Athila4 of *Arabidopsis* and *Calypso* of Soybean Define a Lineage of Endogenous Plant Retroviruses

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The *Athila* retroelements of *Arabidopsis thaliana* encode a putative *envelope* gene, suggesting that they are infectious retroviruses. Because most insertions are highly degenerate, we undertooka comprehensive analysis of the *A. thaliana* genome sequence to discern their conserved features. One family (*Athila4*) was identified whose members are largely intact and share >94% nucleotide identity. As a basis for comparison, related elements (the *Calypso* elements) were characterized from soybean. Consensus *Calypso* and *Athila4* elements are 12–14 kb in length and have long terminal repeats of 1.3–1.8 kb. Gag and Pol are encoded on a single open reading frame (ORF) of 1801 (*Calypso*) and 1911 (*Athila4*) amino acids. Following the Gag-Pol ORF are noncoding regions of ∼0.7 and 2 kb, which, respectively, flank the *env*-like gene. The *env-like ORF* begins with a putative splice acceptor site and encodes a protein with a predicted central transmembrane domain, similar to retroviral *env* genes. RNA of *Athila* elements was detected in an *A. thaliana* strain with decreased DNA methylation (*ddm1*). Additionally, a PCR survey identified related reverse transcriptases in diverse angiosperm genomes. Their ubiquitous nature and the potential for horizontal transfer by infection implicates these endogenous retroviruses as important vehicles for plant genome evolution.

Retrotransposons and retroviruses (collectively referred to as retroelements) replicate by a common mechanism of reverse transcription (for review, see Coffin et al. 1997). Retroelement genomes are delimited by direct long terminal repeats (LTRs), and they encode *gag* and *pol* genes, whose products form a particulate replication intermediate wherein reverse transcription takes place. The primary distinguishing feature between the retrotransposons and retroviruses is that the latter have a third gene called *envelope* (*env*). *env* encodes a transmembrane protein that associates with the cell membrane. The replication intermediate buds from the cell as a membrane-bound virion, and Env extends from the virion surface and interacts with cellular receptors to mediate infection.

Phylogenetic relationships based on reverse transcriptase amino acid sequences identify six distinct lineages of retroelements (Xiong and Eickbush 1990; Malik 2000). One of these—the vertebrate retroviruses—encodes *env* genes and is infectious. The five remaining groups are comprised mostly of retrotransposons and include the well-studied Ty1-*copia* (*Pseudoviridae*) and Ty3-*gypsy* (*Metaviridae*) elements (van Regenmortel et al. 2000), the so-called DIRS1 and BEL groups, and the caulimoviruses (Malik et al. 2000). With the exception of the caulimoviruses and the sparsely populated DIRS1 group, some members of each lineage encode open reading frames (ORFs) with *env*-like features—most notably transmembrane domains. These include a large number of invertebrate Ty3-*gypsy* elements (e.g., *gypsy*, *17.6*, *297*, and *ZAM* from *Drosophila melanogaster*; *TOM* from *Drosophila ananassae*; *TED* from *Trichoplusia ni*; *Yoyo* from *Ceratitis capitata*; for review, see Lerat and Capy 1999), two Ty1-*copia* elements from

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plants (i.e., *SIRE-1* from *Glycine max* [soybean] and *Endovir* from *A. thaliana*; Laten et al. 1998; Kapitonov and Jurka 1999; Peterson-Burch et al. 2000), and several BEL group elements (e.g., *Tas* from *Ascaris lumbricoides* and *Cer7* from *Caenorhabditis elegans*; Felder et al. 1994; Bowen and McDonald 1999). Analyses of *env*-like genes from the various retroelement groups suggests that *env* was independently acquired from viruses multiple times during evolution. The *env*-like ORFs of several insect Ty3-*gypsy* elements are closely related to *env* of the bacculoviruses, and for some *Cer* elements, the *env*-like gene is related to *env* of the phleboviruses (Malik et al. 2000). Despite the widespread presence of *env*-like ORFs and their similarity to known viral *env* genes, *gypsy* of *D. melanogaster* is the only known retroelement outside of the retroviruses for which Env is known to play a role in infection (Kim et al. 1994; Song et al. 1994).

In our analysis of the *A. thaliana* genome sequence, we determined that *Athila*—a degenerate, centromere-associated retroelement (Pelissier et al. 1995, 1996; Copenhaver et al. 1999)—is a Ty3-*gypsy* group retrotransposon with an *env*-like ORF (Wright and Voytas 1998). A related element was also described in *Pisum sativum* (pea) called *Cyclops-2* (Chavanne et al. 1998). Because *Cyclops-2* was less degenerate than *Athila* and prevalent in related legumes, we sought potential functional homologs in soybean. The soybean elements, called *Calypso*, encode an *env*-like gene that shares 29% amino acid identity to the corresponding gene of *Cyclops-2* (Peterson-Burch et al. 2000). This suggests that the *env*-like ORF has evolved under functional constraint and likely plays a role in the life cycle of these elements. For simplicity, we refer to *Athila* and related retroelements as endogenous retroviruses, with the understanding that the biological role of their *env*like genes remains to be determined. The sequence degeneracy of the endogenous plant retroviruses described to date has frustrated attempts to define their structural features. However, further characterization of the soybean *Calypso* elements and completion of the *A. thaliana* genome sequence has enabled us to construct consensus elements that likely approximate functional elements. Here we report a detailed

description of these endogenous retroviruses and provide evidence of their widespread distribution in higher plants.

RESULTS

Athila Elements of *A. thaliana*

To further characterize the *A. thaliana Athila* elements, reverse transcriptases from all Ty3-*gypsy* elements were recovered from the *A. thaliana* genome sequence (Initiative 2000). BLAST searches (Altschul et al. 1990) were performed with reverse transcriptases from *Athila1-1*, *Tat4-1*, and *Tma3-1*, three divergent *A. thaliana* Ty3-*gypsy* elements (Fig. 1; Wright and Voytas 1998). Additional BLAST searches were performed with the most divergent retroelement sequences recovered. A total of 191 unique reverse transcriptases were identified. These were aligned, and when necessary, conservative changes were made to correct frameshift mutations. A phylogenetic tree was generated by the neighbor-joining method (Fig. 1; Saitou and Nei 1987). The elements clustered into three distinct clades designated the classic, *Tat*, and *Athila* lineages.

Phylogenetic analysis further resolved the *Athila* elements into clades, which we designated as distinct families (Fig. 1). These included the previously described *Athila1* family (Wright and Voytas 1998) and six additional families, designated *Athila4-Athila9*. The *Athila*, *Athila2*, and *Athila3* families are not included in the tree, because they have deletions of reverse transcriptase (Pelissier et al. 1995; Wright and Voytas 1998). Elements in four of the seven families had potential coding regions flanking reverse transcriptase and discernible LTRs (*Athila1*, *Athila4*, *Athila5*, and *Athila6*). Relatively intact insertions were given species designations (e.g., *Athila1-1*, Fig. 1). The *Athila4* family was the largest and included 22 members. Six of these (designated *Athila4-1* to *Athila4-6*) were ∼14 kb in length and had LTRs of ∼1.8 kb (Fig. 2). *Athila4-3* and *Athila4-4* were organized in tandem and shared a central LTR. The tandem *Athila4-3*/*Athila4-4* insertion and the individual *Athila4* elements were flanked by 5-bp target-site duplications (data not shown). In pairwise comparisons, the six *Athila4* elements averaged 94% nucleotide identity across their entirety. Despite this high degree of sequence identity, *gag* and *pol* were broken by stop codons and frameshifts.

Calypso Elements of Soybean

In the initial description of the *Cyclops-2* element from pea, related DNA sequences (based on Southern hybridizations) were found to be abundant in other legumes, including soybean (Chavanne et al. 1998). *Cyclops-2* homologs were recovered from soybean by screening a genomic λ phage library using the *Cyclops-2* reverse transcriptase as a hybridization probe. Sixty-three hybridizing phage were characterized, 35 of which were unique based on restriction endonuclease mapping (data not shown). Each of these latter clones was partially sequenced, and 24 had identifiable amino acid sequence similarity to *Cyclops-2* and *Athila* (data not shown)*.* The coding regions of these 24 elements, however, were replete with stop codons, frameshifts, deletions, and insertions. Five of the least degenerate elements (designated *Calypso1-1*, *Calypso2-1*, *Calypso3-1*, *Calypso4-1*, and *Calypso5-1*) were sequenced (Fig. 3). Despite being highly degenerate, each had discernable features such as LTRs and coding regions with similarity to *gag*, *pol*, and the *env*-like gene of *Cyclops-2*. In the case of *Calypso2- 1*, the 5' LTR depicted in Figure 3 is the 3' LTR of a second *Calypso* element that inserted within *Calypso2-1*. *Calypso5-1*

Figure 1 A neighbor-joining tree of reverse transcriptases from *Arabidopsis thaliana* Ty3/*gypsy* retroelements. The tree was generated from amino acid sequences encompassing the seven conserved domains that define reverse transcriptase (Xiong and Eickbush 1990). Each major group (classic, *Tat*, and *Athila*) is labeled. Numbers along the branches indicate bootstrap support for 100 replicates. Arrows indicate the most recent common ancestor for each of the three lineages. Reverse transcriptases from *Tma3-1*, *Tat4-1*, and *Athila1-1* were used in BLAST searches and are noted with an asterisk. BAC numbers for reverse transcriptases are in clockwise order, beginning with the branch marked number 1: T5E15, F23C08-2, MTO24, T16I21, T12H3, F23H14-4, T5M2/T17H1, F23H14-3, F9B22, T12K4, T14K23, T26P13-1, T5E15-2, T14A11, F13E17, F3D18, *Tma3-1* 9D12, MKD10, T1J24, MXE2, F28I8, T13P21/T6B13/F26C24-1-1, F23H14-4-4, F7F23, T5L23, T13H18/F3K12/F14P14-1, F9M13-2, F3H7, MJI6-1, F8D11, T32N15, T32B20, F17L24-2, F28J9-1, F5J5, F5K24, T14K23-2, T7B9-2, T10P12-3, F14I23, F11I2, T1O13, T29A4- 3, F17L24, F19C17, T14A11-5, T27D20, T9F8/T4E14, T7B9-3, F15O4-2, MYM9, F7P1-2, F22D12, K11J14-2, F9M13, F6H5-2, T17A11/F15O11, T12K4-3, MTO24-2, T32E20, F12P23/T4D8, T10P12, T21C14, T13H18/F3K12/F14P14-2, F18P9, MJI6-2, F23H6, T8M17, F7K2, MOD1, T13P21/T6B13/F26C24-2, F26G5-2, T5N23, T21B14, K11J14, F1D9, MXC9, MVA11, F16J10/T20G20/T3P4, F28I8/F1O13, T17A11/F15O11-2, F19I11, T5L23, F6H5, F7M19, F1M23-4, F7I20, T24G3, F8N14, F8N14-2, F23M2, T15J14/F15A23, T4P3, T1O16/T13P21-1, *Tat4-1* MXA21, T24H24, T12O21, F26H6, F7K15, F4M19, F5M15, T19K21/T17A11, F23H14-1-1, F12P23, T15D5, T19K21/T17A11-2, K11J14, T17A11/F15O11, F28H19, T24H24, F9M13-1, T27D6, F28L22, T7B9-1, F24C20/T27D6, F6H8, F15O11/F14O4/T26C18, F23C08-2, F3D18, F28L21, F1M23-2, F23C08, T13P21, F19N02, T5H22, F1M23-3, F17M07, F14G16, MIL15, F10A2, K2K18, T6L9, T3J11, F15K19/T13H18, *Athila1* (*Athila1-3* F4I4, *Athila1-1* MXI10), *Athila6* (F13C19, T32E20-2, F21I2- 2, F7I20, F23H6-2, F5K7/F18P14/F28N16, F3D18-2, F26B15, F9M8, *Athila6-2* F28N5, *Athila6-1* T9E19), *Athila4* (T32O2-2, T25N22/ T13E11, *Athila4-3* F7F22, *Athila4-4* F7F22, F28L21, F9M8-2, *Athila4-1* F1404, F7P3, *Athila4-2* MED5, *Athila4-10* T26P13, *Athila4-7* F7B19/ T15D2, T32O2, F10A2, *Athila4-6* F1809, *Athila4-9* T15D2 *Athila4-5* F7K15, T26P13-2, T29A4, MIF6, T14A16, F21I2, *Athila4-8* T6L9, F7B19/T15D9), *Athila9* (T13D4,T13H18/F3K12/F14P14, F28H19, *Athila9-1* T24G23, T7B9, F14C23, F10A2-2, T27B3), *Athila8* (F17L24/ F9B22), *Athila7* (T3B16, F21A17), *Athila5* (F1M23, T22B15, T19K21/ T17A11, *Athila5-1* F23M2, F23C08, *Athila5-1* F14B16, T5E15-3, T32A11, T18O6/F7E22/F23M2, F11B20).

Figure 2 Structural organization of *Arabidopsis thaliana Athila4* elements. Boxes with filled triangles represent LTRs. Open boxes represent coding sequences, and they are offset to indicate changes in reading frame. Vertical thin lines represent stop codons. Horizontal thin lines represent noncoding sequences. The shaded region identifies the coding region for reverse transcriptase. Shaded boxes indicate the putative *envelope* (*env*) gene. The accession number, the BAC designator, and the position within the BAC for each *Athila4* element are as follows: *Athila4-1*: AC007209, F1404, 33315 to 47208; *Athila4- 2*: AB026642, MED5, 3448 to 17452; *Athila4-3* and *Athila4-4*: AC007534, F7F22, 88613 to 114709; *Athila4-5*: AL353871, F7K15, 86117 to 99436; *Athila4-6*: AF296831, F1809, 38836 to 52851.

contained an insertion within its reverse transcriptase of 1.8 kb, with flanking 5-bp target-site duplications and end sequences suggesting it is a retroelement solo LTR (Fig. 3; data not shown). Despite the high level of sequence degeneracy, the reverse transcriptases of the five *Calypso* elements shared, on average, 81% amino acid identity.

Features of *Athila4* and *Calypso* Elements

For most retroelements, the region adjacent to the 5' LTR is complementary to a cellular tRNA and serves as the site for priming minus-strand DNA synthesis. The primer binding site (PBS) of *Athila4* and *Calypso* is complementary to the 3' end of the aspartic acid tRNA for the GAC codon from *A. thaliana* and soybean (Fig. 4a; Waldron et al. 1985; Wright and Voytas 1998). Complementarity begins at variable positions from the boundary of the 5' LTR, and extends for 13 bases for the *Athila4* elements and for 18 or 19 bases for given *Calypso* elements. For most retroelements, a stretch of purines adjacent to the 3' LTR serves as the priming site for plus-strand DNA synthesis. A polypurine tract (PPT) is found at this location in *Athila4* and *Calypso*, and all of the endogenous plant retroviruses share a conserved core consensus sequence (TTTGGGGG), as well as less conserved flanking sequences (Fig. 4B). A second PPT motif (PPT1) is found after the *env*-like gene. The two PPTs delimit a large noncoding region, which in *Athila* averages ∼2 kb in length (see Figs. 2, 3). A second noncoding region lies between *gag-pol* and the *env*-like gene and approximates 0.7 kb.

Because of the large number of frameshifts and stop codons in *Calypso* coding sequences, a quasiconsensus *Calypso* element was generated. Additionally, a strict *Athila4* consensus sequence was generated, which was possible because of the high degree of sequence homogeneity. Figure 5A depicts the structural organization of these consensus elements, as well as *Cyclops-2* from pea (Chavanne et al. 1998) and three partially sequenced homologs: *Diaspora* from soybean, *BAGY-*2 from barley (Shirasu et al. 2000), and a degenerate element from rice that we identified from the rice genome sequence data. The consensus *Athila4* and *Calypso* elements encode Gag and Pol on a single ORF of 1911 and 1801 amino acids, respectively. These coding regions were aligned with Gag-Pol of *Cyclops-2*, and the percent amino acid identity was plotted along their entirety (Fig. 5A). The first third of the ORFs shares ∼20% amino acid identity, and we define this region as Gag (∼600 amino acid [aa], Fig. 5A). The *Calypso* and *Cyclops-2* Gag proteins encode a conserved finger domain characteristic of retrotransposon and retroviral nucleocapsid proteins (Fig. 5B). This motif is not present in any of the other elements examined. A block of ∼110 amino acid residues is conserved near the N terminus of Gag, suggesting a conserved function. Similarity to this region can be detected in the sequence of *Diaspora* and the rice element but not in *BAGY*-2 (data not shown).

Following Gag is a motif (LI/CDLGA) that we believe is the active site of an aspartic acid protease (Fig. 5B). We define protease as the region of ∼40% amino acid identity that spans

> 10 11

12 13 kb

Figure 3 Structural organization of the *Calypso* elements. Elements are depicted as described in the Fig. 2 legend. A *Calypso* element has inserted into *Calypso2-1*. The left-most LTR depicted belongs to this element; whereas, the right LTR belongs to *Calypso2-1*. *Calypso5-1* has what appears to be a solo *Calypso* LTR within reverse transcriptase, which is indicated by the thick horizontal line.

А	PBS		PPT ₁	PPT
$Athi1a1-1$ $Athila2-1$ $Athi1a4-1$ Athila5-1 Athila6-1 Calypsol-1 Calypso5-1 Cyclops ₂ Rice $BAGY-2$	CA]AUUUGGCGCCGUUGCCGGUUGCAAUAG CA]AUUUGGCGCCGUUGCCAAUUGGGUGUUU CA]AUUUGGCGCCGUUGCCAAUUGGGUGUUU CA1AAUUUGGCGCCGUUGCCGAAGUUCUAUUU CA] AUUUGGCGCCGUUGCCAAUUGGGUGUUU CA1AGUUUUUGGCGCCGUUGUCGGGGAUUUUGUU CA]UUUAUGGCGCCGUUGUCGGGGAACUGGAU CAIAAADGGCGCCGUUGCCGGGGAUUGUGAA CA1UUUCUGGCGCCGUUGUCGGGGAACGGUUG UA AGACUCUUUUCUGGCGCCGUUGCCGGGGAGGGAAGC	$Athilal-3$ $Athi1a2-1$ $Athila3-1$ Athila4-1 $Athila5-1$ Athila6-1 Calypso2-1 Calypso2-1#2 Calypso4-1 Cyclops ₂ Rice	CATGCTCTTTTTGGGGGAGGTG TGATTTAAGTTTGGGGGAGGGC TGATTTAAGTTTGGGGGAGGGT TGATTTAAGTTTGGGGGAGGGC and the contract of the second contract of the contract of the contract of the contract of the contract of the TGATTTAAGTTTGGGGGAGGGC AGGTCTTAATTTGGGGGATTTT AGGCCTTATTTTGGGGGATTTT TGTTCCAAATTTGGGGGAGACA TTATGTACTTTTGGGGTGTTTT TTGAATAAGTGTGGGGGAGATA	GATAAGTTTGGGGGAGT. [TG ACTAAGTTTGGGGGAGT. [TG ATTAAGTTTGGGGGAGT. [TG ACTAAGTTTGGGGGAGT. ITG GCTAAGTCCGGGGGAGT. [TG ACTAAGTTTGGGGGAGT. ITG GTTTCATTTGGGGAATTT [TG GTTTAATTTGGGGGATTT [TG TTTAAATTTGGGGGAGTA [TG AATCTAGTTGGGGAGAGT [TG CTAAGTGTGGCGGGGTTT [TG
Athila4-1	CA1AUUUGGCGCCGUUGCCAAUUGGGUGUUU			
ASP tRNA	ACCGCGGCAACGGCCCCUAGCUUGGGCCC			
	Calvpso1-1 CAlAGUUUUUGGCGCCGUUGUCGGGGAUUUUGUU			

Figure 4 Priming sites for reverse transcription. LTR sequences are underlined and square brackets denote the LTR boundaries. (*A*) The primer binding site (PBS) is complementary to the 3' end of the ASP tRNA. Complementary sequences are shaded, including those that form G:U base pairs. (B) Polypurine tracts following the *envelope* (env)-like ORF (PPT1) and near the 3' LTR (PPT) are characterized by the shaded, conserved core motif.

∼300 amino acid residues between Gag and reverse transcriptase (depicted in light gray, Fig. 5A). Although we do not know the precise boundaries of protease, this region is considerably larger than the proteases of retrotransposons and retroviruses (e.g., 181 aa for Ty1, 99 aa for HIV; Merkulov et al. 1996; Coffin et al. 1997). Following protease is ∼520 amino acids that comprise reverse transcriptase. Reverse transcriptase shares ∼68% amino acid identity among elements. All seven conserved amino acid sequence domains characteristic of retroviral and retrotransposon reverse transcriptases are

Figure 5 Gag-Pol of the *Athila* group elements. (*A*) The structural organization of the *Athila4* and *Calypso* consensus elements are shown along with individualrelated elements from pea (*Cyclops-2*; Chavanne et al. 1998), barley (*BAGY-2*; Shirasu et al. 2000), rice (accession number AP003054, 36238–53391 bp; an inserted element was removed for the diagram), and soybean (*Diaspora*, accession number AF095730). Below *Cyclops-2* is a graph depicting amino acid identity along the length of Gag-Pol. Light gray depicts protease (PR), medium gray depicts reverse transcriptase (RT), and dark gray depicts integrase (IN). All other aspects of the figure are as described in the Fig. 2 legend. (*B*) Amino acid sequence signatures of Gag-Pol. The sequence domains are described in the text. Motifs are shown that define the seven conserved domains of reverse transcriptase (Xiong and Eickbush 1990). For integrase, the zinc-binding domain is shown, as are signatures for the DD35E domain (Fayet et al. 1990) and the GPY/F motif (Malik and Eickbush 1999).

evident (depicted in gray, Fig. 5A). The remainder of Gag-Pol constitutes an ∼450 amino acid integrase (depicted in dark gray, Fig. 5A). In addition to the conserved N-terminal zincbinding motif and the DD35E motif of the catalytic domain, integrase has a C-terminal extension with a GPY/F module (Fig 5B; Malik and Eickbush 1999). The GPY/F module is found in some retroviral and Ty3/*gypsy* element integrases and is thought to bind DNA. Integrase shares ∼64% amino acid identity among *Athila4*, *Calypso*, and *Cyclops-2*.

Features of the *env*-like Gene

After *gag* and *pol* and between the two noncoding regions, the *Athila4* and *Calypso* consensus elements encode ORFs of 619 and 420 amino acids, respectively (Fig. 6A). Recognizable *env*like ORFs are also found in members of the *Athila*, *Athila1* thru *Athila6*, and *Athila9* families (data not shown). The *env*-like ORFs of *Athila2*, *Athila3*, *Athila4*, and *Athila6* share an average of 69% amino acid sequence identity in pairwise comparisons (data not shown). The *Athila1* and *Athila5* elements are divergent (Fig. 1), and their *env*-like ORFs do not align well with the other *Athila* families. The consensus *Calypso env*-like gene shares 29% amino acid sequence identity to the *env*-like gene of *Cyclops-2* (Peterson-Burch et al. 2000). Between the pea/ soybean and *A. thaliana* elements, no significant amino acid sequence similarity was observed.

Retroviral Env proteins are typically transported through the endomembrane system, where they are proteolytically cleaved to generate surface (SU) and transmembrane (TM) proteins prior to being released on the cell surface (Coffin et al. 1997). Targeting to the endomembrane system is mediated by a signal sequence at the N terminus of Env. The N termini of the *Calypso* and *Cyclops-2* Env-like proteins are basic in nature (Fig. 6B). Additionally, the N termini of *Athila4* and *Cyclops-2* are serine-rich. The program PSORT predicts a variety of destinations for the Env-like proteins within the cell (Nakai and Kanehisa 1992). The most confident predictions are for *Calypso2-1* and *Athila4-1*, which suggest targeting to the plasma membrane (70% confidence) and endoplasmic reticulum (85% confidence), respectively.

At the cell surface, the retroviral TM protein spans the plasma membrane. We previously reported a predicted trans-

Figure 6 Features of the *Athila* group *env*-like ORFs. (*A*) Generalized organization of *env*-like ORFs from the *Arabidopsis thaliana Athila* group elements, the soybean *Calypso* elements, *Cyclops-2* of pea, *gypsy* of *Drosophila melanogaster*, and HIV1. The open boxes indicate ORFs. Arrowheads indicate signal sequences. Black boxes indicate transmembrane domains. Vertical lines within boxes denote stop codons. The first methionine within each ORF is indicated by a short line. (B) Amino acid sequences of putative signal sequences at the N termini and predicted transmembrane domains. (*C*) TMpred output for the *Athila4* consensus *envelope* (*env*)-like ORF with and without a frameshift at the C terminus. Values above 500 (on the *X*-axis) are significant and indicate likely transmembrane domains (Hofmann and Stoffel 1993). The *Y*-axis indicates amino acid sequence position. (*D*) Sequences at the putative splice acceptor site of the *Arabidopsis thaliana Athila* and *Calypso* elements. The confidence level indicates the output for NetGene 2 (Brunak et al. 1991; Hebsgaard et al. 1996). The first methionine in the *env*-like ORF is in bold.

membrane domain in the *env*-like ORFs of several *Athila* elements (*Athila*, *Athila1*, *Athila2*, and *Athila3*, Wright and Voytas 1998). The *Athila4* consensus *env*-like ORF also encodes a transmembrane domain (TM1, Fig. 6A–C), to which the program TMpred assigns a score of 2006 (scores above 500 are considered significant; Hofmann and Stoffel 1993). Similarly, a transmembrane domain is predicted near the center of the *Calypso env*-like ORF (TMpred value of 947; Fig. 6A,B and data not shown). The *Cyclops-2 env*-like protein has a potential transmembrane domain at a similar location, but at a reduced confidence level relative to the other elements (TMpred value of 650; Fig. 6A,B and data not shown).

In our analysis of the *Athila4 env*-like gene, we noticed the potential to encode additional transmembrane domains after the stop codon. Strong transmembrane domains were predicted in either the same frame as the *env*-like ORF (TM2, Fig. 6A–C) or in the +1 frame (TM3, Fig. 6A–C). These potential coding regions extend the *env*-like ORF to the first polypurine tract (PPT1) and are conserved among some element families (Fig. 6B). Small ORFs with predicted transmembrane domains are also found at the end of the *Calypso* and *Cyclops-2 env*-like ORFs. In the consensus *Calypso* element, the ORF is in $a - 1$ frame, although the degree of degeneracy among *Calypso* elements reduces confidence in this reading frame assignment. Unfortunately, sequences between *Athila* families were too divergent to ascertain whether the short ORFs are evolving as coding sequences based on frequencies of synonymous versus nonsynonymous substitutions (data not shown).

Retroviral *env* genes are typically expressed from a spliced, subgenomic mRNA (Coffin et al. 1997). The *Calypso env*-like ORF has a predicted splice-site acceptor sequence located at the first methionine, to which the program Net-Gene2 assigns a confidence level of 100% (Fig. 6D; Brunak et al. 1991; Hebsgaard et al. 1996). Although there are other favorable splice acceptors in the vicinity of the *Calypso env*like ORF, only the putative acceptor at the first methionine is conserved (Fig. 6D). For the *Athila* elements, a number of possible splice acceptors are present near the beginning of the *env*-like gene, one of which is located just before the first methionine and is consistently predicted with a high level of confidence (>94%, Fig. 6D). In the animal retroviruses, the splice-site donor is typically located near the 5' LTR or within Gag. Of the several possible donors in these regions, none are well conserved between element families (data not shown).

Distribution of Endogenous Retroviruses in Plants

To assess the distribution of the endogenous retroviruses, a set of degenerate primers was designed based on conserved sequences flanking the seven domains of the *Athila4*, *Cyclops-2*, and *Calypso* reverse transcriptases. Genomic DNAs were surveyed by PCR from 18 plant species, including several dicots (*Gossypium hirsutum*, cotton; *Platanus occidentalis*, sycamore; *Lycopersicon esculentum*, tomato; *Solanum tuberosum*, potato; and *Nicotiana tabacum*, tobacco), old-world monocots (*Oryza sativa*, rice; *Avena sativa*, oat; *Secale cereale*, rye; *Hordeum vulgare*, barley; *Triticum aestivum*, wheat; and *Sorghum bicolor*, sorghum), new-world monocots (*Zea mays*, corn; *Zea mays* ssp., *Parviglumis*, teosinte; a *Tripsicum* species), and a gymnosperm (*Pinus coulteri,* pine). *A. thaliana*, soybean, and pea, served as positive controls. PCR products were cloned and at least three independent clones were sequenced from each species. Most of the PCR products from dicots and old-world monocots encoded reverse transcriptases that shared >60% amino acid identity. In contrast, the new-world monocots and the single gymnosperm surveyed only yielded reverse transcriptases from more distantly related elements (data not shown). The dicot reverse transcriptases had numerous stop codons and insertions/deletions; whereas, sequences from the old-world monocots were considerably less degenerate. The most intact reverse transcriptases were from oat, rye, and barley, which shared >85% nucleotide identity across species. All nucleotide and amino acid sequences were aligned, making it possible to identify and correct frameshifts. A neighbor-joining tree was constructed from these reverse transcriptases and representative *Tat* elements were used as an outgroup (Fig. 7). The endogenous retroviruses clustered on a single branch, and with few exceptions (e.g., *Diaspora* from soybean), elements from a single species clustered together.

Athila4 Elements Are Expressed in a Methylation-Deficient Strain

The *A. thaliana Athila* elements are preferentially located within heterochromatin flanking the centromeres (Pelissier et al. 1996; Initiative 2000). These regions contain repeated sequences that are methylated and likely transcriptionally quiescent (Jeddeloh et al. 1998; Consortium 2000). Some *Athila* group elements and retrotransposons are expressed in genetic backgrounds, such as *ddm1*, which have reduced levels of DNA methylation (Hirochika et al. 2000; Steimer et al. 2000; Lindroth et al. 2001). We sought *Athila4* mRNAs by RT-PCR in *ddm1* backgrounds, using five different *Athila4* primers and a poly(T) primer/adaptor. Fifteen separate *Athila* cDNAs were cloned and sequenced: eight were *Athila4* elements, four were *Athila6* elements, and three could not be easily assigned to a family because of sequence degeneracy (Fig. 8). No transcripts were recovered from a wild-type strain. All 15 transcripts terminated within a 200-bp window of a consensus *Athila* LTR. One *Athila4* cDNA was primed with a *gag* primer and was 8.4 kb in length. A portion of this clone (1.8 kb) was sequenced and matched *Athila4-6*, except for a single base change, which could be the result of a PCR-induced error. No spliced transcripts were detected.

DISCUSSION

We previously reported that the *A. thaliana Athila* retroelements have a novel feature—a putative *env* gene that may enable them to be infectious (Wright and Voytas 1998). Homologs of *Athila* elements have been described in other plant species (e.g., *Cyclops-2* of pea, Chavanne et al. 1998; *BAGY*-2 of barley, Shirasu et al. 2000), all of which are replete with deletions, rearrangements, or stop codons. To ascertain conserved features of these endogenous plant retroviruses, we analyzed *Athila* elements in the completed *A. thaliana* genome sequence. We also recovered *Athila* homologs from soybean—the so-called *Calypso* elements. By generating consensus sequences from degenerate insertions, we were able to identify features that likely define a functional element.

Shared Features Among Plant Endogenous Retroviruses

The characterized plant endogenous retroviruses range from 12 to 14 kb in length and have LTRs ranging from 1.3 to 1.8 kb, among the largest LTRs described to date for Ty3/*gypsy* elements. Like many plant retroelements, Gag and Pol are encoded on a single ORF. One striking feature among *Athila4*,

Figure 7 Neighbor-joining tree based on amino acid sequences of *Tat* and *Athila* group reverse transcriptases. The tree is rooted to six elements from the *Tat* group (from top to bottom): *Tat4-1*, F26H6, *Cinful*, *Vulgar*, *Rire2*, and *Grande1-4*). The numbers on the branches indicate bootstrap support for 100 replicates. Brackets indicate the species from which the elements originated. Branches indicating previously described elements are labeled. Unlabeled branches represent reverse transcriptases from the following species (from top to bottom): sycamore (*syc4-3*, *syc2-3*, *syc4-2*, *syc4-7*); cotton (*cot8-7*, *cot5- 3*, *cot8-6*); lily (*TfcII* elements AF219190, AF219217, AF219200); rice (*rice2-17*, *rice5-2*, *rice2-10*, *rice1*, *rice*); sorghum (*sorg4-3*, *sorg5-6*, *sorg5-4*, *sorg5-5*, *sorg5-2*, *sorg5-8*); tobacco, tomato, and potato (*tom4-10*, *tom10-16*, *tob1*, *tob2-2*, *tob5-3*, *tob4-1*, *tom4-4*, *pot8-8*, *pot8-3*, *pot8-4*, *pot8-5*, *tom10-4*, *pot8-10*, *pot5-1*); oat (*oat6-7*, *oat6- 1*, *oat6-8*); rye (*rye5-4*, *rye5-2*, *rye3-4*, *rye4-4*); barley (*bar7*, *bar2-12*, *bar2-19*, *bar2-4*, *BAGY-2*); wheat (*wheat5-3*, *wheat8-11*, *wheat8-2*, *wheat3-1*, *wheat8-5*); *Arabidopsis thaliana* (*La9-5*, T17A11, *hau8-3*, *Athila5-2* F23M2, *La9-7*, *La9-9*, *Athila1-1 La8-1*, *La9-6*, T16I21, T13D4, *Athila9-1* T24G23, F18P14, *hau7-2*, T13E11, F7B19, *hau8-4*); pea and favabean (*pea1*, *favabean1*, *favabean2*, *pea8-1*, *pea9-1*); soybean (*will8-2*, *hark5*, *will*, L859-3, *hark2*, *will2*, *hark5-1*, L859-6, L859- 2, L852will8-3, L858-2).

Calypso, and *Cyclops-2* is the high degree of sequence conservation of *pol*. Between these elements, reverse transcriptase and integrase, respectively, share ∼68% and 64% amino acid identity. Because reverse transcription is error prone and often leads to accelerated rates of sequence evolution (Gabriel and Mules 1999), this suggests that either Pol is under very tight functional constraints or that the elements have invaded their plant hosts relatively recently. The phylogenetic tree of *A. thaliana* Ty3/*gypsy* elements provides some support for the recent acquisition of *Athila* elements. The short branch lengths supporting the *Athila* and *Tat* element groups suggest they share a more recent common ancestor relative to classic Ty3/*gypsy* element families (see arrows in Fig. 1). Because the *Athila* elements encode an *env*-like ORF, horizontal transfer by infection is one possibility for the apparent difference in their evolutionary history.

In contrast to *pol*, *gag* shows higher levels of sequence divergence. This is typical of retroelement *gag* genes, whose products carry out structural roles. Nonetheless, *Calypso* and *Cyclops-2* Gag have conserved finger motifs characteristic of nucleocapsid proteins, and all three elements have a conserved domain near the Gag N terminus. Gag averages 675 amino acid residues (measured from the first methionine to the active site of protease), which is larger than most classic plant Ty3/*gypsy* element Gag proteins (e.g., *Reina*, 482 aa; Avramova et al. 1996). If the endogenous retroviruses are in-

Figure 8 Transcription termination sites of *Arabidopsis thaliana Athila* group elements. RNA was isolated from a methylation-deficient strain (*ddm1*) and amplified by RT-PCR using an *Athila4* primer and a poly(T) primer/adaptor. PCR products were cloned and sequenced. At the top of the diagram is a generic *Athila* LTR with the region denoted wherein the transcripts terminate. Clones are listed that match the *Athila6* and *Athila4* elements. The numbers next to the arrows indicate the base position for transcription termination sites within the LTR. Clones were recovered with variable termination sites, as indicated by the gaps in the alignment. Three clones (pDW820, pDW825, and pDW780) match highly degenerate, unclassified *Athila* elements, and the numbers for the BACs that carry these elements are provided.

fectious, Gag may carry out functions related to transmission. Many plant viruses encode movement proteins that transport viral nucleic acids from cell to cell (Ghoshroy et al. 1997) or factors that facilitate spread by insect vectors (Woolston et al. 1983). These proteins are typically not well-conserved, and no similarity to the Gag proteins of the endogenous retroviruses is evident.

Another characteristic feature of the endogenous retroviruses is the presence of two large noncoding regions that flank the *env*-like ORF. The upstream region approximates 0.7 kb, and the downstream region approximates 2 kb. In most retroelements, noncoding sequences are very small, and it is generally thought that extraneous sequences are lost to maximize the amount of genetic information that can be encoded within an element. The conservation of noncoding domains among the endogenous retroviruses suggests they play a role in replication. Possibilities include regulating gene expression (either transcription or translation) or facilitating expression of the *env*-like ORF (e.g., in splicing or in enabling internal ribosome entry). Of the two noncoding regions, the 3' region is flanked by conserved polypurine tracts (PPTs), which might serve as priming sites for plus-strand DNA synthesis during reverse transcription. Multiple PPTs are found in other retroelements such as Ty1 and HIV, although in these elements, the upstream PPT resides within *pol* (Hungnes et al. 1993; Heyman et al. 1995). A third, small noncoding region is also found between the 5' LTR and the start of the gag-pol ORF. This region carries the putative primer binding site (PBS) for minus-strand DNA synthesis, which is complementary to an Asp tRNA. This is a distinguishing feature of the endogenous retroviruses, for the classic Ty3/*gypsy* elements and the Ty1/ *copia* group elements have PBSs complementary to initiator Met tRNAs, and the Tat elements have PBSs complementary to Asn, Lys, and Arg tRNAs (Wright and Voytas 1998; D.A. Wright, unpublished observation).

The *env*-Like ORF and Its Potential Role in Infection

We previously concluded that the *env*-like genes of the endogenous retroviruses likely play a functional role in replication, based on sequence conservation between the *Cyclops-2* and *Calypso env*-like genes (Peterson-Burch et al. 2000). With the availability of the *A. thaliana* genome sequence, additional *Athila env*-like genes made it possible to discern conserved features. Computer models predict the Env-like proteins are expressed from a spliced subgenomic mRNA. The Env-like proteins are also predicted to encode a central transmembrane domain. Env-like proteins of animal retroviruses often have both central and C-terminal transmembrane domains, the latter of which anchors Env within the endoplasmic reticulum. In most endogenous plant retroviruses, there is a short ORF after the *env*-like gene that is predicted to encode a transmembrane domain and could serve an anchoring role. Expression of the short ORF as part of Env would require read-through of a stop codon. Alternatively, because a transmembrane domain is also encoded in adjacent reading frames, ribosomal frameshifting may be employed. Attempts to determine if this region is evolving as a coding sequence were not productive because of the high degree of sequence divergence between element families. As other endogenous plant retroviruses are identified, it will be of interest to determine whether they too have this short transmembrane domain-encoding ORF. A functional element will be required to determine experimentally whether it has a biological role.

If the endogenous retroviruses are infectious, then the Env-like protein is likely important in this process. During infection by retroviruses, Env facilitates the merging of the membrane-bound virion with the target cell. The plant cell wall poses an obstacle to membrane-mediated infection. Nonetheless, enveloped plant viruses do exist, including members of *Bunyaviridae* and the *Rhabdoviridae* (van Regenmortel et al. 2000). These viruses bud from the endomembrane system and accumulate in the cell until a feeding invertebrate ingests them and carries them to another plant. Recent work has shown that some animal retrotransposons have acquired *env* genes from viruses (Malik 2000). For example, the *env* gene of the *D. melanogaster gypsy* element is related to *env* of the baculoviruses and was likely acquired by *gypsy* through transduction. To date, however, we have not identified similarity between the *env*-like ORFs of the endogenous plant retroviruses and those of viruses or other genes in the databases. It should be mentioned that some plant Ty1/ *copia* group retrotransposons have *env*-like ORFs (Laten et al. 1998, 1999; Kapitonov and Jurka 1999; Peterson-Burch et al. 2000). These genes are unrelated to the *env*-like genes of *Athila* and its homologs, but they are predicted to be transmembrane proteins. It is tempting to speculate that *env*-like genes play a similar role in both groups of elements.

Distribution of Endogenous Retroviruses in Plants

Using a PCR-based assay, we found that endogenous retroviruses are widely distributed among angiosperms. The recovered reverse transcriptases were strikingly similar and shared >60% amino acid identity. This high degree of sequence conservation belied the fact that most carried mutations, the exception being elements from cereals, namely oat, rye, and barley. The integrity of the cereal reverse transcriptases implies that these elements have undergone more recent episodes of replication, and to date, they are the best candidates for functional endogenous retroviruses. Elements were not recovered from a gymnosperm (pine) and the three newworld monocot species tested (corn, teosinte, and tripsicum). It may be that the endogenous retroviruses are not present in the genomes of these plants or that they are divergent and cannot be amplified by the primers. Phylogenetic analyses of the reverse transcriptases indicated that, with few exceptions, the relationships among the elements reflected relationships among their hosts. This suggests that either the endogenous retroviruses are inherited vertically or if they are viruses, they have a limited host range. As more plant genomes are characterized in greater detail, it will be of interest to determine whether high levels of sequence conservation is a general feature of the endogenous plant retroviruses. This will help address the question as to whether or not they are young retroelements relative to the classic Ty3/*gypsy* elements.

Expression and Activity of *A. thaliana Athila* Group Elements

Most *A. thaliana Athila* elements are located within centromeric heterochromatin, which is typically highly methylated (Vongs et al. 1993; Pelissier et al. 1996; Copenhaver et al. 1999). Methylation is thought to control transposable element activity (Yoder et al. 1997; Martienssen 1998), and several recent studies in plants have shown that decreases in DNA methylation are associated with increased transposable element activity (Hirochika et al. 2000; Lindroth et al. 2001; Miura et al. 2001; Singer et al. 2001). Of particular relevance

to this study, truncated *Athila* transcripts have been reported in strains with *mom1* mutations, which derepress transcriptionally silent loci (Amedeo et al. 2000; Steimer et al. 2000).

We performed RT-PCR on RNAs isolated from *ddm1* plants and were able to amplify cDNA from *Athila4* and *Athila6* elements, two of the most intact *Athila* families. Transcripts terminated at a similar position within the LTR, thereby defining the LTR R/U5 boundary. cDNA as large as 8.4 kb was recovered; however, no spliced messages were identified. Although *Athila* elements are expressed in *ddm1* backgrounds, they are probably not replicating because of sequence degeneracy. For future studies, it will be important to identify a functional *Athila* group element. We envision two approaches for how this might be accomplished: 1) a consensus *Athila4* element could be constructed or 2) elements could be further characterized from species such as the small grains that appear to have structurally intact elements. The identification of a replication-competent *Athila* group element will be necessary to test the hypothesis that these elements are infectious plant retroviruses. If this proves to be the case, the *Athila* group elements may be useful as vectors for gene transfer and the genetic modification of plants.

METHODS

DNA Manipulations and Filter Hybridizations

A soybean genomic λ phage library (Chen et al. 1998) was screened with a reverse transcriptase probe under low stringency conditions (50°C with a 1% SDS wash; Ausubel et al. 1987). The probe was obtained by PCR amplification of *Pisum sativum* DNA using primers based on the *Cyclops-2* reverse transcriptase (DVO701 5'-CCG-TCA-TCC-GGA-ATG-ACA-AGG-ATG and DVO702 5-ACG-GAT-GAG-CCT-TTG-CTT-CGA-ATC). Phage subclones were sequenced by primer walking. Genomic DNAs from 18 plant species (see Results) were surveyed by PCR to identify *Athila*-group reverse transcriptases. DNAs were prepared using genomic tips and protocols supplied by Qiagen. Degenerate primers were designed based on two conserved amino acid sequence motifs flanking the seven core domains of reverse transcriptase (Xiong and Eickbush 1990; VRKEVLKL, DVO1197 5'-GTG-CGN-AAR-GAR-GTN-NTN-AAR-YT, and FIKDFSKV, DVO1198 5-AAC-YTT-NGW-RAA-RTC-YTT-DAT-RAA). PCR was performed in 50 µL reactions with ∼100 ng genomic DNA, 3 µmole of each primer, 2.5 units Taq DNA polymerase, 1X Taq buffer (Promega), and 2.5 mM $MgCl₂$. PCR was performed for 30 cycles under the following conditions: 92°C for 20 sec, 50°C for 30 sec, and 72°C for 90 sec. The PCR products were purified on low-melting agarose gels and cloned into T-vector prepared from pBluescript II KS- (Hadjeb and Berkowitz 1996). *Athila*group reverse transcriptases were sequenced in their entirety from vector-based primers.

Sequence Analysis

DNA Sequence analysis was performed using the GCG software package (Devereux et al. 1984), DNA Strider 1.2 (Marck 1991), and the BLAST search tool (Altschul et al. 1990). Phylogenetic relationships were determined by the neighborjoining distance algorithm using PAUP v4.0 beta 4a (Saitou and Nei 1987; Swofford 1991) and were based on reverse transcriptase amino acid sequences that had been aligned with CLUSTALX v1.63b (Thompson et al. 1994). Transmembrane helices were identified using the PHDhtm program and TMpred (Hofmann and Stoffel 1993; Rost et al. 1995). Splicesite analysis was performed with NetGene2 (Brunak et al. 1991; Hebsgaard et al. 1996). All DNA sequences have been submitted to the DDBS/EMBL/GenBank databases. The *Ca-* *lypso* elements are under accession numbers AF186182, AF186183, AF186184, AF186185, and AF186186. BAC or P1 clone numbers for the Ty3/*gypsy* reverse transcriptases are listed in the Figure 1 legend. Accession numbers for the *Athila4* elements are listed in the Figure 2 legend. The accession numbers of the *Athila*-group reverse transcriptases from various species are AF378012 to AF378081. Additional details regarding these sequences can also be found at our Web site (http://www.public.iastate.edu/∼voytas/).

RT-PCR

Total RNA was isolated from *A. thaliana ddm1* plants using the PUREscript RNA isolation kit (Gentra Systems, Inc.). RNA was annealed to the primer DVO1247, which is a poly(T) oligo with a specific tail (5-GGA-CTT-CAG-GAC-TGC-TTG-ACA-AAG-T₃₀). First-strand DNA synthesis was performed at 42° C for 2 h using Superscript II reverse transcriptase and the manufacturer's protocol (GIBCO BRL). RNase activity was inhibited by the addition of Super RNase IN per the manufacturer's instructions (Ambion). PCR was carried out using the Expand Long Template PCR System (Roche Molecular Biochemicals) with *Athila*-element-specific primers, along with DVO1248, which is specific to the tail of DVO1247.

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