# Recurrent Horizontal Transfers of *Chapaev* Transposons in Diverse Invertebrate and Vertebrate Animals

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### **Abstract**

Horizontal transfer (HT) of a transposable element (TE) into a new genome is regarded as an important force to drive genome variation and biological innovation. In addition, HT also plays an important role in the persistence of TEs in eukaryotic genomes. Here, we provide the first documented example for the repeated HT of three families of *Chapaev* transposons in a wide range of animal species, including mammals, reptiles, jawed fishes, lampreys, insects, and in an insect bracovirus. Multiple alignments of the *Chapaev* transposons identified in these species revealed extremely high levels of nucleotide sequence identity (79–99%), which are inconsistent with vertical evolution given the deep divergence time separating these host species. Rather, the discontinuous distribution amongst species and lack of purifying selection acting on these transposons strongly suggest that they were independently and horizontally transferred into these species lineages. The detection of *Chapaev* transposons in an insect bracovirus indicated that these viruses might act as a possible vector for the horizontal spread of *Chapaev* transposons. One of the *Chapaev* families was also shared by lampreys and some of their common hosts (such as sturgeon and paddlefish), which suggested that parasite—host interaction might facilitate HTs.

Key words: horizontal transfer, Chapaev, viruses, parasite-host interaction.

#### Introduction

Transposable elements (TEs) are fragments of DNA that can move from one place to a new genomic location in their hosts and often make up a large fraction of eukaryotic genomes (Feschotte and Pritham 2007). TEs are divided into two classes based on transposition mechanisms: Class I or RNA elements transpose via reverse transcription of an RNA intermediate; Class II or DNA elements transpose via a DNA intermediate and most do so using a so-called "cut and paste" mechanism (Craig et al. 2002). Horizontal transfer (HT), known as the exchange of genetic material between isolated species, plays an important role in transposon biology and genome evolution (Schaack et al. 2010; Wallau et al. 2012; Ivancevic et al. 2013). The P element of Drosophila was the first TE shown to have been introduced via HT (Daniels et al. 1990). Virtually all major types of TEs have been shown to be capable of HT in a wide variety of eukaryotes (Bartolomé et al. 2009; Schaack et al. 2010; Thomas et al. 2010; Wallau et al. 2012; Ivancevic et al. 2013). However, the majority of reported horizontal transposon transfers involves drosophilid flies (Schaack et al. 2010). Meanwhile, the extent of this phenomenon remains unclear and the molecular mechanisms underlying HT remain largely mysterious. Two facilitating mechanisms have received support recently: One is host–parasite relationships (Yoshiyama et al. 2001; Gilbert et al. 2010); the other is that DNA viruses can act as transposon vectors (Fleming and Summers 1991; Jehle et al. 1998; Turnbull and Webb 2002; Marquez and Pritham 2010; Schaack et al. 2010; Dupuy et al. 2011; Gilbert et al. 2014).

Chapaev transposons represent a relatively new superfamily of DNA transposons, which were first identified in 2007 (Kapitonov and Jurka 2007) and recently affiliated to a larger "megafamily" widespread in eukaryotes dubbed CMC for Chapaev–Mirage–CACTA (Yuan and Wessler 2011). Hallmarks of the Chapaev transposons are generally 3- to 4-bp target site duplication (TSD), terminal-inverted repeats

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with the invariable "5-CAC and GTG-3" termini (Kapitonov and Jurka 2007; Yuan and Wessler 2011). As in most DNA transposons, Chapaev transposases are characterized by the presence of a conserved "DDE" motif in the predicted catalytic domain as well as additional conserved residues diagnostic of the CMC group (Yuan and Wessler 2011). However, the biology and evolution of *Chapaev* transposons remain largely uncharacterized. To expand our knowledge on Chapaev transposons, we have carried out a detailed analysis of the characteristics and evolution of three Chapaev families, which we identified in a variety of invertebrate and vertebrate species, as well as in an insect bracovirus. We provide evidence that the widespread taxonomic distribution of these elements is the result of multiple HT events likely facilitated by both parasitism and viruses.

## **Materials and Methods**

#### **Animal Materials**

For the silkworm Bombyx mori, strain Dazao was obtained from the State Key Laboratory of Silkworm Genome Biology (China) and its DNA extraction was based on the standard techniques (Nagaraja and Nagaraju 1995). Asian Swallowtail Papilio xuthus was purchased from Shanghai Qiuyu Biotechnology Co., Ltd (China). DNA or tissue samples of the Arctic lamprey Lethenteron camtschaticum, the Pacific bluefin tuna Thunnus orientalis, the channel catfish Ictalurus punctatus, the turnip sawfly Athalia rosae, and the lizard Anolis carolinensis were kindly provided by related researchers (please see Acknowledgments for details). Then, their total DNAs were extracted using TIANamp Genomic DNA Kit (TIANGEN). Meanwhile, quality of DNAs extracted from these species was examined on 1% agarose gel electrophoresis.

#### **DNA** Collection

The assembled *B. mori* genome sequence was downloaded from Silkworm Genome Database (SilkDB version 2, http:// www.silkdb.org/silkdb/doc/download.html, last accessed June 10, 2014). The Monarch butterfly Danaus plexippus genome resource (version 2) was obtained from MonarchBase (Zhan and Reppert 2013) through the web site at http://monarchbase.umassmed.edu/resource.html (last accessed June 10, 2014). The triatomine bug Rhodnius prolixus genomic supercontig sequences were downloaded from VectorBase (Lawson et al. 2009) at http://www.vectorbase. org (last accessed June 10, 2014). The whole-genome shotgun (WGS) sequences of the sea lamprey Petromyzon marinus, Arctic lamprey, Pacific bluefin tuna, dragonfly Ladona fulva, turnip sawfly, tenrec Echinops telfairi, and lizard were downloaded from the National Center for Biotechnology http://www.ncbi.nlm.nih.gov/, Information (NCBI, accessed June 10, 2014).

Identification and Copy Number Calculation of Chapaev Elements

A previously uncharacterized Chapaev element (named Garfield\_BM) was discovered in the silkworm genome when proteins of Chapaev elements from Repbase (Jurka et al. 2005) were used as gueries in tBLASTn (default parameters) (Altschul et al. 1990) searches against the draft genome assembly of the silkworm (Zhang H-H, Zhang Z, unpublished data). The sequences of Merrow PM and Conan ET (see Nomenclature for details) were obtained from Repbase (Kapitonov and Jurka 2007). Then, their nucleotide sequences were used as initial gueries (BLASTn [Altschul et al. 1990] using default parameters) to find these Chapaev elements in other genomes available at the NCBI, including nucleotide collection (nr/nt), genome survey sequences (GSS), expressed sequence tag (EST), high throughput genomic sequences (HTGS), and the WGS databases (as of September 2013) (Thomas et al. 2010). They were considered in a species if hits were >80% identical to the guery over at least 300 bp because Merrow transposons identified in all teleost fishes were guite short (from 300 to 800 bp) (table 1).

In order to determine the boundary of these elements, the best hits identified in a species (for which genome sequences were available) were blasted using BLASTn (Altschul et al. 1990) against each genome. Then, these retrieved sequences (identity and coverage >80% of the guery sequences) were extracted with 500-bp flanking sequences using our Perl script, and they were aligned using MUSCLE (Edgar 2004) to determine their boundary. In addition, copies (4–50) in each species (supplementary table S1, Supplementary Material online) were also aligned using MUSCLE, and their consensus sequences were reconstructed using the above multiple alignments in each genome using DAMBE (Xia and Xie 2001) after gaps were removed. If one genome sequence contained highly fragmented copies or low copy number (<3), the best hit represented the consensus sequence. Also, if these Chapaev elements were identified in a nonsequenced species, the best hit identified in this species was used as the consensus sequence. If these transposons identified in one species were chimaeric, they were excluded from the following analysis.

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Next, we used these respective consensus sequences to mask each genome in which Merrow, Garfield, and Conan were identified to estimate copy number. All blast hits with more than 100 bp and 80% identity were used to calculate copy number. Because there are many chimaeric copies in the tenrec genome, only elements that were at least 40% coverage to the consensus sequence were considered in estimating copy number. Three miniature inverted-repeat transposable elements (MITEs) derived from Merrow, Garfield, and Conan were also discovered in Pacific bluefin tuna, silkworm, and lizard. As MITEs are generally <600 bp, size and sequence homogeneity (Feschotte et al. 2002), their copy numbers were calculated based on the following criteria: 1) All

Table 1
Characteristics of *Merrow, Garfield,* and *Conan* Newly Identified in This Study

Group	Common Names	TE Names	Length (bp)	Сору	Exon1 (aa)	Exon2 (aa)	Representatives
				Number			
Species							
Merrow							
Petromyzon marinus	Sea lamprey	Merrow_PM <sup>a</sup>	2,451	>254	78	484	AEFG01041997
Lethenteron camtschaticum	Arctic lamprey	Merrow_LC	2,455	>249	78	484	KF965286
Acipenser transmontanus	White sturgeon	Merrow_AT	807	n/d	_	188	DR976541
Ictalurus punctatus	Channel catfish	Merrow_IP	833	n/d	_	_	KF965284
Ictalurus furcatus	Blue catfish	Merrow_IF	661	n/d	_	_	FD224147
Polyodon spathula	Mississippi paddlefish	Merrow_PS	304	n/d	_	_	JX448770
Thunnus orientalis	Pacific bluefin tuna	MerrowN1_TO	577	18	_	_	KF965282
Ladona fulva	Dragonfly	Merrow_LF	1,922	105	75	233	APVN01033993
Garfield							
Bombyx mori	Silkworm	Garfield_BM	2,289	>7	78	478	AADK01000850
		GarfieldN1_BM	468	97	_	_	KF965283
Rhodnius prolixus	Triatomine bug	Garfield_RP	2,289	16	78	478	ACPB02036275
Cotesia sesamiae Mombasa bracovirus	Viruses	Garfield_MB	2,291	n/d	77	477	EF710639
Papilio xuthus	Asian Swallowtail	Garfield_PX	1,254	n/d	_	149	KF965285
Athalia rosae	Turnip sawfly	Garfield_AR	1,328	2	_	306	KF965288
Danaus plexippus	Monarch butterfly	Garfield_DP	2,092	3	78	478	AGBW01002745
Conan							
Echinops telfairi	Tenrec	Conan_ET <sup>a</sup>	1,865	88	461	_	AAIY02038089
Anolis carolinensis	Lizard	Conan_AC	3,363	1	_	_	AAWZ02011613
		ConanN1_AC <sup>a</sup>	443	40	_	_	KF965287
		ConanN2_AC	295	281	_	_	KF965289

Note.—n/d, not determined, as the data were obtained from sequences deposited in the nucleotide collection (nr/nt) database, EST database, GSS database or HTGS database. —, Not found.

fragments showed more than 80% identity and coverage to their consensus sequences and 2) fragments were considered to be a single insertion when they were separated by less than 200 bp (Granzotto et al. 2011). Meanwhile, there were two subfamilies of MITEs identified in lizard (ConanN1\_AC and ConanN2\_AC) and the length of ConanN1\_AC was about 150 bp longer than that of ConanN2\_AC. Therefore, fragments of ConanN2\_AC were assigned to be a single copy when they were separated by less than 100 bp.

# Sequence Analysis

Potential open reading frame of *Chapaev* elements used in this study was predicted using FGENESH (http://linux1.soft berry.com/berry.phtml, last accessed June 10, 2014), GENSCAN (http://genes.mit.edu/GENSCAN.html, last accessed June 10, 2014), or getorf in EMBOSS-6.3.1 package (Rice et al. 2000) with the default parameters. Multiple alignments of these elements were created by MUSCLE (Edgar 2004). Shading and minor manual refinements of these aligned sequences were deduced using Genedoc (Nicholas et al. 1997). Each pairwise identity was calculated by Bioedit (Hall 1999) after all ambiguous and gapped sites were removed.

We also downloaded Chapaev3-like transposons, which were deposited in Repbase Update (Jurka et al. 2005) at Genetic Information Research Institute (http://www.girinst. org, last accessed June 10, 2014). Then, their nucleotide sequences were used as queries to do BLASTN (default parameters) (Altschul et al. 1990) against their respective available whole-genome sequences. All full-length or nearly full-length copies of each element were extracted with 100-bp flanking sequences using our Perl script. These sequences were aligned using MUSCLE to determine their TSD. Chapaev elements that only had highly fragment copies in their host genome were not included in this analysis. Chapaev paralogous empty sites were identified using the similar method described by previous studies (Marguez and Pritham 2010). Chapaev orthologous sites were determined by synteny analysis of 5,000 bp flanking these Chapaev transposons insertion sites.

Four data sets (one amino acid transposase sequence data set and three nucleotide sequence data sets) were created and used for phylogenetic analyses. The amino acid database consisted of *Chapaev* elements obtained from Repbase and discovered during the course of this study to determine the phylogenetic diversity of the *Chapaev* transposons. The other three databases of nucleotide sequences were created by respective full-length or nearly full-length copies of *Merrow*,

<sup>&</sup>lt;sup>a</sup>Chapaev transposons deposited in Repbase, and other Chapaev transposons were newly identified in this study.

Garfield, and Conan identified in this study to determine the relationship of copies of the same Chapaev element in different hosts.

The amino acid data set was aligned using MUSCLE (Edgar 2004), and a neighbor-joining tree was constructed using MEGA4 (pairwise deletion, Poisson correction model, 1,000 bootstrap replicates; Tamura et al. 2007). The remaining three databases were also aligned using the multiple sequence alignment program MUSCLE (Edgar 2004), and all ambiguous sites were manually excluded because there were a few ambiguous sites whereas most transposons from different species were aligned. Then, the best-suited nucleotide substitution models for these data were selected using Akaike information criterion (AIC) in Modeltest3.6 (Posada and Crandall 1998). The best-suited nucleotide substitution models for Merrow. Garfield and Conan were K81uf+G, HKY+G, and TVM+G, respectively. Then, phylogenetic trees were created using MrBayes 3.1.2 software (Ronquist and Huelsenbeck 2003) until the values of the average standard deviation of split frequencies were stably below 0.01.

# Age Analyses and Relative Insertion Periods

The timing of amplification of transposons in each species could be estimated by calculating the sequence divergence between copies and the ancestral sequence and by applying its neutral mutation rate (Waterston et al. 2002; Pace and Feschotte 2007). To estimate age of each copies of Merrow, Garfield and Conan, only copies spanning at least 50% of their consensus sequence were used in this analysis (Pagan et al. 2010). Then, they were aligned using MUSCLE (Edgar 2004), and the amount of nucleotide substitution (k) between each insertion and its respective consensus was estimated using Kimura 2-parameter distance method (Kimura 1980). Then, the insertion time of each element was estimated by the formula T = k/2r (Li 1997), where T corresponds to the insertion time in millions of years, k corresponds to the number of nucleotide substitutions per site, and r corresponds to the neutral mutation rate of the species lineage. If we accept that the elements from each other within a genome evolve neutrally since their insertion, the rate of neutral evolution available for their host nuclear genes might be employed. We used the neutral mutation rates for tenrec  $(2.9173 \times 10^{-9})$ /site/year; Pace et al. 2008), sea lamprey  $(1.9 \times 10^{-9})$ /site/year; Kuraku and Kuratani 2006), and Arctic lamprey  $(1.9 \times 10^{-9})$ /site/year; Kuraku and Kuratani 2006). Because a neutral mutation rate is not available for silkworm, we applied an estimated mutation rate previously published in Lepidoptera  $(1.909 \times 10^{-8})$ /site/year; Simonsen et al. 2011). Because there is no reliable neutral mutation rate available for other species or their close related taxa, these species were not included in this analysis. The phylogenetic tree of species in this study was based on Timetree of life (Hedges et al. 2006) and Taxonomy in NCBI. Divergence times of species were taken from the literature (Douzery et al. 2004; Peterson et al. 2004; Hedges et al. 2006; Kuraku and Kuratani 2006; Wiegmann et al. 2009). Divergence times between the channel catfish and the blue catfish Ictalurus furcatus, between the tobacco hornworm Manduca sexta and the silkworm, and between turnip sawfly and the ants are unknown. Therefore, their divergence times depicted in the phylogenetic tree were only for illustrative purposes.

# Testing for Purifying Selection

To test for purifying selection, codon alignments of *Merrow* and *Garfield* as well as elongation factor 1-alpha (*EF-1* $\alpha$ ) genes of their hosts were created using PAL2NAL software (http:// www.bork.embl.de/pal2nal/, last accessed June 10, 2014; Suyama et al. 2006). Because there are no complete coding sequences of transposase of Conan transposon in A. carolinensis (Conan AC) due to stop codons or frameshifts. Conan transposon was not included in this analysis. Then, synonymous (ds) and nonsynonymous (dn) divergences between them, as well as their ratio (dn/ds) were calculated using the SNAP tool in the HIV Sequence Database (http://www.hiv.lanl. gov, last accessed June 10, 2014; Korber 2002). Codon bias as determined by the effective number of codon ( $N_c$ ) value was computed using CodonW (Wright 1990).

In addition, multiple alignments of 40-50 copies (at least 50% coverage to their consensus sequences) GarfieldN1 BM (extracted from silkworm), Conan ET (extracted from tenrec), ConanN1\_AC, and ConanN2\_AC (extracted from lizard) were used to built neighbor-joining tree in MEGA 4 (Tamura et al. 2007), with p-distance model, pairwise deletion and 1,000 bootstrap replicates.

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# Polymerase Chain Reaction and Sequencing of Merrow, Garfield, and Conan

To validate the presence of Merrow, Garfield, and Conan identified computationally, their polymerase chain reaction (PCR) primers were designed using their flanking or internal sequences (supplementary table S2, Supplementary Material online). PCR was carried out with an initial denaturation step of 4 min at 95 °C followed by 32-35 cycles of 40 s at 95 °C, 40 s at 55–58 °C, and 2 m at 72 °C. Then, PCR products were run in 1% agarose gels in 1× Tris acetate-ethylenediaminetetraacetic acid buffer and visualized under UV light. Purified PCR products were cloned into PMD-19 cloning vector (TaKaRa). One random clone of each species was selected and sequenced.

# Nomenclature

We note that the *Merrow* identified in the sea lamprey P. marinus and the Conan identified in the tenrec E. telfairi and lizard A. carolinensis were previously designated with different names in Repbase (Kapitonov and Jurka 2007). However, the Repbase nomenclature for these elements

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was a potential source of confusion. For example, the sea lamprey Merrow family has been named Chapaev3-1 PM in Repbase, whereas the Conan families described in tenrec and lizard have been named corresponding Chapaev3-1\_ET and Chapaev3-3N1 AC. The Repbase nomenclature would seem to imply that Chapaev3-1\_PM and Chapaev3-1\_ET are more closely related to each other than to Chapaev3-3N1 AC. However, our results clearly show that Chapaev3-1 ET and Chapaev3-3N1\_AC belong to the same family (Conan) whereas Chapaev3-1\_PM falls within a distinct family (Merrow) (fig. 1 and table 1). Furthermore, our result shows that these families were not restricted to these species but are also present in many other species (table 1). Thus for simplicity and clarity, we decided to introduce the corresponding names Merrow and Conan for these two families. To the best of our knowledge, no members of the Garfield family have been characterized previously or deposited in Repbase.

Widespread Invasions by Chapaev Transposons

## **Results and Discussion**

Identification and Characterization of *Merrow*, *Garfield*, and *Conan* 

While investigating DNA transposons in the assembled genome of the silkworm B. mori, we discovered a previously uncharacterized family of transposon we designated as Garfield BM. A consensus sequence for Garfield BM was reconstructed by aligning multiple copies extracted from the B. mori genome assembly. The consensus length is 2,289 bp long and is predicted to contain two exons encoding a 556 amino acid (aa) transposase (Tpase) (supplementary fig. S1, Supplementary Material online, and table 1). The Tpase displays three highly conserved motifs [C(2)C, LH, and H(4)H] characteristic of the Chapaev superfamily of transposons (Yuan and Wessler 2011). Phylogenetic analysis based on a multiple alignment with representatives of the Chapaev transposases available in Repbase (fig. 1) places the silkworm transposon within the Chapaev3 subgroup (Kapitonov and Jurka 2007). In addition, the silkworm Chapaev elements were associated with a 3-bp putative TSD of 5'-TWA-3' consensus sequence (supplementary table S3, Supplementary Material online). We also observed that the nucleotide adjacent to the apparent TSD was always an "A" on the 5'-end and a "T" on the 3'-end (supplementary fig. S2, Supplementary Material online). To determine if these characteristics are shared with other members of the Chapaev3 group, we also analyzed the insertion bias of Chapaev3-like elements deposited in Repbase (Jurka et al. 2005) and found that all Chapaev3 transposons examined were also flanked by "TWA" TSDs and inserted between "A" and "T" (supplementary table S3 and fig. S2, Supplementary Material online). Furthermore, paralogous empty sites (i.e., homologous sites identified within the same genome but lacking the transposon insertion) confirmed that *Chapaev3*-like elements create a "TWA" TSD upon insertion (supplementary fig. S3, Supplementary Material online).

It has been documented that DNA transposons are capable of invading a variety of species by means of HT (Schaack et al. 2010). Because the level of sequence similarity between Chapaev3 transposases from widely diverged animal species appeared to be inconsistent with the phylogenetic relationships of their hosts (Kapitonov and Jurka 2007), we carried out a detailed investigation of the taxonomic distribution and evolution of Merrow, Garfield, and Conan. We used their consensus sequences as queries in BLASTn (Altschul et al. 1990) searches of all NCBI databases. These searches yielded highly significant hits (e value ranging from 0 to  $6 \times e^{-120}$ ) in a wide range of animal species and in an insect bracovirus (table 1). To rule out database artifacts or contamination, we sought to obtain experimental validation for the presence of these transposons in several of these species by PCR amplification from genomic DNA using primers internal or flanking one of these transposons followed by sequencing of cloned PCR products. We were able to obtain genomic DNA for seven animal species and for all of them confirmed the presence of Merrow, Garfield, and Conan we detected in the corresponding whole-genome assemblies (fig. 2) (GenBank accession numbers KF965282-KF965289).

For each species, we then reconstructed consensus ancestral sequences of Merrow, Garfield, and Conan (see Materials and Methods). Besides consensus sequences of Merrow\_PM and Conan\_ET, we also note that another consensus sequence (ConanN1\_AC) has been deposited previously in Repbase (Kapitonov and Jurka 2007). To the best of our knowledge, all other Merrow, Garfield, and Conan were newly identified in this study (fig. 1 and table 1). Phylogenetic analyses of consensus Tpase sequences confirmed that Merrow, Garfield, and Conan represent three distinct families within the Chapaev3 group (fig. 1). The phylogenetic analysis also suggested that Merrow and Garfield were more closely related to each other and might descend from a relatively recent common ancestor (fig. 1). Indeed, both Merrow and Garfield Tpases are encoded by two exons (fig. 3) whereas most other Chapaev3 Tpases appear to be encoded by a single exon (Kapitonov and Jurka 2007; data not shown). However, pairwise sequence similarity between any Merrow and Garfield consensus sequences was still no greater than 66% at the nucleotide level, suggesting that they represent distinct transposon families (Wicker et al. 2007).

Nonautonomous Elements Derived from *Garfield* and *Conan* 

MITEs are a group of nonautonomous elements, which was first discovered in maize (Bureau and Wessler 1992). Generally, MITEs originate from a particular deletion derivative of an autonomous DNA transposon that is subsequently

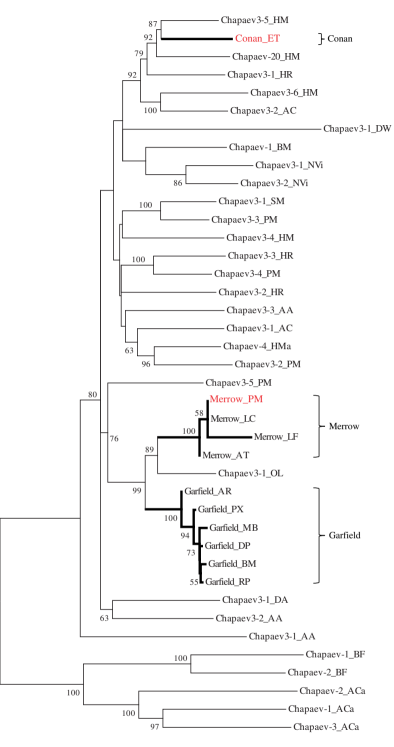


Fig. 1.—Phylogenetic relationships of Chapaev3 transposases. The neighbor-joining tree was constructed using MEGA4 (pairwise deletion, Poisson correction model, 1,000 bootstrap replications) based on a multiple alignment of amino sequences of the Chapaev3 transposases with five representatives (Chapaev1\_BF, Chapaev2\_BF, Chapaev1\_ACa, Chapaev2\_ACa, and Chapaev3\_ACa) obtained from Repbase as an outgroup. Bootstrap values > 50% were shown. Clusters of Merrow, Garfield, and Conan were displayed using thick line. Merrow\_PM and Conan\_ET were deposited in Repbase, and they were shown using red color. However, others were newly identified in this study. Species abbreviations: HM/HMa, Hydra magnipapillata; ET, Echinops telfairi; HR, Helobdella robusta; AC, Anolis carolinensis; DW, Drosophila willistoni; BM, Bombyx mori; NVi, Nasonia vitripennis; SM, Schmidtea mediterranea; PM, Petromyzon marinus; AA, Aedes aegypti; LC, Lethenteron camtschaticum; LF, Ladona fulva; AT, Acipenser transmontanus; OL, Oryzias latipes; AR, Athalia rosae; PX, Papilio xuthus; MB, Cotesia sesamiae Mombasa bracovirus; DP, Danaus plexippus; RP, Rhodnius prolixus; DA, Drosophila ananassae; BF, Branchiostoma floridae; ACa, Aplysia californica.

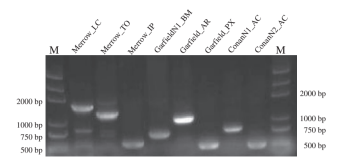


Fig. 2.—Experimental verification of the presence of *Merrow*, *Garfield*, and *Conan* identified in this study. PCR fragments of the expected sizes were obtained from species studied. All PCR products were confirmed by cloning and sequencing. "M" represents the marker. Species abbreviations: LC, *Lethenteron camtschaticum*; TO, *Thunnus orientalis*; IP, *Ictalurus punctatus*; BM, *Bombyx mori*; AR, *Athalia rosae*; PX, *Papilio xuthus*; AC, *Anolis carolinensis*.

amplified to high copy number to form an homogenous subfamily of nonautonomous elements (Feschotte and Pritham 2007). In this study, we found that two MITE families were direct internal deletion derivatives of one full-length Chapaev transposon in the silkworm and lizard (supplementary fig. S4, Supplementary Material online). Our results also showed that ConanN1\_AC and ConanN2\_AC were two relatively old subfamilies of lizard nonautonomous transposons as most of their copies were 80-90% identity to their consensus sequences. This is consistent with a previous proposal that members of the Chapaev DNA transposon superfamily have long been transpositionally inactive in the anole lizard (Novick et al. 2010). By contrast, GarfieldN1 BM seems to have experienced a recent burst transposition in the silkworm as all copies shared more than 91% identity to their ancestral sequence (data not shown). This level of divergence would imply a peak of amplification of GarfieldN1 BM at about 0.8-1.6 Ma (supplementary fig. S5, Supplementary Material online) based on neutral substitution rates previously estimated for lepidopterans (Simonsen et al. 2011). The presence of these MITEs in silkworm and anole lizard was experimentally validated by PCR using their flanking sequences to design primers (fig. 2; GenBank accession numbers KF965283, KF965287, and KF965289).

# Evidence for HTs

Multiple alignments of *Merrow, Garfield*, and *Conan* identified in this study revealed a strikingly high level of interspecific sequence identity (79–99%). Importantly, the level of nucleotide sequence identity is not only limited to coding regions but also extended to noncoding regions of the elements (supplementary fig. S6, Supplementary Material online). In many cases, the level of nucleotide sequence identity of these transposons is unexpectedly high when considering the deep

divergence of their host species (fig. 4). For example, Merrow identified in lampreys and teleost fishes as well as dragonfly shared 87-98% pairwise sequence identity. However, lampreys and jawed fishes diverged approximately 500 Ma and they shared a last common ancestor with dragonfly more than 700 Ma (supplementary table S3, Supplementary Material online; Hedges et al. 2006). A similarly elevated level of sequence identity (87–99%) of Garfield and Conan identified in different insect orders (Lepidoptera, Hymenoptera, and Hemiptera), insect viruses, and vertebrates was also observed (Tables S5 and S6). These insect orders diverged from each other more than 300 Ma (Hedges et al. 2006). In addition, tenrec and lizard diverged from a common ancestor approximately 300 Ma (fig. 4; Hedges et al. 2006). Thus, the extreme level of sequence similarity of Merrow, Garfield, and Conan across such distant species strongly suggests that these transposons invaded their hosts through repeated HT events.

To obtain additional evidence supporting HT, we investigated whether some ancestral copies of Merrow, Garfield, and Conan could be found at orthologous genomic positions in those species in which they were identified. The results showed that none of these transposons was present at orthologous positions in the species studied, with the exception of Arctic lamprey and sea lamprey, where full-length Merrow transposons were found at orthologous positions and therefore must have inserted before the divergence of these lampreys (supplementary fig. S7, Supplementary Material online). Next, we examined the taxonomic distribution of these three transposons and found that it was highly discontinuous and inconsistent with the phylogeny of their host species. For example, Garfield was only identified in turnip sawfly, but it was undetectable in the genome of 10 other species of hymenopteran insects (Zhang et al. 2013). Similarly, Conan was present in the tenrec (an afrotherian mammal) and the anole lizard, but it was not found in any of the dozens of other mammalian and reptilian genomes currently available in the databases.

Several additional lines of evidence rule out the possibility that Merrow, Garfield, and Conan were vertically inherited from the last common ancestor of these species. First, the topology of the phylogenetic tree of Merrow, Garfield, and Conan is incongruent with that of the host species (fig. 1 and supplementary fig. S8, Supplementary Material online). Furthermore, Merrow identified in closely related teleost fishes showed higher level of nucleotide sequence divergence than those from lampreys and teleost fishes, which diverged approximately 500 Ma (supplementary table S4, Supplementary Material online). Similarly, Garfield identified within insects of Lepidoptera exhibited higher sequence divergence at the nucleotide level than Garfield transposons from Lepidoptera and other insect orders (Hymenoptera and Hemiptera) (supplementary table S5, Supplementary Material online). In addition, we found no evidence that purifying

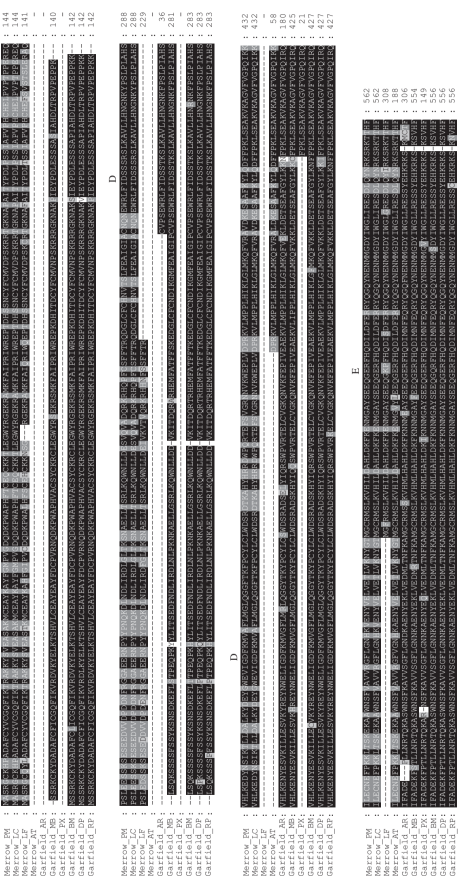


Fig. 3.—Multiple alignments of consensus sequences of Merrow and Garfield transposases. An alignment of these elements was created by MUSCLE. Shading and minor manual refinements of these aligned sequences were deduced using Genedoc. The "DDE" catalytic core encoded by their transposases was also shown. Species abbreviations: PM, Petromyzon marinus; LC, Lethenteron camtschaticum; Ladona fulva; AT, Acipenser transmontanus; AR, Athalia rosae; MB, Cotesia sesamiae Mombasa bracovirus; PX, Papilio xuthus; BM, Bombyx mori, DP, Danaus plexippus; RP, Rhodnius prolixus.

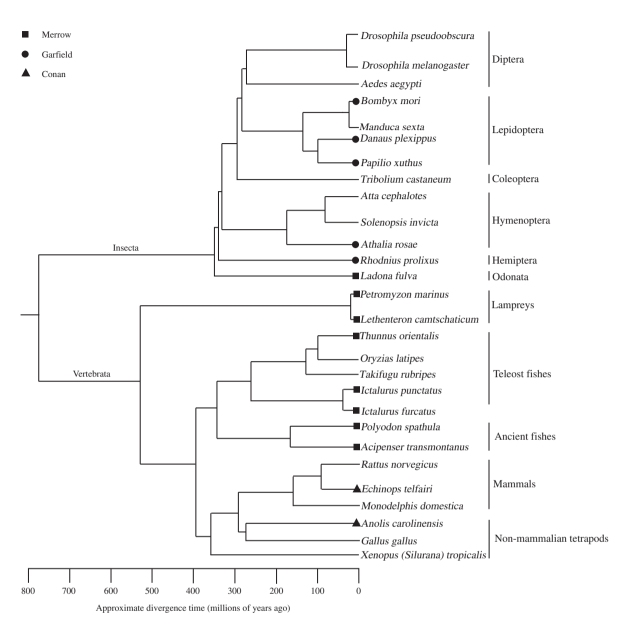


Fig. 4.—Schematic representation of a phylogenetic tree of animal lineages, estimated divergence times (Ma) and species distribution of *Merrow*, *Garfield*, and *Conan* identified in this study. The inferred HT events of these three different families were denoted by solid square, solid circles, and solid triangle, respectively.

selection and codon bias could account for the high level of conservation of *Merrow*, *Garfield*, and *Conan* identified in such widely divergent species. Phylogenetic analysis of these transposons obtained from each species showed a star-like shape, an indicative of a single rapid amplification from one master element followed by the accumulation of discrete mutations in each copy (supplementary fig. S9, Supplementary Material online). This evolutionary pattern is consistent with the neutral evolution typical of DNA transposons (Hartl et al. 1997; Feschotte and Pritham 2007). For all autonomous *Merrow* and *Garfield*, we found that the level of synonymous

divergence (ds) between species was considerably lower than that expected between such highly diverged taxa. For example, lampreys and jawed fishes separated more than 500 Ma (Hedges et al. 2006), yet the ds between lampreys and the white sturgeon *Acipenser transmontanus Merrow* consensus was 0.0088 (supplementary table S7, Supplementary Material online). Similarly, the ds values between *Garfield* consensus sequences of insects were all lower than 0.0718. Importantly, dn/ds for *Merrow* and *Garfield* varied from 0.2632 to 1.3068, consistent with low to no significant purifying selection acting on these transposons (supplementary

table S7, Supplementary Material online). Furthermore, ds values for a well conserved housekeeping gene such as the elongation factor gene  $EF-1\alpha$  were at least ten times higher (from 0.5696 to 3.4936) than those based on the sequences of the corresponding Garfield (from 0.0092 to 0.0718) (Tables S7 and S8). Together, these data suggested that strong purifying selection was not responsible for the high level of sequence identity of these transposons (at least for Garfield) across these widely diverged species. Codon bias as determined by the effective number of codon  $(N_c)$  value is known to represent a potential source of selective constraint on synonymous nucleotides (Wright 1990). N<sub>c</sub> values varied from 21 (one codon per aa—high bias) to 61 (all codons used equally—no bias) (Wright 1990). N<sub>c</sub> values for all transposons identified in this study was 49–59 (supplementary table S9, Supplementary Material online), suggesting that codon bias was also not responsible for the observed high sequence identity. Finally, an inferred insertion period of Merrow, Garfield, and Conan postdated the radiation of two species where these transposons resided (supplementary fig. S5, Supplementary Material online). The only exception to this pattern was the inferred amplification time (10-17 Ma) of Merrow in lampreys, which fell within the divergence time between these two species (10–30 Ma; Kuraku and Kuratani 2006), and was consistent with the findings that Merrow elements occupy orthologous positions in the two lampreys (see above and supplementary fig. S7, Supplementary Material online). Interestingly, our dating (26–36 Ma) for the invasion of Conan\_ET in the tenrec fell within the range inferred for the HT cases previously reported for several hAT transposons in diverse tetrapods, including the tenrec (15–46 Ma; Pace et al. 2008; Gilbert et al. 2010). Tenrecs are confined to Africa (Poux et al. 2005) and the anole lizard (which acquired a nearly identical Conan element) most likely has been endemic to South America (Roughgarden 1995). As the African and American continents separated much earlier (>65 Ma; Marshall et al. 1979) than the inferred introduction of Conan in the tenrec lineage, these observations suggest that Conan underwent a transoceanic movement, similarly to and around the same time as other widely horizontally transferred transposons (Pace et al. 2008; Gilbert et al. 2010). Interestingly, the Garfield element identified in Cotesia sesamiae Mombasa bracovirus (EF710639) was seemingly fulllength and capable to encode an apparently intact Tpase (fig. 3), suggesting that it might represent a recent acquisition by this insect virus.

Together, these data indicate that the most plausible scenario to explain the distribution of *Merrow*, *Garfield*, and *Conan* examined in this study is that these transposons were transferred horizontally into multiple species lineages and subsequently expanded within each genome. HT events between insects and insect bracoviruses (Thomas et al. 2010), between lampreys and teleost fishes (Kuraku et al. 2012), and between the tenrec and lizard (Pace et al. 2008; Gilbert et al.

2010) have been previously described. Thus, we speculate that these taxa have a higher propensity for exchanging genetic material

We also note that HT of *Conan* between tenrec and lizard has been alluded to in Repbase Reports (Kapitonov and Jurka 2007). However, to our knowledge HTs of *Merrow* and *Garfield* have not been reported previously. The clear phylogenetic separation of these three families of *Chapaev* transposons (fig. 1) indicates that these three families have been independently transferred into multiple hosts.

# Possible Vectors and Factors Facilitating HT

Merrow, Garfield, and Conan were identified in such a wide range of species (including not only lampreys, jawed fishes, lizard, tenrec but also silkworm, two distant butterflies, turnip sawfly, triatomine bug, dragonfly, and a bracovirus), suggesting that multiple vectors and mechanisms might be involved in the HTs of these transposons. Garfield discovered in the bracovirus was of particular interest because bracoviruses might represent an ideal vector for the horizontal spread of these transposons among species. These viruses create an obligatory relationship with parasitic wasps, and they only replicate in the ovary cells of wasps. Then, fully formed viral particles in the wasp ovary are injected into the lepidopteran larvae by the wasps. Thus, the intimate association between the parasitoid and their lepidopteran hosts might provide ample opportunity for the HTs of transposons. Indeed, there have been several documented examples of exchanging genetic materials between DNA viruses and their insect hosts (Fleming and Summers 1991; Jehle et al. 1998; Marguez and Pritham 2010; Thomas et al. 2010; Dupuy et al. 2011; Gilbert et al. 2014).

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Another interesting finding was the identification of nearly identical *Merrow* in lampreys and teleost fishes (table 1). Lampreys are opportunistic parasitic feeders that attach to large fish using their cup-like mouth to suck their blood and body fluids. The exchange of large amounts of blood between lampreys and their fish hosts during this parasitic interaction might provide a potential route for the horizontal spread of transposons, as suggested previously for *Tc1*-like transposons (Kuraku et al. 2012). Interestingly, lampreys are known to commonly parasitize sturgeons (Patrick et al. 2009) and paddlefish (Hardisty and Potter 1971), two species that harbor *Merrow* elements nearly identical to those of lampreys (table 1).

# **Supplementary Material**

Supplementary tables S1–S9 and figures S1–S9 are available at *Genome Biology and Evolution* online (http://www.gbe.oxfordjournals.org/).

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# **Literature Cited**

- Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. 1990. Basic local alignment search tool. J Mol Biol. 215(3):403–410.
- Bartolomé C, Bello X, Maside X. 2009. Widespread evidence for horizontal transfer of transposable elements across *Drosophila* genomes. Genome Biol. 10:R22.
- Bureau TE, Wessler SR. 1992. Tourist: a large family of small inverted repeat elements frequently associated with maize genes. Plant Cell 4:1283–1294.
- Craig , NL, Cragie , R, Gellert , M, Lambowitz , AM, editors. 2002. Mobile DNA II. Washington (DC): American Society for Microbiology.
- Daniels SB, Peterson KR, Strausbaugh LD, Kidwell MG, Chovnick A. 1990. Evidence for horizontal transmission of the P transposable element between *Drosophila* species. Genetics 124:339–355.
- Douzery EJ, Snell EA, Bapteste E, Delsuc F, Philippe H. 2004. The timing of eukaryotic evolution: does a relaxed molecular clock reconcile proteins and fossils? Proc Natl Acad Sci U S A. 101:15386–15391.
- Dupuy C, Periquet G, Serbielle C, Bézier A, Louis F, Drezen JM. 2011. Transfer of a chromosomal Maverick to endogenous bracovirus in a parasitoid wasp. Genetica 139:489–496.
- Edgar RC. 2004. MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Res. 32:1792–1797.
- Feschotte C, Jiang N, Wessler SR. 2002. Plant transposable elements: where genetics meets genomics. Nat Rev Genet. 3:329–341.
- Feschotte C, Pritham EJ. 2007. DNA transposons and the evolution of eukaryotic genomes. Annu Rev Genet. 41:331–368.
- Fleming JG, Summers MD. 1991. Polydnavirus DNA is integrated in the DNA of its parasitoid wasp host. Proc Natl Acad Sci U S A. 88: 9770–9774.
- Gilbert C, et al. 2014. Population genomics supports baculoviruses as vectors of horizontal transfer of insect transposons. Nat Commun. 5: 3348.
- Gilbert C, Schaack S, Pace JK 2nd, Brindley PJ, Feschotte C. 2010. A role for host–parasite interactions in the horizontal transfer of transposons across phyla. Nature 464:1347–1350.
- Granzotto A, Lopes FR, Vieira C, Carareto CM. 2011. Vertical inheritance and bursts of transposition have shaped the evolution of the BS non-LTR retrotransposon in *Drosophila*. Mol Genet Genomics. 286:57–66.
- Hall TA. 1999. BioEdit: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. Nucleic Acids Symp Ser. 41:95–98.

- Hardisty M, Potter I. 1971. The biology of lampreys. In: Hardisty M, Potter I, editors. The general biology of adult lamprey. New York: Academic Press. p. 127–206.
- Hartl DL, Lohe AR, Lozovskaya ER. 1997. Modern thoughts on an ancyent marinere: function, evolution, regulation. Annu Rev Genet. 31: 337–358
- Hedges SB, Dudley J, Kumar S. 2006. TimeTree: a public knowledgebase of divergence times among organisms. Bioinformatics 22:2971–2972.
- Ivancevic AM, Walsh AM, Kortschak RD, Adelson DL. 2013. Jumping the fine LINE between species: horizontal transfer of transposable elements in animals catalyses genome evolution. Bioessays 35: 1071–1082.
- Jehle JA, Nickel A, Vlak JM, Backhaus H. 1998. Horizontal escape of the novel Tc1-like lepidopteran transposon TCp3.2 into *Cydia pomonella* granulovirus. J Mol Evol. 46:215–224.
- Jurka J, et al. 2005. Repbase Update, a database of eukaryotic repetitive elements. Cytogenet Genome Res. 110:462–467.
- Kapitonov VV, Jurka J. 2007. Chapaev—a novel superfamily of DNA transposons. Repbase Rep. 7:774–781.
- Kimura M. 1980. A simple method for estimating evolutionary rates of base substitutions through comparative studies of nucleotide-sequences. J Mol Evol. 16:111–120.
- Korber B. 2002. HIV signature and sequence variation analysis. In: Rodrigo AG Jr, Learn GH, editors. Computational analysis of HIV molecular sequences. Dordrecht (The Netherlands): Kluwer Academic Publishers. p. 55–74.
- Kuraku S, Kuratani S. 2006. Time scale for cyclostome evolution inferred with a phylogenetic diagnosis of hagfish and lamprey cDNA sequences. Zoolog Sci. 23:1053–1064.
- Kuraku S, Qiu H, Meyer A. 2012. Horizontal transfers of Tc1 elements between teleost fishes and their vertebrate parasites, lampreys. Genome Biol Evol. 4:817–824.
- Lawson D, et al. 2009. VectorBase: a data resource for invertebrate vector genomics. Nucleic Acids Res. 37:D583–D587.
- Li WH. 1997. Molecular evolution. Sunderland (MA): Sinauer Associates. Marquez CP, Pritham EJ. 2010. Phantom, a new subclass of mutator DNA transposons found in insect viruses and widely distributed in animals. Genetics 185:1507–1517.
- Marshall LG, Butler RF, Drake RE, Curtis GH, Tedford RH. 1979. Calibration of the great American interchange. Science 204:272–279.
- Nagaraja GM, Nagaraju J. 1995. Genome fingerprinting of the silkworm, *Bombyx mori*, using random arbitrary primers. Electrophoresis 16: 1633–1638.
- Nicholas KB, Nicholas HB, Deerfield DW. 1997. GeneDoc: analysis and visualization of genetic variation. EMBNEW News 4:14.
- Novick PA, Smith JD, Floumanhaft M, Ray DA, Boissinot S. 2010. The evolution and diversity of DNA transposons in the genome of the lizard *Anolis carolinensis*. Genome Biol Evol. 3:1–14.
- Pace JK 2nd, Feschotte C. 2007. The evolutionary history of human DNA transposons: evidence for intense activity in the primate lineage. Genome Res. 17:422–432.
- Pace JK 2nd, Gilbert C, Clark MS, Feschotte C. 2008. Repeated horizontal transfer of a DNA transposon in mammals and other tetrapods. Proc Natl Acad Sci U S A. 105:17023–17028.
- Pagan HJ, Smith JD, Hubley RM, Ray DA. 2010. PiggyBac-ing on a primate genome: novel elements, recent activity and horizontal transfer. Genome Biol Evol. 2:293–303.
- Patrick H, Sutton T, Swink W. 2009. Lethality of sea lamprey parasitism on lake sturgeon. Trans Am Fish Soc. 138:1065–1075.
- Peterson KJ, et al. 2004. Estimating metazoan divergence times with a molecular clock. Proc Natl Acad Sci U S A. 101:6536–6541.
- Posada D, Crandall KA. 1998. MODELTEST: testing the model of DNA substitution. Bioinformatics 14:817–818.

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Poux C, et al. 2005. Asynchronous colonization of Madagascar by the four endemic clades of primates, tenrecs, carnivores, and rodents as inferred from nuclear genes. Syst Biol. 54:719–730.

- Rice P, Longden I, Bleasby A. 2000. EMBOSS: the European molecular biology open software suite. Trends Genet. 16:276–277.
- Ronquist F, Huelsenbeck JP. 2003. MrBayes 3: Bayesian phylogenetic inference under mixed models. Bioinformatics 19:1572–1574.
- Roughgarden J, editor. 1995. Anolis lizards of the Caribbean. Ecology, evolution and plate tectonics. New York: Oxford University Press
- Schaack S, Gilbert C, Feschotte C. 2010. Promiscuous DNA: horizontal transfer of transposable elements and why it matters for eukaryotic evolution. Trends Ecol Evol. 25:537–546.
- Simonsen T, et al. 2011. Phylogenetics and divergence times of Papilioninae (Lepidoptera) with special reference to the enigmatic genera *Teinopalpus* and *Meandrusa*. Cladistics 27:113–137.
- Suyama M, Torrents D, Bork P. 2006. PAL2NAL: robust conversion of protein sequence alignments into the corresponding codon alignments. Nucleic Acids Res. 34:W609–W612.
- Tamura K, Dudley J, Nei M, Kumar S. 2007. MEGA4: Molecular Evolutionary Genetics Analysis (MEGA) software version 4.0. Mol Biol Evol. 24:1596–1599.
- Thomas J, Schaack S, Pritham EJ. 2010. Pervasive horizontal transfer of rolling-circle transposons among animals. Genome Biol Evol. 2: 656–664
- Turnbull M, Webb B. 2002. Perspectives on polydnavirus origins and evolution. Adv Virus Res. 58:203–254.

- Waterston RH, et al. 2002. Initial sequencing and comparative analysis of the mouse genome. Nature 420:520–562.
- Wallau GL, Ortiz MF, Loreto EL. 2012. Horizontal transposon transfer in eukarya: detection, bias, and perspectives. Genome Biol Evol. 4: 689–699.
- Wicker T, et al. 2007. A unified classification system for eukaryotic transposable elements. Nat Rev Genet. 8:973–982.
- Wiegmann BM, et al. 2009. Single-copy nuclear genes resolve the phylogeny of the holometabolous insects. BMC Biol. 7:34.
- Wright F. 1990. The "effective number of codons" used in a gene. Gene 87:23–29.
- Xia X, Xie Z. 2001. DAMBE: software package for data analysis in molecular biology and evolution. J Hered. 92:371–373.
- Yoshiyama M, et al. 2001. Possible horizontal transfer of a transposable element from host to parasitoid. Mol Biol Evol. 18:1952–1958.
- Yuan YW, Wessler SR. 2011. The catalytic domain of all eukaryotic cut-and-paste transposase superfamilies. Proc Natl Acad Sci U S A. 108:7884–7889.
- Zhan S, Reppert SM. 2013. MonarchBase: the monarch butterfly genome database. Nucleic Acids Res. 41:D758–D763.
- Zhang HH, Xu HE, Shen YH, Han MJ, Zhang Z. 2013. The origin and evolution of six miniature inverted-repeat transposable elements in *Bombyx mori* and *Rhodnius prolixus*. Genome Biol Evol. 5: 2020–2031.

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