Expanding the plant genome editing toolbox with recently developed CRISPR–Cas systems

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Abstract

Since its first appearance, CRISPR–Cas9 has been developed extensively as a programmable genome-editing tool, opening a new era in plant genome engineering. However, CRISPR–Cas9 still has some drawbacks, such as limitations of the protospacer-adjacent motif (PAM) sequence, target specificity, and the large size of the cas9 gene. To combat invading bacterial phages and plasmid DNAs, bacteria and archaea have diverse and unexplored CRISPR–Cas systems, which have the potential to be developed as a useful genome editing tools. Recently, discovery and characterization of additional CRISPR–Cas systems have been reported. Among them, several CRISPR–Cas systems have been applied successfully to plant and human genome editing. For example, several groups have achieved genome editing using CRISPR–Cas type I-D and type I-E systems, which had never been applied for genome editing previously. In addition to higher specificity and recognition of different PAM sequences, recently developed CRISPR–Cas systems often provide unique characteristics that differ from well-known Cas proteins such as Cas9 and Cas12a. For example, type I CRISPR–Cas10 induces small indels and bi-directional long-range deletions ranging up to 7.2 kb in tomatoes (Solanum lycopersicum L.). Type IV CRISPR–Cas13 targets RNA, not double-strand DNA, enabling highly specific knockdown of target genes. In this article, we review the development of CRISPR–Cas systems, focusing especially on their application to plant genome engineering. Recent CRISPR–Cas tools are helping expand our plant genome engineering toolbox.

Introduction

The rapid progress of plant genome editing technologies has brought about a dramatic evolution in plant research and plant breeding. The most general and well-known genome editing tool is clustered regularly interspaced short palindromic repeats-CRISPR associated protein 9 (CRISPR–Cas9), which applies a programmable RNA-guided Cas9 endonuclease (Osakabe and Osakabe, 2015; Wang et al., 2016; Wada et al., 2020) to edit target sequences. Especially, CRISPR–Cas9 from Streptococcus pyogenes (SpyCas9) has been applied for genome editing in many organisms (Jinek et al., 2012; Cong et al., 2013; Mali et al., 2013; Osakabe and Osakabe, 2015; Wang et al., 2016; Jaganathan et al., 2018; Wada et al., 2020). The main drawbacks of CRISPR–Cas9 are the limitations of available protospacer-adjacent motif (PAM) sequences, target specificity, and its large size, which prevents delivery by virus-based vectors (Yang et al., 2021). To overcome these problems, engineering of SpyCas9 (Kleinstiver et al., 2015, 2016; Slaymaker et al., 2016; Chen et al., 2017; Hu et al., 2018; Lee et al., 2018; Nishimasu et al., 2020).
ADVENTS

• Type I CRISPR–Cas systems, which recognize longer sequences than Cas9 or Cas12a, have been applied for genome editing and transcriptional control.
• Type I CRISPR–Cas3 has been applied to transcriptional control and genome editing in human cells but not yet in plant cells.
• Type I CRISPR–Cas10 induces bi-directional long-range deletions and small indels in human and plant cells.
• Type V CRISPR–Cas12b has been applied to genome editing in monocot and dicot plants with high specificity at target sites, inducing deletions larger than those induced by Cas9.
• Type IV CRISPR–Cas13, which can target RNA, has been used for knockdown of target gene expression and has also provided RNA virus interference activity in plants.

In 2019, several studies reported the successful application of Type I CRISPR–Cas to transcriptional control and gene editing in human cells (Cameron et al., 2019; Dolan et al., 2019; Morisaka et al., 2019; Pickar-Oliver et al., 2019; Chen et al., 2020). Representative structures of Class I Cascade-effectors and Class II inactive Cas-effectors used for transcriptional control are shown in Figure 2. Pickar-Oliver et al. (2019) modified gene expression of a target gene by tethering activation (human acetyltransferase p300) or repression (Krüppel-associated box, KRAB) domains to Escherichia coli type I-E Cascade (EcCascade) or Listeria monocytogenes Finland_1998 type I-B Cascade (LmoCascade) in human cells. They found that tethering of p300 to Cas8e, Cse2, Cas5e, or Cse6 could induce gene activation without abrogating complex formation. By tethering p300 or KRAB domains to Cas6e, gene expression could be modified with high efficiency and high specificity (Pickar-Oliver et al., 2019). Similarly, type I-F Cascade has also been applied for transcriptional control in human cells (Chen et al., 2020). Interestingly, Chen et al. (2020) demonstrated that tethering of transcription activation domain VP64-p65-Rta (VPR) to Csy3 (Cas7 equivalent) of Pseudomonas aeruginosa type I-F
Cascade (PaeCascade) could activate expression of the target gene with high specificity and more efficiently than dCas9-VPR, dAsCas12a-VPR, or EcoCascade-VPR. Expression of PaeCascade containing PaeCsy3-VPR resulted in accumulation of six copies of Csy3-VPR at the target site (a Csy3 protein per every 6 nt of the crRNA). The extended crRNA also recruited more Csy3-VPR to target site, resulting in enhanced gene activation (Chen et al., 2020). Interestingly, expression of type I-B LmoCascade containing LmoCas7-VPR failed to induce gene activation (Pickar-Oliver et al., 2019), suggesting that optimization of linker sequence is required or that the effects of Cas7-VPR on transcriptional control differ depending on the CRISPR–Cas system used. In addition, type I-F PaeCascade needs fewer Cas proteins to function than the type I-E EcoCascade (four Cas proteins versus five Cas proteins). Requirement for fewer Cas components makes the vector system simpler, contributing to the development of easy-to-use genome manipulation tools.

For genome editing, Cameron et al. (2019) fused a dimerization-dependent, nonspecific FokI nuclease domain to Cas8e and found that the FokI-Cascade from Pseudomonas sp. S-6–2 (PseCascade) using a paired gRNA showed higher genome editing efficiency (up to 50%) than other FokI-Cascade complexes such as FokI-EcCascade. Specificity of the FokI-Cascade was analyzed by GUIDE-seq. While more than 250 off-targets were recovered and their read counts ranged from 11 to 41,733 reads/421,646 total reads in Cas9-treated samples, at most 2 off-target sites were detected, and their reads were only 20 reads/10,757 total reads and 35 reads/10,757 total reads, respectively in any of the FokI-PseCascade paired gRNAs-treated samples. These results indicate the high specificity of FokI-Cascade.

Genome editing using full components of type I-E CRISPR–Cas (including Cas3e) was reported by three groups in 2019 (Dolan et al., 2019; Morisaka et al., 2019; Pickar-Oliver et al., 2019). Dolan et al. (2019) achieved genome

**Table 1** General characteristics of each type of CRISPR–Cas system

| Class | Type | Subtype | Signature Gene | Target | crRNA | Representative Cas Proteins Related to Each Step |
|-------|------|---------|----------------|--------|-------|-----------------------------------------------|
| I     | Type I | 7       | cas3           | DNA    | Single crRNA | Cas6, Cas5, Cas7, Cas8, Cas11 |
| I     | Type III | (d³)    | cas3           | DNA    | Single crRNA | Cas6, Cas5, Cas7 |
| I     | Type IV  | 3       | csf1           | –      | –      | Cas6, Cas5, Cas7, Cas10 |
| II    | Type II  | 3       | cas9           | DNA    | tracrRNA:crRNA | Cas9, Cas12 |
| II    | Type V   | 10      | cas12          | DNA    | Single crRNA/tracrRNA | Cas12 |
| II    | Type VI  | 4       | cas13          | RNA    | Single crRNA | Cas13 |

*aClassification is based on Makarova et al. (2020).
*bA cas3 gene is present as a signature gene in type I-D but cleavage is performed by Cas10, not Cas3.
*cSome type IV CRISPR–Cas systems lack a csf1 gene.

**Box 1. CLASSIFICATION OF CRISPR–CAS SYSTEMS**

Bacteria and archaea have a variety of CRISPR–Cas systems that function naturally as an adaptive and heritable immune system (Makarova et al., 2020). In the current classification, CRISPR–Cas has been classified into 2 classes, 6 types, and 33 subtypes based on the gene organization of CRISPR–Cas loci, the presence of the signature cas gene, sequence similarity and phylogenetic analysis of conserved Cas proteins (Makarova et al., 2020). Class is defined by an organization of effector proteins. Class I CRISPR–Cas systems (types I, III, and IV) have a multi-effector complex termed CRISPR-associated complex for antiviral defense (Cascade), whereas Class II CRISPR–Cas systems (types II, V, and VI) has a single multi-domain effector Cas protein (Makarova et al., 2020; Figure 1; Table 1). Each type is represented by the presence of signature proteins: Cas3 in Type I, Cas9 in Type II, Cas10 in Type III, Csfl in Type IV, Cas12 in Type V, and Cas13 in Type VI. Classification of subtype has been more complicated (for details, please refer to a review by Makarova et al., 2020). Recently, we identified that Cas10 is an important component in TiD (Osakabe et al., 2020, 2021). This diversification of CRISPR–Cas results from evolution in an intensive arms race between bacteria and their phage foes (Hampton et al., 2020). Each type and subtype has different characteristics in terms of recognition of polynucleotides (DNA or RNA), organization of cas genes, target recognition and cleavage mechanisms (Makarova et al., 2020; Table 1).
editing using *Thermobifida fusca* type I-E CRISPR–Cas. They purified the TfCascade and Cas3 complex and delivered them as a ribonucleoprotein (RNP) into human embryonic stem cells, resulting in successful induction of mutations at target sites. Interestingly, type I-E CRISPR–Cas induced long-range genomic deletions of up to 100 kb at target sites—a characteristic unique to type I-E CRISPR–Cas and different from the mutation patterns induced by Cas9. The deletions were unidirectional, toward the region upstream of the PAM sequence. Cameron et al. (2019) has also successfully induced mutations using a full PseCascade–Cas3 complex in human cells. Their strategy involved plasmid DNA-based delivery of the full PseCascade–Cas3 complex into human HEK293 cells. The mutation patterns were similar to those reported by Dolan et al. (2019). Morisaka et al. (2019) also delivered EcCascade–Cas3 into human HEK293T cells by using plasmid DNA, and achieved genome editing in human cells. A comprehensive analysis of mutation patterns demonstrated that type I-E CRISPR–Cas can induce unidirectional long deletions with high specificity. In addition, they indicated that type I-E CRISPR–Cas can knock out target genes efficiently and can also be used for knock-in of a DNA fragment via the homology-directed repair (HDR) pathway. By achieving exon skipping of the dystrophin gene in patient-induced pluripotent stem cells, they demonstrated the potential of type I-E CRISPR–Cas for future therapeutic applications (Morisaka et al., 2019).

On the other hand, the application of type I-E CRISPR–Cas to plant genome engineering has been limited to date. The only application of type I-E CRISPR–Cas in plant cells has been in transcriptional control in *Zea mays* (Young et al., 2019; Figure 3A). Young et al. (2019) optimized and engineered type I-E CRISPR–Cas from *Streptococcus thermophilus* to plant genome engineering has been limited to date.

Figure 1 Representative structures of CRISPR–Cas systems. Cas proteins with crRNAs are shown with double-strand DNAs in (A)–(F) and with single-strand RNA in (G). A, Class I type I-D CRISPR–Cas10. Cas10 and Cas3 bind to the Cascade complex containing Cas5, Cas6, Cas7, and crRNA, then Cas10 cleaves and digests dsDNA at target site. B, Class I type I-E CRISPR–Cas3. Cas3 is recruited to the Cascade complex containing Cas5, Cas6, Cas7, Cas8, Cas11, and crRNA, then digests dsDNA at target sites. C, Class II type II CRISPR–Cas9. Single Cas9 protein forms a complex with crRNA annealing to tracrRNA and cleaves dsDNAs at target sites, producing blunt ends. D, Class II type V CRISPR–Cas12a. Single Cas12a protein forms a complex with crRNA and cleaves dsDNAs at target site, producing cohesive ends. E, Class II type V CRISPR–Cas12b. Single Cas12b protein forms a complex with crRNA annealing to tracrRNA and cleaves dsDNAs at target site, producing cohesive ends. F, Class II type V CRISPR–CasΦ. Single CasΦ protein forms a complex with crRNA and cleaves dsDNA at target sites, producing cohesive ends. G, Class II type VI CRISPR–Cas13. Single Cas13 protein forms a complex with crRNA, and cleaves single strand RNA of target gene. PFS, protospacer flanking sequence.
A Cascade complex (without Cas protein containing a nuclease domain) | Effector domains

Transcriptional control
epigenetic modifications
base editing etc.

target site

B Catalytically inactive Cas protein | Effector domains

Transcriptional control
Epigenetic modifications
base editing etc.

target site

Figure 2 Representative structure of Cas proteins with effector proteins. Class I Cascade complex fused to an effector domain (A) and Class II catalytically inactive Cas protein fused to an effector domain (B) are shown. These engineered Cas proteins have been applied for transcriptional control, epigenetic modification, base editing, etc. at target sites. Effector domains can be fused to any Cas proteins of Cascade complex (A) whereas an effector domain can be fused to one Cas protein (B).

Class I type I-D CRISPR–Cas

Recently, we characterized type I-D (TiD) CRISPR–Cas loci from *Microcystis aeruginosa*, and successfully developed a genome editing tool, which we named TiD, based on this system (Osakabe et al., 2020, 2021; Figure 1A; Figure 3B). Several groups have reported in vitro DNA binding and cleavage capability of several type I-D systems from bacteria such as *Sulfolobus islandicus* and *Synechocystis sp. PCC 6803* (Manav et al., 2020, McBride et al., 2020, Lin et al., 2020); however, its applications to genome editing in eukaryotic cells have not been reported. TiD consists of five Cas proteins (Cas 3d, 5d, 6d, 7d, and 10d) and a crRNA that recognizes a 35-nt or 36-nt target sequence. Among TiD Cas proteins, Cas10d is a unique protein not found in other type I CRISPR–Cas systems. Generally, Cas3 has a histidine–aspartate (HD) nuclease domain and functions as a nuclease. However, in the TiD system, it is Cas10d that instead has the HD nuclease domain. Cas10 is known as a signature protein of the type III CRISPR–Cas family, but the Cas10d in TiD has highly diverged in comparison with its type III counterparts. The Cas10d HD domain is rather similar to the Cas3d HD domains of type I-B, -C, -E, and -F. Therefore, although TiD is a unique system that possesses both types I and III signature genes, it is assigned to type I. The presence of an HD domain in Cas10d raised the hypothesis that Cas10d plays a role as a nuclease. Hence, we analyzed ssDNA nuclease activity in vitro and the results indeed indicated that Cas10d, but not Cas3d, has ssDNA nuclease activity (Osakabe et al., 2020). Both these Cas proteins also showed ATPase activity, suggesting that they function as a helicase to unwind the dsDNA. TiD recognizes 5′-GTH-3′ as a PAM together with the following 35- or 36-nt target sequences. Like other type I-based systems, recognition of a longer target sequence (35 or 36 nt) than that of Cas9 (20 nt) suggests that TiD has higher specificity than CRISPR–Cas9. Using TiD, we have successfully induced mutations at target genes in human cells (Osakabe et al., 2021). Interestingly, our results showed that the mutation patterns produced by TiD in both human cells and plants were different from those induced by known Cas proteins such as Cas9, Cas12a, and Cas3: TiD introduced not only small insertion/deletion but also long-range deletions (ranging from 2.5 kb to 18.5 kb), and its direction was not uni-directional, but bi-directional (Osakabe et al., 2020, 2021). Although the mutation patterns induced by genome editing would depend on the host organisms and the presence of active repair pathways, TiD would be a tool that can induce both small indels and long deletions.
whereas majority of mutations induced by CRISPR–Cas9 and Cas12 are small indels (they also induce long deletions in some cases), and type I-E CRISPR–Cas can induce only unidirectional long deletions. These differences can be attributed to the function of a unique nuclease protein, Cas10d.

We optimized TiD for genome editing in plant cells by using plant-cell specific-promoters (CaMV35S, Parsley Ubiquitin 4-2) for expression of codon-optimized cas genes and the AtU6-26 promoter for expression of crRNA (Figure 4). Using plant-optimized TiD vectors targeting tomato INDOLE-3-ACETIC ACID9 (SlIAA9, an important factor for parthenocarpy) and RIPENING INHIBITOR (an important factor for fruit ripening) genes, we introduced mutations successfully into tomato plants (S. lycopersicum L., cv. Micro-Tom and Ailsa Craig) (Osakabe et al., 2020). As with the mutation patterns induced in human cells, short indels and long deletions were detected in callus and shoots. Small indels were induced in 64% of transgenic calli. Long-range deletions were also detected in transgenic calli and shoots. We detected up to 7.2 kb of bi-directional long-range deletions around the target sequence. Sequence analysis of regenerated shoots indicated that the transgenic shoots included 100% mutated DNA sequence, revealing that TiD could induce bi-allelic mutations in the T0 generation.

Figure 3 Application of recently developed CRISPR–Cas tools to plant genome engineering. (A) Type I-E SthCascade-CBF1 for targeted gene activation in Z. mays (Young et al., 2019). B, TiD for targeted mutagenesis in tomatoes (Osakabe et al., 2020). C, Type V CRISPR–AaCas12b for targeted mutagenesis in rice (Ming et al., 2020). D, RNA targeting using type VI CRISPR–Cas13d in N. benthamiana (Mahas and Mahfouz, 2018). Sth, S. thermophilus DGCC7710.
SlIAA9 knockout phenotypes (seedless fruit, changes in leaf morphology) were also observed. To identify the basic characteristics of TiD as a genome editing tool, we searched on-target sites and off-target candidate sequences with 0–5 mismatches against Arabidopsis, rice (*O. sativa* L.) and tomato whole genome for TiD (Osakabe et al., 2020). The results indicated that more on-target sites for TiD exist in tomato and Arabidopsis than for Cas9, but there are fewer in rice. For off-target sites, TiD has clearly fewer off-target candidate sites than Cas9 in these plant genomes, suggesting an advantage for TiD as a highly specific genome editing tool in plant cells. Off-target mutations, including short indels and long deletions, were also not detected from SlIAA9 knockout plants, and the on-target mutations were transmitted to the next generation (Osakabe et al., 2020). These results show that TiD is a useful and unique alternative genome editing tool for both human and plant genome editing. Although further improvements and research is needed, such as identifying regulatory control mechanisms, long-range deletions would allow TiD to be used for chromosome engineering, further expanding what is possible in plant genome engineering.

Class II type V-B CRISPR–Cas12b (C2c1)

CRISPR–Cas type V features an RNA-guided effector protein, Cas12, which contains RuvC domain. Distinct architectures and the diverged RuvC sequences of Cas12 proteins suggest functional diversity (Yan et al., 2019). Identified Cas12 proteins indicate a range of functional activities, including targeting and collateral cleavage of single-strand RNA (ssRNA) and DNA, as well as dsDNA nicking and cleavage (Yan et al., 2019). CRISPR–Cas12a is a well-studied tool and we refer the reader to some excellent detailed reviews of its application (Zetsche et al., 2015; Alok et al., 2020) rather than covering this topic here. On the other hand, CRISPR–Cas12b (formerly known as C2c1) is an alternative type V CRISPR–Cas system with a dual-RNA-guided endonuclease, meaning that it requires two kinds of RNA (crRNA and trans-activating crRNA [tracrRNA]) for its nuclease function, whereas Cas12a requires only crRNA (Shmakov et al., 2015; Table 1; Figure 1, D and E). Cas12b has a conserved RuvC nuclease domain and Nuc domain that has no similarity with Cas12a. CRISPR–Cas12b is an attractive tool because Cas12b produces a long staggered end distal to the PAM, and it is smaller than Cas9 (Teng et al., 2018). However, its optimal temperature for DNA cleavage of Cas12b is generally higher than 40°C, suggesting that it is not suitable for genome editing in mammalian and plant cells. In 2018, Teng et al. (2018) identified a Cas12b from *Alicyclobacillus acidi-philus* (AaCas12b) that can maintain optimal nuclease activity at 31–59°C. Using AaCas12b, they successfully induced mutations at target genes in human and mouse cells (Teng et al. 2018). Mutation efficiency by AaCas12b in human cells was 21.5%. Multiplex genome editing using four gRNAs was also achieved, with a mutation efficiency of 3%–20% depending on the target site. Gene activation using a nuclease-deficient mutant of AaCas12b was also possible, although activation was not as strong as when using dSpyCas9. AaCas12b did not induce mutations at 88 predicted off-target sites, while SpyCas9 induced 3 out of 82 predicted off-target sites, indicating that AaCas12b has higher specificity than SpyCas9. Cas12b putative orthologs were also identified and their interchangeability between Cas12b effectors and dual-RNAs derived from other CRISPR–Cas12b systems revealed (Teng et al., 2019).

Strecker et al. (2019a, 2019b) performed protein engineering of Cas12b to change the optimal DNA cleavage...
temperature to 37°C. They identified mesophilic Bacillus hisashii Cas12b and produced a gain-of-function mutant, BhCas12b v4, that exhibited increased dsDNA cleavage activity and reduced nickase activity to the nontarget DNA strand at 37°C. BhCas12b v4 induced prominent larger deletions of 5–15 bp at target sites. Mutation efficiency was comparable to, or a little lower than, that of SpyCas9, depending on the target site. Also, no off-target cleavages were detected in BhCas12b v4-treated samples while SpyCas9-treated samples included off-target mutations, indicating the higher specificity of BhCas12b v4 when compared with SpyCas9 (similar to AaCas12b).

For the application of Cas12b to plant genome editing, Ming et al. (2020) compared Cas12b proteins from various bacteria in monocot rice (O. sativa): Cas12b from Alicyclobacillus acidoterrestris (Aac), A. acidiphilus (Aa), Bacillus thermoamyllovoran (Bth), and B. hisashii. Comparison of genome editing efficiency suggested that AaCas12b is a more efficient genome editing tool than other Cas12b proteins in rice. AaCas12b and AacCas12b recognized VTTV (V: A, C, G) PAMs, preferentially ATT and GTTG PAM. In particular, AaCas12b induced mutations with high efficiency (>50%) at ATTA, ATTC, and GTTG PAM in rice protoplasts, although the mutation efficiency was very different depending on the target site. AaCas12b generated biallelic mutants with an efficiency of 30%–46%. The deletions occurred at about 12–24 bp distal to the PAM site, and they were larger than those induced by Cas9 (Figure 3C). Moreover, inactivated AaCas12b variants with transcriptional repression/activation domains have controlled the gene expression of targeted genes successfully. These results reveal Cas12b as a promising tool for genome editing in rice.

Because cotton (Gossypium hirsutum) is resistant to high temperature, it raised the possibility that heat-inducible AaCas12b could be used for genome editing in cotton. Wang et al. (2020) assessed this possibility, testing various temperature conditions and durations during the callus induction stage after Agrobacterium tumefaciens-infection of cotton. They found that exposure of explants to 45°C for 4 d resulted in highest genome editing efficiency (17.4%) with little adverse effect on cotton cell culture. AaCas12b induced deletions of 1–16 bp, with the majority ranging from 9 to 14 bp. This is larger than the average size of deletions induced by SpyCas9, which is consistent with results in human cells (Teng et al. 2018). Since cotton is an allotetraploid plant, derived from the ancestral hybridization of two diploid genome (A and D), multiple copies exist in almost all genes. Interestingly, mutations were induced in the GhCLA gene in the Dt sub-genome more efficiently than in the GhCLA gene in the At sub-genome. This suggests that differences in chromatin structure affect genome editing efficiency by Cas12b in cotton. (Wang et al. 2020). Off-target mutations were not detected in genome-edited cotton.

The third application of CRISPR–Cas12b in plants is genome editing of the dicot Arabidopsis (Wu et al., 2020).
transgenes into host genomic DNA, but there is a limit to the size of DNA that can be packaged. Thus, CRISPR–CasΦ would be useful for genome editing via virus-based vectors.

**Class II type VI CRISPR–Cas13**

Type VI CRISPR–Cas is a unique system that can recognize and cleave single-stranded RNA, but not double-stranded DNA by a signature protein, Cas13 (Shmakov et al., 2015, 2017; Figure 1G). Cas13 has two structurally distinct higher eukaryote and prokaryote nucleotide-binding (HEPN) domains, by which Cas13 can target and process precrRNA into mature and functional crRNAs. Cas13 proteins have been divided into four subtypes (Cas13a, b, c, and d; Shmakov et al., 2017). In 2016, using experiments in vitro and in *E. coli*, Abudayyeh et al. (2016) characterized a Cas13a protein (formerly named C2c2) from *Leptotrichia shahii* (LshCas13a) as an RNA-guided RNA-targeting effector. They demonstrated that Cas13a has a collateral RNase activity, which cleaves RNAs nonspecifically once activated, in vitro and in bacteria cells. Abudayyeh et al. (2017) first demonstrated that *Leptotrichia wadei* Cas13a (LwCas13a) can knockdown targeted RNA with higher specificity than RNAi in human cells. No collateral RNase activity was detected in human cells. Other than LshCas13a and LwCas13a, various Cas13 putative orthologs have been identified and applied to RNA editing (Shmakov et al., 2015; Cox et al., 2017; Yan et al., 2018; Konerman et al., 2018; Mahas and Mahfouz, 2018; Xu et al., 2021). Cas13 proteins have also been engineered by inactivating HEPN domains and by fusing effector domains for several applications such as live cell imaging of RNA, base editing, and nucleic acid detection (Abudayyeh et al., 2017; Cox et al., 2017; Gootenberg et al., 2017).

RNA targeting by CRISPR–Cas13 is a promising approach for plant research (Wolter and Puchta, 2018). Abudayyeh et al. (2017) successfully targeted three different genes (EPSPS, HCT, and PDS genes) in rice protoplasts using LwaCas13a. More than 50% knockdown was achieved with seven out of nine gRNAs, showing that LwaCas13a can efficiently disrupt the cytoplasmic RNA pool in plants. In addition, CRISPR–Cas13 technology has offered an approach to combat plant RNA viruses. RNA viruses are the most common type of plant virus, and many plant DNA viruses have an RNA intermediate form in their life cycle (Roossinck 2003). Using codon-optimized LshCas13a, Aman et al. (2018a, 2018b) demonstrated that CRISPR–Cas13a can interfere with Turnip mosaic virus (TuMV) in *Nicotiana benthamiana* and Arabidopsis. A reduction of up to 50% of green fluorescent protein (GFP) signals derived from GFP-TuMV was observed, and the reduction level differed depending on the RNA target, suggesting that RNA accessibility affects Cas13a activity. The effectiveness of CRISPR–LshCas13a to combat virus infection was also shown in a monocot plant, rice, using Southern rice black-streaked dwarf virus and Rice Stripe Mosaic Virus (Zhang et al. 2019). Furthermore, Mahas and Mahfouz (2018) identified a Cas13 variant showing the most efficient RNA virus interference activity in planta in *N. benthamiana* (Figure 3D). They identified CasRX (Cas13d from *Ruminococcus flavefaciens*) as the most robust and specific Cas13 variant in plant cells, as is also the case in mammalian cells (Konermann et al., 2018). CasRX significantly reduced GFP signal expressed from targeted virus with no collateral activity in plant cells. Simultaneous targeting of two different RNA viruses was also achieved with high specificity. RNA knockdown and editing by CRISPR–Cas13 would open new approaches for plant research and provide new tools to combat plant viruses.

**Future perspectives**

In this review, we have summarized the current CRISPR–Cas toolbox, which has expanded dramatically with recently identified CRISPR–Cas systems. The diversity of CRISPR–Cas is surprising for example, the recently identified CasΦ derived from bacteriophage genomes is only half the size of Cas9 (Pausch et al., 2020). These recently discovered CRISPR–Cas systems have different unique characteristics compared with generally used Cas9 and Cas12a for plant genome editing (summarized in Table 2). However, the discovery of additional CRISPR–Cas tools raises some questions (see “Outstanding Questions”). For example, the applicability of several of the more recent CRISPR–Cas systems to plant genome editing has not yet been validated in plant cells. For example, the transposon-associated CRISPR–Cas system, which can insert DNA fragments at the target site, has been validated only in *E. coli* cells (Klompe et al., 2019; Strecker et al., 2019a, 2019b). Importantly, unlike the preceding Cas9 and Cas12a technologies, these recently identified CRISPR–Cas systems have not yet been engineered extensively, suggesting that there is still room for improvement of these tools to facilitate their application to plant genome editing.

In the field of plant genome editing, the development of new tools is not sufficient to achieve efficient generation of genome-edited plants. There are plant-specific problems with plant genome editing; it generally requires tissue culture, it has low HDR efficiency, and it often takes time to establish genome-edited homozygous lines, especially when optimization in nonmodel plants and/or in polyploid plants is necessary. Codon and vector optimization would also be needed depending on the plant species used. For polyploid plants, multiple copies of target genes have to be mutated to establish knockout plants. In addition, establishment of transgene-free genome-edited plants (null-segregant) is also preferable to avoid off-target risks, and to alleviate concerns about genome-edited plants (Wada et al., 2020). Recently, strategies to overcome some of these conventional problems have been reported: plant gene editing through de novo induction of meristems (Maher et al., 2020), efficient gene targeting in Arabidopsis (Miki et al., 2018), and genome editing during haploid induction (Kelliher et al., 2019), etc. Combined with these technological developments, plant genome engineering will become faster, more efficient, and more precise.
| CRISPR–Cas System | Origin | Target Cells | Purpose | PAM/PFS | Brief Description | References |
|-------------------|--------|--------------|---------|---------|------------------|------------|
| Type I-E CRISPR–Cas3 | *S. thermophilus* | *Z. mays* | Gene activation | AA | Gene activation was achieved at a similar level as dCas9-CBF1 in *Z. mays*. | Young et al. (2019) |
| Type I-D CRISPR–Cas10 (TiD) | *M. aeruginosa* | *S. lycopersicum* L. | Genome editing | GTT, GTC | TiD induced small indels and large deletions (~7.5 kb) at target sites in tomatoes. | Osakabe et al. (2020) |
| Type V-B CRISPR–Cas12b | *A. acidoterrestris*, *A. acidiphilus*, and *B. thermoamylovorans* | *B. hisashii* | Genome editing | VTTV (V: A, C, G) | Targeted mutagenesis in monocot rice was achieved with high specificity using AaCas12b, which was more efficient than other Cas12b. Deletions (4–14 bp) were larger than those induced by Cas9. | Ming et al. (2020) |
| Type V-B CRISPR–Cas12b | *A. acidoterrestris* | *G. hirsutum* | Genome editing | TTN | Genome editing of cottons, which is resistant to high temperature, were achieved using AaCas12b. | Wang et al. (2020) |
| Type V-B CRISPR–Cas12b | *Bacillus* sp. V3-13 and *B. hisashii* | *A. thaliana* | Genome editing | ATTA, ATTG | Genome editing of dicot Arabidopsis was achieved using BvCas12b and BhCas12b v4. | Wu et al. 2020 |
| Type V CRISPR–CasΦ | Bacteriophage genomes | *A. thaliana* | Genome editing | TBN (B: G, T, and C) | Small CasΦ has induced mutations into target genes in Arabidopsis protoplasts. | Pausch et al. (2020) |
| Type VI CRISPR–Cas13a | *L. wadei* | *O. sativa* | RNA targeting | no | More than 50% knockdown of target gene was achieved by LwaCas13 in rice. | Abudayyeh et al. (2017) |
| Type VI CRISPR–Cas13a | *L. shahii* | *N. benthamiana* and Arabidopsis | RNA targeting | A, U, and C | LshCas13a produced interference against TuMV expressing GFP in *N. benthamiana* and Arabidopsis. | Aman et al. (2018a, 2018b) |
| Type VI CRISPR–Cas13a | *L. shahii* | *O. sativa*, *N. benthamiana* | RNA targeting | No descriptions | LshCas13a inhibited virus infection in monocot and dicot plants. | Zhang et al. (2019) |
| Type VI CRISPR–Cas13d | *R. flavocaciens* | *N. benthamiana* | RNA targeting | No descriptions | CasRx indicated robust interference efficiencies with high specificity in *N. benthamiana*. | Mahas and Mahfouz (2018) |
OUTSTANDING QUESTIONS

- Can recently discovered CRISPR–Cas systems be developed as useful plant genome editing tools?
- How can modified CRISPR–Cas systems be combined with other technologies to facilitate plant genome engineering?
- What do recently developed CRISPR–Cas tools bring to plant genome engineering?

In the near future, we will be able to choose the most suitable genome editing strategy from a toolbox full of various kinds of unique tools, depending on the experimental purposes, the nature of the target gene, and the plant species. New tools will also expand what is possible, ranging from precise modifications of plant genomes, gene expressions, and epigenomes to chromosomal manipulation. For example, Schmidt et al. (2020) achieved a targeted inversion of a 1.1-Mb heterochromatic knob (hk45S) in Arabidopsis Col-0 chromosome 4, which resulted in the restoration of crossover between the short arms of Col-0 and Ler-1 chromosome 4. Translocations between different chromosomes at targeted sites have also been achieved using CRISPR–Cas9 by Beying et al. (2020). These achievements suggest that manipulation of genetic linkages by chromosome engineering for plant breeding is now possible. Newly developed technologies are set to further accelerate expansion of the possibilities of plant genome editing.

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