Status and Scenario of Genome Editing Device CRISPR-Cas9 in Crop Advancement

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Authors’ contributions

This work was carried out in collaboration among all authors. Authors MSR and ST designed the review concept, literature searches and wrote the first draft of the review article. Author NT, MKT and Sharad Tiwari managed the detail analysis, revision and final compilation of the review article. All authors read and approved the final manuscript.

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ABSTRACT

Plant breeding systems have a extensive antiquity and they had been employed meanwhile the commencement of subjugation of the initial agricultural crop plants. Subsequently, innumerable novel procedures have and are being advanced to added upsurge the profitable worth and harvest of plants. The newest crop enhancement method recognized as genome editing is a method which empowers accurate alteration of the crop genome via bashing out unwanted genes or permitting genes to advance novel occupation. Genome sequencing of many plants and advancement in genome editing methodologies has unlocked prospects to breed advantageous traits. Innovations in genome editing machineries for instance have created it feasible for biotechnologists to mark a specific gene of consideration more proficiently. The first-generation CRISPR/Cas9 genome editing entails modest conniving and cloning approaches, which can be accessible and applicable for various guide RNAs to edit various positions in the genome of targeted organism. It is more willingly recognized in the marketplace economically. The tradition of genome editing has verified to be reimbursements and theatres an encouraging part in upcoming crop development endeavors. So, in presented review article, it is intended to emphasize the advancement and usage of genome editing procedures, in regard, the CRISPR/Cas9 for crop improvement.

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1. INTRODUCTION

CRISPR (Clustered Regularly Interspaced Short Palindromic Repeat) is bacteria immune arrangement through which it can resist against virus attack [1]. Once a virus annexes a bacterium the CRISPR DNA of bacteria produces one or two small RNAs named cRNA and tracer RNA. The produced RNAs predestined to Cas proteins and deliberate campuses that has ability to cut the DNA of the virus with entering intension and these mechanisms defend the bacteria [2]. In 2012, a novel system was discovered to exclusively alter the DNA order of any creature with excessive affluence. The two RNAs, crRNA and tracer RNA pair up with Cas9 protein and directed to the aimed DNA, with the possibility of the corresponding hybridization of bases of the crRNA and the target DNA. Further, Cas9cleaves both the DNA stands [3]. This cleavage happens at extremely specific position that's imposed by the sequence in crRNA molecule. Application of CRISPR-Cas-9 for the purpose of genome editing in plants is one among the leading promptly promising techniques in bioscience since it's becoming manipulator responsive device for advancement of non-transgenic genome editing [4]. CRISPR-Cas-9 is simpler, cost effective, faster, and extremely competent in editing genome even at multiplex level [5]. CRISPR genome editing technology facilitates gene-splicing in a targeted living organism at the position of DNA where replacement, deletion, or insertion took place within the genome. Currently, three types of endonucleases are being applied in the experiments based on plant genome editing. Zinc finger nucleases (ZFNs), transcription activator-like effector nucleases (TALENs), and CRISPR-Cas9 [6-9]. Examples are available on application of ZFN approach for editing lengthly leap of double stranded DNA [10,11].

Genome editing, free from DNA can be done with the help of protoplast-mediated transformation and particle bombardment Woo et al. [12]. Furthermore, Malnoy et al. [13] successfully inserted mutation with the use of CRISPR/Cas9 RNP into protoplasts of grape as well as apple. An effectual, regenerable arrangement from protoplast and somatic embryos [14-16] is pre-requisite. Regrettably, this is not accessible for numerous agriculturally imperative higher plant species [17,18]. The invention of genome editing system especially CRISPR/Cas9 has opened the windows for multidimensional research in animal as well as plants [19]. However, the acronym CRISPR was first devised in 2002 by Jansen et al. [20] and was first examined in the downstream of *Escherichia coli*ap [21]. Further improvements in this field were taken in the year 2005, when the homology of these non-repetitive sequences was identified with foreign DNA sequences obtained from plasmids and phages. Consequently, the mechanism of homology-dependent cleavage was investigated for genome editing and the technology of CRISPR/Cas9 cleavage ‘arrived’ as a capable genome editing tool [22,23].

2. CLUSTERED REGULARLY INTERSpaced PALindromic REPEATS (CRISPR/CAS9) BASED GENOME EDITING METHOD

Different components are involved in the CRISPR cleavage approach. The first component is a 20-nucleotide synthetic gRNA sequence with the ability to bind with target DNA. The second component is Cas9 nuclease enzyme, with the ability to cut 3-4 bases after the protospacer adjacent motif. This Cas9 nuclease consists of two important parts as RuvC-like domains and an HNH domain. Both of these domains have abilities to cut one strand of DNA. The use of CRISPR in plants and animals has been widely documented in more than 5000 articles. The steps involve: (i) identification of specific sequence, (ii) synthesis of a single gRNA (sgRNA), (iii) selection of suitable vector for cloning the sgRNA, (iv) transformation (v) screening of transformants and (vi) final validation. A laboratory with basic facilities for plant transformation can be utilized for genome editing related experiments. This may be due to the straight forward steps in CRISPR/Cas9 mediated (Fig. 1 & Fig.2) genome editing (CMGE). CRISPR/Cas9 approach has proven its efficiency in the last half decade, owing to their ease of use.
Fig. 1. Overview of CRISPR-Cas 9 genome editing

Fig. 2. Illustration of involvement of steps in CRISPR/Cas9 based genome editing

Fig. 3. Presentation of applications of CRISPR/Cas-mediated cis-engineering in plants
3. BENEFITS OF CRISPR-CAS SYSTEM

- Modest cost.
- It doesn’t involve any protein engineering step.
- The simple mechanism of the CRISPR nuclease makes it superior toll for genome editing.
- The main practical advantage is that it may be applied to edit several genes simultaneously.
- Free software exists to guide RNA to focus on any preferred gene.

4. LIMITATIONS

- It is ambiguous, for instance, how specific the guide RNAs are for just the genes they’re alleged to target.
- There are often considerable off-target consequences.
- Non-target DNA resembling the guide RNA can be, activated, or deactivated.
- Release is a huge contest

5. APPLICATION OF CRISPR TECHNIQUE IN CROP IMPROVEMENT

CRISPR/Cas9 gene editing technology is being used in different crops. However, till date most successful experiments have been done in about 20 crop species [24]. These crops were targeted for the improvement of important traits including yield. Development of high yielding, biotic and abiotic stress tolerant/resistant crops is another target. Most of the research papers based on mechanism of knock out specific genes involved in tolerance against abiotic and biotic stresses are available publically. These publications also describe applications of CRISPR/Cas9 in enhancement of tolerance processes. Genome editing approaches demonstrated their significance in the development of biotic stress resistant crops [25]. In this sequence, crops with tolerance against various abiotic stresses like drought and salinity have also been improved with the applications of this technology.

Table 1. List of CRISPR-Cas9 studies for plant improvement

| Crop     | Gene(s) targeted          | Traits targeted          | Obtained results                  | References |
|----------|---------------------------|--------------------------|-----------------------------------|------------|
| Cassava  | Phytoene desaturase       | Trial for CRISPR         | Examination of phenotype          | [90]       |
| Cassava  | elf4E, nCBP-1 and nCBP-2  | CBSD resistance          | Resistance to CBSD                | [91]       |
| Cotton   | CLCuD IR and Rep          | Cotton leaf curl disease | Plants with reduced infection     | [92]       |
| Cotton   | Green fluorescent protein | Phenotypic classification| Insertion of targeted DNA sequence| [93]       |
| Cotton   | GhMYB25-like A and GhMYB25-like D | Fiber development | Fiber development improved | [94]       |
| Rice     | OsSWEET11, OsSWEET14      | Bacterial blight resistance | Bacterial blight resistant plants | [95]       |
| Maize    | Zmzb7                     | Albino related gene      | Albino plant                      | [96]       |
| Maize    | ZmTMS5                    | Male sterility           | Knockout the gene                 | [97]       |
| Maize    | ARGOS8                    | Hybrid development       | Hybrids produced                  | [98]       |
| Rice     | OsERF922                  | Resistance to rice blast | Resistance enhanced               | [99]       |
| Rice     | HTD1, GS3, GW2, GN1A      | Agronomic prospective    | Mutants with improved seed yield  | [100]      |
| Wheat    | TaMLO-A1, TaMLO-B1 and TaMLO-D1 | Resistance to powdery mildew | No infection in edited plants | [101]      |
| Wheat    | TaDREB2 and TaERF3        | Initial trial            | Abiotic stress tolerance          | [102]      |
| Wheat    | TaMLO                     | Resistance against powdery mildew | Resistant plants | [62]       |
| Wheat    | (TaGW2 and TaGASR7)       | Ribonucleoproteins       | Transgene-free plants            | [103]      |
| Soybean  | GmFEI2 and GmSHR          | Hairy root system        | Successful examination of use of hairy root system | [72]       |
| Soybean  | GmFT2a and GmFT4          | Flowering time           | Successfully induced single base substitution | [104]      |
| Groundnut| FAD 2                     | Oleic acid content       | Three mutations identified        | [105]      |
Among genome editing technologies, application of CRISPR-Cas9 have been published by various research groups. In this sequence reports are available on enhancement of metabolic pathways, development of crops against fungal, bacterial, and viral diseases, or cold, drought, and salt stresses (Table 1). Literatures are also available on improvement of nutritionally superior crops with higher yield, and grain quality in various crops. Apart from these, few works are reported on production of haploid seeds. Among these improvements, few may be detailed here as thermo-sensitive genic male sterility in maize [26] and wheat [27], nutritional parameters in sorghum and wheat [28,29], tolerance or resistance against diseases [30,31], and herbicide resistance [32,33].

Due to the simplicity of CRISPR-Cas9 method, it may be applied for the modifications of any targeted gene in plants. This approach provides faster results in comparison to the other approaches. Some of the features of this approach make it superior over others as its simplicity, effectiveness and low cost. This technology is being adopted for the improvement of neglected crops also [34,35]. CRISPR-Cas9 applied to knock down the gene responsible for granule-bound starch synthase (GBSS) in potato plants. It has given tremendous results in just one cycle of transfection with the production of amylopectin starch. The improvement of this trait in potato has given it high commercial values [36]. In cucumber, the activity of eukaryotic translation initiation factor gene elF4E turned off [37]. Engineering for viral and other pathogen resistant in crops have been done successfully [38,39,40]. These experiments demonstrate the efficiency and ability of genome editing technology in the field of crop improvement. This technology is not only suitable for cereals, but also for horticultural and other groups of crops.

6. IMPROVEMENTS IN YIELD AND QUALITY OF CROPS USING CRISPR TECHNOLOGY

**Vegetables**

**Tomato**

Use of genome editing technology has been demonstrated in vegetables including tomato. Application of CRISPR/Cas9 in improvement of tomato was published by Brooks et al. [41]. Other researchers have detailed about applications of genome editing in tomato [42]. Works on use of CRISPR/Cpf1(Cas12a)- a new addition to the CRISPR/Cas genome editing systems [43] are also in progress for the improvement of tomato crop. Apart from this CRISPR/Cpf1 system is now being employed in a wide range of plant species. The utilization of CRISPR/Cpf1 improved genome editing effectiveness in tomato [44]. Use of multiplex sgRNAs given better results instead of single sgRNAs in tomato [45]. CRISPR/Cas9 increased anthocyanin synthesis gene so, that developed tomato plants can be easily identified by their colour [46,47]. For the first time in tomato, two hormone signalling genes DELLA and ETR1 were edited successfully [48]. Edited acetolactate synthase (ALS) gene resulted in chlorsulfuron resistance [49,50]. Since the introduction of genome editing techniques, particularly CRISPR/Cas9, in tomato, multiple traits have been improved [51]. Researchers used CRISPR/Cas9 technology for edition of promoter sequence in tomato [52].

For the first time, food that has been genome-edited using CRISPR–Cas9 technology has been sold on the open market. Since September 2021, Sanatech Seed, based in Tokyo, has been selling Sicilian Rouge tomatoes directly to Japanese consumers, which have been genetically modified to have high quantities of γ-aminobutyric acid (GABA). When taken orally, GABA, according to the company, can help lower blood pressure and enhance relaxation (https://www.nature.com/articles/d41587-021-00026-2).

**Cereals**

**Rice**

New breeding technology such as genome editing has played significant role in the improvement of rice. Rice was the first crop to use the CRISPR/Cas system [53] for its improvement. Most recent improvements in rice [54,55,56] have been published by research groups. CRISPR/Cas9 technology has been applied on genes governing rice grain appearance and quality. Genes including Gnp1a, which affects grain amount, and GS3, which controls grain length in rice, have recently been effectively edited [57]. Edited plants produced longer grain and higher thousand grain weights. In other study, genome editing of GW2, GW5, and TGW6 genes enhanced grain size [58]. Various reports are available on relations of
genes with important characteristics of rice grain. Wang et al. [59] used numerous CRISPR sgRNAs to efficiently remove the segments of the dense and erect panicles in rice. Further, gene that controls grain length were edited to raise thousand kernel weights considerably [60].

**Wheat**

Similar to rice crop, wheat grain length and width were increased by knocking down GW2, that is responsible for grain weight [61]. In an experiment, Shan et al. [62] applied a CRISPR/Cas9 approach to edit the TaMLO gene in wheat (Mildew resistance locus O). The first reported application of CRISPR-Cas9 in wheat was development of resistance to the powdery mildew disease [63]. An experiment was conducted by Wang et al. [61] on utilization of CRISPR/Cas9 for enhancement of seed size in wheat crop. During their experiment they worked on TaGW2, a gene thought to be a negative seed size regulator. In some other experiments, the system has been used to generate low-gluten wheat [64], to increase grain weight [65], for meiotic homologous crossover [66], TaQsd1 gene for postharvest sprouting reduction [67], TaMTL and CENH3 [68] for haploid plant. Furthermore, a specific mutation of TaSBEIIa resulted in high amyllose wheat with significantly greater starch content [69]. These applications of CRISPR/Cas9 in wheat indicate the scope of utilization of genome editing technology to boost wheat yields and grain quality.

**Maize**

Genome editing for dwarf gene in maize was done to obtain semidwarf seedling of maize. CRISPR/Cas9 technology was applied to edit GA20ox3 gene and targeted results were obtained [70]. Second experiment to edit PSY1 gene in maize was done to alter carotenoid production [71]. Development of Fast-Flowering Mini-Maize (FFMM) Lines A and B may help in reduction of generation time. McCaw et al. [72] used an Agrobacterium-mediated conventional transformation technique to successfully introduce CRISPR-Cas9 reagents into immature embryos, resulting in expected transgenic and mutant lines of maize.

**Oilseed crop**

**Soybean**

In the first attempt of genome editing in soybean [73], two DICER-LIKE genes, DCL4a and DCL4b were targeted. Furthermore, two fatty acid desaturase genes, FAD2-1A and FAD2-1B, were altered to generate a high oleic acid soybean variety [74]. Various research groups utilized CRISPR/Cas platform to edit different beneficial genes in soybean. Mutation efficiency was also evaluated in these experiments [75-79]. In an experiment, Du et al. [80] examined the effectiveness of TALENs and CRISPR/Cas9 in modifications of two phytoene desaturase genes (GmPDS11 and GmPDS18) and proved the better performance of CRISPR/Cas9 between both approaches. Furthermore, researchers focused on alteration of a range of GmU6 promoters for driving sgRNA production in soybean and Arabidopsis thaliana. The results revealed that the GmU6-8 and GmU6-10 promoters were the most efficient, resulting in improved editing efficiency [81].

CRISPR-Cas9 method was applied to mutate GmFT2a, a soybean integrator by Cai et al. [82]. However, late flowering was observed in the matured soybean plants with increased vegetative growth. This technology has also improved the profile of soybean seed oil [83] as well as the unpleasant flavour of soybean seed products [84]. Alterations in the content of isoflavones and resistance to soybean mosaic virus [85] have also been achieved after utilization of genome editing technology in soybean.

**Horticultural crops**

**Banana**

There are several examples of utilization of genome editing approaches in improvement of banana crop. Among various beneficial bioactive compounds, Carotenoids affect physiological processes in plants [86]. In this sequence, phytoene desaturase genes i.e. RAS-PDS1 and RAS-PDS2 were modified using CRISPR/Cas9 in banana. The production of ethylene should be considered during development of postharvest preservation technology in banana. Reduction in ethylene production could be very efficient for slowing the ripening process [87]. To achieve these targets, researchers utilized CRISPR/Cas9 for generation of modified MaACO1 plants. Resultant mutant fruits were noticed with decreased ethylene synthesis which has positive effects on slow ripening. These findings suggest that MaACO1 is a good candidate for using the CRISPR/Cas9-mediated editing approach to produce fruit with a longer shelf life. Newly
produced germplasm would significantly reduce postharvest losses and improve the commercial value of the banana sector by prolonging the shelf life of banana fruit [88]. Tripathi et al. [89] recently compiled the known reports on CRISPR/Cas9-based genome editing in banana crop.

**Cassava**

Odipio et al. [90] used CRISPR/Cas9 technology to create MePDS mutants in cassava plants with 22 to 47% success rate. Cassava brown streak disease (CBSD), is a serious problem in East and Central Africa, as well as in West Africa. To overcome this virus CRISPR/Cas9-mediated genome editing was employed to construct ncbp-1, ncbp-2, and ncbp-1/ncbp-2 mutants in a recent experiment done by Gomez et al. [91]. When challenged with CBSV, ncbp-1/ncbp-2 mutants showed delayed and attenuated CBSD aerial symptoms, as well as decreased the intensity and incidence of storage root necrosis. The successful application of genome editing approaches in cassava confirms the possibility of modifications in multiple genes in cassava with disease tolerance/resistance.

**7. PROSPECTS OF CRISPR-CAS9**

- It will be necessary to identify a suitable delivery method for future experiments on crops. The CRISPR/Cas9 system is too vast to be packed into viral vectors due altering its size. Crop improvement will be aided by a small CRISPR system.
- It is important to develop such mechanism that should focus on targeted genes with high efficiency. This will be helpful to achieve the target.
- It is important to gain a better understanding of the hazards associated with this technology. Antimicrobials based on CRISPR-Cas could have unintended consequences: If a strain is eliminated from a population, or if the removal of a specific plasmid affects its growth or metabolism, other, potentially more clinically hazardous species may overtake it.
- During deployment of gene-editing systems in the environment, it is important to exercise on all aspects and dangers. It has been argued that revised nongovernmental rules on the release of CRISPR-Cas and other gene-editing structures that can be used to alter the genetic material of environmental populations, as well as guidance and law from national and international agencies, are needed.
- Adaptation of technically approved guidelines may be followed by community engagement.
- To tackle the off-targeting problem with CRISPR, researchers are now combining the technology with Cas-clover. Cas-CLOVERTM is a revolutionary gene-editing tool that is a true dimeric system, unlike standard single-guided Cas9 and dual-guided Cas9-nickase systems. Cas-CLOVER exhibits great fidelity with no off-targets while keeping high editing efficiency.

**8. CONCLUSION**

Research works on use of genome editing technology to refine the CRISPR/Cas9 protocols in order to make them more user-friendly are being carried out. Researchers are also focused to make this technology widely accessible for research and practical applications in tropical areas, in order to have a stronger impact on agriculture in such places. With the growing use of CRISPR, we will be able to find an alternative to transgenic technology and, hopefully, put an end to the moral difficulties surrounding transgenic crops. New CRISPR Cas-clover approaches are frequently used to address the problem of off-targeting. CRISPR remains an upmarket approach to embrace in India, but its application can still lead to significant agricultural improvement.

**COMPETING INTERESTS**

Authors have declared that no competing interests exist.

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