Application of CRISPR/Cas9 Technology in the Genetic Improvement of Rice Yield and Quality Characters

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Abstract

Rice (*Oryza sativa* L.) is one of the most important food crops in the world. With the continuous increase of global population and the continuous improvement of people's living standards, it has become the focus of attention of rice breeders and botanists to improve the yield and quality of rice. In the new genome editing technology, the CRISPR/Cas system can accurately and specifically edit the target genome region. In recent years, CRISPR/Cas9 technology has made important progress in studying the function of new genes, increasing rice yield and improving rice quality. Therefore, this paper mainly reviewed the application of CRISPR/Cas9 technology in the genetic improvement of rice yield and rice quality in recent years, and looked forward to the application prospect of CRISPR/Cas9 technology in rice genetic breeding, which will provide important reference for the cultivation of new high-quality and high-yield rice varieties.

**Keywords:** rice, CRISPR/Cas9, yield character, quality character, genetic improvement

1. Introduction

Rice (*Oryza sativa* L.) is one of the most important food crops in the world. It is the main food for more than 3 billion people in more than 100 countries and regions (Kusano et al., 2015; Peng et al., 2016a). The world population is expected to reach 10 billion by 2050. At that time, the global demand for agricultural products will increase by about 70%, and the annual grain output will increase from the current 2.1 billion tons to about 3 billion tons (Romero and Gatica-Arias, 2019). With the continuous growth of the world population, the continuous reduction of cultivated land area, the gradual deterioration of ecological environment and other adverse factors, there is a greater risk for the sustainable and stable supply of global food. It is urgent to further improve the yield of rice to ensure global food security (He, 2014). In recent decades, with the continuous improvement of people's living standards, the demand for high-quality rice is also increasing (Peng et al., 2019). Therefore, increasing rice yield and improving rice quality are the focus of attention of rice breeders and botanists.

In recent years, various new technologies and methods have been gradually applied to the genetic improvement of main food crops, especially in the field of gene editing technology. New genome editing techniques have been developed to replace the previous techniques such as random mutagenesis, naturally occurring mutations and classical breeding techniques because these methodologies are time consuming and take too long to obtain individuals with the desired phenotype. New editing technique such as CRISPR/Cas9 has been successfully used in major crops and model plants due to its simplicity, adaptability, and high precision (Ma et al., 2015a Xu et al., 2016). CRISPR/Cas9 system is highly desirable for achieving the goal of rice grain quality improvement with more precision and higher efficiency. Whereas,
the functions of various components of the rice genome ultimately enable the production of higher yielding varieties with better quality rice grains (Xi et al., 2015). In this review, we will focus on the use of CRISPR/Cas technique to improve desirable traits in rice, such as tolerance to biotic and abiotic stresses as well as different parameters involved in determining yield and quality. In addition, we will discuss the problems and prospects of this technique in rice.

2. CRISPR/Cas9 Technology

Gene editing technology is the technology of targeted modification of specific DNA sequence in genome, which includes gene targeting technology established in 1980s and a variety of new and efficient DNA targeting endonuclease technologies developed in recent years, such as ZFNs (zinc finger nucleases), TALEN (transcription activator-like effector nucleases) and CRISPR/Cas9 (clustered regularly interleaved short palindrome repeats) (Doudna et al., 2014; Romero and Gatica-Arias, 2019). In 1987, CRISPR/Cas was accidentally discovered in *E. coli* as an adaptive immune system capable of degrading exogenous DNA (Ishino et al., 1987). Later, it was also reported in other bacteria, including *Shigella dysentery*, *Salmonella enteritidis* and *Mycobacterium tuberculosis* (Nakata et al., 1989). But it wasn't until 2012 that anyone realized the potential of using CRISPR / Cas system for genome editing (Jinek et al., 2012).

DNA targeted endonuclease technology can repair damaged DNA by homologous recombination or non homologous terminal connection by breaking double strands of DNA at the target site of genome, so as to achieve targeted knockout, knockin and gene modification of target site (Peng et al., 2019). The immune memory of CRISPR/Cas system is stored in the form of foreign genome interval sequence inserted into CRISPR array (Koonin et al., 2017). These spacer proteins, together with Cas proteins, serve as a monitoring system for the recognition and degradation of foreign nucleic acids. There are three stages in this process. The first stage, known as adaptive, immune, or spacer acquisition, involves the recognition and integration of foreign DNA (spacer) to CRISPR sites. The viral or plasmid DNA sequence in the spacer is called the primary spacer. Generally, there is a short conservative sequence near the original spacer, which is called the adjacent module of the original spacer (PAM). The second stage is the expression of system. A primary transcript (precrRNA) was transcribed from the CRISPR locus and processed into a small CRISPR RNAs (crRNA). In the final stage, known as interference or immunity, crRNAs, together with trans activated crRNA (tracrRNA), form ribonucleoprotein complexes with Cas proteins. The complex recognizes foreign DNA by base pairing and degrades it (Bhaya et al., 2011). The latest crispr-cas9 technology is based on the reconstruction of the immune system found in bacteria and archaea. Through RNA mediated Cas9 protein to recognize the target DNA sequence and cause DNA double strand breaks. Because of its simple and efficient characteristics, it has been widely used in the process of plant genetic improvement.

There are two essential components of genome editing using CRISPR/Cas system: DNA endonuclease and customizable single RNA (sgRNA) (Figure 1.). Cas9 has a double leaf structure, a large globular recognition leaf (REC) and a small nuclease leaf (NUC). There are
two nuclease domains, RuvC and HNH, which each cut a specific DNA strand. SgRNA is a small non-coding RNA, which is a fusion of crRNA and tracrRNA (Jinek et al., 2012). CRISPR/Cas system can be used to edit any sequence in any organism genome. The only necessary condition is that the target sequence must be adjacent to the PAM sequence. The 5’-NGG-3’ recognition sequence of *Streptococcus pyogenes* Cas9 is PAM. However, there are other Cas proteins with different PAM sequences, such as Cpf1 (or Cas12) which recognize 5’- TTTN-3’ or 5’- TTN-3’ as PAM. In addition, new Cas9 variants (VQR, EQR and VRER) have been developed to identify alternative PAM, thus increasing the possibility of modifying any target sequence in the genome (Anders et al., 2016). The first report describing application of the CRISPR system to rice and wheat plants was published at 2013 (Shan et al., 2013). Since then, the system has been successfully used to edit the plant genome, and has been widely used in the genetic improvement process of rice and other major food crops.

An important feature of CRISPR/Cas9 system is that it can transact on its target, so that it can isolate the mutation agents (CRISPR protein/gene and sgRNA) from the sequence and modify them as required (Ricroch et al., 2017). In order to achieve this, it is necessary to obtain stable production of CRISPR/Cas mutant without CRISPR/Cas expression box in the final plant. This can be achieved by a variety of methods, such as genetic isolation, transient expression of CRISPR components, or transformation of ribonucleoprotein complexes (Pyott et al., 2016; Zhang et al., 2016; Liang et al., 2017). The simplest way to achieve this goal in rice is genetic isolation of transgenic plants in the next generation, which is the most commonly used method to obtain non-transgenic plants. Of course, the transformation of ribonucleoprotein complex can also be realized in rice (Woo et al., 2015). Therefore, compared with other gene editing technologies, CRISPR/Cas gene editing system has the advantages of simple operation and high efficiency, which is widely used in plants.
Figure 1. CRISPR/Cas9 system components and pathways

A. Components required for genome editing using the CRISPR/Cas9 system: a DNA endonuclease (the most commonly used is the Cas9 protein from *Streptococcus pyogenes*) and a customizable single guide RNA (sgRNA). B. Non-homologous end-joining (NHEJ) repair pathway. C. Homologous recombination (HR) repair pathway. PAM, Protopacer adjacent motif (Fernando and Andrés, 2019).

3. Advantages of CRISPR / Cas9 Technology

At present, ZFN (zinc finger nuclease), TALEN (transcription activator-like effector nucleus) and CRISPR/Cas (the clustered regular interspace short palindromic) are the main genomic editing technologies repeat, all of which can precisely replace, insert and knock out plant genome, but ZFN and TALEN technology are relatively time-consuming and laborious (Liu, 2019). Compared with the two technologies, CRISPR/Cas9 technology has the advantages of high editing efficiency, flexibility, low cost and easy operation (Table 1), and has become the mainstream gene editing technology widely used at present (Liu, 2019).
Table 1. Comparison of three gene editing techniques

<table>
<thead>
<tr>
<th>Editing technology</th>
<th>Recognition mode</th>
<th>Vector construction</th>
<th>Advantage</th>
<th>Insufficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZFNs</td>
<td>Specific recognition protein</td>
<td>Complex, need to be redesigned and synthesized</td>
<td>A ZFD can specifically recognize three consecutive nucleotides in DNA strand</td>
<td>Time consuming, expensive, difficult to operate, low specificity and serious Miss problem</td>
</tr>
<tr>
<td>TALENs</td>
<td>Specific recognition protein</td>
<td>Simple, redesigned and synthesized</td>
<td>High specificity, low cost and high efficiency</td>
<td>Time consuming, expensive, cumbersome carrier construction, and low editing efficiency</td>
</tr>
<tr>
<td>CRIDPR/Cas</td>
<td>Homologous recombination or DNA repair</td>
<td>Simple, cutting target DNA complementary and non complementary chains</td>
<td>Editing efficiency is high, flexible, cheap, easy to operate, and exogenous genes are easy to remove</td>
<td>General specificity, obvious Miss effect, limited by PAM recognition site</td>
</tr>
</tbody>
</table>

One of the most attractive features of CRISPR/Cas9 system is its multiple editing ability, which leads to multiple gene mutations in the same transformation event, thus reducing the production time and cost of new plant varieties. This characteristic has been applied to the simultaneous mutation of the whole rice gene family. For example, eight sgRNAs targeting four members of mitogen activated protein kinase gene family were designed. Each pair targets two genomic sites within a single gene site, with an interval of 350-750 bp, so deletion sequences can be observed in the edited cells. A large number of sgRNAs were generated from a single CIS trans gene by using an endogenous tRNA processing system, which precisely cut both ends of the tRNA precursor. After protoplast transformation, the analysis of the target gene showed that the chromosome segments of the four target loci were effectively removed, and the mutation frequency was 4%-20%. When Agrobacterium tumefaciens transformed embryogenic callus, mutations occurred in 8 loci, among which 2 segments were missing (Xie et al., 2015). By using this system, we can obtain mutants with single, double and four genes simultaneously. The mutation efficiency of single mutant is 86% - 100%, that of double mutant is 67% - 100%, and that of four mutant is 86%. The phenotype analysis of the mutants showed that there were severe dwarfing and sterile mpk1 mutants and embryo development defects in the homozygous mpk1 seeds from heterozygous parents. On the contrary, heterozygous mpk6 mutant plants cannot produce homozygous mpk6 seeds at all (Minkenberg et al., 2017). Therefore, CRISPR/Cas9 system can play an important role in the research of rice multi gene mutation.
Another important advantage of CRISPR/Cas9 in the application of major food crops is that in the first generation, transgenic plants can be obtained through the strategy of isolation. In rice, there are several examples of T1 generation plants without transgene mutation (Li et al., 2016a; Wang et al., 2016b). Another way to obtain transgenic plants is to use ribonucleoprotein complex (Cas9 and sgRNAs) for DNA free genome editing. In rice protoplasts, about 20% mutation frequency was obtained by combining Cas9 with sgRNA targeting cytochrome P450 gene (Woo et al., 2015). Therefore, it promotes the application of CRISPR/Cas9 in major food crops and may be further commercialized.

At present, CRISPR/Cas9 is mainly used in plant genome editing, including gene function research and crop genetic improvement. The editing forms can be divided into four aspects: knockout of functional genes, insertion or replacement of genes (fragments), single base editing and gene expression regulation. In order to obtain T1 generation transgenic plants without exogenous genes, a new rice genetic transformation system has been developed. That is to say, using a pair of suicide genes in CRISPR/Cas9 system can effectively kill all pollen and embryos containing CRISPR/Cas produced by T0 plants. One is the gene for bacterial Barney's enzyme, which encodes a toxic protein that kills plant cells. The gene is controlled by REG2 promoter and expressed in early embryonic development. CRISPR/Cas9 system contains CMS2 gene, which encodes rice male gametophyte specific lethal protein under CaMV 35S promoter. In this way, T0 plants and CRISPR/Cas9 systems containing these expression boxes will produce toxic proteins that kill male gametophytes and embryos, and only produce non transgenic seeds. This structure is named CRISPR (TKC), a genetically modified killer (He et al., 2018). In order to test the system, a sgRNA targeting LAZY1 gene was designed, because the LAZY1 mutant has a large tillering angle and can easily identify the mutant individuals. After Agrobacterium tumefaciens transformed rice callus, 65 T0 plants were obtained, 29 of them were lazy type. No TKC or CRISPR/Cas system was found in 10 T0 plants. The target site of each T1 plant contains mutations. These plants are either homozygous or double allelic, showing 100% efficiency of eliminating the target gene (He et al., 2018). Therefore, CRISPR/Cas9 can effectively edit the target gene in plant genome and produce transgenic plants without carrying foreign gene sequence. CRISPR/Cas9 gene editing technology has developed rapidly in just a few years due to its advantages of simple operation, high editing efficiency, support for multi-target editing and various editing forms, and has been widely used in a variety of organisms, making an important contribution to the study of gene function and the inheritance and improvement of important traits of animals and plants. At present, CRISPR/Cas9 technology has also been applied to a variety of plants, including Arabidopsis (Li et al., 2018a), rice (Macovei et al., 2018), wheat (Triticum aestivum) (Okada et al., 2019), maize (Lee et al., 2018), soybean (Glycine max) (Li et al., 2015), sorghum (Che et al., 2018), cotton (Gossypium hirsutum L.) (Li et al., 2018b; Qin et al., 2019), rapeseed (Brassica napus L.) (Braatz et al., 2017), barley (Hordeum vulgare L.) (Lawrenson et al., 2015), Nicotiana benthamiana (Jansing et al., 2018), tomato (Solanum lycopersicum L.) (Ortigosa et al., 2019), potato (Solanum tuberosum) (Enciso-Rodriguez et al., 2019), sweet orange (Citrus sinensis L.) (Peng et al., 2017), cucumber (Cucumis sativus L.) (Chandrasekaran et al., 2016), wild cabbage (Brassica oleracea L.) (Lawrenson et al., 2015), wild legume (Lotus japonicus L.) (Wang et al., 2016a), lettuce (Lactuca sativa L.)
(Woo et al., 2015), *Medicago truncatula* (Michno et al., 2015), *Marchantia polymorpha* (Sugano et al., 2014), tobacco (*Nicotiana tabacum* L.) (Baltes et al., 2014), *Nicotiana attenuate* (Woo et al., 2015), Petuniahybrida (*Nicotiana attenuate*) (Subburaj et al., 2016), grape (*Vitis vinifera* L.) (Wang et al., 2018), apple (*Malus pumila*) (Malnoy et al., 2016), tropical staplcassava (*Manihot esculenta*) (Odipio et al., 2017), watermelon (*Citrullus lanatus*) (Tian et al., 2017), and others. Especially in the process of genetic improvement of important characters in rice, a series of important progress has been made (Table 2).

Table 2. Examples of rice genetic improvement using CRISPR/Cas9

<table>
<thead>
<tr>
<th>Functional gene</th>
<th>Editing mode</th>
<th>Improved trait</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>ERF922</td>
<td>Knockout</td>
<td>Rice blast resistance</td>
<td>Wang et al., 2016c</td>
</tr>
<tr>
<td>SWEET13</td>
<td>Knockout</td>
<td>Bacterial blight resistance</td>
<td>Zhou et al., 2015</td>
</tr>
<tr>
<td>Nramp5</td>
<td>Knockout</td>
<td>Low Cd accumulation</td>
<td>Tang et al., 2017</td>
</tr>
<tr>
<td>SaF/SaM</td>
<td>Knockout</td>
<td>Hybrid compatibility</td>
<td>Xie et al., 2017</td>
</tr>
<tr>
<td>Sc</td>
<td>Knockout</td>
<td>Hybrid compatibility</td>
<td>Shen et al., 2017a</td>
</tr>
<tr>
<td>SITPR/S1A4/S1A6</td>
<td>Knockout</td>
<td>Hybrid compatibility</td>
<td>Xie et al., 2017; Xie et al., 2019</td>
</tr>
<tr>
<td>CSA</td>
<td>Knockout</td>
<td>Reverse-photosensitive sterility</td>
<td>Li et al., 2016b</td>
</tr>
<tr>
<td>TMS5</td>
<td>Knockout</td>
<td>Thermo-sensitive sterility</td>
<td>Zhou et al., 2016</td>
</tr>
<tr>
<td>Hd2/Hd4/Hd5</td>
<td>Knockout</td>
<td>Early maturity</td>
<td>Li et al., 2017</td>
</tr>
<tr>
<td>DEP1/EP3</td>
<td>Knockout</td>
<td>Erect panicle</td>
<td>Li et al., 2016b; Shen et al., 2017b</td>
</tr>
<tr>
<td>Gn1a</td>
<td>Knockout</td>
<td>Increasing grain number</td>
<td>Li et al., 2016a; Shen et al., 2017a</td>
</tr>
<tr>
<td>GS3</td>
<td>Knockout</td>
<td>Increasing grain size</td>
<td>Li et al., 2016b; Shen et al., 2017b</td>
</tr>
<tr>
<td>GW2/GW5/TGW6</td>
<td>Knockout</td>
<td>Increasing grain weight</td>
<td>Shen et al., 2017b; Xu et al., 2016</td>
</tr>
<tr>
<td>SBEIIb</td>
<td>Knockout</td>
<td>High amylose starch</td>
<td>Sun et al., 2017</td>
</tr>
<tr>
<td>Waxy</td>
<td>Knockout</td>
<td>Low amylose starch</td>
<td>Ma et al., 2015a; Zhang et al., 2018</td>
</tr>
<tr>
<td>BADH2</td>
<td>Knockout</td>
<td>Enhancing fragrance</td>
<td>Shen et al., 2017b</td>
</tr>
<tr>
<td>ALS</td>
<td>Replace</td>
<td>Herbicide resistance</td>
<td>Sun et al., 2016</td>
</tr>
<tr>
<td>EPSPS</td>
<td>Replace</td>
<td>Herbicide resistance</td>
<td>Li et al., 2016b</td>
</tr>
<tr>
<td>ACC</td>
<td>Single-base editing</td>
<td>Herbicide resistance</td>
<td>Li et al., 2018c</td>
</tr>
<tr>
<td>SLR1</td>
<td>Single-base editing</td>
<td>Reducing plant height</td>
<td>Lu and Zhu, 2017</td>
</tr>
</tbody>
</table>
4. Application of CRISPR/Cas9 Technology in Rice Yield Genetic Improvement

The yield of rice is mainly determined by the number of effective panicles, the number of grains per panicle and the weight of 1000 grains. Using gene editing technology, we have successfully improved the regulatory genes of rice yield traits, including \( GS3, \ DEPGS5, \ GW2, \ GN1 \) and \( TGW6 \) (Zhang et al., 2017). Li et al. (2018c) used CRISPR/Cas gene editing system to edit \( Gnla \) (OsCKX2), \( DEP1, \ GS3 \) and \( IAP1 \). These genes regulate grain number, ear structure, grain size and plant structure respectively (Li et al., 2016a). Using CRISPR/Cas9 system to edit \( CCD7 \) gene, Butt et al. obtained a multi Tillering Mutant Rice (Butt et al., 2018). Li et al. (2017) used CRISPR/Cas9 mediated multi gene vector to edit three negative growth regulating genes (\( Hd2, \ Hd4 \) and \( Hd5 \)) at the same time, and obtained early maturing rice mutants. The storage characteristics of rice seeds are mainly regulated by lipoxygenase (LOXs). Three genes (\( LOX1, \ LOX2 \) and \( LOX3 \)) in 14 \( LOX \) gene families negatively regulate the storage properties of rice seeds. The application of targeted mutation \( LOX3 \) gene of TALENs improved the storage tolerance of rice seeds (Ma et al., 2015b). Therefore, using CRISPR/Cas9 technology to edit the negative regulatory genes of rice yield can improve the yield of rice.

A method combining pedigree analysis, whole genome sequencing and CRISPR/Cas9 technology can be used to identify genes that have important contribution to yield and other complex quantitative traits on a large scale (Huang et al., Huang 2018). For example, 30 parents and progenies of rice variety IR8 were sequenced, and 57 genes remained in all high-yielding lines were screened by Cas9 or dCas9 gene knockout or knockout for gene editing. The first exon of \( Gn1a \) and \( GS3 \) and the third exon of \( DEP1 \) and \( IAP1 \) were selected as targets of sgRNAs, the mutation frequency of \( Gn1a, \ DEP1, \ GS3 \) and \( IAP1 \) were 42.5%, 67.5%, 57.5% and 27.5% respectively. The plant height, panicle length and the number of flowers per panicle increased by about 90% compared with the control. The results showed that the plant height and ear length of \( dep1 \) mutant decreased by 20% compared with the control, but the number of flowers per ear increased by 50% compared with the control. \( GS3 \) mutant showed larger grain size and longer awn on shell. Finally, mutations in \( IAP1 \) produce three phenotypes based on the nature of the mutations (Li et al., 2016b). SgRNA was designed to mutate miR156 target site. If the mutation does not cause frame shift or change miR156 target, the phenotype is the same as wild type. If the mutation changed the target position of miR156, but did not produce the frame shift of protein, then the mutant plant has 2-4 tillers, less than the wild type 7-10 tillers, but the plant height, flower number and ear length were increased than the wild type.

Rice yield is a complex quantitative trait, which is affected by many factors, including some important agronomic traits and QTLs genes. CRISPR/Cas9 mediated multiple genome editing system is one of the most attractive features of the technology, especially in the editing of QTLs to improve rice yield. Xu et al. (2016) edited three QTLs genes related to grain weight, \( GW2, \ GW5 \) and \( TGW6 \). They found that the functional characteristics of these three genes are good, and the mutation of any of them will lead to a significant increase in grain weight. They used CRISPR/Cas9 system to mutate three grain weight related genes \( GW2, \ GW5 \) and \( TGW6 \) at the same time, and found that rice mutants with 1000 grain weight
increased by 29.3% compared with wild type. In addition, GW5, TGW6 and GW2 mutants had larger grain size than wild type. The length, width and 1000 grain weight of double mutants increased by 12%, 8% and 13%, respectively. The grain length, grain width and 1000 grain weight of the three mutants increased by about 25%, 20% and 28%, respectively. Therefore, CRISPR/Cas9 mediated multiple genome editing system is suitable for the rapid generation and aggregation of beneficial alleles in rice, which is conducive to rice yield.

Using CRISPR/Cas technology to knock out the precise region of the target gene may also improve the yield of rice. Compared with wild type, the deletion of 625 bp in DEP1 gene gave dense erect panicles higher grain number and lower plant height (Huang et al., 2009). With this in mind, Wang et al. (2017) designed four sgRNAs (S1-S4) to achieve the same deletion in indica rice. These four sgRNAs are distributed in four different structures: S1/S4, S1/S3, S2/S4 and S1/S2/S3/S4. In 96 T0 events of each structure, the deletion frequency of 430 bp target is as high as 21%, and that of 10 kb target is as high as 9%. dep1 mutants produced by CRISPR/Cas showed ideal phenotypes (dense erect panicles and reduced plant height). Hence, high yield of rice can be achieved by using CRISPR/Cas9 technology to knock out the specific location of the target gene.

5. Application of CRISPR/Cas9 Technology in Genetic Improvement of Rice Quality

An important goal of rice genetic improvement is to breed new rice varieties with high quality, high yield and high resistance. Generally, rice quality includes appearance quality, nutrition quality, processing quality and cooking and eating quality. Amylose content is the most important index of rice quality for cooking and eating quality. Seeds with high levels of amylose are good sources of resistant starch, a special type of starch that is not digested in the stomach or small intestine but is passed directly to the large intestine. Resistant starch helps improve human health and reduce the risk of non-communicable diseases (Regina et al., 2006). The content of amylose and resistant starch in rice could be increased by changing the branching enzyme (SBEs) of starch. The CRISPR/Cas technology was used to edit the SBEIIb gene expressed in rice grains. After the transformation of rice callus mediated by agrobacterium-mediated, 22 of the 30 transgenic plants mutated in the target region, producing heterozygous, double allelic and homozygous mutants at frequencies of 6.7%, 36.6% and 26.7%, respectively (Sun et al., 2017). No off-target effect was detected in the predicted sites. After the isolation of the non-transgenic mutant lines, the content of amylose and resistant starch in sbeII mutant were increased significantly.

The amylose content in rice seeds is mainly controlled by the Waxy dominant gene, by using the CRISPR/Cas9 technique, we can obtain the mutant of Waxy function deficiency. Through the design of a sgRNA targeting the first exon of Waxy gene, in the callus of two rice varieties (Xiushui 134 和 Wuyunjing 7), T0 transformant had high mutagenesis efficiency (82%~87%) with the help of soil agrobacteria-mediated genetic transformation, and about 4%-15% of the mutants were homozygous (Zhang et al., 2018). Interestingly, the analysis of agronomic traits of T1-generation plants showed that there were no significant differences in plant height, panicle grain number, panicle number per plant, plot yield, grain width, grain length and 1000-grain weight between the wild type and the mutant. But the amylose content of the
Waxy mutant was significantly lower than that of the wild type rice (Zhang et al., 2018). By constructing the expression of CRISPR/Cas9 with pGK03-Wx-gRNA (target 1 and 2 in exon 1 and exon 2 of Wx gene, respectively), the engineering bacteria EHA105 was used to genetically transform the super rice Chujing 27, and 9 independent T₀-generation transformed lines were screened by hygromycin, with 5 lines at target 1 (L1~L5) with mutation frequency of 100% and 4 lines at target 2 (L6~L9) with mutation frequency of 75%. Compared with the wild type, the expression of RNA level Wx gene in mutant lines significantly decreased, and the amylose content in rice significantly decreased (Wang, 2018). Waxy strains of indica rice and japonica rice were obtained by using CRISPR/Cas9 technology to knock out the gene Waxy that controlled amylose content (Ma et al., 2015b). This provides important information for the cultivation of new glutinous varieties of indica rice and japonica rice.

By using CRISPR/Cas9 technology, site-directed editing of PUL gene regulating defunct enzyme in rice was carried out, and the allelic variation of PUL with important breeding value was obtained. By knockout of OsPUL, it was found that the amylose content of T₀-18 decreased by 18.70% and the protein content increased by 22.02% (Feng et al., 2019). Further analysis of mutants revealed that the decrease of GW8 expression in the mutants weakened its inhibition on GL7, which increased the expression of GL7 and led to an increase in grain length. By constructing the knockout expression vector of FAD2-1 gene by CRISPR/Cas9, 6 independent mutant lines were produced by means of agrobacterium-mediated genetic transformation of rice callus (Feng et al., 2019). After the offspring were evaluated to confirm the mutation, the fatty acid spectrum of T₂ series was analyzed (Abe et al., 2018). It was found that in the fad2-1 homozygous mutant, the oleic acid content increased to more than twice that of the wild type, and no linoleic acid was detected. Therefore, the knockout of FAD2-1 gene by CRISPR/Cas9 can produce an increase in oleic acid content in rice seeds.

The field of fragrant rice, favored by the majority of consumers, has a broad market prospect. The CRISPR/Cas9 technology was used to edit the aroma gene OsBADH2 of rice, and the mutant of fragrant rice (Shao et al., 2015). Shao people through the CRISPR/CAS9 technology such as high energy of the fragrance of flowers in 11 Badh2 gene editing, obtained in the derivative of the T₁ generation of a carrier to cut out the skeleton and insert a bases on exon 1 T of mutant materials, the material in the Badh2 significantly lower in the RNA expression level, determination of flavoring substances in results showed that the mutant materials significantly increased (Shao et al., 2017). In addition, the yield and cooking taste quality of wild type and T₂ type rice were examined and analyzed, and it was found that there was no significant difference between the two groups except tiller number and seed setting rate ($P<0.05$). The scent japonica rice varieties farmers 425 as test material, build the two CRISPR/Cas9 Badh2 gene knockout carrier on fixed-point editor, the first carrier (pYLCRISPR/ Cas9-B1-gRNA) of two targets located in exon 2 and 3, respectively the second carrier (pYLCRISPR/ Cas9-B2-gRNA) on the two targets are in exon 2, succeeded T₀ homozygous plants, and the T₀-T₁ generation plants DNA element detection, a total of 8 homozygous mutant strains with different mutation types and without transgenic ingredients were obtained (Sun et al., 2019). At the same time, the rice seeds of these 8 homozygous
mutant lines were tested for aroma. The results showed that the 8 $Badh2$ lines all had different degree of aroma, and the 3 lines edited by pYLCRISPR/Cas9-B1-gRNA were more fragrant.

Using the CRISPR/Cas9 editing technology, in view of water rice protein $GluA$ gene exon 4 of regional design sequence specific gene specificity knockout, gumming point are successfully obtained two mutant genes $GluA3$ materials, these two mutant materials $GluA3$ gene expression levels were significantly lower in RNA, protein translation early termination (Zhou et al., 2019). Among them, the molecular weight of GluA3 protein of M1 mutants changed from 56.03 ku to 51.13 ku, while that of M2 mutants changed to 51.94 ku, leading to the functional change of GluA3 protein in mutants, and the content of GluA3 in rice showed a downward trend. Therefore, the use of CRISPR/Cas9 technology for precise gene editing of important functional genes controlling rice quality traits can achieve targeted functional gene knockout, fixed-point insertion or replacement of genes (fragments), single-base editing and gene expression regulation, and thus achieve genetic improvement of rice quality traits.

6. Problems and Prospects of CRISPR/Cas9 Technology

CRISPR/Cas9 technology has been widely used in many plants because of its advantages such as simple operation, high editing efficiency, support for multi-target editing and various editing forms. It has made an important contribution to the genetic improvement of important characters of rice and other major food crops. Until now, CRISPR/Cas9 system is considered as the best choice of transgenic technology in plant species, but there are still some limitations in practical application (Figure 2). In recent years, in order to improve the editing efficiency and reliability of CRISPR system, a lot of research on the improvement of the system, and new CRISPR/Cas variant have emerged. For example, spCas9-NG, base editing, xCas9, etc. (Manghwar et al., 2019). The key problems and advantages of CRISPR/Cas9 technology are as follows:

1. CRISPR/Cas9 system is large in size and low in editing efficiency, so it is not suitable to be packaged as a virus carrier and transported to somatic tissues. Therefore, CRISPR/Cas9 system is required to have high editing efficiency (Hakim et al., 2019).

2. SpCas9 needs a 5’-NGG-3’ PAM next to a 20-nt DNA target sequence, in which it only recognizes NGG PAM sites, which limits its effectiveness compared with the new CRISPR/Cas variant. However, the xCas9 variant has higher targeting efficiency, higher DNA specificity, lower Miss activity and extensive PAM compatibility (e.g., with NG, GAT and GAA) (Hakim et al., 2019).

3. CRISPR/Cas9 can introduce multiple random non targeted mutations into the genome (Zhang et al., 2015). However, the new CRISPR/Cas variant improves the editing efficiency of the target base in the sequence of interest by identifying different PAMs (Hua et al., 2019). CRISPR/Cas9 introduced mutations at nonspecific sites with similar but not identical homology to the target site.

4. CRISPR/Cas9 needs an Agrobacterium mediated transformation system to produce the mutant, which is not only costly, but also time-consuming. In the future, it is possible to
further improve the editing efficiency of CRISPR system by using genome editing system without tissue culture.

(5) The main problem of industrialization of crops produced by CRISPR/Cas9 technology is the strict supervision and control of genetically modified crops in various countries.

Figure 2. Limitations of current CRISPR/Cas9 system

Abbreviation: Cas9, CRISPR-associated nuclease 9; CRISPR, clustered regularly interspaced short palindromic repeat; HDR, homology-directed repair; PAM, protospacer adjacent motif (Manghwar et al., 2019).

Since CRISPR/Cas genome editing system was first implemented in 2013, with the continuous progress of technology and large-scale sequencing technology, it is possible to create new rice varieties with specific site directed mutations without non targeted mutations (Romero et al., 2019). However, the application of genome editing is still insufficient, which restricts the further application of CRISPR/Cas9 in rice and other crops. Therefore, optimizing gene editing technology can further promote the development and application of crop genetic improvement (Chen et al., 2019). For example, to reduce the requirements of CRISPR genome editing system for PAM sequence; to improve the efficiency of gene replacement editing and reduce the Miss effect; to not directly obtain genome edited rice and other crops in early farmland (Chen et al., 2019). In addition, whether gene editing crops belong to genetically modified organisms has not been clearly determined in the world, and its regulatory standards are controversial (Liu, 2019). Therefore, it is necessary to strengthen the supervision and regulation construction of gene editing crops. It is necessary to organize the scientific community, managers, industry and other communities to discuss, formulate practical laws and regulations, strictly monitor their application scope and degree, and develop the gene editing and breeding technology in accordance with the safety standards.

Genomic editing technology can create genetically modified crops. CRISPR/Cas9 and related genomic editing tools have indeed brought revolutionary changes in rice improvement, which is of great significance to meet and ensure people's demand for high quality and yield of rice
in the future. However, there are still many technical difficulties in CRISPR based plant genome editing technology system that have not been solved yet. To develop a more efficient and accurate gene editing system and optimize mutation detection technology are still the direction of future efforts.

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