A review on advanced methods in plant gene targeting

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A B S T R A C T

Plant genetic engineering is one of the most significant tools implemented in the modern molecular crop breeding techniques. The conventional approaches of plant genetic transformation include \textit{Agrobacterium tumefaciens}, particle bombardment, DNA uptake into protoplast. The transgenic events derived by these methods carry the transgenes that are integrated at random sites in the plant genome. Novel techniques that mediate integration of foreign genes at specific pre-determined locations circumvent many problems associated with the existing methods of gene transfer. The recent years have witnessed the emergence of gene targeting techniques by employing zinc finger nucleases (ZFNs), transcription activator-like effector nucleases (TALENs), and clustered regularly interspaced short palindrome repeats (CRISPR). The present review focuses on the various approaches and their performance of plant gene targeting and suggests future directions in the important areas of plant molecular biology.

1. Introduction

The genetic transformation in plants has revolutionized agriculture by facilitating the introduction of foreign genes into the agronomically and horticulturally important species. This technology leads to the expression of novel traits such as pest resistance, disease resistance, and quality improvement. The transgenic plants are generated based on the genetic transformation techniques mediated by \textit{Agrobacterium tumefaciens}, particle bombardment, and DNA uptake into protoplast. The transgene integration, mediated by these techniques takes place at random sites in the plant genome. The position of genomic integration and the complexity of the integrated DNA influence the level of transgene expression [58,39,21]. Also, the transgenes inserted at random positions may lead to redundant mutations because of its insertion in the active plant genes [37]. The development of techniques that mediate transfer and integration of the foreign genes at specific pre-determined locations obviates many complications associated with the existing gene transfer methods. The introduction of foreign genes via Gene Targeting (GT), which is based on the Homologous Recombination (HR), offers many advantages such as precision gene integration, single copy transgene insertion, and high expression of...
the transgenes. It allows the construction of ‘safer’ transgenic crops, with no unknown ‘position’ effects due to random integration.

GT is a genetic technique that uses HR to alter a specific DNA sequence in an endogenous gene at its original locus in the genome. Paszkowski et al. [56] integrated antibiotic-tolerant gene into the tobacco genome using GT originally. Various HR-dependent approaches have also successfully targeted genes in plants [33,75,80].

In this review, we systematically reviewed the performance of various methods and approaches about the introduction of plant gene targeting. The HR, site-specific recombination, Zinc Finger Nucleases (ZFNs), Transcription Activator-Like Effector Nucleases (TALENs), and clustered regularly interspaced short palindromic repeats (CRISPR) has taken into consideration.

2. Genome editing tools

2.1. Homologous recombination

HR is a DNA maintenance pathway that protects the chromosomes against damages involves both the DNA strands, such as Double Strand Breaks (DSBs) and interstrand crosslinks. HR, illegitimate recombination or Non-Homologous End Joining (NHEJ), and Single-Strand Annealing (SSA) are the three different ways of for-repair process. GT has been widely used in mice and yeast, and its efficiency in plants was not sufficient for the routine applications [60,24]. Different methods were tested for increasing the GT efficiency in plants. Moreover, DSBs are created by a rare enzyme, I-SceI, which could improve the homologous integration frequency at the target site [61]. Such strategy entails transgenic efficiency and successful deployment of GT in mammals that maximized gradually as illustrated by [36]. Voytas [80] stated that DSBs induction at specific genome locations was effective in enhancing the efficiency of GT in plants.

Genome editing, or genome editing with engineered nucleases (GEEN) is a type of genetic engineering to insert, delete or replace DNA in the genome of an organism using engineered nucleases, or “molecular scissors.” These nucleases create site-specific double-strand breaks (DSBs) at desired locations in the genome. The induced DSBs are repaired by non homologous end-joining (NHEJ) or homologous recombination (HR), resulting in targeted mutations.

There are currently 3 types of engineered nucleases being used: zinc finger nucleases (ZFNs), transcription activator-like effector-based nucleases (TALEN), and the CRISPR-Cas system. Fig. 1 illustrates the structure and mode of action of these nucleases and Table 1 shows the comparison between these technologies.

2.2. Zinc finger nucleases

ZFNs are restriction enzymes with Zinc Finger (ZF) domains that recognize a particular sequence of DNA, fused to the nuclease domain of restriction enzyme FokI [38]. Since the domain of ZF could be engineered to focus on novel sequences of DNA, ZFNs were exploited to engineer the endogenous genome loci, particularly in the eukaryotic systems, [10]. According to [45], in the case of ZFNs, one module of DNA-binding involves nearly 30 amino acids and identifies 3 nucleotides integrating module of DNA-binding. It allows the recognition of 9–18 bps of DNA sequences. ZFNs were used for creating breaks in the site-specific chromosome, particularly in the absence of pre-engineered sites for the target [3,76].

The development of ZFN-mediated GT provides Molecular Biologists with the ability to modify the plant genomes site-specific and permanent via homology-directed repair of a targeted genomic DSB. ZFNs can be used to induce DSBs in specific DNA sequences and thereby promote site-specific HR and targeted genomic manipulation. ZFNs have a DNA recognition domain that involves an array of Cy52-His2 ZF. ZFs recognize and bind to particular nucleotide triplets. Various ZFs can be combined together for generating DNA-binding arrays that would identify the expanded sequence patterns with high affinity and greater specificity [15,55], and [68]. The gene constructs were made from the custom-designed ZFNs which were designed to cut at specific DNA sequences at a preselected locus in the plant genome. This was due to the efficiency and directiveness of the ZFs for a broad range of DNA sequences. A site-specific ZF endonuclease has been successfully employed to induce site-specific mutations by Non-Homologous End Joining in Arabidopsis [46].

By convention, the targeted genome modification (TGM) was frequently performed using the synthetic domain of ZFNs that is fused to a cleavage domain of FokI [77,11]. ZFNs were used to modify endogenous genes in various organisms, cell types, and plant species including Arabidopsis [54,82], soybean [18], maize [70] and tobacco [74]. Most of the constraints on the application of ZFN encompass the limited quantity of existing target sites, more effects on context dependence between the low targeting specificity and efficiency, repeat units and the effects of frequent off-target caused partially by the non-specific binding of DNA [20].

In Drosophila, the engineered ZFNs have identified the yellow gene that was observed in the larvae in the presence of the donor DNA. It was either joined elsewhere in Drosophila genome or seen as free-floating molecules released from the chromosomes of Drosophila by FLP recombinase [3].

Bozas et al. [6] studied the genetic analysis of ZFNs-induced GT in Drosophila. Using ZFNs for cleaving the target in the
chromosome, high frequencies of GT in the germ line of Drosophila was targeted. Both local mutagenesis through NHEJ and replacement of the gene through HR are simulated by targeting the cleavage. In this review, we focused on the mechanism related to the processes of applying materials for ry or rosy locus. HR-dropped frequency was significant in homozygous flies particularly for mutations in okr (Rad54) or spnA (Rad51) genes, and two components of invasion-mediated Synthesis-Dependent Strand Annealing (SDSA) pathway. When SSA was blocked by using circular donor DNA, HR was fully removed. This further show that the majority of the NHEG products, were produced in a lig4-dependent process. When both lig-4 and spnA were mutated and provided with a circular donor, the frequency of ry mutations was still high, and no products of HR could be observed. Further, it was stated that the local mutations given in such circumstances required an alternative, like lig-4 mechanism, for independent end-joining. These outcomes indicate the types of repair operating pathways for DSB in the mentioned GT systems. It was also found that the

![Fig. 1. Schematic representation of Genome Editing Tools.](image-url)

| Activity               | ZFN          | TALEN             | CRISPR             |
|------------------------|--------------|-------------------|--------------------|
| Recognition            | Protein-DNA  | Protein-DNA       | RNA-DNA            |
| DNA targeting specificity determinant | Zinc-finger proteins | Transcription activator-like effectors | RNA |
| Nuclease                | Fok1         | Fok1              | Cas9               |
| Target sequences        | 2 - 12 nucleotide and up | 2 - 16 nucleotide and up | Cas9 |
| Construct size          | (1 kb) 2     | (3 kb) 2          | Nearly 20 nucleotide |
| Construct              | Zinc finger sequence specifically recognizing | Protein sequence specific to binding a nucleotide sequence linked to Fok1 | A 20nt crRNA fused to tracrRNA and Cas9 endonuclease |
| Cost and time involved in assembly | 3 bp sequence linked to Fok1 | Protein sequence specific to binding a nucleotide sequence linked to Fok1 | A 20nt crRNA fused to tracrRNA and Cas9 endonuclease |
| Multiplexing            | No           | No                | Capable            |
| Success Rate            | Low          | High              | High               |

Table 1
Comparison of Genome Editing Technologies.

- **ZFN**
  - Recognition: Protein-DNA
  - DNA targeting specificity determinant: Zinc-finger proteins
  - Nuclease: Fok1
  - Target sequences: 2 - 12 nucleotide and up
  - Construct size: (1 kb) 2
  - Construct: Zinc finger sequence specifically recognizing 3 bp sequence linked to Fok1
  - Cost and time involved in assembly: Very expensive and time consuming
  - Multiplexing: No
  - Success Rate: Low

- **TALEN**
  - Recognition: Protein-DNA
  - DNA targeting specificity determinant: Transcription activator-like effectors
  - Nuclease: Fok1
  - Target sequences: 2 - 16 nucleotide and up
  - Construct size: (3 kb) 2
  - Construct: Protein sequence specific to binding a nucleotide sequence linked to Fok1
  - Cost and time involved in assembly: Very expensive and time consuming
  - Multiplexing: No
  - Success Rate: High

- **CRISPR**
  - Recognition: RNA-DNA
  - DNA targeting specificity determinant: RNA
  - Nuclease: Cas9
  - Target sequences: 4.2 kb (Cas9) + 0.1 kb (RNA)
  - Construct size: 23 kb
  - Construct: A 20nt crRNA fused to tracrRNA and Cas9 endonuclease
  - Cost and time involved in assembly: Low cost and minimum time
  - Multiplexing: Capable
  - Success Rate: High
results could be biased toward gene replacement by disabling the main pathway of NHEJ and moving toward simple mutagenesis by interfering with the HR process.

2.3. Transcription activator-like effector nucleases

TALENs have been developed as an alternative to ZFNs, particularly for targeted genome modification, and have indicated high capability for precise manipulation of the genome [16]. Similar to ZFNs, TALENs comprise an engineered oriented domain of TALE (Transcription Activator-Like Effectors) DNA binding and cleavage domain of FokI. The customizable binding domain of TALE DNA consisted of certain approximately identical repeat arrays in tandem that could target any provided sequence based on simple RVD (repeat variable di-residue) code for nucleotide recognition [4,5]. In the past, the modification of TALEN-mediated genome was widely accepted in yeast [43], rat [73,72], fruit fly [4], human pluripotent and somatic cells [51,27], nematode [81], livestock [9], silkworm [47], plants [69,48,83], zebrafish [67,30,2,8,19,52], Xenopus embryo [41], and many other organisms.

TALENs are the integration of the cleavage domain of FokI and binding domains of DNA derived from the TALE proteins. TALEs are the naturally occurring proteins from the plant genus of pathogenic bacteria namely Xanthomonas. [83] pointed out the methods of plant genome targeted modification using TALENs. Further, it was stated that the methods were optimized using protoplasts of Nicotiana tabacum targeting the acetolactate synthase gene. TALENs comprise DNA-binding domains involving in a series of 33–35 amino acid repeats, with each domain recognizing a single base pair. Therefore, a minimum of only 4 types of the module in DNA-binding are required for recognizing C, T, A, and G. Also, it was noted that the single-base identification of TALE-DNA binding domain repeats provided greater flexibility than design when compared to the triplet-restricted ZFNs as mentioned by [44]. Thus, it could be stated that TALENs are physically larger when compared to ZFNs by identifying the similar number of nucleotides.

TALENs have evolved as the reagent of choice for effectively changing the eukaryotic genomes in a targeted fashion [44,73,1]. Although TALEN have been shown to perform at high efficiency in many human cell lines and animal species, but its use in plant genome modification was shown only in 3 species (rice, tobacco and Arabidopsis). Moreover, various studies have used TALENs for creating mutations, especially for NHEJ [13,4,8,42]. It has been shown that the transient assay in the protoplast is an accurate, rapid, and reliable method for assessing the nuclease activity in tobacco and Arabidopsis [82,83]. The repeat arrays in TALEN were 1st cloned into the expression vector pZH051, for assessing the nuclease activity in the protoplasts. The transient assay in the protoplast was developed in both Brachypodium and rice. Encoding constructs for TALEN were introduced into the protoplasts using Polyethylene glycol (PEG). After two days of incubation, the genomic DNA was developed for each sample, and DNA fragments entailed each target site that was amplified by the PCR. The PCR products were digested with restriction enzymes and separated by agarose gel electrophoresis. The PCR amplicons were cloned into the T-A cloning vector, and nearly 30–50 individual clones were examined for mutations by DNA sequencing. TALEN-induced mutations frequencies at each target site in endogenous protoplasts were analyzed. The positive correlations of the nuclease activities in the protoplasts and changed calli in tobacco have been also discussed as illustrated by [83].

2.4. CRISPR/Cas

Targeted Genome Engineering (TGE) was developed as an alternative to traditional transgenic and plant breeding methods for enhancing productivity and ensuring sustainable production. TALENs and ZFNs were used, particularly in genome mutation at specific loci, but such systems need 2 novel binding proteins for DNA flanking a sequence of choice with a C-terminal module for FokI nuclease. Thus, such methods have been mostly accepted by the plant research community. Further, a new method was developed to enhance the efficiency of Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR)/CRISPR-associated (Cas) type II prokaryotic system as an alternative for genome engineering [71]. Various studies mentioned that the ability for reprogramming specificity of CRISPR/Cas endonuclease using customizable small non-coding RNAs had formed the stage for editing applications of a novel genome [49,12,31,14,35,34]. The system is based on Cas9 nuclease and an engineered single guide RNA (sgRNA) that gives a targeted sequence for nucleic acid [12,34,14]. According to [83], similar to TALENs and ZFNs, CRISPR technology has become one of the new plant breeding techniques. Such techniques make it feasible to introduce modifications in the plant genome that are non-distinguishable from those introduced by the traditional breeding and physical or chemical mutagenesis [14].

In plants, CRISPR/Cas9 system was deployed using the transient expression systems that enabled rapid optimization and execution of the method. The applied transient assays in plant research are protoplast and leaf tissue transformation using the method of agro-infiltration. Both methods were used for sgRNA and Cas9. The benefit of employing the protoplast strategy is the probability of achieving high gene co-expression gene even from the isolate plasmids. The separation of protoplasts from the plant tissue needs enzymatic digestion and cell wall elimination. The procedure could be time-consuming as the protoplast cultures are prone and fragile to contamination. An alternative procedure is the agro-infiltration assay that functions on whole plants as well as it is time saving. Such system works by A. tumefaciens strains infiltration carrying a binary plasmid that is comprised of candidate genes for expressing. Gene co-expression efficiency by agro-infiltration assay was lower when compared with the protoplasts, and integrating various genes of choice in one vector is possible.

Target specificity is a significant problem for all technologies for genome editing encompassing CRISPR/Cas. Numerous studies have analyzed the specificity of CRISPR/Cas system in human cells and in vitro [10,22,29,49,57]. The major outcome is that 3’ end of the guide sequence inside the sgRNA particularly indicates the target specificity of CRISPR/Cas system. This result is consistent with the previous studies by other researchers [35,17,14]. The mismatches between the guide sequence and DNA target of the sgRNA situated inside the final 8–10 bp of 20 bp sequence in target mostly remove the recognition of the target by Cas9 whereas mismatches towards the 5’ target end are better tolerated [17,53].

3. Conclusion

In this study, we discussed various approaches and methods regarding GT. The extensive growth and development in the plant GT have been summarized. This review indicates the significance of this research in developing the plant GT methods.

The reviewed findings indicate that one of the most efficient means for improving the HR frequency is to develop a break in the chromosome at the target site. The use of SSR gives the most straightforward method for chromosome excision. The removal of the marker genes from genetically modified and commercial plants is of specific interest because it would deliver a new generation of products for transgenic plants. It was noticed that ZFNs were exploited particularly in the eukaryotic systems to engineer endogenous genome loci. The TALENs have been developed as the reagent of choice for effectively changing eukaryotes genomes.
They have a high efficiency in many human cell lines and animal species, whereas only 3 plant species showed modification due to TALENs. The CRISPR technology is also one of new plant breeding techniques.

The difficulty in employing the transformation techniques in the case of higher plants at a higher frequency in order to get GT events has not been resolved yet. Arabidopsis plants are readily transformed by dipping plants with Agrobacterium strains carrying the transgene. This simple procedure has not been applied successfully in major crop species yet. The efforts required for transforming and screening higher plants for GT events are tedious. The development of the GT technology represents a crucial step in improving our understanding of single gene functions in its genome background by gene knocking. Moreover, it has the potential to increase the public acceptance of the plant gene modification by molecular techniques.

References

[1] Baker M. Nat Methods 2012;9:23–6.
[2] Bedell VM, Wang Y, Campbell JM, Poshusta TL, Starker CG, et al. Nature 2012;491:114–8.
[3] Bibikova M, Beurnier K, Trautman JK, Carroll D. Science 2003;300:764.
[4] Bogdanove AJ, Schornack S, Lahaye T. Curr Opin Plant Biol 2010;13:394–401.
[5] Bogdanove AJ, Voytas DF. Science 2011;333:1843–6.
[6] Bozas A, Beurnier KJ, Trautman JK, Carroll D. Genetics 2009;182(3):641–51.
[7] Britt AB, May GD. Trends Plant Sci 2003;8:90–5.
[8] Cade L, Reyno D, Hwang WY, Tsai SQ, Patel S, Khayter C, et al. Nucleic Acids Res 2012;40:8001–10.
[9] Carlson DF, Tan W, Lilluc SG, Bozas A, Beumer KJ, Trautman JK. Proc Natl Acad Sci USA 2012;109:17382–7.
[10] Carroll D. Gene Ther 2006;15:1463–8.
[11] Carroll D. Genetics 2011;188:773–82.
[12] Carroll D. Nat Biotechnol 2013;31:227–9.
[13] Cermak T, Doyle EL, Christian M, Wang L, Zhang Y, Schmidt C, et al. Genetics 2010;186:757–61.
[14] Cong L, Ran FA, Cox D, Lin S, Barbeton R. Nat Biotechnol 2013:31:822–6.
[15] DeFrancesco L. Nat Biotechnol 2011;29:681–4.
[16] DeGrazia G, Jorgensen SE, Wiedenheft B, Charpentier E. Nature 2013;495:51–8.
[17] Deveraux M, Grunis D, McKeithan T. Science 2003;300:764.
[18] Dinjek M, Chylinski K, Fonfara I, Hauer M, Doudna JA, Charpentier E. Science 2012;337:816–22.
[19] Joung JK, Sander JD. Nat Rev Mol Cell Biol 2013;14:49–55.
[20] Kim S, Veena Gelin SB. Plant J 2007;51:779–91.
[21] Kim YG, Cho J, Chandrasegaran S. Proc Natl Acad Sci USA 1996;93:1156–60.
[22] Kohli A, Twyman RM, Abrances R, Wegel E, Stoger E, Christou P. Plant Mol Biol 2003;52:247–58.
[23] Lee KJ, Lund P, Lowe K, Dunsuir P. Plant Cell 1990:2:415–25.
[24] Li L, Piatek MJ, Atef A, Piatek A, Wibowo A, Fang X, et al. Nat Biotechnol 2012;30:407–16.
[25] Li T, Huang S, Zhao X, Wright DA, Carpenter S, Spalding MH, et al. Nucleic Acids Res 2011;39:6315–25.
[26] Liu J, Li C, Wu H, et al. J Genet Genomics 2012;39:209–15.
[27] Liu Q, Segal DJ, Ghara JB, Barbas CF. Proc Natl Acad Sci USA 1997;94:5525–30.
[28] Lloyd A, Plaisier CL, Carroll D, Drews GN. Proc Natl Acad Sci USA 2005;102:2332–7.
[29] Ma S, San M. Zhang, F, Wang, Y, Liu, Y, Liu, H, Xu, et al. PLoS one. 7 (2012)e 45035.
[30] Mahfouz MM, Li L, Shamimuzzaman M, Wibowo A, Fang X, Zhu JK. Proc Natl Acad Sci USA 2011;108:2623–8.
[31] Mali P, Aach J, Stranges PB, Esselt KM, Moosburner M, Kosuri S, et al. Nat Biotechnol 2013;31:833–8.
[32] Mengiste T, Revenkova E, Becichold N, Paszkowski J. EMBO J 1999;18:4505–12.
[33] Miller JC, Tan S, Qiao G, Barlow KA, Wang J, Xia DF, et al. Nat Biotechnol 2011;29:143–8.
[34] Moore FE. PLoS One 2012;7:e37877.
[35] Nekrasov V, Stasakwicz B, Weigel D, Jones JD, Kamoun S. Nat Biotechnol 2013;31:693–7.
[36] Osakabe K, Osakabe Y, Toki S. Proc Natl Acad Sci USA 2010;107:12034–9.
[37] Pabo CO, Peisach E, Grant RA. Annu Rev Biochem 2001;70:313–40.
[38] Puchaszkowski J, Baur M, Boguci A, Potrykus I, EMBO J 1988;7:4021–6.
[39] Pattanyak V, Lin S, Gulinger JP, Ma E, Doudna JA, Liu DR. Nat Biotechnol 2013;31:839–43.
[40] Peach C, Velen J. Plant Mol Biol 1991;17:49–60.
[41] Puchta H. Trends Plant Sci 1998;3:77–8.
[42] Puchta H. Plant Mol Biol 2002;48:73–182.
[43] Puchta H, Duinon B, Hohn B. Nucleic Acids Res 1993;21:5034–40.
[44] Puchta H, Duinon B, Hohn B. Proc Natl Acad Sci USA 1996;93:5055–60.
[45] Puchta H. J Exp Bot 2005;56:1–14.
[46] Reiss B, Schwertt I, Kopchen K, Wendeler E, Schell J, Puchta H. Proc Natl Acad Sci USA 2000;97:3358–63.
[47] Reiss B. Int Rev Cytol 2003;228:85–139.
[48] Rong YS, Golic KG. Genetics 2003;165:1831–42.
[49] Sander JD, Cade L, Khayter C, Reyon D, Peterson RT, Joung JK, Yeh JR. Nat Biotechnol 2011;29:697–8.
[50] Segal DJ, Beeler RR, Blancafot P, Driever B, Effertz K, Huber A, et al. Biochemistry 2003:42:2137–48.
[51] Shan Q, Wang Y, Chen K, Liang Z, Li J, Zhang Y, et al. Mol Plant 2013:6:1365–8.
[52] Shulka VK, Doyon Y, Miller JC, Shulka VK, deKelver RC, Mohle EA, et al. Nature 2009:437–41.
[53] Sorek R, Avrimec C, Wiedenheft B. Annu Rev Biochem 2013;82:237–66.
[54] Tessson L, Nat Biotechnol 2011:29:695–6.
[55] Tong C, Huang G, Ashton C, Wu H, Yang E. Genet 2012;39:275–80.
[56] Townsend JR, Wright DA, Winerfy RJ, Fu F, Maeder ML, Joung JK, et al. Nature 2009:459:442–5.
[57] Tzfuia T, White C. Trends Biotechnol 2005;23:567–9.
[58] Urvonov FD, Miller JC, Lee YL, Beausaume CR, Rock JM, Augustus S, et al. Nature 2005:435:2.
[59] Urvonov FD, Rebar EJ, Holmes MC, Zhang HS, Gregory PD. Nat Rev Genet 2010;11:636–46.
[60] Vergunst AC, Hooykaas PJ. Crit Rev Plant Sci 1999;18:1–31.
[61] Voytas DF. Annu. Rev. Plant Biol. 2013:64:327–50.
[62] Wood AJ. Science 2011:331:307.
[63] Zhang F, Maeder ML, Unger-Wallace E, Hoshaw JP, Reyon D, Christian M, et al. Proc Natl Acad Sci USA 2010;107:12028–33.
[64] Zhang Y, Zhang F, Li X, Baller JA, Qi Y, Starker CG, et al. Plant Physiol 2013:161:20–7.