# The Origin and Evolution of Six Miniature Inverted-Repeat Transposable Elements in Bombyx mori and Rhodnius prolixus 

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#### Abstract

Miniature inverted-repeat transposable elements (MITEs) are a specific group of nonautonomous DNA transposons, and they are distributed in a wide range of hosts. However, the origin and evolutionary history of MITEs in eukaryotic genomes remain unclear. In this study, six MITEs were identified in the silkworm (Bombyx mori). Five elements are grouped into four known superfamilies of DNA transposons, and one represents a novel class of MITEs. Unexpectedly, six similar MITEs are also present in the triatomine bug (Rhodnius prolixus) that diverged from the common ancestor with the silkworm about 370 Ma . However, they show different lengths in two species, suggesting that they are different derivatives of progenitor transposons. Three direct progenitor transposons (Sola1, hobo/Ac/Tam [hAT], and Ginger2) are also identified in some other organisms, and several lines of evidence suggested that these autonomous elements might have been independently and horizontally transferred into their hosts. Furthermore, it is speculated that the twisted-wing parasites may be the candidate vectors for these horizontal transfers. The data presented in this study provide some new insights into the origin and evolutionary history of MITEs in the silkworm and triatomine bug.


Key words: MITEs, origin, evolution, Bombyx mori, Rhodnius prolixus.

## Introduction

Transposons are discrete DNA fragments, and they are characterized by an inherent property of transposition and replication within their host genomes (Kidwell and Lisch 2001). An increasing number of studies have demonstrated that transposons are a major force to drive the evolution of genome complexity (Kazazian 2004).

Transposons can be classified into two classes (DNA and RNA transposons) based on their transposition intermediates. DNA transposons are a group of transposons that use a "cut-and-paste" mechanism to transpose (Wicker et al. 2007). They are characterized by terminal inverted repeats (TIRs) and target site duplication (TSD), and their autonomous copies encode a transposase that is responsible for their mobility. In the last decades, a specific group of nonautonomous elements that is derived from the autonomous DNA transposons has been found in plants (Bureau and Wessler
1992). These elements, named miniature inverted-repeat transposable elements (MITEs), are also flanked by TIRs and TSD. There are also other common features, including small size, $\mathrm{A}+\mathrm{T}$ richness, high copy number, potential stable secondary structure, a lack of transposase, and size and sequence homogeneity (Feschotte et al. 2002; Han et al. 2010). The most important feature distinguishing them from typical defective elements is their high copy number and structural homogeneity (Feschotte and Pritham 2007). MITEs can be classified into different superfamilies based on their similar structure (such as TIRs or TSD). The distribution of MITEs in the genome is obviously associated with genes (Oki et al. 2008). These results suggested that MITEs might have great effects on gene regulation and genome evolution. For example, MITEs may upregulate (MITEs containing regulatory motifs) or downregulate (MITE-derived small RNAs) the expression levels of host genes where they reside (Yang et al. 2005; Oki et al.

[^0]2008; Kuang et al. 2009). However, the origin and evolutionary history of MITEs in insects are poorly understood.

The taxonomic distribution of DNA transposons suggests that they are almost present in the whole eukaryotic tree of life (Feschotte and Pritham 2007). However, transposons should be eliminated by drift and/or selection in their original lineages. Horizontal transfer (HT) between isolated species seems to be the most important means by which transposons can avoid inevitable stochastic loss (Schaack et al. 2010). Besides, HT of a transposon into a new genome is also regarded as an important force to drive genome variation and biological innovation. Furthermore, almost all transposons have the ability of HT (Schaack et al. 2010). The classic documented examples of transposon HT are the $P$ and $T c 1 /$ mariner transposons of Drosophila (Daniels et al. 1990; Maruyama and Hartl 1991). Although 218 convincing cases of HT of transposons are described in invertebrates, vertebrates, and plants, the molecular mechanisms of most transposon HT have not been elucidated (Schaack et al. 2010). Recently, it has been demonstrated that the intimacy of parasitism might facilitate the HT of transposon (Yoshiyama et al. 2001; Laha et al. 2007; Gilbert et al. 2010).

Previously, we identified a novel cluster of the hobo/Ac/ Tam (hAT) superfamily in Bombyx mori and Rhodnius prolixus and found that this novel element generated MITEs in B. mori rather than in $R$. prolixus (Zhang et al. 2013). It is likely that this element invaded into their hosts independently by HT (Zhang et al. 2013). Here, we report the characterization and classification of six MITEs identified in both B. mori and $R$. prolixus. Meanwhile, the direct or related progenitors of these MITEs were also found in some other organisms. Moreover, our results indicated that progenitors of these MITEs might have been independently and horizontally transferred into their hosts for a long time. Finally, a model is provided for the origin and evolutionary history of these MITEs in the silkworm and the triatomine bug.

## Materials and Methods

## Data Resources

Bombyx mori assembled genomic sequences, expressed sequence tag (EST) sequences, tRNAs, 5 S rRNA, and 7SL RNA were downloaded from Silkworm Genome Database (SilkDB version 2, http://www.silkdb.org/silkdb/doc/download.html, last accessed October 29, 2013). Meanwhile, the full-length cDNA sequences of the silkworm were downloaded from the National Center for Biotechnology Information (NCBI) (accession numbers: AK377185-AK388575). Rhodnius prolixus genomic supercontig sequences, EST sequences, and transcript sequences were downloaded from VectorBase (Lawson et al. 2009). Both Manduca sexta and Me. moldrzyki whole genome shotgun (WGS) sequences were downloaded from NCBI (accession numbers: AIXA01000000 and AGDA00000000). Mengenilla moldrzyki EST sequences were downloaded from
a data repository website (http://datadryad.org/resource /doi:10.5061/dryad.ts058, last accessed October 29, 2013). Heliconius melpomene genomic sequences were downloaded from Butterfly Genome Database (http://www.butterflygen ome.org/, last accessed October 29, 2013). Leaf-cutter ant Atta cephalotes genome sequence and transcript sequences were downloaded from Hymenoptera Genome Database (http://hymenopteragenome.org/, last accessed October 29, 2013). Megachile rotundata and Hydra magnipapillata genomic sequences, and mRNA sequences of Meg. rotundata were obtained from NCBI (ftp://ftp.ncbi.nlm.nih.gov/genomes/, last accessed October 29, 2013). Transposase sequences of other known Sola, hAT, and Ginger transposons were obtained from Repbase (http://www.girinst.org/repbase/update/browse.php, last accessed October 29, 2013).

## Identification and Copy Number Determination of MITEs

Six nonautonomous DNA transposable elements (bm_648, Bmori_800.383, bm_701, bm_1364, bm_295 and bm_248) were obtained from BmTEdb (Xu et al. 2013). Their copies were extracted with 500 bp flanking sequences using our Perl script. Then, these retrieved sequences were aligned using MUSCLE (Edgar 2004) to determine the exact boundary of these transposons. Then, we reconstructed their consensus sequences by multiple alignments of their copies in the silkworm genome using DAMBE (Xia and Xie 2001). To determine the distribution of these transposons, their consensus sequences were used as queries in BlastN searches against the NCBI genome databases of the species for which a complete or near complete genome is available. A transposon was considered to be present in a species if any copy was at least 80\% identity at the nucleotide level to the consensus over at least $50 \%$ of its length. Consensus sequences for each transposon family identified in other species were also constructed based on multiple alignments of copies using DAMBE (Xia and Xie 2001). For families with only truncated copies or a few copies $(<10)$ in the genome, the best hit was chosen as consensus sequence. For families with uncertainty in its internal regions, their reliable $5^{\prime}$ and $3^{\prime}$ terminal regions were used to construct consensus sequence (Gilbert et al. 2010).

To estimate copy number of each transposon family, consensus sequences or representatives were used to mask all genomes where these transposons families were found using BlastN. Blast hits longer than 100 bp and at least 80\% identity at the nucleotide level to their consensus were used to calculate copy number (Gilbert et al. 2010). For MITEs, one copy was defined as full length according to the following criteria: $>90 \%$ nucleotide identity to the consensus, covering $>80 \%$ of the consensus and including complete TIRs (Han et al. 2010). As Novel_NA was not flanked by TIRs, copies that share at least $90 \%$ identity at the nucleotide level to the consensus and cover at least $90 \%$ of its length were considered as intact elements.

## Identification of Autonomous DNA Transposons and Evolutionary Analysis

When we used all MITEs identified in the silkworm as queries to do BlastN search against NCBI nt database, autonomous candidates responsible for the spread of two families (Sola1_NA and hAT1_NA) were found. Then, their potential transposases were predicted using getorf in EMBOSS-6.3.1 package (Rice et al. 2000). To determine the distributions of these two autonomous families, their sequences were used as queries in BlastN searches against the NCBI genome databases of the species for which a complete or near complete genome is available. A transposon was considered to be present in a species if any copy was at least $80 \%$ identity at the nucleotide level to the consensus over at least 50\% of its length.

For other four families, we identified their potential autonomous families using extensive analysis in the silkworm and the triatomine bug. Briefly, we searched the large DNA fragments that shared similar terminal sequences to the above four families. All DNA fragments ranging from $1,000 \mathrm{bp}$ to 10 kb flanked by the termini of these four families were obtained. Then, their potential transposases were predicted using getorf in EMBOSS-6.3.1 package (Rice et al. 2000). These candidates with open reading frame (ORF) were used as queries in BlastN searches against the NCBI genome databases, and the same standard mentioned earlier was used.

To determine the relationships of autonomous Sola1, hAT1, and Ginger2 elements identified in this study and known elements from other organisms (downloaded from Repbase, http://www.girinst.org/repbase/update/browse.php, last accessed October 29, 2013), we aligned amino acid sequences of their transposases using MUSCLE (Edgar 2004). Gap sites and variable regions were excluded, and then MEGA 4 (Tamura et al. 2007) was used to build phylogenetic trees.

## Sequence Analysis

The frequency distribution of sequence identities of MITEs in B. mori and $R$. prolixus was also determined. The percentage identity was based on the similarity between each copy and the respective consensus. Only copies longer than 100 bp were considered. Finally, we investigated insertion site preference of these six MITEs in B. mori and $R$. prolixus. To reflect their insertion preferences exactly, only copies flanked by intact TSDs were included in the analysis. Sequence logos were created by WebLogo (Crooks et al. 2004) based on 30 nt ( 15 upstream and 15 downstream) flanking the insertion site.

Sequences of three most conserved host genes, heat shock cognate 70 (Hsc70-4), Tubulin beta-3 (tub3), and elongation factor 1 alpha (EF1 $\alpha$ ), were used in the comparison with transposon $K a / K s$, with the purpose of testing HT hypothesis. Their accession numbers were listed in supplementary table S1, Supplementary Material online. Pairwise alignments of these
genes were created using MUSCLE (Edgar 2004), and each pairwise identity was calculated by Bioedit (Hall 1999). Codon alignments of hAT1 and these host genes were constructed by PAL2NAL software (Suyama et al. 2006). Then, the Ka/Ks ratio was calculated using DnaSP v5 (Librado and Rozas 2009).

## Polymerase Chain Reaction Analysis

Genomic DNA of the silkworm strain Dazao was obtained from State Key Laboratory of Silkworm Genome Biology (China). Genomic DNA of $R$. prolixus was kindly provided by Dr Ricardo N Araujo (Laboratório de Fisiologia de Insetos Hematófagos, Brasil). Polymerase chain reaction (PCR) primers for all MITEs identified in $B$. mori and $R$. prolixus were designed based on their flanking regions. The list of primers and expected product sizes were given in supplementary table S2, Supplementary Material online. PCRs were performed using the following thermal profiles: initial denaturation at $94^{\circ} \mathrm{C}$ for 4 min and then 30-35 cycles ( 32 cycles for Sola1_NA_RP and hAT1_NA_RP, 35 cycles for hAT2_NA_RP, and 30 cycles for other families in $B$. mori and $R$. prolixus) of $94^{\circ} \mathrm{C}$ for 40 s , 48$55^{\circ} \mathrm{C}$ for $50 \mathrm{~s} \quad\left(48^{\circ} \mathrm{C}\right.$ for hAT2_NA_RP, $50^{\circ} \mathrm{C}$ for Novel_NA_BM and $55^{\circ} \mathrm{C}$ for the rest families), and $72^{\circ} \mathrm{C}$ for 1 min , PCR products were run in $1 \%$ agarose gels in $1 \times$ Tris acetate-EDTA (TAE) buffer and visualized under UV light.

## Results and Discussion

## Identification and Analysis of Six MITEs in B. mori and R. prolixus

Six families of nonautonomous transposon families were obtained from BmTEdb (Xu et al. 2013). These reference sequences were not classified in BmTEdb. However, we demonstrated that five families were grouped into Sola-like, hATlike, Ginger-like, and Tc1/Mariner superfamilies based on their similarities to the characteristic features (TIRs and TSDs) of these superfamilies (table 1), and the last one may represent a novel class of transposons (please see the below results). We also demonstrated that these families may be MITE-like elements. (Each MITE identified in this study is named as $X \_N A \_\# \$$, where $X$ represents a superfamily, NA represents nonautonomous elements, \# is a letter representing the first letter of genus, and $\$$ is a letter representing the first letter of species. Each autonomous element is named as $\left.X_{-} \# \$.\right)$ as they were highly homogeneous in sequence and lack encoding sequences. For example, the majority (about 65\%) of members of Sola1_NA_BM were 300-340 bp in length (data not shown).

Unexpectedly, all these families are also identified in R. prolixus (supplementary table S 3 , Supplementary Material online) when we used the above silkworm transposons as queries to perform BlastN search against the $R$. prolixus genome. However, they showed different lengths in both species (fig. 1). This suggested that they were different

Table 1
Characteristics of Sola1_NA, hAT1_NA, Ginger2_NA, Novel_NA, pogo_NA, hAT2_NA, and their Autonomous Elements

| Group | Length (bp) | TSD | TIR | Copy <br> Number | Accession Number of Representatives |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Species |  |  |  |  |  |
| Sola1_NA |  |  |  |  |  |
| B. mori | 334 | AWWT | GGACCGTTTATTTGAAAGTGGTGTAAGTCCATTT | 83 (41) | KF558364 |
| R. prolixus | 709 | AWDT | GGACCGTTTATITTGAAAGTGGTGTAAG | 105 (40) | KF558358 |
|  | 1,489 ${ }^{\text {a }}$ | ND | ND | 1 | GL562232 1089812386 - |
| H. melpomene | 248 | AWDT | GGACCGTTTATTTTGAGAGTGGTGTAAGTCCATTT | 263 (228) | CAEZ01006881 1305413301 - |
| Danaus plexippus | 194 | ND | ND | 1 | AGBW01008816 62356428 + |
| M. sexta | 222 | ND | ND | 1 | AIXA01005774 1972919950 - |
| A. cephalotes | 2,313 ${ }^{\text {a }}$ | ND | GACCATTTATTCTGAAAGT | 6 | ADTU01013967 58948206 + |
| Meg. rotundata | 2,778 ${ }^{\text {a }}$ | ND | GGACCGTTTATTTGAAAGTGGTGCAAGTCCATTT | 69 | AFJA01007769 1760520382 - |
| hAT1_NA |  |  |  |  |  |
| B. mori | 546 | 8 bp | CAGTGATTCCCAAA | 51 (32) | KF558365 |
| R. prolixus | 619 | 8 bp | CAGTGATTCCCAAA | 27 (25) | KF558359 |
|  | 2,889 ${ }^{\text {a }}$ | 8 bp | CAGTGATTCCCAAA | 2 | GL563087 19810271983915 - |
| H. melpomene | 415 | 8 bp | CAGTGATTCCCAAA | 13 (4) | CAEZO1008298 75167940 - |
| M. sexta | 795 | 8 bp | CAGTGATTCCCAAA | 21 (15) | AIXA01005484 4711347907 + |
| Hy. magnipapillata | 2,916 ${ }^{\text {a }}$ | 8 bp | CAGTGATTCCCAAA | 93 | NW_002143368 803010961 - |
| Me. moldrzyki | 1,573 ${ }^{\text {a }}$ | ND | ND | ND | AGDA01034126 11573 + |
| Ginger2_NA |  |  |  |  |  |
| B. mori | 531 | WRTH | TGTTACGGTAAAAGTA | >120 (28) | KF558366 |
|  | 2,442 ${ }^{\text {a }}$ | NA | TGTTACGGTAGAAGTA | NA | BABH01026869 1285315294 + |
| R. prolixus | 651 | KHTM | TGTTACGGTAAAAGTA | 40 (25) | KF558360 |
|  | 3,767 ${ }^{\text {a }}$ | TATA | TGTTACGGTAAAAGTA | >1 | GL563081 872372876138 - |
| Me. moldrzyki | 599 | ND | TGTTACGGTAAAAGTA | 57 | AGDA01093000 32643854 - |
| Novel_NA |  |  |  |  |  |
| B. mori | 892 | TT | - | 37 (11) | KF558367 |
| Danaus plexippus | 678 | ND | - | 2 | AGBW01012094 7275073427 + |
| R. prolixus | 803 | T | - | 881 (604) | KF558361 |
| pogo_NA |  |  |  |  |  |
| B. mori | 270 | TA | CAGTAAAAGCTCTTTATTCGCG | >484 (15) | KF558368 |
| R. prolixus | 748 | TA | CAGTAAAAGCTCTITATTCGCG | 297 (197) | KF558362 |
| M. sexta | 851 | TA | CAGTAGAAGCTCTTTATTCG | 1 | AIXA01008424 1939620246 + |
| Danaus plexippus | 1,030 | TA | CAGTAAAATCTCTTTATTCGCG | 12 (3) | AGBW01000094 26273651 + |
| hAT2_NA |  |  |  |  |  |
| B. mori | 310 | 8 bp | CGGTGTTTCCCAAC | 224 (29) | KF558369 |
| R. prolixus | 956 | 8 bp | CGGTGTTTCCCAAC | 126 (53) | KF558363 |
| H. melpomene | 522 | 8 bp | CGGTGTTTCCCAAC | 10 (1) | CAEZ01008877 2047021038 + |
| M. sexta | 277 | 8 bp | CAGTGTTTCCCAAC | 78 (28) | AIXA01004312 6329263566 + |
| Me. moldrzyki | 236 | ND | ND | 5 | AGDA01059403 80658300 - |

Note.-W, A/T; R, A/G; D, AT/G; K, T/G; H, A/CT; M, A/C. ND, not determined. The number in bracket is the number of full-length copies in each MITE family. ${ }^{\mathrm{a}}$ The potential autonomous elements.
derivatives of the same progenitor transposons in $B$. mori and R. prolixus. A pairwise comparison of these transposons in both species revealed an extremely high identity (89-99.4\%) at the nucleotide level over the whole length (fig. 1). In R. prolixus, all these families also seemed to be MITEs because of their sequence and size homogeneity. The presence of these six MITEs in both species was confirmed by PCR (fig. 2).

To better understand their evolutionary history, we compared copy numbers of each family between B. mori and $R$. prolixus. Copy numbers of five families (Sola1_NA, hAT1_NA, Ginger2_NA, pogo_NA, and hAT2_NA) were from 27 to several hundreds, and these families had almost
the same copy number in both species (table 1). In contrast, Novel_NA family had a very larger copy number in $R$. prolixus than B. mori (881 vs. 37). In addition, most copies of Novel_NA_RP (604/881) were the verified full-length elements, suggesting that this family might experience a burst of transposition in $R$. prolixus. A similar phenomenon was also observed in another family (pogo_NA_RP). Indeed, star shape trees of Novel_NA_RP and pogo_NA_RP also suggested that they might have experienced one round of burst amplification (supplementary fig. S1, Supplementary Material online).

To further clarify the evolutionary history of these families, the distributions of percentage identity scores between copies


Fig. 1.-Structure and the homologous regions of six MITEs identified in the silkworm and triatomine bug. Black triangles are TIRs, and white rectangles are homologous regions of each transposon in both species. Gray regions represent the variable area of the Novel_NA family. The corresponding names and percentages of identity are shown on the left. The right numbers are the size of each consensus sequence.


Fig. 2.-Experimental verification of the presence of six MITEs identified in the silkworm ( $A$ ) and triatomine bug ( $B$ ). PCR fragments of expected sizes were obtained from both species. " $M$ " indicates marker, and the left numbers show their corresponding sizes.
and corresponding consensus or representatives were estimated (supplementary fig. S2, Supplementary Material online). The results showed that copies of six families (except Sola1_NA) were more conserved in $R$. prolixus than
B. mori. For example, R. prolixus Ginger2_NA family had an identity distribution with a peak at identity $99 \%$. However, Ginger2_NA family in B. mori had a distribution with a peak at identity $93 \%$, and it also included a long tail of more
divergent copies. It appears that these families might have experienced bursts of transposition in the recent past in R. prolixus but in the more distant past in B. mori. However, it should be noted that these observations may not reflect the exact evolutionary histories of these transposons because they may have different nucleotide substitution rates in two different host species.

Finally, we analyzed insertion site preferences of these families. Sola1_NA in B. mori and R. prolixus showed insertion preferences for the AWWT (W: AT) and AWDT (D: AT/G) tetranucleotide, respectively (supplementary fig. S3, Supplementary Material online). This is a hallmark of the Sola1 transposons (Bao et al. 2009). HAT1_NA and hAT2_NA tended to integrate in sequences having conserved TA in positions +4 and +5 (supplementary fig. S3, Supplementary Material online). However, hAT1_NA also showed an additional preference for integration in sequence having a conserved A in position +2 of their TSD in $B$. mori. Ginger2_NA family preferentially targeted 4-bp AT-rich sequences, TATA, an important characteristic of Ginger2 elements (Bao et al. 2010). Novel_NA family had a preferential insertion of T-rich sequences. Pogo_NA had strict insertion specificity for TA dinucleotide.

In summary, MITEs described here were strikingly similar to each other in B. mori and $R$. prolixus. They showed similar TIRs, high sequence identities, and similar targets for integration. All these data indicated that each family identified in both species originated from the same ancestral transposon and could be recognized as the same transposon evolving in two very different genomic backgrounds.

## Annotation of Novel_NA in B. mori and R. prolixus

Classification of transposons is complex and difficult because of their complicated and diverse structures. Therefore, classification of a transposon needs several lines of evidence. Novel_NA identified in this study did not have TIRs and did not show similarity to any known transposons. However, the characteristics of nonautonomous nature, small size, and high copy number suggested that it might be a short interspersed element (SINE)-like or MITE-like element. Here, we demonstrated that Novel_NA was a MITE-like element based on three lines of evidence.

First, SINEs originate from tRNA, 5 S rRNA, or 7 S rRNA genes as a result of their $5^{\prime}$ sequences sharing homology with these genes (Frenkel et al. 2004). Moreover, SINEs also harbor an RNA pol III promoter that is necessary for their transposition. These are two major characteristics that distinguish SINEs from other nonautonomous transposable elements. However, Novel_NA elements lacked a RNA polymerase III (pol III) promoter and did not show homology with any tRNA, 5S rRNA, or 7S rRNA genes of the silkworm. This suggested that Novel_NA might not be a SINE-like element. Recently, large-scale full-length cDNA of the
silkworm was determined (Suetsugu et al. 2013). A full-length Novel_NA, which was amplified by PCR from the silkworm genomic DNA (fig. 2A), was not correspondingly detected by using its sequence as a query in BlastN search against the above cDNA database. This suggested that the propagation of Novel_NA might not occur by reverse transcription.

Second, many MITEs identified in insects tend to generate $\mathrm{A}+\mathrm{T}$ biased TSD during their integration. For instance, HzMINE-2 elements from Helicoverpa zea and DINE-1 elements from Drosophila generated $\Pi$ dinucleotide TSD during their integration (Yang and Barbash 2008; Coates et al. 2011). In particular, the HzMINE-2 and DINE-1 insertion sites were flanked by similar duplications (TITT or WTTT). These nucleotide sequences provided the recognition sites where they integrated, and resulted in the creation of a $\Pi$ TSD. Interestingly, Novel_NA was also characterized by the presence of $\Pi \Pi T$ duplications (fig. 3). These findings suggested that Novel_NA might use the same integration mechanism as that of HzMINE-2 and DINE-1 elements.

Third, Novel_NA showed similar secondary structure to those of HzMINE-2 and DINE-1. They contain a $14 \mathrm{bp} 5^{\prime}-$ SIR (ACGTGAATACGTCT) that pairs with a reverse complementary $3^{\prime}$-SIR (AGACGTATTCACGT) and a 8 bp stem-loop structure downstream of the $3^{\prime}$-SIR (fig. 3). In addition, $5^{\prime}$ and $3^{\prime}$ boundaries of Novel_NA in $R$. prolixus are composed of TG and GT, and they are similar to those of HzMINE-2 and DINE-1 (TA and GT). This suggested that Novel_NA together with HzMINE-2 and DINE-1 might represent a novel class of MITEs in insects as a result of their unusual features distinguishing them from traditional MITEs. Moreover, their common features suggested that they could evolve from a common ancestor.

## Autonomous Transposon Progenitors of MITEs

Genetic and structural features of MITEs imply that they originated from full-length cognate autonomous DNA transposons (Feschotte and Pritham 2007). To our surprise, potential autonomous elements responsible for the spread of three families (Sola1_NA, hAT1_NA, and Ginger2_NA) were found (table 1). For Sola1_NA, the overall size of these significant hits was 2,313, 1,489, and 2,778 bp in $R$. prolixus, A. cephalotes, and Meg. rotundata, respectively (table 1 and supplementary fig. S4A, Supplementary Material online). Such a size was not expected for a so-called miniature transposon. Interestingly, one intact ORF (630 aa), which did not have premature stop codon was clearly identified for transposon in Meg. rotundata. It suggested that it might be a source of functional protein and was still active in Meg. rotundata. However, it was difficult for us to determine the exact full-length transposase encoding by this transposon in $A$. cephalotes and $R$. prolixus due to stop codons or frameshifts (supplementary fig. S4A, Supplementary Material online). These elements in different hosts revealed high

The 5 'terminal region of Novel_NA in B. mori and $R$. prolixus

## 5'-SIR

| scaffold1054 nscaf2801 | : GATAGTTATAATTTTGACGTGAATACGTCTTTAAAATTACTTCCATAGGGACGCAATCCAAAAAAACGAATGATAACAAAATCGGGTG |
| :---: | :---: |
| nscaf2902 | : TTAGCGGCAATTATTGACGTGAATACGTCTTTAAAATTACTTCCATAGGGAGGCAATCCAAAAAAACGAATGATAACAAAATCGGGGG |
| nscaf3052 | : AACGCCGCGATTTTTGACGTGAATGCGTCTTTAAAATTACTTCCATAGGGACGCAATCTAAAAAAACGAATGATAACAAAATCGGGGG |
| nscaf3066 | : AAATCTTTGGATTTTGACGTGAATACGTCTTTAAAATTACTTCTATAGGGAGGCAATCCAAAAAAACGAATGATAACAAAATCGGGGG |
| nscaf2838 | : ACTGTAATATTTTTTGACGTGAATACGTCTTTAAAATTACTTCCATAGGGACGCAATCCAAAAAAACGAATGATAACGAAATCGGGGG |
| scaffold1550 | : ACTGTAATATTTTTTGACGTGAATACGTCTTTAAAATTACTTCCATAGGGACGCAATCCAAAAAAACGAATGATAACGAAATCGGGGG |
| nscaf3053 | : ACAAACATTTTTTTTGACGTGAATACGTCTTTAAAATTACTTCCATAGGGACGCAATCCAAAAAAACGAATGATAACAAAATCGGGGG |
| GL563178 | : TCGAAGGATTTTTTTGACGTGAATACGTCTTTACATTTGCTATCATATGGAGGCAATCCAAAAAACCTAATGATAACAAAATCGGGGG |
| GL563097 | : TAATTTTCAATTTTTGACGTGAATACGTCTTTAAATTTGCTTTCATATGGAAGCAATCCGAAAAACCTAATGATAACAAAATCGGGGG |
| GL563082 | : GGAATTTTTCATTTTGACGTGAATACGTCTTTAAATTTGCTATCATATGGAGGCAATCCAAAAAAACTAATGATAACAAAATTGGGGG |
| GL563055 | : AAAGGGCTACTTTTTGACGTGAATCCGTCTTTAAATTTGCTATCATATGGAGGCAATCCAAAAAACCTAATGATAACAAAATCGGGGG |
| GL562501 | : CGTATTAAGTTTTTTGACGTGAATACGTCTTTAAATTTGCTATCATATGGAGGCAATCCAAAAAACCTAATGATAACAAAATCAGGGG |
| GL558139 | : CGACAGCACAGTTTTGACGTGCATACGTCTTTAAATTTGCTATCATATGGAGGCAATCCAAAAAACCTAATGATAACAAAATCGGGGG |
| GL563163 | : AGCGATGCAGTTTTTGACGTGAATACGTCTTTAAATTTGCTATCATATGGAGGCAATCCAAAAAACCTAATGATAACAAAATCGGGGG |
| GL563154 | : CCCAATGGTTCTTTTGACGTGAATACGTCTTTAAATTTGCTATCATATGGAGGCAATCCAAAAAACCTAATGATAACAAAATCGGGGG |
| GL563021 | : ATAATTTAATTTTTTGACGTGAATACGTCTTTAAATTTGCTATCATATGGAGGCAATCCAAAAAGCCTAATGATAACAAAATCGGGGG |
| GL563038 | AAAAATTAAATTTTTGACGTGAATACGTCTTTAAATTTGCTATCATATGGAGGC-----CTAAAACCTAATGATAACAAAATCGGGGG |

## Species

B. mori
R. prolixus
accession
5710-4856 204405-203625 8366622-8365702 733182-734098 923709-924632 645280-644317 772-1735 3317-2396 4470004-4470934 262609-261674 2825343-2824398 320504-321457
554396-555342
24491-25438
1589765-1588830
519973-519020
151332-150362
905636-904712

Fig. 3.-Alignment of $5^{\prime}$ and $3^{\prime}$ terminal regions from full-length Novel_NA in the silkworm and triatomine bug. Black color shows their subterminal inverted repeat (SIR) and gray indicates their $3^{\prime}$ stem loop. The underlines represent the potential target sites of Novel_NA. Accession numbers of Novel_NA transposons in their genomes are also shown.
sequence identity ( $>85.8 \%$ ), suggesting that they should belong to the same family (supplementary fig. S4A and table S3, Supplementary Material online). For hAT1_NA, its autonomous elements were 2,889, 2,916, and 1,573 bp in length in R. prolixus, Hy. magnipapillata and Me. moldrzyki, respectively (supplementary fig. $S 4 B$, Supplementary Material online). Their potential coding region could be determined. An alignment of this element from different hosts revealed a very high level of identity ( $>88.2 \%$ ) over the whole length of the consensus sequences, suggesting these elements descended from the same active ancestral element (supplementary fig. $S 4 B$ and table S3, Supplementary Material online). For Ginger2_NA, its autonomous element in B. mori and R. prolixus was 2,442 and 3,767 bp, respectively. However, we failed to identify a significant coding region for Ginger2_RP in the $R$. prolixus genome (supplementary fig. S4C, Supplementary Material online). By contrast, a significant coding region of Ginger2_BM was found when we used its nucleotide sequence to perform translated search against the Repbase (http://www.girinst.org/censor/index.php, last accessed October 29, 2013). Meanwhile, an alignment of these two elements showed that their similarity (93.2\%
pairwise nucleotide identity) is restricted to the terminal regions (supplementary fig. S4C and table S3, Supplementary Material online). However, their internal regions do not show any similarity. Therefore, we inferred that the internal region of Ginger2_RP was foreign sequences. Phylogenetic analyses of these elements with other known transposons indicated that they were grouped into the corresponding superfamily (supplementary fig. S5, Supplementary Material online). Multiple alignments of nonautonomous elements and their master elements suggested that they originated by internal deletions from master elements (fig. 4A-C).

Although no direct autonomous element of pogo_NA has been found so far, TIRs of this family shared some similarity to Emigrant (fig. 4D), a MITE family identified in Arabidopsis thaliana (Casacuberta et al. 1998), and to a MITE family Wujin from yellow fever mosquito Aedes aegypti (Tu 1997). This indicated that pogo_NA and these MITEs could be grouped into the same MITE family. Especially, its TIRs also had some similarity to an autonomous pogo-like element, the human Tigger1 element (Robertson 1996). Therefore, our results supported the hypothesis that pogo-like elements had a strong tendency to generate MITEs (Feschotte and

## A



D


Fig. 4.-Homologies in $5^{\prime}$ and $3^{\prime}$ terminal regions between Sola1_NA (A), hAT1_NA (B), hAT2_NA (B), Ginger2_NA (C), pogo_NA (D), and their potential progenitor transposons. Alignment of the terminal sequences was done using MUSCLE. TIRs of the Tigger1, Wujin, and Emigrant families are from Feschotte and Mouchès (2000). "-" indicates a gap, and conserved sites are in white type on a green/blue/red/yellow background.

Mouches 2000). We also failed to find the autonomous elements for hAT2_NA family. However, TIR similarity (11/14) between hAT1_NA and hAT2_NA suggested that they may also belong to the same family (fig. $4 B$ ). Moreover, they tended to integrate in similar target sequences (supplementary fig. S3, Supplementary Material online). All these results indicated that hAT2_NA might use the transposition machinery of autonomous elements similar to that for hAT1_NA.

## Evidence for HT

The screening of complete or near-complete genome sequences available in NCBI revealed that three highly similar autonomous elements (Sola1, hAT1, and Ginger2) were present in six invertebrate genomes, including the silkworm (B. mori), the triatomine bug ( $R$. prolixus), the leaf-cutter ant (A. cephalotes), the alfalfa leafcutting bee (Meg. rotundata), the twisted-wing parasite (Me. moldrzyki), and the freshwater hydrozoan (Hy. magnipapillata) (table 1 and fig. 5).

The autonomous Sola1 family, Sola1, existed in the triatomine bug, the leaf-cutter ant and the alfalfa leaf-cutting bee. In each of the species, a representative sequence was recovered because we were not able to construct their complete consensus sequences as a result of low copy number or fragmentation in their genomes. The nucleotide sequence identities between these transposons were unexpectedly high (>85.8 for all interspecific pairwise comparisons). This
reflects that these transposons might have been horizontally transferred into these hosts because these three host species are distantly related (fig. 5). For instance, there was 93.9\% sequence identity between the triatomine bug Sola1 (Hemiptera, Reduviidae) and the alfalfa leafcutting bee Sola1 (Hymenoptera, Megachilidae), and their hosts diverged about 370 Ma (Hedges et al. 2006). Surprisingly, Sola1 elements from closely related alfalfa leaf-cutting bee and leaf-cutter ant (Hymenoptera, Formicidae) showed higher level of nucleotide sequence divergence than Sola1 transposons from the triatomine bug and the alfalfa leaf-cutting bee (supplementary table S3 and fig. S4A, Supplementary Material online). High sequence identity of Sola1 transposons from divergent species and lower sequence identity of these elements from closely related species suggested horizontal transmission for Sola 1 family.

The autonomous hAT family, hAT1, was present in the triatomine bug, the twisted-wing parasite, and the freshwater hydrozoan Hy. magnipapillata (table 1 and fig. 5). The hAT1 family in the Hy. magnipapillata genome was the only family whose consensus sequence could be reconstructed. However, this family in other two species appeared in partial copies or low copy number, and their representatives were chosen as respective consensuses. The nucleotide identity between the sequences from the triatomine bug and the twistedwing parasite was extremely high (97\%), suggesting TH (supplementary table S3 and fig. S4B, Supplementary


Fig. 5.-_The taxonomic distribution of Sola1_NA, hAT1_NA, Ginger2_NA, Novel_NA, pogo_NA, hAT2_NA, and their autonomous elements among invertebrate genomes for which complete or near-complete genome sequences are available. Presence of these transposon families in each lineage are denoted by rectangles.

Material online). Indeed, their hosts diverged from the common ancestor of Hemiptera and Strepsiptera about 370 Ma (Hedges et al. 2006). Similar patterns of sequence identity of hAT1 were observed between the fresh water hydrozoan and the twisted-wing parasite/the triatomine bug (88-92\%). These species belong to different phyla (Cnidaria
and Arthropoda) that diverged from their common ancestor $>800 \mathrm{Ma}$ (Hedges et al. 2006). Therefore, it was most likely that $h A T 1$ elements might have been horizontally transferred into different host species.

The autonomous Ginger family, Ginger2, was only identified in the silkworm and the triatomine bug (table 1 and
fig. 5). The member of this family appeared in partial or truncated copies in their respective hosts, which made it difficult to reconstruct reliable consensus sequences. The alignment of the Ginger2 sequences from the silkworm and the triatomine bug showed an extremely high level of nucleotide identity (93.2\%) across a region of 1,217 bp (supplementary table S3 and fig. S4C, Supplementary Material online) despite the silkworm and the triatomine bug split at 370 Ma . This suggested that Ginger family might have been horizontally transferred into two host species.

There are generally three criteria used to detect HT events: 1) high sequence similarity of TEs from divergent taxa, 2) topological differences between TE and host phylogenies, and 3) discontinuous distribution of a transposon among closely related taxa (Schaack et al. 2010). Although the strikingly high level of sequence identity of the above three autonomous elements among such highly divergent invertebrates provided the strongest evidence to support the hypothesis that they had transferred horizontally into these lineages, such deduction only based on sequence identity may be problematic because we did not consider variable evolutionary rates that had been shown for some transposons (Malik et al. 1999). Therefore, making HT conclusion of these transposons should be cautious.

To obtain more evidence for HTs of these transposons, we investigated the phylogenetic distributions of the Sola1, hAT1, and Ginger2 families. Our results showed that they were patchy distributions in species (fig. 5). If these transposons had been vertically transferred from a common ancestor, it was difficult to explain the fact that there were as many as 11, 7, and 7 independent losses of the Sola1, hAT1, and Ginger2 families, respectively. For example, Sola1 family was identified only in $A$. cephalotes genome, it was absent in other six closely related sequenced ant genomes, including another leaf-cutter ant Acromyrmex echinatior, which shared a common ancestor with $A$. cephalotes about 8-12 Ma (Schultz and Brady 2008). Similar phenomenon was also observed for hAT1 family. It was present in the fresh water hydrozoan (Cnidaria) and insects (Arthropoda) but absent in Nematoda and Deuterostomes (data not shown). Additionally, it showed the discontinuous distribution across different insect species (fig. 5). Finally, the distribution of Ginger2 family was also patchy, because it was found only in two insect species (the silkworm and the triatomine bug).

A high level of sequence identity of these transposons among distantly related species may also result from intense purifying selection. To directly compare with the coding genes from host species, three most conserved genes (Hsc70-4, tub3, and EF1a) were examined. We retrieved full-length CDS sequences of these genes (except EF1a from Me. moldrzyki) from their hosts (supplementary table S1, Supplementary Material online). $K a / K s$, a ratio of the nonsynonymous and synonymous changes between two sequences, was calculated by DnaSP v5 (Librado and Rozas 2009). For Sola1 and Ginger2
families, we were not able to determine their exact coding sequences in at least two species, and they were not included in this analysis (supplementary fig. S4, Supplementary Material online). The frequency of nonsynonymous changes for the hAT1 families varied from 0.0295 to 0.1588 , and the frequency of synonymous changes ranged from 0.0290 to 0.0574 (supplementary table S3, Supplementary Material online). Importantly, $\mathrm{Ka} / \mathrm{Ks}$ for this family ranged from 0.2509 to 0.9831 for all interspecific comparisons, which is consistent with very low to no significant purifying selection. However, DnaSP v5 did not give an output for Ks for almost all pairwise comparisons of Hsc70-4 and tub3 (supplementary table S3, Supplementary Material online) possibly because synonymous changes were saturated. Meanwhile, EF1a comparisons were in accord with high selection pressure because the ratio Ka/Ks varied from 0.0067 to 0.0240 . Therefore, these results suggested that high level of sequence identity of the hAT1 family did not result from intense purifying selection.

Taking the above evidence together, HT was the only reasonable explanation for such a high level of sequence identity among these transposons in highly divergent host species. This was consistent with previous studies in which the authors provided sufficient evidence supporting HTs of the transposons in the silkworm and the triatimine bug (Thomas et al. 2010; Zhang et al. 2013).

## Mechanism for Transfer

Strepsiptera is an order of insect parasitoids that infect at least 35 families of insects belonging to seven orders, including Thysanura, Blattodea, Mantodea, Orthoptera, Heteroptera, Hymenoptera, and Diptera (Kathirithamby 1989). Meanwhile, some species of Strepsiptera are not completely host specific, and some of them might parasitize several genera and species (Kathirithamby 1989). During the process of parasitism, they obtain nutrients from a single host (Kathirithamby 2009). This order has two major groups: Stylopidia and Mengenillidia. Free living of female adult is one of the most important characters that distinguish Mengenillidae from other families (Kathirithamby 1989). Mengenilla moldrzyki is a newly discovered species and belongs to Mengenillidae group in the strepsipteran order (Pohl et al. 2012). However, its direct host was unknown.

Three families (hAT1, Ginger2_NA, and hAT2_NA) were identified in Me. moldrzyki (the twisted-wing parasite). These families shared an extremely high level of sequence identity with homologs in the silkworm, the triatomine bug, and the freshwater hydrozoan, which diverged from a common ancestor at least 300 Ma (table 1 and fig. 5). Several previous studies had demonstrated that the intimacy of parasite-host interaction might facilitate HTs of transposons (Yoshiyama et al. 2001; Laha et al. 2007; Gilbert et al. 2010). Although we could not directly connect the twisted-wing


Fig. 6.-A model for the origin and evolution of MITEs in the silkworm and triatomine bug. Gray regions indicate the transposase of autonomous elements, and blue triangles are TIRs. The potential processes involved in the origin and evolution of MITEs in the silkworm and in the triatomine bug. 1. An ancestor transposon was first introduced into the ancestor of Lepidopteran insects by HT. Then, many internal deletion copies (including MITEs) of this autonomous element are generated. Meanwhile, this autonomous copy might be stochastically lost or fossilized in the ancestor genome. Finally, the silkworm MITEs are obtained from its ancestor through vertical inheritance. 2. An ancestor transposon was horizontally transferred into the triatomine bug genome. Then, kinds of newly internal deletion versions of this autonomous element were generated in this species. It is speculated that the twisted-wing parasites may be the candidate vectors for these transfers.
parasite with insects studied, these transposons identified in this insect parasitoid might also suggest that their transfers were likely facilitated by parasitism. Afterall, Me. moldrzyki is the only species of Strepsiptera whose complete genome has been sequenced (Niehuis et al. 2012). Meanwhile, there were only 33 insect species whose complete or near-complete genomes were available (fig. 5). This means that our present view of the phylogenetic distribution of these transposons may be biased by the sampling of insect genome projects. Therefore, to have more insights into the origin and evolutionary history of these transposons, it is better to focus on a systematic analysis of the phylogenetic distribution of these transposons in future. To the best of our knowledge, this was the first time to identify transposons in the order Strepsiptera and to report the phenomenon of HT of transposons in these entomophagous parasitoids. Moreover, the published complete genome of Me. moldrzyki will also provide a good opportunity for studying the effect of parasite-host interaction on the evolution of transposons and speciation.

## A Model for the Origin and Evolutionary History of MITEs in the Silkworm and the Triatomine Bug

Using MITEs identified in the silkworm as queries, five families were also found in recent published genomes of three other Lepidpoteran insects (Danaus plexippus, H. melpomene, and M. sexta) (Zhan et al. 2011; Heliconius Genome Consortium, 2012) (table 1 and fig. 5). This suggested that these MITEs in
the silkworm were vertically inherited. Thus, it is most likely that the progenitor elements of these MITEs might have been horizontally introduced into the ancestor of lepidopteran insects and MITEs were generated within the ancestor genome. Then the silkworm obtained MITEs from its ancestor through vertical inheritance (fig. 6). Meanwhile, the progenitor elements of the MITEs have been horizontally invaded into the triatomine bug genome and then generated the MITEs within its genome (fig. 6). Furthermore, we speculate that the twisted-wing parasites may be the candidate vectors for these HTs (fig. 6). In summary, our results provide some new insights into the origin and evolutionary history of the MITEs in B. mori and $R$. prolixus.

## Supplementary Material

Supplementary tables S1-S3 and figures S1-S5 are available at Genome Biology and Evolution online (http://www.gbe. oxfordjournals.org/).

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