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Frequency and distance of transposition of a modified *Dissociation* element in transgenic tobacco

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Effective transposon tagging with the Ac/Ds system in heterologous plant species relies on the accomplishment of a potentially high transposon-induced mutation frequency. The primary parameters that determine the mutation frequency include the transposition frequency and the transposition distance. In addition, the development of a generally applicable transposon tagging strategy requires predictable transposition behaviour. We systematically analysed Ds transposition frequencies and Ds transposition distances in tobacco. An artificial Ds element was engineered with reporter genes that allowed transposon excision and integration to be monitored visually. To analyse the variability of Ds transposition between different tobacco lines, eight single copy T-DNA transformants were selected. For *trans*-activation of the Ds elements, different Ac lines were used carrying an unmodified Ac^+ element, an immobilized sAc element and a stable Ac element under the control of a heterologous chalcone synthase (*chsA*) promoter. With all Ac elements, each Ds line showed characteristic and heritable variegation patterns at the seedling level. Similar Ds line-specificity was observed for the frequency by which Dstranspositions were germinally transmitted, as well as for the distances of the Ds transposition frequencies (0.37%) and high incidences of independent transposition (83%). The majority of these Ds elements (58%) transposed to genetically closed linked sites (≤ 10 cM).

Keywords: heterologous transposon tagging; Ac/Ds system; visual transposition assays; position effects; Nicotiana tabacum

Introduction

The Activator (Ac)/Dissociation (Ds) transposon system from maize (McClintock, 1951) provides a powerful means for gene tagging in heterologous plant species. The autonomous Ac element consists of a transposase gene and discrete termini that are required for transposition to occur (Kunze et al., 1987; Coupland et al., 1988, 1989). Most Ds elements are Ac-derivatives with internal deletions in the transposase gene and, consequently, are non-autonomous. The Ac/Ds transposition behaviour in heterologous plant species has been shown to represent that in maize (Hehl and Baker, 1990; Jones et al., 1990). Transposition takes place non-replicatively, is dependent

*To whom correspondence should be addressed at the Sainsbury Laboratory, John Innes Centre, Colney Lane, Norwich NR4 7UH, UK (Fax: +44(0)1603 250024) of the transposase gene dosage, occurs predominantly to genetically linked sites and upon integration an 8 bp duplication of the target site is created. One advantage of transposon tagging over other approaches to isolate genes with unknown products, such as T-DNA tagging and mapbased cloning, is that no genetic complementation analysis is required: the authenticity of the insertion mutant can be verified by reversion analysis (Walbot, 1992). Unstable phenotypes caused by germinal or somatic reversion to wild-type present evidence that the gene is mutated by a transposon. Part of the gene flanking the transposon can readily be cloned by inverse PCR (Earp *et al.*, 1990) or plasmid rescue (Rommens *et al.*, 1992a) and subsequently, can directly be used for the isolation of the corresponding complementary DNA.

Heterologous Ac/Ds transposon tagging strategies have been developed in a variety of plant species (Haring *et*

al., 1991; Van der Biezen et al., 1994). The Ac components, functioning as transposase donors, were stabilized by deleting part of the cis-required sequences. The regulatory sequences of the transposase genes were often modified to increase the effectiveness of Ds transactivation. The constructs with the Ds components were equipped with selectable or detectable markers to monitor transposition in plants. By non-targeted transposon tagging, genes were tagged that corresponded to any phenotype that was correlated with a transposon insertion (Bancroft et al., 1993; Long et al., 1993; Chuck et al., 1993; Van der Biezen et al., 1996). The effectiveness of these experiments depended on the selection efficiency of plants that carried transposition events. By targeted transposon tagging, the plant crosses were designed to isolate one specific gene-of-interest (Jones et al., 1994; Whitham et al., 1994; James et al., 1995; Lawrence et al., 1995). The efficiency of this procedure depended on the transposon activity and the genetic distance of the transposons to the target genes. Close linkage between the original transposon location and the target gene increased the mutation frequency considerably.

With respect to the mutagenic potential of the Ac/Dssystem, we recently used a mathematical approach to dissect the transposition behaviour into its components (Van der Biezen et al., 1994). To predict statistically the transposon-induced mutation frequency in heterologous hosts, the contribution of the involved parameters was estimated. In this paper, we systematically investigated the two major parameters that determine the efficiency of transposon tagging, including the frequency of somatic and germinal Ds transposition and the Ds transposition distance. Also, the variation of these transposition parameters between different tobacco lines was analysed. To these ends, a T-DNA construct with an artificial Ds element was designed comprising plasmid rescue sequences and reporter genes that allowed excision and reinsertion to be visually monitored. Eight transgenic tobacco lines were selected harbouring single copy T-DNA insertions. Following trans-activation of Ds by three different Ac constructs, plants with unique, germinally transmitted stable transposed Ds elements were identified and the transposition distances determined by segregation analysis of the reporter genes. The Ds lines showed characteristic transposition behaviour for the timing and frequency of Ds excision during embryogenesis, for the germinal Ds transposition frequencies, and for the distance of Ds transpositions. Furthermore, seedling assays showed that Ds was much more active during embryogenesis than at mature stages. In mature plants the Ds transpositions took place late in development, resulting in low germinal transposition frequencies and high incidences of independent transposition events. The Ds elements had a strong preference for transposition to genetically closely linked sites.

Materials and methods

Recombinant cloning of Albatross

To construct an artificial Ds element with the short 5' and 3' termini of Ac (Behrens et al., 1984), PCR amplification was employed using the act3a/act3b primers (1-194) and the act5a/act5b primers (4234-4565). Ac elements contain imperfect terminal repeats (TIRs, ^T/_cAGGGATGAAA), while Ds elements have perfect TIRs (both terminal bases are Ts). To maintain the homology with natural Ds elements, the act5a primer was designed such that it changed the terminal C into a T. Both PCR fragments were cloned in pSU20 (Bartolomé et al., 1991) with the TIRs directed towards each other resulting in plasmid pEB10. The DNA sequence of the 0.5 kb Ds insert of pEB10 was verified by double-strand dideoxy automatic sequencing (Applied Biosystems Model 373A). The pSU20 vector was chosen for potential application of plasmid rescue of Ds-flanking plant DNA because of the small size (2.3 kb), the compatibility with other plasmids for persistence in bacterial hosts (p15A origin of replication), and the presence of the bacterial chloramphenicol resistance gene (CAT). The pEB10 plasmid was linearized by cutting between the TIRs and ligated between the CaMV-35S promoter and the TP:SPEC gene from pSLJ2524 (Scofield et al., 1994). Subsequently, the total fragment was inserted between the T-DNA borders of pMOG22 (Mogen, Leiden, Netherlands), from which the 35S:HPT gene was deleted first. Finally, the NPTII gene (McBride and Summerfelt, 1990) was inserted between pSU20 and the 5' Ds terminus. The total T-DNA construct was named Albatross (Fig. 1) and contained a Ds element of 5.3 kb that is of comparable size to Activator (Ac)elements (4.6 kb). Recombinant clones and binary vectors were constructed by standard procedures (Sambrook et al., 1989) in Escherichia coli strain JM101 and transferred to Agrobacterium tumefaciens strain LBA4404 (Hoekema et al., 1983). Details of the cloning procedures are available on request.

Plant materials, plant transformations and plant crosses

SLJ10512B tobacco plants carried a single T-DNA insertion with a stabilized *Activator* element (*sAc*) of which the 177 terminal bp at the 3' end had been deleted, a *GUS* reporter gene and a *NPTII* gene to select transformants (Scofield *et al.*, 1992). Plants with the *Ac* transposase construct driven by the flower-specific chalcone synthase A (*chsA*) promoter from *Petunia hybrida* (*chs:Ac*) were described by Rommens *et al.* (1992b), and the plants with the wild-type (unmodified) and transpositionally active *Ac* element (*Ac*⁺) were described by Haring *et al.* (1989).

Tobacco Petit Havana SR_1 leaf discs were transformed with the T-DNAs of *Albatross* and pSLJ4731 (Jones *et al.*, 1992) by *Agrobacterium*-mediated transformation



Fig. 1. Map of the Albatross T-DNA comprising an artificial Ds element of 5.3 kb (shown in black). The Ds element contained (1) the short 5' (331 bp) and 3' (194 bp) termini of Ac, (2) a bacterial chloramphenicol resistance gene (CAT) and a bacterial origin of replication (p15A) permitting plasmid rescue of flanking plant DNA, and (3) a neomycin phosphotransferase gene (NPTII) for kanamycin resistance that functioned as plant transformation marker and Ds insertion marker. NPTH was regulated by the promoter and the terminator of the mannopine synthase gene (mas5':NPTII:mas3'). To monitor transposon excision, Ds was cloned between the cauliflower mosaic virus CaMV 35S promoter and the spectinomycin resistance gene (SPEC). For cell-autonomous action, the SPEC gene was preceded by a chloroplast transit peptide sequence (TP) from P. hybrida and concluded by the nopaline synthase termination sequence (35S:TP:SPEC:nos3'). For transfer to the plant genome the transposon-construct was flanked by the left-(LB) and right borders (RB) of the T-DNA of Agrobacterium. The positions (kb) of the EcoRI (E) and HindIII (H) restriction enzyme recognition sites are shown. The annealing sites of the oligonucleotides used as primers in polymerase chain reactions (PCR) are presented as arrows above the construct. Hybridization probes used in Southern analyses are shown as boxes below the construct.

(Horsch *et al.*, 1985). Kanamycin-resistant (100 mg 1^{-1}) *Albatross* transformants were designated *Alb* followed by a number for each independent transformant and a letter to distinguish siblings.

All crosses were performed in the greenhouse using standard emasculation and pollination techniques. Only the results of crosses with plants that were homozygous for the respective T-DNAs are reported. The genotypes were determined by testing self progenies for segregation of kanamycin (Km) resistance in seedling assays. The F₁s were made by crossing the Ac parents (sAc, chs:Ac, Ac^+) with the *Albatross* lines as the pistillate parents. The F_1 plants were hemizygous for Ac and Ds containing T-DNAs and were outcrossed to produce OC_1 progenies. Reciprocal outcrosses were made; however, generally the non-transgenic Petit Havana SR1 cultivar was used as the staminate parent. To analyse the inheritance of (independent) transposition events in the OC_1 , the position of each outcross on the specific branch, the specific inflorescence and the specific flower was recorded (Keller *et al.*, 1993). Finally, the selected OC_1 individuals were self-pollinated to produce OC_1S_1 families. Plants were grown in pots under standard Dutch greenhouse conditions (20°C, 60% humidity, 3 klx).

For aseptic assays, seeds were dipped in 70% ethanol for 30 s, soaked in 1% NaClO (w/v) for 20 min and rinsed in sterile water. Following this sterilisation procedure, 100–600 seedlings were plated in Petri dishes containing Murashige and Skoog (MS) medium pH 5.7 with 1.5% sucrose, 0.8% agar (Difco), spectinomycin (100 mg1⁻¹) and/or kanamycin (200 mg1⁻¹) (Duchefa) and placed in a climate cabin under standard conditions (16 h, 3 klx, 22°C). After 2–3 weeks the antibiotic resistant and sensitive seedlings were counted, and the selected seedlings subjected to GUS analysis and grown in soil to mature plants for genetic and molecular analysis.

Histochemical GUS analyses were done by placing the roots of seedlings in 1 mM 5-bromo-4-chloro-3-indolyl- β -D-glucuronide (X-gluc) in 50 mM NaHPO₄ pH 7. The seedlings were maintained at 100% humidity to prevent dehydration and placed at 25°C for 2–4 h. Seedlings containing the *GUS* gene showed blue coloration of the roots. Because seedlings showing no GUS activity were selected for further analysis, the absence of the *sAc* T-DNA (SLJ10512) was verified in leaf tissue extracts from greenhouse grown plants. First, fluorometric GUS analyses with 1 mM 4-methyl-umbelliferyl- β -D-glucuronide (MUG) were done (Jefferson *et al.*, 1987) and, secondly genomic DNA was subjected to PCR with *Ac*-specific primers (*ac6/ac7*).

Tobacco seedlings that were germinated on medium containing spectinomycin (Sp) developed bleached chloroplasts and, as a consequence, had white (W) cotyledons. However, when transformed with the SPEC gene (aadA), seedlings appeared fully green (FG) on this medium. For cell-autonomous action, SPEC was preceded by a chloroplast transit peptide sequence (TP) from P. hybrida (Scofield et al., 1994). The Ds element cloned between the promoter and TP:SPEC prevented transcription of this gene but expression was restored following excision of Ds. Somatic excisions resulted in variegation (V) patterns of Sp-resistant green spots on the Sp-sensitive white background of the cotyledons. The somatic excision patterns were classified according to the degree of green variegation: fully green seedlings were totally green (FG); highly variegated (HV) seedlings were almost totally green with some small white flecks on the surface lamina or in inner cell layers; medium variegated (MV) seedlings had 3-10 distinct green spots; and lowly variegated (LV) seedlings had 1-2 small green spots. Germinally transmitted Ds excisions resulted in seedlings carrying the event in all cells and hence were fully green (FG) when germinated in the presence of Sp.

In seedling assays, kanamycin (Km) resistance was indicative of the presence of the *Ds* element (*NPTII*) and Sp resistance was indicative of the presence of the

Albatross T-DNA (SPEC) from which a Ds element had been excised (empty donor site, EDS). In medium with Sp and Km, the presence of both markers resulted in fully green (FG) seedlings, the absence of both markers resulted in white (W) seedlings. Seedlings with only an EDS T-DNA and no Ds (Sp^R/Km^S) developed yellow (Y) cotyledons on Sp/Km medium. As control for the transposition reporter genes and the phenotypic seedling assays, various tobacco lines were used that carried none, either one, or both markers. Self seedlings of these transformants were germinated in the presence of both Sp and Km. Seedlings with the pSLJ4731 construct (SPEC and NPTII; Jones et al., 1992) were fully green (FG) (Sp^R/Km^R); pEB34 plants (pMOG22 with SPEC from pSLJ4731) were yellow (Y) (Sp^R/Km^S); Albatross and SLJ10512 lines were white (W.) (Sp^S/Km^R); and seedlings of the non-transgenic cultivar Petit Havana SR1 were also white (W) (Sp^{S}/Km^{S}) .

Calculation of the transposition frequencies and the transposition distance

Somatic excision frequencies in F₁ progenies (all hemizygous for both and Ac and Ds T-DNAs) were calculated by dividing the number of seedlings showing Sp resistance (FG or V) by the total number of seedlings. The number of FG seedlings in OC1 progenies was a measure for the germinal excision frequency and was based on the proportion that inherited the Ds T-DNA, i.e. half of the total population. Germinal transposition events (i.e. excision followed by reinsertion) resulted in FG OC1 seedlings when germinated on double-selective Sp/Km medium. However, FG seedlings (Sp^R/Km^R) were not necessarily the consequence of germinal transposition, because Km resistance could also be conferred by the sAc T-DNA (SLJ10512) which segregated in OC_1 progenies and, moreover, Sp resistance could also be the result of somatic excision. To select seedlings indisputably harbouring a germinally excised and reinserted Ds element, FG seedlings were selected that did not contain the SLJ10512 T-DNA (sAc:: NPTII:: GUS) by analysis of GUS activity. In this way, somatic excision was excluded and Km resistance could only be conferred by the NPTH gene within the Ds element. This approach had the concomitant advantage that stable insertions were selected which allowed reliable determination of the transposition distances in self progenies (OC_1S_1) . To avoid underestimation of the germinal transposition frequencies, selection against sAc required that this SLJ10512 T-DNA was unlinked (in repulsion) to the Albatross T-DNA. Segregation analysis of OC₁ progenies by Km and Sp/Km assays indeed indicated that all eight Albatross T-DNAs were unlinked to the sAc T-DNA (SLJ10512). For the calculation of the germinal transposition frequencies only half of the FG seedlings could unambiguously be recognized (1/2 without)sAc), and therefore, the proportion of FG/GUS⁻ seedlings was based on half of the number of seedlings that potentially could inherit a Ds element (i.e. half of the total seedlings), thus 1/4 of the total OC_1 population. This OC_1 population size was based on the number of variegated (V) seedlings which was 1/4 of the total number of OC_1 seedlings (see Results). Consequently, the germinal transposition frequency was calculated by (No of FG & $GUS^-)/(No \text{ of } V)$. In OC_1 progenies of Alb15, the sizes of the OC_1 populations were determined by counting all seedlings.

The distance of transposition was expressed as the recombination frequency between the Ds element and the original donor site, the Albatross T-DNA, converted to centiMorgans (cM). Recombination between the Ds element and the T-DNA resulted in separation of the respective NPTII (Km^R) and SPEC (Sp^R) markers which were scored in OC1S1 progenies. However, only half of the recombinants could be recognised on Sp/Km medium: those that contained the T-DNA and no Ds element $(Sp^R/$ Km^S) and, consequently, developed yellow (Y) cotyledons. The other class of recombinants only carried the Ds element and no T-DNA (Sp^S/Km^R), and hence developed white (W) cotyledons which could not be distinguished from non-transgenic white (W) seedlings (Sp^{S}/Km^{S}) . Therefore, the proportion of recombinants (Y) was based on half of the total OC_1S_1 population. When significant linkage was observed (P < 0.01 if $\chi^2 > 11.32$), the recombinant fractions (r) were calculated by (No of Y)/ $(\frac{1}{2} \times \text{total No})$. Subsequently, the genetic distances were converted to cM using Haldane's mapping function: $(-\ln(1-2r)/2) \times 100$.

DNA isolation and Southern analysis

DNA isolations and Southern hybridizations were done as described before (Van der Biezen *et al.*, 1995). Five radiolabelled probes were used: SPEC (*XhoI–Bam*HI, 1.1 kb) and 35S (*EcoRI–XhoI*; 1.3 kb) from pSLJ2524 (Jones *et al.* 1992), pSU20 (total plasmid, 2.3 kb; Bartolomé *et al.*, 1991), NPTII (*EcoRI*; 1.0 kb) from pCGN1547 (McBride and Summerfelt, 1990) and CHSA (*SaII–NcoI*; 2.4 kb) from VIP119 (Van der Meer *et al.*, 1990).

Polymerase chain reaction (PCR)

PCRs were done as described before (Van der Biezen et al., 1996). The primer sequences were: act5a 5'-GCA-GATCTTAGGGATGAAAGTAGGATGGG-3', act5b 5'-GCAAGCTTCTCGAGGGGAGAGAGGCAGAGCAGCG-3', act3a 5'-GCAAGCTTCCGAACAAAAACGGTCGG-3', act3b 5'-GCAAGCTTCCGAACAAAAATACCGGT-TCCCG-3', ac6 5'-CGCTTGTTCCATGATGACC-3', ac7 5'-GCTGAAGCCTCTTCTAGTCGG-3', 35s 5'-ATCTC-CACTGACGTAAGGGATGACG-3' and spec 5'-AGTT-GAGTCGATACTTCGGCGATCAC-3' (Isogen Bioscience Amsterdam).

Results

Plants with single copy Ds containing T-DNA insertions

The *Ds* element contained the kanamycin (Km) resistance gene *NPTII* for selection of T-DNA or transposon insertions, and was cloned between the *CaMV-35S* promoter and the structural part of the spectinomycin (Sp) resistance gene *TP:SPEC* (Fig. 1). The reporter genes allowed visual monitoring of *Ds* excision and integration in seedlings by simultaneous Sp/Km assays. First, the transposition capacity of the artificial Ds element was tested following *Agrobacterium*-mediated transfer of the *Albatross* (*Alb*) T-DNA to a transgenic tobacco line (SLJ10512B) that already contained a *sAc* element. Subsequently, the calluses and shoots were regenerated in the presence of Sp. Tissue that developed green spots and sectors (Sp^R) on the white background (Sp^S) indicated *sAc*-induced excision of Ds from the T-DNA (Fig. 2A).

To generate stable *Alb* lines, additional *Agrobacterium* transformations were performed using a non-transgenic



Fig. 2. Visual assays for monitoring Ds excision, Ds integration and the presence of sAc. (A) Regeneration assay for functional analysis of Ds transposition. Leafdiscs from a sAc-containing plant were transformed with the Albatross T-DNA by Agrobacterium and calluses and shoots were regenerated in spectinomycin (Sp) medium. Green variegation (Sp^R) indicated excision from the spectinomycin resistance gene (SPEC). (B) GUS analysis for the presence of the SLJ10512 constructs carrying sAc. The roots of Sp^R/Km^R OC1 seedlings carrying the T-DNA stained blue after treatment with X-gluc. (C-J) Somatic excision phenotypes of double hemizygous Ac/Ds F₁ seedlings derived from the eight Albatross lines crossed with plants with the immobile sAc construct and germinated on medium with Sp: (C) Alb1, (D) Alb2, (E) Alb3, (F) Alb10, (G) Alb15, (H) Alb17, (I) Alb19 and (J) Alb20. (K) V1 excision phenotype of double hemizygous F_1 seedlings derived from Alb2 and the chs:Ac line grown on Sp medium. (L) Distance of Ds transposition. The OC1S1 seedlings were germinated on Sp/Km medium and segregated for a transposed Ds element marked by NPTII for Km^{R} , and the donor site T-DNA marked by SPEC for Sp^R. The distance of transposition (cM) was based on the frequency of recombination between Ds and the T-DNA. The Sp^R/Km^R seedlings were fully green (FG), all Sp^S seedlings were white (W), and recombination between Ds and the T-DNA resulted in Sp^R/Km^S seedlings that were yellow (Y).

tobacco line. In subsequent self progenies of the primary transformants, the number and zygosity of Alb T-DNA inserts (NPTII) were determined by segregation analysis of Km resistance. Eight independent stable Alb transformants with single homozygous T-DNA loci were selected. Southern hybridizations were performed to analyse molecularly the integrity of the T-DNAs. Several enzyme (EcoRI, Hind III)/probe (35S, pSU20, NPTII, SPEC) combinations were used (Hind III/35S analysis shown in Fig. 3, other data not shown). All eight transformants harboured one intact T-DNA copy, whereas Alb1 and Alb15 contained an additional truncated left part of the T-DNA hybridizing to the 35S probe but not to the other probes. The eight single T-DNA copy Albatross transformants allowed subsequent systematic analysis of Ds transposition.

Somatic excision of Ds

For *trans*-activation of the *Ds* elements, the *Albatross* lines were crossed with plants that carried different *Ac* transposase constructs: (1) A stabilized *Ac* element (*sAc*, Scofield *et al.*, 1992), (2) an unmodified (wild-type) and transpositionally active *Ac* element (Ac^+ , Haring *et al.*, 1989), and (3) a stable *Ac* element in which the

transcription of the transposase gene was driven by a heterologous chalcone synthase promoter (chs:Ac, Rommens et al., 1992b). Three homozygous siblings of each Albatross line were crossed with plants carrying homozygous Ac constructs and the F_1 seeds (3-6 capsules per plant) germinated on Sp medium (Fig. 4). The variegation patterns of the seedlings were classified according to the degree of Sp resistance in fully green (FG), highly variegated (HV), medium variegated (MV) and lowly variegated (LV) classes (Fig. 2C-J). Two novel Ds variegation patterns were observed that were unique for plants with the chs:Ac construct. One pattern, designated V1, consisted of 1-3 large green spots of almost the size of the cotyledon (Fig. 2K). The second chs:Ac-induced variegation pattern (V2) consisted of many (10-20) homogeneously scattered small spots.

For each of the eight different *Albatross* lines, characteristic Sp variegation patterns were observed for a particular Ac construct. Although most classes (FG, HV, MV, LV) were represented in all progenies, the lines differed in the proportions of each class. Progeny seedlings from different siblings of a particular *Albatross* line showed remarkable consistency in the composition of the variegation classes. In addition, OC₁ progenies of



Fig. 3. Southern analyses of parental *Albatross* lines and their F_1 and OC_1 progenies carrying somatic and germinal *Ds* transpositions. Left panel: *Hin*dIII digested DNA of eight primary *Albatross* transformants hybridized with the radiolabelled 35S probe. NT was a non-transgenic plant. Differences in intensity of the bands were due to differences in DNA quantities. Right panel: *Eco*RI digested DNA isolated from *Alb* × *sAc* progenies and hybridized with the pSU20 probe. The seven F_1 plants showing somatic transposition (lanes 1–7) were derived from *Alb*1 (lanes 1–3) and *Alb*2 (lanes 4–7) and selected in Sp assays as variegated (V; lanes 1–5) or fully green (FG; lanes 6 and 7) seedlings. The positions of the *Ds* element at full donor sites (i.e. the T-DNA) for the two *Albatross* lines are indicated by triangles. Bands at other positions indicate transposed *Ds* elements. The five OC₁ plants (lanes 9–13) showing germinal transposition were derived from the *Alb*2 line and selected as FG seedlings in Sp/Km assays and lacked GUS activity (Sp^R/Km^R/GUS⁻). Bands at different positions indicate transposed *Ds* elements.

Frequency and distance of Ds transposition

PARENTAL LINES



Fig. 4. Crossing scheme used to analyse somatic and germinal Ds transposition frequencies and the distance of Ds transposition. Eight single-copy homozygous parental Albatross lines (P) were selected and crossed to lines that were homozygous for different Ac constructs. The progeny seedlings (F_1) were hemizygous for both the Ds and the Ac elements and, therefore, all could potentially be variegated when germinated on Sp medium. Following outcrossing to a non-transgenic line (NT), the progenies (OC_1) were grown on double selective Sp/Km medium and subjected to GUS analysis. Sp^R/Km^R seedlings were selected that did not contain the sAc construct (GUS⁻) and, hence, carried a germinally transmitted excised and reinserted Ds element. Subsequently, self progenies (OC_1S_1) were analysed for the segregation of the SPEC (Sp^R) and NPTII (Km^R) genes. The frequency of recombination between SPEC and NPTII represented the genetic distance between the empty donor site T-DNA (EDS) and Ds, respectively, and hence, represented the distance of transposition. FG = fully green, V = variegated, W = white, Y = yellow.

each *Albatross* line showed identical variegation patterns as the F_1 , demonstrating that the line specificities of the variegation patterns were heritable (see next section). The low internal variations in the proportion of the variegation classes per *Albatross* line allowed calculation of the mean proportions of each class which represented somatic excision frequencies (Fig. 5).

With *Alb*15 as the exception, the majority of the *Albatross* progenies from sAc and Ac^+ crosses consisted of FG and V (any variegated class) seedlings. Only small

fractions (0-11%) of the total seedlings were white (W), indicating that *trans*-activation of the *Ds* elements by these Ac constructs was generally very effective. Most of the Albatross progeny trans-activated by the sAc element consisted of seedlings of the MV class ($81\% \pm 3\%$). The sAc crosses with Alb2 and Alb15 were exceptions in that these progenies had a majority of HV (91%) and W seedlings (76%), respectively. Remarkably, the same phenomenon was observed when the Ac^+ element was used for trans-activation of Ds; however, the degree of variegation was higher. With Ac^+ , the majority of progenies consisted of HV seedlings $(50\% \pm 2\%)$, whereas Alb2 had mainly FG seedlings (91%) and Alb15 progenies mostly MV (45%) and LV seedlings (44%). The chs:Ac element induced variegation patterns unlike those caused by transposase genes that were transcribed by the original Ac promoter. Most F_1 progenies contained both chs: Ac-induced variegation classes V1 ($34\% \pm 6\%$) and V2 ($33\% \pm 5\%$). Also with this construct the progenies of Alb2 and Alb15 showed extreme high (100% V1) and low (0% V1 and 0% V2) levels of *trans*-activation, respectively. The variegation specificity of the Albatross lines was observed in F1 and OC_1 progenies with all three Ac constructs. The Alb2 line was shown to contain a Ds element that was most sensitive to *trans*-activation by any of the three elements. while the Alb15 line carried a Ds element that responded lowest to trans-activation. As concluded from the degree of variegation, the Ac^+ element was generally more effective in *trans*-activating the Ds elements than the sAc element. However, excluding the low responding Alb15 line, the progenies from the Ac^+ element showed a higher proportion of white (W) seedlings $(5\% \pm 2\%)$ compared to those with the sAc element (0.4% \pm 0.7%). This might be the result of the loss of the transposase source due to excision of Ac^+ and subsequent failed reinsertion, or due to segregation of Ac^+ after transposition to unlinked sites. Both possibilities did not apply to the immobile sAc element.

We used molecular analysis to determine whether excision of Ds from the SPEC gene corresponded to the phenotypes observed in the seedling assays (FG, V, W). To that end, DNA was isolated from progenies of each type of cross and subjected to PCR and Southern analysis such that empty donor site (EDS) fragments and full donor site (FDS) fragments could potentially be shown. FDS fragments were demonstrated in 32 mature F₁ plants that were V or FG at the seedling stage by PCR (act5b/ spec primers: 0.6 kb) and by Southern hybridization (*Hind* III/SPEC probe: 6.5 kb fragments; *Eco* RI/pSU20 probe: ≥3.3 kb fragments, Fig. 3, right panel). However, the expected EDS fragments (2.6 kb) could only occasionally be demonstrated by Southern hybridization (3 out 32) or by PCR (5 out 32) with the 35s/spec primers (0.5 kb) (data not shown). In contrast, PCR analysis with

Somatic Transposition



Fig. 5. Frequencies of somatic *Ds* excision from the T-DNAs of eight *Albatross* lines *trans*-activated by *sAc* presented in stack columns. The double hemizygous F_1 seedlings were germinated in the presence of Sp and were classified by the degree of green (Sp^R) variegation. Seedlings with uniform green cotyledons were designated fully green (FG); seedlings with almost entirely green cotyledons but with one or more small white flecks were designated as highly variegated (HV); seedlings with 3–10 distinct green spots were classified as medium variegated (MV); and seedlings that were almost totally white but had 1–2 small green spots were designated as lowly variegated (LV). Seedlings with out any visual green variegation were white (W). Seedlings with *chs:Ac*-induced variegation patterns were designated V1 when 1–3 large green spots of almost the size of the cotyledon were present, or classified as V2 when the cotyledons were homogeneously scattered by many (10–20) small spots. Each column represents the results of three F_1 plants, of which 3–6 capsules per plant were taken with 100–200 seeds per capsule.

the same primer set on DNA isolated from V and FG seedlings, instead of mature plants, indeed showed in all 24 cases the expected EDS fragments. Apparently, the high degree of somatic Ds excision during the formation of the cotyledons in embryogenesis, revealed by the Sp seedling assays, did not coincide with the low degree of somatic Ds excision in mature plants. This conclusion is in line with the observation that the degree of Sp resistant variegation at the plant level, revealed by regeneration Sp assays (Fig. 2A), was generally much lower than that observed at the seedling level (Figs 2C–J).

Germinal transposition of Ds

Outcrossing the double hemizygous Ac/Ds F₁ plants with the non-transgenic line resulted in the OC₁ progenies (Fig. 4). For each *Albatross* line, 3–12 F₁ siblings were used for the outcrosses and 3–13 pollinations were made per plant. The OC₁ seedlings were germinated on double selective medium, containing both Sp and Km. These Sp/ Km assays showed similar proportions of variegation classes as observed in the F₁, indicating that the variegation specificity of the *Albatross* lines was heritable (data not shown). Also in the OC₁, the *trans*-activation of *Ds* was generally very effective because in most *Albatross* progenies the expected 1/4 of the seedlings (carrying both Ac and Ds) was variegated. As in the F₁, *Alb*15 crossed with the *sAc* line was the only exception in that only a small fraction of the OC₁ seedlings was LV (<5%) and the others were W.

Transposition events in F₁ cell lineages that had been transmitted through the germline to the OC_1 resulted in FG seedlings when germinated on Sp/Km medium. The frequency of germinal transposition could only unmistakeably be calculated when FG seedlings (Sp^{R}/Km^{R}) were selected that lacked the Ac transposase source and, therefore, could not be the consequence of somatic excision, nor could Km resistance be conferred by the Ac T-DNA. This procedure was followed only for Ds elements trans-activated by the sAc element (Fig. 6). For molecular analysis, DNA was isolated from mature OC_1 plants that were FG (Sp^R/Km^R) at the seedling stage and that lacked the sAc element (GUS⁻). PCR with 35s/ spec primers showed the EDS fragments and, as expected, the FDS fragments (act5b/spec) and the Ac-specific fragments (ac6/ac7) could not be demonstrated. Progeny testing by Sp/Km seedling assays showed that 136 of the 145 OC₁ plants segregated for Sp and Km resistance in the OC_1S_1 (see next section). Southern analysis showed that nine OC_1 plants did not contain a Ds element (NPTII) and apparently escaped the selection. The majority (78%) of the FG seedlings (Sp^R/Km^R) were GUS^+ (Table 1). It would be expected that seedlings with germinal Ds transpositions segregated independently (1:1) from the sAc T-DNA (GUS). Therefore, the surplus of FG/GUS⁺ seedlings was most likely the result of somatic excision. Similar proportions (on average, 1%) of somatically-induced FG seedlings had been observed in the F_1 (Fig. 5). The Alb2 and Alb15 progenies showed the highest (0.95%) and the lowest (0.15%) germinal transposition frequencies, respectively (Table 1). The same Albatross lines showed similar results in the somatic transposition frequencies. The possibility that the low average germinal Ds transposition frequency (0.37%) of all *Albatross* lines *trans*-activated by the *sAc* element was caused by a high incidence of failed Ds reinsertions was tested by seedling assays with only Sp in



Fig. 6. Frequency of germinal *Ds* transpositions from the T-DNAs of eight *Albatross* lines *trans*-activated by *sAc* as determined in outcrossed progenies (OC₁). Sp^R/Km^R seedlings were selected that did not contain the *sAc* T-DNA (GUS⁻) and, hence, unmistakably carried a germinally transmitted excised and reinserted *Ds* element. The results are shown for 6–12 siblings per *Albatross* line (13 000–28 000 seeds).

Ds line	Number of F_1 plants	Number of OC1 capsules	FG seedlings ^a		T7 1 1	Germinal transposition	
			GUS-	GUS^+	Variegated seedlings ^b	frequency $(\% \pm sd)$	
Alb1	6	70	18	80	6,906	0.26 ± 0.06	
Alb2	12	34	30	100	3,173	0.95 ± 0.17	
Alb3	6	49	21	42	4,782	0.44 ± 0.09	
Alb10	6	44	18	53	5,169	0.35 ± 0.08	
Alb15	6	47	7	18	4,602 ^b	0.15 ± 0.05	
Alb17	12	33	13	51	3,211	0.41 ± 0.11	
Alb19	6	49	14	80	6,026	0.23 ± 0.06	
Alb20	12	36	15	65	3,308	0.45 ± 0.12	
Overall	66	362	136	489	37,177	0.37 ± 0.03	

Table 1. Summary of the germinal Ds transposition frequencies in eight *Albatross* lines *trans*-activated by the immobile sAc element

^aFor unambiguous identification of OC_1 plants that carried germinally transmitted transposed *Ds* elements, fully green (FG) seedlings (Sp^R/Km^R) were identified that did not contain the *sAc* T-DNA (GUS⁻); ^bthe number of seedlings that potentially could carry a germinally transmitted transposed *Ds* element (1/2 of the OC_1) was based on the number of variegated seedlings (1/4 of the OC_1); the population sizes of *Alb*15 progenies were determined by counting all seedlings.

the medium. However, the numbers of FG seedlings were not significantly higher than with Km present in the medium, indicating that Ds excision was generally coupled to reinsertion. Additionally, the reciprocal outcrosses showed no differences in the frequency of transmission of transposed Ds elements through the female or male gametes (data not shown). The Ac^+ line generally produced much higher frequencies of FG seedlings (1%-100%) than most of the sAc progenies (0.3%-2.0%). Similarly as in the F₁, the OC₁ progenies of $Alb15 \times Ac^+$ showed a low frequency of FG seedlings (1.4%). In progenies of the chs:Ac line only seedlings with the V1 and V2 phenotypes were observed. Out of an estimated number of 10,000 seedlings (3 F_1 siblings for each 8 Albatross lines, 3 outcrosses per plant), no FG seedlings were observed. Therefore, it is concluded that trans-activation of the Albatross lines by the chs:Ac element did not result in germinal transposition.

Germinal transposition frequencies do not necessarily reflect the proportion of unique Ds transposition events. Clonal propagation of transposition events in somatic F₁ tissues prior to the formation of gametes could lead to transmission of that single event to several OC₁ seedlings. To determine the incidence of independent unique Ds transpositions, the reproductive cell lineages were followed by recording the position of each pollinated F₁ flower. The occurrence of independent germinal Ds transposition was established by investigating whether Ds elements were inserted into different DNA restriction fragments. To analyse these new insertion site (NIS) fragments, DNA was digested (*Eco* RI or *Hind* III) and analysed by Southern hybridization with Ds-specific probes (pSU20 or NPTII) (Fig. 3, right panel). NIS fragments of different sizes indicated that the Ds elements were reinserted at different positions in the genome. Out of the 136 OC1 plants with germinally transmitted transposed Ds elements, 18 were obtained that were derived from F_1 plants that generated only one such OC_1 plant and, by definition, carried unique transposition events. Comparison of NIS fragments between siblings showed that similar NIS fragments were only observed for OC_1 plants that were derived from the same seed capsule, i.e. F_1 flower. Twenty OC₁ plants (20/ 118 = 17%) had similar NIS fragments as one or two siblings and, therefore, in principle carried the same premeiotic transposition event. Thus, in total, 116 OC1 plants were identified that carried unique Ds transposition events. Three of these contained two NIS fragments and, hence, contained two transposed Ds elements.

Distance of Ds transposition

A total of 113 OC_1 plants was identified that carried single, unique, transposed and germinally transmitted Ds elements. To determine the distance of transposition of these Ds elements, OC_1S_1 progenies (200–400 seedlings) were analysed for segregation of Km and Sp resistance that were conferred by the Ds element (NPTII) and the T-DNA (SPEC), respectively (Fig. 4). If significant linkage was observed (p < 0.01), recombinant fractions were calculated and converted to genetic distance in cM. Each Albatross line showed differences in the distribution of transposed Ds elements (Fig. 7). The transposition patterns mainly varied in the dispersal of loosely linked transpositions. Notwithstanding the variation between Ds lines, generally each pattern showed similar characteristics (Table 2). In most *Albatross* lines the majority of *Ds* elements transposed closely linked (0 cM) and unlinked





Fig. 7. Distance of transposition from the T-DNAs of eight *Albatross* lines *trans*-activated by *sAc* as determined in OC_1S_1 progenies. Distances were expressed in cM using Haldane's mapping function to convert the recombination frequencies between the *Ds* elements and the original donor site T-DNAs (P < 0.01). The *Ds* element was marked by the *NPTII* gene and the T-DNA was marked by the *SPEC* gene. The segregation of *Ds* and the T-DNA was analysed in Sp/Km assays, where Sp^R/Km^R seedlings were fully green (FG), Sp^S were white (W), and Sp^R/Km^S seedlings, carrying a recombination event, were yellow (Y).

Table 2. Summary of the distance of Ds transpositions in eight Albatross lines trans-activated by the immobile sAc element

Ds line		Linked transpositions ^a				Unlinked transpositions		
	Number of unique transposed Dss	Mean distance Number % (cM±sd) Range (cM)				Number	%	-
Alb1	13	11	85	3.4 ± 2.4	0.0-15.3	2	15	
Alb2	24	19	79	12.6 ± 3.0	0.0-26.9	5	21	
Alb3	18	14	78	3.6 ± 2.4	0.0-15.8	4	22	
Alb10	17	14	82	3.5 ± 2.4	0.0 - 17.4	3	18	
Alb15	6	6	100	2.9 ± 2.0	0.0-9.3	0	0	
Alb17	10	5	50	5.8 ± 2.2	1.6-4.9	5	50	
Alb19	11	11	100	6.6 ± 3.3	0.0-33.0	0	0	
Alb20	14	11	79	7.4 ± 3.4	0.0-37.7	3	21	
Overall	113	91	81	6.4 ± 2.9	0.0-37.7	22	19	

^aIf significant linkage was observed ($p = \langle 0.01 \text{ if } \chi^2 \rangle 11.32$) recombinant fractions were converted to cM by Haldane's mapping function.

(>40 cM) to the original T-DNA. The exceptional line was *Alb2* of which the majority of transposed *Ds* elements were mapped at positions between 2–27 cM. The average frequencies of linked (<40 cM, 81%) and unlinked (19%) transpositions were comparable in most *Albatross* lines. The mean transposition distance of all *Albatross* lines was

6.4 cM; of which the *Alb*2 line significantly deviated with 12.7 cM. The overall frequency distribution of all 113 transposition distances showed that 35% could not be separated from the donor site, and that the number of transposed *Ds* elements gradually decreased with increasing distances of transposition (Fig. 8).





Discussion

Somatic excision of Ds

We systematically characterized the behaviour of an artificial Ds element in eight tobacco lines trans-activated by three different Ac elements. Since the Ds elements remained immobile until fertilization, the timing and frequency of somatic excisions during embryogenesis could efficiently be monitored visually in seedling assays. The high proportions of seedlings with excised Ds elements showed that trans-activation during cotyledon development was generally very effective. Furthermore, each Ds line showed characteristic spectinomycin resistant (Sp^R) variegation patterns that were inherited to following generations. The lines differed in the proportion of seedlings with Sp^R green spots of different size and number, which were a consequence of the timing and frequency of Ds excisions during development of the cotyledons, respectively. Variability in Ds excision patterns has also been observed in Arabidopsis and lettuce (Bancroft and Dean, 1993a; Yang et al., 1993). The unmodified (wild-type) and transpositionally active Ac^+ element proved to be more effective in trans-activating the Ds elements than the immobile sAc element. This positive effect on the Ds transposition frequency could be the result of an increased number of Ac^+ copies, or could be due to a more *trans*-active Ac^+ element (Jones et al., 1991). The lower trans-activation effectiveness of the sAc element was probably not the result of the sequence modification because the 177 bp terminal deletion does not include the transposase gene (Keller et al., 1993).

The *chs*:Ac element induced different *Ds* variegation patterns (V1, V2) than the Ac elements that were under the control of the endogenous promoter. The V1 class

consisted of seedlings with 1-3 large green spots as a consequence of Ds excision in a few cells early in cotyledon development. These spots arose from a few progenitor Sp^R cells in the center of the cotyledon and did not include the edges. The V2 class, on the other hand, consisted of a large number of small green spots resulting from Ds excisions in many cells late in cotyledon development. The occurrence of two types of chsA promoter-induced variegation patterns suggested that this promoter was active twice during cotyledon development. However, seedlings were only identified that showed either the V1 or the V2 pattern, indicating that during development of an individual seedling the chsA promoter was active only once. Alternatively, the V1 pattern could be the consequence of a few early Ds excisions until high transposase levels became inhibitory (Scofield et al., 1993). The activity of the chsA promoter in P. hvbrida seedlings has been demonstrated by GUS fusions (Koes et al., 1990; Van der Meer et al., 1990). Using other *promoter:Ac* transposase gene fusions (nos:Ac, ocs:Ac, CaMV-35S:Ac) in experiments with similar visual Ds excision assays (streptomycin resistance, SPT) distinct variegation patterns have also been observed, unlike those caused by the immobile sAc or Ac^+ elements (Scofield *et al.*, 1992).

Germinal transposition of Ds

In contrast to developing seedlings, Ds excisions took place infrequently in mature plants. Most likely, the low somatic Ds activity resulted in accompanying low frequencies by which transposed Dss were germinally transmitted (0.37%). This suggestion is supported by the observation that the same Ds lines showed the highest (Alb2) and lowest (Alb15) values in both somatic and germinal Ds transposition frequencies. Previously, it has been shown that *trans*-activation of another *Ds* line by the same and other sAc lines resulted in similar low germinal transposition frequencies (0.38%, Scofield et al., 1992). The high incidence of independent transposition events (83%) indicates a late timing of transposition, i.e. during development of the generative meristem, prior to gametogenesis. Following unlinked transposition of Ds, only half of the progeny seedlings (Sp^R/Km^R) carrying that event could be visually identified by the fully green (FG) appearance, and the others (Sp^S/Km^R) were white (W). Therefore, the phenotypic identification of seedlings carrying transposed Ds elements included a selection bias for genetically linked transpositions. On the basis of the proportions of unlinked and loosely linked Ds transposition in 113 of the 136 selected seedlings (Table 2), it was estimated that a maximum of 30 seedlings with unlinked transpositions could have escaped the selection. Consequently, incorporating these non-selected seedlings would increase the overall germinal transposition frequency to 0.45% ([30 + 136]/37177; Table 1).

Frequency and distance of Ds transposition

The trans-activation of Ds by the chs:Ac element did not result in germinally transmitted transposition events. Most likely the chsA promoter did not express the Ac transposase gene in germinal cell lineages (L2) because it is predominantly active in L1 tissues (Koes et al., 1990; Van der Meer et al., 1990). No differences were observed in progenies from the reciprocal outcrosses, excluding differences in the activity of the chsA promoter between male and female organs. In other promoter: Ac fusion experiments in tobacco, the heterologous promoters induced distinct germinal Ds excision frequencies: 7% for the CaMV-35S:Ac fusion, 0.87% for the ocs:Ac fusion and, 0.09% for the nos: Ac fusion (Scofield et al., 1992). Remarkably, in contrast to these low germinal transposition frequencies of Ds elements (0.0–7%), much higher germinal excision frequencies (4-80%) were reported for wild-type Ac^+ elements in tobacco. However, comparison of the transposition rates is complicated by the different Ac dosages, and different transposition assays applied in these studies (Hehl and Baker, 1990; Jones et al., 1991; Fitzmaurice et al., 1992; Keller et al., 1993). In most other heterologous hosts, the activities of Ac^+ and Dscannot be compared directly because the regulatory sequences of the Ac component were often modified when used in combination with Ds (Bancroft et al., 1992; Ellis et al., 1992; Yang et al., 1993).

Distance of Ds transposition

Although the 113 Ds transpositions from the eight T-DNA loci in tobacco showed a general pattern, their distributions varied in the dispersal of loosely linked transpositions. This variability in transposition pattern has been observed before for Ac in tobacco and for Ds in Arabidopsis (Dooner et al., 1991; Bancroft and Dean, 1993b). Following simultaneous selection for seedlings with transposed Ds elements, half of the seedlings with unlinked transpositions were not identified, and as a consequence, the proportions of unlinked transpositions were possibly underestimated. However, the Ds elements in all tobacco lines showed a strong tendency for transposition to genetically closely linked sites. On average, 81% of the Ds elements transposed within 40 cM, and 58% transposed within a 10 cM distance. Similar observations have been made for Ac in tobacco: 72% within 40 cM, 45% within 10 cM (Jones et al., 1990; Dooner et al., 1991). In addition, the frequencies of linked (<40 cM) Ds transposition distances in Arabidopsis and tomato, also determined by segregation analysis of the T-DNA and the Ds, were comparable: 68% and 84%, respectively (Bancroft and Dean, 1993b; Carroll et al., 1995).

Implications of this study

For the further development of transposon tagging strategies in heterologous hosts using the Ac/Ds

system, the parameters involved should be optimized. To increase the transposon-induced mutation frequency, high germinal Ds transposition frequencies should be established by modification of the Ac transposase sources. Higher transposase levels produced by using promoters lead to increased transheterologous position frequencies up to a certain threshold when transposition is inhibited (Scofield et al., 1993). With respect to these dosage effects, transposase expression should be regulated within a narrow range. Simultaneously. high frequencies of independent transpositions should be achieved by inducing transposition in late germline cells. Therefore, Ds elements should be trans-activated in the generative meristem just prior to the formation of gametes by regulated expression of the Ac transposase using tissuespecific or inducible promoters (Balcells et al., 1994; Charng et al., 1995). The Ds lines with the visual transposition reporters that were generated and characterised in this study will be useful for testing novel Ac constructs.

In transposon tagging experiments with particular Ds lines, the variability in the frequency and distances of germinal Ds transpositions can result in unexpected lower mutation frequencies. Variation in the expression of transgenes is presumed to be attributed to position effects due to differences in the integration site of the T-DNA. Supposedly, the chromatin conformation at the site of integration influences the binding of transcription factors and other factors. Indeed, the presence of chromatin matrix-associated regions (MARs) around the transgenes has been shown to reduce the variability of their expression in tobacco (Mlnàrovà et al., 1994). It is suggested that the variability in Ds transposition behaviour is caused by differences in the timing and frequency of binding of the Ac transposase and other factors due to the effects of the genomic position. This hypothesis may be tested by cloning MARs around the Ds elements. Likewise, the establishment of artificial chromatin domains around the T-DNA loci may result in the Ds elements being less susceptible to influences of chromatin in the topological vicinity of the integration site.

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