Methylation of chloroplast DNA does not affect viability and maternal inheritance in tobacco and may provide a strategy towards transgene containment

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Abstract We report the integration of a type II restriction-methylase, mFokI, into the tobacco chloroplast genome and we demonstrate that the introduced enzyme effectively directs the methylation of its target sequence in vivo and does not affect maternal inheritance. We further report the transformation of tobacco with an E. coli dcm methylase targeted to plastids and we demonstrate efficient cytosine methylation of the plastid genome. Both adenosine methylation of FokI sites and cytosine methylation of dcm sites appeared phenotypically neutral. The ability to tolerate such plastid genome methylation is a pre-requisite for a proposed plant transgene containment system. In such a system, a chloroplast located, maternally inherited restriction methylase would provide protection from a nuclear-encoded, plastid targeted restriction endonuclease. As plastids are not paternally inherited in most crop species, pollen from such plants would carry the endonuclease transgene but not the corresponding methylase; the consequence of this should be containment of all nuclear transgenes, as pollination will only be viable in crosses to the appropriate transplastomic maternal background.

Keywords Chloroplast transformation · DNA methylation · Gene containment · Transplastomic

Introduction

The presence of low levels of methyl cytosine and adenosine within plastid DNA of liverwort (Takio et al. 1994), pea (Ohta et al. 1991), sycamore (Ngernprasirtsiri et al. 1988), maize (Gauly and Kössel 1989), and tomato (Kobayashi et al. 1990) have been reported, and an association with plant development has been proposed (Kobayashi et al. 1990). However, other groups have not always been able to duplicate such results (Marano and Carrillo 1991) and it has not been possible to detect reported methylation using bisulfite sequencing (Fojtová et al. 2001). This has led to the proposal that much of the reported methylation may be an artifact due to the use of inappropriate restriction enzymes for its detection, primarily EcoRII (Fojtová et al. 2001).

An exception to this is the chloroplast DNA of the unicellular green alga, Chlamydomonas reinhardtii, where extensive cytosine methylation is found (Sager 1972). There are two Chlamydomonas mating types, and hypermethylation of chloroplast DNA is associated with the maternal mating type (Sager and Ramani 1973; Burton et al. 1979). Under normal conditions, progeny inherit chloroplast DNA from the mt+ parent and that from the paternal mating type (mt−) is lost. Methylation of the chloroplast DNA is directed by a nuclear encoded, chloroplast targeted methylase, and constitutive expression of this gene in transgenic mt− cells leads to methylation of the paternal chloroplast genome and a high frequency of chloroplasts being inherited from the paternal parent (Nishiyama et al. 2004). It was originally proposed that this uniparental inheritance could be brought about by a restriction/modification system acting to protect maternal and to digest non-methylated paternal chloroplast DNA (Sager and Lane 1972).
However, more recent studies demonstrate that methylation is not necessary for protection from nucleases and support a model in which it promotes differential chloroplast DNA replication in germinating zygotes (Umen and Goodenough 2001).

In most crop plants, plastids are only inherited maternally; this is primarily a result of exclusion from the pollen generative cell, although examples of rare paternal transmission have been reported (Ruf et al. 2007; Svab and Maliga 2007). We postulated that bacterial restriction/methylation systems could be adapted and used to engineer transgenic plants in order to provide a mechanism for preventing transgenes from outcrossing to non-transgenic crops or wild relatives. Restriction-modification systems are believed to have evolved as a natural defence mechanism to protect bacteria against invading bacteriophages. The restriction endonuclease recognises short, specific nucleotide sequences in foreign DNA entering the bacterial cell and cuts at or near the recognised site. At the same time, the corresponding restriction methylase provides protection for the host’s DNA by adding a methyl group to the same target sequence. Three classes of restriction/modification systems are recognised in bacteria but only the type II system has the sequence specific methylation and sequence specific cleavage activities as separate enzyme functions (reviewed in Murray 2000; Bickle and Kruger 1993). Such systems have the potential to be modified to provide a method for transgene containment in plants. If a gene encoding a bacterial restriction endonuclease were to be equipped with a chloroplast import sequence and used for plant transformation, then the endonuclease would be anticipated to cleave the plastid genome at its target sites—destroying the chloroplast and leading to cell death. However, integration of a sequence specific methylase transgene into the plastid genome should result in methylation at target sites, conferring resistance to the imported nuclease, provided that all sites are methylated and that methylation of plastid DNA is tolerated by the plant. The potential number of such containment systems is theoretically in the hundreds, only being limited by the number of restriction endonuclease/methylase pairs available with recognition sites within the chloroplast genome.

The FokI restriction/modification system is the best characterised member of a small subset of the type II group in which the endonuclease activity is conferred by a monomer rather than two polypeptides acting as a homodimer (Kita et al. 1989). We anticipated that a monomer might be more readily taken up by the chloroplast import apparatus and for this reason the FokI system was chosen for engineering for plant expression. The FokI recognition sequence is GGATG which occurs 237 times in the tobacco chloroplast genome. Protection from digestion by FokI endonuclease (RFokI) is conferred by the FokI methylase (MFokI) which methylates the central adenosine within the recognition sequence (Sugisaki et al. 1989). As a first step in the creation of an outcross isolation system, we report the generation of homoplasmic plants carrying the MFokI transgene and demonstrate methylation at most, but not all, of the cognate sites. No deleterious effects on plant growth or plastid inheritance were observed.

To test whether cytosine methylation of the plastid genome could also be tolerated by the plants we transformed wild type plants with an E. coli dcm methylase transgene (Marinus and Morris 1973) which we equipped with a plastid import leader sequence. Extensive methylation of target sites was observed and again, no obvious deleterious effects on plant growth were seen. The demonstration of the successful insertion of a restriction methylase into the chloroplast genome and the ability of transgenic plants to tolerate extensive methylation of both adenosines and cytosines within their plastid genome is a critical first step in engineering restriction enzyme based containment systems.

Materials and methods

Plant material

Tobacco (N. tabacum cv. Petit Havana) plants were grown in MSR3 media (Waters et al. 2004). Transplastomic lines were rooted and propagated on MSR3 supplemented with 500 μg ml⁻¹ of spectinomycin. Nuclear transformants were grown on the same media supplemented with 50 μg ml⁻¹ kanamycin.

Construction of the chloroplast transformation vector

The plastid transformation vector pFaadMe2t was based upon pFaadAI ΔSN, kindly provided by Eibl et al. (1999). The pFaadAI ΔSN vector includes a sequence corresponding to nucleotides 114,279–116,171 of the tobacco plastid genome for facilitating integration through homologous recombination, an aadA resistance marker under the control of the 16S-rDNA promoter, a 25 bp ribosome binding site synthesised to match the corresponding sequence of the tobacco rbcL gene, and the 3'-end (450 bp) of the C. reinhardtii rbcL gene. In order to create pFaadMet2 the mFokI gene was amplified by PCR from the bacterium F. okeanokoites using the oligonucleotides 5'-MFokI CCACCGCGAGGAGGTAAATGAGATTATTG and 3'-MFokI GGAGCATGCAGATTATTGGGAGCAAG and the amplified product ligated into the pTag PCR cloning vector (R&D Systems, UK). The resulting plasmid was subsequently digested with Sphi to
release a fragment encoding the methylase as a 1.9 kb fragment which was then ligated into the \textit{SphI} site of \textit{pFaadASN} between the \textit{aadA} gene and the 3' end of \textit{rbcL} to create \textit{pFaadMet2} (Fig. 1a). \textit{E. coli} clones were functionally selected through the ability of plasmid DNA to withstand digestion with \textit{FokI} endonuclease.

**Generation of transplastomic plants**

Leaves were placed with abaxial side up on RMOP medium (Svab et al. 1990) in a petri dish. Preparation of and bombardment with particles was performed as described by Bock (1998) except gold particles (0.6 \( \mu \)m) were prepared instead of tungsten. Biolistics was carried out using the DuPont PDS1000He biolistic gun and 1,100 psi rupture disks (Bio-Rad). After bombardment the leaves were incubated at 25°C and 1 to 2 days later the leaves were cut into sections (5 mm \( \times \) 5 mm) and transferred to RMOP medium containing 500 \( \mu \)g ml\(^{-1}\) of spectinomycin dyhydrochlororide. Green calli that formed on leaf segments were subcultured onto the same selective medium. Shoots growing leaves were then re-callused a second time and subsequent shoots rooted on MS medium.

**Construction of \textit{dcm} methylase binary vectors and plant transformation**

Plasmid \textit{pBSRBCS} contains the petunia SSU611 ribulose bisphosphate carboxylase small subunit (\textit{rbcS}) chloroplast targeting sequence (Dean et al. 1987) cloned into \textit{pBlue-script II SK+} (Stratagene, UK). This sequence is derived from plasmid \textit{pDHERYB1} (Fray et al. 1999) and contains \textit{NeoI} and \textit{SphI} sites overlapping the initiating ATG codon and the cleavage site of the encoded chloroplast transit peptide fragment, respectively.

Genes encoding proteins to be targeted to chloroplasts can be cloned into \textit{pBSRBCS} using the \textit{SphI} site for generating the ATG codon of the desired gene in frame with the \textit{RBCS} sequence.

Plasmid \textit{pBC35dcm} was designed to direct the constitutive expression of a plastid targeted \textit{E. coli} \textit{Dcm} methylase in \textit{planta}. The \textit{dcm} gene was PCR-amplified from the \textit{E. coli} strain DH5\( \alpha \) with oligonucleotides \textit{dcmF} (TGGCATGCAGGAATAATATGCAGTAAC) and \textit{dcmR} (AGATCTAGATTATCGTGACTGACCAGCTGTTG) introducing an \textit{SphI} site overlapping the ATG codon at the 5' end and a \textit{BamHI} site at the 3' end. The amplified fragment was cloned into the \textit{SphI} and \textit{BamHI} sites of the \textit{pBSRBCS}. The \textit{RBCS-dcm} cassette was excised as a \textit{KpnI/XbaI} fragment and ligated to \textit{pBC35} cut with the same enzymes to give the plasmid \textit{pBC35dcm}. \textit{pBC35} is a \textit{pBIN19} binary vector derivative (Bevan 1984), into which the Cauliflower Mosaic Virus (CaMV) 35S promoter, multiple cloning site and CaMV 35S terminator cassette PCR amplified from \textit{pDH51} (Pietrzak et al. 1986), was cloned between the \textit{EcoRI} and \textit{HindIII} sites (R.G. Fray, unpublished).

Binary vectors were transferred to \textit{Agrobacterium tumefaciens} strain LBA4404 and used to transform sterile tobacco leaves as previously described (Fray et al. 1999).

**DNA blot analysis**

About 50 \( \mu \)g (or 1.5 \( \mu \)g for the analysis of chloroplast DNA) of total cellular DNA was digested with appropriate restriction enzymes, electrophoresed on 0.8% agarose gels and transferred to Hybond N+ (Amersham, UK) membranes using a standard neutral transfer procedure (Sambrook et al. 1989). DNA probes were random prime labelled with [\( x \)-32P]dCTP using a RediprimeTM II (Amersham, UK) kit. Probes were hybridised to the filters in: 10\( \times \) Denhardt's, 1% SDS, 5\( \times \) SSPE, 1 mg ml\(^{-1}\) sonicated salmon sperm DNA at 65°C. Membrane washings were carried out with 2\( \times \), 1\( \times \) and 0.1\( \times \) SSPE, 0.1% SDS at 65°C, 10 min each. Signals were detected using autoradiographic film (KODAK X-OMAT AR). All probes were cloned, fully sequenced and gel purified prior to use. Homoplasomy was confirmed using a 937 bp \textit{ndhF} probe equivalent to fragment 114,316–115,253 of the chloroplast genome. The same probe was used to assess susceptibility to \textit{FokI} digestion in the short single copy (SSC) sequence. For further cpDNA methylation analysis of \textit{FokI} sites in

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**Fig. 1** Chloroplast transformation with the \textit{FokI} methylase. The transformation vector \textit{pFaadMet2} targets transgene insertion to the small single copy region at a site proximal to the \textit{ndhF} gene (a). Insertion at this site results in an increase in size of 3,278 bp for the \textit{BglII} fragment containing the \textit{ndhF} probe sequence (b). Following Southern analysis, the absence of the smaller band derived from the wild type (WT) plastome indicates that all six transplastomic plants are homoplastic (lanes 1–6)
the large single copy (LSC), a 1,268 bp fragment of the accD gene corresponding to nucleotides 59,057–60,325 of the chloroplast genome was used. To assay FokI sites in the inverted repeat sequences (IR), probe chIRE (nucleotides 95,327–96,322 IRB, 13,729–146,304 IRA) was used.

For analysis of sites affected by Dcm methylation, four fragments were generated to probe ScrFI cut cpDNA. Two fragments were used to assay ScrFI sites in the LSC–a 800 bp rbcL fragment (nucleotides 57,730–58,531) and a 1,066 bp chlA fragment (nucleotides 25,042–26,108 of the chloroplast genome). A 995 bp chlB probe (nucleotides 27,148–28,143) was used to assay ScrFI sites in the inverted repeats.

**Results**

Generation and analysis of transplastomic mFokI plants

Plastid transformation experiments with pFaadMet2 resulted in several spectinomycin resistant lines. After further rounds of selection, the homoplastic state was confirmed for six of these by Southern analysis (Fig. 1b). MfokI methylates the sequence GGATG, which occurs 237 times in the plastid genome. To test for the presence of such methylation, total DNA from wild type and transplastomic plants was cut with BglII, and the resulting fragments analysed by Southern blotting. Probes were chosen such that multiple FokI sites could be assayed in the small single copy (SSC), large single copy (LSC) and inverted repeat (IR) regions of the plastid genome.

Southern analysis using the ndhF probe adjacent to the transgene insertion site in the SSC, detects hybridizing fragments of 4,746 bp and 8,024 bp in the wild type and transplastomic plants, respectively following BglII digestion alone (Figs. 1b, 2a). There are four FokI sites within the wild type BglII fragment and a further two within the transgene sequence. Digestion with both BglII and FokI gives the expected hybridizing bands of 2,531 bp and 1,255 bp for wild type. However, in the case of the transplastomic plants, digestion with both enzymes gives a single band indistinguishable from that produced by a BglII digestion alone, indicating that all six FokI recognition sites are protected from FokI digestion in the transplastomic DNA (Fig. 2a). Using the accD probe, susceptibility to digestion of two FokI sites within the LSC was assayed. In both wild type and transplastomic plants, digestion with BglII results in a hybridizing fragment of 2,666 bp. Upon digestion with both BglII and FokI, the two predicted hybridizing bands of 1,287 bp and 578 bp are seen in the digestion of the wild type DNA, but for the transplastomic DNA, the hybridizing band remains 2,666 bp in size, indicating that both FokI sites are protected (Fig. 2b). Using probe chIRE from the IR, a single hybridizing band of 1,685 bp is obtained following BglII digestion of both wild type and transplastomic DNA. Following digestion with both BglII and FokI, additional cleavage at the three FokI sites should give hybridizing bands of 673 bp, 414 bp and 223 bp. In the wild type, the double digest gave a single band of 673 bp, consistent with complete digestion by these enzymes (the two smaller bands being run off the end of the gel) (Fig. 2c). When the transplastomic DNA is digested with both enzymes, bands of 1,685 bp (no FokI digestion) and 673 bp (full digestion) are seen. A number of additional digestion products are evident and their derivation can be explained as follows: 1,310 bp (A + B + C), 1,271 bp (B + C + D), 1,048 bp (C + D), 896 bp (B + C) B BglII, F FokI, WT wild type, T transplastomic

**Fig. 2** Southern analysis indicates that FokI sites are protected from digestion to varying degrees in transplastomic plants. a Digestion with BglII gives ndhF hybridizing fragments of 4,746 bp and 8,024 bp for the wild type (WT) and transplastomic (T) plants respectively. Digestion with both BglII and FokI gives bands of 2,531 and 1,255 bp for wild type but protection of all six FokI sites in the transplastomic plant means that the 8,024 bp fragment is not digested further. b FokI sites within the LSC were assayed using the accD probe indicated. In both wild type and transplastomic plants, digestion with BglII gives a fragment of 2,666 bp. Digestion with both BglII and FokI, gives the two predicted bands of 1,287 and 578 bp for wild type, but protection of the two FokI sites leaves the 2,666 bp fragment unaltered in the transplastomic plant. c FokI sites within the IR were tested using the chIRE probe indicated. Digestion of both wild type and transplastomic cpDNA with BglII gives a single band of 1,685 bp. Digestion of wild type with both BglII and FokI gives a band of 673 (the smaller 414 bp and 223 bp fragments have been run out of the gel). Digestion of transplastomic DNA with both enzymes gives bands of 1,685 (no FokI digestion) and 673 bp (full digestion).

Transplastomic plants expressing the mFokI methylase grew well with no obvious deleterious phenotypes observed as a result of MfokI induced adenosine cpDNA methylation. Reciprocal crosses were performed between transplastomic FaadMet2 and wild type plants and seeds were sown on MSR3 media supplemented with spectinomycin. 100% of germinating seeds were spectinomycin resistant when FaadMet2 was the maternal parent (Fig. 3a) and no spectinomycin resistant progeny were seen when it was the pollen donor (Fig. 3b). Two independent FaadMet2 lines were tested in this way and more than 10,000 seeds assayed in total. Complete methylation of FokI sites hybridizing to the SSC accD probe was confirmed for three of the spectinomycin resistant progeny (not shown).
Generation and analysis of dcm transgenic plants

To test the effect of cytosine methylation, plants were transformed with a gene encoding the E. coli Dcm methylase equipped with a plastid import sequence. The dcm gene product methylates cytosines within the target sequence CCWGG (W = A or T). There are 171 such sites within the tobacco chloroplast genome. Transformed plants expressing the dcm methylase construct were analysed for the presence of cpDNA methylation by digestion with ScrFI. This enzyme cuts at the sequence CCNGG but is sensitive to cytosine methylation. Thus, in the presence of full Dcm methylation, CCCGG and CCGGG will remain susceptible to digestion. An rbcL probe (in the LSC) was chosen such that in the absence of methylation a fragment of 797 bp should result from digestion with ScrFI, whilst protection of a CCWGG site will result in a hybridising fragment of 1,810 bp (Fig. 4a). Extensive or complete methylation was seen at the assayed sites. A further eight sites were assayed in the LSC using chlA as probe. In the absence of Dcm methylation, digestion of wild type DNA with ScrFI gives chlA hybridizing bands of 505, 461, 360 as well as a 27 bp fragment (not detected). Protection of all eight CCWGG sites in Dcm methylation lines should result in two hybridizing fragments of 874 and 2,738 bp. The presence of additional intermediate bands may indicate partial methylation.

No growth defects were seen as a result of this cytosine methylation.

Discussion

We have shown that bacterial restriction methylases can be expressed in the plastid environment and despite extensive adenosine or cytosine methylation of the chloroplast genome, plants appeared phenotypically normal. For an outbreeding suppression system based upon bacterial restriction/modification enzymes to work effectively, the entire target sites in the plastid genome should be protected from the endonuclease activity. In the mFokI transplastomic plants, apparently full target methylation was seen at eight sites within the assayed BglII fragments of SSC and LSC. However, incomplete methylation occurred at the three sites within the BglII fragment from the inverted repeats. This incomplete methylation in the IR may simply be a result of stochometry (there are two copies of the IR sequence for every one copy of the LSC sequence), or it may be an indication that sequences within the IR are less accessible to the methylase—perhaps due to their relative proximity to the IR sequences involved in initiating DNA replication (Kunnimalaiyaan et al. 1997).

For the transgenic plants expressing a nuclear encoded, plastid targeted Dcm methylase, a total of 23 Dcm target sites were tested by ScrFI digestion. The presence of the predicted higher molecular weight bands from the transgenic plants indicates that all the assayed sites are normally methylated. However, the faint bands of intermediate size seen when using the IR and one of the LSC probes may indicate that methylation is incomplete in some DNA molecules (Fig. 4).
For an out crossing inhibition scheme based upon bacterial restriction systems to work effectively, full methylation of all target sites will be required. Although extensive methylation was observed in both the FokI and Dcm plants, some sites were not fully protected. Thus, it may be necessary to test a number of alternative methylase
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