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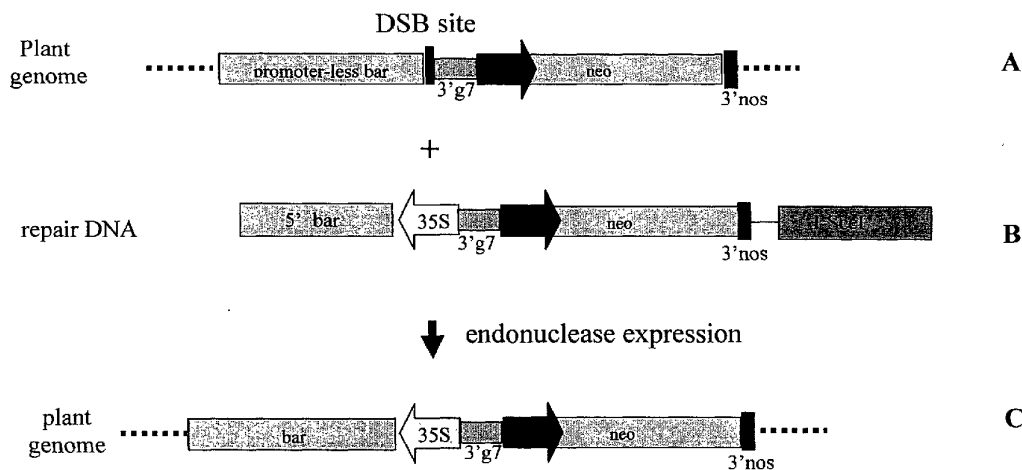
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(54) Title: IMPROVED TARGETED DNA INSERTION IN PLANTS



(57) Abstract: Methods and means are provided to improve targeted DNA insertion in plants using rare-cleaving "double stranded break" inducing enzymes. Also provided are improved I-SceI encoding nucleotide sequences.

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Improved targeted DNA insertion in plants.

Field of the invention

The current invention relates to the field of molecular plant biology, more specific to the field of plant genome engineering. Methods are provided for the directed introduction of a foreign DNA fragment at a preselected insertion site in the genome of a plant. Plants containing the foreign DNA inserted at a particular site can now be obtained at a higher frequency and with greater accuracy than is possible with the currently available targeted DNA insertion methods. Moreover, in a large proportion of the resulting plants, the foreign DNA has only been inserted at the preselected insertion site, without the foreign DNA also having been inserted randomly at other locations in the plant's genome. The methods of the invention are thus an improvement, both quantitatively and qualitatively, over the prior art methods. Also provided are chimeric genes, plasmids, vectors and other means to be used in the methods of the invention.

Background art

The first generation of transgenic plants in the early 80's of last century by *Agrobacterium* mediated transformation technology, has spurred the development of other methods to introduce a foreign DNA of interest or a transgene into the genome of a plant, such as PEG mediated DNA uptake in protoplast, microprojectile bombardment, silicon whisker mediated transformation etc.

All the plant transformation methods, however, have in common that the transgenes incorporated in the plant genome are integrated in a random fashion and in unpredictable copy number. Frequently, the transgenes can be integrated in the form of repeats, either of the whole transgene or of parts thereof. Such a complex integration pattern may influence the expression level of the transgenes, e.g. by destruction of the transcribed RNA through posttranscriptional gene silencing mechanisms or by inducing methylation of the introduced DNA, thereby downregulating the transcriptional activity on the transgene. Also, the integration site per se can influence the level of expression of the transgene. The

combination of these factors results in a wide variation in the level of expression of the transgenes or foreign DNA of interest among different transgenic plant cell and plant lines. Moreover, the integration of the foreign DNA of interest may have a disruptive effect on the region of the genome where the integration occurs, and can influence or disturb the normal function of that target region, thereby leading to, often undesirable, side-effects.

Therefore, whenever the effect of introduction of a particular foreign DNA into a plant is investigated, it is required that a large number of transgenic plant lines are generated and analysed in order to obtain significant results. Likewise, in the generation of transgenic crop plants, where a particular DNA of interest is introduced in plants to provide the transgenic plant with a desired, known phenotype, a large population of independently created transgenic plant lines or so-called events is created, to allow the selection of those plant lines with optimal expression of the transgenes, and with minimal, or no, side-effects on the overall phenotype of the transgenic plant. Particularly in this field, it would be advantageous if this trial-and-error process could be replaced by a more directed approach, in view of the burdensome regulatory requirements and high costs associated with the repeated field trials required for the elimination of the unwanted transgenic events. Furthermore, it will be clear that the possibility of targeted DNA insertion would also be beneficial in the process of so-called transgene stacking.

The need to control transgene integration in plants has been recognized early on, and several methods have been developed in an effort to meet this need (for a review see Kumar and Fladung, 2001, *Trends in Plant Science*, 6, pp155-159). These methods mostly rely on homologous recombination-based transgene integration, a strategy which has been successfully applied in prokaryotes and lower eukaryotes (see e.g. EP0317509 or the corresponding publication by Paszkowski *et al.*, 1988, *EMBO J.*, 7, pp4021-4026). However, for plants, the predominant mechanism for transgene integration is based on illegitimate recombination which involves little homology between the recombining DNA strands. A major challenge in this area is therefore the detection of the rare homologous recombination events, which are masked by the far more efficient integration of the introduced foreign DNA via illegitimate recombination.

One way of solving this problem is by selecting against the integration events that have occurred by illegitimate recombination, such as exemplified in WO94/17176.

Another way of solving the problem is by activation of the target locus and/or repair or donor DNA through the induction of double stranded DNA breaks via rare-cutting endonucleases, such as I-SceI. This technique has been shown to increase the frequency of homologous recombination by at least two orders of magnitude using *Agrobacteria* to deliver the repair DNA to the plant cells (Puchta *et al.*, 1996, *Proc. Natl. Acad. Sci. U.S.A.*, 93, pp5055-5060; Chilton and Que, *Plant Physiol.*, 2003).

WO96/14408 describes an isolated DNA encoding the enzyme I-SceI. This DNA sequence can be incorporated in cloning and expression vectors, transformed cell lines and transgenic animals. The vectors are useful in gene mapping and site-directed insertion of genes.

WO00/46386 describes methods of modifying, repairing, attenuating and inactivating a gene or other chromosomal DNA in a cell through I-SceI double strand break. Also disclosed are methods of treating or prophylaxis of a genetic disease in an individual in need thereof. Further disclosed are chimeric restriction endonucleases.

However, there still remains a need for improving the frequency of targeted insertion of a foreign DNA in the genome of a eukaryotic cell, particularly in the genome of a plant cell. These and other problems are solved as described hereinafter in the different detailed embodiments of the invention, as well as in the claims.

Summary of the invention

In one embodiment, the invention provides a method for introducing a foreign DNA of interest, which may be flanked by a DNA region having at least 80% sequence identity to a DNA region flanking a preselected site, into a preselected site, such as an I-SceI site of a genome of a plant cell, such as a maize cell comprising the steps of

- (a) inducing a double stranded DNA break at the preselected site in the genome of the cell, e.g by introducing an I-SceI encoding gene;

(b) introducing the foreign DNA of interest into the plant cell ;
characterized in that the foreign DNA is delivered by direct DNA transfer which may be accomplished by bombardment of microprojectiles coated with the foreign DNA of interest. The I-SceI encoding gene can comprise a nucleotide sequence encoding the amino acid sequence of SEQ ID No 1, wherein said nucleotide sequence has a GC content of about 50% to about 60%, provided that

- i) the nucleotide sequence does not comprise a nucleotide sequence selected from the group consisting of GATAAT, TATAAA, AATATA, AATATT, GATAAA, AATGAA, AATAAG, AATAAA, AATAAT, AACCAA, ATATAA, AATCAA, ATACTA, ATAAAA, ATGAAA, AAGCAT, ATTAAT, ATACAT, AAAATA, ATTAAT, AATTAA, AATACA and CATAAA;
- ii) the nucleotide does not comprise a nucleotide sequence selected from the group consisting of CCAAT, ATTGG, GCAAT and ATTGC;
- iii) the nucleotide sequence does not comprise a sequence selected from the group consisting of ATTTA, AAGGT, AGGTA, GGTA or GCAGG;
- iv) the nucleotide sequence does not comprise a GC stretch consisting of 7 consecutive nucleotides selected from the group of G or C;
- v) the nucleotide sequence does not comprise a AT stretch consisting of 5 consecutive nucleotides selected from the group of A or T; and
- vi) the nucleotide sequence does not comprise the codons TTA, CTA, ATA, GTA, TCG, CCG, ACG and GCG. An example of such an I-SceI encoding gene comprises the nucleotide sequence of SEQ ID 4.

The plant cell may be incubated in a plant phenolic compound prior to step a).

In another embodiment, the invention relates to a method for introducing a foreign DNA of interest into a preselected site of a genome of a plant cell comprising the steps of

- (a) inducing a double stranded DNA break at the preselected site in the genome of the cell ;
- (b) introducing the foreign DNA of interest into the plant cell ;
characterized in that the double stranded DNA break is introduced by a rare cutting endonuclease encoded by a nucleotide sequence wherein said nucleotide sequence has a GC content of about 50% to about 60%, provided that

- i) the nucleotide sequence does not comprise a nucleotide sequence selected from the group consisting of GATAAT, TATAAA, AATATA, AATATT, GATAAA, AATGAA, AATAAG, AATAAA, AATAAT, AACCAA, ATATAA, AATCAA, ATACTA, ATAAAA, ATGAAA, AAGCAT, ATTAAT, ATACAT, AAAATA, ATTAAA, AATTAA, AATACA and CATAAA;
- ii) the nucleotide does not comprise a nucleotide sequence selected from the group consisting of CCAAT, ATTGG, GCAAT and ATTGC;
- iii) the nucleotide sequence does not comprise a sequence selected from the group consisting of ATTTA, AAGGT, AGGTA, GGTA or GCAGG;
- iv) the nucleotide sequence does not comprise a GC stretch consisting of 7 consecutive nucleotides selected from the group of G or C;
- v) the nucleotide sequence does not comprise a AT stretch consisting of 5 consecutive nucleotides selected from the group of A or T; and
- vi) the nucleotide sequence does not comprise the codons TTA, CTA, ATA, GTA, TCG, CCG, ACG and GCG.

In yet another embodiment, the invention relates to a method for introducing a foreign DNA of interest into a preselected site of a genome of a plant cell comprising the steps of

- (a) inducing a double stranded DNA break at the preselected site in the genome of the cell ;
- (b) introducing the foreign DNA of interest into the plant cell ;

characterized in that prior to step a, the plant cells are incubated in a plant phenolic compound which may be selected from the group of acetosyringone (3,5-dimethoxy-4-hydroxyacetophenone), α -hydroxy-acetosyringone, sinapinic acid (3,5 dimethoxy-4-hydroxycinnamic acid), syringic acid (4-hydroxy-3,5 dimethoxybenzoic acid), ferulic acid (4-hydroxy-3-methoxycinnamic acid), catechol (1,2-dihydroxybenzene), p-hydroxybenzoic acid (4-hydroxybenzoic acid), β -resorcylic acid (2,4 dihydroxybenzoic acid), protocatechuic acid (3,4-dihydroxybenzoic acid), pyrogalllic acid (2,3,4 -trihydroxybenzoic acid), gallic acid (3,4,5-trihydroxybenzoic acid) and vanillin (3-methoxy-4-hydroxybenzaldehyde).

The invention also provides an isolated DNA fragment comprising a nucleotide sequence encoding the amino acid sequence of SEQ ID No 1, wherein the nucleotide sequence has a GC content of about 50% to about 60%, provided that

- i) the nucleotide sequence does not comprise a nucleotide sequence selected from the group consisting of GATAAT, TATAAA, AATATA, AATATT, GATAAA, AATGAA, AATAAG, AATAAA, AATAAT, AACCAA, ATATAA, AATCAA, ATACTA, ATAAAA, ATGAAA, AAGCAT, ATTAAT, ATACAT, AAAATA, ATTAAA, AATTAA, AATACA and CATAAA;
- ii) the nucleotide does not comprise a nucleotide sequence selected from the group consisting of CCAAT, ATTGG, GCAAT and ATTGC;
- iii) the nucleotide sequence does not comprise a sequence selected from the group consisting of ATTTA, AAGGT, AGGTA, GGTA or GCAGG;
- iv) the nucleotide sequence does not comprise a GC stretch consisting of 7 consecutive nucleotides selected from the group of G or C;
- v) the nucleotide sequence does not comprise a AT stretch consisting of 5 consecutive nucleotides selected from the group of A or T; and
- vi) codons of said nucleotide sequence coding for leucine (Leu), isoleucine (Ile), valine (Val), serine (Ser), proline (Pro), threonine (Thr), alanine (Ala) do not comprise TA or GC duplets in positions 2 and 3 of said codons.

The invention also provides an isolated DNA sequence comprising the nucleotide sequence of SEQ ID No 4, as well as chimeric gene comprising the isolated DNA fragment according to the invention operably linked to a plant-expressible promoter and the use of such a chimeric gene to insert a foreign DNA into an I-SceI recognition site in the genome of a plant.

In yet another embodiment of the invention, a method is provided for introducing a foreign DNA of interest into a preselected site of a genome of a plant cell comprising the steps of

- a) inducing a double stranded DNA break at the preselected site in the genome of the cell by a rare cutting endonuclease
 - b) introducing the foreign DNA of interest into the plant cell ;
- characterized in that said endonuclease comprises a nuclear localization signal.

Brief description of the figures

Table 1 represents the possible trinucleotide (codon) choices for a synthetic I-SceI coding region (see also the nucleotide sequence in SEQ ID No 2).

Table 2 represents preferred possible trinucleotide choices for a synthetic I-SceI coding region (see also the nucleotide sequence in SEQ ID No 3).

Figure 1: Schematic representation of the target locus (A) and the repair DNA (B) used in the assay for homologous recombination mediated targeted DNA insertion. The target locus after recombination is also represented (C). DSB site: double stranded DNA break site; 3'g7:transcription termination and polyadenylation signal of *A. tumefaciens* gene 7; neo: plant expressible neomycin phosphotransferase; 35S: promoter of the CaMV 35S transcript; 5' bar : DNA region encoding the amino terminal portion of the phosphinotricin acetyltransferase; 3'nos: transcription termination and polyadenylation signal of *A. tumefaciens* nopaline synthetase gene; Pnos: promoter of the nopaline synthetase gene of *A. tumefaciens*; 3'ocs: 3' transcription termination and polyadenylation signal of the octopine synthetase gene of *A. tumefaciens*.

Detailed description

The current invention is based on the following findings:

- a) Introduction into the plant cells of the foreign DNA to be inserted via direct DNA transfer, particularly microprojectile bombardment, unexpectedly increased the frequency of targeted insertion events. All of the obtained insertion events were targeted DNA insertion events, which occurred at the site of the induced double stranded DNA break. Moreover all of these targeted insertion events appeared to be exact recombination events between the provided sequence homology flanking the double stranded DNA break. Only about half of these events had an additional insertion of the foreign DNA at a site different from the site of the induced double stranded DNA break.
- b) Induction of the double stranded DNA break by transient expression of a rare-cutting double stranded break inducing endonuclease, such as I-SceI, encoded by chimeric gene comprising a synthetic coding region for a rare-cutting endonuclease such as I-SceI designed according to a preselected set of rules surprisingly increased the quality of the resulting targeted DNA insertion events (i.e. the frequency of perfectly targeted DNA insertion events). Furthermore, the endonuclease had been equipped with a nuclear localization signal.
- c) Preincubation of the target cells in a plant phenolic compound, such as acetosyringone, further increased the frequency of targeted insertion at double stranded DNA breaks induced in the genome of a plant cell.

Any of the above findings, either alone or in combination, improves the frequency with which homologous recombination based targeted insertion events can be obtained, as well as the quality of the recovered events.

Thus, in one aspect, the invention relates to a method for introducing a foreign DNA of interest into a preselected site of a genome of a plant cell comprising the steps of

- (a) inducing a double stranded DNA break at the preselected site in the genome of the cell ;
 - (b) introducing the foreign DNA of interest into the plant cell ;
- characterized in that the foreign DNA is delivered by direct DNA transfer.

As used herein "direct DNA transfer" is any method of DNA introduction into plant cells which does not involve the use of natural *Agrobacterium spp.* which is capable of

introducing DNA into plant cells. This includes methods well known in the art such as introduction of DNA by electroporation into protoplasts, introduction of DNA by electroporation into intact plant cells or partially degraded tissues or plant cells, introduction of DNA through the action of agents such as PEG and the like, into protoplasts, and particularly bombardment with DNA coated microprojectiles. Introduction of DNA by direct transfer into plant cells differs from *Agrobacterium*-mediated DNA introduction at least in that double stranded DNA enters the plant cell, in that the entering DNA is not coated with any protein, and in that the amount of DNA entering the plant cell may be considerably greater. Furthermore, DNA introduced by direct transfer methods, such as the introduced chimeric gene encoding a double stranded DNA break inducing endonuclease, may be more amenable to transcription, resulting in a better timing of the induction of the double stranded DNA break. Although not intending to limit the invention to a particular mode of action, it is thought that the efficient homology-recombination-based insertion of repair DNA or foreign DNA in the genome of a plant cell may be due to a combination of any of these parameters.

Conveniently, the double stranded DNA break may be induced at the preselected site by transient expression after introduction of a plant-expressible gene encoding a rare cleaving double stranded break inducing enzyme. As set forth elsewhere in this document, I-SceI may be used for that purpose to introduce a foreign DNA at an I-SceI recognition site. However, it will be immediately clear to the person skilled in the art that also other double stranded break inducing enzymes can be used to insert the foreign DNA at their respective recognition sites. A list of rare cleaving DSB inducing enzymes and their respective recognition sites is provided in Table I of WO 03/004659 (pages 17 to 20) (incorporated herein by reference). Furthermore, methods are available to design custom-tailored rare-cleaving endonucleases that recognize basically any target nucleotide sequence of choice. Such methods have been described e.g. in WO 03/080809, WO94/18313 or WO95/09233 and in Isalan *et al.*, 2001, *Nature Biotechnology* 19, 656- 660; Liu *et al.* 1997, *Proc. Natl. Acad. Sci. USA* 94, 5525-5530.)

Thus, as used herein "a preselected site" indicates a particular nucleotide sequence in the plant nuclear genome at which location it is desired to insert the foreign DNA. A person skilled in the art would be perfectly able to either choose a double stranded DNA break

inducing (“DSBI”) enzyme recognizing the selected target nucleotide sequence or engineer such a DSBI endonuclease. Alternatively, a DSBI endonuclease recognition site may be introduced into the plant genome using any conventional transformation method or by conventional breeding using a plant line having a DSBI endonuclease recognition site in its genome, and any desired foreign DNA may afterwards be introduced into that previously introduced preselected target site.

The double stranded DNA break may be induced conveniently by transient introduction of a plant-expressible chimeric gene comprising a plant-expressible promoter region operably linked to a DNA region encoding a double stranded break inducing enzyme. The DNA region encoding a double stranded break inducing enzyme may be a synthetic DNA region, such as but not limited to, a synthetic DNA region whereby the codons are chosen according to the design scheme as described elsewhere in this application for I-SceI encoding regions.

The double stranded break inducing enzyme may comprise, but need not comprise, a nuclear localization signal (NLS) [Raikhel, *Plant Physiol.* 100: 1627-1632 (1992) and references therein], such as the NLS of SV40 large T-antigen [Kalderon *et al.* *Cell* 39: 499-509 (1984)]. The nuclear localization signal may be located anywhere in the protein, but is conveniently located at the N-terminal end of the protein. The nuclear localization signal may replace one or more of the amino acids of the double stranded break inducing enzyme.

As used herein “foreign DNA of interest” indicates any DNA fragment which one may want to introduce at the preselected site. Although it is not strictly required, the foreign DNA of interest may be flanked by at least one nucleotide sequence region having homology to a DNA region flanking the preselected site. The foreign DNA of interest may be flanked at both sites by DNA regions having homology to both DNA regions flanking the preselected site. Thus the repair DNA molecule(s) introduced into the plant cell may comprise a foreign DNA flanked by one or two flanking sequences having homology to the DNA regions respectively upstream or downstream the preselected site. This allows to better control the insertion of the foreign DNA. Indeed, integration by homologous recombination will allow precise joining of the foreign DNA fragment to the plant nuclear genome up to the nucleotide level.

The flanking nucleotide sequences may vary in length, and should be at least about 10 nucleotides in length. However, the flanking region may be as long as is practically possible (e.g. up to about 100-150 kb such as complete bacterial artificial chromosomes (BACs)). Preferably, the flanking region will be about 50 bp to about 2000 bp. Moreover, the regions flanking the foreign DNA of interest need not be identical to the DNA regions flanking the preselected site and may have between about 80% to about 100% sequence identity, preferably about 95% to about 100% sequence identity with the DNA regions flanking the preselected site. The longer the flanking region, the less stringent the requirement for homology. Furthermore, it is preferred that the sequence identity is as high as practically possible in the vicinity of the location of exact insertion of the foreign DNA.

Moreover, the regions flanking the foreign DNA of interest need not have homology to the regions immediately flanking the preselected site, but may have homology to a DNA region of the nuclear genome further remote from that preselected site. Insertion of the foreign DNA will then result in a removal of the target DNA between the preselected insertion site and the DNA region of homology. In other words, the target DNA located between the homology regions will be substituted for the foreign DNA of interest.

For the purpose of this invention, the "sequence identity" of two related nucleotide or amino acid sequences, expressed as a percentage, refers to the number of positions in the two optimally aligned sequences which have identical residues (x100) divided by the number of positions compared. A gap, i.e. a position in an alignment where a residue is present in one sequence but not in the other, is regarded as a position with non-identical residues. The alignment of the two sequences is performed by the Needleman and Wunsch algorithm (Needleman and Wunsch 1970) Computer-assisted sequence alignment, can be conveniently performed using standard software program such as GAP which is part of the Wisconsin Package Version 10.1 (Genetics Computer Group, Madison, Wisconsin, USA) using the default scoring matrix with a gap creation penalty of 50 and a gap extension penalty of 3.

In another aspect, the invention relates to a modified I-SceI encoding DNA fragment, and the use thereof to efficiently introduce a foreign DNA of interest into a preselected site of a

genome of a plant cell, whereby the modified I-SceI encoding DNA fragment has a nucleotide sequence which has been designed to fulfill the following criteria:

- a) the nucleotide sequence encodes a functional I-SceI endonuclease, such as an I-SceI endonuclease having the amino acid sequence as provided in SEQ ID No 1.
- b) the nucleotide sequence has a GC content of about 50% to about 60%
- c) the nucleotide sequence does not comprise a nucleotide sequence selected from the group consisting of GATAAT, TATAAA, AATATA, AATATT, GATAAA, AATGAA, AATAAG, AATAAA, AATAAT, AACCAA, ATATAA, AATCAA, ATACTA, ATAAAA, ATGAAA, AAGCAT, ATTAAT, ATACAT, AAAATA, ATTAAA, AATTAA, AATACA and CATAAA;
- d) the nucleotide does not comprise a nucleotide sequence selected from the group consisting of CCAAT, ATTGG, GCAAT and ATTGC;
- e) the nucleotide sequence does not comprise a sequence selected from the group consisting of ATTTA, AAGGT, AGGTA, GGTA or GCAGG;
- f) the nucleotide sequence does not comprise a GC stretch consisting of 7 consecutive nucleotides selected from the group of G or C;
- g) the nucleotide sequence does not comprise a GC stretch consisting of 5 consecutive nucleotides selected from the group of A or T; and
- h) the nucleotide sequence does not comprise codons coding for Leu, Ile, Val, Ser, Pro, Thr, Ala that comprise TA or CG duplets in positions 2 and 3 (i.e. the nucleotide sequence does not comprise the codons TTA, CTA, ATA, GTA, TCG, CCG, ACG and GCG).

I-SceI is a site-specific endonuclease, responsible for intron mobility in mitochondria in *Saccharomyces cerevisiae*. The enzyme is encoded by the optional intron Sc LSU.1 of the 21S rRNA gene and initiates a double stranded DNA break at the intron insertion site generating a 4 bp staggered cut with 3'OH overhangs. The recognition site of I-SceI endonuclease extends over an 18 bp non-symmetrical sequence (Colleaux *et al.* 1988 *Proc. Natl. Acad. Sci. USA* 85: 6022-6026). The amino acid sequence for I-SceI and a universal code equivalent of the mitochondrial I-SceI gene have been provided by e.g. WO 96/14408.

WO 96/14408 discloses that the following variants of I-SceI protein are still functional:

- positions 1 to 10 can be deleted
- position 36: Gly (G) is tolerated
- position 40: Met (M) or Val (V) are tolerated
- position 41: Ser (S) or Asn (N) are tolerated
- position 43: Ala (A) is tolerated
- position 46: Val (V) or N (Asn) are tolerated
- position 91: Ala (A) is tolerated
- positions 123 and 156: Leu (L) is tolerated
- position 223 : Ala (A) and Ser (S) are tolerated

and synthetic nucleotide sequences encoding such variant I-SceI enzymes can also be designed and used in accordance with the current invention.

A nucleotide sequence encoding the amino acid sequence of I-SceI, wherein the amino-terminally located 4 amino acids have been replaced by a nuclear localization signal (SEQ ID 1) thus consist of 244 trinucleotides which can be represented as R1 through R244. For each of these positions between 1 and 6 possible choices of trinucleotides encoding the same amino acid are possible. Table 1 sets forth the possible choices for the trinucleotides encoding the amino acid sequence of SEQ ID 1 and provides for the structural requirements (either conditional or absolute) which allow to avoid inclusion into the synthetic DNA sequence the above mentioned "forbidden nucleotide sequences". Also provided is the nucleotide sequence of the contiguous trinucleotides in UIPAC code.

As used herein, the symbols of the UIPAC code have their usual meaning i.e. N= A or C or G or T; R= A or G; Y= C or T; B= C or G or T (not A); V= A or C or G (not T); D= A or G or T (not C); H=A or C or T (not G); K= G or T; M= A or C; S= G or C; W=A or T.

Thus in one embodiment of the invention, an isolated synthetic DNA fragment is provided which comprises a nucleotide sequence as set forth in SEQ ID No 2, wherein the codons are chosen among the choices provided in such a way as to obtain a nucleotide sequence with an overall GC content of about 50% to about 60%, preferably about 54%-55% provided that the nucleotide sequence from position 28 to position 30 is not AAG; if the nucleotide sequence from position 34 to position 36 is AAT then the nucleotide sequence from position 37 to

position 39 is not ATT or ATA; if the nucleotide sequence from position 34 to position 36 is AAC then the nucleotide sequence from position 37 to position 39 is not ATT simultaneously with the nucleotide sequence from position 40 to position 42 being AAA; if the nucleotide sequence from position 34 to position 36 is AAC then the nucleotide sequence from position 37 to position 39 is not ATA; if the nucleotide sequence from position 37 to position 39 is ATT or ATA then the nucleotide sequence from position 40 to 42 is not AAA; the nucleotide sequence from position 49 to position 51 is not CAA; the nucleotide sequence from position 52 to position 54 is not GTA; the codons from the nucleotide sequence from position 58 to position 63 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; if the nucleotide sequence from position 67 to position 69 is CCC then the nucleotide sequence from position 70 to position 72 is not AAT; if the nucleotide sequence from position 76 to position 78 is AAA then the nucleotide sequence from position 79 to position 81 is not TTG simultaneously with the nucleotide sequence from position 82 to 84 being CTN; if the nucleotide sequence from position 79 to position 81 is TTA or CTA then the nucleotide sequence from position 82 to position 84 is not TTA; the nucleotide sequence from position 88 to position 90 is not GAA; if the nucleotide sequence from position 91 to position 93 is TAT, then the nucleotide sequence from position 94 to position 96 is not AAA; if the nucleotide sequence from position from position 97 to position 99 is TCC or TCG or AGC then the nucleotide sequence from position 100 to 102 is not CCA simultaneously with the nucleotide sequence from position 103 to 105 being TTR; if the nucleotide sequence from position 100 to 102 is CAA then the nucleotide sequence from position 103 to 105 is not TTA; if the nucleotide sequence from position 109 to position 111 is GAA then the nucleotide sequence from 112 to 114 is not TTA; if the nucleotide sequence from position 115 to 117 is AAT then the nucleotide sequence from position 118 to position 120 is not ATT or ATA; if the nucleotide sequence from position 121 to 123 is GAG then the nucleotide sequence from position 124 to position 126; the nucleotide sequence from position 133 to 135 is not GCA; the nucleotide sequence from position 139 to position 141 is not ATT; if the nucleotide sequence from position 142 to position 144 is GGA then the nucleotide sequence from position 145 to position 147 is not TTA; if the nucleotide sequence from position 145 to position 147 is TTA then the nucleotide sequence from position 148 to position 150 is not ATA simultaneously with the nucleotide sequence from position 151 to 153 being TTR; if the nucleotide sequence

from position 145 to position 147 is CTA then the nucleotide sequence from position 148 to position 150 is not ATA simultaneously with the nucleotide sequence from position 151 to 153 being TTR; if the nucleotide sequence from position 148 to position 150 is ATA then the nucleotide sequence from position 151 to position 153 is not CTA or TTG; if the nucleotide sequence from position 160 to position 162 is GCA then the nucleotide sequence from position 163 to position 165 is not TAC; if the nucleotide sequence from position 163 to position 165 is TAT then the nucleotide sequence from position 166 to position 168 is not ATA simultaneously with the nucleotide sequence from position 169 to position 171 being AGR; the codons from the nucleotide sequence from position 172 to position 177 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise GCAGG; the codons from the nucleotide sequence from position 178 to position 186 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise AGGTA; if the nucleotide sequence from position 193 to position 195 is TAT, then the nucleotide sequence from position 196 to position 198 is not TGC; the nucleotide sequence from position 202 to position 204 is not CAA; the nucleotide sequence from position 217 to position 219 is not AAT; if the nucleotide sequence from position 220 to position 222 is AAA then the nucleotide sequence from position 223 to position 225 is not GCA; if the nucleotide sequence from position 223 to position 225 is GCA then the nucleotide sequence from position 226 to position 228 is not TAC; if the nucleotide sequence from position 253 to position 255 is GAC, then the nucleotide sequence from position 256 to position 258 is not CAA; if the nucleotide sequence from position 277 to position 279 is CAT, then the nucleotide sequence from position 280 to position 282 is not AAA; the codons from the nucleotide sequence from position 298 to position 303 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; if the nucleotide sequence from position 304 to position 306 is GGC then the nucleotide sequence from position 307 to position 309 is not AAT; the codons from the nucleotide sequence from position 307 to position 312 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; the codons from the nucleotide sequence from position 334 to position 342 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; if the nucleotide sequence from position 340 to position 342 is AAG then the nucleotide sequence from position 343 to

345 is not CAT; if the nucleotide position from position 346 to position 348 is CAA then the nucleotide sequence from position 349 to position 351 is not GCA; the codons from the nucleotide sequence from position 349 to position 357 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; the nucleotide sequence from position 355 to position 357 is not AAT; if the nucleotide sequence from position 358 to position 360 is AAA then the nucleotide sequence from position 361 to 363 is not TTG; if the nucleotide sequence from position 364 to position 366 is GCC then the nucleotide sequence from position 367 to position 369 is not AAT; the codons from the nucleotide sequence from position 367 to position 378 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; if the nucleotide sequence from position 382 to position 384 is AAT then the nucleotide sequence from position 385 to position 387 is not AAT; the nucleotide sequence from position 385 to position 387 is not AAT; if the nucleotide sequence from position 400 to 402 is CCC, then the nucleotide sequence from position 403 to 405 is not AAT; if the nucleotide sequence from position 403 to 405 is AAT, then the nucleotide sequence from position 406 to 408 is not AAT; the codons from the nucleotide sequence from position 406 to position 411 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; the codons from the nucleotide sequence from position 421 to position 426 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; the nucleotide sequence from position 430 to position 432 is not CCA; if the nucleotide sequence from position 436 to position 438 is TCA then the nucleotide sequence from position 439 to position 441 is not TTG; the nucleotide sequence from position 445 to position 447 is not TAT; the nucleotide sequence from position 481 to 483 is not AAT; if the nucleotide sequence from position 484 to position 486 is AAA, then the nucleotide sequence from position 487 to position 489 is not AAT simultaneously with the nucleotide sequence from position 490 to position 492 being AGY; if the nucleotide sequence from position 490 to position 492 is TCA, then the nucleotide sequence from position 493 to position 495 is not ACC simultaneously with the nucleotide sequence from position 496 to 498 being AAY; if the nucleotide sequence from position 493 to position 495 is ACC, then the nucleotide sequence from position 496 to 498 is not AAT; the nucleotide sequence from position 496 to position 498 is not AAT; if the nucleotide sequence from position 499 to

position 501 is AAA then the nucleotide sequence from position 502 to position 504 is not TCA or AGC; if the nucleotide sequence from position 508 to position 510 is GTA, then the nucleotide sequence from position 511 to 513 is not TTA; if the nucleotide sequence from position 514 to position 516 is AAT then the nucleotide sequence from position 517 to position 519 is not ACA; if the nucleotide sequence from position 517 to position 519 is ACC or ACG, then the nucleotide sequence from position 520 to position 522 is not CAA simultaneously with the nucleotide sequence from position 523 to position 525 being TCN; the codons from the nucleotide sequence from position 523 to position 531 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; if the nucleotide sequence from position 544 to position 546 is GAA then the nucleotide sequence from position 547 to position 549 is not TAT, simultaneously with the nucleotide sequence from position 550 to position 552 being TTR; the codons from the nucleotide sequence from position 547 to position 552 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; if the nucleotide sequence from position 559 to position 561 is GGA then the nucleotide sequence from position 562 to position 564 is not TTG simultaneously with the nucleotide sequence from position 565 to 567 being CGN; if the nucleotide sequence from position 565 to position 567 is CGC then the nucleotide sequence from position 568 to position 570 is not AAT; the nucleotide sequence from position 568 to position 570 is not AAT; if the nucleotide sequence from position 574 to position 576 is TTC then the nucleotide sequence from position 577 to position 579 is not CAA simultaneously with the nucleotide sequence from position 580 to position 582 being TTR; if the nucleotide sequence from position 577 to position 579 is CAA then the nucleotide sequence from position 580 to position 582 is not TTA; if the nucleotide sequence from position 583 to position 585 is AAT the nucleotide sequence from position 586 to 588 is not TGC; the nucleotide sequence from position 595 to position 597 is not AAA; if the nucleotide sequence from position 598 to position 600 is ATT then the nucleotide sequence from position 601 to position 603 is not AAT; the nucleotide sequence from position 598 to position 600 is not ATA; the nucleotide sequence from position 601 to position 603 is not AAT; if the nucleotide sequence from position 604 to position 606 is AAA then the nucleotide sequence from position 607 to position 609 is not AAT; the nucleotide sequence from position 607 to position 609 is not AAT; the nucleotide sequence from position 613 to position 615 is not CCA; if the

nucleotide sequence from position 613 to position 615 is CCG, then the nucleotide sequence from position 616 to position 618 is not ATA; if the nucleotide sequence from position 616 to the nucleotide at position 618 is ATA, then the nucleotide sequence from position 619 to 621 is not ATA; if the nucleotide sequence from position 619 to position 621 is ATA, then the nucleotide sequence from position 622 to position 624 is not TAC; the nucleotide sequence from position 619 to position 621 is not ATT; the codons from the nucleotide sequence from position 640 to position 645 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; if the nucleotide sequence from position 643 to position 645 is TTA then the nucleotide sequence from position 646 to position 648 is not ATA; if the nucleotide sequence from position 643 to position 645 is CTA then the nucleotide sequence from position 646 to position 648 is not ATA; the codons from the nucleotide sequence from position 655 to position 660 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; if the nucleotide sequence from position 658 to 660 is TTA or CTA then the nucleotide sequence from position 661 to position 663 is not ATT or ATC; the nucleotide sequence from position 661 to position 663 is not ATA; if the nucleotide sequence from position 661 to position 663 is ATT then the nucleotide sequence from position 664 to position 666 is not AAA; the codons from the nucleotide sequence from position 670 to position 675 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise ATTTA; if the nucleotide sequence from position 691 to position 693 is TAT then the nucleotide sequence from position 694 to position 696 is not AAA; if the nucleotide sequence from position 694 to position 696 is AAA then the nucleotide sequence from position 697 to position 699 is not TTG; if the nucleotide sequence from position 700 to position 702 is CCC then the nucleotide sequence from position 703 to position 705 is not AAT; if the nucleotide sequence from position 703 to position 705 is AAT then the nucleotide sequence from position 706 to position 708 is not ACA or ACT; if the nucleotide sequence from position 706 to position 708 is ACA then the nucleotide sequence from position 709 to 711 is not ATA simultaneously with the nucleotide sequence from position 712 to position 714 being AGY; the nucleotide sequence does not comprise the codons TTA, CTA, ATA, GTA, TCG, CCG, ACG and GCG; said nucleotide sequence does not comprise a GC stretch consisting of 7 consecutive nucleotides selected

from the group of G or C; and the nucleotide sequence does not comprise a AT stretch consisting of 5 consecutive nucleotides selected from the group of A or T.

A preferred group of synthetic nucleotide sequences is set forth in Table 2 and corresponds to an isolated synthetic DNA fragment is provided which comprises a nucleotide sequence as set forth in SEQ ID No 3, wherein the codons are chosen among the choices provided in such a way as to obtain a nucleotide sequence with an overall GC content of about 50% to about 60%, preferably about 54%-55% provided that if the nucleotide sequence from position 121 to position 123 is GAG then the nucleotide sequence from position 124 to 126 is not CAA; if the nucleotide sequence from position 253 to position 255 is GAC then the nucleotide sequence from position 256 to 258 is not CAA; if the nucleotide sequence from position 277 to position 279 is CAT then the nucleotide sequence from position 280 to 282 is not AAA; if the nucleotide sequence from position 340 to position 342 is AAG then the nucleotide sequence from position 343 to position 345 is not CAT; if the nucleotide sequence from position 490 to position 492 is TCA then the nucleotide sequence from position 493 to position 495 is not ACC; if the nucleotide sequence from position 499 to position 501 is AAA then the nucleotide sequence from position 502 to 504 is not TCA or AGC; if the nucleotide sequence from position 517 to position 519 is ACC then the nucleotide sequence from position 520 to position 522 is not CAA simultaneous with the nucleotide sequence from position 523 to 525 being TCN; if the nucleotide sequence from position 661 to position 663 is ATT then the nucleotide sequence from position 664 to position 666 is not AAA; the codons from the nucleotide sequence from position 7 to position 15 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven contiguous nucleotides from the group of G or C; the codons from the nucleotide sequence from position 61 to position 69 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven contiguous nucleotides from the group of G or C; the codons from the nucleotide sequence from position 130 to position 138 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven contiguous nucleotides from the group of G or C; the codons from the nucleotide sequence from position 268 to position 279 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven

contiguous nucleotides from the group of G or C; the codons from the nucleotide sequence from position 322 to position 333 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven contiguous nucleotides from the group of G or C; the codons from the nucleotide sequence from position 460 to position 468 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven contiguous nucleotides from the group of G or C; the codons from the nucleotide sequence from position 13 to position 27 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; the codons from the nucleotide sequence from position 37 to position 48 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; the codons from the nucleotide sequence from position 184 to position 192 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; the codons from the nucleotide sequence from position 214 to position 219 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; the codons from the nucleotide sequence from position 277 to position 285 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; and the codons from the nucleotide sequence from position 388 to position 396 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; the codons from the nucleotide sequence from position 466 to position 474 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; the codons from the nucleotide sequence from position 484 to position 489 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; the codons from the nucleotide sequence from position 571 to position 576 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the

group of A or T; the codons from the nucleotide sequence from position 598 to position 603 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; the codons from the nucleotide sequence from position 604 to position 609 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; the codons from the nucleotide sequence from position 613 to position 621 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; the codons from the nucleotide sequence from position 646 to position 651 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; the codons from the nucleotide sequence from position 661 to position 666 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; and the codons from the nucleotide sequence from position 706 to position 714 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T.

The nucleotide sequence of SEQ ID No 4 is an example of such a synthetic nucleotide sequence encoding an I-SceI endonuclease which does no longer contain any of the nucleotide sequences or codons to be avoided. However, it will be clear that a person skilled in the art can readily obtain a similar sequence encoding I-SceI by replacing one or more (between two to twenty) of the nucleotides to be chosen for any of the alternatives provided in the nucleotide sequence of SEQ ID 3 (excluding any of the forbidden combinations described in the preceding paragraph) and use it to obtain a similar effect.

For expression in plant cell, the synthetic DNA fragments encoding I-SceI may be operably linked to a plant expressible promoter in order to obtain a plant expressible chimeric gene.

A person skilled in the art will immediately recognize that for this aspect of the invention, it is not required that the repair DNA and/or the DSBI endonuclease encoding DNA are

introduced into the plant cell by direct DNA transfer methods, but that the DNA may thus also be introduced into plant cells by *Agrobacterium*-mediated transformation methods as are available in the art.

In yet another aspect, the invention relates to a method for introducing a foreign DNA of interest into a preselected site of a genome of a plant cell comprising the steps of

- (a) inducing a double stranded break at the preselected site in the genome of the cell ;
- (b) introducing the foreign DNA of interest into the plant cell ;

characterized in that prior to step (a), the plant cells are incubated in a plant phenolic compound.

"Plant phenolic compounds" or "plant phenolics" suitable for the invention are those substituted phenolic molecules which are capable to induce a positive chemotactic response, particularly those who are capable to induce increased vir gene expression in a Ti-plasmid containing *Agrobacterium* sp., particularly a Ti-plasmid containing *Agrobacterium tumefaciens*. Methods to measure chemotactic response towards plant phenolic compounds have been described by Ashby *et al.* (1988 *J. Bacteriol.* **170**: 4181-4187) and methods to measure induction of vir gene expression are also well known (Stachel *et al.*, 1985 *Nature* **318**: 624-629 ; Bolton *et al.* 1986 *Science* **232**: 983-985). Preferred plant phenolic compounds are those found in wound exudates of plant cells. One of the best known plant phenolic compounds is acetosyringone, which is present in a number of wounded and intact cells of various plants, albeit it in different concentrations. However, acetosyringone (3,5-dimethoxy-4-hydroxyacetophenone) is not the only plant phenolic which can induce the expression of vir genes. Other examples are α -hydroxy-acetosyringone, sinapinic acid (3,5 dimethoxy-4-hydroxycinnamic acid), syringic acid (4-hydroxy-3,5 dimethoxybenzoic acid), ferulic acid (4-hydroxy-3-methoxycinnamic acid), catechol (1,2-dihydroxybenzene), p-hydroxybenzoic acid (4-hydroxybenzoic acid), β -resorcylic acid (2,4 dihydroxybenzoic acid), protocatechuic acid (3,4-dihydroxybenzoic acid), pyrogallol (2,3,4 - trihydroxybenzoic acid), gallic acid (3,4,5-trihydroxybenzoic acid) and vanillin (3-methoxy-4-hydroxybenzaldehyde). As used herein, the mentioned molecules are referred to as plant phenolic compounds. Plant phenolic compounds can be added to the plant culture medium either alone or in combination with other plant phenolic compounds. Although not intending

to limit the invention to a particular mode of action, it is thought that the apparent stimulating effect of these plant phenolics on cell division (and thus also genome replication) may be enhancing targeted insertion of foreign DNA.

Plant cells are preferably incubated in plant phenolic compound for about one week, although it is expected incubation for about one or two days in or on a plant phenolic compound will be sufficient. Plant cells should be incubated for a time sufficient to stimulate cell division. According to Guivarc'h *et al.* (1993, *Protoplasma* 174: 10-18) such effect may already be obtained by incubation of plant cells for as little as 10 minutes.

The above mentioned improved methods for homologous recombination based targeted DNA insertion may also be applied to improve the quality of the transgenic plant cells and plants obtained by direct DNA transfer methods, particularly by microprojectile bombardment. It is well known in the art that introduction of DNA by microprojectile bombardment frequently leads to complex integration patterns of the introduced DNA (integration of multiple copies of the foreign DNA of interest, either complete or partial, generation of repeat structures). Nevertheless, some plant genotypes or varieties may be more amenable to transformation using microprojectile bombardment than to transformation using e.g. *Agrobacterium tumefaciens*. It would thus be advantageous if the quality of the transgenic plant cells or plants obtained through microprojectile bombardment could be improved, i.e. if the pattern of integration of the foreign DNA could be influenced to be simpler.

The above mentioned finding that introduction of foreign DNA through microprojectile bombardment in the presence of an induced double stranded DNA break in the nuclear genome, whereby the foreign DNA has homology to the sequences flanking the double stranded DNA break frequently (about 50% of the obtained events) leads to simple integration patterns (single copy insertion in a predictable way and no insertion of additional fragments of the foreign DNA) provides the basis for a method of simplifying the complexity of insertion of foreign DNA in the nuclear genome of plant cells.

Thus the invention also relates to a method of producing a transgenic plant by microprojectile bombardment comprising the steps of

- (a) inducing a double stranded DNA break at a preselected site in the genome of a cell a plant, in accordance with the methods described elsewhere in this document or available in the art; and
- (b) introducing the foreign DNA of interest into the plant cell by microprojectile bombardment wherein said foreign DNA of interest is flanked by two DNA regions having at least 80% sequence identity to the DNA regions flanking the preselected site in the genome of the plant.

A significant portion of the transgenic plant population thus obtained will have a simple integration pattern of the foreign DNA in the genome of the plant cells, more particularly a significant portion of the transgenic plants will only have a one copy insertion of the foreign DNA, exactly between the two DNA regions flanking the preselected site in the genome of the plant. This portion is higher than the population of transgenic plants with simple integration patterns, when the plants are obtained by simple microprojectile bombardment without inducing a double stranded DNA break, and without providing the foreign DNA with homology to the genomic regions flanking the preselected site.

In a convenient embodiment of the invention, the target plant cell comprises in its genome a marker gene, flanked by two recognition sites for a rare-cleaving double stranded DNA break inducing endonuclease, one on each side. This marker DNA may be introduced in the genome of the plant cell of interest using any method of transformation, or may have been introduced into the genome of a plant cell of another plant line or variety (such a as a plant line or variety easy amenable to transformation) and introduced into the plant cell of interest by classical breeding techniques. Preferably, the population of transgenic plants or plant cells comprising a marker gene flanked by two recognition sites for a rare-cleaving double stranded break inducing endonuclease has been analysed for the expression pattern of the marker gene (such as high expression, temporally or spatially regulated expression) and the plant lines with the desired expression pattern identified. Production of a transgenic plant by microprojectile bombardment comprising the steps of

- (a) inducing a double stranded DNA break at a preselected site in the genome of a cell of a plant, in accordance with the methods described elsewhere in this document or available in the art; and
- (b) introducing the foreign DNA of interest into the plant cell by microprojectile bombardment wherein said foreign DNA of interest is flanked by two DNA regions having at least 80% sequence identity to the DNA regions flanking the preselected site in the genome of the plant;

will lead to transgenic plant cells and plants wherein the marker gene has been replaced by the foreign DNA of interest.

The marker gene may be any selectable or a screenable plant-expressible marker gene, which is preferably a conventional chimeric marker gene. The chimeric marker gene can comprise a marker DNA that is under the control of, and operatively linked at its 5' end to, a promoter, preferably a constitutive plant-expressible promoter, such as a CaMV 35S promoter, or a light inducible promoter such as the promoter of the gene encoding the small subunit of Rubisco; and operatively linked at its 3' end to suitable plant transcription termination and polyadenylation signals. The marker DNA preferably encodes an RNA, protein or polypeptide which, when expressed in the cells of a plant, allows such cells to be readily separated from those cells in which the marker DNA is not expressed. The choice of the marker DNA is not critical, and any suitable marker DNA can be selected in a well known manner. For example, a marker DNA can encode a protein that provides a distinguishable color to the transformed plant cell, such as the A1 gene (Meyer *et al.* (1987), *Nature* 330: 677), can encode a fluorescent protein [Chalfie *et al.*, *Science* 263: 802-805 (1994); Crameri *et al.*, *Nature Biotechnology* 14: 315-319 (1996)], can encode a protein that provides herbicide resistance to the transformed plant cell, such as the bar gene, encoding PAT which provides resistance to phosphinothricin (EP 0242246), or can encode a protein that provides antibiotic resistance to the transformed cells, such as the aac(6') gene, encoding GAT which provides resistance to gentamycin (WO 94/01560). Such selectable marker gene generally encodes a protein that confers to the cell resistance to an antibiotic or other chemical compound that is normally toxic for the cells. In plants the selectable marker gene may thus also encode a protein that confers resistance to a herbicide, such as a herbicide comprising a glutamine synthetase inhibitor (e.g. phosphinothricin) as an active ingredient. An example of

such genes are genes encoding phosphinothricin acetyl transferase such as the *sfr* or *sfrv* genes (EP 242236; EP 242246; De Block *et al.*, 1987 *EMBO J.* 6: 2513-2518).

The introduced repair DNA may further comprise a marker gene that allows to better discriminate between integration by homologous recombination at the preselected site and the integration elsewhere in the genome. Such marker genes are available in the art and include marker genes whereby the absence of the marker gene can be positively selected for under selective conditions (e.g. *codA*, cytosine deaminase from *E. coli* conferring sensitivity to 5-fluoro cytosine, Perera *et al.* 1993 *Plant Mol. Biol.* 23, 793; Stougaard (1993) *Plant J.*: 755). The repair DNA needs to comprise the marker gene in such a way that integration of the repair DNA into the nuclear genome in a random way results in the presence of the marker gene whereas the integration of the repair DNA by homologous recombination results in the absence of the marker gene.

It will be immediately clear that the same results can also be obtained using only one preselected site at which to induce the double stranded break, which is located in or near a marker gene. The flanking regions of homology are then preferably chosen in such way as to either inactivate the marker gene, or delete the marker gene and substitute for the foreign DNA to be inserted.

It will be appreciated that the means and methods of the invention are particularly useful for corn, but may also be used in other plants with similar effects, particularly in cereal plants including wheat, oat, barley, rye, rice, turfgrass, sorghum, millet or sugarcane plants. The methods of the invention can also be applied to any plant including but not limited to cotton, tobacco, canola, oilseed rape, soybean, vegetables, potatoes, *Lemna* spp., *Nicotiana* spp., *Arabidopsis*, alfalfa, barley, bean, corn, cotton, flax, pea, rape, rice, rye, safflower, sorghum, soybean, sunflower, tobacco, wheat, asparagus, beet, broccoli, cabbage, carrot, cauliflower, celery, cucumber, eggplant, lettuce, onion, oilseed rape, pepper, potato, pumpkin, radish, spinach, squash, tomato, zucchini, almond, apple, apricot, banana, blackberry, blueberry, cacao, cherry, coconut, cranberry, date, grape, grapefruit, guava, kiwi, lemon, lime, mango, melon, nectarine, orange, papaya, passion fruit, peach, peanut, pear, pineapple, pistachio, plum, raspberry, strawberry, tangerine, walnut and watermelon.

It is also an object of the invention to provide plant cells and plants comprising foreign DNA molecules inserted at preselected sites, according to the methods of the invention. Gametes, seeds, embryos, either zygotic or somatic, progeny or hybrids of plants comprising the targeted DNA insertion events, which are produced by traditional breeding methods are also included within the scope of the present invention.

The plants obtained by the methods described herein may be further crossed by traditional breeding techniques with other plants to obtain progeny plants comprising the targeted DNA insertion events obtained according to the present invention.

The following non-limiting Examples describe the design of a modified I-SceI encoding chimeric gene, and the use thereof to insert foreign DNA into a preselected site of the plant genome.

Unless stated otherwise in the Examples, all recombinant DNA techniques are carried out according to standard protocols as described in Sambrook *et al.* (1989) *Molecular Cloning: A Laboratory Manual*, Second Edition, Cold Spring Harbor Laboratory Press, NY and in Volumes 1 and 2 of Ausubel *et al.* (1994) *Current Protocols in Molecular Biology*, Current Protocols, USA. Standard materials and methods for plant molecular work are described in *Plant Molecular Biology Labfax* (1993) by R.D.D. Croy, jointly published by BIOS Scientific Publications Ltd (UK) and Blackwell Scientific Publications, UK. Other references for standard molecular biology techniques include Sambrook and Russell (2001) *Molecular Cloning: A Laboratory Manual*, Third Edition, Cold Spring Harbor Laboratory Press, NY, Volumes I and II of Brown (1998) *Molecular Biology LabFax*, Second Edition, Academic Press (UK). Standard materials and methods for polymerase chain reactions can be found in Dieffenbach and Dveksler (1995) *PCR Primer: A Laboratory Manual*, Cold Spring Harbor Laboratory Press, and in McPherson *et al.* (2000) *PCR - Basics: From Background to Bench*, First Edition, Springer Verlag, Germany.

Throughout the description and Examples, reference is made to the following sequences:

SEQ ID No 1: amino acid sequence of a chimeric I-SceI comprising a nuclear localization signal linked to a I-SceI protein lacking the 4 amino-terminal amino acids.

SEQ ID No 2: nucleotide sequence of I-SceI coding region (UIPAC code).

SEQ ID No 3: nucleotide sequence of synthetic I-SceI coding region (UIPAC code).

SEQ ID No 4: nucleotide sequence of synthetic I-SceI coding region.

SEQ ID No 5: nucleotide sequence of the T-DNA of pTTAM78 (target locus).

SEQ ID No 6: nucleotide sequence of the T-DNA of pTTA82(repair DNA).

SEQ ID No 7: nucleotide sequence of pCV78.

Table 1. (corresponding to SEQ ID 2)

Trinucleotide	AA	Possible trinucleotides	UIPAC code	PROVISIO
R1	M	ATG	ATG	
R2	A	GCA GCC GCG GCT	GCN	
R3	K	AAA AAG	AAR	
R4	P	CCA CCC CCG CCT	CCN	
R5	P	CCA CCC CCG CCT	CCN	
R6	K	AAA AAG	AAR	
R7	K	AAA AAG	AAR	
R8	K	AAA AAG	AAR	
R9	R	AGA AGG CGA CGC CGG CGT	AGR or CGN	
R10	K	AAA AAG	AAR	NOT AAG
R11	V	GTA GTC GTG GTT	GTN	
R12	N	AAC AAT	AAY	IF R12 AAT NOT (R13 ATT OR R13 ATA). IF R12 AAC NOT (R13 ATT AND R14 AAA) IF R12 AAC NOT R13 ATA
R13	I	ATA ATC ATT	ATH	IF R13 ATT NOT R14 AAA IF R13 ATA NOT R14 AAA
R14	K	AAA AAG	AAR	
R15	K	AAA AAG	AAR	
R16	N	AAC AAT	AAY	
R17	Q	CAA CAG	CAR	NOT CAA
R18	V	GTA GTC GTG GTT	GTN	NOT GTA
R19	M	ATG	ATG	
R20	N	AAC AAT	AAY	AVOID ATTTA
R21	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R22	G	GGA GGC GGG GGT	GGN	
R23	P	CCA CCC CCG CCT	CCN	IF R23 CCC NOT R24 AAT
R24	N	AAC AAT	AAY	
R25	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	
R26	K	AAA AAG	AAR	IF R26 AAA NOT (R27 TTG AND R28 CTN)
R27	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	IF R27 (TTA OR CTA) NOT R28 TTA
R28	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R29	K	AAA AAG	AAR	
R30	E	GAA GAG	GAR	NOT GAA
R31	Y	TAC TAT	TAY	IF R31 TAT NOT R32 AAA

Trinucleotide	AA	Possible trinucleotides	UIPAC code	PROVISIO
R32	K	AAA AAG	AAR	
R33	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	IF R33 (TCC OR TCG OR AGC) NOT (R34 CAA AND R35 TTR)
R34	Q	CAA CAG	CAR	IF R34 CAA NOT R35 TTA
R35	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R36	I	ATA ATC ATT	ATH	
R37	E	GAA GAG	GAR	IF R37 GAA NOT R38 TTA
R38	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R39	N	AAC AAT	AAY	IF R39 AAT NOT R40 (ATT OR ATA)
R40	I	ATA ATC ATT	ATH	
R41	E	GAA GAG	GAR	IF R41 GAG NOT R42 CAA
R42	Q	CAA CAG	CAR	
R43	F	TTC TTT	TTY	
R44	E	GAA GAG	GAR	
R45	A	GCA GCC GCG GCT	GCN	NOT GCA
R46	G	GGA GGC GGG GGT	GCN	
R47	I	ATA ATC ATT	ATH	NOT ATT
R48	G	GGA GGC GGG GGT	GCN	IF R48 GGA NOT R49 TTA
R49	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	IF R49 TTA NOT (R50 ATA AND R51 TTR) IF R49 CTA NOT (R50 ATA AND R51 TTR) IF R50 ATA NOT R51 (CTA OR TTG)
R50	I	ATA ATC ATT	ATH	
R51	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R52	G	GGA GGC GGG GGT	GCN	
R53	D	GAC GAT	GAY	
R54	A	GCA GCC GCG GCT	GCN	IF R54 GCA NOT R55 TAC
R55	Y	TAC TAT	TAY	IF R55 TAT NOT (R56 ATA AND R57 AGR)
R56	I	ATA ATC ATT	ATH	
R57	R	AGA AGG CGA CGC CGG CGT	AGR or CGN	
R58	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	AVOID GCAGG
R59	R	AGA AGG CGA CGC CGG CGT	AGR or CGN	
R60	D	GAC GAT	GAY	
R61	E	GAA GAG	GAR	AVOID AAGGT
R62	G	GGA GGC GGG GGT	GCN	
R63	K	AAA AAG	AAR	
R64	T	ACA ACC ACG ACT	ACN	
R65	Y	TAC TAT	TAY	IF R65 TAT NOT R66 TGC

Trinucleotide	AA	Possible trinucleotides	UIPAC code	PROVISIO
R66	C	TGC TGT	TGY	
R67	M	ATG	ATG	
R68	Q	CAA CAG	CAR	NOT CAA
R69	F	TTC TTT	TTY	
R70	E	GAA GAG	GAR	
R71	W	TGG	TGG	
R72	K	AAA AAG	AAR	
R73	N	AAC AAT	AAY	NOT AAT
R74	K	AAA AAG	AAR	IF R74 AAA NOT R75 GCA
R75	A	GCA GCC GCG GCT	GCN	IF R75 GCA NOT R76 TAC
R76	Y	TAC TAT	TAY	
R77	M	ATG	ATG	
R78	D	GAC GAT	GAY	
R79	H	CAC CAT	CAY	
R80	V	GTA GTC GTG GTT	GTN	
R81	C	TGC TGT	TGY	
R82	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R83	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R84	Y	TAC TAT	TAY	
R85	D	GAC GAT	GAY	IF R85 GAC NOT R86 CAA
R86	Q	CAA CAG	CAR	
R87	W	TGG	TGG	
R88	V	GTA GTC GTG GTT	GTN	
R89	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R90	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	
R91	P	CCA CCC CCG CCT	CCN	
R92	P	CCA CCC CCG CCT	CCN	
R93	H	CAC CAT	CAY	IF R93 CAT NOT R94 AAA
R94	K	AAA AAG	AAR	
R95	K	AAA AAG	AAR	
R96	E	GAA GAG	GAR	
R97	R	AGA AGG CGA CGC CGG CGT	AGR or CGN	
R98	V	GTA GTC GTG GTT	GTN	
R99	N	AAC AAT	AAY	
R100	H	CAC CAT	CAY	AVOID ATTTA
R101	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	

Trinucleotide	AA	Possible trinucleotides	UIPAC code	PROVISIO
R102	G	GGA GGC GGG GGT	GCN	IF R102 GGC NOT R103 AAT
R103	N	AAC AAT	AAY	AVOID ATTTA
R104	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R105	V	GTA GTC GTG GTT	GCN	
R106	I	ATA ATC ATT	ATH	
R107	T	ACA ACC ACG ACT	ACN	
R108	W	TGG	TGG	
R109	G	GGA GGC GGG GGT	GCN	
R110	A	GCA GCC GCG GCT	GCN	
R111	Q	CAA CAG	CAR	
R112	T	ACA ACC ACG ACT	ACN	AVOID ATTTA
R113	F	TTC TTT	TTY	
R114	K	AAA AAG	AAR	IF R114 AAG NOT R115 CAT
R115	H	CAC CAT	CAY	
R116	Q	CAA CAG	CAR	IF R116 CAA NOT R117 GCA
R117	A	GCA GCC GCG GCT	GCN	AVOID ATTTA
R118	F	TTC TTT	TTY	
R119	N	AAC AAT	AAY	NOT AAT
R120	K	AAA AAG	AAR	IF R120 AAA NOT R121 TTG
R121	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R122	A	GCA GCC GCG GCT	GCN	IF R122 GCC NOT R123 AAT
R123	N	AAC AAT	AAY	AVOID ATTTA
R124	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R125	F	TTC TTT	TTY	
R126	I	ATA ATC ATT	ATH	
R127	V	GTA GTC GTG GTT	GCN	
R128	N	AAC AAT	AAY	IF R128 AAT NOT R129 AAT
R129	N	AAC AAT	AAY	NOT AAT
R130	K	AAA AAG	AAR	
R131	K	AAA AAG	AAR	
R132	T	ACA ACC ACG ACT	ACN	
R133	I	ATA ATC ATT	ATH	
R134	P	CCA CCC CCG CCT	CCN	IF R134 CCC NOT R135 AAT
R135	N	AAC AAT	AAY	IF R135 AAT NOT R136 AAT
R136	N	AAC AAT	AAY	AVOID ATTTA
R137	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	

Trinucleotide	AA	Possible trinucleotides	UIPAC code	PROVISIO
R138	V	GTA GTC GTG GTT	GTN	
R139	E	GAA GAG	GAR	
R140	N	AAC AAT	AAY	
R141	Y	TAC TAT	TAY	AVOID ATTTA
R142	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R143	T	ACA ACC ACG ACT	ACN	
R144	P	CCA CCC CCG CCT	CCN	NOT CCA
R145	M	ATG	ATG	
R146	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	IF R146 TCA NOT R147 TTG
R147	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R148	A	GCA GCC GCG GCT	GCN	
R149	Y	TAC TAT	TAY	NOT TAT
R150	W	TGG	TGG	
R151	F	TTC TTT	TTY	
R152	M	ATG	ATG	
R153	D	GAC GAT	GAY	
R154	D	GAC GAT	GAY	
R155	G	GGA GGC GGG GGT	GCN	
R156	G	GGA GGC GGG GGT	GCN	
R157	K	AAA AAG	AAR	
R158	W	TGG	TGG	
R159	D	GAC GAT	GAY	
R160	Y	TAC TAT	TAY	
R161	N	AAC AAT	AAY	NOT AAT
R162	K	AAA AAG	AAR	IF R162 AAA NOT (R163 AAT AND R164 AGY)
R163	N	AAC AAT	AAY	
R164	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	IF R164 TCA NOT (R165 ACC AND R166 AAY)
R165	T	ACA ACC ACG ACT	ACN	IF R165 ACC NOT R166 AAT
R166	N	AAC AAT	AAY	NOT AAT
R167	K	AAA AAG	AAR	IF R167 AAA R168 NOT TCA OR R168 NOT AGC
R168	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	
R169	I	ATA ATC ATT	ATH	
R170	V	GTA GTC GTG GTT	GTN	IF R170 GTA NOT R171TTA
R171	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R172	N	AAC AAT	AAY	IF R172 AAT NOT R173 ACA
R173	T	ACA ACC ACG ACT	ACN	IF R173 (ACC OR ACG) NOT (R174 CAA AND

Trinucleotide	AA	Possible trinucleotides	UIPAC code	PROVISIO
R174	Q	CAA CAG	CAR	R175 TCN)
R175	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	AVOID ATTTA
R176	F	TTC TTT	TTY	
R177	T	ACA ACC ACG ACT	ACN	
R178	F	TTC TTT	TTY	
R179	E	GAA GAG	GAR	
R180	E	GAA GAG	GAR	
R181	V	GTA GTC GTG GTT	GTN	
R182	E	GAA GAG	GAR	IF R182 GAA NOT (R183 TAT AND R184 TTR)
R183	Y	TAC TAT	TAY	AVOID ATTTA
R184	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R185	V	GTA GTC GTG GTT	GTN	
R186	K	AAA AAG	AAR	
R187	G	GGA GGC GGG GGT	GGN	IF R187 GGA NOT (R188 TTG AND R189 CGN)
R188	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R189	R	AGA AGG CGA CGC CGG CGT	AGR or CGN	IF R189 CGC NOT R190 AAT
R190	N	AAC AAT	AAY	NOT AAT
R191	K	AAA AAG	AAR	
R192	F	TTC TTT	TTY	IF R192 TTC NOT (R193 CAA AND R194 TTR)
R193	Q	CAA CAG	CAR	IF R193 CAA NOT R194 TTA
R194	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R195	N	AAC AAT	AAY	IF R195 AAT NOT R196 TGC
R196	C	TGC TGT	TGY	
R197	Y	TAC TAT	TAY	
R198	V	GTA GTC GTG GTT	GTN	
R199	K	AAA AAG	AAR	NOT AAA
R200	I	ATA ATC ATT	ATH	IF R200 ATT NOT R201 AAT
R201	N	AAC AAT	AAY	NOT ATA
R202	K	AAA AAG	AAR	NOT AAT
R203	N	AAC AAT	AAY	IF R202 AAA NOT R203 AAT
R204	K	AAA AAG	AAR	NOT AAT
R205	P	CCA CCC CCG CCT	CCN	NOT CCA
R206	I	ATA ATC ATT	ATH	IF R205 CCG NOT R206 ATA
				IF R206 ATA NOT R207 ATA

Trinucleotide	AA	Possible trinucleotides	UIPAC code	PROVISIO
R207	I	ATA ATC ATT	ATH	IF R207 ATA NOT R208 TAC NOT ATT
R208	Y	TAC TAT	TAY	
R209	I	ATA ATC ATT	ATH	
R210	D	GAC GAT	GAY	
R211	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	
R212	M	ATG	ATG	
R213	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	
R214	Y	TAC TAT	TAY	AVOID ATTTA
R215	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	IF R215 (TTA OR CTA) NOT R216 ATA
R216	I	ATA ATC ATT	ATH	
R217	F	TTC TTT	TTY	
R218	Y	TAC TAT	TAY	
R219	N	AAC AAT	AAY	AVOID ATTTA
R220	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	IF R220 (TTA OR CTA) NOT R221 ATT IF R220 (TTA OR CTA) NOT R221 ATC
R221	I	ATA ATC ATT	ATH	IF R221 ATT NOT R222 AAA NOT ATA
R222	K	AAA AAG	AAR	
R223	P	CCA CCC CCG CCT	CCN	
R224	Y	TAC TAT	TAY	AVOID ATTTA
R225	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R226	I	ATA ATC ATT	ATH	
R227	P	CCA CCC CCG CCT	CCN	
R228	Q	CAA CAG	CAR	
R229	M	ATG	ATG	
R230	M	ATG	ATG	
R231	Y	TAC TAT	TAY	IF R231TAT NOT R232 AAA
R232	K	AAA AAG	AAR	IF R232 AAA NOT R233 TTG
R233	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R234	P	CCA CCC CCG CCT	CCN	IF 234 CCC NOT R235 AAT
R235	N	AAC AAT	AAY	IF R235 AAT NOT R236 ACA IF R235 AAT NOT R236 ACT
R236	T	ACA ACC ACG ACT	ACN	IF R236 ACA NOT (R237 ATA AND R238 AGY)
R237	I	ATA ATC ATT	ATH	
R238	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	

Trinucleotide	AA	Possible trinucleotides	UIPAC code	PROVISIO
R239	S	AGC AGT TCA TCC TCG TCT	AGY or TCN	
R240	E	GAA GAG	GAR	
R241	T	ACA ACC ACG ACT	ACN	
R242	F	TTC TTT	TTY	
R243	L	TTA TTG CTA CTC CTG CTT	TTR or CTN	
R244	K	AAA AAG	AAR	

Table 2. (corresponding to SEQ ID No 3)

Trinucleotide	AA	Choices	UIPAC	PROVISIO	Exemplified I-SceI (SEQ ID No 4)
R1	M	ATG	ATG		ATG
R2	A	GCC GCT	GCY		GCC
R3	K	AAA AAG	AAR		AAG
R4	P	CCA CCC CCT	CCH		CCT
R5	P	CCA CCC CCT	CCH		CCC
R6	K	AAA AAG	AAR		AAG
R7	K	AAA AAG	AAR		AAG
R8	K	AAA AAG	AAR		AAG
R9	R	AGA CGC CGG	AGA or CGS		CGC
R10	K	AAA	AAA		AAA
R11	V	GTC GTG	GTS		GTG
R12	N	AAC	AAC		AAC
R13	I	ATC ATT	ATY		ATC
R14	K	AAA AAG	AAR		AAG
R15	K	AAA AAG	AAR		AAG
R16	N	AAC	AAC		AAC
R17	Q	CAG	CAG		CAG
R18	V	GTC GTG	GTS		GTG
R19	M	ATG	ATG		ATG
R20	N	AAC	AAC		AAC
R21	L	CTC CTG	CTS		CTG
R22	G	GGC GGA	GGM		GGA
R23	P	CCA CCC CCT	CCH		CCT
R24	N	AAC	AAC		AAC

Trinucleotide	AA	Choices	UIPAC	PROVISIO	Exemplified I-SceI (SEQ ID No 4)
R25	S	AGC TCA TCC	AGC or TCM		AGC
R26	K	AAA AAG	AAR		AAG
R27	L	CTC CTG	CTS		CTC
R28	L	CTC CTG	CTS		CTG
R29	K	AAA AAG	AAR		AAG
R30	E	GAG	GAG		GAG
R31	Y	TAC	TAC		TAC
R32	K	AAA AAG	AAR		AAG
R33	S	AGC TCA TCC	AGC or TCM		AGC
R34	Q	CAA CAG	CAR		CAG
R35	L	CTC CTG	CTS		CTG
R36	I	ATC ATT	ATY		ATC
R37	E	GAA GAG	GAR		GAA
R38	L	CTC CTG	CTS		CTG
R39	N	AAC	AAC		AAC
R40	I	ATC ATT	ATY		ATC
R41	E	GAA GAG	GAR	IF R41 GAG NOT R42 CAA	GAG
R42	Q	CAA CAG	CAR		CAG
R43	F	TTC	TTC		TTC
R44	E	GAA GAG	GAR		GAA
R45	A	GCC GCT	GCY		GCT
R46	G	GGC GGA	GGM		GGC
R47	I	ATC	ATC		ATC
R48	G	GGC GGA	GGM		GGC
R49	L	CTC CTG	CTS		CTG
R50	I	ATC ATT	ATY		ATC
R51	L	CTC CTG	CTS		CTG
R52	G	GGC GGA	GGM		GGC
R53	D	GAC GAT	GAY		GAT
R54	A	GCC GCT	GCY		GCC
R55	Y	TAC	TAC		TAC
R56	I	ATC ATT	ATY		ATC
R57	R	AGA CGC CGG	AGA or CGS		AGA
R58	S	AGC TCA TCC	AGC or TCM		TCC
R59	R	AGA CGC CGG	AGA or CGS		CGG

Trinucleotide	AA	Choices	UIPAC	PROVISIO	Exemplified I-SceI (SEQ ID No 4)
R60	D	GAC GAT	GAY		GAC
R61	E	GAA GAG	GAR		GAA
R62	G	GGC GGA	GGM		GGC
R63	K	AAA AAG	AAR		AAG
R64	T	ACC ACT	ACY		ACC
R65	Y	TAC	TAC		TAC
R66	C	TGC TGT	TGY		TGC
R67	M	ATG	ATG		ATG
R68	Q	CAG	CAG		CAG
R69	F	TTC	TTC		TTC
R70	E	GAA GAG	GAR		GAG
R71	W	TGG	TGG		TGG
R72	K	AAA AAG	AAR		AAG
R73	N	AAC	AAC		AAC
R74	K	AAA AAG	AAR		AAG
R75	A	GCC GCT	GCY		GCC
R76	Y	TAC	TAC		TAC
R77	M	ATG	ATG		ATG
R78	D	GAC GAT	GAY		GAC
R79	H	CAC CAT	CAY		CAC
R80	V	GTC GTG	GTS		GTG
R81	C	TGC TGT	TGY		TGT
R82	L	CTC CTG	CTS		CTG
R83	L	GTC CTG	CTS		CTG
R84	Y	TAC	TAC		TAC
R85	D	GAC GAT	GAY	IF R85 GAC NOT R86 CAA	GAC
R86	Q	CAA CAG	CAR		CAG
R87	W	TGG	TGG		TGG
R88	V	GTC GTG	GTS		GTC
R89	L	CTC CTG	CTS		CTG
R90	S	AGC TCA TCC	AGC or TCM		AGC
R91	P	CCA CCC CCT	CCH		CCT
R92	P	CCA CCC CCT	CCH		CCT
R93	H	CAC CAT	CAY	IF R93 CAT NOT R94 AAA	CAC
R94	K	AAA AAG	AAR		AAG

Trinucleotide	AA	Choices	UIPAC	PROVISIO	Exemplified I-SceI (SEQ ID No 4)
R95	K	AAA AAG	AAR		AAG
R96	E	GAA GAG	GAR		GAG
R97	R	AGA CGC CGG	AGA or CGS		CGC
R98	V	GTC GTG	GTS		GTG
R99	N	AAC	AAC		AAC
R100	H	CAC CAT	CAY		CAT
R101	L	CTC CTG	CTS		CTG
R102	G	GGC GGA	GGM		GGC
R103	N	AAC	AAC		AAC
R104	L	CTC CTG	CTS		CTC
R105	V	GTC GTG	GTS		GTG
R106	I	ATC ATT	ATY		ATC
R107	T	ACC ACT	ACY		ACC
R108	W	TGG	TGG		TGG
R109	G	GGC GGA	GGM		GGA
R110	A	GCC GCT	GCY		GCC
R111	Q	CAA CAG	CAR		CAG
R112	T	ACC ACT	ACY		ACC
R113	F	TTC	TTC		TTC
R114	K	AAA AAG	AAR	IF R114 AAG NOT R115 CAT	AAG
R115	H	CAC CAT	CAY		CAC
R116	Q	CAA CAG	CAR		CAG
R117	A	GCC GCT	GCY		GCC
R118	F	TTC	TTC		TTC
R119	N	AAC	AAC		AAC
R120	K	AAA AAG	AAR		AAG
R121	L	CTC CTG	CTS		CTG
R122	A	GCC GCT	GCS		GCC
R123	N	AAC	AAC		AAC
R124	L	CTC CTG	CTS		CTG
R125	F	TTC	TTC		TTC
R126	I	ATC ATT	ATY		ATC
R127	V	GTC GTG	CTS		GTG
R128	N	AAC	AAC		AAC
R129	N	AAC	AAC		AAC

Trinucleotide	AA	Choices	UIPAC	PROVISIO	Exemplified I-SceI (SEQ ID No 4)
R130	K	AAA AAG	AAR		AAG
R131	K	AAA AAG	AAR		AAG
R132	T	ACC ACT	ACY		ACC
R133	I	ATC ATT	ATY		ATC
R134	P	CCA CCC CCT	CCH		CCC
R135	N	AAC	AAC		AAC
R136	N	AAC	AAC		AAC
R137	L	CTC CTG	CTS		CTC
R138	V	GTC GTG	GTS		GTG
R139	E	GAA GAG	GAR		GAG
R140	N	AAC	AAC		AAC
R141	Y	TAC	TAC		TAC
R142	L	CTC CTG	CTS		CTC
R143	T	ACC ACT	ACY		ACT
R144	P	CCC CCT	CCY		CCC
R145	M	ATG	ATG		ATG
R146	S	AGC TCA TCC	AGC or TCM		AGC
R147	L	CTC CTG	CTS		CTG
R148	A	GCC GCT	GCY		GCC
R149	Y	TAC	TAC		TAC
R150	W	TGG	TGG		TGG
R151	F	TTC	TTC		TTC
R152	M	ATG	ATG		ATG
R153	D	GAC GAT	GAY		GAC
R154	D	GAC GAT	GAY		GAC
R155	G	GGC GGA	GGM		GGA
R156	G	GGC GGA	GGM		GGC
R157	K	AAA AAG	AAR		AAG
R158	W	TGG	TGG		TGG
R159	D	GAC GAT	GAY		GAC
R160	Y	TAC	TAC		TAC
R161	N	AAC	AAC		AAC
R162	K	AAA AAG	AAR		AAG
R163	N	AAC	AAC		AAC
R164	S	AGC TCA TCC	AGC or TCM	IF R164 TCA NOT R165 ACC	AGC

Trinucleotide	AA	Choices	UIPAC	PROVISIO	Exemplified I-Scel (SEQ ID No 4)
R165	T	ACC ACT	ACY		ACC
R166	N	AAC	AAC		AAC
R167	K	AAA AAG	AAR	IF R167 AAA R168 NOT TCA OR R168 NOT AGC	AAG
R168	S	AGC TCA TCC	AGC or TCM		TCA
R169	I	ATC ATT	ATY		ATT
R170	V	GTC GTG	GTS		GTG
R171	L	CTC CTG	CTS		CTG
R172	N	AAC	AAC		AAC
R173	T	ACC ACT	ACY	IF R173 ACC NOT (R174 CAA AND R175 TCN)	ACC
R174	Q	CAA CAG	CAR		CAA
R175	S	AGC TCA TCC	AGC or TCM		AGC
R176	F	TTC	TTC		TTC
R177	T	ACC ACT	ACY		ACC
R178	F	TTC	TTC		TTC
R179	E	GAA GAG	GAR		GAA
R180	E	GAA GAG	GAR		GAA
R181	V	GTC GTG	GTS		GTG
R182	E	GAA GAG	GAR		GAG
R183	Y	TAC	TAC		TAC
R184	L	CTC CTG	CTS		CTC
R185	V	GTC GTG	GTS		GTC
R186	K	AAA AAG	AAR		AAG
R187	G	GGC GGA	GGM		GGC
R188	L	CTC CTG	CTS		CTG
R189	R	AGA CGC CGG	AGA or CGS		CGC
R190	N	AAC	AAC		AAC
R191	K	AAA AAG	AAR		AAG
R192	F	TTC	TTC		TTC
R193	Q	CAA CAG	CAR		CAG
R194	L	CTC CTG	CTS		CTG
R195	N	AAC	AAC		AAC
R196	C	TGC TGT	TGY		TGC
R197	Y	TAC	TAC		TAC

Trinucleotide	AA	Choices	UIPAC	PROVISIO	Exemplified I-SceI (SEQ ID No 4)
R198	V	GTC GTG	GTS		GTG
R199	K	AAG	AAG		AAG
R200	I	ATC ATT	ATY		ATC
R201	N	AAC	AAC		AAC
R202	K	AAA AAG	AAR		AAG
R203	N	AAC	AAC		AAC
R204	K	AAA AAG	AAR		AAG
R205	P	CCC CCT	CCY		CCT
R206	I	ATC ATT	ATY		ATC
R207	I	ATC	ATC		ATC
R208	Y	TAC	TAC		TAC
R209	I	ATC ATT	ATY		ATC
R210	D	GAC GAT	GAY		GAC
R211	S	AGC TCA TCC	AGC or TCM		AGC
R212	M	ATG	ATG		ATG
R213	S	AGC TCA TCC	AGC or TCM		AGC
R214	Y	TAC	TAC		TAC
R215	L	CTC CTG	CTS		CTG
R216	I	ATC ATT	ATY		ATC
R217	F	TTC	TTC		TTC
R218	Y	TAC	TAC		TAC
R219	N	AAC	AAC		AAC
R220	L	CTC CTG	CTS		CTG
R221	I	ATC ATT	ATY	IF R221 ATT NOT R222 AAA	ATC
R222	K	AAA AAG	AAR		AAG
R223	P	CCA CCC CCT	CCH		CCA
R224	Y	TAC	TAC		TAC
R225	L	CTC CTG	CTS		CTG
R226	I	ATC ATT	ATY		ATC
R227	P	CCA CCC CCT	CCH		CCT
R228	Q	CAA CAG	CAR		CAG
R229	M	ATG	ATG		ATG
R230	M	ATG	ATG		ATG
R231	Y	TAC	TAC		TAC
R232	K	AAA AAG	AAR		AAG

Trinucleotide	AA	Choices	UIPAC	PROVISIO	Exemplified I-Scel (SEQ ID No 4)
R233	L	CTC CTG	CTS		CTG
R234	P	CCA CCC CCT	CCH		CCC
R235	N	AAC	AAC		AAC
R236	T	ACC ACT	ACY		ACC
R237	I	ATC ATT	ATY		ATC
R238	S	AGC TCA TCC	AGC or TCM		AGC
R239	S	AGC TCA TCC	AGC or TCM		AGC
R240	E	GAA GAG	GAR		GAG
R241	T	ACC ACT	ACY		ACC
R242	F	TTC	TTC		TTC
R243	L	CTC CTG	CTS		CTG
R244	K	AAA AAG	AAR		AAG

Examples

Example I: Design, synthesis and analysis of a plant expressible chimeric gene encoding I-SceI.

The coding region of I-SceI wherein the 4 aminoterminal amino acids have been replaced by a nuclear localization signal was optimized using the following process:

1. Change the codons to the most preferred codon usage for maize without altering the amino acid sequence of I-SceI protein, using the Synergy Geneoptimizer™;
2. Adjust the sequence to create or eliminate specific restriction sites to exchange the synthetic I-SceI coding region with the universal code I-SceI gene;
3. Eliminate all GC stretches longer than 6 bp and AT stretches longer than 4 bp to avoid formation of secondary RNA structures than can effect pre-mRNA splicing
4. Avoid CG and TA duplets in codon positions 2 and 3;
5. Avoid other regulatory elements such as possible premature polyadenylation signals (GATAAT, TATAAA, AATATA, AATATT, GATAAA, AATGAA, AATAAG, AATAAA, AATAAT, AACCAA, ATATAA, AATCAA, ATACTA, ATAAAA, ATGAAA, AAGCAT, ATTAAT, ATACAT, AAAATA, ATTAAA, AATTAA, AATACA and CATAAA), cryptic intron splice sites (AAGGTAAGT and TGCAGG), ATTTA pentamers and CCAAT box sequences (CCAAT, ATTGG, CGAAT and ATTGC);
6. Recheck if the adapted coding region fulfill all of the above mentioned criteria.

A possible example of such a nucleotide sequence is represented in SEQ ID No 4. A synthetic DNA fragment having the nucleotide sequence of SEQ ID No 4 was synthesized and operably linked to a CaMV35S promoter and a CaMV35S 3' termination and polyadenylation signal (yielding plasmid pCV78; SEQ ID No 7).

The synthetic I-SceI coding region was also cloned into a bacterial expression vector (as a fusion protein allowing protein enrichment on amylose beads). The capacity of semi-purified I-SceI protein to cleave in vitro a plasmid containing an I-SceI recognition site was verified.

Example 2. Isolation of maize cell lines containing a promoterless bar gene preceded by an I-SceI site.

In order to develop an assay for double stranded DNA break induced homology-mediated recombination, maize cell suspensions were isolated that contained a promoterless bar gene preceded by an I-SceI recognition site integrated in the nuclear genome in single copy. Upon double stranded DNA break induction through delivery of an I-SceI endonuclease encoding plant expressible chimeric gene, and co-delivery of repair DNA comprising a CaMV 35S promoter operably linked to the 5' end of the bar gene, the 35S promoter may be inserted through homology mediated targeted DNA insertion, resulting in a functional bar gene allowing resistance to phosphinotricin (PPT). The assay is schematically represented in Figure 1.

The target locus was constructed by operably linking through conventional cloning techniques the following DNA regions

- a) a 3' end termination and polyadenylation signal from the nopaline synthetase gene
- b) a promoter-less bar encoding DNA region
- c) a DNA region comprising an I-SceI recognition site
- d) a 3' end termination and polyadenylation signal from *A. tumefaciens* gene 7 (3'g7)
- e) a plant expressible neomycin resistance gene comprising a nopaline synthetase promoter, a neomycine phosphotransferase gene, and a 3' ocs signal.

This DNA region was inserted in a T-DNA vector between the T-DNA borders. The T-DNA vector was designated pTTAM78 (for nucleotide sequence of the T-DNA see SEQ ID No 5)

The T-DNA vector was used directly to transform protoplasts of corn according to the methods described in EP 0 469 273, using a He89-derived corn cell suspension. The T-DNA vector was also introduced into *Agrobacterium tumefaciens* C58C1Rif(pEHA101) and the resulting *Agrobacterium* was used to transform an He89-derived cell line. A number of target lines were identified that contained a single copy of the target locus construct pTTAM78, such as T24 (obtained by protoplast transformation) and lines 14-1 and 1-20 (obtained by *Agrobacterium* mediated transformation)

Cell suspensions were established from these target lines in N6M cell suspension medium, and grown in the light on a shaker (120 rpm) at 25°C. Suspensions were subcultured every week.

Example 3: Homology based targeted insertion.

The repair DNA pTTA82 is a T-DNA vector containing between the T-DNA borders the following operably linked DNA regions:

- a) a DNA region encoding only the aminoterminal part of the bar gene
- b) a DNA region comprising a partial I-SceI recognition site (13 nucleotides located at the 5' end of the recognition site)
- c) a CaMV 35S promoter region
- d) a DNA region comprising a partial I-SceI recognition site (9 nucleotides located at the 3' end of the recognition site)
- e) a 3' end termination and polyadenylation signal from *A.tumefaciens* gene 7 (3'g7)
- f) a chimeric plant expressible neomycine resistance gene
- g) a defective I-SceI endonuclease encoding gene under control of a CaMV 35S promoter

The nucleotide sequence of the T-DNA of pTTA82 is represented in SEQ ID NO 6.

This repair DNA was co-delivered with pCV78 (see Example 1) by particle bombardment into suspension derived cells which were plated on filter paper as a thin layer. The filter paper was plated on Mahq1 VII substrate.

The DNA was bombarded into the cells using a PDS-1000/He Biolistics device. Microcarrier preparation and coating of DNA onto microcarriers was essentially as described by Sanford *et al.* 1992. Particle bombardment parameters were: target distance of 9cm; bombardment pressure of 1350 psi, gap distance of ¼'' and macrocarrier flight distance of 11 cm. Immediately after bombardment the tissue was transferred onto non-selective Mhi1VII substrate. As a control for successful delivery of DNA by particle bombardment, the three

target lines were also bombarded with microcarriers coated with plasmid DNA comprising a chimeric bar gene under the control of a CaMV35S promoter (pRVA52).

Four days after bombardment, the filters were transferred onto Mh1 VII substrate supplemented with 25 mg/L PPT or on Ahx1.5VIIino1000 substrate supplemented with 50 mg/L PPT.

Fourteen days later, the filters were transferred onto fresh Mh1 VII medium with 10 mg/L PPT for the target lines T24 and 14-1 and Mh1 VII substrate with 25 mg/L PPT for target line 1-20.

Two weeks later, potential targeted insertion events were scored based on their resistance to PPT. These PPT resistant events were also positive in the Liberty Link Corn Leaf/Seed test (Strategic Diagnostics Inc.).

Number of PPT resistant calli 38 days after bombardment:

Target line	pRVA52		pTTA82+pCV78	
	Total number of PPT ^R events	Mean number of PPT ^R events/petridish	Total number of PPT ^R events	Mean number of PPT ^R events/petridish
1-20	75	25	115	7.6
14-1	37	12.3	38	2.2
24	40	13.3	2	0.13

The PPT resistant events were further subcultured on Mh1 VII substrate containing 10 mg/L PPT and callus material was used for molecular analysis. Twenty independent candidate TSI were analyzed by Southern analysis using the 35S promoter and the 3' end termination and polyadenylation signal from the nopaline synthase gene as a probe. Based on the size of the expected fragment, all events appeared to be perfect targeted sequence insertion events. Moreover, further analysis of about half of the targeted sequence insertion events did not

show additional non-targeted integration of either the repair DNA or the I-SceI encoding DNA.

Sequence analysis of DNA amplified from eight of the targeted insertion events demonstrated that these events were indeed perfect homologous recombination based TSI events.

Based on these data, the ratio of homologous recombination based DNA insertion versus the “normal” illegitimate recombination varies from about 30% for 1-20 to about 17% for 14-1 and to about 1% for 24.

When using vectors similar to the ones described in Puchta *et al*, 1996 (*supra*) delivered by electroporation to tobacco protoplasts in the presence of I-SceI induced double stranded DNA breaks, the ratio of homologous recombination based DNA insertion versus normal insertion was about 15%. However, only one of out of 33 characterized events was a homology-mediated targeted sequence insertion event whereby the homologous recombination was perfect at both sides of the double stranded break.

Using the vectors from Example 2, but with a “universal code I-SceI construct” comprising a nuclear localization signal, the ratio of HR based DNA insertion versus normal insertion varied between 0.032% and 16% for different target lines, both using electroporation or *Agrobacterium* mediated DNA delivery. The relative frequency of perfect targeted insertion events differed between the different target lines, and varied from 8 to 70% for electroporation mediated DNA delivery and between 73 to 90% for *Agrobacterium* mediated DNA delivery.

Example 4. Acetosyringone pre-incubation improves the frequency of recovery of targeted insertion events.

One week before bombardment as described in Example 3, cell suspensions were either diluted in N6M medium or in LSIDhy1.5 medium supplemented with 200 μ M

acetosyringone. Otherwise, the method as described in Example 3 was employed. As can be seen from the results summarized in the following table, preincubation of the cells to be transformed with acetosyringone had a beneficial effect on the recovery of targeted PPT resistant insertion events.

Target line	Preincubation with acetosyringone		No preincubation	
	Total number of PPT ^R events	Mean number of PPT ^R events/petridish	Total number of PPT ^R events	Mean number of PPT ^R events/petridish
1-20	89	7.6	26	3.7
14-1	32	3.6	6	0.75
24	0	0	2	0.3

Example 5: DSB-mediated targeted sequence insertion in maize by Agrobacterium-mediated delivery of repair DNA.

To analyze DSB-mediated targeted sequence insertion in maize, whereby the repair DNA is delivered by Agrobacterium-mediated transformation, T-DNA vectors were constructed similar to pTTA82 (see Example 3), wherein the defective I-SceI was replaced by the synthetic I-SceI encoding gene of Example 1. The T-DNA vector further contained a copy of the *Agrobacterium tumefaciens virG* and *virC* (pTCV83) or *virG*, *virC* and *virB* (pTCV87) outside the T-DNA borders. These T-DNA vectors were inserted into LBA4404, containing the helper Ti-plasmid pAL4404, yielding *Agrobacterium* strains A4995 and A 4996 respectively.

Suspension cultures of the target cell lines of Example 2, as well as other target cell lines obtained in a similar way as described in Example 2, were co-cultivated with the Agrobacterium strains, and plated thereafter on a number of plates. The number of platings was determined by the density of the cell suspension. As a control for the transformation efficiency, the cell suspension were co-cultivated in a parallel experiment with an Agrobacterium strain LBA4404 containing helper Ti-plasmid pAL4404 and a T-DNA vector

with a chimeric phosphinotricin resistance gene (bar gene) under control of a CaMV 35S vector. The T-DNA vector further contained a copy of the *Agrobacterium tumefaciens* *virG*, *virC* and *virB* genes, outside the T-DNA border. The results of four different independent experiments are summarized in the tables below:

Agrobacterium experiment I:

Target line	Control		A4495	
	N° of platings	N° of transformants	N° of platings ⁽¹⁾	N° of TSI events
T24	26	10	32	0
T26	36	44	36	1
14-1	20	18	28	0
T1 F155	26	7	24	0

Agrobacterium experiment II:

Target line	Control		A4495	
	N° of platings	N° of transformants	N° of platings ⁽¹⁾	N° of TSI events
1-20	18	~200	27	11
T79	24	~480	24	6
T66	26	73	31	0
T5	28	35	18	0
T1 F154	22	65	16	1

Agrobacterium experiment III:

Target line	Control		A4496	
	N° of platings	N° of transformants	N° of platings ⁽¹⁾	N° of TSI events
T24	50	~2250	30	1
T26	44	~220	32	1
14-1	20	~1020	13	1
T1 F155	33	~1870	32	0

Agrobacterium experiment IV:

Target line	A3970		A4496	
	N° of platings	N° of transformants	N° of platings ⁽¹⁾	N° of TSI events
T1 F154			28	1
T5	12	~600	28	1
T66			28	0
T79			24	0
1-20	18	~400	40	9

Thus, it is clear that, while Agrobacterium-mediated repair DNA delivery is clearly feasible, the frequency of Targeted Sequence Insertion (TSI) events is lower in comparison with particle bombardment-mediated repair DNA delivery. Southern analysis performed on 23 putative TSI events showed that 20 TSI events are perfect, based on the size of the fragment. However, in contrast with the events obtained by microprojectile bombardment as in Example 3, only 6 events out of 20 did not contain additional inserts of the repair DNA, 9 events did contain 1 to 3 additional inserts of the repair DNA, and 5 events contained many additional inserts of the repair DNA.

Particle bombardment mediated delivery of repair DNA also results in better quality of DSB mediated TSI events compared to delivery of repair DNA by Agrobacterium. This is in contrast for particle bombardment mediated delivery of "normal transforming DNA" which is characterized by the lesser quality of the transformants (complex integration pattern) in comparison with Agrobacterium-mediated transformation.

This indicates that the quality of transformants obtained by particle bombardment or other direct DNA delivery methods can be improved by DSB mediated insertion of sequences. This result is also confirmed by the following experiment: upon DSB mediated targeted sequence insertion of a 35S promoter, in absence of flanking sequences with homology to the target locus in the repair DNA, we observed that upon electroporation-mediated delivery of repair DNA, only a minority of the TSI events did contain additional non-targeted insertions of 35S promoter (2 TSI events out of 16 analyzed TSI events show additional at random insertion(s) of the 35S promoter). In contrast random insertion of the 35S promoter was considerably higher in TSI events obtained by Agrobacterium mediated delivery of the 35S promoter (17 out 22 analyzed TSI events showed additional at random insertion(s) of the 35S promoter).

Example 6: Media composition

Mahq1VII: N6 medium (Chu *et al.* 1975) supplemented with 100mg/L casein hydrolysate, 6 mM L-proline, 0.5g/L 2-(N-morpholino)ethanesulfonic acid (MES), 0.2M mannitol, 0.2M sorbitol, 2% sucrose, 1mg/L 2,4-dichlorophenoxy acetic acid (2,4-D), adjusted to pH5.8, solidified with 2,5 g/L Gelrite®.

Mhi1VII: N6 medium (Chu *et al.* 1975) supplemented with 0.5g/L 2-(N-morpholino)ethanesulfonic acid (MES), 0.2M mannitol, 2% sucrose, 1mg/L 2,4-dichlorophenoxy acetic acid (2,4-D), adjusted to pH5.8 solidified with 2,5 g/L Gelrite®.

Mh1VII: idem to Mhi1VII substrate but without 0.2 M mannitol.

Ahx1.5VIIino1000: MS salts, supplemented with 1000mg/L myo-inositol, 0.1 mg/L thiamine-HCl, 0.5mg/L nicotinic acid, 0.5mg/L pyridoxine-HCl, 0.5g/L MES, 30g/L sucrose, 10g/L glucose, 1.5mg/L 2,4-D, adjusted to pH 5.8 solidified with 2,5 g/L Gelrite®.

LSIDhy1.5: MS salts supplemented with 0.5mg/L nicotinic acid, 0.5mg/L pyridoxine-HCl, 1mg/L thiamine-HCl, 100mg/L myo-inositol, 6mM L-proline, 0.5g/L MES, 20g/L sucrose, 10g/L glucose, 1.5mg/L 2.4-D, adjusted to pH 5.2.

N6M: macro elements: 2830mg/L KNO₃; 433mg/L (NH₄)₂SO₄; 166mg/L CaCl₂·2H₂O; 250 mg/L MgSO₄·7H₂O; 400mg/L KH₂PO₄; 37.3mg/L Na₂EDTA; 27.3mg/L FeSO₄·7H₂O, MS micro elements, 500mg/L Bactotrypton, 0.5g/L MES, 1mg/L thiamin-HCl, 0.5mg/L nicotinic acid; 0.5mg/L pyridoxin-HCl, 2mg/L glycin, 100mg/L myo-inositol, 3% sucrose, 0.5mg/L 2.4-D, adjusted to pH5.8.

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Throughout this specification and the claims which follow, unless the context requires otherwise, the word “comprise”, and variations such as “comprises” and “comprising”, will be understood to imply the inclusion of a stated integer or step or group of integers or steps but not the exclusion of any other integer or step or group of integers or steps.

5

The reference in this specification to any prior publication (or information derived from it), or to any matter which is known, is not, and should not be taken as an acknowledgment or admission or any form of suggestion that that prior publication (or information derived from it) or known matter forms part of the common general knowledge in the field of endeavour to which this specification relates.

10

THE CLAIMS DEFINING THE INVENTION ARE AS FOLLOWS:

1. An isolated DNA fragment comprising the nucleotide sequence of SEQ ID No 4 or any nucleotide sequence obtained by replacing one to twenty nucleotides of SEQ ID No. 4
5 for any of the alternatives provided in the nucleotide sequence of SEQ ID No 3, in such a way that the GC content of said nucleotide sequence is about 50 to about 60%, provided that
 - a) if the nucleotide sequence from position 121 to position 123 is GAG then the nucleotide sequence from position 124 to 126 is not CAA;
 - 10 b) if the nucleotide sequence from position 253 to position 255 is GAC then the nucleotide sequence from position 256 to 258 is not CAA;
 - c) if the nucleotide sequence from position 277 to position 279 is CAT then the nucleotide sequence from position 280 to 282 is not AAA;
 - d) if the nucleotide sequence from position 340 to position 342 is AAG then the
15 nucleotide sequence from position 343 to position 345 is not CAT;
 - e) if the nucleotide sequence from position 490 to position 492 is TCA then the nucleotide sequence from position 493 to position 495 is not ACC;
 - f) if the nucleotide sequence from position 499 to position 501 is AAA then the nucleotide sequence from position 502 to 504 is not TCA or AGC;
 - 20 g) if the nucleotide sequence from position 517 to position 519 is ACC then the nucleotide sequence from position 520 to position 522 is not CAA simultaneous with the nucleotide sequence from position 523 to 525 being TCN;
 - h) if the nucleotide sequence from position 661 to position 663 is ATT then the nucleotide sequence from position 664 to position 666 is not AAA;
 - 25 i) the codons from the nucleotide sequence from position 7 to position 15 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven contiguous nucleotides from the group of G or C;
 - j) the codons from the nucleotide sequence from position 61 to position 69 are
30 chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven contiguous nucleotides from the group of G

or C;

k) the codons from the nucleotide sequence from position 130 to position 138 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven contiguous nucleotides from the group of G

5 or C;

l) the codons from the nucleotide sequence from position 268 to position 279 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven contiguous nucleotides from the group of G or C;

10 m) the codons from the nucleotide sequence from position 322 to position 333 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven contiguous nucleotides from the group of G or C;

15 n) the codons from the nucleotide sequence from position 460 to position 468 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of seven contiguous nucleotides from the group of G or C;

20 o) the codons from the nucleotide sequence from position 13 to position 27 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

25 p) the codons from the nucleotide sequence from position 37 to position 48 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

q) the codons from the nucleotide sequence from position 184 to position 192 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

30 r) the codons from the nucleotide sequence from position 214 to position 219 are chosen according to the choices provided in such a way that the resulting nucleotide

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sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

5 s) the codons from the nucleotide sequence from position 277 to position 285 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

10 t) the codons from the nucleotide sequence from position 388 to position 396 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

u) the codons from the nucleotide sequence from position 466 to position 474 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

15 v) the codons from the nucleotide sequence from position 484 to position 489 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

20 w) the codons from the nucleotide sequence from position 571 to position 576 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

25 x) the codons from the nucleotide sequence from position 598 to position 603 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

30 y) the codons from the nucleotide sequence from position 604 to position 609 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

z) the codons from the nucleotide sequence from position 613 to position 621 are

chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

5 aa) the codons from the nucleotide sequence from position 646 to position 651 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T;

10 bb) the codons from the nucleotide sequence from position 661 to position 666 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T; and

15 cc) the codons from the nucleotide sequence from position 706 to position 714 are chosen according to the choices provided in such a way that the resulting nucleotide sequence does not comprise a stretch of five contiguous nucleotides from the group of A or T.

2. An isolated DNA fragment according to claim 1, characterized in that it contains a nucleotide sequence differing from the nucleotide sequence of SEQ ID No 4 in only one position.

20

3. An isolated DNA fragment according to claim 1, characterized in that it contains a nucleotide sequence differing from the nucleotide sequence of SEQ ID No 4 in only ten positions.

25 4. An isolated DNA fragment comprising the nucleotide sequence of SEQ ID No 4.

5. A chimeric gene comprising the isolated DNA fragment according to any one of claims 1 to 4 operably linked to a plant-expressible promoter.

30 6. Use of a chimeric gene according to claim 5 to insert a foreign DNA into an I-SceI recognition site in the genome of a plant.

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7. A method for introducing a foreign DNA of interest into a preselected site of a genome of a plant cell comprising the steps of
- a) inducing a double stranded DNA break at the preselected site in the genome of the cell;
 - b) introducing the foreign DNA of interest into the plant cell; characterized in that the double stranded DNA break is introduced by an I-Sce I endonuclease encoded by an isolated DNA fragment according to any one of claims 1-4.
8. The method of claim 7, characterized in that said endonuclease comprises a nuclear localization signal.
9. The method of claim 7 or 8, wherein the foreign DNA of interest is introduced into said plant cell by direct DNA transfer.
10. The method of any one of claims 7 to 9, wherein said direct DNA transfer is accomplished by bombardment of microprojectiles coated with the foreign DNA of interest.
11. The method of any one of claims 7 to 10, wherein said foreign DNA of interest is flanked by a DNA region having at least 80% sequence identity to a DNA region flanking the preselected site.
12. The method of any one of claims 7 to 11, wherein said double stranded DNA break is induced by introduction of an I-SceI encoding gene.
13. The method of any one of claims 7 to 12, whereby the plant cell is a maize cell.
14. The method of claim 13, wherein the maize cell is comprised within a cell suspension.

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15. The method of any one of claims 7 to 14, whereby said plant cell is incubated in a plant phenolic compound prior to step a).

16. The method of claim 15, wherein said plant phenolic compound is acetosyringone.

5

17. The isolated DNA fragment of claim 1, the chimeric gene of claim 5 or a use thereof, or the method of claim 7, substantially as herein described with reference to the Examples and the accompanying drawing.

1/1

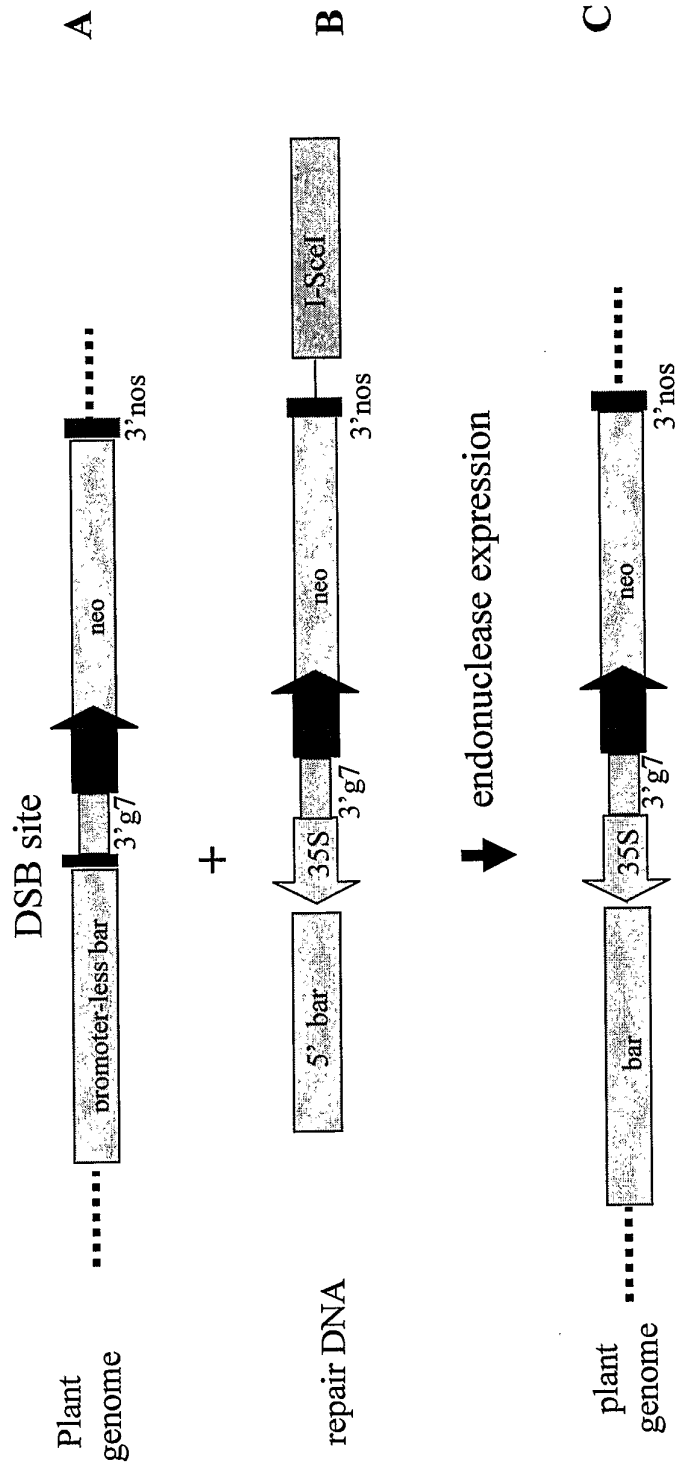


Figure 1

Thr Phe Glu Glu Val Glu Tyr Leu Val Lys Gly Leu Arg Asn Lys Phe
 180 185 190

Gln Leu Asn Cys Tyr Val Lys Ile Asn Lys Asn Lys Pro Ile Ile Tyr
 195 200 205

Ile Asp Ser Met Ser Tyr Leu Ile Phe Tyr Asn Leu Ile Lys Pro Tyr
 210 215 220

Leu Ile Pro Gln Met Met Tyr Lys Leu Pro Asn Thr Ile Ser Ser Glu
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garcarttyg argcnggnat hggncnath ctngnggayg cntayathcg ntcncnggay 180
garggnaara cntaytgyat gcarttygar tggaaraaya argcntayat ggaycaygtm 240
tgyctnctnt aygaycartg ggtncntnctn ccncncaya araargarcg ngtnaaycay 300
ctnggnaayc tngtnathac ntgggngcn caracnttya arcaycargc nttyaayaar 360
ctngcnaayc tnttyathgt naayaayaar aaracnathc cnaayaayct ngtngaraay 420
tayctnacnc cnatgtcnc ntngntaytg ttyatggayg ayggnggnaa rtgggaytay 480
aayaaraayt cnacnaayaa rctnathgtm ctnaayacnc artcnyyac nttygargar 540
gtngartayc tngtnaargg nctncgnaay aarttyarc tnaaytgyta ygtnaarath 600
aayaaraaya arccnathat htayathgay tcnatgtcnc ayctnathht ytayaayctn 660
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 garggmaara cytactgyat gcagttcgar tggaaraaca argcytacat ggaycaygts 240
 tgyctstctst acgaycartg ggtsctstcm cchchcaya araargarcg sgtsaaccay 300
 ctsggmaacc tsqtsatyac ytgggmgcy caracyttca arcaycargc yttcaacaar 360
 ctsgcsaacc tsttcatyct saacaacaar aaracyatyc chaacaacct sgtsgaraac 420
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 aacaaraact cmacyaaciaa rctmatygtc ctsaacacyc artcmttcac yttcgargar 540
 gtsgartacc tsqtsaargg mctscgsaac aarttccarc tsaactgyta cgtsaagaty 600
 aacaaraaca arccyatyat ctacatygay tcmatgtcmt acctsatytt ctacaaccts 660
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gaaggcaaga cctactgcat gcagttcgag tggagaaca aggcctacat ggaccacgtg     240
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aacaagaaca agcctatcat ctacatcgac agcatgagct acctgatctt ctacaacctg     660
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