Components of the Arabidopsis mRNA Decapping Complex Are Required for Early Seedling Development

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To understand the mechanisms controlling vein patterning in Arabidopsis thaliana, we analyzed two phenotypically similar mutants, varicose (vcs) and trident (tdt). We had previously identified VCS, and recently, human VCS was shown to function in mRNA decapping. Here, we report that TDT encodes the mRNA-decapping enzyme. VCS and TDT function together in small cytoplasmic foci that appear to be processing bodies. To understand the developmental requirements for mRNA decapping, we characterized the vcs and tdt phenotypes. These mutants were small and chlorotic, with severe defects in shoot apical meristem formation and cotyledon vein patterning. Many capped mRNAs accumulated in tdt and vcs mutants, but surprisingly, some mRNAs were specifically depleted. In addition, loss of decapping arrested the decay of some mRNAs, while others showed either modest or no decay defects, suggesting that mRNAs may show specificity for particular decay pathways (3′ to 5′ and 5′ to 3′). Furthermore, the severe block to postembryonic development in vcs and tdt and the accompanying accumulation of embryonic mRNAs indicate that decapping is important for the embryo-to-seedling developmental transition.

INTRODUCTION

RNA decay is an essential process that allows rapid changes in a cell’s gene expression profile, such as in response to environmental and developmental signals. Decay of an mRNA can be initiated by internal cleavage, or RNA decay can proceed from either its 5′ or its 3′ end. A large number of studies have elucidated the importance of microRNA (miRNA)–directed RNA cleavage for development in both animal and plant systems (Reinhart et al., 2002; Carrington and Ambros, 2003). However, much less is known about the 5′ to 3′ and 3′ to 5′ decay pathways and their possible roles in the development of multicellular organisms.

In both mammalian and yeast cells, initiation of bulk mRNA decay typically requires the removal of the 3′ poly(A) tail (Couttet et al., 1997; reviewed in Parker and Song, 2004). The deadenylated mRNA can then enter one of two decay pathways: either the cytoplasmic exosome complex can extend decay from the 3′ end; or the 5′ cap can be cleaved (decapping), after which the XRN1 exoribonuclease hydrolyzes the RNA from its 5′ end. The decapping pathway is also used by specialized mRNA decay pathways. For example, deadenylation-independent decapping is used in yeast both for autoregulation of the RPS28B RNA and for nonsense-mediated decay, in which mRNAs with a premature stop codon are selectively degraded (Muhlrad and Parker, 1994; Badis et al., 2004). These RNA decay pathways have been characterized extensively in yeast, and studies using mammalian cells revealed that the mRNA-decapping pathway is largely conserved.

Decapping of mRNA in yeast is performed by DCP2, a NUDIX/mutT domain–containing protein whose activity is stimulated by interaction with a second protein, DCP1 (Beelman et al., 1996; Dunckley and Parker, 1999). In mammalian cells, decapping requires an additional protein, called either Ge-1 or HEDLS, which facilitates the binding of hDCP1 and hDCP2 (Fenger-Gron et al., 2005; Yu et al., 2005). No HEDLS homolog was found in the yeast genome, indicating that there are differences between the decapping complexes in animals and yeast.

Yeast and mammalian decapping enzymes localize to dispersed cytoplasmic foci called processing bodies (P-bodies) (van Dijk et al., 2002; Sheth and Parker, 2003). In addition to roles in RNA decay, P-bodies are also important for sequestering translationally arrested mRNAs (Liu et al., 2005a, 2005b; Sen and Blau, 2005), and in yeast, sequestered RNAs can exit the P-body and return to polysomes (Brengues et al., 2005). Thus, the P-body is emerging as a dynamic structure with functions in multiple aspects of RNA metabolism and control.

In plants, bulk mRNA decay pathways and their possible roles in development are not well known. One indication that these pathways might have developmental roles comes from observations of Arabidopsis thaliana poly(A) ribonuclease mutants, which show defects in embryogenesis (Chiba et al., 2004; Reverdatto et al., 2004). However, the relative importance of 5′ versus 3′ decay and the specificity of mRNA targets for each pathway are largely unknown.

The 5′ to 3′ mRNA decay pathway in plants has been studied using mutants with defects in the Arabidopsis XRN4 exoribonuclease. In contrast with yeast and animals, Arabidopsis does not

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appear to contain an XRN1-like enzyme but instead has three XRN2-like enzymes (XRN2 to XRN4), and XRN4 appears to encode the functional 5' exoribonuclease for the decay of cytoplasmic decapped RNAs (Kastenmayer and Green, 2000). Reverse genetics, ethylene insensitivity, and overexpression suppressor screens have led to the identification of xrn4 mutants (Gazzani et al., 2004; Souret et al., 2004; Olmedo et al., 2006; Potuschak et al., 2006), xrn4 mutants accumulate a small set of decapped RNAs, including fragments produced following the cleavage of some miRNA targets and RNAs encoding the ethylene-signaling components EBF1 and EBF2. These observations have led to the question of whether the 5' to 3' mRNA decay pathway in plants serves just for bulk decay or whether it might also be used for regulated destabilization of specific mRNA classes.

To identify genes that are required for normal leaf development, we are analyzing mutants with defects in leaf vein patterning. The vascular system is composed of two tissues, xylem and phloem, that are organized together into veins. Vascular tissues arise early in organogenesis, and leaves and cotyledons produce stereotypical patterns of dispersed veins. A large body of data supports a role for auxin as a positive signal reviewed in Sieburth and Deyholos, 2004; Van Norman et al., 2004; Clay and Nelson, 2005; Koizumi et al., 2005; Sieburth and Deyholos, 2006). One study identified VCS homolog from humans. In one study, hVCS (called Ge-1) mutants showed defects. Cotyledons of wild-type and vcs-1 mutants produce broad leaves with a pointed apex and a modestly diminished pattern of veins; at 22°C, they produce narrow leaves with a small number of abnormally thick veins; and when grown at 29°C, vcs leaves are highly reduced and contain only a single very thick vein. Molecular characterization of the lesions in these vcs mutants are consistent with their being null alleles. A highly similar gene, VARICOSE-RELATED (VCR), is located <1 kb from VCS. Although we could detect no phenotype for vcr mutants, we speculated that activity of this gene might be responsible for the suppressed phenotypes we observed at low growth temperatures. Recently, two studies characterized the VCS homolog from humans. In one study, hVCS (called Ge-1) was shown to colocalize with hDCP2 (Yu et al., 2005). DCP2 encodes the mRNA-decaping enzyme, which is located in P-bodies; this localization of hVCS suggested that VCS might also function in mRNA storage and decay. Another study found that interactions between hDCP2 and hDCP1 depended on the presence of hVCS (called HEDLS) (Fenger-Gron et al., 2005). These studies raised the possibility that the developmental defects in vcs mutants might arise due to defects in the assembly of the mRNA-decaping complex.

One strategy for analyzing the role of the mRNA-decaping pathway is to engage in reverse genetics and screen for temperature-sensitive mutants. To gain insight into the molecular mechanisms controlling vein patterning, we are characterizing Arabidopsis mutants with vein patterning defects. We previously characterized five vcs alleles (vcs-1 to vcs-5) that were isolated in the Landsberg erecta (Ler) background and characterized vcs-1 in detail (Deyholos et al., 2003). All five produced identical temperature-dependent phenotypes, with increasingly severe phenotypes occurring when the mutants were grown at intermediate and high temperatures. We have now obtained additional vcs alleles from the SALK T-DNA insertion collection (Alonso et al., 2003), which were generated in the Columbia (Col-0) background. These insertions segregated for mutants with a more severe phenotype than vcs-1. We determined that one insertion allele, vcs-7, was an RNA null mutant (see Supplemental Figure 1 online), so we characterized this mutant in detail. When grown at 22°C, vcs-7 mutants had short roots, small chlorotic cotyledons, and typically failed to produce leaf primordia. By contrast, vcs-1 mutants were green, had slightly longer roots, and produced narrow pointed leaves (Figure 1A; see Supplemental Figure 2 online). In addition, vcs-7 mutants produced cotyledons that expanded slowly and accumulated anthocyanins around their margins, hypocotyls that were short and slightly swollen, and root hairs that were short and swollen. These traits were similar to those we reported previously for vcs-1 grown at 29°C (Deyholos et al., 2003). The vcs-7 mutant also showed cotyledon vein pattern defects. Cotyledons of wild-type and vcs-1 plants contain a primary vein and four secondary veins that form between two and four closed loops, whereas in vcs-7, the cotyledon secondary veins were mostly free-ending, resulting in a fork-like appearance (Figure 1A). This vcs-7 phenotype is similar to that of the recently described vcs-6 allele (Xu et al., 2006).

The vcs-7 mutant also differed from vcs-1 in its developmental response to growth temperature. The vcs-1 mutant, when grown at 16°C, produced broad pointed leaves with modestly disrupted vein patterns (Deyholos et al., 2003). By contrast, growth of vcs-7 at 16°C resulted in only modest suppression. For example, among 33 vcs-7 mutants grown at 16°C, 8 produced no visible leaf primordia, 15 produced numerous very small structures that resembled leaf primordia, and 10 produced small leaves (see Supplemental Figure 3 online). These data suggest that most, but not all, of the low-temperature suppression of the vcs phenotype was abolished in the vcs-7 mutant.

To determine why vcs-7 and vcs-1 mutants showed such different phenotypes, we constructed vcs-1/vcs-7 transheterozygotes. The transheterozygote was indistinguishable from vcs-1; it produced green organs, a longer root, and leaves that expanded to become broad and pointed (Figure 1B). This result indicated that vcs suppression arose due to a dominant activity in the Landsberg accessions. A likely candidate for low-temperature suppression was VCR. Mutations in VCR result in no observable
phenotype (Deyholos et al., 2003), and because VCR and VCS are separated by <1 kb, we have been unable to assess redundancy genetically. However, we were able to test whether the suppression of vcs-7 was linked to the VCS locus. To do this, we examined the mutant F2 progeny from crosses between vcs-7 heterozygotes and Ler and between vcs-1 heterozygotes and Col-0. The F2 progeny from both crosses segregated for both severe (vcs-7-like) and suppressed (vcs-1-like) mutants in ratios consistent with two or more unlinked suppressor loci. Because Ler suppressors were not linked to VCS, we conclude that VCR does not contribute substantially to low-temperature suppression. These observations, together with the transheterozygote phenotype, indicate that unlinked dominant loci within the Ler genome suppress the vcs phenotype.

**tdt Mutants Appear Similar to vcs Mutants**

Another mutant in our collection, tdt, had a severe phenotype similar to that of vcs-7, but it mapped to a different chromosomal location. Like vcs-7, tdt mutants produced no leaves, short roots with short and swollen root hairs, chlorotic cotyledons that accumulated anthocyanins around their margins, and a short and swollen hypocotyl (Figure 1A). In addition, the cotyledon-hypocotyl junction was extremely fragile, which caused cotyledons to frequently fall off, even with gentle handling. We attribute this fragility to a pinched region that we often observed at the proximal end of the cotyledon petiole (see Supplemental Figure 4 online).

Vascular defects in tdt mutants were also similar to those of vcs-7 in that cotyledon secondary veins mostly failed to form closed loops (Figure 1A; see Supplemental Figure 1 online). However, tdt mutants also showed a novel vascular phenotype within the vascular transition. The vascular transition is a region where the root-like vascular organization of the lower hypocotyl shifts to a shoot-like vascular organization (Esau, 1965). In tdt-1 mutants, this region ranged from being devoid of detectable xylem to containing xylem but with poorly aligned veins. This defect showed incomplete penetrance, but it was most prevalent when seedlings were germinated at higher temperatures (Table 1). By contrast, not only did we never observe a loss of vascular tissue within the vcs transition zone (n > 500 vcs-7 mutants), but...
vcs-1 (and other alleles in the Landsberg background) instead accumulate extra ectopic tracheary elements within their cotyledons and the upper hypocotyl (Deyholos et al., 2003).

As with vcs-7, tdt was isolated in the Col-0 accession. To further explore possible similarities between vcs and tdt, we examined whether the phenotype of tdt mutants could also be suppressed by unlinked loci in Ler. Following a cross between tdt heterozygotes and Ler, the F2 mutants showed both suppressed and severe phenotypes (data not shown). We compared leaf development in vcs-1 (in the Ler accession) with leaf development of tdt after four sequential crosses to Ler (tdt MC4). As with vcs-1, tdt MC4 produced leaf primordia (Figure 1C). Furthermore, there was much greater leaf development in both vcs-1 and tdt MC4 when grown at 16°C, while growth at 22 and 29°C resulted in progressively more severe defects in leaf development. However, tdt MC4 showed less extensive leaf development than that shown by vcs-1. Although some details of the vcs and tdt phenotypes differed, the major phenotypic defects were similar, which suggested that VCS and TDT might function in related pathways.

Table 1. Prevalence of the tdt-1 Transition Zone Defect Depends on Germination Temperature

<table>
<thead>
<tr>
<th>Growth Temperature</th>
<th>Vascular Transition Zone Defect</th>
<th>n</th>
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<tbody>
<tr>
<td>16°C</td>
<td>36%</td>
<td>86</td>
</tr>
<tr>
<td>22°C</td>
<td>69%</td>
<td>42</td>
</tr>
<tr>
<td>29°C</td>
<td>94%</td>
<td>85</td>
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To gain insight into the leaf initiation defects in tdt and vcs-7 mutants, we used confocal microscopy to analyze shoot apical meristem (SAM) structure. In the wild type, the SAM arises during embryogenesis; in the mature embryo, it appears as a dome composed of three well-defined layers of small, dense cells (Barton and Poethig, 1993). Among the embryos derived from a tdt-1 heterozygote, 25% were clearly distinguishable by their misshapen cells (5 of 23). These embryos contained small dense cells in the SAM region; however, they were not organized into distinct layers and they did not appear as a dome (Figure 2A). The vcs-7 embryos were also clearly distinguished from their siblings based on their SAM morphology (seven vcs-7 embryos out of 25 embryos examined). These vcs-7 embryos contained small dense cells in the SAM region; however, this region was not dome-shaped, nor were the cells organized into distinct layers (Figure 2A).

To determine how the SAM changed following germination, we also examined SAM of 3-d-old seedlings. The wild-type 3-d-old SAM appeared in its characteristic dome of layered cells and was accompanied by leaf primordia (Figure 2A). By contrast, the tdt 3-d-old seedlings still showed disorganized cell layers and no leaf primordia (n = 13). The SAM of the vcs-7 3-d-old seedlings varied: some were modestly rounded and layered (Figure 2A), while others were more similar to the tdt embryonic SAM; however, none had leaf primordia (n = 11). Thus, the severe defects in tdt and vcs SAMs might preclude the initiation of leaf primordia.

Figure 2. SAM and Vascular Defects in tdt and vcs-7.

(A) Confocal analysis of SAM organization in mature embryos and 3-d-old seedlings. Arrows indicate ectopic differentiating tracheary elements. Bars = 10 μm.

(B) Reporter gene expression in cotyledon and hypocotyl transition zones (TZ) of the wild type (Col-0) and tdt-1. AthB8:GUS confers expression in provascular, procambial, and vascular cells; VH1:GUS confers strong expression in provascular, procambial, and phloem cells; DR5:GUS confers expression in cells responding to auxin. Bars = 100 μm.
Vascular Defects in the \textit{tdt} Transition Zone Arise Early

To understand the developmental basis for the \textit{tdt} mutant’s hypocotyl vascular defects, we characterized hypocotyl vascular development using well-characterized \(\beta\)-glucuronidase (GUS) reporters. We first used the AthB8:GUS reporter, which confers expression to provascular, procambial, and differentiated vascular tissues (Baima et al., 1995; Scarpella et al., 2004). To determine whether these tissues were present in the \textit{tdt} transition zone, we compared GUS staining patterns of AthB8:GUS in the wild type with that of AthB8:GUS in \textit{tdt} seedlings. In the wild type, we observed strong vascular-associated GUS staining (Figure 2B). In the \textit{tdt} mutants, we observed strong AthB8:GUS expression in places where vascular tissues were present. However, in the \textit{tdt} transition zone, the extent of GUS staining correlated with the extent of the vascular defect, and in individuals lacking transition zone vascular tissues, AthB8:GUS conferred no GUS staining. This observation suggested that TDT was required prior to provascular or procambial specification.

We also examined vascular development using the VH1:GUS reporter, which confers expression to provascular, procambial, and phloem cells (Clay and Nelson, 2002). We observed strong vascular expression in the wild type, including through the transition zone (Figure 2B). However, as with AthB8:GUS, the extent of GUS staining in the \textit{tdt-1} mutant transition zone correlated with the extent of transition zone vascular development. In \textit{tdt-1} mutants devoid of transition zone xylem, there was no VH1:GUS staining. These observations corroborated the AthB8:GUS expression patterns and suggested that the loss of xylem in the \textit{tdt-1} transition zone was accompanied by a loss of phloem.

Finally, we analyzed GUS staining patterns conferred by the auxin-responsive DR5:GUS reporter (Ulmasov et al., 1997), which is useful for detecting provascular and procambial cells (Mattsson et al., 2003). In \textit{tdt-1} mutants, we detected strong DR5:GUS expression at the proximal termini of cotyledon vascular strands but no expression within the transition zone (Figure 2B). This observation suggested that auxin was transported normally from the cotyledon but that transport pathways might terminate at the transition zone. Together, these reporter expression patterns suggested that vascular defects in the \textit{tdt} transition zone arose early, prior to provascular cell specification.

\textit{TDT Encodes the Arabidopsis Homolog of DCP2}

We mapped \textit{TDT} to a 65-gene interval near the top of chromosome 5. We then screened T-DNA insertion lines (Alonso et al., 2003) mapping to genes within this interval, looking for mutants with a similar phenotype. Only one line, SALK_000519, segregated for a similar mutant, and this mutant was indistinguishable from \textit{tdt-1}. Crosses between \textit{tdt-1} and the Salk line (heterozygotes) produced 25% mutant seeds (59 wild type and 18 \textit{tdt} \( \times \) \textit{2} = 0.108, \( df = 1 \)), critical value = 3.841 at 95%), indicating that these were allelic mutants. SALK_000519 (\textit{tdt-2}) contained a T-DNA insertion in exon 3 of gene At5g13570. We sequenced this gene from \textit{tdt-1} mutants and identified a 50-bp deletion within exon 5 (Figure 3A). The \textit{tdt-1} deletion occurred just after Val-176 and is

\begin{figure}[h]
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\includegraphics[width=\textwidth]{figure3.png}
\caption{\textit{TDT} Encodes the mRNA-Decapping Enzyme.}
\end{figure}

(A) Diagram of the \textit{TDT} gene. Boxes correspond to exons, and the 5' end is to the left. Light gray boxes correspond to the DCP2 domain, and dark gray boxes correspond to the NUDIX domain. Locations of the T-DNA insertion and the lesion identified in \textit{tdt-1} are indicated.

(B) Alignment of DCP2 and NUDIX domains of \textit{TDT} with those of \textit{Saccharomyces cerevisiae} (Sc), \textit{Schizosaccharomyces pombe} (Sp), and \textit{Homo sapiens} (Hs). Arrows indicate conserved E residues that are required for catalytic activity (all are conserved in \textit{TDT}), and asterisks indicate residues required for interaction with DCP1 (She et al., 2006). Solid gray underlining indicates the DCP2 domain, and the dashed underline indicates the NUDIX domain.
predicted to result in a truncated protein with 17 novel C-terminal amino acids.

To determine whether we had identified the correct gene, we tested both a 35S:green fluorescent protein (GFP):TDT transgene and a genomic clone containing the TDT gene for their ability to rescue tdt-1 mutants. We observed full rescue of tdt-1 mutants by introduction of the TDT genomic clone (observed for 12 independent lines). For example, line 29 was identified by PCR as heterozygous at the TDT locus. Its self-pollinated progeny segregated 348 phenotypically wild-type seedlings and 19 tdt-1 mutants (18.3:1), which is consistent with the 15:1 ratio expected for full rescue ($\chi^2 = 0.742$; critical value $= 3.841$ at 95%). The full rescue by the TDT genomic clone indicated that we had identified the correct gene. We observed partial rescue of the tdt mutant in lines carrying the 35S:GFP:TDT transgene. We followed the development of two homozygous tdt-1 transgenic seedlings. Both plants produced leaves; however, they were stunted, male-sterile, and showed limited female fertility. The partial rescue of tdt-1 by the GFP:TDT construct further confirmed that we had identified the correct gene.

TDT was predicted to encode a 374–amino acid protein with N-terminal DCP2 and NUDIX domains (Figure 3A). These two domains are highly conserved among mRNA-decapping enzymes (Figure 3B). The TDT N terminus, through the NUDIX domain, shared 49% identity (69% similarity) with the Schizosaccharomyces pombe sequence, 41% identity (65% similarity) with Homo sapiens, and 37% identity (62% similarity) with Saccharomyces cerevisiae. The C-terminal domains of these proteins are much less similar: both the S. cerevisiae and S. pombe proteins are much longer (741 and 970 amino acids, respectively, compared with 138 amino acids in TDT).

Studies of the yeast mRNA-decapping enzyme have identified three invariant Gln residues that are essential for catalytic activity (She et al., 2006); all three of these residues are present in the predicted TDT sequence (Figure 3B), and the lesions for tdt-1 and tdt-2 predict abolishing one and all three of these residues, respectively. Thus, the identical phenotypes of tdt-1 and tdt-2 are consistent with molecular predictions of a loss of enzyme activity. Characterization of the yeast DCP2 protein also led to the identification of three amino acids that are required for interaction with DCP1 (She et al., 2006). The Arabidopsis TDT sequence has mostly retained this motif; two of the amino acids are identical, and the third shows a conserved change (Figure 3B). The molecular identification of TDT and VCS as potential components of the same complex is consistent with their similar mutant phenotypes.

TDT and VCS Function in the Same Pathway and Can Physically Interact

In animal cells, Ge-1/HEDLS/vVCS provides a scaffold that stabilizes interactions between hDCP1 and hDCP2, and this stabilization is required for decapping activity (Fenger-Gron et al., 2005). Sequence identity between TDT and DCP2 and between VCS and Ge-1/HEDLS suggested that these Arabidopsis proteins might also function together in mRNA decay. To test this possibility, we analyzed tdt-1 vcs-7 double mutants and tested whether we could detect interactions between the VCS and TDT proteins.
the supply of mRNA destined for decay. To determine whether GFP:TDT localization was also sensitive to cycloheximide, we performed confocal analysis of cycloheximide-treated seedlings. Following a short cycloheximide treatment, no cytoplasmic foci were observed (Figure 4F); instead, we observed a cytoplasmic GFP signal similar to that of 35S:GFP. This loss of cytoplasmic foci following cycloheximide treatment supports the punctate pattern of GFP:TDT corresponding to P-bodies.

In mammalian cells, hVCS/Ge-1/HDLS is required for the assembly of the decapping complex. To test whether VCS was required for normal localization of TDT, we crossed GFP:TDT to vcs-7 and examined the intracellular pattern of GFP in mutant roots. We examined >25 GFP-positive vcs-7 mutants from three different crosses, and in every one, we found very weak and diffuse GFP expression (Figure 4G), while the phenotypically wild-type siblings still showed punctate spots within the cytoplasm. Incubation of the vcs-7 seedlings in the proteasome inhibitor MG132 resulted in slightly brighter, but still diffuse, GFP signal (Figure 4I), while a control incubation in DMSO (Figure 4H) had no discernible effect. The same construct in wild-type plants showed no altered pattern of GFP:TDT localization when incubated in DMSO or MG132 (Figures 4J to 4L). These data indicated that VCS is required for the normal localization of GFP:TDT and that in the absence of VCS, TDT is unstable. Together, these observations support the notion these two proteins functioning together in a complex.

RNA Decay Is Altered in vcs and tdt Mutants

The molecular characterization of vcs and tdt led to the prediction that these mutants might have RNA decay defects. To test this prediction, we compared decay rates for four seedling-expressed mRNAs among the wild type, vcs-7, and tdt-1. We assayed RNA decay by comparing relative levels of specific seedling-expressed mRNAs following cordycepin-induced transcriptional arrest (Gutiérrez et al., 2002).

We analyzed the decay of BLUE MICROPYLAR END3 (BME3; At3g54810) RNA, which encodes a GATA-type zinc finger transcription factor that is required for germination (Gutiérrez et al., 2002; Liu et al., 2005c). This RNA was previously reported as unstable, and our decay assays confirmed its instability in the wild type (Figure 5A). In both tdt-1 and vcs-7, this mRNA’s decay rate was approximately half that of the wild type. Reduction of decay in tdt and vcs suggested that the normal pathway for this mRNA’s decay includes decapping. However, the fact that some decay still occurred suggests either that the loss of TDT and VCS did not provide a complete block to decapping or that BME3 mRNA’s normal decay also uses another pathway. There is no evidence that this mRNA, or any included in this part of our investigation, is the target of a small RNA. Thus, the mostly likely candidate for an alternative decay pathway is exosome-mediated 3'-5' decay.

We also analyzed the decay of MYB4 (At4g38620) mRNA. This mRNA showed similar decay rates in the wild type, tdt-1, and vcs-7 (Figure 5A). These similar mRNA decay rates suggested that decay of this mRNA might not require decapping; instead, it might specifically use the exosomal 3'-5' pathway.

We also analyzed the decay of RNA encoding SCARFACE (SFC) and BYPASS1 (BPS1). SFC encodes an ARF-GAP protein
that is required for normal vesicle trafficking between the plasma membrane and the endosome and is required for normal vein patterning and root growth (Sieburth et al., 2006). BPS1 encodes a plant-specific protein that is required to prevent the root from producing a novel graft-transmissible apocarotenoid that arrests shoot growth (Van Norman et al., 2004; Van Norman and Sieburth, 2007). Both of these RNAs showed decay in the shoot growth (Van Norman et al., 2004; Van Norman and Sieburth, 2007). Both of these RNAs showed decay in the shoot growth (Van Norman et al., 2004; Van Norman and Sieburth, 2007). Both of these RNAs showed decay in the shoot growth (Van Norman et al., 2004; Van Norman and Sieburth, 2007).

Because real-time RT-PCR detects relative mRNA levels, the apparent increase of SFC and BPS1 mRNAs in tdt-1 mutants indicates a slower rate of decay than that of the internal control (ACTIN2). These data indicate a stringent requirement for de-capping for the decay of both of these mRNAs. The decay of SFC and BPS1 mRNA in vcs-7 mutants was also strongly reduced, but not to the same extent as in tdt-1 mutants. The simplest explanation for the decay of these RNAs in vcs-7 is that VCR might support the limited formation of a functional decapping complex in a few specialized cell types.

These data reveal that the requirement for the decapping complex for mRNA decay depends on the specific mRNA.

RNA Cleavage Targeted by miRNAs and trans-Acting Silencing RNAs Does Not Require VCS or TDT

In yeast and animal cells, P-bodies have been implicated in both RNA decay and the sequestration of translationally arrested mRNAs (reviewed in Bruno and Wilkinson, 2006). In animal cells, P-bodies also contain both ARGONAUTE2 (AGO2) and GW182 (Liu et al., 2005a; Behm-Ansmant et al., 2006), where they are likely to function in translational arrest of miRNA targets. Recently, though, GW182 has also been implicated in miRNA-targeted degradation of specific mRNAs (Behm-Ansmant et al., 2006), which suggested that miRNA-directed cleavage of mRNA targets might also occur in P-bodies.

To address whether VCS and/or TDT were required for the normal cleavage of miRNA targets, we used real-time RT-PCR to assess relative levels of several different uncleaved miRNA targets (Figure 5B). We reasoned that if miRNA-directed cleavage was disrupted, then we would observe elevated levels of targets in tdt-1 and vcs-7 mutants. By contrast, we found that vcs-7 and tdt-1 mutants accumulated either similar levels (AGO1 and DCL1) or reduced levels (ARF8 and ARF17) of miRNA targets. The lack of elevated miRNA targets in the mutants indicated that neither miRNA biogenesis nor target cleavage required TDT or VCS.

We also considered ARF3 and ARF4 as possible RNA substrates for VCS and TDT, because these RNAs are regulated by trans-acting silencing RNAs, and this regulation is important for normal leaf development (Adenot et al., 2006; Fahlgren et al., 2006; Garcia et al., 2006). We used real-time RT-PCR to compare the levels of uncleaved ARF3 and ARF4 RNAs and to detect unprocessed TAS3 RNA. We found that the relative levels of these RNAs were decreased in tdt and vcs mutants (Figure 5C). Although it is not readily apparent why these RNAs are reduced, the observation that their levels are not elevated indicates that neither the cleavage of ARF3 or ARF4 target RNAs nor the processing of TAS3 RNA required TDT or VCS.

Because of their roles in leaf and vascular development, we were particularly interested in whether the class III homeodomain leucine zipper (HD-ZIPIII) genes were misexpressed. The HD-ZIPIII RNAs are miRNA targets, and members of this family (PHB, PHV, REV, AthB15/CNA, and AthB8) play both positive and negative roles in vascular patterning (McConnell and Barton, 1998; Emery et al., 2003; Prigge et al., 2004). Real-time RT-PCR analysis revealed that uncleaved HD-ZIPIII RNAs were generally at normal levels, with the exception of REV and AthB8 in tdt-1 (Figure 5D), and we confirmed these results by real-time RT-PCR using alternative primers (also flanking the cleavage site), sequencing reaction products, and by analysis of microarray data.

In these graphs, data for the wild type are represented by black bars, data for tdt-1 by gray bars, and data for vcs-7 by white bars. Four-day-old seedlings are represented in (A), and 3-d-old seedlings are represented in (B) to (D). Error bars indicate SD; n = 4.

(A) RNA decay monitored by analyzing the relative expression of four mRNAs following treatment with the transcription inhibitor cordycepin (0, 45, or 90 min after the addition of cordycepin).

(B) Relative expression levels of select miRNA target RNAs in 3-d-old seedlings grown at 22°C. In tdt-1 and vcs-7, these miRNA target RNAs show levels that are either similar to, or lower than, that of the wild type.

(C) Analysis of two trans-acting silencing RNA targets (ARF3 and ARF4) and the TAS3 RNA.

(D) Relative expression levels of the HD-ZIPIII RNAs in tdt-1 and vcs-7 reveal no consistent pattern of mRNA overaccumulation in tdt-1 and vcs-7 mutants.

**Figure 5.** *vcs* and *tdt* Mutants Show Defects in RNA Decay and Aberrant Patterns of RNA Accumulation.
(see below). The mostly unchanged levels of these miRNA targets further support TDT and VCS not being required for miRNA-directed cleavage of target RNAs. Our data are at odds, however, with a recent analysis by Xu et al. (2006), who found PHB, AthB8, PHV, and REV to be highly elevated in both tdt/dcp2 and vcs mutants. Possible explanations for the difference between our results include developmental timing (our study used 4-d-old seedlings to minimize developmental differences, while the Xu et al. [2006] study used 6-d-old seedlings) or the part of the RNA examined. We assayed uncleaved HD-ZIPIII products, while the portion of these RNAs analyzed by Xu et al. (2006) was not described. It is possible that the cleavage fragments accumulate in tdt and vcs mutants, as 3' fragments of miRNA-cleaved target RNAs have also been shown to accumulate in xrn4 mutants (Souret et al., 2004).

Global Analysis of mRNA Profiles in tdt Mutants

To gain broader insight into how the loss of mRNA decapping affected RNA levels, we performed microarray experiments to compare RNA profiles of tdt-1 and wild-type plants. To minimize the extent to which tdt's developmental defects would confound the results, we used RNA from 3-d-old seedlings. This analysis revealed a large number of mRNAs that were elevated in tdt mutants, and Supplemental Table 1 online lists the 142 mRNAs elevated by fivefold or higher. These elevated RNAs derived from many different functional classes, including signaling molecules, transcription factors, transporters, and metabolism-related. Two classes that were particularly well represented are embryo-expressed RNAs and those encoding heat-shock proteins.

To test whether these microarray results accurately reflected tdt-1 mRNA levels, and to explore whether RNA levels were affected similarly in vcs-7, we performed real-time RT-PCR for six of these elevated mRNAs (Figure 6A). This analysis confirmed that each of these RNAs was elevated in tdt-1 and revealed even higher levels of accumulation than those reported by the microarray. Furthermore, each of these six RNAs was also elevated in vcs-7, although four of them showed more modest increases than that observed in tdt-1.

Molecular characterization of VCS and TDT led to the prediction that accumulated mRNAs would retain their 5' cap. To determine whether this was the case, we assessed whether two of the RNAs that accumulated in tdt-1 and vcs-7 were capped. To do this, we used a modified rapid amplification of cDNA ends (RACE) assay and compared RNA ligation to an RNA primer before and after removal of the 5' cap with Tobacco Acid Phosphatase (TAP) (Gazzani et al., 2004). Ligation products could only be detected in tdt and vcs RNAs that had been treated with TAP (Figure 7), indicating that in the untreated samples, the 5' cap blocked ligation. These data provide strong evidence that VCS and TDT encode components of the mRNA-decapping machinery.

Our microarray analysis of tdt mutants also revealed a set of mRNAs that were present at lower levels in tdt-1, and Supplemental Table 2 online lists the 52 genes whose RNA levels were reduced by fivefold or more. These RNAs correspond to a wide range of functional groups, including signaling, transport (e.g., iron transport–related), and transcription factors. We had not predicted finding this group of severely reduced RNAs, and to characterize them further we examined four in detail using real-time RT-PCR. This analysis confirmed the very low expression of these RNAs in tdt-1 and revealed either similar or more severely depleted RNA levels in vcs-7 (Figure 6B).

We compared the tdt overexpressed and underexpressed mRNAs with the 26 RNAs found to be elevated in xrn4 mutants (Souret et al., 2004; Olmedo et al., 2006). None of the RNAs
elevated in *xrn4* mutants was among our list of fivefold-overexpressed RNAs, and one, At4g00780, was among the *tdt* underexpressed mRNAs. The lack of congruence between gene expression changes in *xrn4* and *tdt* RNAs was surprising, as models for mRNA decay depict these genes as acting in sequential steps of the same pathway (Figure 8).

We noticed that many of the overexpressed mRNAs in *tdt* mutants encode embryogenesis-related proteins, such as Late Embryogenesis Early (LEA) and embryo-expressed heat shock proteins (Hughes and Galau, 1989; Wehmeyer and Vierling, 2000; Schmid et al., 2005). To further explore the connection with embryo-expressed mRNAs, we also analyzed the level of *LEAFY COTYLEDON1* (*LEC1*) RNA. *LEC1* encodes an embryo-expressed transcription factor that is sufficient to induce embryogenesis (Lotan et al., 1998). Using real-time RT-PCR, we found *LEC1* RNA levels to be elevated by 30-fold in *tdt-1* and by 12-fold in *vcs-7*. Elevated levels of translatable *LEC1* in *vcs* and *tdt* mutant seedlings might contribute to the many overexpressed embryo-related RNAs, similar to the misregulation of *LEC1* in *pickle* mutants that has also been linked to the expression of embryonic traits in mutant seedlings (Ogas et al., 1999).

**DISCUSSION**

Signal transduction pathways, whether communicating developmental or environmental information, typically result in altered gene expression. Although most commonly we think of these changes as occurring at the level of transcription, full responsiveness to a new signal is also likely to require removal of the earlier products of gene expression. One example of this comes from controlled proteolysis conferred by F-box proteins, for example the rapid proteolysis of AUX/IAA proteins following auxin binding to the TIR1 F-box protein (reviewed in Quint and Gray, 2006).

RNA decay can also be important for responses to both developmental and environmental signals. The best-known examples come from miRNA-directed RNA cleavage. For example, miR165/166 targets a family of five genes, known collectively as HD-ZIPIIIs, for cleavage. Loss of cleavage due either to mutations at target cleavage sites or to ectopic miRNA expression causes severe developmental defects (McConnell and Barton, 1998; Emery et al., 2003; Prigge et al., 2004; Kim et al., 2005; Figure 7.

**Figure 7.** *tdt-1* and *vcs-7* Accumulate Capped mRNAs.

Capped RNAs were assayed by ligation to a 5’ RNA adapter either before (−) or after (+) treatment with TAP to remove the cap. We detected an abundant product only in the mutants, and only after TAP treatment, confirming that these abundant mRNAs were capped. ctl indicates a reaction without template addition, and (G) indicates genomic DNA supplied as the template.

**Figure 8.** Model for mRNA Bulk Decay Pathways.

Two key features of an mRNA in the cytoplasm are the 5’-5’ 7-methylguanosine (7MG) cap and the 3’ poly(A) tail. Bulk mRNA decay is typically initiated by deadenylation. The deadenylated mRNA can then undergo further decay from its 3’ end by the activity of the exosome. Alternatively, the deadenylated mRNA can have its 5’ cap removed (decapped), followed by further decay from its 5’ end by XRN4 activity.
mRNA Substrates Show Specificity for 3’ and 5’ Decay Pathways

The mRNA bulk decay pathways, worked out in yeast and animals, feature deadenylation followed by either 3’ to 5’ exoribonuclease activity (via the exosome) or decapping, followed by 5’ to 3’ exoribonuclease activity, which in yeast is performed by XRN1 (Figure 8). Our analyses of RNA decay in tdt and vcs mutants found that some mRNAs showed no decay, suggesting obligate decay via decapping. Other mRNAs showed a reduced decay rate, suggesting that these mRNAs normally decay using both pathways, and still others retained normal rates of decay in vcs and tdt, suggesting that their normal decay uses the 3’ to 5’ exosomal pathway. Thus, our data reveal that only some mRNAs show the promiscuous use of both decay pathways and suggest the presence of RNA populations that specifically decay via the 5’ to 3’ and the 3’ to 5’ decay pathways.

How specific mRNA substrates are targeted to a particular mRNA decay pathway is largely unknown. The most likely scenario is that proteins binding to specific mRNA sequences provide targeting. Some conserved sequence elements in 3’ untranslated regions have been observed, for example, in the auxin-induced SAUR genes (McClure et al., 1989), but no obviously conserved sequences were found among the unstable Arabidopsis RNAs (Gutiérrez et al., 2002).

Following decapping, a 5’ to 3’ exoribonuclease leads to RNA hydrolysis (Figure 8). In yeast, this activity is performed by XRN1 (Larimer and Stevens, 1990); however, in Arabidopsis, no XRN1-like gene has been identified, and instead cytoplasmic 5’ to 3’ exoribonuclease activity is thought to be performed by XRN4 (Kastenmayer and Green, 2000). Indeed, several different screens have led to the characterization of xrn4 mutants and have shown that xrn4 mutants accumulate some RNAs (Gazzani et al., 2004; Souret et al., 2004; Olmedo et al., 2006; Potuschak et al., 2006).

This model for bulk decay predicts that vcs, tdt, and xrn4 should accumulate the same set of mRNAs, albeit capped in vcs and tdt mutants and decapped in xrn4. However, we found that tdt/dcp2 mutants had many more RNAs with elevated levels than were found for xrn4 mutants (Souret et al., 2004; Olmedo et al., 2006), and we also found that some mRNAs were at severely reduced levels in tdt, which was not reported for xrn4. The more modest effect on RNA in xrn4 mutants agrees with that mutant’s mild phenotype but is difficult to reconcile with the severe phenotype of decapping mutants and the prevailing model for bulk RNA decay (Figure 8). One possibility is that another, as yet unidentified, enzyme can also carry out 5’ to 3’ exoribonuclease activity. Alternatively, the phenotypic differences might arise due to the presence or absence of the cap; uncapped mRNAs accumulating in xrn4 might be silenced, while capped mRNAs accumulating in vcs and tdt might be translated, leading to secondary effects on gene expression. Finally, the difference might be related to P-bodies themselves. In addition to RNA decay, P-bodies also function in translational regulation (Bren-gues et al., 2005; Liu et al., 2005a, 2005b; Sen and Blau, 2005). In xrn1 mutants, P-bodies appear abnormal, yet they still form, whereas in vcs (and presumably tdt), P-bodies fail to form. If, like animal and yeast cells, plant P-bodies also function in translational regulation, then presumably vcs and tdt, but not xrn4, might regain the expression of translationally arrested mRNAs. This possibility might not only contribute to the severe tdt and vcs phenotypes but also might account for the unexplained reduction of uncleaved mRNA and trans-acting silencing RNA targets found in these mutants. If these RNAs are normally translationally regulated in P-bodies, then the loss of P-bodies in mutants might expose these RNAs to other decay pathways.

Uncovering Natural Variation in mRNA Decay Pathways

Natural variation has been shown for a variety of pathways, including flowering time and light responses (Johanson et al., 2000; Maloof et al., 2001). We found that vcs and tdt phenotypes were both strongly influenced by genetic background. Both vcs and tdt showed a suppressed phenotype when in the Ler accession, and this suppression is conferred by a dominant activity that is genetically unlinked to VCS or TDT.

Why both mutants are suppressed in Ler (but to different extents) remains an outstanding question. We first considered that Ler could have higher activity of genes related to VCS and TDT; however, TDT/DCP2 appears to have no close homologs, and VCR, the closest VCS homolog, is tightly linked to VCS and so inconsistent with an unlinked suppressor. Alternatively, Ler might express higher levels of an alternative scaffold, thus partially restoring decapping activity.

Suppression might also arise due to relief from the symptoms of decapping defects. We found that many capped mRNAs overaccumulate in vcs-7 and tdt-1. If the severe phenotype arises due to these abundant capped mRNAs, then any activity that reduces RNA levels (e.g., increased exosomal activity) might suppress the vcs and tdt phenotypes. It is also possible that only a few of the overaccumulated capped mRNAs cause the severe phenotype, and a suppressed phenotype might arise in a genotype in which these particular genes had lower transcription rates. We also found a set of mRNAs with lower than wild-type levels in vcs and tdt, and it is also possible that these reduced mRNAs are what leads to the severe vcs and tdt phenotypes. If so, perhaps in Ler, expression of these genes is not so severely depressed. We have begun analyzing the Ler suppressors and have found that the major Ler suppressors for vcs and tdt map to different loci.

Vascular Defects in Decapping Mutants

The striking vascular defects shown by both vcs and tdt mutants indicate that vascular patterning requires a normal mRNA decapping complex and presumably mRNA decay. Establishment of vein patterns has been linked to the movement of auxin through responsive cells, although genetic approaches have
also revealed the importance of molecules such as sterols, polyamines, and inositol (1,4,5) triphosphate (reviewed in Fukuda, 2004; Sieburth and Deyholos, 2006). The flexibility of a set of cells to adopt a vascular fate can also be limited by differentiation (Scarpella et al., 2004), and vascular defects in vcs and tdt might also arise due to a lack of responsiveness to inductive signals. In tdt mutants, vascular defects include the vascular transition. The vascular transition is a region within the upper hypocotyl connecting the root-like vascular system of the lower hypocotyl with the shoot’s vascular system. This transition entails both a modification of vein position and a change in cell type organization—the lower hypocotyl shows exarch xylem with an alternating arrangement of xylem and phloem, whereas the shoot has endarch xylem and collaterally arranged vascular tissues (Esau, 1965). The developmental basis for the vascular transition is largely unknown, although proposals include it being simply one part of a single system or the vascular transition being the connection site for two separately established systems. Although careful anatomical analyses using Arabidopsis and other systems have revealed nearly simultaneous differentiation through this zone (Busse and Evert, 1999), the requirement for TDT/DCP2 for the establishment of a normal vascular transition might reflect an initially discontinuous origin of the embryo’s vascular system.

**Developmental Roles for mRNA Decapping?**

The severe seedling defects in vcs and tdt mutants, together with the specificity shown by some mRNAs for decay using decapping, suggest that mRNA decapping might be used for developmental regulation of mRNA levels. Good candidates for developmentally regulated decapping are the heat-stress RNAs that are expressed during normal seed development (Wehmeyer and Vierling, 2000; Kotak et al., 2007). Recently, the levels of three heat-stress RNAs were shown to rise through seed development and then fall precipitously within hours of imbibition (Kotak et al., 2007), suggesting that imbibition might activate an mRNA decay pathway. We found these same three RNAs retained in 3-d-old tdt seedlings (two are listed in Table 1, and Hsp101 was reported as elevated fourfold in our microarray analysis). These results suggest that the imbibition-triggered decline of these RNAs requires mRNA decapping.

RNA decay directed by miRNA and trans-acting silencing RNA has been documented extensively in plants, and in zebrafish, decay via miRNA-directed deadenylation is required for normal progression through embryogenesis (Giraldez et al., 2006). Our data indicate that in plants, the developmental transition to early seedling development appears to be particularly sensitive to mRNA-decapping defects. Future characterization of decapping substrate specificity should help us to distinguish between developmental and housekeeping roles of this RNA decay process.

**METHODS**

**Plant Growth and Microscopy**

Growth of Arabidopsis thaliana plants, tissue preparation for light microscopy, and GUS staining were performed as described previously (Deyholos et al., 2003). The tdt mutant in the Landsberg background was derived from four sequential crosses to Ler, and vcs-7 corresponds to SALK_032031. Oligonucleotide sequences for genotyping are listed in Supplemental Table 3 online.

Confocal analysis for observation of embryo and seedling SAMs followed the aniline blue staining and tissue preparation methods described by Bougourd et al. (2000). The tdt and vcs mutants were easily identified by their misshapen cells. We attribute these misshapen cells to defects arising as a consequence of the tdt genotype, rather than fixation artifacts, because in replicative experiments phenotypically wild-type sibling embryos fixed alongside tdt (within the same tube) looked normal. Confocal analysis for GFP analysis used live roots of 4-d-old seedlings grown vertically on growth medium. To test the effect of cycloheximide, 4-d-old seedlings were incubated in liquid medium (0.5× Murashige and Skoog salts, 1% sucrose, and 0.5 g/L MES, pH 6.8) supplemented with 200 μg/mL cycloheximide (Sigma-Aldrich) and shaken for 80 min. Treatment with MG132 (Sigma-Aldrich) used a 50 μM solution and shaking for 60 min. We used a Zeiss 510 Meta laser scanning confocal microscope as described previously (Sieburth et al., 2006).

Analysis of transition zone defects in tdt-1 grown at different temperatuers was performed on cleared tissue examined under a dark field using magnification up to ×90. The transition zone was considered defective if the primary vein of at least one cotyledon did not have differentiated xylem that connected with the hypocotyl vasculature.

Seedling images were obtained using an Olympus SZX12 microscope. Vascular patterns were observed using dark-field base, and tissue was fixed in Carnoy’s (3:1 ethanol:acetic acid) and cleared by incubation in saturated chloral hydrate (Sigma-Aldrich). Details of cellular anatomy and GUS staining patterns were examined using an Olympus BX-50 microscope.

**Isolation of tdt and Molecular Analyses of Mutants**

The tdt-1 mutant was isolated in a small pilot screen of T-DNA-mutagenized Col-0 seeds. This mutant was selected for analysis because it lacked leaves and its cotyledons showed a forked vein pattern. Subsequent analysis showed that the lesion was not linked to the T-DNA, so conventional mapping approaches were used to identify the TDT gene. The sequences of mapping primers are available upon request. The SALK lines corresponding to tdt-2 and vcs-7 were sequenced to verify their T-DNA insertion sites. We sequenced the entire gene from tdt-1, and we identified only one lesion, a 50-nucleotide deletion that was also readily detectable by PCR of genomic DNA.

**Yeast Two-Hybrid Analysis**

Directed yeast two-hybrid analyses followed the Clontech protocols. Full-length coding regions of VCS and TDT were amplified from seedling cDNA, sequenced, and cloned into both bait (pGADT7) and prey (pGBKT7) vectors.

**Quantitative RT-PCR Analyses**

We isolated total RNA using the Agilent RNA Isolation Mini kit or the Qiagen RNeasy Plant Mini kit (for the cordycepin experiments) and performed randomly primed cDNA synthesis with the Promega reverse transcription system. We performed real-time RT-PCR using the Roche Lightcycler and FastStart DNA Master SYBR Green 1 master mix. Each real-time RT-PCR experiment included internal controls (ACTIN2) for each template. Data shown are averages of four to six reactions and were confirmed by independent experiments. HD-ZIPIII expression was also confirmed by reactions using separately designed oligonucleotides. To determine the relative steady state level of RNAs, we calculated the relative expression $\frac{\text{surface area of} \ 	ext{actin} \ 	imes \text{target}{\text{actin}}}{\text{average} \ 	ext{ACTIN2} \ 	imes \text{target}{\text{ACTIN2}}}$ and converted this number to a percentage. We tested each reaction’s specificity using the machine’s standard melt curve method and also
analyzed product size by gel electrophoresis. Sequences of oligonucleotides used as primers are available in Supplemental Table 3 online. Some primer sequences were taken from Vazquez et al. (2004), Vaucheret et al. (2004), and Li et al. (2005); these primers are listed in Supplemental Table 3 online, along with sequences of primers designed by us.

**RACE Analysis of Capped mRNAs**

To determine whether accumulating mRNAs were capped, RNA ligase-mediated (RLM) RACE was performed using the First Choice RLM-RACE Analysis of Capped mRNAs kit from Ambion. Total RNA was isolated from 4-d-old Arabidopsis seedlings (Qiagen RNeasy kit). Col-0, tdt-1, and vcs-7 RNAs (1 μg) were ligated to the 5′ RACE Adapter (5′-CUGAUUGCGAUAUGAA-CACUGCGUUUGCCUUGAUAU-3′) either prior to or following treatment with TAP to remove the 5′ cap. Random decamers were used to prime the reverse transcription reaction, and PCR was performed using a primer specific to the RACE adapter (5′ RACE Outer Primer) and gene-specific primers.

**GFP-TDT Fusion and TDT Rescue Constructs**

The TDT rescue construct encompassed a 6019-bp PstI-EcoRI fragment from BAC T8114, which was cloned into pCAMBIA1300. The plant DNA contained the entire At5g13570 coding region, the entire upstream (5′) and downstream (3′) regions, and small portions of the upstream and downstream genes. The GFP:TDT construct was made by placing the TDT cDNA coding region (clone U61209) between the EcoRI and BamHI cloning sites of pEGAD (Cutler et al., 2000). Constructs were sequenced prior to transformation into plants. Both constructs were introduced to wild type (Col-0) and tdt-1 heterozygotes using the floral dip method (Clough and Bent, 1998). Transgenic seeds were isolated based on their resistance (hygromycin for the pCAMBIA vector and BASTA for the pEGAD vector), and the GFP:TDT lines were introduced into the vcs-7 mutant background by standard genetic methods.

**Analysis of tdt Rescue**

We analyzed nine independently generated lines that segregated for tdt, and all showed similar ratios of phenotypically normal to mutant plants (~15:1). Data presented in Results are from transgenic rescue line 29.

**Microarray Analysis**

Total RNA was isolated from 3-d-old wild-type and tdt-1 seedlings using the Agilent Plant RNA Isolation Mini kit. RNA samples (1 μg) were submitted to the University of Utah Microarray Core Facility and went through two steps of RNA quality control prior to the labeling protocol, which consisted of analysis of the sample on an Agilent bioanalyzer and validation of nucleic acid concentration on a NanoDrop spectrophotometer. Samples were labeled using the Agilent Low RNA Input Linear Amplification kit. RNA labeling involved first-strand reverse transcription primed using an oligo(dT)/T7 polymerase promoter sequence, second-strand cDNA synthesis, and generation of fluorescently labeled cRNA in an in vitro transcription reaction performed in the presence of Cy3-CTP or Cy5-CTP. Labeled samples were hybridized to the Arabidopsis 3 oligo expression array (44K) using the Agilent Gene Expression Hybridization kit and were incubated overnight (~17 h) at 60°C. The microarray slide was washed (6 × SSC [1 × SSC is 0.15 M NaCl and 0.015 M sodium citrate] and 0.005% lauryl sarcosine at room temperature for 1 min, 0.6 × SSC and 0.005% lauryl sarcosine at room temperature for 1 min, and a 5-s acetonitrile dip) and scanned using the Agilent G2565BA microarray scanner. Three biological replicates were performed, with a total of three arrays hybridized. One of these arrays was hybridized with reversed dyes.

Data from the scanned microarrays were obtained using Agilent Feature Extraction Software, version 8.5 (http://www.chem.agilent.com/Scripts/PDS.asp?iPage=2547). The quality of the data was assessed by visual inspection of M versus A plots. Some intensity-dependent bias was noted and was corrected in the Lowess-normalized data set generated by Agilent Feature Extraction. Control spots and spots for which there were data from two or fewer arrays were filtered from the data set. All pairwise differentially expressed genes were identified using SAM software (Tusher et al., 2001) using the data from the remaining 34,347 Arabidopsis probe sets. A false discovery rate parameter of 1% was used for the one-class SAM analysis. Average fold change for each gene was calculated from the log ratio data. Genes from the array were cross-referenced to The Arabidopsis Information Resource (http://www.Arabidopsis.org) using systematic names provided by Agilent. A single annotation was assigned to each gene using The Arabidopsis Information Resource annotations and the Gene Ontology biological process. All generated annotations were manually curated. All data from microarray experiments have been deposited in the Gene Expression Omnibus database under accession number GSE7359 (www.ncbi.nlm.nih.gov/geo).

**Accession Numbers**

Sequence data from this article can be found in the GenBank/EMBL data libraries under the following accession numbers: TDT (At1g53150), VCS (At3g13300), VCR (At3g13290), ACT2 (At3g18780), LEC1 (At1g21970), LEA (At1g52690), Hsp17.4 (At3g46250), At HSP90.1 (At5g52640), At HSP70 (At2g32120), HSP26.5 (At1g52560), HSP23.5-M (At5g51440), ARF4 (At3g60450), ARF3 (At2g33860), TAZ3 (At3g17185), PHB (At2g34710), REV (At5g06690), AthB15/CNA (At1g52150), PHV (At1g30490), AthB8 (At4g32880), SFC (At1g33000), BPS1 (At1g01550), BME3 (At3g54810), MYB4 (At1g22640), DCL1 (At1g91040), ARF8 (At5g37020), ARF17 (At1g77850), and AGO1 (At1g48410). Microarray data have been deposited in the Gene Expression Omnibus site with accession number GSE7359.

**Supplemental Data**

The following materials are available in the online version of this article.

Supplemental Figure 1. vcs-7 Is an RNA Null Allele.

Supplemental Figure 2. Phenotypes of vcs-1, vcs-7, and tdt-1.

Supplemental Figure 3. Low-Temperature Growth Confers Minimal Suppression to the vcs-7 Allele.

Supplemental Figure 4. Pinched Appearance of tdt Cotyledon Petioles.

Supplemental Table 1. Genes with at Least a Fivefold Increased RNA Level in tdt-1 Mutants.

Supplemental Table 2. Genes with at Least a Fivefold Decreased RNA Level in tdt-2 Mutants.

Supplemental Table 3. Oligonucleotide Sequences of Primers Used for PCR, Real-Time RT-PCR, and RACE in This Study.

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REFERENCES


the miRNA pathway are crucial for plant development. Genes Dev. 18: 1187–1197.


Components of the Arabidopsis mRNA Decapping Complex Are Required for Early Seedling Development

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