Creation of a potato mutant lacking the starch branching enzyme gene *StSBE3* that was generated by genome editing using the CRISPR/dMac3-Cas9 system

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Abstract  The potato tuber starch trait is changed depending on the composition of amylose and amylopectin. The amount of amylopectin is determined by the activity of the starch branching enzymes SBE1, SBE2, and SBE3 in potato. SBE3, a homolog of rice BEI, is a major gene that is abundant in tubers. In this study, we created mutants of the potato *SBE3* gene using CRISPR/Cas9 attached to the translation enhancer dMac3. Potato has a tetraploid genome, and a four-allele mutant of the *SBE3* gene is desired. Mutations in the *SBE3* gene were found in 89 of 126 transformants of potato plants. Among these mutants, 10 lines contained four mutant *SBE3* genes, indicating that 8% efficiency of target mutagenesis was achieved. These mutants grew normally, similar to the wild-type plant, and yielded sufficient amounts of tubers. The potato starch in these tubers was similar to that of the rice BEI mutant. Western blot analysis revealed the defective production of SBE3 in the mutant tubers, suggesting that these transformants were loss-of-function mutants of *SBE3*.

Key words: amylopectin, CRISPR/Cas9, potato (*Solanum tuberosum* L.), starch, starch-branching enzyme (SBE).

Introduction

Storage starch in plants consists of amylose and amylopectin, the composition ratio of which alters starch properties. In many plant species, mutants showing altered amylose contents are known. Granule-bound starch synthase (GBSS) is a major enzyme involved in amylose biosynthesis (Nelson and Rines 1962). Mutants lacking this enzyme produce amylose-free starch. Mutants lacking the *GBSS* gene in rice show a waxy phenotype (Sano 1984). Similar mutants are also known in other plant species, such as maize (Shure et al. 1983), barley (Rohde et al. 1988), and sorghum (Pedersen et al. 2005).

Amylopectin is synthesized in concert by the activities of soluble starch synthase (SS) and starch branching enzyme (BE) (Guan et al. 1995). Mutants deficient in starch branching enzymes show aberrations in amylopectin. A rice mutant lacking the starch branching enzyme BEIIb, known as *ae*, shows elongated amylose chains with an increase in amylose content (Nishi et al. 2001; Yano et al. 1985). The mutant lacking BEI showed no significant change in the apparent amylose content, but the structure of amylopectin was altered because of a decrease in long chains and an increase in short chains of amylopectin. In addition, the endosperm starch of this mutant displayed altered properties of gelatinization (Satoh et al. 2003).

Potato, *Solanum tuberosum* L., is one of the most important crops in the world. There have been many attempts to breed potato mutants showing altered starch traits. However, the traditional breeding of potato is quite difficult because potato has a strong vegetative reproduction phase in addition to its polyploidy. Especially, use of recessive alleles is challenging because of its polyploid nature. To date, some genetically modified potatoes showing desired traits have been established. To obtain amylose-free potato starch, a knockdown transformant was created by introduction of the antisense gene of *GBSSI* (Visser et al. 1991).

Abbreviations: PCR, polymerase chain reaction; gRNA, guide RNA.
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Recently, genome editing technology has been applied to potato. A mutant lacking the production of a glycoalkaloid has been created (Sawai et al. 2014). However, it is difficult to establish a deficient potato mutant because potato has a tetraploid genome. Therefore, to introduce mutations in all four alleles for each gene in the genome, a powerful genome editing tool is required. We found a translational enhancer, dMac3, that strongly elevates the translation efficiency of the downstream ORF (Aoki et al. 2014). We established an improved CRISPR/Cas9 system using dMac3 and applied it to create a potato mutant lacking GBSS activity (Kusano et al. 2018). We confirmed that this mutant shows the trait of low amylose content in tuber starch (Kusano et al. 2018).

Many kinds of starch branching enzymes are found in potato. Among them, SBE1, SBE2, and SBE3 are known as the major enzymes involved in the biosynthesis of amylopectin (Larsson et al. 1996; Van Harsselaar et al. 2017). SBE2 is the counterpart of rice BEIIB and is considered to be involved in short chain amylopectin. A transformant harboring an antisense gene to SBE2 has been produced, and it exhibits an amylose extension trait similar to the rice ae mutant (Jobling et al. 1999). A mutant in which genes encoding SBE1 and SBE2 have been disrupted was reported and resulted in the severe amylose extension phenotype (Tuncel et al. 2019).

Three genes were identified as homologs of the rice BEI gene. Two genes, SBE1.1 and SBE1.2, have been reported as potato homologs due to their homology to *Arabidopsis* SBE1. Sequences of the potato SBE1 and SBE3 genes also have been published. However, potato SBE1 is not homologous to rice BEI because of low sequence similarity. Potato SBE3 shows high similarity to rice BEI. In addition, SBE1.1 and SBE1.2 are identical to that of potato SBE3 except for their C-terminal regions. Currently, SBE3 is considered the homolog of rice BEI and is a major starch branching enzyme gene that is commonly expressed in developing potato tuber (Van Harsselaar et al. 2017).

In this study, to elucidate the function of potato SBE3, we created a genome-edited potato mutant in which mutations occur in all four alleles of the SBE3 gene and the evaluation of the traits of the obtained mutant.

**Materials and methods**

**Plant material, tissue culture and plant growth conditions**

The *Solanum tuberosum* L. cv. Sayaka was used. Potato plants were cultured on medium containing an MS basal salt mixture (Fuji Film Wako, Tokyo, Japan) (Murashige and Skoog 1962), 3% sucrose, and 0.3% Gelrite (Fuji Film Wako) with the pH adjusted to 6.0. Plants were grown under long-day conditions with 16 h light and 8 h dark at 23°C in a growth chamber. The regeneration of potato plants was performed as described previously (Ohnuma et al. 2020). Regenerated potato plants were cultivated using a growth room. They were grown under 16 h light (200–400 µmol m−2 s−1) and 8 h dark conditions at 23°C in a growth chamber. Tubers were harvested from wild-type and transformed potatoes, which were grown at 22°C under 16 h light and 8 h dark conditions in a growth chamber.

**Plasmid construction**

The plasmid for CRISPR/Cas9 was constructed essentially according to Kusano et al. (2018). For construction of the gRNA genes, the DNA fragments corresponding to the target sequences were chemically synthesized. The fragments were inserted into the *BbsI* sites of the appropriate guide RNA vector, pMR203, pMR204, and pMR205 (kindly provided by M. Endo). The resultant gRNA genes were located between the *AtU6*-26 promoter and the gRNA scaffold sequence. They were introduced into pBS_GwIsceI using the multisite-Gateway system (Invitrogen, Carlsbad, CA, USA). The pBS_GwIsceI plasmid had the regions for the Gateway attR1 and attR2 sites lying between the two *I-SceI* sites (Kusano et al. 2018). The resultant plasmid contained the fragments corresponding to the tandem gRNA genes between two *I-SceI* sites. The gene fragments were digested with *I-SceI* and then introduced into the *I-SceI* sites of the pZD-dxCas9 plasmid (Kusano et al. 2018) to generate CRISPR/Cas9 vectors containing three gRNA genes.

**Plant transformation**

Potato was transformed using the Agrobacterium-mediated procedure using the *A. tumefaciens* EHA105 strain according to Yamada et al. (2004) and Kusano et al. (2018). Stem internodes were isolated from germ-free potato plants grown in a clean environment for 3 weeks and cut into approximately 1 cm pieces. The pieces were infected with *Agrobacterium* harboring an appropriate plasmid. Callus induction and plant regeneration were performed on a plate of 3CSZR medium (Sheeran and Bevan 1988) containing 0.3% Gelrite (Fuji Film Wako) supplemented with 5 mg l−1 hygromycin B (Fuji Film Wako) and cultured for 2 months. Regenerated plants were examined by detection of the transgene by PCR using the Cas9-S (5′-GCG GTA AGA ATA GAA TCT GTT AT-3′) and Cas9-T (5′-GACGCGCTATAGAATTTCCAA-3′) primers, which amplified part of the *Cas9* gene sequence. Genome edited potato plants were cultivated in a green room.

**Cleaved amplified polymorphic sequence (CAPS) analysis**

Genomic DNA was prepared from potato leaves using the RED Extract N-Amp™ Plant PCR kit (Sigma-Aldrich, St. Louis, USA). The 1.0-kb region in the potato SBE3 gene, in which the sequences of the target sites of the guide RNAs were contained, was PCR-amplified using the StSBE3_CAPS_Fw (5′-CTA TGA GAC GGA TAG TTG AGA ATG TG-3′) and StSBE3_CAPS_Rv (5′-GCC CAG GTG CCA CCT AAA TG-3′) primers that were derived from the regions in introns 1 and 3. The
amplified fragment was digested with BamHI. The wild-type gene has a BamHI site in the corresponding region. Therefore, we identified mutant alleles as those where the amplified fragment was no longer digested by BamHI. The number of mutant alleles among the four genes was estimated by the ratio of undigested fragments to digested fragments.

**DNA sequence analysis**
The 0.5-kb region of the potato SBE3 gene that contained the target sites of the guide RNAs was PCR-amplified using the StsSBE3_seq_Fw (5′-TGA TAT AAA CTG TGT GCA AAG CAA-3′) and StsSBE3_seq_Rv (5′-GGA GTG TCA GTG GCA AAG CAA-3′) primers. Then, the amplified fragment was inserted into the pTA2 vector (TOYOBO, Osaka, Japan). The resultant plasmids were used for nucleotide sequence analysis.

**Determination of the properties of tuber starch**
Starch granules were prepared from potato tubers according to Noda et al. (2004). Sliced sections of potato tuber were dipped into 0.4% potassium iodide-0.12% iodide solution. Photographs were taken after washing with water. The amylose contents in the potato tubers were analyzed according to a previous paper (Noda et al. 2004). The data were statistically analyzed using Dunnett’s multiple comparison test.

**Protein extraction, SDS-PAGE and western blot analysis**
Sectioned tissues of potato tuber (150 mg) were frozen in liquid nitrogen and ground into a fine powder using an SK mill (Tokken Inc., Kashiwa, Japan). Then, 400 µl of the protein extraction buffer (50 mM Tris-HCl, pH 6.8, 8 M urea, 4% SDS, 20% glycerol, 1 mM DTT, 0.01% bromophenol blue) was added to the powdered tuber. The mixture was well combined and centrifuged at 20,000 g for 15 min at room temperature. The supernatant solution was collected as a crude protein fraction. This fraction (10 µl) was used for 7.5% SDS-polyacrylamide gel electrophoresis and subjected to western blot analysis using antiserum raised against rice BE1, which was prepared from developing rice seeds (Kawasaki et al. 1996). SDS-PAGE and western blot analysis were performed according to Wakasa et al. 2020. For this analysis, a 5,000-fold dilution of the antiserum was used. Protein interaction with the antibody was detected using the anti-rabbit IgG HPR-linked secondary antibody (GE Healthcare, Chalfont Saint Giles, UK). Signals were detected with Pierce™ ECL Plus western blotting Substrate (Thermo Fishers Scientific, Waltham, MA, USA).

**Results**

**Determination of the nucleotide sequences of the potato SBE3 gene and construction of the CRISPR/Cas9 gene targeted to this gene**
The nucleotide sequence of the potato SBE3 gene of *Solanum phureja* has been registered in the genome database SpudDB (Acc. no. PGSC0003DMG400009981). This gene consists of 15 exons and is predicted to encode a protein containing the transit peptide for translocation to amyloplasts (Van Harsselaar et al. 2017). The region encoding the mature protein started from the third exon. For target mutagenesis to the SBE3 gene of *S. tuberosum*, we selected three regions, gRNA-1, gRNA-2 and gRNA-3, corresponding to the sequences within the third exon of the SBE3 gene (Figure 1A). These regions were used for the generation of CRISPR/Cas9 guide RNAs. In the region of the target sequences, there was a BamHI site, which facilitated the detection of targeted mutations by CAPS analysis (Figure 1A).

It is considered that the nucleotide sequences of the SBE3 genes are largely different depending on the potato variety. In this study, we used the potato cultivar Sayaka, which is commonly used for genome editing. To determine the nucleotide sequence of the SBE3 gene of Sayaka, we analyzed the sequences around the target sites in the SBE3 gene. The corresponding region was PCR-amplified, and the nucleotide sequences were determined. We found that this region contained a polymorphism in three different nucleotide sequences, named WT-A, WT-B, and WT-C (Supplementary Figure S1). Among the sequences, the nucleotide sequence of WT-A was identical to the registered sequence in the database.

In 18 nucleotide sequences, WT-A, WT-B, and WT-C were repeatedly detected 2, 10, and 6 clones, respectively, suggesting that they may exist in a 1 : 2 : 1 proportion.

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Figure 1. Structure of the potato SBE3 gene and location of the gRNAs used for targeted mutagenesis. (A) Schematic representation of the SBE3 gene. Exons are indicated by boxes. Arrows in the upper panel indicate the position of the PCR primers that were used for amplification of the fragment for the CAPS analysis. The positions of the gRNAs are shown in the lower panel. (B) Nucleotide sequences of the regions corresponding to the gRNAs. The upper panel shows the nucleotide sequences of gRNA-1, gRNA-2, and gRNA-3 used for this experiment. The BamHI site is shown in red letters. The lower panel shows the corresponding nucleotide sequences in WT-C. Polymorphic nucleotides found in the WT-C genome are highlighted. PAM sequences are underlined.
in the four alleles contained in the tetraploid genome of Sayaka. Of the three gRNA target sites, a single nucleotide difference was present in each of the gRNAs in WT-C (Figure 1B), whereas no difference was found in those in WT-B.

We have reported that dMac3 significantly enhanced the translation efficiency of downstream ORFs composed of 161 nt derived from the 5′UTR of OsMac3 mRNA (Aoki et al. 2014). Using this system, we established an improved CRISPR/Cas9 vector system, pZD-dxCas9, which enables effective genome editing of the potato GBSS gene (Kusano et al. 2018). For targeted mutagenesis of the potato SBE3 gene, guide RNAs were chemically synthesized and placed between the AtU6-26 promoter and the chimeric single-guide RNA (gRNA) scaffold (Li et al. 2013; Mali et al. 2013) (Figure 2). The resultant gRNA genes were introduced into the pZD-dxCas9 vector, resulting in the formation of CRISPR/Cas9 vectors containing tandemly located gRNA genes, pZD-StSBE3-dxCas9 (Figure 2).

**Creation of a genome edited potato**

Using pZD-StSBE3-dxCas9, the CRISPR/Cas9 gene was introduced into the stems of potato plants. Among the regenerated potato transformants, mutants of the SBE3 gene were investigated. There is a BamHI site in or near the target sites, and therefore, the occurrence of targeted mutations can be detected by disruption of this BamHI site. The DNA fragment containing the target site was PCR-amplified, and CAPS analysis was carried out by BamHI cleavage.

Potato, *S. tuberosum*, has a tetraploid genome; thus, genome-editing events may occur on each of the four alleles. Introducing CRISPR/Cas9 obtained 126 potato transformants. Mutant genes were detected by the occurrence of a BamHI-uncleaved PCR-amplified fragment.
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The frequency of generated mutations was estimated by the ratio of the uncleaved fragments to the amplified fragments. CAPS analysis indicated that 89 (71%) transformants contained mutant genes that exhibited the uncleaved fragment after \textit{BamH}I digestion (Table 1). Among them, ten transformants (8%) were suggested to be mutants that had mutations in all four alleles of the \textit{SBE3} gene (Table 1). This result suggests that site-specific mutation occurred at the target site in the potato cells in planta.

### Table 1. Estimated allele numbers of mutations in the \textit{SBE3} gene in the transformants.

| Number of mutant alleles estimated | Transformants | Number of mutants | Ratio of mutants (%) |
|-----------------------------------|--------------|-------------------|----------------------|
| 0                                 | 37           | 29                |                      |
| 1                                 | 25           | 20                |                      |
| 2                                 | 34           | 27                |                      |
| 3                                 | 20           | 16                |                      |
| 4                                 | 10           | 8                 |                      |
| Total                             | 126          | 100               |                      |

Sequence analysis of the target sites in the mutant alleles of the \textit{SBE3} gene

We analyzed the nucleotide sequences of the representative mutants that were considered to contain four or three mutant alleles. We detected four different nucleotide sequences in the transformed lines #19, #30, #31, #86, and #99, whereas no wild-type nucleotide sequence was detected in these lines, suggesting that they were four-allele mutants of \textit{SBE3} (Figure 4).

Line #19 contained four different types of mutant sequences, which were considered to be derived from WT-A, WT-B, WT-B, and WT-C because these mutant sequences contained single nucleotide polymorphisms corresponding to these genomes. One mutant sequence contained the two deletion sites of 4 nt and 6 nt, resulting in a 10-nt deletion in total. The other sequences contained 8-nt and 2-nt (10-nt in total) deletions, a 55-nt deletion, and a 59-nt deletion. These results showed that line #19 contained mutations in four alleles of the \textit{SBE3} gene (Figure 4). Among them, two sequences were considered derived from the WT-B sequence, which was predicted to have two copies in the genome (Figure 4). These results suggest that frameshift mutations occurred in these genes (Figure 4).

Similarly, frameshift mutations were found in the \textit{SBE3} gene of the mutant lines #30, #31, #86, and #99. These lines also contained four different mutant genes corresponding to each of the three types of \textit{SBE3} gene alleles (Figure 4; Supplementary Figure S2). Thus, it is suggested that these lines contained four-allele mutations of the \textit{SBE3} gene.

Line #76 contained a wild-type nucleotide sequence along with the three mutant \textit{SBE3} sequences, indicating that this transformant was a three-allele mutant (Figure 4). Line #53 contained five mutant sequences in addition to a wild-type sequence (Figure 4). Line #115 showed 8 mutant sequences, but no wild-type sequence was detected (Figure 4; Supplementary Figure S2).

### Analysis of storage starch in the mutant tubers

The mutant lines were cultivated. These plants grew normally and showed a morphology similar to that of the wild type during vegetative growth (Figure 5A). They produced normally shaped tubers similar to the wild-type plant (Figure 5B). Iodine staining assays showed that the tubers of these mutants stained strongly, similar to the wild-type plants (Figure 5C). From these mutant tubers starch was prepared. An analysis of the amylose content revealed a significant reduction in the amylose content of the mutants compared with that of wild-type starch (Figure 5D).

### Detection of \textit{SBE3} in the mutant lines

Potato \textit{SBE3} is considered the homolog of rice BEI (Supplementary Figure S3) and could be immunologically detected by the antiserum against rice BEI. To detect the \textit{SBE3} protein in tubers, we prepared crude protein fractions from the tubers that were the four-allele mutant lines and compared them with those from the wild-type plant. When the rabbit antiserum raised against rice BEI was used for western blot analysis, a protein of approximately 100 kD in molecular size was specifically detected in these lines, suggesting that they were four-allele mutants of \textit{SBE3} (Figure 4).
detected in the protein fraction of the wild-type tuber (Figure 6), suggesting that this protein corresponded to potato SBE3. In the tubers of the four allele mutant lines #19, #31, #99, and #115, no 100-kD protein was detected. This suggests that SBE3 was deficient in these mutants (Figure 6). These results indicated that these mutants

Figure 4. Nucleotide sequences around the target site in the potato SBE3 gene of the representative mutants. The wild-type nucleotide sequence (WT) is shown in the upper panel of the figure. Genome types A, B, and C indicate those from WT-A, WT-B, and WT-C, respectively. Nucleotide sequences containing the outside region are shown in Supplementary Figure S1. The gRNAs (gRNA-1, gRNA-2, and gRNA-3) are indicated above the figure. The region corresponding to the BamHI site is indicated by red letters. Polymorphic nucleotides between three genomes are shown in blue letters. Numbers with the prefix # indicate the individual transformants. Multiple DNA sequences that were detected were aligned. Gaps indicate a nucleotide deletion in the mutants. Nucleotide insertions are indicated in the appended lines. The nucleotide sequences that are indicated by gaps throughout the entire region in #30, #31, #86, #99, and #115 are shown in Supplementary Figure S2. Nucleotide numbers of deletions and insertions are summarized on the right side. Red-colored numbers in the right side showed the frequency of the nucleotide sequences detected.
Discussion

In this study, we created a potato mutant in which the SBE3 gene was disrupted using a modified CRISPR/Cas9 system containing three gRNAs corresponding to the region in the third exon of the SBE3 gene (Figure 1). Genome editing is a newly developed technology using a sequence-specific artificial nuclease to enable modification of a specific gene (Osakabe and Osakabe 2015). Whereas the conventional breeding method takes a long time to obtain a variety possessing a desired trait, genome editing technology may facilitate rapid establishment of an appropriate mutant with a desired trait (Xiong et al. 2015). CRISPR/Cas9 is a powerful tool to induce a mutation at the target site with a high editing efficiency (Ménoret et al. 2013; Remy et al. 2014).

The transformation of potato has been established using a cultivar, Sayaka, that is commonly used for the creation of genetically modified plants as well as genome-edited plants (Kusano et al. 2016, 2018). Because potato has a tetraploid genome, it is necessary to modify all four alleles of the target gene. Previous studies have reported that efficient genome editing is achieved by a modified CRISPR/Cas9 system assembled with the translational enhancer dMac3 along with three guide RNAs derived from a region close to or inside the target gene (Kusano et al. 2018). We used gRNAs that were derived from the region encoding the mature protein downstream of the transition signal region and expected the loss of function of the SBE3 gene in the mutants.

The nucleotide sequences of the genes of S. tuberosum cv. Sayaka have not been registered in the database. The nucleotide sequences of the guide RNAs were taken based on the registered nucleotide sequences of another Solanum species, S. phureja. The structures of the genes in potato cultivars are known to be largely divergent.
Genome edited potato mutant lacking StSBE3

They were the four-allele mutants. Because frameshift no wild-type sequences. These results confirm that mutations in the SBE3 gene. These mutants contained indel mutations in the SBE3 gene. Among them, mutant lines #19, #30, #31, #86, and #99 contained four mutant and no wild-type sequences. These results confirm that they were the four-allele mutants. Because frameshift mutations occurred in these mutant SBE3 genes, these mutant genes were considered to have defective SBE3 activity (Figure 4). In lines #53 and #115, more than four mutant gene sequences were detected (Figure 4), suggesting chimeric mutant lines.

Potato SBE3 is a homolog of rice BEI. The antiserum raised against rice BEI also detected potato SBE3 (Figure 6). Western blot analysis showed defective production of SBE3 in the tubers of four-allele mutants, suggesting that they were mutants with a loss of function of SBE3 (Figure 6).

The rice mutant lacking BEI (sbe1 mutant) shows an aberrant storage starch module, in which the structure of amylopectin is quite different from that of the wild-type strain, as evidenced by an increased degree of iodine staining and alteration of the chain length distribution but no significant difference in the amylose content (Satoh et al. 2003). There could be similarities between the starch trait caused by the potato sbe3 mutant to that caused by the rice sbe1 mutant. The potato sbe3 mutant showed strong iodine staining of the tuber starch along with a significantly reduced amylose content (Figure 5). We have reported that the rice flo2 mutant, which markedly decreased the BE1 activity, showed a decrease in amylose content (Kawasaki et al. 1996). This suggests that properties of starch are influenced by many factors. These results suggest that the potato sbe3 mutant showed an altered storage starch trait as in the rice sbe1 mutant.

To date, there has been no sbe3 mutant potato. The four-allele mutants of the SBE3 gene (sbe3 mutant) that we created grew normally and produced sufficient amounts of tubers with a morphology similar to that of the wild type (Figure 5). Such a mutant potato is expected to contribute determining the functions of the genes involved in starch biosynthesis. It is suggested that the sbe3 mutant may be applied in agriculture as a potato cultivar.

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Figure 6. Detection of potato SBE3 in the tubers. Total protein was prepared from the tubers of the mutant lines and WT. Proteins were separated by SDS-PAGE (lower panel), and SBE3 was detected by western blot analysis using a rabbit antiserum raised against rice BEI (upper panel). An arrow indicates the band corresponding to SBE3. Size markers are shown to the left. WT: wild-type, Numbers with the prefix #: mutant lines
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