# **Targeted DNA insertion in plants**

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Conventional methods of DNA sequence insertion into plants, using *Agrobacterium*-mediated transformation or microprojectile bombardment, result in the integration of the DNA at random sites in the genome. These plants may exhibit altered agronomic traits as a consequence of disruption or silencing of genes that serve a critical function. Also, genes of interest inserted at random sites are often not expressed at the desired level. For these reasons, targeted DNA insertion at suitable genomic sites in plants is a desirable alternative. In this paper we review approaches of targeted DNA insertion in plant genomes, discuss current technical challenges, and describe promising applications of targeted DNA insertion for crop genetic improvement.

plant genetic engineering | plant genome editing | plant genetics | targeted gene insertion | gene stacking

Providing the world's people with sufficient food and fiber while minimizing the environmental footprint of agriculture is one of the greatest challenges of our time. Genetic improvement of crop plants is an important component of enhancing the sustainability of global agricultural systems (1).

One strategy of introducing agronomically important traits into plants is through genetic engineering (Box 1), which directly manipulates the genetic makeup of the plants using molecular genetic tools (2). Compared with crop improvement methods that rely on cross-pollination, genetic engineering introduces genes encoding desirable traits directly into the plant genome. In addition, genetic engineering can be used to introduce genes from any species, expanding the diversity of agronomically useful traits that can be accessed (3).

Genetic engineering of plants often requires plant transformation (Box 1), which has been established for a wide range of plant species (4). In conventional plant transformation protocols, DNA is delivered into plant cells either via the plant-infecting soil bacterium *Agrobacterium tumefaciens* or by microprojectiles propelled by a particle gun (5). These methods result in varying copies of the DNA inserted at random locations in the host genome, which may negatively alter the plant phenotypes (6, 7). To obtain a genetically engineered plant variety with optimal phenotypes, hundreds of independent transformation events (Box 1) are typically generated (8, 9). These plants are screened for individuals carrying a single-copy insertion with high field performance (8). This pipeline can be labor-intensive and timeconsuming, especially for plant species with long generation times (10).

In contrast to these conventional approaches, insertion of DNA at precharacterized genomic targets increases the chance of creating the desired traits in the resulting plants (11). During the past 30 y, various methods for targeted DNA insertion in plants have been established. Many of these methods have been improved for higher efficiencies and a broader range of genomic sites that can be targeted. In this paper, we review examples of targeted gene insertion in plants. We also discuss the technical challenges and propose strategies to address these hurdles. In the last section, we highlight the potential application of these methods in the context of agricultural production.

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### Methods of Targeted DNA Insertion in Plants

In this section, we provide a historical overview of methods of targeted DNA insertion in plants and provide a few examples for each method. For a more comprehensive list of the published reports of targeted DNA insertion in plants we refer readers to *SI Appendix*, Table S1.

Unaided Homologous Recombination. Pioneering research in the 1980s in mammalian cells demonstrated that exogenous DNA can be targeted to specific sites within the host genome at a low frequency through homologous recombination (HR) (12-15). These discoveries inspired the first demonstration of targeted DNA insertion in the model plant tobacco by Paszkowski et al., in 1988 (16). The researchers isolated protoplasts (Box 1) from tobacco lines carrying a partially deleted selectable marker gene and electroporated these protoplasts with DNA encoding the marker gene carrying a different deletion (16). Because the two deletions were nonoverlapping, HR between the genomic DNA and the donor DNA restored the function of the marker gene (16). The estimated efficiency of HR in this study was 0.5 to  $4.2 \times$ (16), which was comparable to the HR frequency reported  $10^{-}$ in mammalian cells around the same time (15, 17). Similar studies in tobacco and Arabidopsis documented HR-based targeted DNA insertion at similar efficiencies (18-21).

Researchers reported that HR-based targeted DNA insertion strategies often result in inauthentic HR events (21-24). To enrich the true HR-mediated targeted insertion events, two types of positive-negative selection systems were established. In the first system, Risseeuw et al. generated recipient tobacco plants carrying the negative selectable marker gene codA, which encodes a cytosine deaminase and confers lethality in the presence of the chemical compound 5- fluorocytosine (25). The researchers electroporated protoplasts derived from these recipient plants with a plasmid to induce HR in the desired manner, which would insert a kanamycin resistance marker gene and simultaneously disrupt codA (25). By applying kanamycin and 5-fluorocytosine at the same time, the efficiency of targeted insertion was increased to  $5.7 \times 10^{-3}$  (25). However, because this positive-negative selection system relied on the presence of the *codA* gene at the target site, it is not applicable to other genomic

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## Box 1.

### Definitions of some key terms in plant genetic engineering

**Genetic Engineering** Genetic engineering of plants refers to the process of directly manipulating DNA with molecular genetic tools to make changes to the plant genome, often for improved agronomic traits. Examples of genetic engineering in plants include inserting DNA from a different species into the target plant genome and modifying endogenous plant DNA through genome editing.

**Plant Transformation** Plant transformation refers to the method of introducing DNA into the plant genome. The two conventional ways of transforming plants rely on the use of either the bacterial plant pathogen *Agrobacterium tumefaciens* or a particle gun to deliver the DNA to be inserted. These methods insert DNA at random sites in the plant genome.

**Selectable Marker Genes** In plant transformation, selectable marker genes are used to distinguish the cells that have uptaken the delivered DNA from the prevalent untransformed cells. Usually, the expression of a selectable marker gene confers the ability to survive a specific selection pressure. An herbicide or an antibiotic is often used during the selection process to eliminate the untransformed cells. Sometimes, a negative selectable marker gene, which confers lethality, is used to eliminate transformed cells carrying DNA inserted at unintended targets.

**Plant Regeneration** Plant regeneration is the process of inducing the growth and differentiation of multipotent plant cells into whole plants. Successful genetic engineering of plants often requires the regeneration of whole plants from transformed plant cells. Plant regeneration usually involves culturing plant tissue in the presence of specific plant hormones under sterile conditions.

**Transformation Events** Each plant regenerated from an independently transformed plant cell is considered a single transformation event. Independent transformation events vary in the location and the copy number of the DNA insert.

**Protoplasts** Protoplasts are plant cells with the cell wall removed. They can be transformed at high efficiency through methods of direct DNA transfer. However, regenerating plants from transformed protoplasts is often challenging.

targets. In a second system developed by Terada et al., in 2002, a dual selection plasmid carries a positive selectable marker gene (Box 1) flanked by sequence homologous to the genomic target and a diphtheria toxin gene outside the homologous region as a negative marker conferring lethality (26). The desired HR would result in the targeted insertion of only the positive selectable marker gene, whereas random integration of the delivered DNA into the host genome would result in the insertion of both the positive and the negative selectable markers (26). This approach was used to generate insertional mutants at two loci in rice at 1 to 2% efficiencies (26, 27).

Overexpressing genes known to enhance HR can also promote high-efficiency targeted DNA insertion in plants. For example, in 2005, Shaked et al. reported that constitutive expression of the yeast HR-promoting chromatin remodeling gene *Radiation sensitive 54* (*ScRAD54*) in *Arabidopsis* increased the efficiency of targeted DNA insertion by one to two orders of magnitude without altering the phenotypes (28). Notably, the insert contained a promoterless green fluorescent protein (GFP) gene without any selectable marker gene, which suggests that any nucleotide sequence can be used as the insert in this method (28).

Because of the low frequency of intrinsic HR, most of the early methods of targeted DNA insertion that relied solely on this intrinsic process were inefficient and cannot be applied broadly in plants. To overcome the inefficiency, other methods have subsequently been developed, which are described in the following sections.

**Recombinase-Based Methods.** Recombinases recognize specific nucleotide sequences known as recombination sites and activate the swapping of DNA (29). The relative positioning of the two recombination sites determines the outcome of the recombination (*SI Appendix*, Fig. S14). When matching recombination sites are present on both the genomic target and the donor DNA carrying the nucleotide sequence to be inserted, the corresponding recombinase can catalyze the targeted insertion of the donor DNA at the genomic target (*SI Appendix*, Fig. S1 *B* and *C*). In the early 1990s, various recombinase systems were

developed for site-specific gene insertion (30), such as the bacteriophage Cre-*Lox* system (31), the yeast flipase–flipase recognition target (FLP-*FRT*) system (32), and the yeast recombinase-recombination site (R-*RS*) system (33). They were exploited by plant scientists to integrate DNA fragments at defined genomic targets bearing appropriate recombination sites.

Because these recombination reactions are reversible in the presence of the recombinase, inserted DNA can be excised from the genomic target (34) (SI Appendix, Fig. S1A). This creates a challenge for the use of recombinase systems for stable on-target integration of DNA. In 1995, Albert et al. modified the Cre-Lox system to effectively prevent the reversal of the recombinasecatalyzed DNA insertion in tobacco plants (35). First, nucleotide sequences of Lox sites were altered so that the recombination reaction strongly favors one direction (35). Second, they devised strategies in which the amount of the Cre recombinase is reduced after the intended DNA insertion (35). With these improvements, a high proportion of regenerated tobacco plants carried a single-copy insertion at the designated genomic target (35). Additional examples of targeted gene insertion leading to the restoration of a marker gene using recombinase systems have been reported in tobacco (6, 36), Arabidopsis (37-39), soybean (40), rice (41, 42), and maize (43). These studies demonstrated that recombinase systems can be used to induce targeted gene insertion in diverse plant species.

Additional improvements of recombinase-based targeted gene insertion in plants have been made. Positive-negative selection systems have been employed in recombinase-based gene insertion methods to enrich on-target insertion events. For example, the cytokinin biosynthesis gene *isopentyl transferase (ipt)* or the cytosine deaminase gene *codA* have been utilized as negative selectable marker genes because they encode proteins that kill the plant tissue under the proper selective conditions (36, 38). In these experimental setups, undesired insertion of the donor plasmid at random genomic sites would introduce the negative marker gene into the plant genome and abolish the host cell (36, 38). In addition to adopting a positive-negative selection system, Nanto et al. in 2005 also placed special recombination sites on the donor DNA which promote the excision and removal of

randomly integrated donor but not the on-target DNA insert (36). With these improvements, the overall targeting efficiency reached 3% (36). Besides, improvements in plant transformation techniques have also increased the overall efficiency of recombinase-based targeted gene insertion. For example, Anand et al. in 2019 applied an optimized FLP-*FRT* system and the *Wus2-Bbm* maize transformation method (44), achieving targeted DNA insertion at a 7% efficiency in maize (43).

Recombinase-based gene insertion methods can be efficient, but they invariably depend on the availability of recipient lines carrying preintegrated recombination sites. Available genomic targets for DNA insertion are therefore limited. This limitation may be overcome by using more recent gene insertion tools (discussed below) to place recombination sites at a wider range of genomic sites as landing pads for additional targeted DNA insertion using recombinase-based methods (45).

DNA Repair-Based Methods. Genome modification at a given target can be introduced at relatively high frequencies during the repair of DNA double-stranded breaks (DSBs) (46). Cellular mechanisms to repair DSBs can be roughly classified into end joining (EJ) and HR (47, 48). The EJ pathway is further divided into nonhomologous end joining (NHEJ) and microhomologymediated end joining (MMEJ, also known as alternative EJ) (47). During NHEJ, the two broken ends are directly rejoined, sometimes accompanied by sequence modifications at the junction ends, mostly in the form of small insertions or deletions (49). Unlike NHEJ, MMEJ involves the alignment of microhomology sequences (usually less than 16 nucleotides [nt] in length) present on both DNA ends before joining and usually results in the deletion of the nucleotide sequence between the two microhomologies (50). By contrast, the HR repair pathway is generally considered error-free and requires longer homology (usually >20 nt) on both DNA ends (47). HR is rare compared with EJ, especially in somatic plant cells (51, 52). NHEJ is widely considered as the most prevalent repair pathway in plants. NHEJ, MMEJ, and HR have all been successfully exploited for routine targeted DNA insertion (SI Appendix, Fig. S2) in mammalian cells (53, 54). In plants, most reported examples of targeted gene insertion through DNA repair have relied on HR. There have been only a few examples of targeted gene insertion in plants through NHEJ. We will discuss examples associated with both repair pathways in this paper. Notably, a recent study in rice suggests that the efficiencies of sequence deletion resulting from MMEJ and NHEJ are comparable, with the efficiency of MMEJ largely influenced by the availability of microhomologies near the DSB (55). This indicates that MMEJ has the potential to be exploited for targeted gene insertion in plants in the future.

Generating DSBs at defined genomic targets is crucial to efficient DNA insertion at these sites. Site-directed nucleases (SDNs) are enzymes capable of inducing DSBs at genomic targets with specific nucleotide sequences (51, 56). In this section, we highlight examples of targeted DNA insertion in plants achieved using four major SDN platforms: meganucleases, zinc finger nucleases (ZFNs), transcriptional activator-like effector nucleases (TALENs), and CRISPR-Cas. We do not include examples of SDN-mediated sequence replacement in our discussion, for which the readers are directed to two comprehensive reviews (51, 57).

*Meganucleases.* Meganucleases (also known as homing endonucleases) are naturally found in a large number of prokaryotic and eukaryotic organisms. They selectively cleave DNA at genomic targets with specific nucleotide sequences of 14 base pairs (bp) to 40 bp (58). The relatively high targeting specificity made them promising candidates as genome engineering tools in the 1990s. In 1993 and 1996, Puchta et al. demonstrated that DSBs generated by the yeast meganuclease *I-SceI* can increase the frequency of HR at a specific locus by over two orders of

magnitude in tobacco cells (59, 60). In the 1996 study, the researchers established an assay system in tobacco where the targeted DNA insertion by HR would activate a selectable marker gene (60). Codelivery of the donor DNA with a plasmid encoding *I-SceI*, which recognized the insertion target, boosted the HR efficiency from  $10^{-5}$  to over  $10^{-3}$  (60). In a subsequent study of the resulting repair junctions, *Agrobacterium*-delivered DNA encoding *I-SceI* was found to occasionally integrate at the genomic cleavage target of *I-SceI* (61). This discovery led to the application of meganucleases to insert *Agrobacterium*-delivered DNA at designated targets in the plant genome (62, 63). The use of meganucleases for targeted DNA insertion has also been demonstrated in maize (64) and barley (65) through resorting the function of a selectable marker gene.

The reliance on an existing site in the genome that matches the sequence specificity of the nuclease puts constraints on the application of meganucleases for targeted gene insertion in plants. Due to this limitation, meganucleases have largely been replaced by newer molecular tools (discussed below).

**ZFNs**. ZFNs are chimeric nucleases created by the fusion of a DNA-binding domain and a nonspecific DNA cleavage domain, typically derived from the endonuclease FokI (66). The DNA-binding domain consists of multiple zinc finger repeats, each recognizing a distinct nucleotide triplet. By combining various zinc finger repeats, the DNA-binding domain can be programmed to recognize a specific nucleotide sequence of 9 to 18 bases (67). Because the FokI domain can only cut DNA when dimerized, a pair of ZFNs that recognize sites in close proximity are used to cut DNA at the intended genomic target (68). Compared with meganucleases, ZFNs are more flexible SDNs because they can be programmed to target any genomic location.

The use of ZFNs in targeted DNA insertion in plants was first demonstrated in 2005 by Wright et al. in tobacco (69). The researchers codelivered DNA encoding a ZFN and a donor repair template into tobacco cells to repair a defective reporter gene that had previously been integrated into the tobacco genome (69). Cleavage at the defective reporter gene by the ZFN enhanced HR between the target and the repair template, resulting in the insertion of a 600-bp DNA fragment, restoring the function of the reporter gene (69). This proof-of-concept study showed that ZFNs can be used in plants to induce HR and targeted DNA insertion. Shukla et al. in 2009 used ZFNs to insert an herbicide tolerance gene within the maize metabolic gene inositol-1,3,4,5,6-pentakisphosphate 2-kinase (IPK1), disrupting its function (70). This targeted insertion conferred herbicide tolerance and reduced the accumulation of phytate, a component in seeds that can promote mineral deficiencies in humans by impairing the absorption of iron, zinc, and calcium (70).

Stacking multiple traits at a single locus is often desirable because it greatly reduces the breeding efforts. To this end, Ainley et al. in 2013 demonstrated the iterative use of ZFNmediated targeted gene insertion through HR in maize to stack multiple marker genes at a single site (71). Building on the same system, Kumar et al. in 2015 developed a system that simultaneously exchanges selectable markers and integrates new trait genes in maize, enabling nonmarker trait genes to be stacked (72). A similar NHEJ-based gene stacking strategy by the iterative use of ZFN-mediated targeted gene insertion has also been proposed in tobacco and *Arabidopsis* (73). In another effort of stacking genes at a designated genomic target in plants, Bonawitz et al. in 2019 demonstrated the targeted insertion of a 16.2-kb DNA fragment carrying four transgenes into the soybean genome using ZFNs (74).

ZFNs are highly programmable: The amino acids of the zinc finger domain can be adjusted for a variety of genomic targets. Therefore, the application of ZFNs does not rely on the creation of recipient plant lines carrying preintroduced target sites.

Specific zinc finger repeats have been developed for most nucleotide triplets (75). However, the modular combination of these repeats for an effective, sequence-specific ZFN requires laborious screening and optimization (75).

TALENS. Like ZFNs, TALENs are chimeric DNA-cutting enzymes resulting from the fusion of a highly modular DNA binding domain and the FokI nuclease domain (76, 77). The DNA binding domain of a TALEN originates from the transcription activatorlike (TAL) effectors from the bacterial plant pathogen Xanthomonas and consists of up to about 30 near-identical repeats (78, 79). Within each repeat, two variable amino acids dictate the recognition of a particular base on a DNA sequence (80). By joining multiple repeats, a DNA-binding domain that recognizes a specific stretch of DNA is generated (80). A pair of TALENs recognizing genomic targets in proximity can lead to the dimerization of the FokI nuclease domain and induce a DSB (76, 77). TALENs can be engineered to target virtually any given DNA sequence in a relatively simple manner (77), which gives this technology additional flexibility compared with ZFNs. Notably, TALENs is the first genome editing technique used to facilitate immunotherapies by deactivating immune genes which would otherwise cause the infused immune cells to attack the patient (81). Two infants with leukemia have been successfully treated with this therapeutic approach (82), demonstrating the preciseness of TALENs in genome editing.

In 2013, Voytas and coworkers reported the in-frame insertion of a yellow fluorescent protein (YFP) reporter gene into an endogenous gene in tobacco protoplasts by codelivering a plasmid encoding TALENs and a repair template plasmid (83). While no fluorescence was observed when the donor plasmid was delivered alone, about 14% of the cells treated with both plasmids produced the fluorescence, indicating high-efficiency targeted gene insertion through HR (83). The same research group also used TALENs in tomato to insert a constitutive 35S promoter upstream of the anthocyanin synthesis gene ANT1, which led to the accumulation of the pigment anthocyanin in regenerated tomato plants (84). In this study, DNA sequences encoding the TALENs and the repair template were placed on a viral replicon, which increases in copy number when delivered (84, 85). TALENs were also used to demonstrate targeted gene insertion in potato using the viral replicon-based delivery method, restoring the activity of a selectable marker gene or a reporter gene (86). Despite the requirement of protein engineering for every distinct target, the TALENs technology is still valued for plant genome engineering because of its programmability, efficiency, and target specificity (87).

CRISPR-Cas. The CRISPR-Cas platform originates from a prokaryotic adaptive immune system, which provides protection from invading viruses by cutting the viral nucleic acid (88). First established in 2012 as a molecular tool to cut DNA with specific nucleotide sequences (89), the CRISPR-Cas system typically consists of a Cas nuclease and a guide RNA molecule, which directs the Cas to generate DSBs at genomic targets with a defined nucleotide sequence (90). Since the report of the Cas nuclease prototype Streptococcus pyogenes Cas9 (SpCas9) (91), numerous naturally occurring or engineered Cas nucleases with various features have been discovered (92, 93). Target recognition by a Cas nuclease is governed by Watson-Crick base pairing between the programmable section of a guide RNA and the genomic target (89). The recognition specificity can be easily changed by modifying the variable region of the guide RNA, which makes CRISPR-Cas a highly programmable tool. The technology has been adopted in a wide range of applications (94, 95), including targeted gene insertion in many plant species (96).

In 2012, Puchta and colleagues described a strategy for targeted gene insertion in *Arabidopsis* known as *in planta* gene targeting (IPGT) (97). During IPGT, a transgenically expressed SDN simultaneously cuts the intended insertion target in the host genome and a chromosomal transgenic donor, releasing the donor DNA and causes its insertion at the intended genomic target (97). In 2014, the same group demonstrated the use of CRISPR-Cas for IPGT in Arabidopsis by inserting a selectable marker gene at an endogenous locus (98). In this study, the researchers first delivered the CRISPR-Cas gene and the donor DNA to plants by Agrobacterium as a transgenic T-DNA locus to initiate IPGT (98). After identifying the plants carrying the intended targeted insertion, the original T-DNA was removed from the genome through genetic segregation (98). With a slightly different delivery strategy, Zhao et al. in 2016 cotransformed Arabidopsis plants with two separate Agrobacterium strains, which carried the CRISPR-Cas machinery and the donor respectively, and induced IPGT causing the insertion of a GFP reporter gene at an endogenous locus (99). Although the reported insertion efficiency was low (<1%), the inserted DNA at the intended target did not contain any selectable marker gene, making the strategy applicable to the targeted insertion of virtually any DNA sequence (99). To increase the efficiency of targeted insertion of marker-free DNA, Miki et al. in 2018 reported the use of a sequential transformation approach in Arabidopsis to induce IPGT (100). In the first round of transformation, a plant line stably expressing Cas9 was generated as the parental line (100). In the second round of transformation, T-DNA carrying the guide RNA and a GFP donor was delivered to the Cas9-expressing parental line by Agrobacterium to induce IPGT (100). With this method, the researchers increased the insertion frequency to between 6 and 9% without the use of chemical selection (100). The T-DNA inserts can later be removed from the plants carrying the desired DNA insert through genetic segregation.

The amount of the delivered donor DNA affects the efficiency of targeted insertion. Using an *Agrobacterium*-delivered viral replicon system (85) to enrich the donor DNA can increase the efficiency of CRISPR-Cas-mediated targeted gene insertion, as demonstrated in tomato (84, 101), potato (86), wheat (102, 103), and rice (104). Alternatively, particle bombardment, which delivers higher copies of DNA molecules to plant cells than *Agrobacterium* does, has been employed to codeliver the CRISPR-Cas machinery and the repair template into plant tissues. Targeted insertions of selectable marker genes by particle bombardment have been achieved in maize (105), soybean (106), and rice (107).

Targeted insertion of marker-free DNA has been reported in rice (108-111) and maize (112) using particle bombardmentbased delivery methods. Li et al. in 2016 took advantage of the relatively more efficient NHEJ repair pathway to insert a DNA fragment into the intron of the endogenous rice gene EPSPS, which encodes EPSP synthase, the target of the common herbicide glyphosate (108). The inserted DNA altered the amino acid sequence of the gene product, which led to herbicide tolerance in rice (108). The mutations at the imperfect junction ends were embedded within untranslated regions of the intron and thus would not affect the protein-coding sequence (108). In another example, our team exploited the NHEJ repair mechanism to insert a DNA fragment encoding two genes involved in carotenoid biosynthesis at specific rice genomic targets, which were preevaluated for their ability to accommodate large-event mutations without altering plant performance (109). We demonstrated the targeted insertion of this carotenoid biosynthesis cassette at two independent genomic targets and obtained biofortified rice with no observable yield penalty (109). Shi et al. in 2017 applied CRISPR-Cas to insert an active promoter in front of an endogenous maize gene by HR to increase its expression (112). Increased expression of the target gene ARGOS8 led to significantly improved grain yield under drought stress environments in the field (112). Recently, Lu et al. in 2020 demonstrated that the efficiency of NHEJ-mediated gene insertion in rice can

be enhanced by an order of magnitude when the linear donor DNA carries two specific chemical modifications at its ends, including phosphorylation and phosphorothioate linkages (111). The researchers also devised an homology-directed repair (HDR)-based sequence replacement strategy building on the method of high-frequency gene insertion (111). Notably, the methods reported in these studies do not rely on chemical selection of the DNA insert. In principle, they can be used for the site-specific insertion of any DNA sequence.

### **Challenges and Opportunities**

**Increasing On-Target Insertion Frequency.** Currently, targeted DNA insertion in plants is usually inefficient, especially for large DNA fragments. Increasing the efficiency of targeted gene insertion will not only reduce the labor in screening plants but also allow the insertion of marker-free DNA fragments. To meet these goals, the frequency of targeted insertion needs to reach a practical level that allows the identification of plants with the desired insertion among a manageable population size.

Increasing the amount of donor DNA delivered can potentially promote the insertion of the donor DNA at the genomic target. Particle bombardment often delivers more DNA compared with Agrobacterium-mediated transformation but tends to induce more unintended sequence disruptions in the host genome (113). A large amount of donor DNA may be delivered into protoplasts with direct gene transfer methods, but regenerating plants from protoplasts is extremely challenging and remains a bottleneck for most plant species (114). Improvement in plant transformation and plant regeneration methods (Box 1) may overcome the limitations of these methods in the future. Besides, phosphorothioate linkages at the ends of a donor DNA fragment has been shown to increase the stability of the donor, which in turn contributes to increased insertion frequency (111). Also, putting the donor DNA in viral replicons (85) to increase its cellular copy number has been demonstrated as a promising strategy to achieve targeted gene insertion in a variety of plant species (84, 86, 101, 102, 104).

Biochemically tethering the donor DNA fragment to the SDN machinery leads to local donor enrichment at the insertion target, which contributes to increased insertion efficiencies in human cell lines (115, 116). Similar strategies have been used in rice for in-frame insertion of DNA encoding an HA epitope tag to label endogenous rice proteins (110, 117). In these studies, the DNA repair template was either fused with the guide RNA (117) or carried a short T-DNA border sequence, which is attracted to an *Agrobacterium* VirD2 relaxase fused to the Cas9 nuclease (110). These biochemical tethering methods have not yet been tested in other plant species.

The frequency of targeted insertion may also be increased by manipulating DNA repair pathways. DSB repair by NHEJ typically leads to the rejoining of the broken ends of the genomic target without incorporating the donor DNA. Consistently, suppressing the NHEJ repair pathway has been demonstrated to enhance DNA repair through HR in mammalian cells (118-120). Similarly, Qi et al. in 2013 reported that knocking out key NHEJ components such as KU70 or LIG4 in Arabidopsis enhanced the efficiency of HR-mediated targeted DNA insertion by three to sixteen folds (121). Similarly, a loss-of-function mutation in LIG4 in rice was reported to shift the repair pathway from NHEJ toward MMEJ, which can potentially be employed to improve the frequency of targeted gene insertion (122). Increased HR efficiency in Arabidopsis has also been achieved by knocking out the nucleosome assembly gene chromatin assembly factor 1 (CAF-1) (123), or by knocking out the DNA repair-related gene Rad50, whose homolog in yeast is involved in the cellular response to DSBs (124). HR in Arabidopsis can also be promoted by overexpressing the HR repair component hypersensitive to MMS, irradiation and MMC (MIM) (125) or the yeast chromatin

remodeling gene RAD54 (28). These examples demonstrate that repair pathways that favor DNA insertion can be promoted in plants by manipulating genes involved in DNA repair. These manipulations may help to achieve targeted gene insertion at higher efficiencies.

In recent years, the RNA-guided endonuclease Lachnospiraceae bacterium Cas12a (LbCas12a, also known as LbCpf1) has emerged as a promising tool for targeted gene insertion or sequence replacement in plants (107, 126–131). LbCas12a recognizes a T-rich protospacer-adjacent motif (PAM), which allows the nuclease to access AT-rich genomic regions where PAMs for various Cas9 nucleases are underrepresented (132). Furthermore, unlike for SpCas9, the position of the DSB induced by LbCas12a is located outside the critical region of the genomic target recognized by the guide RNA (132). Because of this feature, gene editing by LbCas12a has been hypothesized to favor large deletions, gene insertions, or gene replacements, because these changes would significantly disrupt the target sequence and prevent further cleavage by LbCas12a (132). Consistent with this hypothesis, Vu et al. in 2020 observed that the use of LbCas12a resulted in a higher frequency of targeted gene insertion compared with SpCas9 in one gene insertion experiment in tomato (126). Similarly, Wolter and Puchta in 2019 observed in an experiment in Arabidopsis that LbCas12a resulted in a higher IPTG efficiency compared with Staphylococcus aureus Cas9 (SaCas9) (129). To further test this hypothesis in plants, it is worthwhile comparing LbCas12a and Cas9 in targeted gene insertion experiments involving additional genomic targets in a wider range of plant species.

Studies in mammalian cells (133, 134) and plants (135, 136) have demonstrated that the nucleotide sequence context surrounding the DSB influences the DNA repair pathway utilized. Therefore, the choice of genomic targets for DNA insertion is crucial for high-efficiency targeted insertion in plants. However, there have been few high-throughput analyses of the DSB repair outcomes at diverse genomic sites in plants (137). Additional studies in the future may reveal potential features associated with genomic contexts that favor donor insertion during DSB repair in various plant species. This knowledge would provide guidance to target selection for increased on-target gene insertion frequencies.

Reducing Off-Target Insertions. While targeted gene insertion can occur at a reasonable efficiency in plants, off-target (ectopic) insertion of the donor DNA often arises (23, 24, 97, 109, 138). Sometimes, ectopic insertion events are mistakenly recognized as carrying the on-target DNA insert. For example, the ectopic insertion of a promoterless marker gene may unintendedly lead to its activation (65). Ectopic insertions can also generate falsepositive PCR genotyping results arising from a PCR artifact known as template switching (139), giving the false impression of an on-target insertion (140). Therefore, although selection and PCR are efficient ways of identifying primary transformants, results from these assays should be verified using other methods. Southern blotting, whole-genome sequencing, or Southern-bysequencing (141) are useful for validating the putative insert and assessing the presence of ectopic inserts. Various dualselection systems have also proved effective in eliminating plants carrying ectopic insertions (26, 36). When both ectopic and ontarget inserts exist, genetic segregation can often be used to remove the ectopic inserts.

Off-target insertions can be reduced by using SDNs with increased target specificity. For CRISPR-Cas, this can be achieved by designing guide RNAs with reduced off-target effects (142, 143) and by using Cas nucleases with enhanced specificity (144–146). Besides, using egg cell-specific or early embryospecific promoters to drive the expression of CRISPR-Cas has

been shown to reduce excessive nuclease activity in *Arabidopsis* (100, 147, 148).

In 2019, Liu and coworkers established a novel search-andreplace genome editing platform known as prime editing (149). In prime editing, a Cas9 nickase introduces a single-stranded break at a designated genomic target. Subsequently, a reverse transcriptase tethered to the Cas9 nickase extends the 3' end of the nicked strand of the DNA using a programmable prime editing guide RNA as the template for reverse transcription (149). Using prime editing, the researchers introduced a wide variety of short sequence edits in human cells, including an insertion of up to 3 bp (149). Because no DSB at the editing target is incurred during prime editing, fewer off-target mutations are introduced compared with methods that involve DSB generation (149). Prime editing has been applied in genome editing in wheat (150) and rice (150–156), including the insertion of nucleotide triplets in rice cells (150, 152). As a promising genome editing platform, prime editing may be optimized in the future to achieve targeted insertion of larger DNA fragments.

# Application of Targeted DNA Insertion for Crop Improvement

**Genomic Safe Harbors.** It has been known for decades that the expression of eukaryotic genes is influenced by the surrounding genomic context (6, 157). Appropriate spatial and temporal expression of the inserted genes depends largely on the insertion site, as the silencing of transgenes often occurs (158). In addition, the insertion of exogenous DNA may affect the expression of endogenous genes, especially when the insertion site is within a gene (159). Thus, transgene insertion at random sites through conventional transgenic approaches without proper testing often leads to unintended effects, such as decreased yield (160).

Alternatively, targeted insertion can be applied to insert trait genes into plant genomes at precharacterized sites known as genomic safe harbors, which are known not to interfere with plant performance (109). Defining genomic safe harbors for a crop plant is rewarding but can be laborious because in-depth analyses of plant performance would require multiseason field trials and the assessment of diverse phenotypes (34). Still, potential genomic safe harbors can be quickly identified by characterizing existing transgenic or mutant lines, identifying the insertion sites, measuring the expression of inserted genes, and evaluating plant performance using high-throughput phenomic approaches (161).

**Stacking Multiple-Trait Genes at a Single Locus.** Gene stacking in plants refers to the combination of multiple desirable trait genes, often in an elite cultivar (162). Traditionally, this is achieved through crossbreeding, where plants expressing valuable traits are cross-pollinated and individuals with combined trait genes are identified in the progeny through genetic screens. The lack of genetic linkage among individual transgenes often results in complex genetic segregation patterns. As a result, identifying plants with the desired genetic makeup often requires prolonged screening of large populations.

Alternatively, multiple genes can be positioned at a single genomic safe harbor through targeted gene insertion and crossbred into the desired cultivars as a single genetic locus. The simplified genetic segregation pattern would reduce the number of plants that need to be screened. Using targeted gene insertion for gene stacking has been demonstrated in a number of plant species (72, 163–166) and has the potential to increase the efficiency of plant breeding.

**Marker-Free Insertion.** In conventional plant transformation, DNA insertion into the plant genome is a rare event. Therefore, most transformation protocols rely on chemical selection to eliminate untransformed cells. Accordingly, DNA inserted through these

methods must contain a selectable marker gene (Box 1), which usually remains as part of the inserted DNA in the final product and often triggers additional governmental regulation and public concern (167). In addition, from an engineering perspective, the presence of the marker in the genetic background prevents future insertion of DNA fragments containing the same marker gene. Therefore, it is desirable to generate engineered plants without selectable markers.

Because DNA insertion tends to occur at DSBs, codelivering the donor DNA with an SDN to a plant can increase the frequency of the donor DNA being inserted at the defined genomic target. Screening methods such as PCR can often be used to identify the desired insertion events among plants carrying the SDN. The process does not rely on chemical selection of the intended insert. Although a selectable marker is sometimes used to select plants carrying transgenes encoding the SDN during the delivery process, the marker gene is genetically unlinked to the target locus and can be removed from the final product through genetic segregation. This strategy of inserting marker-free DNA at designated genomic targets has been successfully applied in rice (99, 104, 108–111) and tomato (101) and may be applied to additional crop species with an established transformation protocol.

**Concluding Remarks.** There is an increasing need to engineer complex genetic traits into plants. For instance, in an effort to develop combined-trait corn varieties, four independent DNA inserts were combined through conventional breeding, resulting in corn plants that showed broad-spectrum insect resistance and tolerance to two herbicides at the same time (168). This massive multiyear breeding program was a joint effort of Monsanto Company and Dow AgroSciences, two of the largest agriculture companies in the United States at the time (169). In contrast, the ability to target multiple DNA elements to a single genomic site for trait stacking can simplify the inheritance pattern of the trait genes and thereby reduce the laborious breeding.

With over three decades of advancements in plant genome engineering, targeted insertion of large DNA fragments at defined genomic sites is no longer a dream. However, despite the remarkable progress in targeted insertion in plants, highefficiency targeted insertion of large, marker-free DNA fragments and the recovery of plants with minimal off-target effects is still technically challenging. Targeted gene insertion in plants can be optimized in the future by improving DNA delivery to the plant cells, reducing the off-target effect of SDNs, and shaping the DNA repair mechanism to favor gene insertion.

New molecular tools are being developed at an accelerating rate. Beyond the conventional plant transformation methods, new delivery technologies such as carbon nanotubes (170), viral replicons (85), and de novo meristem induction (171) help overcome the hurdle of gene delivery. In addition, editing platforms based on prime editing (149, 150) or the CRISPR-Cas ribonucleoprotein (172, 173) are promising strategies with reduced off-target effects. Technological advances such as these will contribute to improved efficiency of targeted DNA insertion in plants.

Data Availability. There are no data underlying this work.

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