; SS, AH, and YK: performed site-directed mutagenesis; SS, KA, MH, and NM: analyzed mutations in transgenic soybean; TI: analyzed unintended remaining foreign DNA; SS, JA, and TY: contributed to the writing the manuscript. All authors have read and approved the manuscript.

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Availability of data and materials
The datasets supporting this study are included within the manuscript and its additional files. The vector developed in this study is available from the corresponding author on reasonable request. The datasets of genomic sequence in Kariyutaka and mutants (K2–1–16 and K4–1–37) have been deposited in the DDBJ Sequence Read Archive under the BioProject Accession PRJDB10633.

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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