Applications and potential of genome editing in crop improvement

Yi Zhang1, Karen Massel2, Ian D. Godwin2 and Caixia Gao3,4*

Abstract
Genome-editing tools provide advanced biotechnological techniques that enable the precise and efficient targeted modification of an organism’s genome. Genome-editing systems have been utilized in a wide variety of plant species to characterize gene functions and improve agricultural traits. We describe the current applications of genome editing in plants, focusing on its potential for crop improvement in terms of adaptation, resilience, and end-use. In addition, we review novel breakthroughs that are extending the potential of genome-edited crops and the possibilities of their commercialization. Future prospects for integrating this revolutionary technology with conventional and new-age crop breeding strategies are also discussed.

Introduction
In today’s world, almost one billion people suffer from chronic malnourishment, while at the same time our agricultural systems are degrading, exacerbated by the loss of biodiversity and the increasing uncertainties of climate change [1]. With the global population projected to exceed 9 billion by 2050, contemporary agriculture will face enormous challenges, requiring crops with higher yields and of improved quality, and needing fewer inputs [2]. Although conventional breeding is currently the most widely used approach in crop improvement, it is labor intensive and it usually takes several years to progress from the early stages of screening phenotypes and genotypes to the first crosses into commercial varieties.

Genetically modified (GM) crops that have beneficial traits are produced by the transfer of genes (transgenes) or gene elements of known function into elite crop varieties. Despite the promise that GM crops hold for global food security, their use is affected by largely unsubstantiated health and environmental safety concerns. Government regulatory frameworks that aim to safeguard human and environmental biosafety have led to significant cost barriers to the rapid widespread adoption of new GM traits [3]. As a result, the advantages of GM traits have been restricted to a small number of cultivated crops.

Genome editing is defined as a collection of advanced molecular biology techniques that facilitate precise, efficient, and targeted modifications at genomic loci [4, 5]. Genome editing using zinc-finger nucleases (ZFNs) [6] and transcription activator-like effector nucleases (TALENs) [7] has been around for two decades, but it has recently come under the spotlight through the development of clustered regularly interspaced short palindromic repeats (CRISPR)/Cas systems [8] which provide simplicity and ease of targeted gene editing (Fig. 1a). All of these technologies use typical sequence-specific nucleases (SSNs) that can be induced to recognize specific DNA sequences and to generate double-stranded breaks (DSBs) (Fig. 1a). The plant’s endogenous repair systems fix the DSBs either by non-homologous end joining (NHEJ), which can lead to the insertion or deletion of nucleotides thereby causing gene knockouts, or by homologous recombination (HR), which can cause gene replacements and insertions (Fig. 1a) [9]. Many gene knockout mutants and some gene replacement and insertion mutants have been produced through the use of genome-editing technologies in a wide variety of plants, and many of these mutants have been shown to be useful for crop improvement (Table 1).

The risks involved in altering genomes through the use of genome-editing technology are significantly lower than those associated with GM crops because most edits alter only a few nucleotides, producing changes that are not unlike those found throughout naturally occurring populations [10]. Once the genomic-editing agents have...
segregated out, there is no way to distinguish between a ‘naturally occurring’ mutation and a gene edit. Thus, the introduction of genome editing into modern breeding programs should facilitate rapid and precise crop improvement.

Zinc-finger nucleases

ZFNs are fusions of zinc-finger-based DNA-recognition modules and the DNA-cleavage domain of the FokI restriction enzyme (Fig. 1a). Each individual zinc finger typically recognizes and binds to a nucleotide triplet, and fingers are often assembled into groups to bind to specific DNA sequences [11]. To date, ZFNs have been used to modify Arabidopsis, Nicotiana, maize, petunia, soybean, rapeseed, rice, apple, and fig (reviewed in [12, 13]). In one example of the application of ZFNs to crop breeding, the endogenous maize gene ZmIPK1 was disrupted by insertion of PAT gene cassettes, and this resulted in herbicide tolerance and alteration of the inositol phosphate profile of developing maize seeds [14]. As a proven technology, ZFN-mediated targeted transgene integration was also used for trait stacking in maize, that is for assembling a number of useful traits together to create an even greater potential for crop improvement [15]. Later, Cantos et al. [16] used ZFNs to identify safe regions for gene integration in rice, and these identified sites should serve as reliable loci for further gene insertion and trait stacking. Nevertheless, the design of ZFNs remains a complicated and technically challenging process, and one that often has low efficacy.

Transcription activator-like effector nucleases

Like ZFNs, TALENs are fusions of transcriptional activator-like effector (TALE) repeats and the FokI restriction enzyme (Fig. 1a) [17]. However, each individual TALE repeat targets a single nucleotide, allowing for more flexible target design and increasing the number of potential target sites relative to those that can be targeted by ZFNs. Genome editing by TALENs has been demonstrated in a wide variety of plants including...
| Crop species | Gene editor | Target gene | DNA repair type | Target trait | Reference |
|--------------|-------------|-------------|----------------|-------------|-----------|
| Maize        | ZFNs        | ZmIPK1      | HR             | Herbicide tolerant and phytate reduced maize | [14]      |
| Maize        | ZFNs        | ZmTLP       | HR             | Trait stacking | [15]      |
| Rice         | ZFNs        | OsQQR       | HR             | Trait stacking | [16]      |
| Rice         | TALENs      | OsSWEET14   | NHEJ           | Bacterial blight resistance | [18]      |
| Wheat        | TALENs      | TaMLO       | NHEJ           | Powdery mildew resistance | [19]      |
| Maize        | TALENs      | ZmGL2       | NHEJ           | Reduced epicuticular wax in leaves | [20]      |
| Sugarcane    | TALENs      | COMT        | NHEJ           | Improved cell wall composition | [21]      |
| Sugarcane    | TALENs      | COMT        | NHEJ           | Improved saccharification efficiency | [22]      |
| Soybean      | TALENs      | FAD2-1A, FAD2-1B, FAD3A | NHEJ | High oleic acid contents and low linoleic contents | [23]      |
| Soybean      | TALENs      | FAD2-1A, FAD2-1B, FAD3A | NHEJ | High oleic acid contents | [24]      |
| Potato       | TALENs      | Vnn         | NHEJ           | Minimizing reducing sugars | [25]      |
| Rice         | TALENs      | OsBADH2     | NHEJ           | Fragrant rice | [26]      |
| Maize        | TALENs      | ZmMTL       | NHEJ           | Induction of haploid plants | [27]      |
| *Brassica oleracea* | TALENs | FRIGIDA     | NHEJ           | Flowering earlier | [28]      |
| Tomato       | TALENs      | ANT1        | HR             | Purple tomatoes with high anthocyanin | [29]      |
| Rice         | CRISPR/Cas9 | LAZY1       | NHEJ           | Tiller-spreading | [39]      |
| Rice         | CRISPR/Cas9 | Gn1a, GS3, DEP1 | NHEJ | Enhanced grain number, larger grain size and dense erect panicles | [40]      |
| Wheat        | CRISPR/Cas9 | GW2         | NHEJ           | Increased grain weight and protein content | [41]      |
| Camelina sativa | CRISPR/Cas9 | FAD2       | NHEJ           | Decreased polyunsaturated fatty acids | [42]      |
| Rice         | CRISPR/Cas9 | SBE1b       | NHEJ           | High amyllose content | [43]      |
| Maize        | CRISPR/Cas9 | Wx1         | NHEJ           | High amyllopectin content | [44]      |
| Potato       | CRISPR/Cas9 | Wx1         | NHEJ           | High amyllopectin content | [45]      |
| Wheat        | CRISPR/Cas9 | EDR1        | NHEJ           | Powdery mildew resistance | [46]      |
| Rice         | CRISPR/Cas9 | OsERF922    | NHEJ           | Enhanced rice blast resistance | [47]      |
| Rice         | CRISPR/Cas9 | OsSWEET13   | NHEJ           | Bacterial blight resistance | [48]      |
| Tomato       | CRISPR/Cas9 | SIML02      | NHEJ           | Powdery mildew resistance | [49]      |
| Tomato       | CRISPR/Cas9 | SIJAZ2      | NHEJ           | Bacterial speck resistance | [50]      |
| Grapefruit   | CRISPR/Cas9 | CsLOB1 promoter | NHEJ | Allevitated citrus canker | [51]      |
| Orange       | CRISPR/Cas9 | CsLOB1 promoter | NHEJ | Citrus canker resistance | [52]      |
| Grapefruit   | CRISPR/Cas9 | CsLOB1 promoter | NHEJ | Citrus canker resistance | [53]      |
| Cucumber     | CRISPR/Cas9 | elf4E       | NHEJ           | Virus resistance | [54]      |
| Mushroom     | CRISPR/Cas9 | PPO         | NHEJ           | Anti-browning phenotype | [55]      |
| Tomato       | CRISPR/Cas9 | SP5G        | NHEJ           |Earlier harvest time | [56]      |
| Tomato       | CRISPR/Cas9 | SIAGL6      | NHEJ           | Parthenocarpyn | [57]      |
| Maize        | CRISPR/Cas9 | TMG5        | NHEJ           | Thermosensitive male-sterile | [58]      |
| Rice         | CRISPR/Cas9 | OsMATL      | NHEJ           | Induction of haploid plants | [59]      |
| Tomato       | CRISPR/Cas9 | SP, SP5G, CLV3, WUS, GGP1 | NHEJ | Tomato domestication | [60]      |
| Rice         | CRISPR/Cas9 | ALS         | HR             | Herbicide resistance | [61]      |
| Rice         | CRISPR/Cas9 | ALS         | HR             | Herbicide resistance | [62]      |
| Rice         | CRISPR/Cas9 | ALS         | HR             | Herbicide resistance | [63]      |
| Rice         | CRISPR/Cas9 | ALS         | HR             | Herbicide resistance | [64]      |
Table 1 Crop traits that have been improved by genome-editing techniques (Continued)

| Crop species | Gene editor | Target gene | DNA repair type | Target trait        | Reference |
|--------------|-------------|-------------|-----------------|---------------------|-----------|
| Soybean      | CRISPR/Cas9 | ALS         | HR              | Herbicide resistance | [65]      |
| Maize        | CRISPR/Cas9 | ALS         | HR              | Herbicide resistance | [66]      |
| Potato       | CRISPR/Cas9 | ALS         | HR              | Herbicide resistance | [67]      |
| Flax         | CRISPR/Cas9 | EPSPS       | HR              | Herbicide resistance | [68]      |
| Cassava      | CRISPR/Cas9 | EPSPS       | HR              | Herbicide resistance | [69]      |
| Maize        | CRISPR/Cas9 | ARGOS8      | HR              | Drought stress tolerance | [70]      |

CRISPR clustered regularly interspaced short palindromic repeats, HR homologous recombination, NHEJ non-homologous end joining, TALEN transcription activator-like effector nuclease, ZFN zinc-finger nuclease

Arabidopsis, Nicotiana, Brachypodium, barley, potato, tomato, sugarcane, flax, rapeseed, soybean, rice, maize, and wheat (reviewed in [12, 13]). The first application of TALEN-mediated genome editing in crop improvement was in rice, where the bacterial blight susceptibility gene OsSWEET14 was disrupted and the resulting mutant rice were found to be resistant to bacterial blight [18]. Analogously, TALENs have been used in wheat to knockout three TaMLO homoeologs in order to create powdery mildew-resistant wheat [19]. By knocking out the maize GL2 gene, Char et al. [20] obtained mutants with the glossy phenotype, with reduced epicuticular wax in the leaves and the potential to be surface manured. In sugarcane, cell wall composition and saccharification efficiency have been improved by TALEN-mediated mutagenesis [21, 22].

TALENs can be used to modify the nutritional profiles of crops: soybeans with high oleic acid and low linoleic acid contents were generated by disrupting fatty acid desaturase (FAD) genes, thus improving the shelf life and heat stability of soybean oil [23, 24]. In potato tubers, the accumulation of reducing sugars during cold storage influences the quality of the product, and knocking out the vacuolar invertase (VInv) gene resulted in tubers that had undetectable levels of problematic reducing sugars [25].

Flavor is very important, and fragrant rice has been produced through the use of TALEN technology to disrupt the betaine aldehyde dehydrogenase (BADH2) gene [26]. In addition, the production of haploid plants that inherit chromosomes from only one parent can greatly accelerate plant breeding. Using TALENs to create frame-shift mutations in MATRILINEAL (MTL) in maize, Kelliher et al. [27] obtained haploid maize. Also, by targeting FRIGIDA with TALENs, early-flowering mutants of Brassica oleracea were obtained [28].

Crop improvement by TALEN-mediated gene insertion is well exemplified in the tomato, where incorporating TALENs and donor DNA into geminivirus replicons significantly increased their copy number and hence the efficiency of homologous recombination [29]; a strong promoter was inserted upstream of the gene controlling anthocyanin biosynthesis, and purple tomatoes with high anthocyanin levels were obtained [29]. These examples demonstrate the vast potential of TALEN technology for crop trait improvement. However, the construction of TALE repeats remains a challenge and the efficiency of gene targeting with TALENs is variable.

CRISPR/Cas9 system

CRISPR/Cas systems, especially the type II CRISPR/SpCas9 system from Streptococcus pyogenes, have been developed as versatile genome-editing tools for a wide variety of potential applications (Fig. 1a) [30]. Compared with ZFNs and TALENs, the CRISPR/Cas system is characterized by its simplicity, efficiency, and low cost, and by its ability to target multiple genes [31, 32]. Because of these characteristic features, CRISPR/Cas9 has been rapidly exploited in plants [33–35] and may be an effective solution to a variety of problems in plant breeding [36]. To date, many crops such as rice, maize, wheat, soybean, barley, sorghum, potato, tomato, flax, rapeseed, Camelina, cotton, cucumber, lettuce, grapes, grapefruit, apple, oranges, and watermelon have been edited by this technique (reviewed in [37, 38]). The most frequent application has been in the production of null alleles, or gene knockouts, predominantly achieved by the introduction of small indels that result in frame-shift mutations or by introducing premature stop codons (Fig. 1a).

Yield is a major concern in crop breeding. In rice, when the LAZY1 gene was knocked out by CRISPR/Cas9, a tiller-spreading phenotype was generated, which could increase crop yield under certain circumstances [39]. Li et al. [40] used the CRISPR/Cas9 system to mutate the Gnia, DEPI, and GS3 genes of the rice cultivar Zhonghua11, producing mutants with enhanced grain number, dense erect panicles, and larger grain size, respectively. Grain Weight 2 (GW2) is a key gene in cereal crops, which when disrupted increases grain weight and protein content in wheat [41].

The nutritional profiles of crops can also be improved by CRISPR/Cas9. As in the case of TALEN-mediated knockout in soybean to improve the shelf life and heat stability of soybean oil [23], CRISPR/Cas9 technology
has been used to target FAD2 to improve oleic acid content while decreasing polyunsaturated fatty acids in the emerging oil seed plant Camelina sativa [42]. In rice, Sun et al. [43] used CRISPR/Cas9 technology to generate targeted mutations in SBEIIb, leading to a higher proportion of long chains in amylopectin, which improved the fine structure and nutritional properties of the starch [43]. Using CRISPR/Cas9, DuPont Pioneer (now Corteva Agriscience) knocked out the maize waxy gene Wx1, which encodes the granule-bound starch synthase (GBSS) gene that is responsible for making amylose [44]. In the absence of GBSS expression in the endosperm, amylose was not synthesized, and this created a high amyllopectin (waxy) maize with improved digestibility and the potential for bio-industrial applications [44].

The technology also has been used to improve resistance to biotic stresses. Zhang et al. [46] used CRISPR/Cas9 technology to generate Taedr1 wheat plants by simultaneous modification of the three homoeologs of EDR1. The resulting plants were resistant to powdery mildew and did not show mildew-induced cell death [46]. In rice, enhanced rice blast resistance and bacterial blight resistance were separately obtained by mutagenesis of OsERF922 and OsSWEET13 [47, 48]. Furthermore, powdery mildew-resistant tomatoes were generated by editing SIMLOI [49], and bacterial speck-resistant tomatoes were created by disrupting SlJAZ2 [50]. Citrus canker is a severe disease that is responsible for significant economic losses worldwide, and CsLOB1 is a susceptibility gene for citrus canker. By modifying the CsLOB1 promoter, canker symptoms were alleviated in Duncan grapefruits [51] and Wanjincheng oranges had enhanced resistance to citrus canker [52]. CRISPR/Cas9 technology was later used to disrupt the coding region of CsLOB1 in Duncan grapefruits, resulting in crops that had no canker symptoms [53]. In the cucumber, when the elf4E (eukaryotic translation initiation factor 4E) gene was disrupted, broad virus resistance was generated [54]; the plants were shown to be immune to an Ipomovirus (Cucumber Vein Yellowing Virus) and were resistant to the potyviruses Zucchini yellow mosaic virus and Papaya ring spot mosaic virus-W [54].

Several other traits have been manipulated using CRISPR/Cas9 technologies. Polyphenol oxidase (PPO) is an enzyme that causes browning in many fruits and vegetables. By knocking out this gene, Waltz and coworkers [55] developed a non-browning mushroom. In the tomato, CRISPR/Cas9-engineered mutations in SELF-PRUNING 5G (SPSG) can result in rapid flowering [56], and a mutation in SIAGAMOUS-LIKE 6 (SIAGL6) is responsible for the parthenocarpic phenotype [57]. In maize, when the thermosensitive genic male-sterile 5 gene (TMS5) was knocked out, thermosensitive male-sterile maize was generated [58]. Recently, haploid rice was induced by knockout of OsMATL by CRISPR/Cas9 [59]. Genome-editing techniques can also accelerate the domestication of crops. Using CRISPR/Cas9, Li et al. [60] introduced desirable traits into wild tomato accessions by targeting the coding sequences, cis-regulatory regions, and upstream open reading frames of genes associated with tomato morphology, flower and fruit production, and ascorbic acid synthesis [60].

CRISPR/Cas9-mediated editing by gene insertion and replacement has been used to create herbicide-resistant crops. Herbicide-resistant rice has been developed by a variety of methods such as disrupting DNA ligase 4, which is implicated in NHEJ repair [61], using two single-guide RNAs (sgRNAs) targeting the repair template [62], NHEJ-mediated intron targeting [63], and the use of chimeric single-guide RNAs (sgRNAs) carrying both target site and repair template sequences [64]. When the targeting efficiency was high enough, herbicide-resistant soybean and maize could be created by co-transforming CRISPR/Cas9 and donor DNAs by particle bombardment [65, 66]. Using geminivirus replicons that increase the copy number of CRISPR/Cas9 and a repair template, Butler et al. [67] produced herbicide-resistant potatoes. Moreover, herbicide-resistant flax has been generated using a combination of single-stranded oligonucleotides and CRISPR/Cas9 [68]. Recently, a promoter swap and dual amino-acid substitutions were achieved at the EPSPS locus in cassava, generating glyphosate tolerance [69]. In addition to producing herbicide-resistant crops, CRISPR/Cas9-mediated gene insertion and replacement methods have created drought-resistant properties in maize [70]. The GOS2 promoter confers a moderate level of constitutive expression, and when it was inserted into the 5′-untranslated region of the native ARGOS8 gene, or when it replaced the endogenous ARGOS8 promoter, increased ARGOS8 transcripts were detected and resulted in increased drought tolerance [70].

**Novel technical breakthroughs**

Genome-editing technology already shows great potential in agriculture, but it is still limited by the low efficiency of HR, off-target effects, restrictive protospacer adjacent motif (PAM) sequences, and other issues. Fortunately, novel innovations are continually being added to the genome-editing toolkit to address these limitations.

**Base editing**

To date, HR repair of DSBs using template donor DNA has been found to be much less efficient than template-
free NHEJ, making it difficult to induce single nucleotide substitutions (rather than indels) in plants. However, genome-wide association studies have shown that single-base changes are usually responsible for variations in elite traits in crop plants [71]; hence, efficient techniques for producing precise point mutations in crops are needed urgently. CRISPR/Cas9-mediated base-editing technology is a new genome-editing approach that can accurately convert one DNA base into another, without the use of a DNA repair template [72]. The base-editing technologies employ Cas9 nickase (nCas9) or dead Cas9 (dCas9) fused to an enzyme with base conversion activity. For example, cytidine deaminases convert cytosine (C) to uracil (U), and the latter is treated as thymine (T) in subsequent DNA repair or replication processes, so creating a C→T or T→A substitution (Fig. 1b) [72, 73]. Likewise, adenine deaminases convert adenine (A) to inosine (I), which is treated as guanine (G) by polymerases, creating A→G or G→C substitutions (Fig. 1b) [74]. Cytidine-deaminase-mediated base editing (CBE) has been used in rice, Arabidopsis, wheat, maize, and tomato (reviewed in [75, 76]). Recently, this technology has been used in watermelon and wheat to create herbicide-resistant plants [77, 78].

Adenine-deaminase-mediated base editing (ABE) is more complicated than CBE because no known naturally occurring cytidine deaminases catalyze adenine deamination in DNA rather than RNA. Fortunately, Gaudelli and colleagues [74], using several rounds of directed evolution and protein engineering, were able to develop an efficient ABE. In rice, Yan et al. [79] generated a fluorescence-tracking A to G base editor. Hua et al. [80] also developed an adenine base editor in rice and used it in multiplex base editing. An ABE has also been used with rapeseed protoplasts and in Arabidopsis, and the desired phenotypic alterations and germline transmission were observed in Arabidopsis [81]. Li et al. [82] improved the ABE system for producing edited rice and wheat plants, and generated point mutations within the acetyl-coenzyme A carboxylase (ACC) gene in rice that conferred herbicide-resistance.

In addition to generating point mutations, CBE can also be used to produce nonsense mutations that disrupt genes of interest and knockout their gene functions [83]. CBE is much more specific than conventional SSN-mediated knockout, causing few if any indels. All-in-all, base-editing tools have given genome editing a new dimension, broadening its potential applications by means of nucleotide-specific modifications at specific genomic sites.

**DNA-free genome editing systems**

Conventional genome editing involves the delivery and integration into the host genome of DNA cassettes encoding editing components. Integration occurs at random, and therefore can generate undesirable genetic changes. Even if the DNA cassettes are degraded, the resulting fragments may be integrated and could produce undesirable effects [84]. Prolonged expression of genome-editing tools increases off-target effects in plants since nucleases are abundant in these organisms [19, 26, 85]. Moreover, the introduction of foreign DNA into plant genomes raises regulatory concerns in relation to GM organisms [86]. Therefore, DNA-free genome editing is a groundbreaking technology, producing genetically edited crops with a reduced risk of undesirable off-target mutations, and meeting current and future agriculture demands from both a scientific and regulatory standpoint.

DNA-free genome editing has been accomplished using both protoplast-mediated transformation and particle bombardment. The first successful report of DNA-free genome editing in plants was by Woo and colleagues [87] who transfected CRISPR/Cas9 ribonucleoproteins (RNPs) into protoplasts of Arabidopsis, tobacco, lettuce, and rice. Similarly, Malnoy et al. [88] produced targeted mutations by delivering purified CRISPR/Cas9 RNPs into protoplasts of both grape and apple. Unfortunately, efficient, regenerable protoplast systems are not available for a number of agriculturally important higher crop species, and therefore there has been a search for other DNA-free genome editing methods.

Particle bombardment-mediated DNA-free genome-editing technology has been developed in wheat and maize [89–91]. Both CRISPR/Cas9 RNA and CRISPR/Cas9 RNPs have been delivered into wheat embryos by particle bombardment, and both methods created genome-edited plants [89, 90]. In maize, CRISPR/Cas9 RNPs have been used not only to generate knockout mutants, but also to obtain targeted knockin mutants with the help of single-stranded DNA oligonucleotides [91]. Unlike CRISPR/Cas9 editing with DNA cassettes, CRISPR/Cas9 RNPs cause few if any off-target effects in plants and have a relatively high editing efficiency [90, 91].

Recently, a combination of base editing and DNA-free genome editing has been described in wheat [78], with an average frequency of C-to-T conversion of 1.8%. This development should greatly facilitate both the application of base editing to plant breeding and the commercialization of edited plants.

**CRISPR/Cpf1 system**

The type II CRISPR/SpCas9 system is simple and efficient, but it can only recognize DNA sequences upstream of the appropriate 5′-NGG-3′ PAMs, thus restricting potential target sites. Therefore, Cas9 variants were needed to overcome this limitation. The type V CRISPR/Cpf1 system has demonstrated great potential in this area. Cpf1 recognizes T-rich PAMs and generates cohesive ends with four or five nucleotide overhangs rather than blunt-end breaks,
which complements the characteristics of Cas9 to a large extent (Fig. 1a) [92]. Recently, Cpf1 from *Francisella novicida* (FnCpf1) was used for targeted mutagenesis in tobacco and rice [93], and the Cpf1 ortholog from a Lachnospiraceae bacterium (LbCpf1) generated targeted mutations in rice [94, 95]. A variant AsCpf1 (Cpf1 ortholog from Acidaminococcus sp. BV3L6) demonstrated high genome-editing efficiencies in human cells [96], but was less efficient in rice [97] and in soybean and rice protoplasts [98, 99].

When tested for their ability to induce targeted gene insertions via HR, the FnCpf1 and LbCpf1 nucleases generated precise gene insertions at a target site in rice, at a higher frequency than most other genome-editing nucleases [100]. LbCpf1 has also been used for targeted gene replacement in rice [101]. Recently, to expand the scope of CRISPR/Cpf1-mediated genome editing in rice, Li et al. [102] developed an LbCpf1 (RR) variant that enables the editing and multiplex editing of target genes containing TYCV PAMs.

Like the CRISPR/Cas9 system, the CRISPR/Cpf1 system may be combined with base editing and/or DNA-free genome editing. In fact, CRISPR/Cpf1-mediated DNA-free genome editing has been achieved in rice [98]. As CRISPR/Cpf1-mediated base editing using a T-rich PAM sequence has produced C-to-T conversions in human cells [103], similar applications in crop plants should not be too far in the future.

**Prospects and future directions**

**Multiplexing and trait stacking in crop breeding**

In plants, cellular processes are often regulated by complex genetic networks, and the manipulation of agronomic traits depends on the precise engineering of complex metabolic pathways, which requires the concerted expression of multiple genes. Therefore, molecular tools with the capability to manipulate multiple genes simultaneously are of great value in both basic research and practical applications.

One of the advantages of CRISPR systems over other genome-editing methods is their potential for multiplexing, the simultaneous editing of multiple target sites [31]. Using Golden Gate cloning or the Gibson Assembly method, several groups have assembled multiple sgRNAs into single Cas9/sgRNA expression vectors, in which multiple sgRNAs are driven by separate promoters (reviewed in [104]). Xie et al. [105] have developed a general strategy for producing numerous sgRNAs from a single polycistronic gene. They engineered the endogenous tRNA-processing system, creating a simple and robust platform for expanding the targeting and multiplex editing capability of the CRISPR/Cas9 system. This tRNA-processing system has also been employed for multiplex editing in the CRISPR/Cpf1 system [106]. Cpf1 differs from Cas9 in being a dual nuclease that not only cleaves target DNA but also processes its own CRISPR RNA [107, 108]. Taking advantage of this characteristic, Wang et al. [109] engineered CRISPR/Cpf1 together with a short DR-guide array in rice and demonstrated the feasibility of multiplex gene editing. Multiple sgRNAs can also be used to target a single gene to improve rates of editing in crops that have low transformation or editing efficiencies.

**High-throughput mutant libraries**

Now that the complete genomes of many crops have been sequenced, the challenge of the post-genomic era is to analyze the functions of all crop genes systematically, as most of the genes sequenced to date have unknown functions and may control important agronomic traits. Gene knockout is a frequently used and effective strategy for identifying gene functions; hence, large-scale mutant libraries at the whole-genome level are of great value for functional genomics and for crop improvement.

Genome-wide mutant libraries in rice have been constructed by two teams. Lu et al. [110] designed 88,541 sgRNAs targeting 34,234 genes to create a total of 91,004 targeted loss-of-function mutants. Meng et al. [111] designed 25,604 sgRNAs corresponding to 12,802 genes and obtained more than 14,000 transgenic T0 lines. These two groups selected rice for genome-wide targeted mutagenesis mainly because of its relatively small genome, rich genomic resources, and highly efficient transformation system. As techniques evolve, the construction of mutant libraries in other valued crop species should not be too long delayed.

**Gene regulation**

Besides gene knockouts and knockins, genome editing tools can also be used to regulate gene expression. Gene regulation mainly involves the repression and activation of genes and is often achieved by fusing transcriptional repressors or activators to the DNA-binding domains of genome-editing constructs (such as zinc finger protein (ZFP), TALE, or dCas9), thereby targeting the regulatory regions of endogenous genes [112]. In rapeseed, the VP16 transcriptional activation domain was fused to ZFP, which binds to the DNA sequence downstream of the transcription start site of *KASII* genes. Mutants in which *KASII* was activated displayed the desirable agronomic trait of decreased levels of palmitic acid and total saturated fatty acid [113]. CRISPR/Cas9 can also be used to repress or activate the transcription of plant genes by combining catalytically inactive dCas9 with sgRNAs that target specific promoter sequences [114, 115]. Furthermore, both AsCpf1 and LbCpf1 have been used to repress transcription in *Arabidopsis*, thus underlining the great promise of Cpf1 for modulating plant transcriptomes [99].
Recently, CRISPR/Cas9 technology has been used for crop improvement by altering the cis-regulatory control of quantitative trait loci. Rodriguez-Leal et al. [116] used CRISPR/Cas9 to mutate the SICLV3 promoters in tomato and produced hundreds of regulatory mutations. In this way, they could systematically assess the association of cis-regulatory regions with phenotypic traits, which should be helpful in enhancing tomato breeding. Zhang et al. [117] reported that endogenous plant upstream open reading frames (uORFs) could be edited by CRISPR/Cas9 technology to modulate the translation of mRNAs. Targeting the uORF of *LsGGP2* generated a mutant lettuce with improved tolerance to oxidative stress and increased ascorbate content [117]. This strategy provides a generalizable, efficient method for manipulating the translation of mRNAs, which can be applied to dissect biological mechanisms and improve crops.

Unlike applications that aim primarily to alter DNA sequences, the effects of genome editing on gene regulation act at the transcript level, and could be used to reveal the function of many non-canonical RNAs that are related to crop improvement. As most non-coding transcripts are nuclear and lack open reading frames, genome editing that modulates transcription directly is optimally suited to interrogating the function of such RNAs.

**Fig. 2** a) The network of genome editing methods and the corresponding genome editing tools. b) Flow chart illustrating the successive steps in plant genome editing, and the estimated time needed for each step. RNP ribonucleoprotein, TALEN transcription activator-like effector nuclease, ZFN zinc-finger nuclease
Conclusions
Over the past several decades, traditional breeding that depends on access to plant populations with sufficient variability has made great contributions to agriculture. However, this variability is mainly derived from spontaneous mutations or from mutations that are induced by chemical mutagens or physical irradiation. Such mutations are usually rare and occur at random. Moreover, many types of variation might not arise in elite varieties, and consequently time-consuming, laborious breeding programs are needed to introduce desirable alleles into elite crops. By contrast, genome editing as an advanced molecular biology technique can produce precisely targeted modifications in any crop [4, 5].

In this review, we have described the current applications of three standard genome-editing techniques for crop improvement, and have introduced the relatively new base-editing and CRISPR/Cpf1 systems, which also have great potential in agriculture. Given the availability of a variety of genome-editing tools with different applications (Fig. 2a), it is important to consider the optimal system for a given species and purpose. Once appropriate genome-editing tools have been selected, the target sequences are designed and introduced into the most suitable vectors, and the appropriate genetic cargo (DNA, RNA, or RNPs) for delivery is selected (Fig. 2b). After the genetic cargo has entered the target plant cells, the target sequences will be modified, and edited calli will be regenerated and will ultimately give rise to edited plants (Fig. 2b).

It may well be that protoplast-based systems are not readily available, or even possible, in a species of choice. Furthermore, regeneration by tissue culture may be difficult or limited to a few model genotypes. In these cases, it may be beneficial to design methodologies that do not require regeneration, such as the use of pollen, or the use of immature embryos that can be coaxled to germinate in vitro. With the progress already made in the development of genome-editing tools and the development of new breakthroughs, genome editing promises to play a key role in speeding up crop breeding and in meeting the ever-increasing global demand for food. Moreover, the exigencies of climate change call for great flexibility and innovation in crop resilience and production systems. In addition, we must take into account government regulations and consumer acceptance around the use of these new breeding technologies.

Abbreviations
ABE: Adenine deaminase-mediated base editing; AsCpf1: Cpf1 ortholog from Acidaminococcus sp. BV3L6; CBE: Cytidine deaminase-mediated base editing; CRISPR: Clustered regularly interspaced short palindromic repeats; dCas9: Dead Cas9; DSB: Double-stranded break; FAD: fatty acid desaturase; FnCpf1: Cpf1 from Francisella novicida; GBSS: granule-bound starch synthase; GM: Genetically modified; HR: Homologous recombination; LbcCpf1: Cpf1 from Lachnospiraceae bacterium; NHEJ: Non-homologous end joining; PAM: Protospacer adjacent motif; RNP: Ribonucleoprotein; ssRNA: Single-guide RNA; SSN: Sequence-specific nuclease; TALE: Transcriptional activator-like effector; TALEN: Transcription activator-like effector nuclease; ZFN: Zinc-finger nuclease; ZFP: Zinc finger protein

Acknowledgements
We apologize to those colleagues whose work is not cited due to restrictions on the number of references. We thank Yanpeng Wang (Institute of Genetics and Developmental Biology, CAS) for help with the figures. CG was supported by the National Natural Science Foundation of China (317688103), and YZ by the Shandong Natural Science Foundation (ZR2018BC016).

Authors’ contributions
YZ drafted the first version of the manuscript, KM and IDG revised the manuscript, and CG designed and revised the manuscript. All authors read and approved the final manuscript.

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

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Author details
1Shandong Provincial Key Laboratory of Plant Stress, College of Life Science, Shandong Normal University, Jinan 250014, China. 2The University of Queensland, School of Agriculture and Food Sciences, St Lucia, QLD 4072, Australia. 3The State Key Laboratory of Plant Cell and Chromosome Engineering, and Center for Genome Editing, Institute of Genetics and Developmental Biology, Chinese Academy of Sciences, Beijing 100101, China. 4University of Chinese Academy of Sciences, Beijing 100049, China.

Published online: 30 November 2018

References
1. Foley JA, Ramankutty N, Brauman KA, Cassidy ES, Gerber JS, Johnston M, et al. Solutions for a cultivated planet. Nature. 2011;478:337–42.
2. Tilman D, Balzer C, Hill J, Befort BL. Global food demand and the sustainable intensification of agriculture. Proc Natl Acad Sci U S A. 2011;108:20260–4.
3. Prado JR, Segers G, Voelker T, Carson D, Dobert R, Phillips J, et al. Genetically engineered crops: from idea to product. Ann Rev Plant Biol. 2014;65:769–90.
4. Chen K, Gao C. Targeted genome modification technologies and their applications in crop improvements. Plant Cell Rep. 2013;33:575–83.
5. Gao C. Genome editing in crops: from bench to field. Natl Sci Rev. 2015;2:13–5.
6. Kim YG, Cha J, Chandrasegaran S. Hybrid restriction enzymes zinc finger fusions to FokI cleavage domain. Proc Natl Acad Sci U S A. 1996;93:11560–6.
7. Christian M, Cermak T, Doyle EL, Schmidt C, Zhang F, Hummel A, et al. Targeting DNA double-strand breaks with TAL effector nucleases. Genetics. 2010;186:757–61.
8. Jinek M, Chylinski K, Foiafa J, Hauer M, Doudna JA, Charpentier E. A programmable dual-RNA-guided DNA endonuclease in adaptive bacterial immunity. Science. 2012;337:816–21.
9. Symington LS, Gautier J. Double-strand break end resection and repair pathway choice. Annu Rev Genet. 2011;45:247–71.
10. Voytas DF, Gao C. Precision genome engineering and agriculture: opportunities and regulatory challenges. PLoS Biol. 2014;12:e1001877.
11. Urnov F, Rebar E, Holmes M, Zhang H, Gregory P. Genome editing with engineered zinc-finger nucleases. Nat Rev Genet. 2010;11:636–46.
12. Martinez-Fortün J, Phillips DW, Jones HD. Potential impact of genome editing in world agriculture. Emerg Top Life Sci. 2017;1:117–33.
13. Ran Y, Liang Z, Gao C. Current and future editing reagent delivery systems for plant genome editing. Sci China Life Sci. 2017;60:490–505.
14. Shukla VK, Doyon Y, Miller JC, DeKelver RC, Moehle EA, Worden SE, et al. Precise genome modification in the crop species Zea mays using zinc-finger nucleases. Nature. 2009;459:437–41.
15. Ankel-Werleid S, Saito Y, Welter ME, Murray MG, Zeiter B, Amora R, et al. Trait stacking via targeted genome editing. Plant Biotechnol. 2013;11:1126–34.
16. Cantos C, Francisco P, Trijatmiko KR, Slanet-Loedin I, Chadha-Mohanty PK. Identification of “safe harbor” loci in indica rice genome by harnessing the property of zinc-finger nucleases to induce DNA damage and repair. Front Plant Sci. 2014;5:302.
17. Boch J, Scholze H, Schornack S, Landgraf A, Hahn S, Kay S, et al. Breaking the code of DNA binding specificity of TAL-type III effectors. Science. 2009;326:1509–12.

18. Li T, Liu B, Spalding MH, Weeks DP, Yang B. High-efficiency TALEN-based gene editing produces disease-resistant rice. Nat Biotechnol. 2012;30:390–2.

19. Wang Y, Cheng X, Shan Q, Zhang Y, Liu J, Gao C, Qiu J. Simultaneous editing of three homoalleles in hexaploid bread wheat confers herbicide resistance to powdery mildew. Nat Biotechnol. 2014;32:947–51.

20. Char SN, Unger-Wallace E, Frame B, Briggs SA, Main M, Spalding MH, et al. Heritable site-specific mutagenesis using TALENs in maize. Plant Biotechnol J. 2015;13:1002–10.

21. Jung JH, Altipeter F. TALEN-mediated targeted mutagenesis of the caffeic acid O-methyltransferase in highly polyplaid sugarcane improves cell wall composition for production of bioethanol. Plant Mol Biol. 2016;92:131–42.

22. Kannan B, Jung JH, Mosley GW, Lee SM, Altipeter F. TALEN-mediated targeted mutagenesis of more than 100 COMT copies/alleles in highly polyplaid sugarcane improves saccharification efficiency without compromising biomass yield. Plant Biotechnol J. 2018;16:856–66.

23. Haun W, Coffman A, Clasen BM, Stoddard TJ, Luo S, Ayre E, et al. Improved soybean oil quality by targeted mutagenesis of the fatty acid desaturase 2 gene family. Plant Biotechnol J. 2014;12:934–40.

24. Demorest ZL, Coffman A, Baltes NJ, Stoddard TJ, Clasen BM, Luo S, et al. Direct stacking of sequence-specific nuclease-induced mutations to produce high oleic and low linolenic soybean oil. BMC Plant Biol. 2016;16:225.

25. Clasen BM, Stoddard TJ, Luo S, Demorest ZL, Li J, Cedrone F, et al. Improving cold storage and processing traits in potato through targeted gene knockout. Plant Biotechnol J. 2016;14:169–76.

26. Shan Q, Zhang Y, Chen K, Zhang K, Gao C. Creation of fragrant rice by gene knockout. Plant Biotechnol J. 2016;14:1291–3.

27. Kelliher T, Starr D, Richbourg L, Chintamanani S, Delzer B, Nuccio ML, et al. Multiplex genome editing using Cas9 RNA-guided endonuclease. Nat Biotechnol. 2013;31:691–3.

28. Sun Z, Li N, Huang G, Xu J, Pan Y, Wang Z, et al. Site-specific gene targeting using transcription activator-like effector (TALE)-based nuclease in Brassica oleracea. J Integr Plant Biol. 2013;55:1092–103.

29. Čermák T, Baltes NJ, Čepek R, Gómez-Elvira D, Voytas DF. High-frequency, precise modification of the tomato genome. Genome Biol. 2015;16:232.

30. Hsu PD, Lander ES, Zhang F. Development and applications of CRISPR-Cas9 human genome engineering via Cas9 RNA-guided endonuclease. J Genet Genomics. 2016;43:251–62.

31. Mali P, Yang L, Ersfeld K, Aach J, Guell M, DiCarlo JE, et al. RNA-guided human genome engineering via Cas9. Science. 2013;339:823–30.

32. Li JF, Norville JE, Aach J, McCormack M, Zhang D, Bush J, et al. Multiplex and homologous recombination-mediated genome editing in Arabidopsis and Nicotiana benthamiana using guide RNA and Cas9. Nat Biotechnol. 2013;31:688–91.

33. Nekrasov V, Staskawicz B, Weigel D, Jones JDG, Kamoun S. Breaking the rules of the human genome engineering via Cas9. Science. 2013;339:819–23.

34. Waltz E. Gene-edited CRISPR mushroom escapes US regulation. Nature. 2016;532:293.

35. Nekrasov V, Wang C, Win J, Lanz C, Weigel D, Kamoun S. Rapid generation of a transgene-free powdery mildew resistant tomato by genome deletion. Sci Rep. 2017;7:482.

36. Ortizoa A, Gimenez-Ibanez S, Leonhardt N, Solano R. Design of a bacterial speck resistant tomato by CRISPR/Cas9-mediated editing of S/JAZ2. Plant Biotechnol J. 2018;doi:10.1111/pbi.13006.

37. Ha J, Orbovic V, Jones BJ, Wang N. Modification of the PthA4 effector binding elements in type I Cas9L1 promoter using Cas9/gRNA to produce transgenic Duncan grapefruit alleviating Xcc/PhoA64Cas10L13 infection. Plant Biotechnol J. 2016;14:1291–301.

38. Peng A, Chen S, Lei T, Xu L, He Y, Wu L, et al. Engineering canker resistant plants through CRISPR/Cas9-targeted editing of the susceptibility gene CsLOBL1 promoter in citrus. Plant Biotechnol J. 2017;10:1011–3.

39. Ja H, Zhang Y, Orbovic V, Xu J, White FF, Jones JB, Wang N. Genome editing of the disease susceptibility gene CsLOBL7 in citrus confers resistance to citrus canker. Plant Biotechnol J. 2017;15:817–23.

40. Changdasekaran J, Brumun M, Wolf D, Leibman D, Klapc C, Pearlmann M, et al. Development of broad virus resistance in non-transgenic cucumber using CRISPR/Cas9 technology. Mol Plant Pathol. 2016;17:1140–53.

41. Waltz E. Gene-edited CRISPR mushroom escapes US regulation. Nature. 2016;532:293.

42. Yao L, Zhang Y, Liu C, Liu Y, Wang Y, Liang D, et al. Significant enhancement of fatty acid composition in seeds of the all-hexaploid, Camelina sativa, using CRISPR/Cas9 gene editing. Plant Biotechnol J. 2017;15:648–57.

43. Sun Y, Jiao G, Liu Z, Zhang X, Li J, Guo X, et al. Generation of high-amylose rice through CRISPR/Cas9-mediated targeted mutagenesis of starch branching enzymes. Front Plant Sci. 2017;8:298.

44. Butt H, Eid A, Ali Z, Atia MAM, Mokhtar MM, Hassan N, et al. Efficient targeted mutagenesis in tetraploid potato (Solanum tuberosum) by transient CRISPR-Cas9 expression in protoplasts. Plant Cell Rep. 2017;36:117–28.

45. Zhang Y, Bai Y, Wu G, Zou S, Chen Y, Gao C, Tang D. Simultaneous modification of three homoalleles of TaDRE1 by gene editing enhances powdery mildew resistance in wheat. Plant J. 2017;91:714.

46. Jiang WZ, Henry IM, Lynagh PG, Comai L, Cahoon EB, Weeks DP. Significant enhancement of fatty acid composition in seeds of the all-hexaploid, Camelina sativa, using CRISPR/Cas9 gene editing. Plant Biotechnol J. 2017;15:648–57.

47. Sun Y, Jiao G, Liu Z, Zhang X, Li J, Guo X, et al. Generation of high-amylose rice through CRISPR/Cas9-mediated targeted mutagenesis of starch branching enzymes. Front Plant Sci. 2017;8:298.

48. Peerus. DuPont Announces Intentions to Commercialize First CRISPR-Cas Product. Press Release. 18 April 2018. https://www.pioneer.com/home/site/about-news-media/news-releases/templateCONTENT/guid1D8BF7B11117-9A56-EOB6-3EA6F85AAE92. Accessed 8 Nov 2018.

49. Anderson M, Tureson H, Nicola A, Fält AS, Samuelsson M, Hofvander P. Efficient targeted mutagenesis in tetraploid potato (Solanum tuberosum) by transient CRISPR-Cas9 expression in protoplasts. Plant Cell Rep. 2017;36:117–28.

50. Zhang Y, Bai Y, Wu G, Zou S, Chen Y, Gao C, Tang D. Simultaneous modification of three homoalleles of TaDRE1 by gene editing enhances powdery mildew resistance in wheat. Plant J. 2017;91:714.

51. Jiang WZ, Henry IM, Lynagh PG, Comai L, Cahoon EB, Weeks DP. Significant enhancement of fatty acid composition in seeds of the all-hexaploid, Camelina sativa, using CRISPR/Cas9 gene editing. Plant Biotechnol J. 2017;15:648–57.

52. Sun Y, Jiao G, Liu Z, Zhang X, Li J, Guo X, et al. Generation of high-amylose rice through CRISPR/Cas9-mediated targeted mutagenesis of starch branching enzymes. Front Plant Sci. 2017;8:298.

53. Boch J, Scholze H, Schornack S, Landgraf A, Hahn S, Kay S, et al. Breaking the code of DNA binding specificity of TAL-type III effectors. Science. 2009;326:1509–12.

54. Li T, Liu B, Spalding MH, Weeks DP, Yang B. High-efficiency TALEN-based gene editing produces disease-resistant rice. Nat Biotechnol. 2012;30:390–2.

55. Wang Y, Cheng X, Shan Q, Zhang Y, Liu J, Gao C, Qiu J. Simultaneous editing of three homoalleles in hexaploid bread wheat confers herbicide resistance to powdery mildew. Nat Biotechnol. 2014;32:947–51.

56. Char SN, Unger-Wallace E, Frame B, Briggs SA, Main M, Spalding MH, et al. Heritable site-specific mutagenesis using TALENs in maize. Plant Biotechnol J. 2015;13:1002–10.

57. Jung JH, Altipeter F. TALEN mediated targeted mutagenesis of the caffeic acid O-methyltransferase in highly polyplaid sugarcane improves cell wall composition for production of bioethanol. Plant Mol Biol. 2016;92:131–42.
66. Svitasev S, Young JK, Schwartz C, Gao H, Fakko SC, Cigan AM. Targeted mutagenesis, precise gene editing, and site-specific gene insertion in maize using Cas9 and guide RNA. Plant Physiol. 2015;169:931–45.

67. Butler NM, Baltes NJ, Voytas DF, Douches DS. Geminivirus-mediated genome editing in potato (Solanum tuberosum L.) using sequence-specific nucleases. Front Plant Sci. 2016;7:1045.

68. Sauer NJ, Zrnčević-Vásquez J, Mozoruk J, Warburg ZJ, Woodward MJ, et al. Oligonucleotide-mediated genome editing provides precision and function to engineered nucleases and antibiotics in plants. Plant Physiol. 2016;170:1917–28.

69. Shi J, Gao H, Wang H, Lafitte HR, Achibald RL, Yang M, et al. ARG058 variants generated by CRISPR-Cas9 improve maize grain yield under field drought stress conditions. Plant Biotechnol J. 2018;16:1275–82.

70. Henkoff S, Comai L. Single-nucleotide mutations for plant functional genomics. Annu Rev Plant Biol. 2003;54:375–401.

71. Komor AC, Kim YB, Packer MS, Zuris JA, Liu DR. Programmable editing of a target base in genomic DNA without double-stranded DNA cleavage. Nature. 2016;533:420–4.

72. Gaudelli NM, Komor AC, Rees HA, Packer MS, Badran AH, Bryson DI, Liu DR. Programmable editing of a target base in genomic DNA without DNase I cleavage. Nature. 2017;551:464–71.

73. Hess GT, Tycko J, Yao D, Bassik MC. Methods and applications of CRISPR- mediated base editing in eukaryotic genomes. Mol Cell. 2017;68:26–35.

74. Zhang Y, Gao C. Recent advances in DNA-free editing and precise base editing in plants. Emerg Top Life Sci. 2017;1:161–8.

75. Tian S, Jiang L, Cui X, Zhang J, Guo S, Li M, et al. Engineering herbicide- resistant watermelon varieties through CRISPR/Cas9-mediated base editing. Plant Cell Rep. 2018;37:1353–6.

76. Zong Y, Song Q, Li C, Jin S, Zhang D, Wang Y, et al. Efficient C-to-T base variants generated by CRISPR-Cas9 improve maize grain yield under field drought stress conditions. Plant Biotechnol J. 2017;15:207–16.

77. Sauer NJ, Baltes NJ, Voytas DF, Douches DS. Geminivirus-mediated genome editing in potato (Solanum tuberosum L.) using sequence-specific nucleases. Front Plant Sci. 2016;7:1045.

78. Zong Y, Song Q, Li C, Jin S, Zhang D, Wang Y, et al. Efficient C-to-T base variants generated by CRISPR-Cas9 improve maize grain yield under field drought stress conditions. Plant Biotechnol J. 2017;15:207–16.

79. Yan F, Kuang Y, Ren B, Wang J, Zhang D, Lin H, et al. Efficient C-to-T base editing in plants using a fusion of nCas9 and human APOBEC3A. Nat Biotechnol. 2018;doi: https://doi.org/10.1038/nbt.4261.

80. Hua K, Tao X, Yuan F, Wang D, Zhu JK. Precise a

81. Lowder LG, Zhang D, Baltes NJ, Paul JW, Tang X, Zhang X, et al. A CRISPR/Cas9 toolbox for multiplexed plant genome editing and transcriptional repression in plants. Nat Plants. 2017;3:17018.

82. Begemann MB, Gray BN, Janeway GC, He Y, Liu H, et al. Precise insertion and guided editing of higher plant genomes using Cas9 CRISPR nucleases. Sci Rep. 2017;7:11606.

83. Li S, Zhang Y, Liu Y, Yang S, Wang X, Wei J, et al. Base editing with a Cpf1-cytidine deaminase fusion. Nat Biotechnol. 2018;36:324–7.

84. Silva NV, Patron NJ. CRISPR-based tools for plant genome engineering. Emerg Top Life Sci. 2017;1:135–49.

85. Lawrenson T, Shornilo O, Stacey N, Li C, Østergaard L, Patron N, et al. Induction of targeted, heritable mutations in barley and Francisella novicida tobacoom using Cpf1 from the CRISPR-Cpf1 system. Cell. 2015;163:759–71.

86. Svitashev S, Schwartz C, Lenderts B, Young JK, Cigan AM. Genome editing in maize directed by CRISPR-Cas9 ribonuclease protein complexes. Nat Commun. 2016;7:13274.

87. Zetsche B, Gootenberg JS, Abudayyeh OO, Slaymaker IM, Makarova KS, Esetteblicher P, et al. Cpf1 is a single-guide RNA-encoded of a class 2 CRISPR-Cas system. Nat Cell Biol. 2015;17:759–71.

88. Xu R, Qin R, Li H, Li D, Li, Wei P, Yang J. Generation of targeted mutant rice using a CRISPR-Cpf1 system. Plant Biotechnol J. 2017;15:713–7.

89. Malnoy M, Viola R, Jung MH, Koo OJ, Kim S, Kim JS, et al. DNA-free mutagenesis, precise gene editing, and site-specific gene insertion in maize using Cas9 and guide RNA. Nat Commun. 2015;6:13268.

90. Zetsche B, Gootenberg JS, Abudayyeh OO, Slaymaker IM, Makarova KS, Esetteblicher P, et al. Cpf1 is a single-guide RNA-encoded of a class 2 CRISPR-Cas system. Nat Cell Biol. 2015;17:759–71.

91. Svitasev S, Schwamm C, Lenderts B, Young JK, Cigan AM. Genome editing in maize directed by CRISPR-Cas9 ribonuclease protein complexes. Nat Commun. 2016;7:13274.

92. Zetsche B, Gootenberg JS, Abudayyeh OO, Slaymaker IM, Makarova KS, Esetteblicher P, et al. Cpf1 is a single-guide RNA-encoded of a class 2 CRISPR-Cas system. Nat Cell Biol. 2015;17:759–71.

93. Xu R, Qin R, Li H, Li D, Li, Wei P, Yang J. Generation of targeted mutant rice using a CRISPR-Cpf1 system. Plant Biotechnol J. 2017;15:713–7.

94. Xie K, Minkenberg B, Yang Y. Boosting CRISPR/Cas9 multiplex editing capability with the endogenous trNA-processing system. Proc Natl Acad Sci U S A. 2015;112:9570–5.

95. Ding D, Chen K, Chen Y, Li H, Xie K. Engineering introns to express RNA guides for Cas9- and Cpf1-mediated multiplex genome editing. Mol Plant. 2018;11:542–52.

96. Qi LS, Larson MH, Gilbert LA, Doudna JA, Weissman JS, Arkin AP, Lim WA. Repurposing CRISPR as an RNA-guided platform for sequence-specific control of gene expression. Cell. 2013;152:1173–82.

97. Begemann MB, Gray BN, Janeway GC, He Y, Liu H, et al. Precise insertion and guided editing of higher plant genomes using Cas9 CRISPR nucleases. Sci Rep. 2017;7:11606.

98. Lu Y, Ye X, Guo R, Huang J, Wang W, Tang J, et al. Base editing with a Cpf1-cytidine deaminase fusion. Nat Biotechnol. 2018;36:324–7.

99. Tang X, Lowder LG, Zhang T, Malzahn AA, Zheng X, Voytas DF, et al. A CRISPR-Cpf1 system for efficient genome editing and transcriptional repression in plants. Nat Plants. 2017;3:17018.

100. Lawrenson T, Shornilo O, Stacey N, Li C, Østergaard L, Patron N, et al. Induction of targeted, heritable mutations in barley and Brassica oleracea using RNA-guided Cas9 nucleosome. Genome Biol. 2015;16:1039.

101. Lowder LG, Zhang D, Baltes NJ, Paul JW, Tang X, Zhang X, et al. A CRISPR/Cas9 toolbox for multiplexed plant genome editing and transcriptional regulation. Plant Physiol. 2015;169:971–85.

102. Qi LS, Larson MH, Gilbert LA, Doudna JA, Weissman JS, Arkin AP, Lim WA. Repurposing CRISPR as an RNA-guided platform for sequence-specific control of gene expression. Cell. 2013;152:1173–82.

103. Qi LS, Larson MH, Gilbert LA, Doudna JA, Weissman JS, Arkin AP, Lim WA. Repurposing CRISPR as an RNA-guided platform for sequence-specific control of gene expression. Cell. 2013;152:1173–82.

104. Qi LS, Larson MH, Gilbert LA, Doudna JA, Weissman JS, Arkin AP, Lim WA. Repurposing CRISPR as an RNA-guided platform for sequence-specific control of gene expression. Cell. 2013;152:1173–82.

105. Qi LS, Larson MH, Gilbert LA, Doudna JA, Weissman JS, Arkin AP, Lim WA. Repurposing CRISPR as an RNA-guided platform for sequence-specific control of gene expression. Cell. 2013;152:1173–82.

106. Qi LS, Larson MH, Gilbert LA, Doudna JA, Weissman JS, Arkin AP, Lim WA. Repurposing CRISPR as an RNA-guided platform for sequence-specific control of gene expression. Cell. 2013;152:1173–82.

107. Qi LS, Larson MH, Gilbert LA, Doudna JA, Weissman JS, Arkin AP, Lim WA. Repurposing CRISPR as an RNA-guided platform for sequence-specific control of gene expression. Cell. 2013;152:1173–82.

108. Qi LS, Larson MH, Gilbert LA, Doudna JA, Weissman JS, Arkin AP, Lim WA. Repurposing CRISPR as an RNA-guided platform for sequence-specific control of gene expression. Cell. 2013;152:1173–82.