The genetic editing of GS3 via CRISPR/Cas9 accelerates the breeding of three-line hybrid rice with superior yield and grain quality

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Abstract Grain size is one of the major traits that determine rice grain yield and quality. The GS3 gene is the first major quantitative trait locus (QTL) that was identified in regulating rice grain length and weight. It was reported that the gs3 allele with a mutation in the organ size regulation (OSR) domain of the GS3 protein produced longer grains. In this study, we used the CRISPR/Cas9 gene editing technology to introduce an edited gs3 allele into our indica maintainer line, Mei1B, to enhance its grain yield and quality. Through molecular analysis and sequencing, a homologous edited-gs3 mutant line without any transgene was obtained in the T1 generation and was named Mei2B. A superior male sterile line Mei2A was generated by backcrossing the cytoplasmic male sterile (CMS) line Mei1A with Mei2B. Mei2B had a higher grain quality and yield compared to its wild-type Mei1B. Its grain length increased by 7.9%, its length/width ratio increased from 3.89 to 4.19, TGW increased by 6.7%, and grain yield per plant increased by 14.9%. In addition, genetic improvement of other quality traits including brown rice length (6.83 mm), brown rice grain length/width ratio (3.61), matched the appearance standards set for traditional Simiao (silk seedling) type cultivars. Two restorer lines were outcrossed to both Mei1A and Mei2A to produce hybrid rice. Compared to two hybrids of Mei1A, the hybrids of Mei2A had longer grains, higher length/width ratio, TGW, and yield per plant. In addition, the hybrids of Mei2A showed a better grain appearance including better translucency, a lower chalky rice rate, and degree of chalkiness than the hybrids of Mei1A. These results demonstrated that the introduction of an elite gs3 allele into Mei1A via CRISPR/Cas9 gene editing technology led to significant genetic improvement of the rice grain. The resultant CMS line Mei2A(gs3) displayed much higher grain quality and yield than the original Mei1A. Therefore, our study demonstrated that the targeted genetic improvement via gene editing
technology can enhance rice breeding, especially the breeding of three-line hybrid rice.

Keywords Grain size · CRISPR/cas9 · Yield · Grain appearance · Three-line hybrid rice

Abbreviations
CMS Cytoplasmic male sterility
GH998 Guanghui 998
TGW Thousand grain weight
LC Low chalkiness
HC High chalkiness
HASV High alkali spreading value
WT Wild type
G715 Gui715
FAM V FAM value
HEX V HEX value
LG Long grain
LASV Low alkali spreading value
SG Short grain
AC Amylose content

Introduction

Rice is a staple cereal crop grown in more than 100 countries across the world and supplies calories for 2/3 of the world’s population. The traits related to grain size (GS) including grain length (GL), grain width (GW), grain thickness (GT), and grain length/width ratio are important factors determining rice yield (Sakamoto and Matsuoka 2008). These traits are also closely related to rice quality characteristics such as grain appearance, processing, cooking, and taste (Xu et al. 1993, 2004; Tan et al. 2000; Li et al. 2004; Song et al. 2007; Shomura et al. 2008; Wang et al. 2008; Wang et al. 2015; Huang et al. 2013; Huang and Qian 2017). During grain development, especially the filling process in big and round grain, it is easy to generate chalkiness because of the long transportation route from the back to the abdomen in the grain itself. Therefore, the transparent and high-quality rice grains are mostly produced in slender grain varieties (Gu et al. 2001; Xu et al. 2004). The market demand for high-quality slender rice has increased recently and has led to research focus for breeders to develop high grain length/width ratio varieties or Simiao-type varieties (fan length ≥6.5 mm and rice length/width ratio ≥3.5, T/GDSMM 001–2019) (Fitzgerald et al. 2009; Wang et al. 2012).

Traditional rice breeding methods significantly depend on the breeder’s experience since complex trait selection can be difficult as well as labor and time consuming. Although MAS (marker-assisted selection) breeding technology greatly increases breeding efficiency, it is somewhat limited when a low recombination rate exists and the genetic drags of the target gene/QTL locus are hard to break. Recently, a genome editing tool derived from an adaptive immune mechanism of microorganisms, the CRISPR-Cas9 (clustered regularly interspaced short palindromic repeats/CRISPR-associated protein9) system, has been successfully applied to plants (Li et al. 2013; Nekrasov et al. 2013; Shan et al. 2013). This technology can precisely modify a plant’s own genes without the introduction of foreign ones (Jiang et al. 2013; Liang et al. 2014; Cai et al. 2015; Lawrenson et al. 2015; Iqbal et al. 2016). Through this gene editing technology, desired traits can be quickly introduced into a target variety, thereby significantly improving the breeding efficiency (Zhang et al. 2016; Tang et al. 2017; Soyk et al. 2017; Kuang et al. 2020).

More than 500 QTL genes related to rice grain size have been identified (http://www.ricedata.cn/ Index. HTM). Among them, 19 QTL genes have been cloned in rice (Chen et al. 2020). Of these cloned QTL genes, GS3, qGL3/GL3.1, TGW3/GL3.3, LGY3, and GS9 are major QTLs, and they have negative effects on controlling grain length (Fan et al. 2006; Qi et al. 2012; Zhang et al. 2012; Liu et al. 2018; Xia et al. 2018; Ying et al. 2018; Zhao et al. 2018). Loss-of-function at the N-terminal organ size regulating (OSR) domain of the GS3 protein made rice produce longer grains. It was found that of 78 varieties, about half of the indica varieties and one-tenth of the japonica varieties contained the long grain allele gs3 at the GS3 locus, indicating that this gene has significant potential for rice production (Fan et al. 2006; Mao et al. 2010). Previous studies have shown that the knockout of the GS3 gene can lead to longer grain lines. These studies mainly focused on the oval-shaped japonica varieties for germplasm creation, but a few focused on long-shaped indica rice (Han et al. 2018), especially indica hybrid rice (Li et al. 2016; Shen et al. 2016 and 2017; Chen et al. 2020).

The three-line hybrid rice system that includes a cytoplasmic male sterile line, a maintainer line, and a restorer line has made a significant contribution to food security in China (Yuan and Tang 1999). Until now, the breeding of three-line hybrid rice still faces great challenges because of the low efficiency in creating excellent maintainer lines
and the high cost of seed production. In particular, when traditional cross-selection methods are used to introduce exogenous elite genes into improved maintainer lines and sterile lines, this is accompanied with the risk of introgressive major or minor restoring genes, which can significantly decrease the efficiency of hybrid rice breeding (Ren et al. 2016). Aimed to test whether the introduction of a gs3 allele into an indica CMS (cytoplasmic male sterile) line Mei1A could improve the grain quality and yield of its hybrids, we focused on editing the GS3 gene of this maintainer line Mei1B using CRISPR/Cas9 technology in this study. Mei1A is a derivative of an elite CMS line of MeiA, whose hybrid varieties are used to dominate the rice industry at the beginning of this century in Guangxi, China, because of the small grain size leading to better quality (Liang et al. 2001). Mei1A inherits many favorable traits from MeiA, including a high yield potential and high GCA (general combination ability). However, due to the disadvantage of its grain length, it is hard to use Mei1A as a hybrid parent to breed a competitive hybrid rice to match the current rice market demand for longer and slender (Simao-type) grains and better taste. In our research, this results showed that the introduction of a gs3 allele into Mei1A via CRISPR/Cas9 technology did improve grain quality and yield of Mei1A’s mutant (Mei2A) and its hybrids. Such a genetic improvement system could rapidly establish superior parental lines from small grain to long grain varieties, thus laying a foundation for sustainable utilization of existing resources.

Materials and methods

Plant materials

Plant materials included an indica maintainer line Mei1B, and its corresponding sterile line Mei1A and the relationship of Mei1B and Mei1A were shown in Fig. 2a. Restorer lines Guanghui 998 (GH998) and Gui 715(G715) were used as the test-cross parents. These plant materials were grown in the transgenic isolation greenhouse of Guangxi Academy of Agricultural Sciences during the entire time of the research.

GS3 genotype detection of Mei1B and its target sequence design

To identify whether Mei1B contains an OSR mutation at the N terminus of the GS3 protein thereby introducing a target sequence mutation, two primer pairs GS3-F1/R1 and GS3-F2/R2 (Table S1) were designed near the exons 1 and 2 of GS3 (Os03g0407400) sequence to amplify DNA fragments of the maintainer line Mei1B. The PCR products were isolated on 1% agarose gel and sequenced. The sequencing results were then compared with the GS3 (Os03g0407400) sequence in the NCBI data base. This confirmed that the exon2 of the GS3 gene in maintainer line Mei1B did not contain a C-A base shift in the OSR domain that caused the long grain mutation. To improve the effectiveness of detecting loss-of-function of the GS3 gene, two gRNA target sites were further designed on exons 1 and 2 of the GS3 gene in Mei1B via the website (http://www.rogenome.net/cas-designer/). They were target 1(cctcgagaatccgatctcgcgg) in the exon 1 and target 2 (tgacgcatactgagcagcgtg) in exon 2. The sequences of these two targets were also screened for off-target effects using the BLAST program from the NCBI website. It showed that there were no matched off-target sequences, indicating an extremely low probability of any off-target effects in our experiment.

Plasmid construction and genetic transformation

ECO31I restriction sites were introduced into the target site sequence of GS3 with two primer pairs GS3-Y1+/GS3-Y1- and GS3-B1+/GS3-B1-, respectively (Table S1). The double-stranded DNA fragments were ligated into the vectors pBWA and pBWD by restriction endonuclease enzyme and ligase. The recombinant plasmid CRISPR-Cas9-GS3 is shown in Fig. S1.

To screen the positive plasmid construct, vector primers Yl-R and Pbw2- (Table S1) were used for PCR amplification of the 1100 bp editing element in the recombinant plasmid. Amplified products were verified by sequencing and then transferred into the Agrobacterium EHA105. The recombinant plasmid CRISPR-Cas9-GS3 was transformed into calli of the rice variety Mei1B, according to the Agrobacterium-mediated transformation method. Finally, hygromycin was used to screen T0 transgenic lines. Positive lines were also identified by PCR amplification using a hygromycin-resistant gene primer pair (Hyg-F/ Hyg-R) and GS3 primers GS3-F1/-R1, GS3-F2/-R2 (Table S1). The PCR products were sequenced. The whole transformation experiment was undertaken by the BioRun company in Wuhan, China.
Grain size measurement of T₀ transgenic lines

Seedlings of transgenic lines and wild-type plants were simultaneously planted at our transgenic greenhouse. Ten fully filled uniform grains of a transgenic line/wild-type plant were harvested at maturity stage and measured for grain length using an automatic grain test instrument (Wanshen SC-G, China).

Establishment of the long-grain maintainer lines and sterile lines

To screen transgene-free mutants, Hyg-F/Hyg-R were used to amplify the DNA template of each individual line in the T₁ generation. Additionally, an individual Mei1B line of the T₁ generation which had the homozygous mutation of OSR significantly increased grain length, and no marker gene was named as Mei2B (Fig. 2b) and was chosen to outcross with rice sterile line Mei1A. After twice backcrossing and with selection, one sterile line with the stable long-grain trait was obtained. These new lines were identified by sequencing to confirm that they had the edited gs3 locus and no transgene component and were named as Mei2A (Fig. 2d).

Testing and trait characterization of potential hybrid rice combinations

The widely used restorer line GH998 together with our own long-grain restorer line G715 was used to test the improved long-grain sterile lines together with the original sterile line for hybrid rice combinations. Ten plants from each hybrid combination and their female parents and five plants from each restorer line were used to investigate the panicle length and grain number per panicle. The grain length and grain width of each line were measured on ten fully filled seeds. One hundred grain weight was randomly measured and then converted into 1000 grain weight. Rice grain quality was assessed following the standards set by the Quality for Cooking Rice Variety NY/T 593–2013. At the same time, fluorescent labeled primers were developed via the penta-primer amplification refractory mutation system (PARMS) (Ye et al. 2001; Zhang et al. 2019; Lu et al. 2020), and primers of genes related to rice grain traits such as GS3, Wx, ALK, and Chalk5 were used to identify the relevant parental genotypes (Fan et al. 2003; Li et al. 2014). The primer information is provided in Table S1. Data analyses were performed using the GraphPad Prism 8 software.

Results

Targeted mutation of the GS3 gene generated three edited gs3 loci

The CRISPR/Cas9 construct with two editing targets of the GS3 gene were introduced into the hybrid maintainer line Mei1B, and 12 edited transgenic plants were obtained. DNA sequencing of the GS3 gene from these 12 transgenic lines revealed that biallelic mutations occurred in three transgenic lines, the two lines (P437-2, P437-13) were homozygous at the target 1 of gs3, and the line P437-6 was homologous at the target 2 of gs3 (Fig. 1). The three T₀ transgenic plants were then self-pollinated to produce the T₁ plants. Grain length and grain width of the three transgenic lines (T₁) were next measured and compared with that of the control (the maintainer line, Mei1B). Phenotypic comparison showed that the average grain length of these three homozygous mutant lines (P437-2, P437-6, P437-13) were longer than 10.00 mm, while the average grain length of the control Mei1B was 9.40 mm, indicating a >5% increase of grain length in the three mutant lines (Fig. 1, Table S2). Sequence alignments of these three homozygous mutants against that of the control Mei1B further showed that 1-base pair (bp) insertion occurred at each target site, inducing gene frame shift of GS3 and then resulting in a premature translation termination (Fig. 1a). Thus, we generated three different edited gs3 loci which regulated the long grain trait of the rice studied.

Identification of transgene-free maintainer line Mei2B with homozygous gs3 mutation

The three transgenic T₀ plants, P437-2, P437-6, and P437-13, were homozygotes at the gs3 locus. We then selected transgene-free plants from the three transgenic plant T₁ progenies via PCR amplification of the hygromycin-resistant gene (Fig. S2). The PCR results identified that a transgene-free plant (P437-6–2) with the nonfunctional gs3 allele and without the selection marker gene from the T₁ offsprings of P437-6.
This transgene-free line with the nonfunctional \(gs3\) allele termed as Mei2B (Fig. 2b, c) would be applied to develop a new long-grain sterile line Mei2A (Fig. 2d, e).

New sterile line Mei2A construction through the transgene-free maintainer line Mei2B

The selection process to develop the male sterile line Mei2A is presented in Fig. 2. Firstly, the maintainer Mei2B carrying the edited \(gs3\) loci was backcrossed to the sterile line Mei1A. Then, sterile lines with a homozygous mutation and significantly increased grain length were selected in the BC<sub>1</sub>F<sub>1</sub> generation. The selected line was next crossed with Mei2B to produce the new sterile line Mei2A (Fig. 2d).

Agronomic trait comparisons between Mei2B with the \(gs3\) mutation and the control line Mei1B with the \(GS3\) allele

To determine how the knockout of the \(GS3\) gene influences rice agronomic traits, the major relevant agronomic traits of maintainers Mei1B and Mei2B were investigated, including the grain length, grain width, ratio of grain length to width, panicle length, grain number per panicle, filled grain number per
Fig. 2 Schematic diagrams of the three-line hybrid rice system and process for converting the mutant maintainer line Mei2B into a mutant male sterile line Mei2A. a Diagram of the three-line hybrid system. It consists of a cytoplasmic sterile (CMS) line and a maintainer line and a restorer line. The CMS line carries both the cytoplasmic sterile gene (S) and the nuclear recessive sterile gene (rr) and is used to produce hybrid seed. The maintainer line carries the cytoplasmic fertile gene (N) and the nuclear recessive sterile gene (rr). It is fertile per se. When it is crossed to the CMS line, their hybrids remain sterile. So, the maintainer line is used to maintain the sterility of CMS and produce sterile seeds for hybrid seed production. The restorer line carries the dominant nuclear fertile gene (RR). When it is crossed to a CMS, its dominant nuclear gene (R) restores the fertility of their hybrids. The male sterile line Mei1A(S(rr)) and its corresponding maintainer line Mei1B (N (rr)) have the only difference in fertility. Male sterile lines rely on maintainer lines for reproduction and improvement. b The process to create the new maintainer line Mei2B carrying homozygous gs3 alleles. c Photographs of the plant, panicle, grain length, and width of Mei1B and Mei2B. d The process to convert the maintainer line Mei2B to superior long-grain sterile line Mei2A. e Photographs of the plant, panicle, and spikelet of Mei1A and Mei2A.
panicle, seed setting rate, 1000 grain weight, effective tiller number, tiller number at the active stage, plant height, and weight per plant (Fig. 3 and Table S3). No significant difference was present between Mei1B and Mei2B in tillering number, grain width, and filled grain number per panicle (Fig. 3b and d). Compared to Mei1B, grain length, 1000 grain weight, and grain number per panicle of Mei2B were increased by 7.9%, 7.7%, and 25.5%, respectively. Although the seed setting rate of Mei2B was lower than Mei1B by 13.6%, its weight per plant was significantly increased by 14.9%. In view of the consistency of agronomic traits between the male sterile lines and their related maintainer lines (apart from, the only difference being in their fertility), this result only presented the histogram of the agronomic trait analysis of the maintainer lines concerned.

Hybrids based on Mei2A carrying the edited gs3 allele had longer grains and yield than hybrids based on Mei1A with GS3 allele.

Four hybrid combinations (Mei2A/GH998, Mei1A/GH998, Mei2A/G715, and Mei1A/G715) were developed by crossing the long-grain sterile line Mei2A and the original sterile line Mei1A with two restorer lines GH998 and G715, respectively. Compared with Mei1A/GH998, the hybrid combination Mei2A/GH998 showed 5.6%, 8.2%, 7.1%, and 15.4% increase in grain length, ratio of grain length to width, 1000 grain weight, and grain weight per plant, respectively. However, there were no significant differences between these two hybrid combinations in grain width, panicle length, plant height, grain number, filled grain number, seed setting rate, and effective tillers (Table S3, Fig. 4). Similarly, there were no significant differences between the combinations of Mei2A/G715 and Mei1A/G715 in grain width, grain number, filled grain number, seed setting rate, plant height, and effective tiller. And Mei2A/G715 presented a significant increase in grain length, ratio of grain length to width, panicle length, 1000 grain weight, and weight per plant than Mei1A/G715 (Fig. 4). These traits were increased by 11.2%, 12.6%, 3.8% and 8.1%, 15.0%, respectively (Table S3, Fig. 4).

Rice quality analysis

Since the grain length and ratio of grain length to width are highly related to the appearance quality of rice grain, its grain length, ratio of grain length to width, translucency grade, chalky rate, and chalkiness degree of the polished rice grain were then

![Fig. 3](image-url) Comparison of agronomic traits between Mei1B with the GS3 allele and Mei2B with the gs3 allele. *, *, and *** indicated the significant difference of studied traits of Mei1B and Mei2B at P levels 0.05, 0.01, and 0.001. Each p value for each trait was obtained from a t test between Mei1B and Mei2B (n=10 for each genotype)
investigated in the hybrid parents and their hybrid combinations (Table S4, Fig. 5). Our results showed that Mei2B(Mei2A) and its combinations (Mei2A/GH998 and Mei2A/G715) had superior or a close appearance quality, compared to Mei1A/Mei1B and its combinations (Mei1A/GH998 and Mei1A/G715) (Fig. 5). In particular, the improved parental line Mei2B(Mei2A, gs3) had a 6.83 mm brown rice grain and the ratio of rice length to width of 3.61, which was close to the Simiao-type standard. We also analyzed the alkali spreading value, gel consistency, and amylose content traits for rice quality (Table S4). Compared with the original Mei1B(Mei1A), the alkali spreading value and gel consistency of the improved line Mei2B(Mei2A) were significantly increased. The two hybrids Mei1A/GH998 and Mei2A/GH998 showed significant difference in amylose content and alkali spreading value (Fig. 5).
In the two G715 combinations, Mei2A/G715 exhibited lower amylose content and higher gel consistency, which indicated that the rice cooking quality of Mei2A/G715 was superior to Mei1A/G715. Although the head rice rate which reflects rice processing quality was decreased slightly in Mei2B(Mei2A) and Mei2A/G715 (Fig. 5), these results indicated that the overall appearance quality and rice qualities of
Mei2B(Mei2A) and its combinations (Mei2A/GH998 and Mei2A/G715) had been improved, especially in Mei2A/G715. In addition, the genotypes of GS3 and three quality-related genes (Wxb, ALK, Chalk5) were also consistent with rice quality testing; see Table 1 and Table S1.

**Discussion**

The traits related to rice yield are important goals in the breeding of new varieties. Previous reports demonstrated that introducing the gs3 gene into rice varieties substantially increased rice grain yield (Li et al. 2016; Shen et al. 2016; Chen et al. 2020). In this study, we found that Mei2B(gs3) had the significant increase in thousand grain weight (TGW) and grain yield per plant, consistent with the previous studies. Even though the seed setting rate of Mei2B(gs3) decreased, this was offset by the increases in grain number per panicle, filled grain per panicle, and TGW, making Mei2B to have a similar yield as Mei1B. The decreased seed setting rate of our improved maintainer line also occurred in previous studies (Shen et al. 2016; Chen et al. 2020). The source-sink-flow theory suggests that the increase in grain length, grain weight, and the total number of grains per panicle could have enhanced the “sink” capacity of the whole grain structure resulting in insufficient photosynthesis in individual plants and thus more empty grains or a lower seed setting rate (Shen et al. 2017). However, in contrast to the hybrids of our original parental line Mei1A(GS3), hybrids from of our improved line Mei2A(gs3) had a similar seed setting rate and an increased grain length, 1000 grain weight, and total yield per plant, thereby ensuring the overall stability of the yield (Table S3). The slender type Simiao or “silk seedling rice” is very popular in Southern China and has a higher market value. Through the introduction of a gs3 allele, the grain length of the improved male sterile line Mei2A aligned with the grain length for Simiao, and this could greatly increase its market competitiveness. Indeed, not only the yield of their F1 was enhanced, but also the rice quality of their F1 reached the highly valued Simiao standard, when Mei2A was combined with a slender restorer G715. In addition, our data also indicated that the edited gs3 allele improved the eating and cooking quality of the male sterile line and its hybrid combinations. Overall, our results demonstrated that the improvement of Mei1A through introducing a gs3 allele via CRISPR/Cas9 technology indeed leads to a better grain appearance and yield in both the male sterile line itself and its hybrid combinations.

Rice quality traits of the combinations are usually determined or influenced by the male sterile line that has been used (Mao et al. 2006). It is important to breed a male sterile line which completely maintains the trait of male sterility. The CRISPR-Cas9-mediated genome editing technology can not only avoid some limitations in traditional crop breeding programs, but it can transfer the desired traits more quickly and accurately in this contemporary crop genetic improvement (Haque et al. 2018; Mishra et al. 2018). In our rice breeding project, the CRISPR/Cas9-mediated genome editing method was used to knock out the GS3 gene in the maintainer line Mei1B, thereby successfully obtaining a gs3 mutant that specifically resulted in a

| Variety | Fluorescence value and Genotype of GS3 | Fluorescence value and Genotype of Wxb | Fluorescence value and Genotype of ALK | Fluorescence value and Genotype of Chalk5 |
|---------|----------------------------------------|----------------------------------------|----------------------------------------|------------------------------------------|
| Mei1B(Mei1A) | GS3(FAM V) | Wxb(Hex V) | ALK(Hex V) | Chalk5(FAM V) |
| Mei2B(Mei2A) | gs3 | Wxb(Hex V) | ALK(Hex V) | Chalk5(FAM V) |
| GH998 | gs3(HEX V) | Wxb(Hex V) | ALK(Hex V) | Chalk5(HEX V) |
| G715 | gs3 (HEX V) | Wxb(Hex V) | ALK(Hex V) | Chalk5(FAM V) |
| Mei1A/GH998 | GS3/gs3 | Wxb | ALK | Chalk5/chalk5 |
| Mei1A/G715 | GS3/gs3 | Wxb | ALK | Chalk5 |
| Mei2A/GH998 | gs3 | Wxb | ALK | Chalk5/chalk5 |
| Mei2A/G715 | gs3 | Wxb | ALK | Chalk5 |
loss of function of the grain size gene. In the T₁ mutant generation of Mei1B, a specific line with an increased grain length and homozygous mutated locus and no transgene was selected to generate the new maintainer line Mei2B. We obtained the new sterile lines Mei2A with the gs3 allele through the improved maintainer line Mei2B and developed new hybrid combinations within 2 years in this study. In contrast, the conventional three-line rice breeding system requires lengthy cycles of backcrossings to evaluate and achieve stable sterile lines. Our study significantly shortened the breeding cycles to release new hybrid rice varieties.

We know that the restoration and conservation relationships of the heterosis within three-line hybrid rice breeding are far more complicated than that of the conventional rice production system, and such complications can restrict the genetic improvements of relevant traits (Zhou 1994; Gong et al. 2020). In our research, a genome editing tool CRISPR-Cas9 was initially adopted to knockout the GS3 gene of the maintainer line, and a molecular marker assisting selection skill was then used to further analyze the exact allelic effect of this target gene and the genotypes of quality-related genes. Detailed genetic information from both phenotyping and genotyping were applied together to guide our selection practices. Consequently, the combination of a molecular marker assisted selection (MAS) with genome editing technology proved to be significantly efficient regarding genetic improvement of the GS3 gene in the three-line hybrid rice production system. However, in this study, we specifically chose a strong restorer line GH998 to further evaluate the restoration effect on the improved male sterile line. Alongside this, another restorer line G715 was also selected and tested to make a long grain hybrid combination. Even though our test combination with the gs3/gs3 allele overall produced a higher yield and quality (Fig. 4 and 5, Table 1). This indicates that the main QTL gene with negative regulation effects should gain more attention when selecting the genotype at the corresponding locus of the restorer lines. The maintainer line with a single gene knockout also had a strong positive restoration relationship with the restorer lines (Fig. 4), ensuring that potential genetic gain on high yield or high general combination ability (GCA) for individual cross combinations tested especially the restorer lines. In the future, joining the gene editing and molecular markers could reduce the challenges posed by phenotyping (Meuwissen et al. 2001; Poland and Rutkoski 2016) and obtain a more accurate genomic prediction in pre-emptive breeding programs (Emebiri et al. 2021).

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Author contribution J H, D Q, and Y P mainly carried out the sequencing, agronomic trait statistics and rice quality analysis. L G carried out the crossing and back-crossing experiments. J H, S L, K L, and G D performed data analysis and manuscript writing. G D and C Z designed the research.

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Availability of data and material The data used in this study are available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate Not applicable.

Consent for publication Not applicable.

Competing interests The authors declare no competing interests.

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References

Cai YP, Chen L, Liu XJ, Sun S, Wu CX, Jiang BJ, Han TF, Hou WS (2015) CRISPR/Cas9-mediated genome editing in soybean hairy roots. PLoS ONE 10:1–13. https://doi.org/10.1371/journal.pone.0136064
Chen YY, Zhu AK, Xue P, Wen XX, Cao YR, Wang BF, Zhang Y, Shah L, Cheng SH, Cao LY, Zhang YX (2020) Effects of GS3 and GL3.1 for grain size editing by CRISPR/Cas9 in rice. Rice Sci 27:405–413. https://doi.org/10.1016/j.rscsci.2019.12.010

Emebiri L, Hildebrand SM, Tan MK, Juliana P, Singh PK, Guenter-Davila G, Singh RP (2021) Pre-emptive breeding against Karnal bunt infection in common wheat: combining genomic and agronomic information to identify suitable parents. Front Plant Sci 12:1–11. https://doi.org/10.3389/fpls.2021.675859

Fan C, Xing Y, Mao H, Lu T, Han B, Xu C, Li X, Zhang QF (2006) GS3, a major QTL for grain length and weight and minor QTL for grain width and thickness in rice, encodes a putative transmembrane protein. Theor Appl Genet 112:1164–1171. https://doi.org/10.1007/s00122-006-0218-1

Fitzgerald MA, McCouch SR, Hall RD (2009) Not just a grain of rice: the quest for quality. Trends Plant Sci 14:133–139

Gao ZY, Zeng DL, Cui X, Zhao YH, Yan MX, Huang DN, Li JY, Qian Q (2003) Map-based cloning of the ALK gene, which controls the gelatinization temperature of rice. Sci China, Ser C Life Sci 46:661–668. https://doi.org/10.1360/03yc0099

Gong R, Huang DQ, Chen YB, Li H, Wang ZD, Zhou DG, Zhao L, Pan YY, Chang YX, Xiang Y, Wang CR, Zhou SC (2020) Comparative metabolomics analysis reveals the variations of eating quality among three high-quality rice cultivars. Mol Breed 40: https://doi.org/10.1007/s11032-020-01192-y

Gu YJ, Xiong F, Wang Z, Chen G, Li WF (2001) A contrast of the endosperm development between rice and wheat. J Nanjing Norm Univ, Nat Sci Ed 24(3): 65–74. (In Chinese with English abstract)

Han Y, Luo DJ, Usman B, Nawaz G, Zhao N, Liu F, Li RB (2018) Development of high yielding gluttonous cytoplasmic male sterile rice (Oryza sativa L.) lines through CRISPR/Cas9 based mutagenesis of Wx and TGW6 and proteomic analysis of anther. Agronomy 8:. https://doi.org/10.3390/agronomy8120290

Haque E, Taniguchi H, Hassan MM, Bhowmik P, KarimM R, Smiech M, Zhao KJ, Islam RM (2018) Application of CRISPR/Cas9 genome editing technology for the improvement of crops cultivated in tropical climates: recent progress, prospects, and challenges. Front Plant Sci 9:1–12. https://doi.org/10.3389/fpls.2018.00617

Hirano HY, Eiiguchi M, Sano Y (1998) A single base change altered the regulation of the waxy gene at the posttranscriptional level during the domestication of rice. Mol Biol Evol 15:978–987. https://doi.org/10.1093/oxfordjournals.molbev.a026013

Huang H, Qian Q (2017) Progress in genetic research of rice grain shape and breeding achievements of long-grain shape and good quality japonica rice. Chinese J Rice Sci 31:665–672. https://doi.org/10.16819/j.1001-7216.2017.7115

Huang HY, Jiang LR, Zheng JS, Wang TS, Wang HC, Huang YM, Hong ZL (2013) Genetic bases of rice grain shape: so many genes, so little known. Trends Plant Sci 18:218–226. https://doi.org/10.1016/j.tpls.2012.11.001

Iqbal Z, Sattar MN, Shafiq M (2016) CRISPR/Cas9: A tool to circumscribe cotton leaf curl disease. Front Plant Sci 7:1–11. https://doi.org/10.3389/fpls.2016.00475

Ishikhi M, Morino K, Nakajima M, Okagaki RJ, Wessler SR, Izawa T, Shimamoto K (1998) A naturally occurring functional allele of the rice waxy locus has a GT to TT mutation at the 5’ splice site of the first intron. Plant J 15:133–138. https://doi.org/10.1046/j.1365-313X.1998.00189.x

Jiang WZ, Zhou HB, Bi HH, Fromm M, Yang B, Weeks DP (2013) Demonstration of CRISPR/Cas9/sgRNA-mediated targeted gene modification in Arabidopsis, tobacco, sorghum and rice. Nucleic Acids Res 41:1–12. https://doi.org/10.1093/nar/gkt780

Kuang Y, Li S, Ren B, Li CD, Qstergaard L, Patron N, Uauy C, Harwood W (2020) Base-editing-mediated artificial evolution of OsALS1 in planta to develop novel herbicide-tolerant rice germplasms. Mol Plant 13:565–572. https://doi.org/10.1007/s13059-020-0101-0

Lawrenson T, Shorinola O, Stacey N, Li CD, Qstergaard L, Patron N, Uauy C, Harwood W (2015) Induction of targeted, heritable mutations in barley and Brassica oleracea using RNA-guided Cas9 nuclease. Genome Biol 16:1–13. https://doi.org/10.1186/s13059-015-0826-7

Li JF, Norville JE, Aach J, McCormack M, Zhang D, Bush J, Church GM, Sheen J (2013) Multiplex and homologous recombination-mediated genome editing in Arabidopsis and Nicotiana benthamiana using guide RNA and Cas9. Nat Biotechnol 31:688–691. https://doi.org/10.1038/nbt.2654

Li JM, Xiao JH, Grandillo S, Jiang LY, Yan ZY, Deng QY, Yuan LP (2004) QTL detection for rice grain quality traits using an interspecific backcross population derived from cultivated Asian (O. sativa L.) and African (O. glaberrima S.) rice. Genome 47:697–704. https://doi.org/10.1139/G04-029

Li M, Li X, Zhou Z, Fang M, Pan X, Lin Q, Luo W, Wu GJ, Li HQ (2016) Reassessment of the four yield-related genes Gln1a, DEP1, GS3, and IPA1 in rice using a CRISPR/Cas9 system. Front Plant Sci 7:1–13. https://doi.org/10.3389/fpls.2016.00377

Li YB, Fan CC, Xing YZ, Fan BC, Xing YZ, Yun P, Luo LJ, Yan B, Peng B, Xie WB, Wang GW, Li XH, Xiao JH, Xu CG, He YQ (2014) Chalk5 encodes a vacuolar H + -translocating pyrophosphatase influencing grain chalkiness in rice. Nat Genet 46:398–404. https://doi.org/10.1038/ng.2923

Li Q, Han R, Wu K et al (2018) G-protein βγ subunits determine grain size through interaction with MADS-domain transcription factors in rice. Nat Commun 9:1–12. https://doi.org/10.1038/s41467-018-03047-9

Mao CX, Wan YZ, Ma GH, Shi YM, Zhou H, Song ZP, Wei SF, Xie LP, Wei SB (2006) Current status analysis of indica CMS line MeiA and its characters analysis. Guangxi Agric Sci 6: 289–291(In Chinese)

Mao CX, Wan YZ, Ma GH, Shi YM, Zhou H, Song ZP, Wei SF, Xie LP, Wei SB (2006) Current status analysis of indica CMS line MeiA and its characters analysis. Guangxi Agric Sci 6: 289–291(In Chinese)

Mol Breeding (2022) 42: 22
Mao HL, Sun SY, Yao JL, Wang CR, Yu SB, Xu CG, Li XH, Zhang QF (2010) Linking differential domain functions of the GS3 protein to natural variation of grain size in rice. Proc Natl Acad Sci U S A 107:19579–19584. https://doi.org/10.1073/pnas.1014491107

Meuwissen THE, Hayes BJ, Goddard ME (2001) Prediction of total genetic value using genome-wide dense marker maps. Genetics 157:1819–1829. https://doi.org/10.1093/genetics/157.4.1819

Mishra R, Joshi RK, Zhao K (2018) Genome editing in rice: recent advances, challenges, and future implications. Front Plant Sci 9: https://doi.org/10.3389/fpls.2018.01361

Nekrasov V, Staskawicz B, Weigel D, Jones JD, Kamoun S (2013) Targeted mutagenesis in the model plant Nicotiana benthamiana using Cas9 RNA-guided endonuclease. Nat Biotechnol 31:691–693. https://doi.org/10.1038/nbt.2655

Poland J, Rutkoski J (2016) Advances and challenges in genomic selection for disease resistance. Annu Rev Phytopathol 54:79–98. https://doi.org/10.1146/annurev-phyto-080615-100056

Qi P, Lin YS, Song XJ, Shen JB, Huang W, Shan JX, Zhu MZ, Jiang LW, Gao JP, Lin HX (2012) The novel quantitative trait locus GL3.1 controls rice grain size and yield by regulating Cyclin-T1.3. Cell Res 22:1666–1680. https://doi.org/10.1038/cr.2012.151

Ren GY, Jin LA, Xie HA (2016) Retrospcetive and perspective on indica three-line hybrid rice breeding research in China. Chin Sci Bull 61(35): 3748–3760. (In Chinese with English abstract)

Sakamoto T, Matsuoka M (2008) Identifying and exploiting grain yield genes in rice. Curr Opin Plant Biol 11:209–214. https://doi.org/10.1016/j.pbi.2008.01.009

Shan QW, Wang YP, Li J, Zhang Y, Chen KL, Liang Z, Zhang K, Liu JX, Xi JJ, Qiu JL, Gao CX (2013) Targeted genome modification of crop plants using a CRISPR-Cas system. Nat Biotechnol 31(8):686–688

Shen L, Wang C, Fu Y, Wang J, Liu Q, Zhang X, Yan C, Qian Q, Wang KJ (2016) QTL editing confers opposing yield performance in different rice varieties. J Integr Plant Biol 60:89–93. https://doi.org/10.1111/1jip.12501

Shen L, Li J, Fu YP, Wang JJ, Hua YF, Jiao ZX, Yan CJ, Wang KJ (2017) Orientation improvement of grain length and grain number in rice by using CRISPR/Cas9 system. Chin J Rice Sci 31(3):223–231 (In Chinese with English abstract)

Shomura A, Izawa T, Ebana K, EbanaK KanegaeH, EbitaniT YM (2008) Deletion in a gene associated with grain size increased yields during rice domestication. Nat Genet 40:1023–1028. https://doi.org/10.1038/ng.169

Song XJ, Huang W, Shi M, Zhu MZ, Lin HX (2007) A QTL for rice grain width and weight encodes a previously unknown RING-type E3 ubiquitin ligase. Nat Genet 39:623–630. https://doi.org/10.1038/ng2014

Soyk S, Muller NA, Park SJ, Schmalenbach I, Jiang K, Hayama R, Zhang L, Van Eck J, Jimenez-Gomez JM, Lippman ZB (2017) Variation in the flowering gene SELF PRUNING 5G promotes day-neutrality and early yield in tomato. Nat Genet 49:162–168. https://doi.org/10.1038/ng.3733

Tan YF, Xing YZ, Li JX, Yu SB, Xu CG, Zhang QF (2000) Genetic bases of appearance quality of rice grains in Shanyou 63, an elite rice hybrid. Theor Appl Genet 101:823–829. https://doi.org/10.1007/s001220051549

Tang L, Mao B, Li Y, Lv QM, Zhang LP, Chen CY, He HJ, Wang WP, Zeng XF, Shao Y, PanYL, Hu YY, Peng Y, Fu XQ, Li HQ, Xia ST, Zhao BR (2017) Knockout of OsNRamp5 using the CRISPR/Cas9 system produces low Cd-accumulating indica rice without compromising yield. Sci Rep 7:1–12. https://doi.org/10.1038/s41598-017-14832-9

Wang E, Wang J, Zhu X, Hao W, Wang LY, Li Q, Zhang LX, He W, Lu BR, Lin HX, Ma H, Zhang GQ, He ZH (2008) Control of rice grain-filling and yield by a gene with a potential signature of domestication. Nat Genet 40:1370–1374. https://doi.org/10.1038/ng.220

Wang S, Li S, Liu Q, Wu K, Zhang JQ, Wang SS, Wang Y, Chen XB, Zhang Y, Gao CX, Wang F, Huang HX, Xu XD (2015) The OsSPL16-GW7 regulatory module determines grain shape and simultaneously improves rice yield and grain quality. Nat Genet 47:949–954. https://doi.org/10.1038/ng.3352

Xia D, Zhou H, Liu R, Dan WH, Li PB, Wu B, Chen JX, Wang LQ, Gao GJ, Zhang QL, He YQ (2018) GL3.3, a novel QTL encoding a GS3/KHAGLY-like kinase, epistatically interacts with GS3 to produce extra-long grains in rice. Mol Plant 11:754–756. https://doi.org/10.1016/j.molp.2018.03.006

Xu ZJ, Chen WF, Ma DR, Lu YN, Zhou SQ, Liu LX (2004) Correlations between rice grain shapes and main qualitative characteristics. Acta Agron Sin 30(9):894–900 (In Chinese with English abstract)

Xu ZJ, Chen WF, Zhang LB (1993) The difference of quality characteristic and their relation, to yield cultivars. J Shenyang Agric Univ 24(3):217–223 (In Chinese with English abstract)

Ye S, Dhillon S, Ke X, Collins AR, Day NM (2001) An efficient procedure for genotyping single nucleotide polymorphisms. Nucleic Acids Res 29:88–88. https://doi.org/10.1093/nar/29.17.e88

Ying JZ, Ma M, Bai C, Huang X, Liu JL, Fan YY, Song XJ (2018) TGW3, a major QTL that negatively modulates grain length and weight in rice. Mol Plant 11:750–753. https://doi.org/10.1016/j.molp.2018.03.007

Yuan LP, Tang CD (1999) Retrospective and current and perspective on hybrid rice breeding. China Rice 1:3–6 (In Chinese)

Zhang B, Zhao N, Liu YY, Jia L, Fu Y, He XX, Liu KF, Xu ZJ, Bao BL (2019) Novel molecular markers for high-throughput sex characterization of Cynoglossus semilaevis. Aquaculture 513:734331. https://doi.org/10.1016/j.aquaculture.2019.734331

Zhong XJ, Wang JF, Huang J, Lan HX, Wang CL, Yin CF, Wu YY, Tang HJ, Qian Q, Li JY, Zhang HS (2012) Rare allele of OsPPKL1 associated with grain length causes extra-large grain and a significant yield increase in rice. Proc Natl Acad Sci U S A 109:21534–21539. https://doi.org/10.1073/pnas.1219776110
Zhang Y, Liang Z, Zong Y, Wang YP, Liu JX, Chen KL, Gao QJL, CX, (2016) Efficient and transgene-free genome editing in wheat through transient expression of CRISPR/Cas9 DNA or RNA. Nat Commun 7:1–8. https://doi.org/10.1038/ncomms12617

Zhao DS, Li QF, Zhang CQ, Zhang C, Yang QQ, Pan LX, Ren XY, Lu J, Gu MH, Liu QQ (2018) GS9 acts as a transcriptional activator to regulate rice grain shape and appearance quality. Nat Commun 9: https://doi.org/10.1038/s41467-018-03616-y

Zhou KL (1994) Breeding of CMS lines in indica hybrid rice. Hybrid Rice 9: 22–26(In Chinese)

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