RNA Target Sequences Promote Spreading of RNA Silencing

Helena Van Houdt, Annick Bleys, and Anna Depicker*

Department of Plant Systems Biology, Flanders Interuniversity Institute for Biotechnology, Ghent University, Karel Lodewijk Ledeganckstraat 35, B–9000 Ghent, Belgium

It is generally recognized that a silencing-inducing locus can efficiently reduce the expression of genes that give rise to transcripts partially homologous to those produced by the silencing-inducing locus (primary targets). Interestingly, the expression of genes that produce transcripts without homology to the silencing-inducing locus (secondary targets) can also be decreased dramatically via transitive RNA silencing. This phenomenon requires primary target RNAs that contain sequences homologous to secondary target RNAs. Sequences upstream from the region homologous to the silencing inducer in the primary target transcripts give rise to approximately 22-nucleotide small RNAs, coinciding with the region homologous to the secondary target. The presence of these small RNAs corresponds with reduced expression of the secondary target whose transcripts are not homologous to the silencing inducer. The data suggest that in transgenic plants, targets of RNA silencing are involved in the expansion of the pool of functional small interfering RNAs. Furthermore, methylation of target genes in sequences without homology to the initial silencing inducer indicates not only that RNA silencing can expand across target RNAs but also that methylation can spread along target genes.

RNA silencing is a conserved mechanism that occurs in various eukaryotic organisms and leads to targeted degradation of RNA sequences homologous to the trigger (for review, see Matzke et al., 2001; Sharp, 2001; Zamore, 2001; Huttvéigner and Zamore, 2002). The potency of double-stranded RNA (dsRNA) in activating RNA silencing was first demonstrated in Caenorhabditis elegans and was designated RNA interference (RNAi; Fire et al., 1998). Posttranscriptional gene silencing (PTGS; for reviews, see Kooter et al., 1999; Vaucheret et al., 2001; Voinnet, 2001), which is mechanistically related to RNAi, can also be efficiently elicited in plants upon introduction of gene constructs that give rise to dsRNA (Hamilton et al., 1998; Waterhouse et al., 1998; Chuang and Meyerowitz, 2000; Smith et al., 2000; Sijen et al., 2001a; Wesley et al., 2001; Stoutjesdijk et al., 2002) or upon viral infection of plants that initiates production of dsRNA replication intermediates (Ruz et al., 1998; Baulcombe, 1999).

Sense and antisense small RNAs (approximately 20–25 nucleotides), homologous with posttranscriptionally silenced sequences, accumulate specifically in various PTGS systems in plants (Hamilton and Baulcombe, 1999). Studies of RNAi in fruitfly (Drosophila melanogaster) revealed that small RNAs result from symmetric processing of the dsRNA (Hammond et al., 2000). Hamilton and Baulcombe (1999) first proposed that these small RNAs correspond to specificity determinants in PTGS and RNAi. These small RNAs have been shown to guide a nuclease complex to cleave single-stranded RNA with complementary sequences in fruitfly embryo lysates (Elbashir et al., 2001a, 2001b). Therefore, the 21- to 23-nucleotide RNAs are referred to as small interfering RNAs (siRNAs) or guide RNAs. Target mRNAs are cut in the center of the region recognized by the complementary guide RNAs (Elbashir et al., 2001a), and mRNAs are cleaved only in the region corresponding to the dsRNA (Zamore et al., 2000).

The amplification of the siRNA signal during RNAi in C. elegans has been investigated and, in addition to trigger-coincident siRNAs, populations of small antisense RNAs have been detected that correspond to regions of the target RNA molecules located upstream of the initial trigger dsRNA, designated secondary siRNAs (Sijen et al., 2001b). The abundance of secondary siRNAs seems to decrease as a function of the distance from the region homologous to the primary trigger. Functionality has been demonstrated by means of a transitive RNAi assay (Sijen et al., 2001b), in which two targets for silencing are provided. Similarly, plant viral vectors carrying part of a transgene elicit the production of transgene-specific, secondary siRNAs upon infection. As a consequence, these plants are protected against infection by an unrelated virus that carries another part of the transgene (Vaistij et al., 2002).

Plant RNA silencing is frequently accompanied by DNA methylation in symmetrical and nonsymmetrical cytosines (Bender, 2001) in transcribed regions of the silenced genes (Ingelbrecht et al., 1994; English et
al., 1996; Sijen et al., 1996; Kovářík et al., 2000; Van Houdt et al., 2000a), although its role is still unclear. Sequence-specific methylation signals consisting of RNA-DNA associations are believed to be involved in methylating silenced genes (Wassenegger, 2000). RNA-directed DNA methylation (RdDM) was first discovered in tobacco (Nicotiana tabacum) plants that contained multimeric genome-integrated copies of a viroid cDNA (Wassenegger et al., 1994). In these plants, specific de novo methylation that is restricted to the cDNA region was detected whenever viroids replicated autonomously (Pelissier et al., 1999). Further, viroid-infected plants accumulate small RNAs identical in size to those found in plants exhibiting PTGS of transgenes (Papaefthimiou et al., 2001). These results suggest that viroid-related RNAs induce methylation of the homologous cDNA copies. However, it remains controversial whether silencing-triggering dsRNA molecules, small guide RNAs, or intermediate RNA products are the signals for methylation of homologous DNA. PTGS induced by viral RNA that carries a short region homologous to the transgene leads to spreading of methylation throughout the transcribed region of the transgene (Jones et al., 1999; Thomas et al., 2001; Vaistij et al., 2002). Direct interaction between the input recombinant virus and the homologous transgene might lead to RdDM, and the progressive degradation of target mRNA (Zamore et al., 2000) could release more fragments, which additively direct methylation throughout the transcribed region of the transgene (Thomas et al., 2001). As an alternative, viral dsRNAs, synthesized by a putative plant RNA-dependent RNA polymerase (RdRp; Dalmay et al., 2000; Mourrain et al., 2000) or its derived siRNAs could mediate RdDM (Vaistij et al., 2002).

Here, we address the question of whether the in trans-silencing capacity of a silencing-inducing transgene locus can be transmitted to target RNA, subsequently able to silence secondary targets in trans. Therefore, we tested whether a posttranscriptionally silenced transgene locus can silence in trans a secondary target, which only produces nonhomologous transcripts; to this end, we created a stepwise homology between the silencing inducer and the secondary target by producing a transcript from a primary target with one region homologous to the silencing inducer and another region homologous to the secondary target mRNA. We investigated the production of small, approximately 22-nucleotide long RNAs, corresponding to target mRNAs and analyzed the methylation status of sequences silenced in trans in the region nonhomologous to the silencing inducer.

RESULTS

A Posttranscriptionally Silenced Inverted Repeat Transgene Locus Can Trigger Silencing of a Reporter Gene Producing Nonhomologous Transcripts

To determine whether the RNA-silencing activity of a silencing-inducing RNA can be transmitted to another RNA sequence by fusing this sequence to part of the silencing-inducing sequence in a single transcript, we studied transgenic tobacco plants with different combinations of three transgene loci (X, Y, and Z). Locus X (Fig. 1) harbors an inverted repeat about the right T-DNA border of T-DNA GVCHS287, carrying a neomycin phosphotransferase II (nptII) gene under control of the cauliflower mosaic virus 35S promoter (CaMV P35S) and the 3′-signaling se-
quences of the chalcone synthase gene (3' chs) of snapdragon (Van Houdt et al., 2000a, 2000b; Fig. 1). The two convergently transcribed nptII genes in locus X produce only very low amounts of the NPTII protein compared with those produced by a single-copy nptII transgene. The invertedly repeated nptII transgenes in locus X had been shown to be posttranscriptionally silenced and methylated in the 3' one-half of the genes (Van Houdt et al., 2000a). Locus Y (Fig. 1) contains a single copy of the T-DNA GUSchsS and harbors a chimeric β-glucuronidase (gus) gene under control of P3SS and 3'chs with an artificial intron in the 5' region of the coding sequence. In tobacco plants hemizygous for locus Y, the gus expression levels are normal (Table I). Locus Z has two or more copies of T-DNA XD610 with a gus gene under control of P3SS and the 3'-untranslated region (UTR) of the nopaline synthase gene (3' nos). In locus Z (Fig. 1), the gus expression is stable (Table I). Here, the in trans-silencing effects between these three transgene loci X, Y, and Z were studied in hybrid transgenic tobacco plants with any possible combination of these three loci under hemizygous condition, obtained by crossing the appropriate parental plants (“Materials and Methods”). In trans-silencing effects were revealed by a reduced GUS activity of particular loci in certain combinations. The results of the GUS activity measurements in protein extracts of different types of hybrid plants are summarized in Table I.

The silenced nptII genes in locus X have previously been demonstrated to silence in trans homologous nptII transgenes (Van Houdt et al., 2000a) and transiently expressed genes with partial transcript homology to locus X-derived nptII transcripts (Van Houdt et al., 2000b). We also observed that the stably expressed gus gene in locus Y, with partial transcript homology to the nptII transcripts of the silencing-inducing locus X (Fig. 1), was silenced efficiently in trans (compare XY with Y; Table I). The homology between the transcripts of X and Y was mainly situated in the 3'-UTR (206 nucleotides), but the 5'-UTR also had a small region of homology (29 nucleotides). This relatively short region of homology between locus X-derived nptII and locus Y-derived gus transcripts was sufficient to degrade very efficiently Y-derived transcripts. To assess the stability of in trans-silencing of Y in XY hybrids, a 4-week-old phosphinothricin-resistant progeny of a self-fertilized XY hybrid was analyzed. Loss of locus X in the progeny plants, as revealed by kanamycin sensitivity, correlated with reactivation of gus expression in locus Y in the expected 1:4 ratio, indicating that the in trans-silenced phenotype is not transmitted to the next generation when the silencing-inducing locus X is absent.

In contrast to the low GUS activity detected in XY hybrids, the GUS activity in XZ hybrids was normal and similar to that in Z plants (Table I). This observation allows us to conclude that the nptII genes of locus X could trigger neither transcriptional silencing of the gus genes in locus Z, although also driven by the CaMV P3SS promoter, nor posttranscriptional silencing of the gus genes in locus Z, which was expected because both loci produce transcripts without significant homology (Fig. 1). These data, in addition to results of run on transcription analyses of locus X-containing plants (Depicker et al., 1996; M. Fojtova and A. Kovar, unpublished data), support the fact that silenced effects in YX hybrid tobacco plants are not triggered by P3SS homology.

When Y and Z loci were combined in so-called YZ hybrids, the tobacco cells produced two types of gus transcripts with a 1,809-nucleotide homologous region in the gus-coding sequence. Both types of gus genes, however, remained normally expressed as reflected in the high GUS activity in YZ hybrids, and displayed a dosage effect, as expected for normally expressed genes (compare YZ with Y and Z; Table I). Thus, the RNA-silencing mechanism was not activated in YZ hybrid tobacco plants. Therefore, it is interesting to observe that upon creation of a stepwise homology between X and Z by the presence of locus Y, the gus expression in locus Z together with that of locus Y was reduced in XY hybrid tobacco plants (compare XYZ with YZ; Table I).

As shown schematically in Figure 2, silencing of the target could be triggered by a stepwise homology that was created between a silenced locus (X) and a nonhomologous target gene (Z) by introducing a chimeric recombinant gene (Y) with one region homologous to the silenced locus (X) and another homologous to the target (Z). We refer to this as a case of transitive silencing in plants in which the silencing inducer (X) and the primary and secondary targets (Y and Z) are all nuclearily expressed transgenes. The

<table>
<thead>
<tr>
<th>Genotype</th>
<th>GUS Activitya</th>
<th>No. of Plants Analyzedb</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>Below detection</td>
<td>1</td>
</tr>
<tr>
<td>Y</td>
<td>368 ± 165</td>
<td>9</td>
</tr>
<tr>
<td>Z</td>
<td>126 ± 30</td>
<td>10</td>
</tr>
<tr>
<td>XY</td>
<td>2 ± 1</td>
<td>4</td>
</tr>
<tr>
<td>XZ</td>
<td>139 ± 35</td>
<td>9</td>
</tr>
<tr>
<td>YZ</td>
<td>477 ± 101</td>
<td>10</td>
</tr>
<tr>
<td>XYZc</td>
<td>4 ± 3</td>
<td>22</td>
</tr>
</tbody>
</table>

aMean ± so. bThe axenically grown plants were analyzed 4 weeks after sowing on Murashige and Skoog medium supplemented with 1% (w/v) Suc in Falcon petri dishes (Becton Dickinson, Bedford, MA). cXYZ plants were grown in the presence of phosphinothricin (10 mg mL⁻¹) and hygromycin (25 mg mL⁻¹); under these conditions, both XYZ plants and YZ plants containing the pNE T-DNA (see “Materials and Methods”) were able to develop. A polymerase chain reaction with X-specific primers was performed to screen for the presence of X.
results imply that the silencing capacity of locus X is transferred to Y sequences upstream of the homology between X and Y.

In Trans-Silenced Loci Give Rise to Approximately 22-Nucleotide Small RNAs

To understand the observation that locus Z is silenced only in XYZ hybrids through a transitive silencing effect and to confirm that silencing capacities of silencing-inducing loci can expand to target genes, we assessed the accumulation of sequence-specific small RNAs. First, we determined whether small RNAs specific for the silencing-inducing locus could be detected. RNA gel blots using hydrolyzed gus RNAs as a probe (Fig. 3A) revealed locus X-specific nptII 22-nucleotide small RNAs in the low-M₉ RNA fraction of plants hemizygous for locus X and of XYZ hybrid tobacco plants. Wild-type SR1 plants (Fig. 3A, lane 1), hybrid lines without the nptII transgene (Fig. 3A, YZ in lanes 6–8), and a transgenic line containing a normally expressed nptII transgene (Fig. 3A, lane 5), which show transitive silencing, but not YZ hybrids, X-hemizygous plants, or wild-type SR1 (Fig. 3B, lanes 6–8), only relatively small hybridizing molecules were detected. RNA gel blots using hydrolyzed target-specific small RNAs. Therefore, we suggest that small RNAs corresponding to 3’chs-UTR region may direct the formation of small RNAs of more upstream-located sequences.

DNA Sequences That Are Nonhomologous to the Silencing-Inducing Locus X Become Methylated upon Inactivation

The silencing-inducing locus X contains two invertedly repeated, convergently transcribed nptII transgenes that are extensively methylated in the center of the repeat (Van Houdt et al., 2000a). In addition, a completely homologous nptII transgene becomes methylated upon in trans silencing by locus X (Van Houdt et al., 2000a). Does in trans methylation rely on sequence homology with the (initial) silencing-inducing locus or can target sequences silenced in trans be involved in producing the methylation signal? To address these questions, we examined the methylation status of several cytosines located in the gus-coding sequences in the genomic DNA of non-silenced YZ and silenced XYZ plants via DNA gel-blot analysis with the methylation-sensitive restriction enzyme HpaII. Figure 4A shows the gus-coding sequences in the T-DNAs of loci Y and Z, the location of non-methylation-sensitive restriction sites HpaI and ScaI, used to delimit the analyzed region of the T-DNA, and of the methylation-sensitive HpaII restriction sites, and the sizes of the digestion products of non-methylated gus transgenes. HpaII recognizes the sequence CCGG and will not cut this sequence in case at least one of the cytosines is methylated. Upon complete digestion with HpaII of the gus transgenes in locus Y and Z, only relatively small hybridizing DNA fragments will be detected, with the largest one being 427 bp (Fig. 4A). This result was obtained with different samples of YZ genomic DNA analyzed in a triple digest ScaI/Hpal/HpaII (Fig. 4B, lanes 5–8). However, the gus probe clearly revealed intense bands of higher Mr in the different XYZ genomic DNA samples than in the YZ samples, indicating strongly enhanced cytosine methylation upon silencing of the gus genes (Fig. 4B, compare lanes 1–4 with 5–8). The size of the hybridizing bands with higher molecular mass (0.7, 1.0, and 1.1 kb; Fig. 4B) suggests the silencing-inducing locus X become methylated upon inactivation.
involvement of targets in spreading of RNA silencing

According to the current model for PTGS or RNA silencing (Kooter et al., 1999; Matzke et al., 2001; Vance and Vaucheret, 2001), dsRNA molecules elicit the activation of the silencing response. Because locus X consists of two invertedly repeated T-DNAs and contains convergently transcribed nptII transgenes (Van Houdt et al., 2000a), we postulate that readthrough over the T-DNA right border sequences results in the production of double-stranded hairpin RNAs that trigger silencing of the nptII transgenes. We show that low-M, RNA fractions of locus X-harboring tobacco plants and not similar fractions of transgenic plants with normal nptII expression contain approximately 22-nucleotide small nptII RNAs, which are probably the product of nptII dsRNA cleavage by a dicer-like RNase. The posttranscriptionally silenced transgene locus X can silence in trans the partially homologous gus transgene in locus Y (Van Houdt et al., 2000a; Table I). The mechanism of in trans silencing is most probably based on the presence of siRNAs that correspond to the region of homology between silencing inducing and target RNA, namely the 3′ region in our analysis. Thus, the 3′-specific small RNAs seem to mark the locus Y-derived gus transcripts for degradation.

How can locus X direct degradation of locus Z-derived gus transcripts in XYZ plants? According to the current model of target RNA degradation (Elbashir et al., 2001a), the likely hypothesis is that siRNAs homologous to locus Z-derived transcripts would be involved. gus-specific small RNAs are readily detected in the low-M, RNA fraction of XYZ plants, but not in similar fractions of non-silenced Z plants. These molecules are candidates to function as siRNAs for sequence-specific degradation of gus transcripts, resulting in the low GUS activity detected in XYZ tobacco plants compared with that in Z plants. In a recent study, the spreading of RNA targeting upon virus-induced gene silencing (VIGS) in Nicotiana benthamiana and Arabidopsis has been described (Vaistij et al., 2002). The effect of VIGS spreads beyond the viral sequences inducing RNA...
Figure 4. Outline of the DNA gel blot to analyze the cytosine methylation status in the gus-coding sequences of XYZ and YZ plants. A, Representation of the analyzed restriction sites. The analyzed region of the gus-containing T-DNAs in locus Y and Z (GUSchs and LXD610, respectively) is represented by a thick black bar for each T-DNA. The relative location of the start and stop codons of the gus gene and the synthetic intron (I) are indicated above the bars. The recognition sites of the methylation-insensitive restriction enzymes HpaI and SapI were used to border the segments for methylation analysis. The functional elements of the gus chimeric genes are indicated by boxes above the black bars (for abbreviations, see Fig. 1). The fragments drawn below the bars are the DNA fragments obtained upon full HpaII digestion of the SapI/HpaI and SapI/SapI fragments of the T-DNAs in locus Y and Z, respectively (size in bp). The sequence used as probe (the full-length gus-coding sequence) is given as a bar just below the Y and Z T-DNA fragments. B, DNA gel blot of XYZ and YZ hybrids. Genomic DNA of four gus-silenced XYZ and four normal gus-expressing YZ tobacco hybrids (13 μg per lane) was cut in triple digests with two methylation-insensitive enzymes HpaI and SapI and with the methylation-sensitive HpaII. The full-length gus-coding sequence was used as probe. The length of size markers is indicated in base pairs. Specifically in XYZ samples, hybridizing bands of higher M are detected.
silencing, because the single-stranded target transcripts are converted to dsRNA by the putative SDE1/SGS2 RdRp (Vaistij et al., 2002). This observation gives an insight into how the detected gus-specific small RNAs, corresponding to the most 3′-800 bp of the gus-coding sequence, might be generated. RdRp-dependent synthesis of gus antisense RNA could be primed by locus X-derived 3′/chs-specific small RNAs on a locus Y-derived transcript, in analogy with the extension of primer siRNAs into a dsRNA product in embryo extracts of fruitfly (Lipardi et al., 2001). As an alternative, because no priming is required in an RdRp-dependent polymerization reaction (Schiebel et al., 1993), a particular feature of the Y-derived gus transcript, such as the mere association with a siRNA-protein complex or partial degradation by an RNA-induced silencing complex (Hammond et al., 2000), could allow it to be recognized by an RdRp as template for synthesis. RNase III-like cleavage of the nascent dsRNA would subsequently give rise to the detected small gus RNAs, which are probably involved in the 3′-5′ spreading of silencing, as revealed by the reduced GUS activity in XYZ hybrids.

Several studies in plants indicate that silencing can also spread to transgene regions downstream of the primary target (5′-3′ spreading; Braunstein et al., 2002; Han and Grierson, 2002; Vaistij et al., 2002). This observation cannot be ascribed to siRNA-primed RdRp-directed synthesis of dsRNA on the basis of transgenic mRNA targets, nor can it be reconciled easily with unprimed RdRp synthesis, which appears to start preferentially at the 3′ terminus of the template (Schiebel et al., 1993).

Although several papers describe the production of secondary siRNAs and the spreading of RNA silencing induced upon viral infection (Braunstein et al., 2002; Vaistij et al., 2002) or by nucleary expressed transgenes (Han and Grierson, 2002), target-specific siRNA production apparently does not occur by default. In tobacco plants transformed with a chimeric transgene comprising sequences encoding gus followed by satellite RNA (satRNA), there is no indication for spreading of siRNA production upon helper virus infection (Wang et al., 2001). Also, phytoene desaturase and ribulose-1,5-bisphosphate carboxylase endogenous transcripts do not serve as templates for secondary siRNA production upon VIGS in Arabidopsis and are therefore not involved in the spreading process of RNA silencing (Vaistij et al., 2002). In summary, the generality, requirements, and characteristics of secondary siRNA production in RNA silencing remain to be determined.

PTGS in plants, resulting in the degradation of homologous RNAs, has frequently been associated with sequence-specific de novo methylation of transcribed sequences of silenced transgenes. Also, in trans-silenced transgenes homologous to a silencing inducer become extensively methylated (Van Houdt et al., 2000a; Béclin et al., 2002). The region of methylation of a silenced transgene, induced upon viral infection, has been confined to the region of homology between the viral genome and the transgene (Jones et al., 1998; Wang et al., 2001). However, in other studies, methylation of a silenced transgene induced upon viral infection spreads into transcribed sequences not corresponding to viral sequences (Jones et al., 1999; Thomas et al., 2001; Vaistij et al., 2002), which has been associated with maintenance of silencing in the absence of the viral inducer (Vaistij et al., 2002). Enhanced cytosine methylation of the gus-coding sequence in XYZ plants showing transitive silencing of the gus genes has been observed, whereas non-silenced gus genes in YZ plants remain hypomethylated. The detected methylation is confined to the 3′ one-half of the gus-transcribed sequences, coinciding with the region that mainly gives rise to the gus-specific small RNAs, probably because it might be the region to be copied first into dsRNA by an RdRp. Further, methylation is only partial, because fragments corresponding to hypomethylated molecules are detected. It remains unclear whether locus Y gus genes are predominantly methylated and thus Y and Z gus genes are discriminated as methylation targets or whether both Y and Z gus genes are partially methylated. Two possible types of interactions could be invoked for de novo methylation of the gus sequences. First, methylation could be RNA directed and induced by interactions of the Y and/or Z gus genes with gus dsRNAs or gus-specific small RNAs. In a second scenario, DNA-DNA interactions between the methylated 3′/chs regions of the nptII genes of the silencing-inducing locus X and target Y could be a signal for methylation of the paired sequences following by spreading of methylation into nonhomologous gus sequences.

The occurrence of transitive silencing will have to be taken into account in case RNA silencing or RNAi is the technique of choice in functional genomic studies to obtain a null mutant phenotype for any particular gene (Nishikura, 2001). It is possible that particular siRNAs produced by a silencing inducer correspond to an identical stretch of nucleotides in a family member of the studied gene, not targeted on purpose, that could initiate a process in which the partially homologous endogenous transcript is used as template to produce secondary siRNAs (Sanders et al., 2002). Therefore, transitive silencing of coordinately expressed genes with highly conserved domains can be anticipated. The design of a dsRNA trigger will be crucial to create a selected gene-specific mutant phenotype. On the other hand, we envision that transitive silencing can be applied in silencing technologies, circumventing laborious construction of inverted repeat transgenes.
MATERIALS AND METHODS

Transgenic Tobacco Lines and Production of Hybrid Plants

The production of plants containing locus X and several characteristics of locus X have been described previously (locus 1 in Van Houdt et al., 2000a, 2000b). The locus Y-containing primary tobacco (Nicotiana tabacum) transformant GUSchlussS29 was obtained via Agrobacterium tumefaciens cotransformation of tobacco cv. Petit Havana SRI leaf discs with the A. tumefaciens strains C58C1Ribusp (pGV2260, pNE) and C58C1Ribusp (pGV2260, pGUSchlussS). The plasmids pNcE, carrying a hygromycin resistance marker (De Buck et al., 1998), and pGUSchlussS, with the phosphinothricin resistance marker (Van Houdt et al., 2000b), have been described previously. Transformant GUSchlussS29 was obtained upon hygromycin selection and, in addition to the pNcE T-DNA insert(s), contained an independently segregating locus, designated locus Y, harboring a single copy of the GUSchlussS T-DNA. The tobacco leaf disc transformation in which the locus Z-containing primary transformant LXD610-2 was generated, has been described previously (De Loose et al., 1995).

Hemizygous X and Z plants were obtained as hybrid progeny by crossing tobacco plants homozygous for locus X (=HOlo1) and homozygous for locus Z (=LXD610-2/9) to wild-type SRI, respectively. Hemizygous Y plants originated from the cross between the hemizygous primary tobacco transformant GUSchlussS29 and SRI and by selecting for the presence of locus Y in the hybrid progeny with phosphinothricin (10 μg mL⁻¹). Hemizygous plants XY and YZ were the hybrid progeny plants of the cross between HOlo1 and GUSchlussS29 and between GUSchlussS29 and LXD610-2/9, respectively, which were selected for the presence of Y. Hemizygous plants XZ were the hybrid progeny of the cross between HOlo1 and LXD610-2/9, and hemizygous plants XYZ were obtained by crossing hemizygous plants XY to LXD610-2/9, which is homozygous for locus Z, and selected for the presence of locus Y. Because the presence of locus X in this hybrid progeny could not be selected for, we screened for hemizygous plants XYZ through a locus X-specific PCR.

Enzymatic Assays

Protein extracts were prepared and GUS activity was measured as described by Van Houdt et al. (2000b).

DNA Gel-Blot Analysis

Genomic DNA from leaf tissue of mature tobacco plants was isolated with the DNeasy plant kit (Qagen, Hilden, Germany). DNA gel-blots hybridization was mainly done as described previously (Van Houdt et al., 1997). Probes were labeled with the Gene Images random prime labeling kit (Amersham Biosciences, Little Chalfont, UK) and detected with the Gene Images CDP-Star module (Amersham Biosciences).

Small RNA Analysis

To detect small RNAs, the procedures described by Hamilton and Baulcombe (1999) and Mette et al. (2000) were adapted. Tobacco leaf tissue was frozen in liquid nitrogen, and total RNA was extracted with TRIzol reagent (Invitrogen, Carlsbad, CA), according to the manufacturer’s instructions. Most of the high-Mᵦ RNAs were precipitated and the lower Mᵦ RNAs were recovered from the supernatant as described. For the different samples analyzed, a similar amount of RNA of the lower Mᵦ, RNA fraction, as estimated on gel, was separated on gel (15% [v/w] polyacrylamide and 7 M urea) and transferred to Hybond N⁺ membranes (Amersham Biosciences) by electroblotting with a Kem en Tec semidry blotter II (BIOzym, Landgraaf, The Netherlands). Size and polarity controls, DNA oligomers were loaded on the same gels. 32P-labeled probes were synthesized in vitro from a linearized plasmid with an SP6/7 transcription kit (Roche Diagnostics, Brussels) and [α-32P]CTP. The probe was hydrolyzed into fragments of approximately 50 nucleotides. Hybridization and washes were performed as described (Hamilton and Baulcombe, 1999; Mette et al., 2000) at 30°C. Labeled membranes were exposed to a phosphor imager screen (Amersham Biosciences).

Distribution of Materials

Upon request, all novel materials described in this publication will be made available in a timely manner for noncommercial purposes, subject to the requisite permission from any third-party owners of all or parts of the material. Obtaining any permissions will be the responsibility of the requestor.

ACKNOWLEDGMENTS

We thank Sylvie De Buck, Gert Van der Auwera, and Frank Van Breusegem for critical reading of the manuscript and helpful comments, the partners of the European Union Biotech Project (no. QLRT-2000–00078) for stimulating discussions, Heide Van Horebeke for generating transformants and performing crosses, Els Van Lerberge for technical assistance, Dries Brabant and Ruben Dario Garcia Perez for analysis of particular hybrid lines, Pongsopha Attasart for her input in the small RNA analysis, and Martine De Cock and Rebecca Verbanck for help with the manuscript and the figures, respectively.

Received June 4, 2002; returned for revision August 3, 2002; accepted October 2, 2002.

LITERATURE CITED

Braunstein TH, Mouri B, Johannessen M, Albrechtsen M (2002) Specific degradation of 3’ regions of GUS mRNA in posttranscriptionally silenced tobacco lines may be related to 5’-3’ spreading of silencing. RNA 8: 1034–1044
Involvement of Targets in Spreading of RNA Silencing


Lipardi C, Wei Q, Paterson BM (2001) RNAi as random degradative PCR: siRNA primers convert mRNAs into dsRNAs that are degraded to generate new siRNAs. Cell 107: 297–307


