



Xu et al., 2019; Gao, 2021; Huang et al., 2021; Li et al., 2021c; Xia et al., 2021; Zhan et al., 2021; Puchta et al., 2022). To date, three major CRISPR/Cas mediated precision genome editing stems have been developed and successfully applied in plants such as homolog -directed DNA repair (HDR)-mediated targeted gene replacement or gene targeting (Sun et al., 2016; Li et al., 2019; Li and Xia, 2020; Lu et al., 2020; Chen et al., 2022a; Puchta et al., 2022), base editing (Komor et al., 2016; Nishida et al., 2016; Gaudelli et al., 2017; Li et al., 2017; Lu and Zhu, 2017; Shimatani et al., 2017; Zong et al., 2017; Hua et al., 2018; Wei et al., 2021; Tian et al., 2022) (Figure 1), and prime editing (An alone et al., 2019; Butt et al., 2020; Jiang et al., 2020; Hua et al., 2020a; Li et al., 2020c; Lin et al., 2020; Tang et al., 2020; Xu et al., 2020a, 2020c; Lu et al., 2021; Wang et al., 2021b; Perroud et al., 2022) (Figure 2). Among these three precise editing technologies, HDR enables the installation or replacement of all kinds of mutations or various lengths of fragments in a predefined manner, representing the holy grail of genome editing. However, although various strategies have been attempted in the past decade (for review, please check Zhan et al., 2021; Puchta et al., 2022; Chen et al., 2022a), HDR remains challenging in plants due to the facts that once the double-strand breaks (DSBs) are generated by CRISPR/Cas nucleases the predominant repair mechanism in cells is nonhomologous end joining (NHEJ) which usually results in random indels, as well as the obstacles in delivery of sufficient donor repair template (DRT) into the vicinity of the DSB and competition with the original DNA strand/fragment to be replaced in plant cells (Li et al., 2019; Lu et al., 2020; for review, please check Li and Xia, 2020; Zhan et al., 2021; Chen et al., 2022a). In contrast, base editing and prime editing are two alternative promising strategies for precise genome editing without a DSB and a DRT. Whereas base editing has emerged as an alternative and effective tool to HDR-mediated gene replacement for precise single base substitution of an allele with a single SNP, facilitating precise gene editing by transition of one single base to another in a programmable manner (Komor et al., 2016; Nishida et al., 2016; Gaudelli et al., 2017) (Figure 1A–C), prime editing enables the installation of all 12 types of base substitutions and small indels, and substantially expands the scope and capabilities of precision genome editing (An alone et al., 2019) (Figure 2A).

Since the development of the first generation of base editor (BE) and prime editor (PE) for base editing and prime editing in mammalian cells (Komor et al., 2016; Nishida et al., 2016; Gaudelli et al., 2017; An alone et al., 2019), diverse strategies have been exploited to optimize these two editors in order to improve the precise editing efficiency and specificity, and to expand targeting scopes in plants (Li et al., 2020d; Molla et al., 2021; Hua et al., 2022). Here, we summarize the latest developments of various BEs and PEs, as well as their applications in plants. We also provide recommendations in selection of the proper BEs or PEs in practical applications in plants. Moreover, we propose the perspectives for further optimization

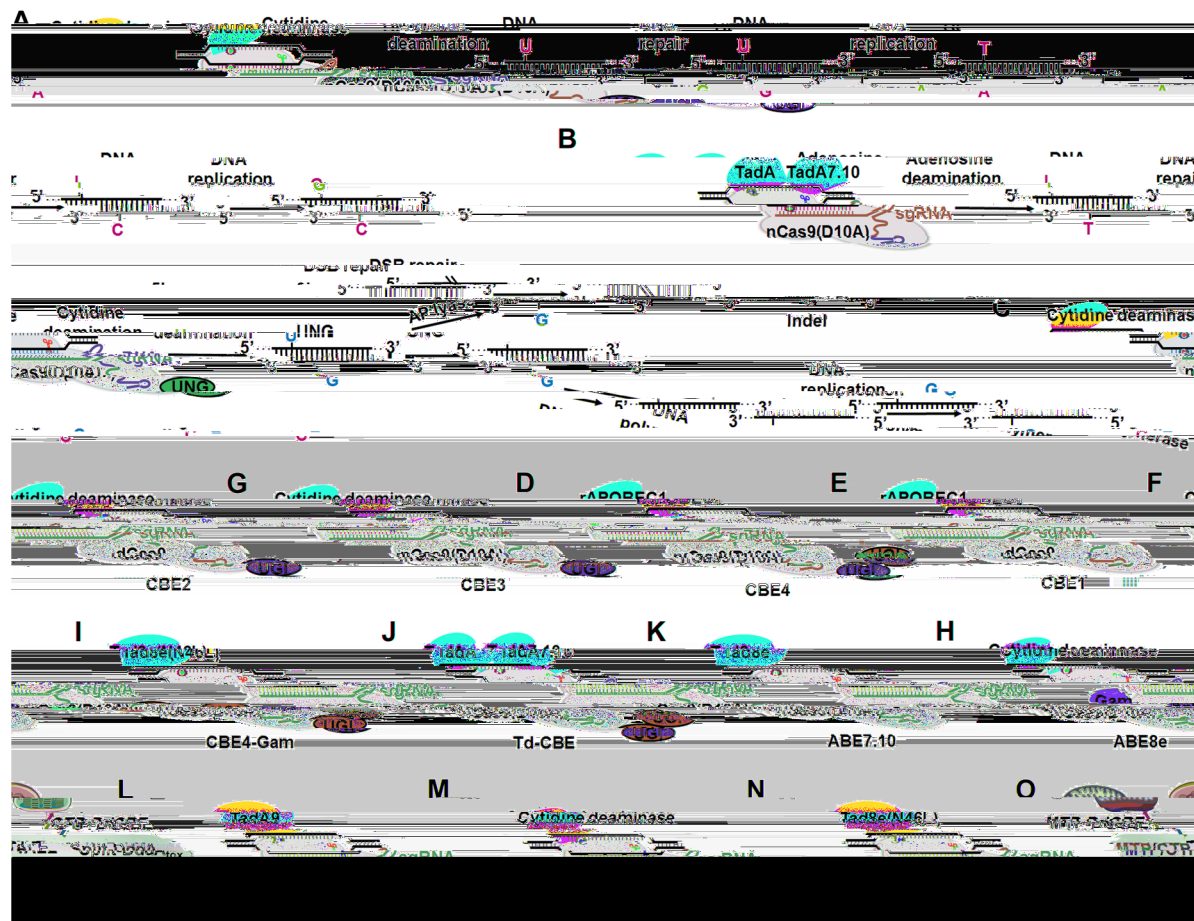
of these two editors. We truly believe this review will provide a valuable clue to the readers on how to select the appropriate BEs and PEs, as well as future perspectives to streamline these two editors into the routine and customized platform for both fundamental biological studies and crop improvement.

BASE EDITORS AND THEIR APPLICATIONS IN PLANTS

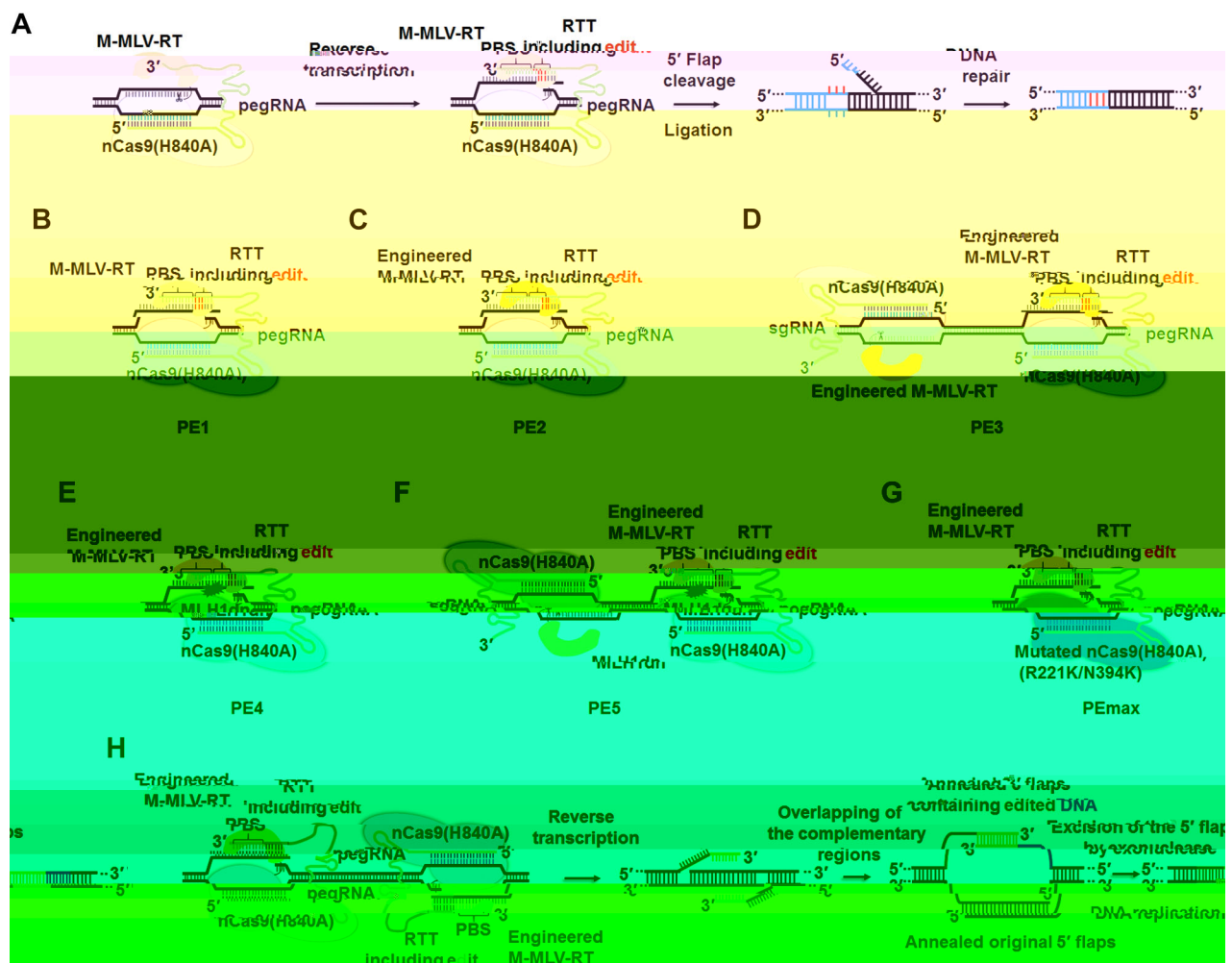
Base editing is a breakthrough technology that can precisely and efficiently achieve single base transition or transversion at target sites without inducing DSBs and the need for a DRT. Three BEs are currently in use: cytosine base editors (CBEs) for C:G to T:A transition (Figure 1A), adenine base editors (ABEs) for A:T to G:C transition (Figure 1B) and C-to-G base editors (CGBEs) for C:G to G:C transversion (Figure 1C). Precise base editing enables a single nucleotide substitution in a specific target gene to generate either loss-of-function or gain-of-function mutations, thus greatly accelerating functional annotation, crop improvement, de novo domestication or directed evolution of target genes in crop plants (Ren et al., 2018; Bharat et al., 2020; Kuang et al., 2020; Zeng et al., 2020; Xu et al., 2021a; Yan et al., 2021; Tan et al., 2022). Since the report of the first generation of CBE and ABE in 2016 (Komor et al., 2016) and in 2017 (Gaudelli et al., 2017) in mammalian cells, respectively, many efforts have been attempted in order to optimize and upgrade these two BEs in plants.

Cytosine base editors

The first-generation CBE was engineered by fusing a rat cytidine deaminase rAPOBEC1 to the N-terminus of an impaired dead Cas9 (dCas9) (Cas9 with D10A and H840A mutations) to generate rAPOBEC1-dCas9 and designated as CBE1 (Komor et al., 2016) (Figure 1D). The substitution of C to T in DNA is created by deaminating the cytosine (C) into uracil (U) in the exposed non-target DNA strand, and the subsequent DNA repair and replication results in C to T base conversion (Figure 1A). The cellular base excision repair (BER) mechanism enables C:G to T:A transition in vivo, while recognizes an G:U base pair as a mismatch. The BER activity eliminates the uracil with the help of uracil N-glycosidase (UNG), resulting in a low efficiency of the CBE1 system (Komor et al., 2016). To improve base editing efficiency, td1Tf.32140TD(519(as)19



() CRISPR/nCas9-mediated c tosine base editing. A c tosine base editor (CBE), which is composed of a catalytic impaired nCas9(D10A) and a c tosine deaminase, binds to the target sequence in the genomic DNA in a guide RNA (gRNA)-programmed manner. The c tosine deaminase catalyzes the deamination of c tosine (C) in a narrow window of the non-target and makes the base change from C to U (uracil) at a target site. U is recognized as thymine (T) during DNA replication, resulting in a C G to T A transition. () CRISPR/nCas9-mediated adenine base editing. An adenine base editor (ABE) is composed of an adenosine deaminase and nCas9(D10A) fusion binding to the target site in a gRNA-programmed manner. The adenosine deaminase catalyzes an A (adenine) to I (inosine) change at the target site. During replication, the original A is replaced with G (guanine). Finally, A T to G C conversion is achieved in the non-target DNA strand. () CRISPR/nCas9-mediated C-to-G base editing. The C-to-G base editor (CGBE) is composed of a c tosine deaminase, nCas9(D10A), and uracil N-glycosylase (UNG), and binds to the target site in a gRNA-programmed manner. The c tosine deaminase catalyzes the deamination of c tosine (C) and makes the base change from C to U (uracil). UNG can remove U from the DNA double strands and an error-prone DNA polymerase replaces G with C at the target site. The C G to G C transversion occurs during DNA replication. As nCas9(D10A) nicks the target strand, a DSB is formed when the abasic site on the non-target strand is converted into a nick by an apurinic or apirimidinic site lase (AP lase). The DSB results in indel formation at the target site. () The first-generation c tosine base editor, CBE1, was engineered by fusing c tosine deaminase, rAPOBEC1 to the N-terminus of a dead Cas9 (dCas9, a mutant of Cas9 containing both D10A and H840A mutations). () The second-generation base editor, CBE2, was engineered by fusing rAPOBEC1 to the N-terminus of dCas9 and fusing a uracil DNA glycosylase inhibitor (UGI) to the C-terminus of dCas9. () The third-generation c tosine base editor, CBE3, was engineered by fusing different deaminases to the N-terminus of nCas9(D10A), and fusing UGI to the C-terminus of nCas9(D10A), respectively. The deaminases that have been successfully applied in plants include rAPOBEC1 (Li et al., 2017; Lu and Zhu, 2017; Zong et al., 2017), PmCDA1 (Shimatani et al., 2017; Zhong et al., 2019), hAID (Ren et al., 2018; Wang et al., 2020a), APOBEC3A (Zong et al., 2018), and evoFENRY (Zeng et al., 2020). () The fourth-generation c tosine base editor, CBE4, was developed by fusing two UGI molecules to the terminus of s



() The CRISPR/nCas9-mediated prime editing system. A prime editor mainly consists of a catalytically impaired nCas9(H840A), a M-MLV-RT (Molone murine leukemia virus reverse transcriptase), and a prime editing guide RNA (pegRNA). pegRNA is composed of three components, including a single-guide RNA (sgRNA) targeting the specific site, a reverse transcription template (RTT) encoding the desired edit, and a primer binding site (PBS) initiating RT. The nCas9(H840A)-M-MLV-RT and pegRNA complex bind to the target sequence in the genomic DNA in a sequence-specific manner. The M-MLV-RT helps the 3' DNA end from the PBS to prime the reverse transcription of an edit-encoding extension from pegRNA directly into the target site. () The first-generation prime editor, PE1, was engineered by fusing a wild M-MLV-RT to the N-terminus of nCas9(H840A). () The second-generation prime editor, PE2, was engineered by fusing an engineered M-MLV-RT with six amino acid mutations to the N-terminus of nCas9(H840A). () The third-generation prime editor, PE3, was engineered by using an additional sgRNA on the non-targeting strand. () The fourth-generation prime editor, PE4, was developed with co-expression of a dominant negative mismatch repair (MMR) protein (MLH1dn) on the basis of PE2. () The fifth-generation prime editor, PE5, was developed with transient co-expression of a dominant negative MMR protein (MLH1dn) on the basis of PE3. () PEmax was engineered by replacing nCas9(H840A) with a mutated version which harbors R221K and N394K mutations. () Overview of the design of twinPE or GRAND editor and the sequence replacement process. The single-strand DNAs (red and blue lines) produced by the paired pegRNAs containing RTTs highlighted in light red and light blue, respectively, bind to each other through their complementary ends highlighted in orange. The original 5' flaps were replaced by the annealed 3' flaps containing the edited DNA following DNA replication and repair.

CBE4, was developed by fusing two UGI molecules to the C-terminal of Cas9 nickase on the basis of CBE3 to enhance the inhibition of UNG (Komor et al., 2017) (Figure 1G). Compared with CBE3, CBE4 not only improves the base editing efficiency but also reduces the frequency of C to A or G transversions by 2.3 times. In addition, bacteriophage Mu Gam protein was added on the basis of CBE4 to construct a BE CBE4-Gam, in order to further improve the product purity and reduce the occurrence of indels (Komor et al., 2017) (Figure 1H).

CBEs, especially CBE3 and CBE4, have been widely used in plants. Initially, a base editing system was developed using a rat APOBEC1 in rice. To validate and test the feasibility of CBEs in plants, by fusing a rat APOBEC1 to the N-terminus of nCas9 (D10A) to form a structure of rAPOBEC1-nCas9 (D10A), two agriculturally important genes of rice, OsNRT1.1B and OsSLR1, were edited at editing efficiencies of 2.7% and 13.3%, respectively (Lu and Zhu, 2017) (Table 1). Simultaneously, three targets in rice, one target (P2)

Table 1. Base editing in plants

Base editors	Structures of various base editors	Species	Target gene	Transformation	Selection	Editing efficiency (%)	Editing window	Trait improvement	References
APOBEC1-CBE2/ CBE3	APOBEC1-XTEN-nCas9	Rice	OsNRT1.1B, OsSLR1	Agrobacterium	H grom cin	2.70–13.30	C4–C8	High nitrogen use efficienc ; Dwarf	Lu and Zhu (2017)
	APOBEC1-XTEN-nCas9-UGI	Rice	OsSBEIIb, OsPDS	Bombardment	H grom cin	20.00	C4–C8	High am lose	Li et al. (2017)
	APOBEC1-XTEN-nCas9/ dCas9-UGI	Rice, Wheat Mai e	OsCDC48, OsNRT1.1B, OsSPL14, TaLOX2, ZmCENH3	Agrobacterium /Bombardment	Herbicide	0–43.48	C3–C9	High nitrogen use efficienc ; High ield	Zong et al. (2017)
	APOBEC1-XTEN-nCas9-UGI	Rice	OsCERK1, OSSERK1, OSSERK2, ipa1, Pi-ta	Agrobacterium	H grom cin	10.50–38.90	C4–C8	High ield; Blast resistance	Ren et al. (2017)
	APOBEC1-XTEN-nCas9-UGI	Wheat	TaALS-P174	Bombardment	Herbicide	33.00–75.00	C4–C8	Herbicide resistance	Zhang et al. (2019)
	APOBEC1-XTEN-nCas9-UGI	Cotton	GhCLA, GhPEBP	Agrobacterium	H grom cin	0–57.78	C4–C8	–	Qin et al. (2020)
hAID-CBE3	hAID-XTEN-nCas9	Rice	OsFLS2, OSAOS1, OsJAR1, OsJAR2, OSCO12, OsPi-D2	Agrobacterium	H grom cin	8.30–73.30	C3–C8	Blast resistance	Ren et al. (2018)
	hAID-nSpCas9-NG/nCas9-UGI	Rice	OsBZR1, OSSERK2	Agrobacterium	H grom cin	4.44–27.08	C3–C8	Enhance fruit qualit	Ren et al. (2019)
	hAID-nScCas9-UGI	Rice	OsMPK9, OsMPK17, OsCPK5, OsMPK15, OsMPK16, OsCPK6, OsCPK7, OsCPK8	Agrobacterium	H grom cin	2.56–97.92 0–95.83	C3–C8	–	Wang et al. (2020a)
	hAID-nSpCas9-UGI	Rice	OsCO12, OsBSR, OsMPK13, OSGS1, OSGSK4	Agrobacterium	H grom cin	26.00–34.15	C3–C8	–	Xu et al. (2021c)
	hAID-XTEN-nSpRY-UGI	Rice	OsAAT, OSCDC48, OsDEP1, OsNRT1, OSOD, TAALS, TaMTL, TaLOX2, STGBSS-T6	Agrobacterium /Bombardment	Herbicide/ H grom cin	0–82.90 1.20–20.00	C1–C17	Herbicide resistance	Zong et al. (2018)
	A3A-XTEN-nCas9-UGI; Gam-XTEN-A3A-nCas9-UGI	Rice , Potato	ALS, DELLA, ETR1.	Agrobacterium	Herbicide/ H grom cin	26.20–53.80	C2–C5	Herbicide resistance	Shimatani et al. (2017)
	nCas9/dCas9-PmCDA1	Rice , Tomato	OsDEP1, OsCDC48, OsGS3, OsPDS	Agrobacterium	H grom cin	0–21.10 3.50–56.30	C2–C5	–	Zhong et al. (2019)
	PmCDA1-xCas9-UGI	Tomato	SiALS	Agrobacterium	H grom cin	32.00	C2–C5	Herbicide resistance	Veillet et al. (2020)
	nCas9-NG-PmCDA1-UGI	potato	OsWaxy, OsEUI1	Agrobacterium	H grom cin	8.3–86.1	C1–C17	Reduced am lose content	Liu et al. (2021b)
	PmCDA1-nScCas9 ⁺ -UGI-UGI	Rice	OsCKX2, OsWaxy, OsEUI1, OsSPL4, OsSPL7, OsSPL14,	Agrobacterium	H grom cin	0–86.30 0–59.40	C4–C12 C2–C5	–	Zeng et al. (2020)

Continued

Table 1. Continued

Base editors	Structures of various base editors	Species	Target gene	Transformation	Selection	Editing efficiency (%)	Editing window	Trait improvement	References
PmCDA1-nCas9-NG-UGI-UGI	PmCDA1-nCas9-NG-UGI-UGI		LF1, OsIAA13, OsMADS57, OsGBSSI			0–66.70	C4–C8		
ABE7.10	TadA-TadA7.10-nCas9	Rice, Wheat	OsALS, OsCDC48, OsAA ^T , OsDEP1, OsACC, OsNRT1.1B, OsEV, OsOD, TADEP1, TaEPSPS, TagW2	Agrobacterium /Bombardment	Herbicide/ H grom cin	3.20–59.10	A4–A8	Herbicide resistance	Li et al. (2018)
	TadA-TadA7.10-nCas9(D10A)		OsSERK2, OsMPK6, OsWRKY45, OsMPK13	Agrobacterium	H grom cin	6.45–62.26	A4–A8	–	Yan et al. (2018)
	TadA-TadA7.10-nCas9	Rice	OsSPL7, OsSPL14, OsSLR1, OsSPL4, OsSPL16, OsSPL17, OsSPL18	Agrobacterium	H grom cin	0–26.00 4.80–61.30	A4–A8	High yield	Hua et al. (2018)
	TadA-TadA7.10-nSpCas9	Rice	OsMPK14, OsCPK9, OsMPK15, OsCPK10	Agrobacterium	H grom cin	50.00–94.12	A4–A8	–	Wang et al. (2020a)
ABE-P1S	TadA7.10-nSpCas9	Rice	OsSERK2, OsSPL14, SLR1, Tms9-1, OsNRT1.1B, OsACC1, OsDEP1	Agrobacterium	H grom cin	4.50–96.30 0–61.10	A1–A12	High yield; Herbicide resistance	Hua et al. (2020b)
	TadA7.10-nSaCas9		OsEPSPS, OsALS, OsWaxy	Agrobacterium	Herbicide/ H grom cin	4.00–100.00 0–100.00	A4–A8	Herbicide resistance	Wei et al. (2021)
ABE8e	TadA8e(V106W)-nCas9	Rice							
	TadA8e(V106W)-nCas9-NG	Rice	OsCOI2, OsBSR, OsMPK13, OsGS1, OsGSK4	Agrobacterium	H grom cin	27.79–93.75	A3–A10	Herbicide resistance	Xu et al. (2021c)
	TadA8e-XTEN-nSpRY	Rice	OsSPL14, OsIAA13, OsSPL7, OsLF1, OsGBSSI, OsCKS2, OsEU1, OsTS	Agrobacterium	H grom cin	0–90.50 0–92.50 0–100.00	A1–A14 A1–A14 A1–A14	–	Tan et al. (2022)
	TadA8e-DBD-nSpG	Rice							
	TadA8e-DBD-nSpRY								
	TadA9-XTEN-nSpRY	Rice	OsMPK6, OsMPK13, OsSERK2, OsWRKY45, OsDEP2, OsETR2, OsGSK4, OsJAR1, OsGS1, OsALS1	Agrobacterium	Herbicide/ H grom cin	0–97.92 0–100.00 0–37.50 0–68.75	A1–A12 A4–A10 A3–A10 A4–A12	–	Yan et al. (2021)
	TadA9-XTEN-nSpCas9	Rice	OsALS, OsBADH2, OsLAZY1, OsPDS	Agrobacterium	Herbicide/ H grom cin	0.40–87.60	C2–C5 A4–A8	–	Xu et al. (2021a)
pDUBE1	TadA8e-nCas9-CDA1-UGI								

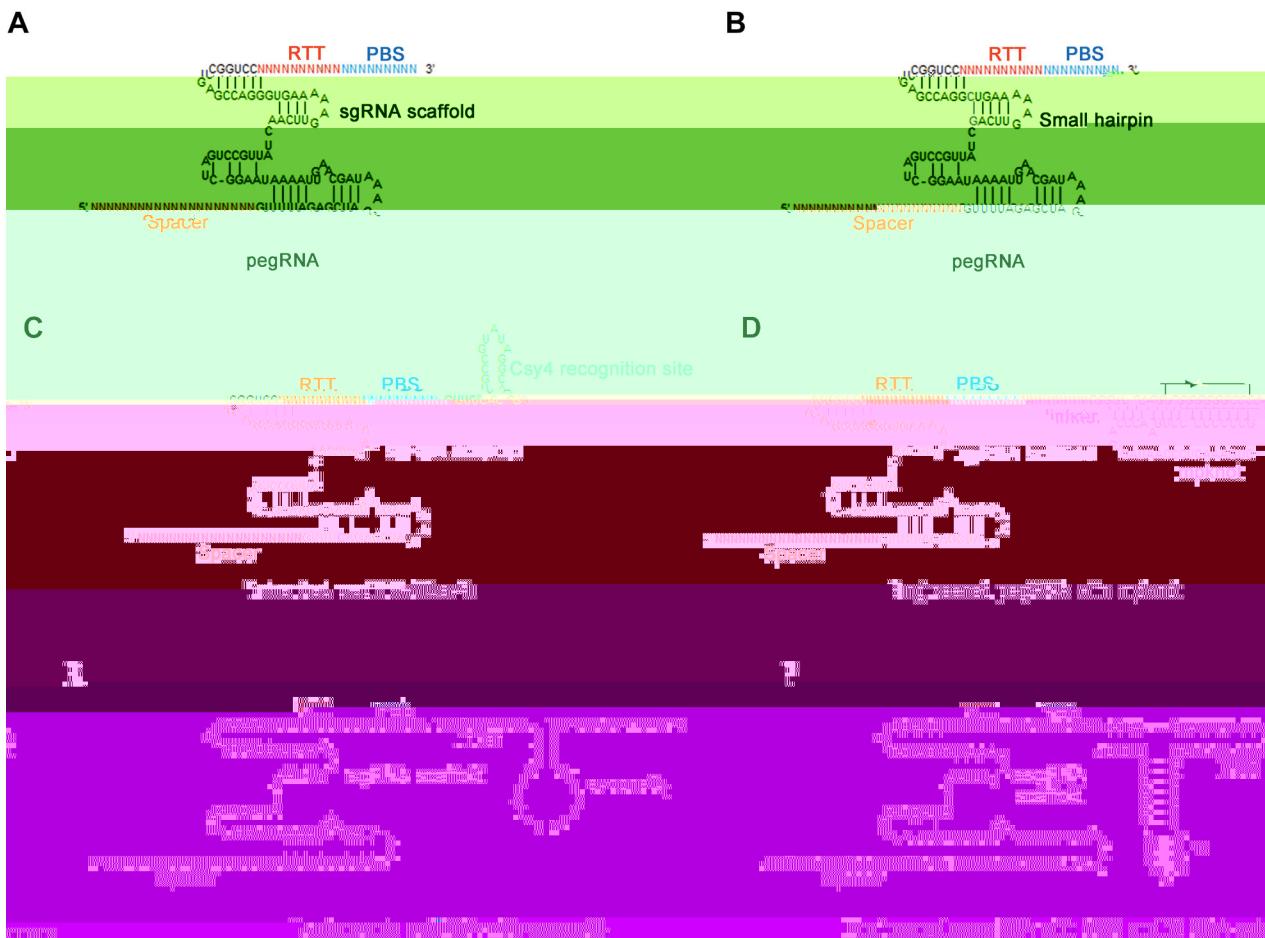
in OsPDS, which encodes a ph toene desaturase, and two targets (S3 and S5) in OsSBELlb, which encodes a starch branching en me llb, were successfull edited in rice b using CBE3 with the efficiencies of 19.2%, 10.5%, and 1.0% at the S5, S3, and P2 targets, respectivel (Li et al., 2017) (Table 1). Meanwhile, targeted C G to T A transitions in OsCDC48, OsSPL14, OsNRT1.1B, TaLOX2, ZmCENH3 genes were achieved at frequencies of up to 43.48% from position 3 to 9 within the protospacer in the genomes of rice, wheat and mai e b using a nCas9-c tidine deaminase fusion (Zong et al., 2017) (Table 1). Although the editing efficiencies of CBEs has been improved to a certain extent, its

enables c^tidine deamination was obtained b^r phage-assisted
continuous evolution (

Selecting the base-edited cells from massive transformed calli for regeneration is time-consuming and labor-intensive during plant tissue culture, especiaall for the low-efficient ed-

dsDNA, resulting in efficient C:G to T:A conversion in the human mitochondrial genome with high target specificity (Mok et al., 2020). Similar research was performed in lettuce (*Lactuca sativa*) and rapeseed (*Brassica napus*) protoplasts with up to 23% efficiencies (Kang et al., 2021). Recently, using the DdCBE linked to a plastid-targeting signal peptide (PTP) of AtRecA1 protein at its N-terminus, three target genes (16s rRNA, *rpoC1*, *psbA*) located in the plastid genome were successfully edited without leaving any foreign genes in either the plastid or nuclear genomes in *Arabidopsis* (Naka et al., 2021) (Figure 1O; Table 1). Furthermore, an efficient DdCBE construct was constructed by fusing a chloroplast transition peptide (CTP) to its N-terminus. This CTP-DdCBE achieved a conserved chloroplast gene chlorophyll A of photosystem I (*psaA*), for C to T transitions in rice chloroplasts (Li et al., 2021b) (Figure 1O; Table 1). The edited *psaA* could potentially improve photosynthetic efficiency and grain yield of crops. The successful implementation of DdCBEs (CRISPR-independent organelle BEs) in plant organelle cells increases the possibility of precise manipulation of organelle genomes for crop improvement (Kang et al., 2021).

Single nucleotide variation is the genetic basis for the improvement of important crop traits. Random mutagenesis by physical or chemical methods has long been applied to improve traits in plants, but it is labor-intensive and time-consuming. The base editing construct can enable the artificial evolution of agri-



() The schematic diagram of a canonical pegRNA. A pegRNA is composed of three components, including a single-guide RNA (sgRNA) targeting the specific site, a reverse transcriptase (RT) template (RTT) encoding the desired edit, and a primer binding site (PBS) initiating RT. The RTT sequence is highlighted in red, the PBS sequence is highlighted in blue, and the spacer sequence is highlighted in dark red. () The schematic diagram of a pegRNA, which has a C/G pair at the bottom of the small hairpin. The C/G base pair is highlighted in purple. () The schematic diagram of a Cs 4-processed pegRNA, which protects the 3' extension from degradation by exonucleases. Cs 4 is a specialized ribonuclease that selects clustered regular interspaced short palindromic repeats (CRISPR) transcripts from the cellular milieu for binding and cleavage. With Cs 4 processing, the hairpin Cs 4 recognition site remains at the 3' end of the pegRNA as an extension. At the same time, mutation of the fourth one of the consecutive uracils (highlighted in purple) was introduced to the scaffold of pegRNA. The Cs 4 recognition site sequence is highlighted in green. () An engineered pegRNA with a structured RNA pseudoknot (mpknot), protects its 3' extension from degradation by exonucleases. The mpknot is a frameshifting pseudoknot from Moloney murine leukemia virus (M-MLV), and it is an endogenous template for the M-MLV-RT from which the RT in canonical prime editors was engineered, raising the possibility that mpknot might help recruit the RT. The mpknot sequence is highlighted in pink. () An engineered pegRNA with a structured RNA pseudoknot evopreQ₁, which protects the 3' extension from degradation by exonucleases. evopreQ₁, as a modified preqeosine1-1 riboswitch aptamer composed of 42 nucleotides (nt) in length, is one of the smallest naturally derived RNA structural motifs with a defined tertiary structure. The evopreQ₁ sequence is highlighted in dark slate. (-) A representative engineered epegRNA with MS2 and f6 RNA aptamers. esgRNA, enhanced sgRNA with modifications highlighted in green. RNA aptamers can recruit their respective effector proteins for efficient gene editing. The MS2 sequence is highlighted in orange. The f6 sequence is highlighted in purple.

degradation of the 3' extension, and eventually improving the efficiency of prime editing by 3- to 4-fold in human cells without increasing off-target editing activity (Nelson et al., 2022) (Figure 3D, E). Extension of this strategy in plants significantly enhanced prime editing efficiency (Jiang et al., 2022b; Li et al., 2022b; Zou et al., 2022) (Table 2). In addition, MS2-based PE (MS2-PE) has also been developed to improve the prime editing efficiency by using RNA aptamers (MS2 and f6) in pegRNA and fusion of their binding protein MCP with the PE2's stem (Figure 3F), and achieved up to 10.1-fold increase in editing efficiency at five of six targets in transgenic rice lines (Chai et al., 2021).

While canonical PEs mainly enable base conversions and installation of small indels (An alone et al., 2019), development of PE capable of knock-in or replacement of large DNA fragments is highly desirable either for gene therapy or crop improvement. Recently, several powerful strategies have been developed to precisely replace, insert, and delete large DNA fragments in human cells, including twinPE (An alone et al., 2022) (Figure 2H), GRAND editing (Wang et al., 2022c) (Figure 2H), PRIME-Del (a prime editing-based method, which induces a deletion using a pair of pegRNAs that target opposite DNA strands) (Choi et al., 2022), and PEDAR (PE-Cas9-based

Table 2. Clustered regularly interspaced short palindromic repeats (CRISPR)/CRISPR-associate protein 9 (Cas9) mediated prime editing in plants

PEs	PE features (PBS length (nt) and RT template length (nt))	Target gene	Editing effinc			Transformation	Selection	Trait improvement	References
			Mutation t pe	Desired (%)	Undesired (%)				
PE2	13, 15–20	OsALS, OsIPA1, OstFB1	2–4 bp Subs, 2 bp Ins	0.00–2.04	NR	Agrobacterium	H grom cin, Bisp ribac sodium	Herbicide resistance, High yield	Butt et al. (2020)
Sp-PE2	13, 13	GFP	2 bp Subs	15.60	NR	Agrobacterium	H grom cin	-	Hua et al. (2020a)
pPE2	10–13, 10–34	HPTII, OsPDS, OsACC, OsWx	1–3 bp Ins, 1 bp Subs	0.00–59.90	NR	Agrobacterium	H grom cin	-	Xu et al. (2020a)
PE2	13, 16; 13, 14–23	OsALS, OsACC	1–3 bp Subs	1.00–7.60	0.00	Agrobacterium	H grom cin	Herbicide resistance	Jiang et al. (2022b)
pPE2	10–13, 11–18	OsPDS, OsACC, OsALS, OsCDC48	1 bp Ins, 1–2 bp Subs	0.00–29.17	NR	Agrobacterium	H grom cin	Herbicide resistance	Li et al. (2022b)
PPE	8, 17	OsALS	2 bp Subs	2.10	NR	Agrobacterium	H grom cin	Herbicide resistance	Zong et al. (2022)
pPE2 (an engineered pegRNA with mpknot)	10–13, 13–18	OsPDS, OsALS, OsCDC48	1 bp Ins, 1–2 bp Subs	10.42–25.00	NR	Agrobacterium	H grom cin	Herbicide resistance	Li et al. (2022b)
pPE2 (an engineered pegRNA with evopreQ _i)	10–13, 11–18	OsPDS, OsACC, OsALS, OsCDC48	1 bp Ins, 1–2 bp Subs	2.08–50.00	NR	Agrobacterium	H grom cin	Herbicide resistance	Li et al. (2022b)
ePPE(replacing M-MLV-RT with M-ΔRNaseH)	8, 17	OsALS	2 bp Subs	11.30	NR	Agrobacterium	H grom cin	Herbicide resistance	Zong et al. (2022)
pZ1WS (derived b the CaMV35S-CmYLCV/U6 composite promoter)	13, 16	ZmALS1, ZmALS2	2–3 bp Subs	4.80–53.20	NR	Agrobacterium	Glufosinate ammonium	Herbicide resistance	Jiang et al. (2020)
PE3	13, 13–16	AP01, GFP, OsALS	1–2 bp Subs	0.00–17.10	NR	Agrobacterium	H grom cin, Bisp ribac sodium	Herbicide resistance	Hua et al. (2020a)
Sp-PE3	13, 28–59	HPTII, OsEPSPS	3–7 bp Subs	2.22–9.38	NR	Bombardment	H grom cin	Herbicide resistance	Li et al. (2020c)

Continued

Table 2. Continued

PEs	PE features (PBS length (nt) and RT template length (nt))	Target gene	Mutation type	Editing effienc.		Transformation	Selection	Trait improvement	References
				Desired (%)	Undesired (%)				
PPE3	10–12, 9–17	OscDC48, OsALS	1–3 bp Subs, 6 bp Del	2.60–21.80	NR	Bombardment	H grom cin	Herbicide resistance	Lin et al. (2020)
pPPE3	13, 10	OsNxr, OsACC	1 bp Subs	0.00–16.70	NR	Agrobacterium	H grom cin	Herbicide resistance	Xu et al. (2020a)
PE-P1	11–14, 14–23	OsDEP1, OsALS, OsACC	1–4 bp Subs	0.00–1.40	0.00	Agrobacterium	H grom cin	Herbicide resistance	Xu et al. (2020c)
PE3	9–13, 9–28	OSSPL14, OsDHDPs, OsNR2	2–3 bp Subs	0.00–1.00	NR	Bombardment	H grom cin	Herbicide resistance	Li et al. (2022a)
PE3	13, 16; 13, 14–23	OsALS, OsACC, OsEPSPS	1–3 bp Subs	1.30–70.30	9.00–37.90	Agrobacterium	H grom cin	Herbicide resistance	Jiang et al. (2022b)
PPE3-unmodified	11–14, 11–18	OSROC, OsALS, OSCDC48, OsDEP1	1–3 bp Subs, 3 bp Ins (Normal)	0.00–2.90	NR	Agrobacterium	H grom cin	Herbicide resistance	Zou et al. (2022)
pPPE3b	13, 10	OsACC	1 bp Subs	6.25	NR	Agrobacterium	H grom cin	Herbicide resistance	Xu et al. (2020a)
PE-P2(nCas9 (H840A)-M-MLV-T2A-hpt)	11–14, 14–23	OsDEP1, OsALS, OsACC	1–4 bp Subs	1.70–26.00	0.00–8.00	Agrobacterium	H grom cin	Herbicide resistance	Xu et al. (2020c)
pCXPE03 (driven by the RPS5A promoter)	14, 17–18	SIGAI, SIALS, SPDS	2 bp Subs, 2 bp Ins	0.00–6.70	NR	Agrobacterium	H grom cin	Herbicide resistance	Lu et al. (2021)
PPE3-evopreQ ₁	11–14, 11–18	OSROC, OsALS, OSCDC48, OsDEP1	1–3 bp Subs, 3 bp Ins (Normal)	2.60–47.50 5.00–60.50 (HT)	NR	Agrobacterium	H grom cin	Herbicide resistance	Zou et al. (2022)
PPE3-mpknot	11–14, 11–18	OSROC, OsALS, OSCDC48, OsDEP1	1–3 bp Subs, 3 bp Ins (Normal)	0.00–4.20 0.00–6.30 (HT)	NR	Agrobacterium	H grom cin	Herbicide resistance	Zou et al. (2022)
PE-P2-RT-S (N-terminal M-MLV + a single desired mutation in RT)	8–14, 13–23	OSGS3, OsALS, OsACC, OsChlk5, OsDEP1, OsWay, OSGRF4, OSSD1, OsEPSPS, Oscold1, OsPSR1	1 bp Subs	0.00–61.40	0.00–15.00	Agrobacterium	H grom cin	-	Xu et al. (2022)
PE-P3-RT-M (C-terminal M-MLV + multiple non monobase mutations in RT)	8–14, 13–23	OSGS3, OsALS, OsACC, OsChlk5, OsDEP1, OsWay, OSGRF4, OSSD1, OsEPSPS, Oscold1, OsPSR1	3–4 bp Subs	0.00–82.60	0.00–18.00	Agrobacterium	H grom cin	-	Xu et al. (2022)

Table 2. Continued

Base editing and prime editing in plants

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PEs	PE features (PBS length (nt) and RT template length (nt))	Target gene	Mutation type	Editing effienc.		Transformation	Selection	Trait improvement	References
				Desired (%)	Undesired (%)				
PE-P3-RT-S C-terminal M-MLV + a single desired mutation in RTT)	8–14, 14–23	OsgS3, OsALS, OsACC, OsChalk5, OsDEP1, OsWaxy	1 bp Subs	0.00–22.70	0.00–15.00	Agrobacterium	H grom cin	-	Xu et al. (2022)
PE-P2-RT-M (N-terminal M-MLV + multiple s non mouse base mutations in RTT)	8–14, 14–23	OsgS3, OsALS, OsACC, OsChalk5, OsDEP1, OsWaxy	3 bp Subs	0.00–26.00	0.00–8.00	Agrobacterium	H grom cin	-	Xu et al. (2022)
PE3-HS (h grom - chinY46'-based)	9–13, 9–28; 13, 19	OssPL14+mhpII, OsDHDPs+mhpII, OsnR2+mhpII	2–3 bp Subs, 1 bp Subs	1.30–2.10	NR	Bombardment	H grom cin	-	Li et al. (2022a)
PE3-AS (OsALS-S6271-based)	9–13, 9–28; 13, 12	OssPL14+OsALS, OsDHDPs+OsALS, OsnR2+OsALS, OssPL14+OsALS +OsDHDPs	2–3 bp Subs, 3 bp Subs	2.40–14.30	NR	Bombardment	H grom cin, BisP ribac sodium	-	Li et al. (2022a)
PE3-DS (PE3-HS +PE3-AS)	9–13, 9–28; 13, 12; 13, 19	OssPL14+OsALS +mhpII, OsDHDPs +OsALS+mhpII, OsnR2+OsALS+ mhpII, OssPL14 +OsALS+OsdHDPs+mhpII, OssPL14+OsALS+OsePSPS+mhpII	2–7 bp Subs, 3 bp Subs, 1 bp Subs	3.20–54.20	NR	Bombardment	H grom cin, BisP ribac sodium	-	Li et al. (2022a)
PE4	13,16; 13, 14–23;	OsALS, OsACC	1–3 bp Subs	5.20–27.10	0.00–2.10	Agrobacterium	H grom cin	Herbicide resistance	Jiang et al. (2022b)
PE5	13,16; 13, 14–23;	OsALS, OsACC, OsEPSPS	1–3 bp Subs	1.60–64.10	6.40–18.30	Agrobacterium	H grom cin	Herbicide resistance	Jiang et al. (2022b)
PEmax	13, 22	OsePSPS	3 bp Subs	37.20–39.80	18.40–21.00	Agrobacterium	H grom cin	Herbicide resistance	Jiang et al. (2022b)

Continued

Table 2. Continued

PEs	PE features (PBS length (nt) and RT template length (nt))	Target gene	Editing efficienc.			Transformation	Selection	Trait improvement	References
			Mutation t pe	Desired (%)	Undesired (%)				
PEmax	13, 22	OsEPSPS	3 bp Subs	38.20-39.50	20.80-20.90	Agrobacterium	H grm cin	Herbicide resistance	Jiang et al. (2022b)

ePPE5max (ePPE3max+ OsmLH1dn)
pPPE2max-evopreQ₁ (PEmax+an)
.....

deletion and repair) method (Jiang et al., 2022a). These prime editing stems were developed by employing similar strategies such as using a pair of designed pegRNAs that target the opposite DNA strands, and the RTTs from the two respective pegRNAs were nonhomologous to the target sites but partially complementary to each other (An alone et al., 2022; Choi et al., 2022; Jiang et al., 2022a; Wang et al., 2022c). Although the above stems for replacement or knock-in of large DNA fragments through prime editing had been successfully applied in mammalian cells, the feasibilities of twinPE, GRAND editing, PRIME-Del, and PEDAR in plant prime editing remain to be investigated in the near future.

Applications of diverse PEs in plants

Prime editing stem substantially expands the scope and capabilities of precision genome editing and holds great promise to introduce precise genome modifications such as SNP and/or small indels into plant genomes to improve agriculturally important traits in crops (Li et al., 2020d). Since the first report of prime editing in mammalian cells in 2019 (An alone et al., 2019), the feasibilities and efficacies of PE2 and PE3 for precise genome editing had soon been investigated in rice (Butt et al., 2020; Hua et al., 2020a; Jiang et al., 2020; Li et al., 2020c; Lin et al., 2020; Tang et al., 2020; Xu et al., 2020a, 2020c), other plant species (Jiang et al., 2020; Lin et al., 2020; Lu et al., 2021), and followed by further optimization to improve their prime editing efficiencies thereafter (Jiang et al., 2020; Li et al., 2020c; Lu et al., 2021; Xu et al., 2022; Li et al., 2022a; Jiang et al., 2022b; Xu et al., 2020a, 2020c) (Table 2). Applications of diverse PEs in plants, their features and editing efficiencies and so forth, are summarized in Table 2.

The feasibility and efficacy of a series of plant codons optimized from PE2 and PE3 were first validated and investigated in rice and wheat protoplasts (Lin et al., 2020; Tang et al., 2020) or stable rice plants (Li et al., 2020c; Xu et al., 2020a, 2020c) almost simultaneously in five laboratories. Except for the intrinsic nature of target genes, various parameters such as PBS length, RT template length, and the position of nicking sgRNA significantly affected the precise editing efficiency of PE2 and PE3 in rice and wheat protoplasts (Lin et al., 2020). The PE2 stem could also induce programmable editing at different genome sites at a frequency of 0% to 31.3% in rice stable lines, suggesting that the efficiency of pPPE2 varied greatly at different genomic sites and with pegRNAs of diverse structures (Xu et al., 2020a). By using the polII promoter Actin to drive the expression of the tandem repeats of polycistronic transfer RNAs to simultaneously produce pegRNA and nicking sgRNA in a PE3, 28 bp and a 59 bp fragments with desired edits were precisely installed into an exogenous inactive hexokinase phosphotransferase (HPT) gene hptII to restore its function, and an endogenous gene OsEPSPS, which encodes a 5-enolpyruvylshikimate-3-phosphate synthase (EPSPS), to generate a novel allele with TAP-IVS mutations (T173I, A174V, and P177S) which confers rice glufosinate resistance in rice stable lines, respectively (Li et al., 2020c).

Furthermore, development of a plant PE b fusion of HPT to the C-terminus of nCas9-M-MLV with a self-cleaving 2A peptide (P2A) linker and paired with an enhanced sgRNA (esgRNA) improved the prime editing efficiency and achieved versatile nucleotide substitutions in rice stable lines (Xu et al., 2020c). Compared with normal PE, this strategy could increase the editing efficiency up to 22-fold at the OsALS-1 site (from 1.20% to 26.00%) (Xu et al., 2020c). Later on, a PE2-based plant PE with a pegRNA of 13-nt PBS and 15-nt RTT was transformed into rice to alter the target codon TGG for Trp548 of the ALS gene generated herbicide resistant rice plants (Butt et al., 2020). At the same time, prime editing of OsIPA1 in rice reduced the number of unproductive tillers and improved rice yield (Butt et al., 2020). An inactive eGFP gene was used as a transgenic reporter. After prime editing, it was restored into a wild-type EGFP sequence with two precise base conversions (T-G and G-C) at efficiencies of 15.60% and 17.10% for PE2 and PE3, respectively (Hua et al., 2020a). Simultaneously, by using a pegRNA designed to introduce a S627N mutation in OsALS, PE3 was also employed to gen-

ALS2 and PDS1 were obtained at the efficiencies of 6.70% and 3.40%, respectively (Lu et al., 2021) (Table 2).

Directed evolution (DE) is a technology of making random mutation(s) in a target gene to generate novel germplasms and enrich genetic diversity (Zhang and Qi, 2019). Currently, base editing can enable artificial evolution of agriculturally important genes in crops to explore novel gene resources and germplasms (Kuang et al., 2020; Li et al., 2020a; Liu et al., 2020; Xu et al., 2021a; Wang et al., 2022b). Compared with base editing, prime editing has greater potential for evolving plant genes, because it can install all types of small genetic modifications that can be harnessed for producing all possible substitutions for key amino acids with improved agronomic performance when combined with a well-designed pegRNA library. For example, a prime editing library-mediated saturation mutagenesis (PLSM) method had been developed to identify 16 types of herbicide resistance-conferring mutations at six different target residues in OsACC1 using a pegRNA library with all possible combinations of substitutions (64 types), which enabled a more comprehensive screening than that achieved by base editing (Xu et al., 2021b). Among the 16 kinds of mutations, three types of mutations were first reported in plants. The PLSM's stem is an alternative approach to create novel germplasms for crop breeding.

RECOMMENDATION OF PROPER BE AND PE FOR BOTH BASIC RESEARCH IN PLANTS AND CROP IMPROVEMENT

According to optimizations and applications of the current BEs and PEs in plants as described above, we recommend to select appropriate BEs and PEs for precise gene editing for both basic biological research in plants and crop improvement. For base editing, we recommend using evoFERNY for CBE (Zeng et al., 2020), and TadA9 or TadA8e-DBD (hTadA8e) for ABE (Yan et al., 2021; Tan et al., 2022) in base editing. In addition, it would be good to select the appropriate Cas protein capable of targeting the region near the desired editing site due to the limitations of PAM sites and the editing windows of various BEs. At present, the suitable choice is to select the editors constructed from these three Cas proteins with broad PAM sites, including Cas9-NG (NGN PAM), ScCas9⁺⁺ (NNG PAM), and SpRY (NNN prefer NRN PAM) (Hua et al., 2019; Ren et al., 2019; Zhong et al., 2019; Wang et al., 2020a; Liu et al., 2021b). Moreover, using a surrogate stem to restore the defective genes into the functional ones encoding antibiotics or herbicides could be more cost-effective and improve the base editing efficiency (Xu et al., 2020b).

For prime editing, we recommend using PEmax or a PE with M-MLV-RT fused to the N-terminus of nCas9(H840A) (Chen et al., 2021; Jiang et al., 2022b; Li et al., 2022b; Xu et al., 2022) (Figure 2). Further, except the intended base

substitutions, introduction of additional multiple-nucleotide substitutions in RTT could stimulate prime editing efficiency (Chen et al., 2021; Li et al., 2022c; Xu et al., 2022). For example, some non-mutant substitutions could be introduced at +1 to +6 positions (counting 3'-base of RTT as position +1), in order to avoid the repeat nicking of edited targets (Xu et al., 2022) (Figure 3). Furthermore, additional structured RNA sequences, such as evopreQ₁ appended to the 3'-end of pegRNA will stabilize the pegRNAs and thus improve the prime editing efficiency (Li et al., 2022b; Nelson et al., 2022; Zou et al., 2022) (Figure 3). Moreover, a strong composite promoter, such as CaMV 35S enhancer+CmYLCV promoter+U6 promoter, could be used to enhance the expression of the pegRNA and thus improve the prime editing efficiency (Jiang et al., 2020). Lastly, using the reporter genes such as antibiotics or herbicides as surrogates to enrich the lines with desired edits improved the prime editing efficiency in a cost-effective and labor-saving way, especially for multiplex prime editing in plants (Li et al., 2022a) (Table 2).

FUTURE PERSPECTIVES FOR FURTHER OPTIMIZATION OF BE AND PE IN PLANTS

Although impressive progresses have been made during the last several years, the following aspects such as optimization of the existing BEs, exploitation of novel BEs, and optimization of PEs to further improve their precise editing efficiencies as well as developing novel PEs capable of installation of larger indels in plants, would be highly desirable in the next few years.

Optimization of the existing BEs and exploitation of novel BEs

To date, CBE and ABE for base transition have been well optimized in terms of improving editing efficiency, expanding the target scope and reducing off-targets. However, for base transversion, such as CGBE for C to G and C to A, the editing efficiency is relatively lower in comparison with other BEs in plants (Koblan et al., 2021; Tian et al., 2022). Thus, it is still necessary to increase the efficiency of CGBE. Most importantly, in order to increase the flexibility of BEs, exploitation of other types of BEs for transversion of A to C (T to G) or A to T (T to A) will certainly be very beneficial in substitution of an base pair into the desired one within the editing window in a target gene of interest in plants. In addition, concerning the base editing window, two aspects are worth of further optimization. (i) Narrow the editing window of BE to a single base, reduce the by-products of unintended editing, for example, a more precise adenine base editor ABE9 (Figure 1L), which was developed recently by introducing two mutations L145T and N108Q in ABE8e, maintained the editing activity and minimized the editing window to position 5–6 in mammalian cells (Chen et al., 2022b). Furthermore, by combining with PAM-less Cas proteins, it will be possible to

achieve accurate single base editing at an target sites in the genome. (ii) Widen the width of the editing window of BE for saturation mutation studies such as de novo domestication or DE to generate novel gene resources or germplasm in plants. For example, fusion of T7 RNA pol merase with different deaminases (c^tidine and adenosine deaminase), substantially widens the mutational spectrum in mammalian cells (Cravens et al., 2021). In addition, engineering BEs fused with additional chromatin modulating peptides, such as pioneer factor SOX2 (SRY-box transcription factor 2), to initiate chromatin unfolding and stimulate transcription, could be a promising strategy to further increase base editing efficacy (Yang et al., 2022).

Optimization of PEs

A series of parameters such as stable and properly folded pegRNAs, effective assembly of the PE-pegRNA complex, and more active reverse transcriptase are essential for efficient prime editing. In PE, the canonical pegRNA consists of a sgRNA, a RTT and a PBS (Figure 2A). PBS and RTT at the 3'-terminal of pegRNA are easily to partially degrade by exoribonucleases inside the cells, resulting in truncated pegRNAs (Feng et al., 2022; Nelson et al., 2022). The truncated pegRNAs can still search and recognize the target sites, but not be able to complete the correct editing due to loss of the PBS or RTT-PBS (Nelson et al., 2022). Adding a

PEs, especially PE, are not widely used or even impossible in the polyploid species and agriculturally important food crops such as common wheat due to its complex hexaploid genome, gene redundancy, as well as relatively lower transformation efficiency (Li et al., 2021c). Third, for base editing and prime editing in different plant species, we suggest using the aforementioned optimized strategies in combination with a stronger promoter to drive the expression of both nCas-deaminase and the sgRNA for BE, or nCas-M-MLV-RT and pegRNA for PE, respectively (Li et al., 2022b). Finally, it is worth noting that the innate nature of target genes may affect the editing outcomes of both BEs and PEs in plants; for example, some genes or targets could only be edited at a very low efficiency or even not be accessible (Hua et al., 2022). Understanding the potential mechanism underlying this phenomenon will certainly benefit the precision genome editing of any targets at will in a user-defined manner in plants. Nevertheless, following the continuous endeavors on optimization of BE and PE as well as engineering a novel generation of BE and PE, we envision that both BEs and PEs will become the routine and customized precise gene editing tools for both plant fundamental research and crop improvement in the near future.

- (2020). Directed evolution of adenine base editors with increased activity and therapeutic application. *Nat. Biotechnol.* 38, 892–900.
- (1999). Unusual transcriptional and translational regulation of the bacteriophage Mu *lambdoid* operon. *Pharmacol. Ther.* 83, 367–388.
- (2015). Variation in NRT1.1B contributes to nitrate-use divergence between rice subspecies. *Nat. Genet.* 47, 834–838.
- (2022). Improvement of base editors and prime editors advances precision genome engineering in plants. *Plant Physiol.* 180, 1795–1810.
- (2020a). Precision genome engineering in rice using prime editing's stem. *Plant Biotechnol. J.* 18, 2167–2169.
- (2019). Genome engineering in rice using Cas9 variants that recognize NG PAM sequences. *Mol. Plant* 12, 1003–1014.
- (2020b). Simplified adenine base editors improve adenine base editing efficiency in rice. *Plant Biotechnol. J.* 18, 770–778.
- (2018). Precise A to G/C base editing in the rice genome. *Mol. Plant* 11, 627–630.
- (2021). CRISPR/Cas9s stems: The link between functional genes and genetic improvement. *Crop J.* 10, 678–687.
- (2021). Adapting CRISPR/Cas9's stem for targeting mitochondrial genome. *Front. Genet.* 12, 627050.
- (2022a). Deletion and replacement of long genomic sequences using prime editing. *Nat. Biotechnol.* 40, 227–234.
- (2022b). Optimized prime editing efficiently generates gli phosate-resistant rice plants carrying homozygous TAP-IVS mutation in EPSPS. *Mol. Plant* 15, 1646–1649.
- (2020). Prime editing efficiently generates W542L and S621I double mutations in two ALS genes in maize. *Genome Biol.* 21, 257.
- (2010). Regulation of OsSPL14 by OsmiR156 defines ideal plant architecture in rice. *Nat. Genet.* 42, 541–544.
- (2019). Cytosine, but not adenine, base editors induce genome-wide off-target mutations in rice. *Science* 365, 292–295.
- (2012). A programmable dual-RNA-guided DNA endonuclease in adaptive bacterial immunity. *Science* 335, 816–821.
- (2021). Chloroplast and mitochondrial DNA editing in plants. *Nat. Plants* 7, 899–905.
- (2019). Curing cytoplasmic male sterility via TALEN-mediated mitochondrial genome editing. *Nat. Plants* 5, 722–730.
- (2021). Predicting the efficiency of prime editing guideRNAs in human cells. *Nat. Biotechnol.* 39, 198–206.
- (2018). CRISPR-Cas guides the future of genetic engineering. *Science* 360, 866–869.
- (2021). Efficient C/G-to-G/C base editors developed using CRISPRi screens, target library analysis, and machine learning. *Nat. Biotechnol.* 39, 1414–1425.
- (2018). Improving cytidine and adenine base editors by expression optimization and ancestral reconstruction. *Nat. Biotechnol.* 36, 843–846.
- (2016). Programmable editing of a target base in genomic DNA without double-stranded DNA cleavage. *Nature* 534, 420–424.
- (2017). Improved base excision repair inhibition and bacteriophage Mu Gam protein yields C:G-to-T:A base editors with higher efficiency and product purity. *Sci. Adv.* 3, eaao4774.
- (2020). Base-editing-mediated artificial evolution of OsALS1 in planta to develop novel herbicide-tolerant rice germplasms. *Mol. Plant* 13, 565–572.
- (2021). CRISPR C-to-G base editors for inducing targeted DNA transversions in human cells. *Nat. Biotechnol.* 39, 41–46.
- (2020a). Targeted, random mutagenesis of plant genes with dual cytosine and adenine base editors. *Nat. Biotechnol.* 38, 875–882.
- (2020b). SWISS: multiplexed orthogonal genome editing in plants with a Cas9 nickase and engineered CRISPR RNA scaffolds. *Genome Biol.* 21, 141.
- (2018). Expanded base editing in rice and wheat using a Cas9-adenosine deaminase fusion. *Genome Biol.* 19, 59.
- (2020c). Precise modifications of both exogenous and endogenous genes in rice by prime editing. *Mol. Plant* 13, 671–674.
- (2022a). Multiplex precision gene editing by a surrogate prime editor in rice. *Mol. Plant* 15, 1077–1080.
- (2022b). Development of a highly efficient prime editor 2's stem in plants. *Genome Biol.* 23, 161.
- (2020d). Toward precision genome editing in crop plants. *Mol. Plant* 13, 811–813.
- (2017). Generation of targeted point mutations in rice by a modified CRISPR/Cas9's stem. *Mol. Plant* 10, 526–529.
- (2021a). Genome editing mediated by SpCas9 variants with broad non-canonical PAM compatibility in plants. *Mol. Plant* 14, 352–360.
- (2021b). High-efficiency plastome base editing in rice with TAL cytosine deaminase. *Mol. Plant* 14, 1412–1414.
- (2019). Precise gene replacement in rice by RNA transcript-templated homologous recombination. *Nat. Biotechnol.* 37, 445–450.

- (2020). Precise gene replacement in plants through CRISPR/Cas genome editing technology: Current status and future perspectives. *aBIOTECH*, 58–73.
- (2021c). Present and future prospects for wheat improvement through genome editing and advanced technologies. *Plant Commun.*, 100211.
- (2022c). High efficient prime editing by introducing same-sense mutations in pegRNA or stabilizing its structure. *Nat. Commun.*, 1669.
- (2021d). The CRISPR/Cas9 revolution continues: From base editing to prime editing in plant science. *J. Genet. Genomics*, 661–670.
- (2001). Translational repression and specific RNA binding by the coat protein of the *Pseudomonas* phage PP7. *J. Biol. Chem.*, 22507–22513.
- (1994). Mutations that increase the affinity of a translational repressor for RNA. *Nucleic Acids Res.*, 3748–3752.
- (2021). High-efficient prime editing with optimized paired pegRNAs in plants. *Nat. Biotechnol.*, 923–927.
- (2020). Prime genome editing in rice and wheat. *Nat. Biotechnol.*, 582–585.
- (2021a). Developing a novel artificial rice germplasm for dinitroaniline herbicide resistance by base editing of OsTubA2. *Plant Biotechnol. J.*, 5–7.
- (2021b). The ScCas9(++) variant expands the CRISPR toolbox for genome editing in plants. *J. Integr. Plant Biol.*, 1611–1619.
- (2020). A CRISPR-Cas9-mediated domain-specific base-editing screen enables functional assessment of ACCase variants in rice. *Plant Biotechnol. J.*, 1845–1847.
- (2021c). Enhancing prime editing by Cas 4-mediated processing of pegRNA. *Cell Res.*, 1134–1136.
- (2020). Targeted, efficient sequence insertion and replacement in rice. *Nat. Biotechnol.*, 1402–1407.
- (2021). Precise genome modification in tomato using an improved prime editing stem. *Plant Biotechnol. J.*, 415–417.
- (2017). Precise editing of a target base in the rice genome using a modified CRISPR/Cas9 stem. *Mol. Plant*, 523–525.
- (2020). CRISPR/Cas stems in genome editing: Methodologies and tools for sgRNA design, off-target evaluation, and strategies to mitigate off-target effects. *Adv. Sci.*, 1902312.
- (2019). Gene editing in plants: Progress and challenges. *Natl. Sci. Rev.*, 421–437.
- (2020). A bacterial Ctidine deaminase toxin enables CRISPR-free mitochondrial base editing. *Nature*, 631–637.
- (2021). Precise plant genome editing using base editors and prime editors. *Nat. Plants*, 1166–1187.
- (2000). RecE/RecT and Redalpha/Redbeta initiate double-stranded break repair by specifically interacting with their respective partners. *Genes Dev.*, 1971–1982.
- (2021). Targeted base editing in the plastid genome of *Arabidopsis thaliana*. *Nat. Plants*, 906–913.
- (2022). Engineered pegRNAs improve prime editing efficiency. *Nat. Biotechnol.*, 402–410.
- (2022). Evolution of an adenine base editor into a small, efficient cytosine base editor with low off-target activity. *Nat. Biotechnol.*, <https://doi.org/10.1038/s41587-022-01533-6>
- (2016). Targeted nucleotide editing using hybrid prokaryotic and vertebrate adaptive immune systems. *Science*, aaf8729.
- (1998). DNA strand invasion promoted by Escherichia coli RecT protein. *J. Biol. Chem.*, 12274–12280.
- (1993). The RNA binding site of bacteriophage MS2 coat protein. *EMBO J.*, 595–600.
- (2022). Prime editing in the model plant *Physcomitrium patens* and its potential in the tetraploid potato. *Plant Sci.*, 111162.
- (2022). Updates on gene editing and its applications. *Plant Physiol.*, 1725–1730.
- (2020). High-efficient and precise base editing of C*G to T*A in the allotetraploid cotton (*Gossypium hirsutum*) genome using a modified CRISPR/Cas9 stem. *Plant Biotechnol. J.*, 45–56.
- (2019). Cas9-NG greatly expands the targeting scope of the genome-editing toolkit by recognizing NG and other atypical PAMs in Rice. *Mol. Plant*, 1015–1026.
- (2017). A CRISPR/Cas9 toolkit for efficient targeted base editing to induce genetic variations in rice. *Sci. China Life Sci.*, 516–519.
- (2018). Improved base editor for efficiently inducing genetic variations in rice with CRISPR/Cas9-guided hyperactive hAID mutant. *Mol. Plant*, 623–626.

- (2022). A super pan-genomic landscape of rice. *Cell Res.*, 32, 878–896.
- (2017). Targeted base editing in rice and tomato using a CRISPR-Cas9 ctidine deaminase fusion. *Nat. Biotechnol.*, 35, 441–443.
- (2017). Deep mutational scanning of *S. pyogenes* Cas9 reveals important functional domains. *Sci. Rep.*, 7, 16836.
- (2016). Engineering herbicide-resistant rice plants through CRISPR/Cas9-mediated homologous recombination of acetolactate synthase. *Mol. Plant*, 9, 628–631.
- (2022). PhiCas9-ABE: A PAM-less/free high-efficiency adenine base editor toolbox with wide target scope in plants. *Plant Biotechnol. J.*, 10, 934–943.
- (2019). Off-target effects and the solution. *Nat. Plants*, 5, 341–342.
- (2020). Plant prime editors enable precise gene editing in rice cells. *Mol. Plant*, 13, 667–670.
- (2022). Efficient C-to-G editing in rice using an optimized base editor. *Plant Biotechnol. J.*, 10, 1238–1240.
- (2020). Expanding the CRISPR toolbox in *P. patens* using SpCas9-NG variant and application for gene and base editing in Solanaceae crops. *Int. J. Mol. Sci.*, 21, 1024.
- (2020). Unconstrained genome targeting with near-PAMless engineered CRISPR-Cas9 variants. *Science*, 368, 290–296.
- (2021a). Microbial single-strand annealing proteins enable CRISPR gene-editing tools with improved knock-in efficiencies and reduced off-target effects. *Nucleic Acids Res.*, 49, e36.
- (2022a). dCas9-based gene editing for cleavage-free genomic knock-in of long sequences. *Nat. Cell Biol.*, 30, 268–278.
- (2022b). Base editing-mediated targeted evolution of ACCase for herbicide-resistant rice mutants. *J. Integr. Plant Biol.*, 64, 2029–2032.
- (2022c). Efficient targeted insertion of large DNA fragments without DNA donors. *Nat. Methods*, 19, 331–340.
- (2018a). A single transcription factor promotes both field and immunity in rice. *Science*, 360, 1026–1028.
- (2021b). Splicing changes and fluorescent tagging with prime editing vectors for plants. *Front. Genome Ed.*, 2, 617553.
- (2020a). Targeted base editing in rice with CRISPR/SgCas9 s stem. *Plant Biotechnol. J.*, 18, 1645–1647.
- (2020b). Cas12a base editors induce efficient and specific editing with low DNA damage response. *Cell Rep.*, 32, 107723.
- (2019). CRISPR technology is revolutionizing the improvement of tomato and other fruit crops. *Hortic. Res.*, 7, 77.
- (2018b). Genomic variation in 3,010 diverse accessions of Asian cultivated rice. *Nature*, 555, 43–49.
- (2021). Efficient generation of homozygous substitutions in rice in one generation utilizing an rAbe8e base editor. *J. Integr. Plant Biol.*, 63, 1595–1599.
- (2021). The power and versatility of genome editing tools in crop improvement. *J. Integr. Plant Biol.*, 63, 1591–1594.
- (2017). CRISPR-GE: A convenient software toolkit for CRISPR-based genome editing. *Mol. Plant*, 10, 1246–1249.
- (2019). Genome editing for horticultural crop improvement. *Hortic. Res.*, 7, 113.
- (2021a). Development of an efficient plant dual C-to-Tosine and adenine editor. *J. Integr. Plant Biol.*, 63, 1600–1605.
- (2020a). Development of plant prime-editing stems for precise genome editing. *Plant Commun.*, 1, 100043.
- (2021b). Identification of herbicide resistance *OsACC1* mutations via *in planta* prime-editing library screening in rice. *Nat. Plants*, 7, 888–892.
- (2020b). Discriminated sgRNAs-based surrogate stems greatly enhances the screening efficiency of plant base-edited cells. *Mol. Plant*, 13, 169–180.
- (2022). A design optimized prime editor with expanded scope and capability in plants. *Nat. Plants*, 8, 45–52.
- (2020c). Versatile nucleotide substitution in plants using an improved prime editing stem. *Mol. Plant*, 13, 675–678.
- (2021c). SpRY greatly expands the genome editing scope in rice with highly flexible PAM recognition. *Genome Biol.*, 22, 6.
- (2021). High-efficiency and multiplex adenine base editing in plants using new TadA variants. *Mol. Plant*, 14, 722–731.
- (2018). Highly efficient A-T to G-C base editing by Cas9-guided tRNA adenosine deaminase in rice. *Mol. Plant*, 11, 631–634.
- (2022). Pioneer factor improves CRISPR-based C-To-G and G-To-T base editing. *Adv. Sci.*, 9, e2202957.

- • • • • (2020). PhieCBEs: Plant high-efficient C-tidine base editors with expanded target range. *Mol. Plant*, 13, 1666–1669.
- • • • • (2021). Genome editing for plant research and crop improvement. *J. Integr. Plant Biol.*, 3, 3–33.
- • • • • (2021). Expanding base editing scope to near-PAMless with engineered CRISPR/Cas9 variants in plants. *Mol. Plant*, 14, 191–194.
- • • • • (2019). Generation of herbicide tolerance traits and a new selectable marker in wheat using base editing. *Nat. Plants*, 5, 480–485.
- • • • • (2019). CRISPR enables directed evolution in plants. *Genome Biol.*, 20, 83.
- • • • • (2021). G1 cos lase base editors enable C-to-A and C-to-G base changes. *Nat. Biotechnol.*, 39, 35–40.