

# Stowaway: A New Family of Inverted Repeat Elements Associated with the Genes of Both Monocotyledonous and Dicotyledonous Plants

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Members of a new inverted repeat element family, named *Stowaway*, have been found in close association with more than 40 monocotyledonous and dicotyledonous plant genes listed in the GenBank and EMBL nucleic acid data bases. *Stowaway* elements are characterized by a conserved terminal inverted repeat, small size, target site specificity (TA), and potential to form stable DNA secondary structures. Some elements are located at the extreme 3' ends of sequenced cDNAs and supply polyadenylation signals to their host genes. Other elements are in the 5' upstream regions of several genes and appear to contain previously identified *cis*-acting regulatory domains. Although the *Stowaway* elements share many structural features with the recently discovered *Tourist* elements, the two families share no significant sequence similarity. Together, the *Stowaway* and *Tourist* families serve to define an important new class of short inverted repeat elements found in possibly all flowering plant genomes.

## INTRODUCTION

The majority of interspersed repetitive DNA in eukaryotes has been suggested to be transposable elements or their remnants (Flavell, 1986). Moreover, some highly repetitive families of transposable elements are frequently associated with genes. Several human gene sequences, for instance, harbor the retroposon *Alu* ( $\sim 10^6$  copies per haploid genome) and the long interspersed nuclear sequence (LINE) L1 ( $\sim 10^5$  copies per haploid genome) (Berg and Howe, 1989). The proximity of transposable elements may influence the expression of the neighboring cellular genes by activating cryptic or supplying *cis*-acting regulatory regions (Clemens, 1987; Paulson et al., 1987; Baumruker et al., 1988; Stavenhagen and Robins, 1988; Banville and Boie, 1989; Chang-Yeh et al., 1991; Goodchild et al., 1992; Maichele et al., 1993).

Flowering plants have genomes that are on average much larger than those of other higher eukaryotes and are thought to have a correspondingly larger number of transposable elements (Bennett and Smith, 1991). Some known plant retrotransposons occur at high copy number in their host genomes (Grandbastien, 1992). The *del2* (dispersed element of lilies) element, for example, constitutes 4% of the lily genome (Leeton and Smyth, 1993). Previously, we have described a recent insertion of a mobile element, *Tourist-Zm1* (*Zea mays*), in a maize *waxy* allele (Bureau and Wessler, 1992). This element was found to be a member of another highly repetitive transposable element family associated with more than 30 wild-type

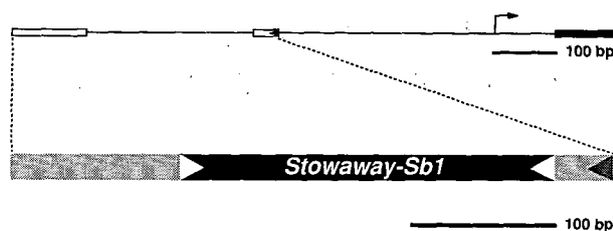
genes of cereal grasses listed in nucleotide data bases (Bureau and Wessler, 1992, 1994). *Tourist* is characterized by terminal inverted repeats (TIRs), small size, target site preference (TAA), and potential to form stable DNA secondary structure. In this report, we describe a new family of transposable elements, named *Stowaway*, which are similar in structure but not in sequence to *Tourist* and are associated not only with listed gene sequences of cereal grasses but also with dicotyledonous plant genes. Furthermore, the fact that some *Stowaway* elements contain previously identified *cis*-acting regulatory regions provides evidence that this new family has contributed to the evolution of host gene expression.

## RESULTS

### Identification of *Stowaway* in Higher Plant Gene Sequences

The *Tourist-Sb5* (*Sorghum bicolor*) element, located at the extreme 5' end of the sorghum phosphoenolpyruvate carboxylase CP21 gene sequence (Lepiniec et al., 1993; Bureau and Wessler, 1994), is interrupted by a 257-bp insertion (Figure 1). The presence of an imperfect TIR and a flanking 2-bp direct repeat (TA) suggests that this insertion, similar to *Tourist*, may be a transposable element. We have named this new element *Stowaway-Sb1*.

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**Figure 1.** Position of *Stowaway-Sb1* within *Tourist-Sb1*.

The disrupted *Tourist-Sb1* (top diagram, open boxes) in the 5' flanking region of the sorghum phosphoenolpyruvate carboxylase CP21 gene (transcription start site, bent arrow; 5' coding sequence, black rectangle) has been expanded to show the position of *Stowaway-Sb1*. Triangles indicate the position of TIRs.

To determine if this element was a member of a larger family, computer-assisted sequence similarity searches of the GenBank (version 77.0) and EMBL (version 34.0) nucleic acid data bases were performed using *Stowaway-Sb1* as a query sequence. As new *Stowaway* elements were identified, these sequences were also used as queries. New *Stowaway* elements were defined as sequences that shared not only significant nucleotide similarity (>60% overall sequence similarity between any two elements), but also other structural features characteristic of the family, including TIR sequence similarity, target site duplication size, secondary structure, and overall length. Members of the *Stowaway* family that were identified in this way are listed in Table 1. Surprisingly, 47 plant sequences were identified as harboring *Stowaway* elements. Although several more degenerate sequences (<60% overall sequence similarity with another element and/or only partial structural similarity with *Stowaway*) were identified, only the best matches are presented. Whereas *Tourist* elements were found only within selected cereal grasses, the *Stowaway* family has a much wider distribution, with members in both monocotyledonous and dicotyledonous plants.

Although sequence similarity between elements ranges from 45 to 85% (Figures 2 and 3 and data not shown), *Stowaway* family members share several other features. First, *Stowaway* elements have a conserved 11-bp TIR with an overall consensus sequence of C<sub>90</sub>T<sub>88</sub>C<sub>93</sub>C<sub>79</sub>C<sub>82</sub>T<sub>92</sub>C<sub>88</sub>C<sub>69</sub>G<sub>77</sub>T<sub>89</sub>T<sub>65</sub> (numbers in subscript refer to the percent occurrence). Second, *Stowaway* elements are small, ranging from 80 to 323 bp. Whereas the monocotyledonous elements are variable in size, the dicotyledonous elements are considerably more homogeneous (248 ± 24 bp). In general, the reported *Stowaway* elements found in monocotyledonous plant genes are more similar to one another than to elements found in dicotyledonous plant genes and vice versa (Figures 2 and 3 and data not shown). Third, *Stowaway* elements are AT rich (72 ± 5%). Fourth, *Stowaway* has a strong target site preference; ~85% of the *Stowaway* elements listed in Table 1 have TA targets (Figures 2 and 3 and data not shown). Among plant transposable elements characterized to date, only *Tourist* and *Stowaway* have target sequence preference (Bureau and Wessler, 1992,

1994). A TA target site sequence is also characteristic for members of the *IS630-Tc1* (transposon of *Caenorhabditis*) family of transposable elements (Doak et al., 1994). There is, however, no significant sequence similarity between *IS630-Tc1* family members and *Stowaway*. Fifth, *Stowaway* elements have a potential to form DNA secondary structures (Table 1; Figure 4). *Stowaway-Zm3*, for example, can be folded into a perfect hairpin except for a 1-bp mismatch. The FB elements of *Drosophila* (Smith and Corces, 1991) and Tc6 of *Caenorhabditis* (Dreyfus and Emmons, 1991) also have the potential to form hairpin-like structures but lack significant sequence similarity with *Stowaway*.

### Evidence for Element Insertion

Although *Stowaway* family members have structural features of transposable elements, it was important to obtain evidence for element mobility in both monocotyledonous and dicotyledonous plants. To this end, two approaches were utilized. *Stowaway-St5* (*Solanum tuberosum*), *Stowaway-Ps1* (*Pisum sativum*), and *Stowaway-Le2* (*Lycopersicon esculentum*) have inserted into dicotyledonous plant genes that belong to gene families (paralogous loci) or are members of a group of homologous genes that have been previously isolated from closely related species (orthologous loci). Alignment of the sequences flanking these elements with paralogous and orthologous loci indicated that these elements correspond exactly with insertion polymorphisms (Figure 5). To provide evidence for element mobility in monocotyledonous plants, polymerase chain reaction (PCR) was employed to amplify introns harboring *Stowaway-Os3* (*Oryza sativa*) and *Stowaway-Zm3* from orthologous loci of closely related species or of cultivars within the same species. In each case, insertion polymorphisms were identified that corresponded precisely to the location of *Stowaway* and a TA target site duplication, thus revealing the site of a relatively recent insertion event (Figure 5).

### *Stowaway* Elements Contain Previously Identified *cis*-Acting Regulatory Domains

A subset of the *Stowaway* elements that are located in the 5' flanking regions of plant genes harbors previously identified *cis*-acting regulatory domains. Sequences in the *Stowaway-Le2* element (located within the promoter of a tomato gene encoding the small subunit of ribulose-1,5-bisphosphate carboxylase), for instance, contain ~50% of the sequences protected in a DNase footprinting assay (Manzara et al., 1993). In addition, *Stowaway-Le1* shares ~80% sequence similarity with *Stowaway-Le2* and occupies ~65% of a negative regulatory domain identified within the LAT59 promoter (−804 to −418, relative to the start of transcription) (Twell et al., 1991). Similarly, two putative embryogenesis-specific nuclear factors bind within the internal sequences of the *Stowaway-Dc2* (*Daucus carota*) element of carrot (Hatzopoulos et al., 1990), and

**Table 1. Stowaway Elements Associated with Plant Gene Sequences Listed in the GenBank and EMBL Data Bases**

Element <sup>a</sup>	Locus Name	Gene Description <sup>b</sup>	Position <sup>c</sup>	Size (bp)	$\Delta^{\circ}G^d$	References <sup>e</sup>
<b>Monocotyledonous Plants</b>						
<i>Os1</i>	OSHSP82A	Heat shock protein 82A	in2 (2150)	150	-58.0	NA
<i>Os2</i>	RICAMYC	$\alpha$ -Amylase C	3' (1667)	245	-70.1	Kim and Wu (1992)
<i>Os3</i>	OSWAXY	Starch synthase	in13 (3049)	234	-77.1	Wang et al. (1990)
<i>Os4</i>	OSWAXY	Starch synthase	3' (4046)	204	-44.5	Wang et al. (1990)
<i>Os5</i>	OSPCNAGEN	PCNA	5' (-432)	122	-47.1	Suzuka et al. (1991)
<i>Os6</i>	OSPCNAGEN	PCNA	5' (-1315)	227	-91.4	Suzuka et al. (1991)
<i>Os7</i>	OSRAMY3A	$\alpha$ -Amylase 3A	in2 (405)	234	-60.7	Sutliff et al. (1991)
<i>Zm1</i>	ZM27KZNB	27-kD zein	3'UTR/3' (830)	163	-24.7	Das et al. (1991)
<i>Zm2</i>	ZMAZ22KD	22-kD zein	3' (4195)	156	-30.0	Thompson et al. (1992)
<i>Zm3</i>	ZMAYSPG	P	in2 (4760)	80	-54.5	Athma et al. (1992)
<i>Zm4</i>	M23537	10-kD zein	3' (591)	153	-27.8	Kirihara et al. (1988)
<i>Zm5</i>	ZMGAPC4	GAP dehydrogenase	in3 (1444)	157	-38.4	Kersanach et al. (1994)
<i>Sh1</i>	SCFSCPEPCD	PEP carboxylase	in6 (2675)	267	-75.1	Albert et al. (1992)
<i>Sb1</i>	SVPEPCGX	PEP carboxylase	5' (-469)	255	-50.7	Lepiniec et al. (1993)
<i>Sb2</i>	SVPEPCGX	PEP carboxylase	in6 (3231)	131	-42.3	Lepiniec et al. (1993)
<i>Ta1</i>	S117442	Metallothionein	5' (-505)	167	-49.9	Kawashima et al. (1992)
<i>Ta2</i>	TAAAM254	$\alpha$ -Amylase 2/54	5' (-458)	100	-40.5	Huttly et al. (1992)
<i>Hv1</i>	HVBKIN12G	Protein kinase	in2 (1747)	81	-30.7	Halford et al. (1992)
<i>Hv2</i>	BLYRCABG	Rubisco activase <i>RcaA</i>	in3 (839)	159	-40.3	Rundle and Zielinski (1991)
<i>Hv3</i>	BLYGLB2	1,3-1,4- $\beta$ -Glucanase	in1 (166)	323	-87.8	Wolf (1991)
<i>Hv4</i>	BLYCLDAA	Cold-regulated protein	3'UTR (604)	49 <sup>f</sup>	ND	Cattivelli and Bartels (1990)
<i>Hv5a</i>	HVPRP1A	Pathogenesis related (Hv-1a)	3'UTR (636)	123 <sup>f</sup>	ND	Bryngelsson and Gréen (1989)
<i>Hv5b</i>	HVPRP1B	Pathogenesis related (Hv-1b)	3'UTR (647)	36 <sup>f</sup>	ND	Bryngelsson and Gréen (1989)
<i>Hv5c</i>	HVPRP1C	Pathogenesis related (Hv-1c)	3'UTR (604)	93 <sup>f</sup>	ND	Bryngelsson and Gréen (1989)
<b>Dicotyledonous Plants</b>						
<i>Ps1</i>	PEALCTN	Lectin	5' (-1296)	275	-64.9	Mandaci and Dobres (1993)
<i>Ps2</i>	PEACAB80	CAB binding protein	5' (-615)	276	-56.5	Timko et al. (1985)
<i>Pc1</i>	PCPR2G	Pathogenesis related (PR2)	5' (-415)	243	-90.0	Van de Löcht et al. (1990)
<i>Dc1</i>	DCDC8	DC8	5' (-514)	253	-65.9	Franz et al. (1989)
<i>Dc2</i>	S47635	DC59	5' (-409)	249	-78.1	Hatzopoulos et al. (1990)
<i>Bn1</i>	BNESPSPG	EPSP synthase	in5 (1835)	220	-43.6	Gasser and Klee (1990)
<i>Bn2</i>	BNAC2PROMO	Cruciferin	5' (-1147)	247	-41.7	Breen and Crouch (1992)
<i>Sa1</i>	SASCHSG	Chalcone synthase	5' (-334)	260	-57.7	Batschauer et al. (1991)
<i>St1</i>	STWIN12G	Wound induced	5' (-139)	259	-53.1	Stanford et al. (1989)
<i>St2</i>	STRBCS1	Rubisco small subunit	in1 (337)	285	-49.2	NA
<i>St3</i>	STPROINI	Proteinase inhibitor	3' (1393)	240	-37.0	Lee and Park (1989)
<i>St4</i>	POTPATA	Patatin	in2 (922)	259	-67.3	Mignery et al. (1988)
<i>St5</i>	STPATP1	Patatin pseudogene	in5 (4378)	226	-34.8	Pikaard et al. (1986)
<i>St6</i>	STPOAC58	Actin	in3 (1426)	167	-41.6	Drouin and Dover (1990)
<i>Le1</i>	LELAT59	Pollen maturation specific	5' (-656)	260	-51.6	Twell et al. (1991)
<i>Le2</i>	LERBSS1	Rubisco small subunit	5' (-275)	251	-40.2	Manzara et al. (1993)
<i>Le3</i>	TOMATPACA	Ca <sup>2+</sup> -ATPase	5' (-114)	290	-65.5	Wimmers et al. (1992)
<i>Le4</i>	TOMPHEAMLY	PAL	5' (-494)	248	-43.1	NA
<i>Nt1</i>	NTT85A	Auxin binding protein	in2 (622)	225	-50.7	NA
<i>Nt2</i>	NTCHN50	Endochitinase	5' (-209)	244	-46.1	Fukuda et al. (1991)
<i>Np1</i>	TOBPMA1A	H <sup>+</sup> -ATPase	in2 (443)	247	-40.2	Perez et al. (1992)
<i>Ns1</i>	NTNIA2	Nitrate reductase	in1 (1041)	249	-46.5	Vaucheret et al. (1989)
<i>Nr1</i>	NRTY8	tRNA-Tyr	5' (-62)	75 <sup>f</sup>	ND	Fuchs et al. (1992)
<i>Ph1</i>	PETEPSP	EPSP synthase	5' (-1025)	243	-39.0	Benfey et al. (1990)
<i>Ph2</i>	PHCHSD	Chalcone synthase D	5' (-115)	221	-39.3	Koes et al. (1989)
<i>Ph3</i>	PHCHSG	Chalcone synthase G	in1 (884)	257	-62.7	Koes et al. (1989)

<sup>a</sup> *Os*, *Oryza sativa* (rice); *Zm*, *Zea mays* (maize); *Sh*, *Saccharum hybrida* (sugarcane); *Sb*, *Sorghum bicolor* (sorghum); *Ta*, *Triticum aestivum* (wheat); *Hv*, *Hordeum vulgare* (barley); *Ps*, *Pisum sativum* (pea); *Pc*, *Petroselinum crispum* (parsley); *Dc*, *Daucus carota* (carrot); *Bn*, *Brassica napus* (rape seed); *Sa*, *Sinapis alba* (mustard); *St*, *Solanum tuberosum* (potato); *Le*, *Lycopersicon esculentum* (tomato); *Nt*, *Nicotiana tabacum* (tobacco); *Np*, *N. plumbaginifolia*; *Ns*, *N. sylvestris*; *Nr*, *N. rustica*; *Ph*, *Petunia hybrida*. Stowaway-*Os3* has been previously referred to as *Tnr1* (transposable element in rice) (Umeda et al., 1991).

<sup>b</sup> PCNA, proliferating cell nuclear antigen; GAP, glyceraldehyde-3-phosphate; PEP, phosphoenolpyruvate; CAB, chlorophyll *a/b* binding protein; ESPS, 5-enolpyruvylshikimate-3-phosphate; PAL, phenylalanine ammonium-lyase.

<sup>c</sup> 5', 5' flanking region; 3', 3' flanking region; in, intron sequence; UTR, untranslated region. Positions (given in parentheses) are relative to the translation start site. The position of Stowaway-*Nr1* is relative to the start of transcription.

<sup>d</sup> Kilocalorie per mole; ND, not determined.

<sup>e</sup> NA, not available.

<sup>f</sup> Only partial sequence available.

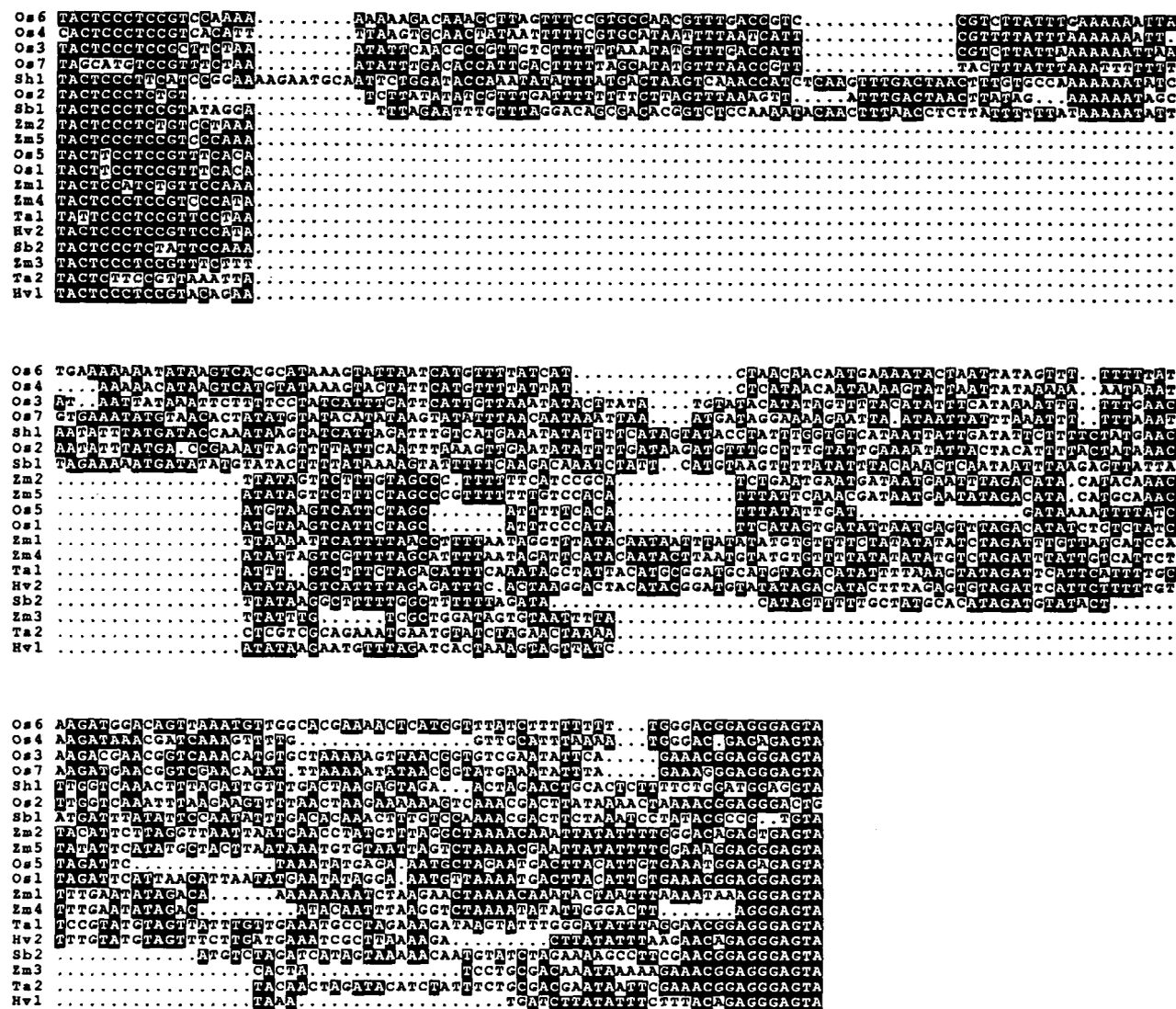


Figure 2. Multiple Sequence Alignments of *Stowaway* Elements Associated with Monocotyledonous Plant Genes.

Two base pairs immediately flanking the termini of each *Stowaway* element were included in the multiple alignment. The alignment is one of many possible optimal multiple sequence alignments and does not necessarily reflect the best pairwise relationship between any two *Stowaway* elements. *Stowaway-Hv3* and truncated elements (see Table 1) were omitted from the alignment. Conserved nucleotides are indicated by white letters on a black background.

*Stowaway-St1* occupies ~45% of an upstream region important for wound inducibility in potato (Stanford et al., 1989, 1990).

In addition to the identification of *Stowaway* among genomic sequences, elements were also found in four barley stress-induced mRNAs (Table 1). Whereas one of these mRNAs was inducible by cold temperature stress (Cattivelli and Bartels, 1990), the remaining three transcripts (Hv-1a, Hv-1b, and Hv-1c) were derived from members of a gene family that encodes pathogenesis-induced thaumatin-like proteins (Bryngelsson and Gr en, 1989). The presence of *Stowaway* at the same position in all three Hv-1 transcripts indicates that insertion predates the amplification of this gene family. Interestingly, each

Hv-1 transcript is polyadenylated at a different site within *Stowaway* sequences (Figure 6). This may indicate either that these related elements (>93% sequence similarity) have multiple sites for polyadenylation or that the minor sequence differences influence poly(A) site selection.

## DISCUSSION

In this report, we describe an element family named *Stowaway*, which was first identified as an insertion in a member of another

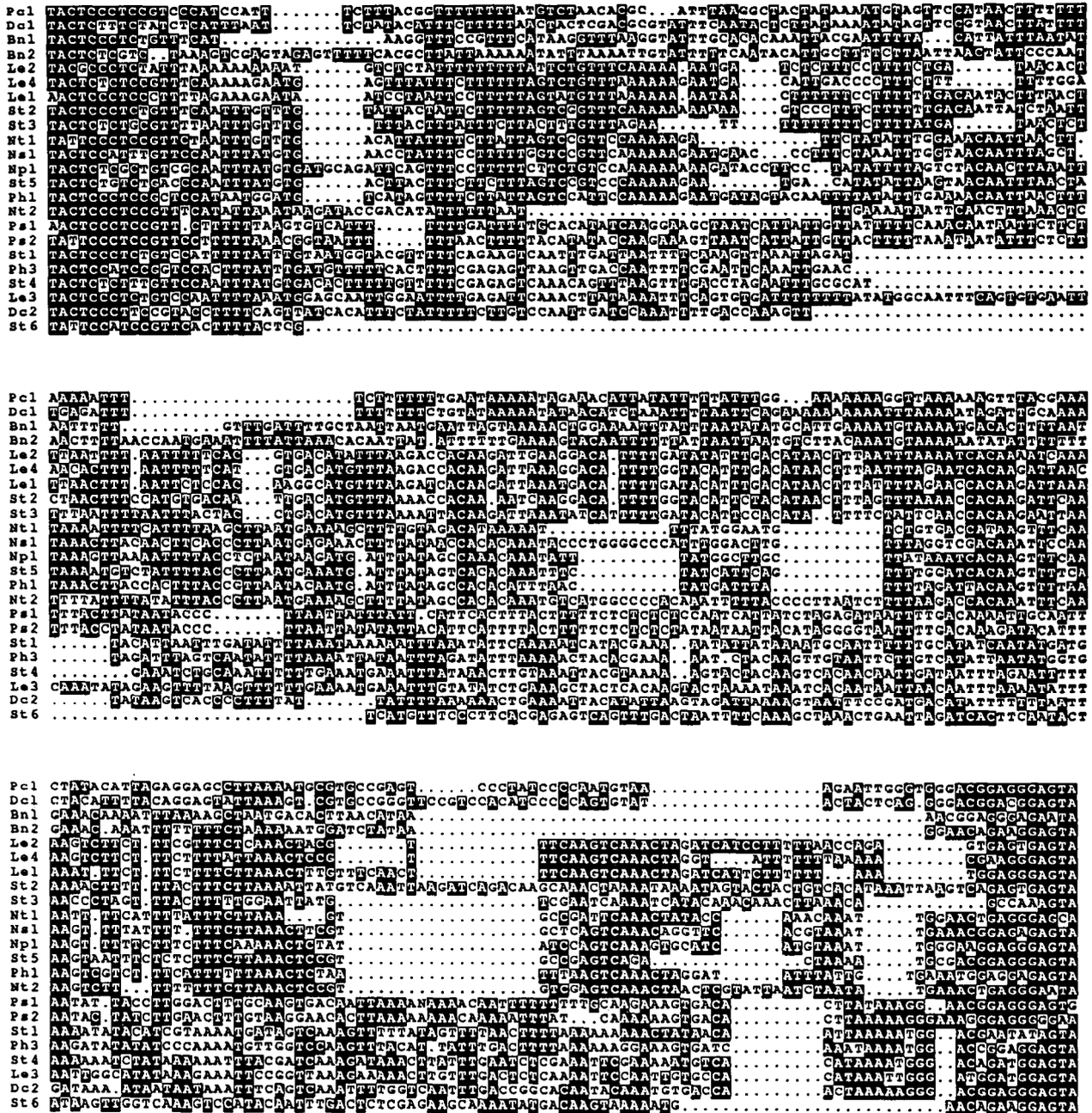
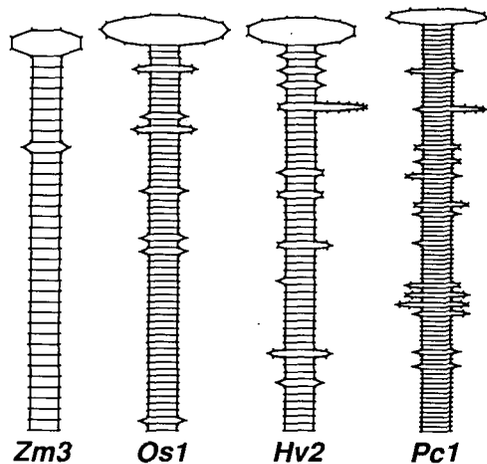


Figure 3. Multiple Sequence Alignment of Stowaway Elements Associated with Dicotyledonous Plant Genes.

Stowaway-Ph2, Stowaway-Sa1, and truncated elements (see Table 1) were omitted from the alignment. See legend to Figure 2. Abbreviations are as given in Table 1.

family of transposable elements, *Tourist*. More than 30 genes from several grasses harbor the transposable element *Tourist*; they are short (113 to 343 bp), have the potential to form DNA secondary structures, are AT rich, and have a preference for insertion at the trinucleotide TAA (Bureau and Wessler, 1992, 1994). *Tourist* elements have a mobile history because a maize mutant *waxy* allele was found to be caused by the recent

insertion of *Tourist-Zm1*. In addition, the locations of *Tourist* elements in other genes correspond with insertion polymorphisms at orthologous loci. Despite the fact that *Stowaway* shares no significant sequence similarity to *Tourist*, these two families of elements have strikingly similar structural features. For instance, *Stowaway* elements are short, are AT rich, have the potential to form DNA secondary structures, and have target



**Figure 4.** Stem-Loop Structures of *Stowaway*-Zm3, -Os1, -Hv2, and -Pc1.

The outline of each folded element is shown (horizontal lines correspond to a base pair). The dinucleotide direct repeats immediately flanking the elements were not included in the predicted DNA secondary structures. *Pc*, *Petroselinum crispum*.

insertion site preference. This suggests that the *Stowaway* and *Tourist* families are members of a larger class of elements that probably transpose by a similar mechanism. The presence of *Tourist* in maize, sorghum, rice, and barley indicates that *Tourist* is probably ubiquitous in the genomes of cereal grasses. Computer-assisted data base searches revealed that *Stowaway* elements are found in the genomes of both monocotyledonous and dicotyledonous plants, indicating that the *Tourist/Stowaway* superfamily of mobile elements is an important component of the genomes of possibly all flowering plants.

Mobility of some *Stowaway* elements is evident by their correspondence to insertion polymorphisms between orthologous and paralogous loci. Such polymorphisms also verify that the *Stowaway* target site is a dinucleotide (preferentially TA). Examples of *Stowaway* mobility given in Figure 5 indicate that element activity has occurred on an evolutionary time scale. For instance, *Stowaway*-Os1 was identified in intron 2 of the heat shock protein 82A gene of domesticated rice, *O. sativa*, and other A genome-type rice species (Table 1 and data not shown). The absence of *Stowaway*-Os1 in non-A-genome-type rice species (Figure 5D) suggests that the *Stowaway*-Os1 insertion corresponds to the approximate divergence date of the A genome (~14 to 17 million years; Dally, 1988). In contrast, it appears that *Stowaway*-Zm1 has transposed into intron 2 of the maize *P* gene much more recently because an insertion polymorphism was identified between orthologous loci of two maize inbred lines (Figure 5E).

The mechanism of mobility of the *Tourist/Stowaway* element superfamily remains unknown. The presence of TIRs is reminiscent of inverted repeat elements that transpose via a DNA intermediate or "cut-and-paste" mechanism (Bureau and

A

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STPATP1  TTTCTTAATATA > St5 < TATAATAGAAAA
STPATP2  TTTCTTAATATA -----TGAAAGGAAA
POTPATA  TTTCTTAATATA -----TGGTAGAAAA
STPATG   TTTCTTAATATA -----TGATAGGAAA
  
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B

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PEACAB80 TATAATTAAC TA > Ps2 < TATATACTAGTT
PEACAB66 TATAATTAAC CA -----TATACTAGTA
  
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C

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LERBSS1  TCTTGTCTATTA > Le2 < TAAAATAT*AAA
STRBCS3  TCTTGTCTATTA -----AAATAT*AAA
  
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D

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Osativa  GTTTGTCTATTA > Os1 < TACTATGAATTA
Opuncat  GTTTGTCTATTA -----CTATGAATTA
Oeichin  ATTTGTTTATTA -----CTATAAATTA
  
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E

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maizW22  TCTATATATATA > Zm3 < TATGACTAGGC
maizMo20 TCTATATATATA -----TGTACTAGGC
teosZmm  TMTATATATATA -----TGTACTAGGC
  
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**Figure 5.** Polymorphisms Corresponding to *Stowaway* Insertions into Dicotyledonous and Monocotyledonous Genes.

(A) *Stowaway*-St5 is located within intron 5 of the potato patatin pseudogene (STPATP1) but not in the corresponding position of three other members of the patatin gene family (STPATP2, POTPATA, and STPATG).

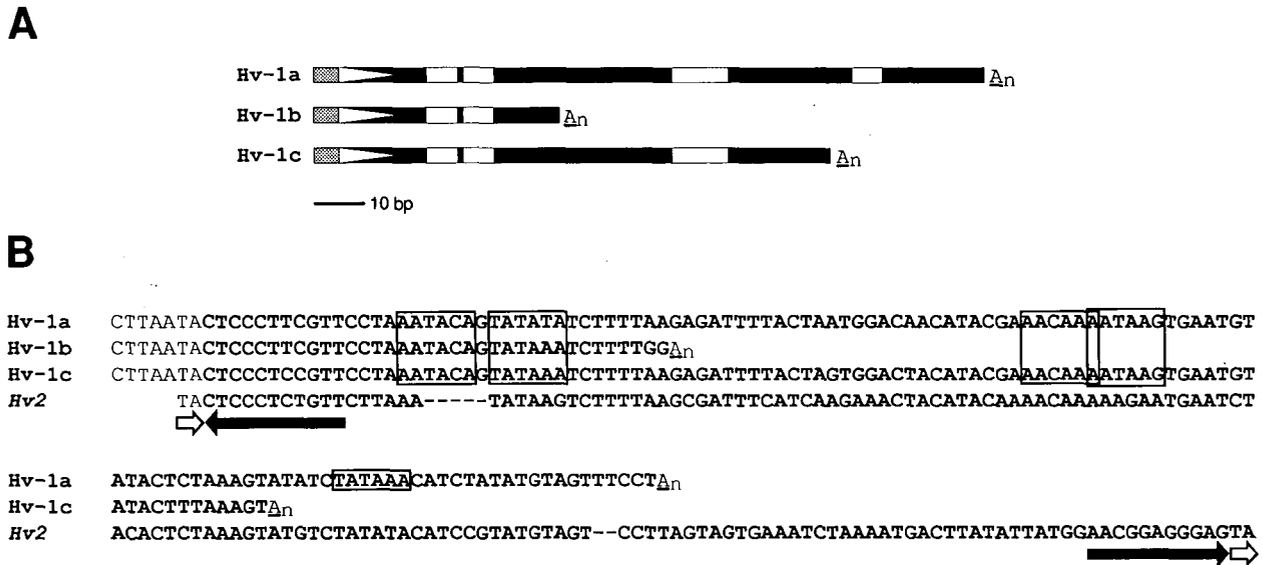
(B) *Stowaway*-Ps2 is located in the 5' flanking region of the pea CAB80 (PEACAB80) gene but not in CAB66 (PEACAB66), another member of the CAB gene family in pea.

(C) *Stowaway*-Le2 is located in the 5' flanking region of the tomato *rbcs1* gene (LERBSS1) but not in the corresponding position of the potato *rbcs3* gene (STRBCS3). An asterisk indicates a short variable region (LERBSS1, 7 bp; STRBCS3, 16 bp).

(D) *Stowaway*-Os1 is located within intron 2 of the heat shock protein 82A gene of *Oryza sativa* (*Osativa*) but not in the wild rice species *O. punctata* (*Opuncat*) and *O. eichingeri* (*Oeichin*).

(E) *Stowaway*-Zm3 is located within intron 2 of the maize *P* gene of the inbred line W22 (maizW22) but not in the Mo20 (maizMo20) inbred line or in the teosinte, *Z. mays* subsp. *mexicana* (teosZmm).

In all cases [(A) through (E)], significant sequence similarity extends past the region delimited. The presumed TA target sites are boxed, and dashed lines indicate gaps corresponding to the *Stowaway* element and one copy of the target site.



**Figure 6.** Polyadenylation Sites of Three Hv-1 Gene Family Members Occur in *Stowaway* Sequences.

(A) Schematic of the 3' ends of three Hv-1 transcripts showing the positions of putative polyadenylation signals (open rectangles) in *Stowaway* sequences (black rectangles with open arrowheads extending to the poly[A] addition sites [A<sub>n</sub>]).

(B) Sequences of the 3' ends of each Hv-1 transcript aligned with the *Stowaway-Hv2* element (located within intron 3 of the barley Rubisco activase gene). The inverted repeat (black arrows), direct repeat (open arrows), putative polyadenylation signals (boxed sequences), and poly(A) tails (A<sub>n</sub>) are indicated. Gaps (dashed lines) were introduced for optimal alignment.

Wessler, 1992, 1994). It cannot be ruled out, however, that *Tourist* and *Stowaway* are solo long terminal repeats (LTRs) because the LTRs of some retroelements also have TIRs, albeit short (~5 bp). Retroelements transpose via an RNA intermediate and do not excise. The absence of detectable excision events in our study does not necessarily support the notion that *Tourist* and *Stowaway* are solo LTRs because excision for these family of elements may be rare and/or precise.

Although *Stowaway* and *Tourist* elements share no significant sequence similarity to other previously reported transposable elements, some aspects of their structures are reminiscent of the IS630-Tc1 transposon superfamily (Berg and Howe, 1989; Dreyfus and Emmons, 1991; Doak et al., 1994). Members of this superfamily share significant transposase sequence similarity and, similar to *Stowaway*, have a TA target sequence preference. Interestingly, the 1.6-kb Tc6 element of *Caenorhabditis*, a Tc1-like element, consists of a 765-bp TIR and has the potential to form inverted repeat DNA secondary structures similar to that of *Tourist* and *Stowaway* elements (Dreyfus and Emmons, 1991). Because no *Stowaway* element identified to date contains a significant open reading frame that would encode a transposase, it would be premature to suggest that *Stowaway* is a member of the IS630-Tc1 superfamily. Furthermore, elements belonging to the IS630-Tc1 superfamily have been found in many diverse species but have not as yet been identified in plants.

The correspondence of *Stowaway* element location with previously identified *cis*-acting regulatory domains provides strong

evidence that these elements have influenced the evolution of normal genes. For instance, *Stowaway-Hv4* and *Stowaway-Hv5* (*Hordeum vulgare*) provide polyadenylation signals and polyadenylation sites for their host genes. In addition, some elements (i.e., *Stowaway-Le1*, *Stowaway-Le2*, *Stowaway-Dc2*, and *Stowaway-St1*) may provide *cis*-acting regulatory regions to downstream genes. Previous reports of transposable elements supplying the *cis*-acting regulatory domains of normal genes are restricted to retrotransposons and retroposons. For example, the retrotransposons LTR-IS/MuRRS (murine retrovirus-related sequence), RTVL-H (retrovirus-like element with a histidine tRNA primer binding site), and THE-1 (transposon-like human element) provide polyadenylation signals for the mouse A1, human PLT (placental LTR terminated), and THE-1-containing genes, respectively (Paulson et al., 1987; Baumruker et al., 1988; Goodchild et al., 1992). The retroposon B2 has been identified in the 3' ends of several mouse transcripts, and, in one example, provides alternative polyadenylation signals for the mouse  $\gamma$ -phosphorylase kinase gene (Clemens, 1987; Maichele et al., 1993). Furthermore, a VL30 (virus-like element encoding 30S RNA)-like retroviral insertion confers androgen sensitivity on the mouse sex-limited protein gene (Stavenhagen and Robins, 1988), and IAP (intracisternal-A particles)-derived solo LTRs supply the promoters of the rat oncomodulin and mouse MIPP (mouse IAP-promoted placental) genes (Banville and Boie, 1989; Chang-Yeh et al., 1991). Because the regulatory requirements of most plant genes that harbor *Stowaway* elements are poorly

characterized, we are uncertain of the extent of element involvement in the regulation of these genes.

Finally, it is important to note the role played by computer-assisted sequence similarity searches in the discovery of the *Stowaway* and *Tourist* families. Because many of these degenerate elements share less than 65% sequence identity over their length, standard hybridization protocols would not have been useful in identifying family members. In light of the enormous amount of sequences currently being generated by several genome projects, similar data base searches will undoubtedly lead to the identification of other elements and provide additional examples of the intimate association of mobile elements and normal genes.

## METHODS

### DNA Sequence Analysis

The UWGCG (University of Wisconsin, Madison, Genetics Computer Group) and IG (IntelliGenetics, Inc., Mountain View, CA) computer program suites were accessed through the BioSciences Computational Resource, University of Georgia, Athens, GA. Data base searches were conducted using the programs FASTDB (IG), FASTA (UWGCG), and BLAST (National Center for Biotechnology Information, National Institutes of Health, Bethesda, MD) (Devereaux et al., 1984; Altschul et al., 1990). Pairwise and multiple alignments of element sequences were performed using the programs GAP and PILEUP (UWGCG), respectively. A gap penalty of 3.0 and gap length penalty of 0.3 were used, and complete elements were compared with their ends weighted. Minimum energy folding of element sequences was performed using the program FOLD (UWGCG) with DNA base pair and stacking energies as described previously (Breslauer et al., 1986). DNA secondary structures were visualized using the program SQUIGGLES (UWGCG).

### DNA Manipulations

*Oryza sativa* (International Rice Research Institute [IRRI], Los Baños, The Philippines, accession number IR25587-109-3-3-3-3), *O. punctata* (IRRI accession number 103006), and *O. eichingeri* (IRRI accession number 101422) genomic DNA was obtained from G. Kochert (University of Georgia, Athens). *Zea mays* subsp. *mexicana* (accession Iltis 28620) germplasm was acquired from J. Doebley (University of Minnesota, St. Paul, MN), and genomic DNA was isolated as previously described (Dellaporta et al., 1983). Oligonucleotides were synthesized corresponding to the sequences within or flanking intron 2 of the following genes. Nucleotide positions relative to the start of translation are given within parentheses. In the rice heat shock protein 82A gene (GenBank locus name OSHSP82A), the primer sequences are 5'-CATCTGGGGAGC-AGCTTGGG-3' (1558 to 1577) and 5'-TGAGGCGGCGCTCTCAAGG-3' (2481 to 2462); in the maize *P* gene (Athma et al., 1992), the primer sequences are 5'-ACACTCGGACCGTGAGAGG-3' (4510 to 4529) and 5'-GAGGTGGCTGGCGATCAGGG-3' (5029–5010). Polymerase chain reaction (PCR) amplification was performed as previously described (Bureau and Wessler, 1992), except that an annealing temperature of 65°C was used. PCR products were checked by agarose gel electrophoresis and cloned into a TA plasmid vector (Invitrogen, San Diego,

CA). Plasmid DNA isolation and dideoxy sequencing were performed using Qiagen (Chatsworth, CA) plasmid miniprep and Sequenase (United States Biochemicals) kits, respectively, as directed by the manufacturers.

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