

Transposable elements in the *Anopheles funestus* transcriptome

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Abstract Transposable elements (TEs) are present in most of the eukaryotic genomes and their impact on genome evolution is increasingly recognized. Although there is extensive information on the TEs present in several eukaryotic genomes, less is known about the expression of these elements at the transcriptome level. Here we present a detailed analysis regarding the expression of TEs in *Anopheles funestus*, the second most important vector of human malaria in Africa. Several transcriptionally active TE families belonging both to Class I and II were identified and characterized. Interestingly, we have identified a full-length putative active element (including the presence of full length TIRs in the genomic sequence) belonging to the *hAT* superfamily, which presents active members in other insect genomes. This work contributes to a comprehensive understanding of the landscape of transposable elements in *A. funestus* transcriptome. Our results reveal that TEs are abundant and diverse in the mosquito and that most of the TE families found in the genome are represented in the mosquito transcriptome, a fact that could indicate activity of these elements. The vast diversity of TEs expressed in *A.*

funestus suggests that there is ongoing amplification of several families in this organism.

Keywords *Anopheles funestus* · Transposable elements · Transcriptome

Abbreviations

CDS	Coding sequence
Db	Database
LTR	Long terminal repeat
MITE	Miniature inverted repeat transposable elements
NLTR	Non-long terminal repeat
ORF	Open reading frame
PSI-Blast	Position-specific iterated BLAST
RB	Repbase
RPS-Blast	Reverse position-specific BLAST
TE	Transposable elements
TIR	Terminal inverted repeat
TSD	Terminal site duplications

Background

Transposable Elements (TEs) are abundant and ancient genetic sequences present in all eukaryotic genomes showing the ability of transposing between different loci. The distribution and abundance of TEs within and between different genomes varies widely, constituting the majority of the DNA content in some species, while in others they represent just a small fraction of the total genomic DNA (Bennetzen et al. 2005; de Koning et al. 2011; Kidwell 2002). In insects, for instance, genome sizes vary from less than 100 Megabases (Mb) to more than 10 Gigabases (Gb) (Gregory et al. 2007). The causes

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and consequences of these differences—sometimes in organisms with very similar degrees of complexity—are yet unknown. However, much of this variation reflects different evolutionary dynamics of TE families present in different genomic contexts and shows the enormous impact that these elements might have had in eukaryotic genomes' evolution.

Although far from a consensus for a universal TE classification system (Piégu et al. 2015), these elements have been classified based on their genetic structure and mode of replication into two main classes—Class I, or retrotransposons, and Class II, or DNA transposons (Finnegan 1989), and further hierarchically into orders, superfamilies, families and subfamilies by Wicker et al. (2007). Accordingly, Class I is composed of five orders: the LTRs, DIRS and Penelope-Like elements and the LINEs and SINEs, or Non-LTRs. These elements replicate through a reverse transcription step of an intermediary RNA and produce new copies in each cycle of replication while Class II transpose directly as a DNA molecule. The last are further divided into two subclasses, subclass 1 the classical 'cut-and-paste' elements (characterized by their terminal inverted repeats), and subclass 2 characterized by a transposition process without double-stranded cleavage (Orders Helitron and Polintons/Maverick). Both Class I and Class II are further classified into several superfamilies, distinguished by large-scale features, such as the structure of protein and noncoding domains, the presence and size of target site duplications (TSD). Superfamilies are further classified into families (also named clades or lineages) defined by DNA sequence conservation and further into subfamilies defined by phylogenetic relationships (Wicker et al. 2007) as well as by the degree of identity among their sequences.

An archetypal TE family can be composed of elements with different degree of activity: some elements having coding capacity, called autonomous elements, and others with inactivating mutations that can still harbor the ability of being mobilized by active and autonomous counterparts, known as non-autonomous elements. Most of the TEs in the present-day genomes are inactive, deteriorated or methylated, a fact that has been related to the evolution of controlling mechanism acting on the TE's mobilization. Probably, due to their mutagenic activity, eukaryote genomes have developed efficient mechanisms to silence them. Inactive elements populate genomes and evolve neutrally until eventually being lost from the genome. TEs can also be co-opted by their host genome; domestication of TE-derived protein coding and regulatory modules has indeed taken place repeatedly in the course of eukaryotic genome evolution (Miller et al. 1997; Casacuberta and Pardue 2005; Kapitonov and Koonin 2015).

Active elements have been reported in many genomes including mosquitoes—as *Herves* in *Anopheles*

(Arensburger et al. 2005), flies—*Drosophila P* element (Kidwell 1985), and L1 in humans (Sassaman et al. 1997).

Numerous studies have been conducted to identify and characterize TEs in insect at the genomic level, such as in *D. melanogaster* (Kaminker et al. 2002), *A. gambiae* (Holt et al. 2002; Fernández-Medina et al. 2011), *Culex quinquefasciatus* (Arensburger et al. 2010; Marsano et al. 2012), *Rhodnius prolixus* (Mesquita et al. 2015), and *Tribolium castaneum* (Wang et al. 2008), among others; however, less is known about these elements at the transcription level (de Araujo et al. 2005; Deloger et al. 2009; Mourier and Willerslev 2010; Iorizzo et al. 2011; Jiang et al. 2012; Sze et al. 2012). TEs are not only relevant due to the bulk of the genome occupied by them, but also due to the impact they have at the transcription level, by inserting into coding or regulatory regions, by influencing alternative mRNA processing or as sources of small regulatory RNAs (Kines and Belancio 2012; Mourier and Willerslev 2010; Cowley and Oakey 2013; de Araujo et al. 2005; Iorizzo et al. 2011; Kelley and Rinn 2012).

Here, we present an analysis of the transposable elements present in the *A. funestus* transcriptome, which together with *A. gambiae*, are the most important vectors for malaria transmission in sub-Saharan Africa. Mosquitoes of different species are among the most relevant invertebrate's vectors for veterinary and human vector borne diseases (VBD). Several *Anophelines* species are responsible for the transmission of Malaria, a disease that caused 212 million cases and 429,000 deaths, in 2015 (World Malaria Report 2015). Of the several measures used to control VBD, the mosquito populations are one of the main targets by the use of insecticides against which, many of them have acquired resistance (Chénais et al. 2012). TE-mediated mechanisms for developing resistance against insecticides in mosquitoes have been reported previously (Mouches et al. 1990, 1991; Darboux et al. 2007). Even if the relevance of TEs to insecticide resistance as a rule is not especially strong, they exemplify relations between the 'TEs landscape' and the appearance of adaptive traits, which is of big importance given the fact that vector control is one of the only control measures that show broad efficacy against diseases such as malaria.

In the malaria mosquito, several transcriptionally active TE families belonging both to Class I and II were identified and characterized. A full-length putative active element (with the presence of full-length TIRs in the genomic sequence) was also identified. This element belongs to the *hAT* superfamily, which also presents active members in other insect genomes (*Herves* in *A. gambiae*, *Hermes* in *Musca domestica* and *A. aegypti* and *hobo* in *Drosophila*) (Arensburger et al. 2005; Warren et al. 1994; O'Brochta et al. 1996; Sarkar et al. 1997; Sheen et al. 1993). The diversity of the TEs in the genomic dataset (Neafsey et al.

2015) has been shown to be higher than our estimation based on expressed sequences, as expected since not all the TEs in the genome will be expressed at the transcriptome level.

Materials and methods

Transcriptome assembling

We have assembled the raw data generated in a *de novo* transcriptome Illumina sequencing approach that used RNA from 30 *A. funestus* adult female individuals 3–5 days old (Crawford et al. 2010) derived from a newly founded colony in Burkin Faso and mRNA sequences obtained from different stages (pupae, larvae and adults from both sexes) of two strains (a field collected and a laboratory strain) and sequenced by the 454 technique (Gregory et al. 2011). The ABySS system—a short-read assembler that can process genome or transcriptome sequence data (Simpson et al. 2009; Birol et al. 2009; Robertson et al. 2010) was then used to assemble the short sequences obtained. As transcriptome samples typically contain transcripts with a wide range of expression levels, and assemblies generated with different k-mer lengths perform differently in capturing transcripts expressed at different levels, it is recommended to use several values for k-mer. We used k values ranging from 25 to 65 and generated a final non-redundant fastA following reassembly of the different fastA files from each k, using a parallelized blast/cap3 pipeline where blastn was used with decreasing word sizes (-w switch) from 300 to 60 to feed sequences to the cap3 assembler (Huang and Madan 1999; Karim et al. 2011). After assembling the sequences the final contig set consisted of 46,398 contigs.

Transposable elements identification and characterization

In order to characterize and identify putative transposable elements sequences, all the expression units generated (46,398 sequences) were used as queries for several BLAST programs performed on different databases (both public and in-house database versions) as follows: (1) NR-LIGHT by Blastx (Altschul et al. 1997): a subset of the non-redundant database (nr-db) from the NCBI, containing approximately 30% of the sequences and based on 106 genera and species, (2) SWISSP protein database by Blastx, (3) Gene Ontology database by Blastx (4) the CDD database by RPSblast (5) the eukaryotic cluster of orthologous groups (KOG) database by RPSblast, (6) the PFAM database by RPSblast, (7) the PRK database for functional annotation of the NCBI by RPSblast, (8) the SMART database by RPSblast, (9) an in-house database called “TE-DATABASE” generated

by Blastx using transposable elements against a subset of the nr-db having the following keywords related to TEs: “transposase”, “transcriptase”, “transposable element”, “retroposon”, “retrotransposon”, (10) an in-house database “TE-CLASS” generated by psi-blast using TE-class specific proteins as queries on the nr-db using the RPSblast, (11) an in-house database “TRANSPOSASE” generated by blastX on the nr-db using the keyword “transposase”, (12) an in-house database “GAG” generated by blastX on the nr-db using the keyword “gag”, (13) an in-house database “RRNA” containing rRNA, (14) an in-house database “MIT-PLA” containing mitochondrial and plastid DNA sequences and (15) “REPBASE” a TE reference database for eukaryotic elements: all the protein sequences from TEs deposited in REPBASE (both the entries with translated ORFs as well as the theoretical translations of those ORFs larger than 200 aa that were not presented as translated sequences in Replibase) were used as queries by a Psiblast on the nr-database in order to generate the TE models that were further used to run rps-Blasts against different sets of query sequences. We have previously used a similar approach in order to characterize the TE present in the genome of *A. gambiae* (Fernández-Medina et al. 2011).

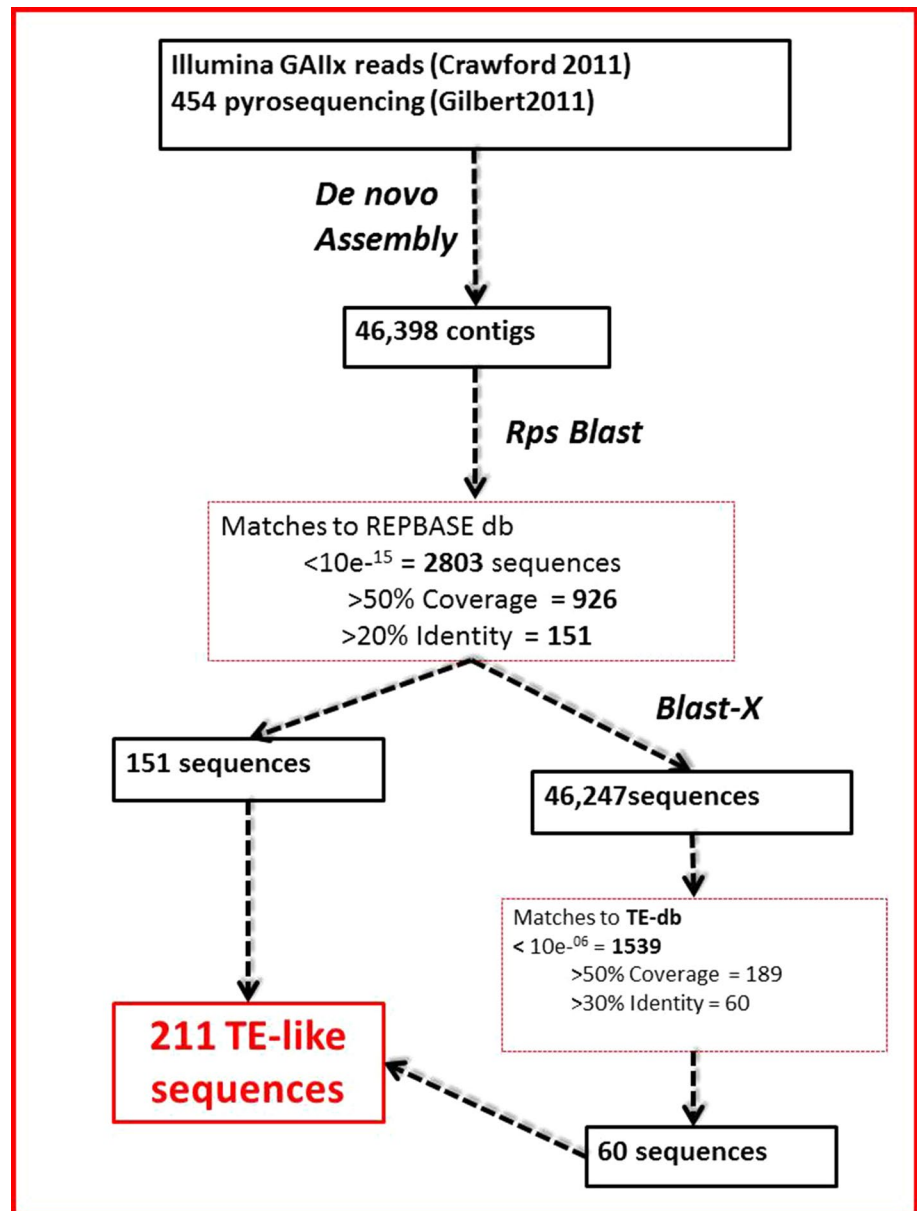
Our objective was to identify and characterize putative active elements being expressed in the *A. funestus* transcriptome. The criteria for selecting putative active transposable element sequences is presented in the workflow shown in Fig. 1. Briefly, 46,398 sequences were screened for identities against the sequences deposited in Replibase (RB) by RPSBlast. 2803 showed e-values $<10e^{-15}$, of which 926 have more than 50% coverage and 151 more than 20% identity with the Replibase elements. The remaining 45,247 sequences were screened by their identities to a TE-database by Blastx. 1539 were selected of which those covering more than 50% of the elements and presenting amino acid identities higher than 30% were included for further analysis. We totally selected 211 sequences presenting similarities to previously described TEs and further analyzed those sequences. All the remaining sequences were excluded of our analysis.

The sequences were further classified according to their TE class and superfamily, and divided in those presenting conserved domains (according to their “Best matches to the CDD” and “Best matched to pfam” databases) and those representing fragments or not showing the presence of functional domains.

Genome mapping

To evaluate the degree of identity of the “de novo” assembled sequences to the recently assembled *An. funestus* genome (Neafsey et al. 2015), we compared the assembled sequences to the assembled genome (version 1.0

Fig. 1 Pipeline used for the identification of TE-like sequences in the transcriptome of *A. funestus*



from Vectorbase) using the blat tool (Li and Durbin 2010), as well as by mapping the raw reads from the Kiribina (NCBI bioproject PRJNA177025) and Folonzo data sets (PRJNA177018) by Blastn (Altschul et al. 1997) (using a word size of 30, allowing for 1 gap, minimum 95% identity and up to 10 mapped reads to different targets if and only if the matches had the same score) to the deduced coding sequences of *An. funestus* added of 211 TE sequences that were “de novo” assembled. The resulting read mapping indicated an average/median coverage depth of 92.7/70.8 fold for the Folonzo data set and 68.5/47.6 fold for the Kiribina data set. To estimate the copy number for each TE or CDS, we divided the CDS or TE fold coverage obtained for each library by the median coverage of the respective library.

Phylogenetic analysis

Sequences belonging to different superfamilies were aligned with canonical sequences using MUSCLE (Edgar 2004). Phylogenetic relationships among the transposable element sequences and canonical sequences from the same superfamily/lineage were explored using neighbor-joining (NJ) and maximum likelihood (ML). The amino acid substitution models were evaluated using MEGA 5.0 (Tamura et al. 2011), the models with the lowest BIC scores (Bayesian Information Criterion) were considered the best to describe the substitution pattern (Tamura et al. 2011). NJ and ML trees were constructed using MEGA 5.0. Bootstrap values for each branch were assessed from 1000 replicates in both cases.

Results and discussion

Transposable elements identification

In order to study the expression of TEs in *A. funestus*, we assembled the 102.6 M Illumina reads from RNAseq generated by Crawford et al. (2010) together with the 375,619 454-pyrosequencing reads generated by Gregory et al. (2011) yielding a total of 46,398 contigs. We further used a pipeline that relies on different algorithms based on the Blast programs, against several databases as subjects (see methodology) in order to identify and characterize the TEs expressed in the mosquito genome (Fig. 1).

The results were compiled in a database (db) of expressed repetitive elements called Afun-TEExcel (Table S1). This database provides information about 211 sequences that were clearly identified as TEs in the mosquito's transcriptome. The information is organized as an Excel spreadsheet with cells containing, in a hyperlinked format, the results obtained after the various analyses performed in the characterization of each TE-like sequence.

We used stringent criteria for the inclusion of a sequence as a putative TE-like transcript (i.e. sequences with highly significant blast matches to Repbase or to TE-CLASS). Therefore, we have obtained a conservative set of expressed, putative active elements.

Our approach relies on the similarity of the transcript sequences to known proteins derived from TEs already characterized in other genomes. The use of RPSBlast, a position specific search engine based on profiles, against a PSIBlast-generated RB profile database as a first approach facilitated the classification and functional annotation of sequences. We identified 2803 sequences with significant matches to a set of TE-specific profiles by RPSBlast, based on all the elements deposited in RB, however, most of these sequences present very small coverage or low identities or, in some cases present high identities to nuclear proteins and were consequently discarded. We further restricted the search to those matches presenting at least 50% coverage (926 sequences) and more than 20% amino acid identity (151 sequences) to known TEs. The remaining sequences (46,247) were further classified according to their matches to the “TE-db” by BlastX. This approach allowed the inclusion of 1539 sequences that were again restricted to those presenting at least 50% coverage (189 sequences) and more than 30% amino acid identity (60 sequences). The resulting 211 sequences (151 plus 60) present e-values in the rps-Blast against RB, smaller than 1×10^{-15} confirming the identity of the sequences to already characterized TEs. However, many of these sequences (118) are not represented by full-length TE-transcripts (i.e. gag-pol, for class I, or transposase, for class II), but by truncated sequences of which 77.9% belong to the LTR order, 14.4% to the

NLTRs and the remaining 7.6% to the class II. These fragments have been clearly characterized as TEs, however, they have not been further analyzed.

Since most of the TEs in the genomes are represented by inactive or truncated copies, the transcripts identified here constitute an under representation of the total TE content in this genome. However, our study shows that TEs belonging to different families and subfamilies are present in this organism, many of which are transcriptionally active.

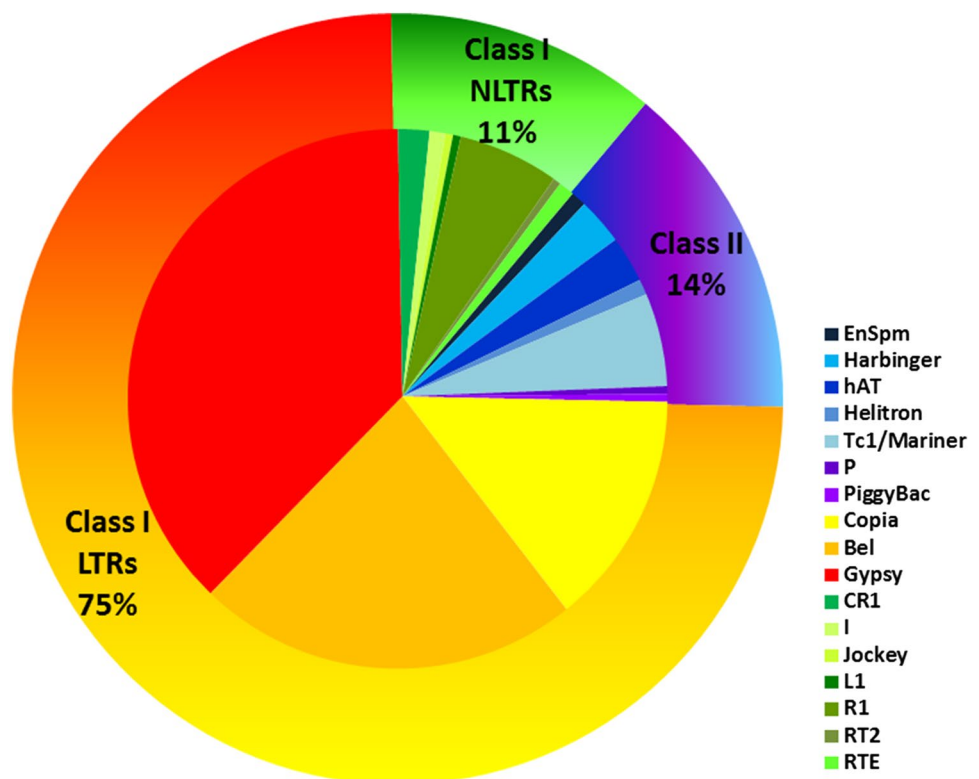
Transposable elements characterization

Transposable Elements representing all the classes/orders and most of the superfamilies previously characterized in insects were found in the *A. funestus* transcriptome, contributing to the mRNA and probably to protein diversity in this mosquito. Overall, the TE-related sequences identified in the transcriptome of *A. funestus* belong to Class I (86%), with a great overrepresentation of the LTR retrotransposons (Fig. 2). The high diversity of TEs being expressed in this organism is also present in other insect genomes, such as the mosquitoes: *A. gambiae*, *A. darlingi* and *Culex quinquefasciatus*, or the fly *D. melanogaster* (Holt et al. 2002; Marinotti et al. 2012; Arensburger et al. 2010; Adams et al. 2000). Although these insects harbor relatively small genomes, they contain many different TE families belonging to most of the TE orders/superfamilies so far identified. Not all the TE diversity found in a given genome would be expressed, in most of the cases very few elements are able to be expressed due to a high degree of deterioration of the elements in modern day genomes. In order to correlate our results with those of the reference genome, we have compared the number of TE families resulting from the analysis of the transcriptome (this study) with those identified in the genome (Neafsey et al. 2015) (Table 1). Most of the families we have identified in the transcriptome have also been identified in the genome. There are some discrepancies though, mainly with the number of families identified in each superfamily or lineage but not in the presence/absence of the superfamilies. The exceptions for this are the class II superfamilies, Transib and Academ, with two and four families identified in the genome that were not identified in the transcriptome and the class I, NLTRs, Ingi, L2 and Ouctast. These differences can be due to the differential expression of the families in the transcriptome.

In our data set, composed of 211 TE-like sequences, we have identified 30 sequences corresponding to the expression of full-length elements, 61 sequences spanning over full-length domains, and 120 sequences representing fragmented domains clearly belonging to TEs, as previously mentioned.

In order to confirm the presence of these TE families in the genome, and to validate the “de novo” assembly,

Fig. 2 Distribution of TE-like sequences in the *A. funestus* transcriptome. The outer chart represents the three main classes/orders of TEs (LTRs, Non-LTRs and Class II) and the inner chart shows the distribution of TE superfamilies within each class/order. The figures are based on the 211 sequences that were characterized in the transcriptome



we selected the sequences corresponding to full-length domains belonging to all the superfamilies identified in the transcriptome and compared them to the assembled *An. funestus* genome. Sixty-two of the 211 sequences had no matches to the assembled genome, while 106 and 133 TE's provided better than 95 and 90% identity to genomic sequences, respectively, indicating that the “de novo” TE assembly appears reasonable. The lack of genomic representation of almost 30% of the identified TE's may reflect the difficulties posed by these repetitive sequences on genome assemblies. To additionally validate the TE sequences, we mapped the genomic raw sequences both from the *Kiribina* and *Folonzo* strains of *A. funestus* (Neafsey et al. 2015) to the 211 TE sequences. The median genomic coverage of each dataset (mapped to 13,714 TE sequences+CDS) was of the same order of magnitude, 47.6 and 70.8 fold, respectively. All the TE-like sequences were identified in the genomic raw data with an average linear coverage of 99.7% in both datasets, including all the TE sequences that had no genomic matches. While we cannot exclude that our “de novo” assembly contain chimeric elements, the matches to genomic sequences as well as the raw reads indicate that the majority are representative of sequences corresponding to bona fide elements present in the genome, or the assembly of closely related elements. The element copy number in each library was estimated by considering the coverage of the TE-like sequences in each set (*Kiribina* and *Folonzo* strains) divided by the median

library coverage to all TE sequences+CDS in order to obtain an average for both libraries. Overall, at the genome level, the Class I contains superfamilies with the highest estimated copy numbers, especially belonging to the NLTR order. Among them, the R1 superfamily is the most abundant with 315 copies spread in the genome (Figure S1).

The diversity of the TEs in the genomic dataset has been shown to be higher than our estimation departing from expressed sequences (Neafsey et al. 2015) an expected fact, since not all the TEs in the genome will be expressed at the transcriptome level. Totally, 4719 L elements were reported in the genome of *A. funestus* of which the *Gypsy* constitute the least represented superfamily in copy number, 786 copies, against 2129 *Copia* and 1328 *Bel-Pao* copies (Neafsey et al. 2015). We also identified several Class II elements, including many families of *Tc-1/mariner*, *hAT*, *PiggyBac* and *Harbinger* superfamilies that were also previously identified at the genomic level (Neafsey et al. 2015).

Class I: LTRs

Four superfamilies within the LTR retrotransposons have been previously described: *Gypsy*, *Bel-Pao*, *Copia* and *DIRS* (for review see Wicker et al. 2007). Each of them has been subsequently classified into different clades/lineages (Copeland et al. 2005; de la Chaux and Wagner 2011; Llorens et al. 2009).

Table 1 Transposable Elements families identified in the transcriptome (this work) and genome (Neafsey et al. 2015) of *A. funestus*

	<i>A. funestus</i> transcriptome	<i>A. funestus</i> genome
Class II		
EnSpm	2	12
<i>Harbinger</i>	6	1
<i>hAT</i>	6	9
<i>Helintron</i>	2	3
<i>Tc1/Mariner</i>	12	10
<i>P</i>	1	3
<i>PiggyBac</i>	1	4
<i>Transib</i>	0	2
<i>Tsessebeii</i>	0	0
<i>Pegassus</i>	0	0
<i>Academ</i>	0	4
Class I		
LTRs		
<i>Copia</i>	30	26
<i>Bel</i>	49	103
<i>Gypsy</i>	78	77
NLTRs		
<i>CR1</i>	4	17
<i>I</i>	2	8
<i>Ingi</i>	0	1
<i>Jockey</i>	1	9
<i>L1</i>	1	7
<i>L2</i>	0	1
<i>Loner</i>	0	0
<i>Outcast</i>	0	7
<i>R1</i>	13	24
<i>RT2</i>	1	0
<i>R4</i>	0	0
<i>RTE</i>	2	6
TOTAL	211	334

In the *A. funestus* transcriptome sequences belonging to the three main superfamilies (*Gypsy*, *Bel-Pao* and *Copia*) were identified. Transcripts belonging to the *Gypsy* superfamily were the most abundant, even if they have been reported as the less abundant LTR superfamily at the genomic level (Neafsey et al. 2015). Of the 78 *Gypsy*-like transcripts, one corresponds to a full-length sequence expressing all the domains in both ORFs 1 and 2: retropepsin, reverse transcriptase (RVT-1), RNaseHI_RT_ty3, and integrase (RVE); 29 contain one or more full-length domains and the remaining transcripts correspond to sequences with truncated domains.

The *Bel-Pao* elements were the second most abundant transcripts identified within the LTR order (49 sequences). Six of them harbor full-length ORFs with all the domains

present in a complete element: DUF1758-1759/RVT-1/PeptA17/RVE and 27 contain full length domains.

Copia elements, were the least numerous LTR transcripts in the transcriptome, although they represent the most numerous superfamily at the genomic level, with an estimation of 2000 copies (Neafsey et al. 2015). Ten out of 30 *Copia* sequences contain full-length domains, and three have all the domains present in a full-length element (GAG- integrase/RVE/RVT_2/RNaseH_Ty1).

Class I: non-LTRs

It is well established that Non-LTR elements create in their replication cycle “Dead-on-arrival” sequences, producing fragments or truncated sequences missing their 5' ends. They have a replicative transposition mechanism that depends on the transcription of the whole element following a reverse transcription step. They are the least represented type of element in the *A. funestus* transcriptome; 24 different transcripts corresponding to seven different clades, representing 11% of all the TE-like transcripts were identified (Fig. 2). However, only the transcripts of the R1 clade present ORFs containing full-length domains and were further analyzed. Sequences representing truncated domains of the *CR1* (4), *I*, *Jockey*, *L1*, *RT2* and *RTE* clades (one sequence in each superfamily) could be identified. The truncated domains correspond to the Exo-Endo-Phosphatase (EEP); the RVT1 and the RH domains. These truncated sequences probably represent incomplete assembled sequences.

Class II: DNA transposons

Most of the previously identified DNA transposons belong to the class of cut-and-paste DNA transposons (Subclass 1), currently represented by 15 superfamilies (Kapitonov and Jurka 2008). The transposases encoded by cut-and-paste DNA transposons are also called DDE/DDD Transposases, due to the universal occurrence of three conserved acidic catalytic residues: two aspartates (D) and one glutamate (E), or three aspartates (DDD).

In the *A. funestus* transcriptome, the Class II elements correspond to 14% of the TE-like sequences, most of which belong to the TIR order. Two transcripts belonging to the *Helintron* order have also been identified. The identified transposase domains belong to different DDE superfamilies of endonucleases including DDE_1 and DDE_3 (from *Tc1-mariner* elements), DDE_4 (from *Harbinger*), the *hAT* family dimerization domain (from *hAT* elements), DDE_Tnp1_7 (from a *Piggybac* element), Tnp_P (from a *P*-element) and the DEAD-like and C-terminal domains of helicases from two *Academ* elements. The most

abundant expressed transposase belongs to the *Tc1/mariner* superfamily.

Phylogenetic analyses

Class I: LTRs

The use of the coding region corresponding to the reverse transcriptase to determine phylogenetic relationships has shown to be adequate for classification purposes (Xiong and Eickbush 1988). We performed a phylogenetic analysis of all the sequences presenting full-length ORFs corresponding to the RT domain, i.e. 10 *Gypsy*, nine *Bel-Pao* and three *Copia* sequences, together with previously published

reference sequences belonging to different insect species. The phylogeny confirmed the sequence classification based in our pipeline for all the LTR sequences at the superfamily level (Fig. 3). Additional phylogenetic analyses were performed for the sequences in each superfamily in order to classify them into lineages and further into families.

The *Gypsy* superfamily The *Gypsy* superfamily in insects has been classified into five different families also called lineages or clades (i.e. *Mag*, *Mdg1*, *Mdg3*, *Gypsy* and *CsRn1*). The 10 *Gypsy* sequences identified in the *A. funestus* transcriptome spanning the whole RT domain (177 aa. positions) were aligned together with reference sequences representing the five *Gypsy* lineages previously described

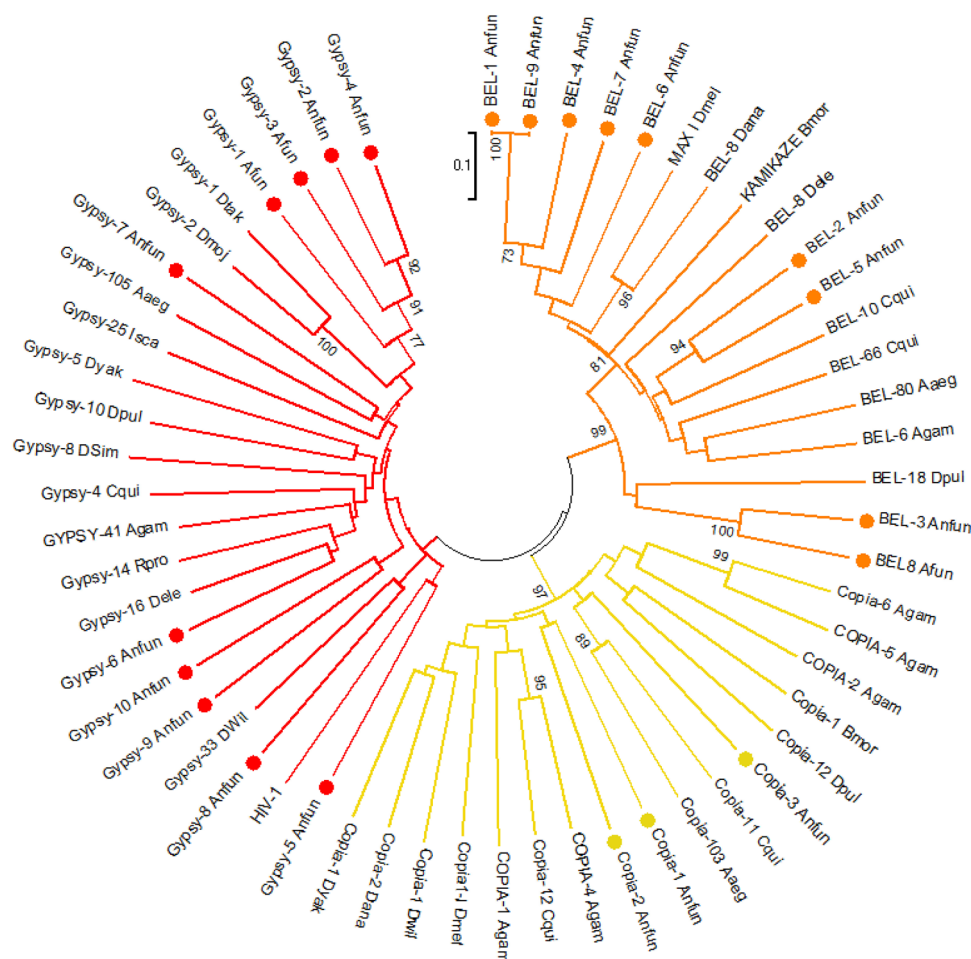


Fig. 3 Phylogenetic relationships of LTR sequences from *A. funestus*. The phylogenetic relationships of 22 L sequences from *A. funestus* plus 35 reference sequences from other insect genomes (accession numbers in Table S3) including sequences from the *gypsy*, *copia*, *Bel-Pao*, and HIV, spanning the RT domain. The phylogeny was inferred using the Neighbor-Joining method (Saitou and Nei 1987). The optimal tree with the sum of branch length=15.51 is shown. The evolutionary distances were computed using the p-distance method. The analysis involved 59 amino acid sequences. All ambiguous positions

were removed for each sequence pair. There were a total of 302 positions in the final dataset. The analyses were conducted in MEGA5 (Tamura et al. 2011). The numbers above the branches indicate the bootstrap value of a total of 1000 resamplings (only values higher than 70 are shown in the Figure). The different LTR superfamilies are coloured as follows: *Gypsy* in red, *Pao-Bel* in green and *Copia* in blue. The *A. funestus*' sequences are highlighted with colored dots corresponding to each superfamily

and three *Bel-Pao* elements as outliers (Fig. 4). A *Gypsy*-like partial element previously described in the genome of *A. funestus*, deposited in RB and called *Afun1* (Cook et al. 2000) was also included in the alignment. This sequence did not cluster with any of the expressed sequences from *A. funestus* and it is 100% identical to *Gypsy35_Agam*, from the *A. gambiae*'s genome. Nine sequences from *A. funestus* clustered together with elements belonging to four different lineages, i.e. *gypsy*, *mag*, *mdg3* and *CsRn1*; sequences clustering with the *mdg1* lineage, were not identified (Fig. 4).

Four sequences belonging to the *CsRn1* lineage (*Gypsy1-4_Afun*) belong to different families (mean p-distance of 40.08%, ranging from 26.3 and 47.4%). The sequences *Gypsy1_Afun*, *Gypsy2_Afun* and *Gypsy4_Afun* share the same family with several sequences from *A.*

gambiae. *Gypsy1* has 94.59% identity with *Gypsy49-AG*; *Gypsy2_Afun* has 85.10% identity with *Gypsy53-AG*, and *gypsy4* has 91.48 and 86.70% identity with *Gypsy52_AG* and *Gypsy2_AG*, respectively. *Gypsy3_Afun*, on the other hand is not clustering with any of the sequences used as references.

Gypsy5_Afun and *Gypsy8_Afun* clustered together with reference sequences from the *mdg3* lineage; however they belong to different families, presenting more than 63% of p-distance among them. *Gypsy8_Afun* clustered in the same family with sequences from *A. gambiae*, with a mean p-distance of 27.11%, while *Gypsy5_Afun* clustered with sequences from *A. aegypti* with a mean p-distance of 33.48% constituting the same family. *Gypsy7_Afun*, *Gypsy9_Afun* and *Gypsy10_Afun* clustered within the

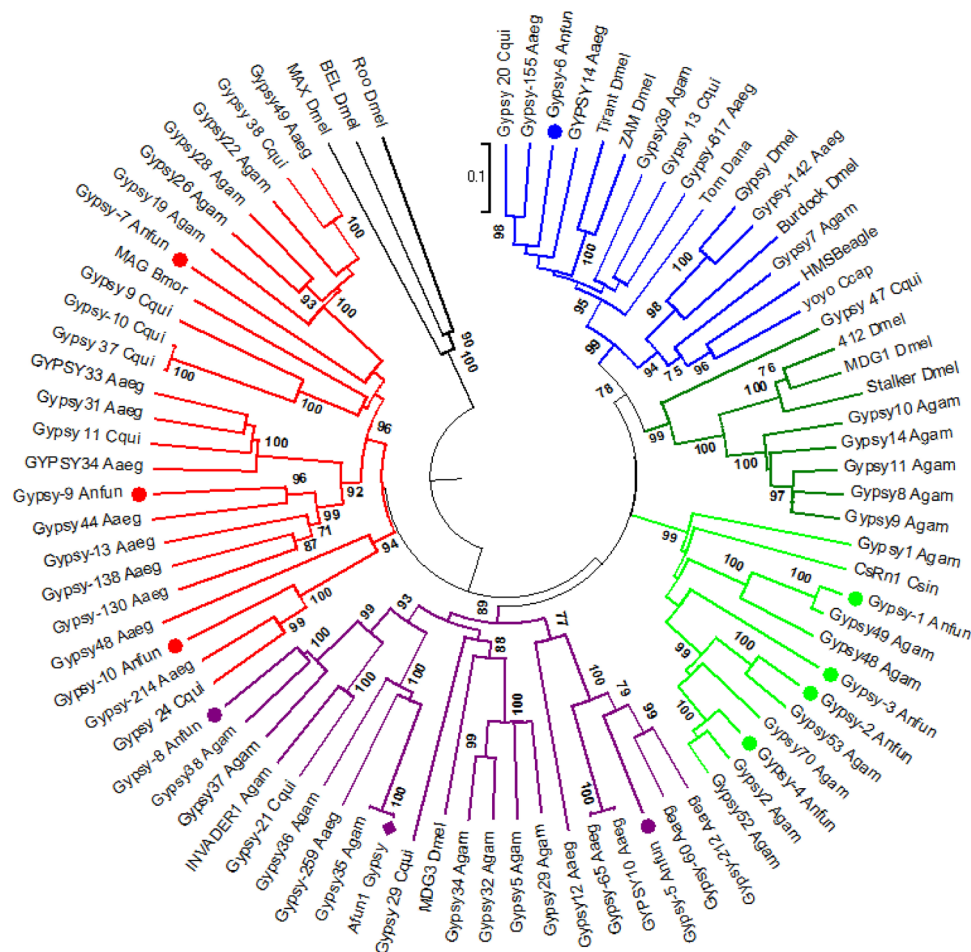


Fig. 4 Phylogenetic relationships of *Gypsy* sequences from *A. funestus*. The phylogenetic relationships of ten sequences from *A. funestus* and 56 reference sequences from other insect genomes (accession numbers in Table S3). The phylogeny was inferred using the Neighbor-Joining method (Saitou and Nei 1987). The optimal tree with the sum of branch length = 15.77 is shown. The evolutionary distances were computed using the p-distance method. The analysis involved 85 amino acid sequences. All ambiguous positions were removed for

each sequence pair. There were a total of 334 positions in the final dataset. The analyses were conducted in MEGA5 (Tamura et al. 2011). The numbers above the branches indicate the bootstrap value of a total of 1000 resamplings (only values higher than 70 are shown in the Figure). The different *Gypsy* lineages are coloured as follows: blue for *Gypsy*, olive-green for *Mdg1*, light-green for *CsRn1*, purple for *Mdg3*, and red for the *Mag* lineage. The *A. funestus*' sequences are highlighted with colored dots corresponding to each lineage

MAG lineage however; they are clearly related to different families within this lineage (mean p-distance 69.2% ranging from 66.8 to 62.7%).

The *Bel-Pao* superfamily The *Bel-Pao* superfamily has been previously classified into seven discrete lineages (*Pao*, *Sinbad*, *Bel*, *Tas*, *Suzu*, *Flow* and *Dan*) which tend to cluster with the host species phylogeny (Copeland et al. 2005; de la Chaux and Wagner 2011). These elements appear to have colonized only the genomes of the kingdom *Animalia*. We performed a phylogenetic analysis of the full-length RT region (213 aa) of nine *Bel-Pao* elements expressed in *A. funestus* and a C-terminal truncated sequence together with reference sequences from other insect genomes (Fig. 5).

The majority of the elements from *A. funestus* preliminary classified as *Bel-Pao* clustered together with the *bel* lineage while two sequences did it with the *Pao* reference sequences, and none of them clustered with sequences from the other lineages in this superfamily. The p-distance between the sequences *Bel-1_Afun*, *Bel-4_Afun*, *Bel-7_Afun* and *Bel-9_Afun* is 37.4%, (ranging from 0.0 to 47.2%) indicating that they belong to the same family. On the other hand, sequences *Bel-2_Afun* and *Bel-5_Afun* (p-distance 29%) belong to the same family than *Bel-6_Afun* (mean p-distance 44.2%).

The *Copia* superfamily We identified three sequences in the *A. funestus* transcriptome containing full-length RT

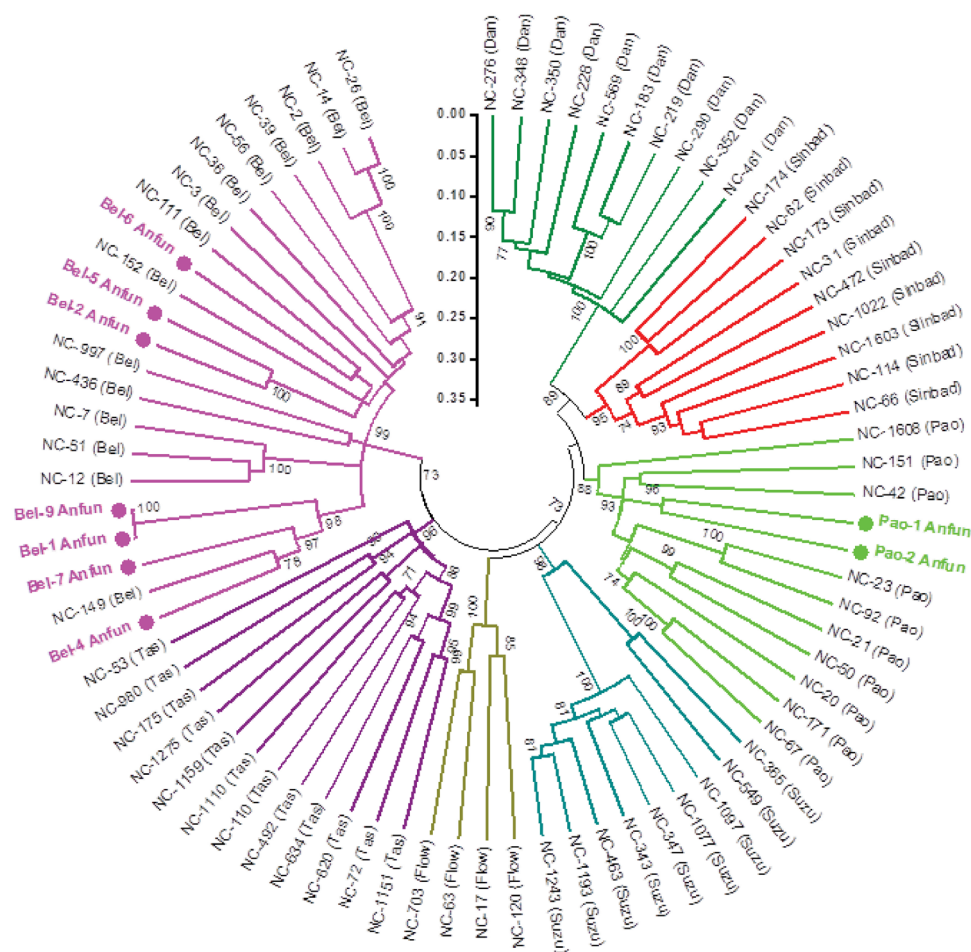


Fig. 5 Phylogenetic relationships of *BellPao* sequences from *A. funestus*. Phylogenetic relationships of nine sequences from *A. funestus* and 69 reference sequences from other insect genomes (accession numbers in Table S3). The phylogeny was inferred using the Neighbor-Joining method (Saitou and Nei 1987). The optimal tree with the sum of branch length=17.56 is shown. The evolutionary distances were computed using the p-distance method. The analysis involved 78 amino acid sequences. All ambiguous positions were removed for each sequence pair. There were a total of 241 positions

in the final dataset. The analyses were conducted in MEGA5 (Tamura et al. 2011). The numbers above the branches indicate the bootstrap value of a total of 1000 resamplings (only values higher than 70 are shown in the Figure). The different *Pao/Bel* lineages are coloured as follows: green for *Dan*, red for *Sinbad*, light-green for *Pao*, turquoise for *Suzu*, olive-green for *Flow*, purple for *Tas* and pink for *Bel*. The *A. funestus*' sequences are highlighted with colored dots corresponding to each lineage

sequences (246 aa) and belonging to the *Copia* superfamily. The *Copia* superfamily in insects has not been previously classified into different families or lineages/clades. We used 62 *copia* reference sequences from other insects (including, mosquitoes, flies and bugs) in order to classify the three *copia* sequences expressed in the *A. funestus* transcriptome. In our phylogeny the reference sequences grouped into, at least, five different lineages (I–V) one of them corresponds to *Drosophila*'s sequences only (indicated in blue in Fig. 6) while the others contain a mixture of sequences belonging to different mosquitoes' species. These sequences clustered within the same major group of sequences and together with sequences obtained from *A. aegypti* and *C. quinquefasciatus*, but none of the families previously characterized in *A. gambiae*. The p-distance for the three sequences from *A. funestus* is 51.0%, indicating that they belong to different families.

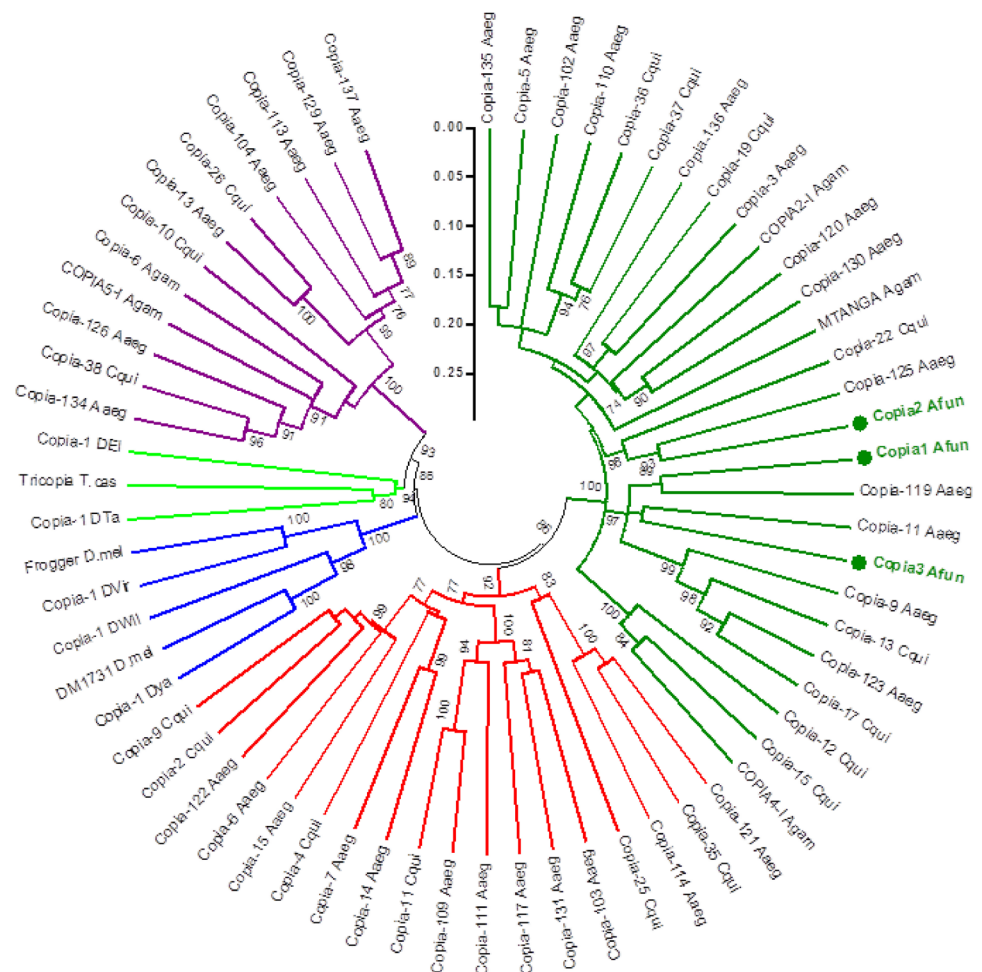
Class I: non-LTR

Several sequences belonging to different clades belonging to the NLTR order were identified in the *A. funestus*

transcriptome, although not all of them correspond to the RT region or represented full-length domains.

Eight sequences (seven belonging to the R, and one to the RTE clade) corresponding to the RT sequence were aligned to reference sequences representing all the clades described in the Non-LTR order. The phylogenetic analysis confirmed the classification obtained after our pipeline was applied (Fig. 7). A phylogenetic analysis including reference sequences belonging to the R1 superfamily in insects was performed (data not shown). The *A. funestus* R1 sequences clustered into four different clusters. The sequences *R1-1,2,3,5,8_Afun* clustered in a node together with sequences from *A. gambiae*. The p-distance among them is 32.5% (ranging from 16.2 to 33.5%), indicating that they belong to the same lineage together with sequences from *A. gambiae*. *R1-9_Afun* (mean p-distance against all the sequences from *A. funestus* 53.5%) and *R1-4_Afun* (mean p-distance against all the sequences from *A. funestus* 55.4%), on the other hand belong to more distant families.

Fig. 6 Phylogenetic relationships of *Copia* sequences from *A. funestus*. Phylogenetic relationships of three sequences from *A. funestus* and 62 reference sequences from other insect genomes (accession numbers in Table S3). The phylogeny was inferred using the Neighbor-Joining method (Saitou and Nei 1987). The optimal tree with the sum of branch length = 12.87 is shown. The evolutionary distances were computed using the p-distance method. The analysis involved 65 amino acid sequences. All ambiguous positions were removed for each sequence pair. There were a total of 268 positions in the final dataset. The analyses were conducted in MEGA5 (Tamura et al. 2011). The numbers above the branches indicate the bootstrap value of a total of 1000 resamplings (only values higher than 70 are shown in the Figure). The different *Copia* lineages are colored. The *A. funestus*' sequences are highlighted with green colored dots



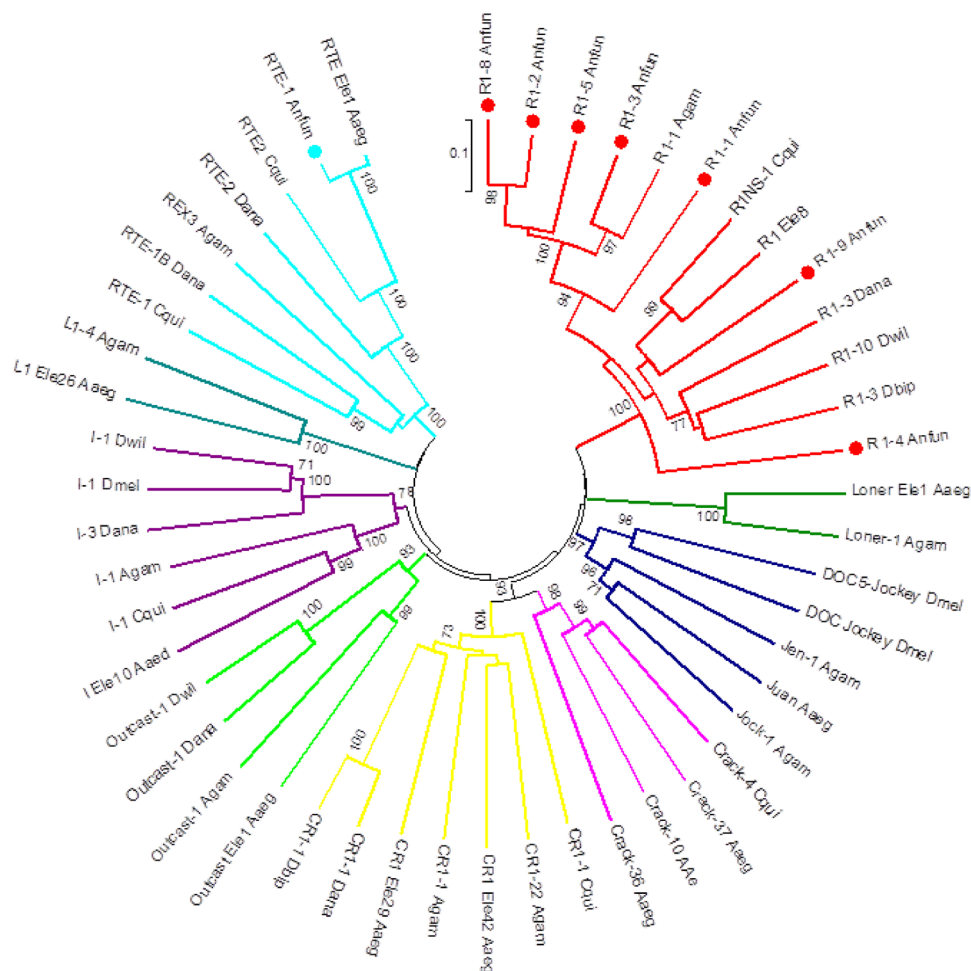


Fig. 7 Phylogenetic relationships of *NLTRs* sequences from *A. funestus*. Phylogenetic relationships of seven sequences from *A. funestus* and 42 reference sequences from other insect genomes (accession numbers in Table S3). The phylogeny was inferred using the Neighbor-Joining method (Saitou and Nei 1987). The optimal tree with the sum of branch length = 12.45 is shown. The evolutionary distances were computed using the p-distance method. The analysis involved 50 amino acid sequences. All ambiguous positions were removed for each sequence pair. There were a total of 316 positions in the final

dataset. The analyses were conducted in MEGA5 (Tamura et al. 2011). The numbers above the branches indicate the bootstrap value of a total of 1000 resamplings (only values higher than 70 are shown in the Figure). The different *NLTR* superfamilies are coloured as follows: red for *R1*, green for *Lones*, blue for *Jockey*, pink for *Crack*, yellow for *CR1*, light-green for *Outcast*, purple for *I*, turquoise for *L1* and light-blue for *RTE*. The *A. funestus*' sequences are highlighted with colored dots corresponding to each superfamily

Class II: DNA transposons

Transposases typically contain two domains: the N-terminal DNA-binding domain (an Helix-turn-Helix domain, known as HTH) (Pietrokovski and Henikoff 1997) and the C-terminal catalytic domain, characterized by the presence of a conservative D(Asp)DE(Glu)/D triad (Brillet et al. 2007) that has been shown to be essential for transposase activity (Lohe et al. 1995). Phylogenetic analyses of this domain in *mariner* elements permitted the classification of these elements into eleven subfamilies: *cecropia*, *irritans*, *mauritiana*, *mellifera*, *capitata* (Robertson 1993; Robertson and MacLeod 1993), *mori* (Robertson and Asplund 1996), *elegans* and *briggsae* (Robertson 2002), *rosa* (Gomulski

et al. 2001), *vertumnana* (Green and Frommer 2001), and *marmoratus* (Bui et al. 2007).

We have used the full-length transposase sequences of four *Tc1/mariner*, three *Pogo* and four *Harbinger* sequences from the *A. funestus* transcriptome in a phylogenetic analysis including reference sequences representing the different already characterized superfamilies (Fig. 8). Two of the sequences clustered together with reference sequences from the *mariner*'s family while three sequences clustered within the *Pogo* family, other two sequences clustered with the *Tc1* family, all with significant bootstrap values. Four sequences clustered together with the *Harbinger* family. The p-distances among the sequences belonging to the *Harbinger* (75.37%), *Pogo* (74.31%) and *Tc1* (71.59%)

Fig. 8 Phylogenetic relationships of DDE/D sequences from *A. funestus*. Phylogenetic relationships of eleven sequences from *A. funestus* and 48 reference sequences from other insect genomes (accession numbers in Table S3). The phylogeny was inferred using the Neighbor-Joining method (Saitou and Nei 1987). The optimal tree with the sum of branch length = 14.37 is shown. The analysis involved 59 amino acid sequences. All ambiguous positions were removed for each sequence pair. There were a total of 285 positions in the final dataset. The analyses were conducted in MEGA5 (Tamura et al. 2011). The numbers above the branches indicate the bootstrap value of a total of 1000 resamplings (only values higher than 70 are shown in the Figure). The different DDE/D superfamilies are coloured as follows: red for *Mariners*, green for *Tc1*, pink for *pogo*, and blue for *Harbinger*. The *A. funestus*' sequences are highlighted with colored dots corresponding to each lineage

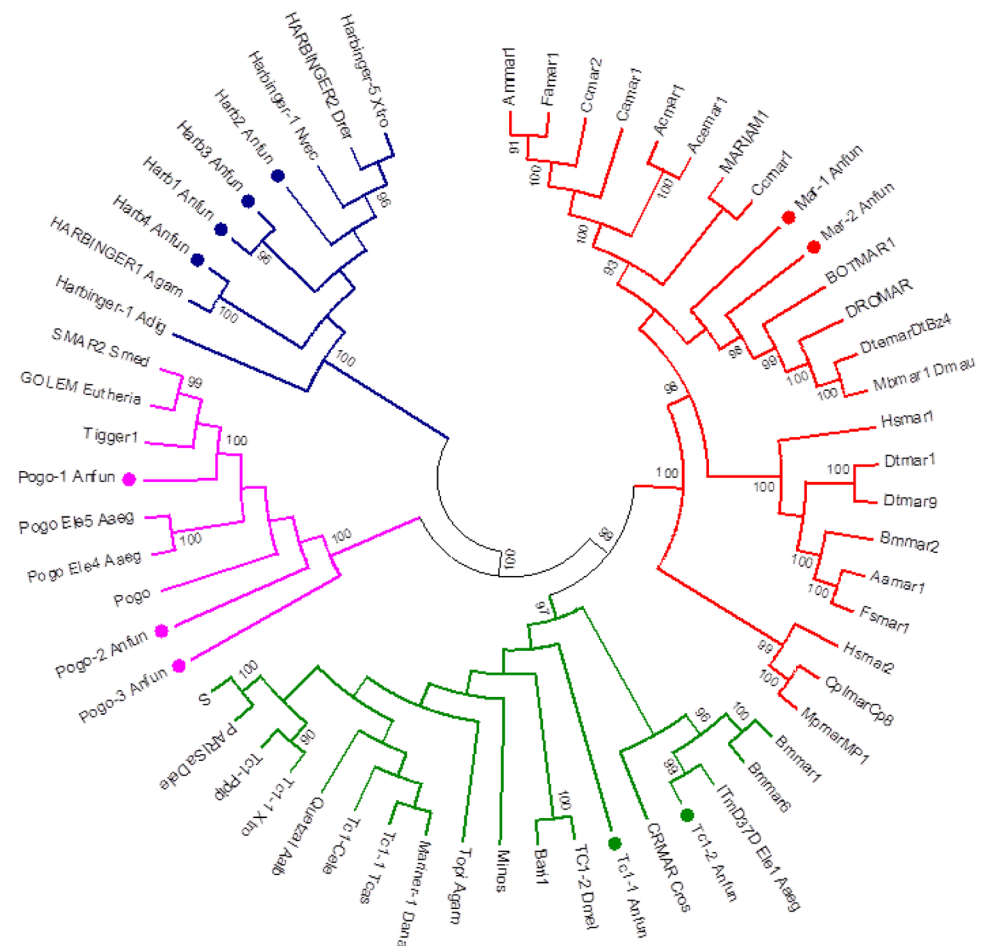


Table 2 Amino-acids present in the DDE/D motif of the *Tc1/mariner* and *pogo* elements

Mar1_Afun DD34D	D ₉₂ D ₄ H ₂₇ P ₁ D
Mar2_Afun GD34N	G ₉₂ D ₄ H ₂₇ P ₁ N
Tc1-1_Afun DD34E	D ₈₅ D ₄ H ₂₇ P ₁ E
Tc1-2_Afun DD37D	D ₉₁ D ₄ H ₃₀ P ₁ D
Pogo1_Afun DD32D	D ₇₁ D ₄ H ₂₅ P ₁ D
Pogo2_Afun AN45D	A ₁₀₄ N ₄ H ₂₄ P ₁₅ D
Pogo3_Afun DD45D	D ₁₀₉ D ₄ H ₂₇ P ₁₅ D

families indicated that they belong to different subfamilies within each family. While the two *mariner* sequences show a lower distance (56.17%) indicating that the sequences do not belong to the same subfamily.

The *Tc1/mariner* elements identified in the *A. funestus* transcriptome contain some divergences from the canonical DDD/E domains. Transposase sequences containing the GD34N and the AN45D were identified. We cannot assure that these sequences result in active transposases (Table 2).

In order to further characterize these two sequences within the *mariner* family, a phylogeny including these two sequences and members representing each of the

characterized *mariner* subfamilies was performed. The phylogeny indicated that the *mariner* sequences belong both to the *mauritanica* subfamily (data not shown) (Robertson and McLeod 1993; Wallau et al. 2014).

The hAT superfamily The *hAT* elements (by *hobo*, *Ac* and *Tam3*) are present in a wide range of plants and animals, including insects (Kempken and Windhofer 2001; Weil and Kunze 2000). Interestingly, these elements have been found in active forms in insects and also, to be active when introduced into divergent insect species, making them suitable as gene drivers. *Hermes* from the housefly, *Musca domestica* (Atkinson et al. 1993), *hobo* from *D. melanogaster* (McGinnis et al. 1983), *Herves* from *A. gambiae* (Arensburger et al. 2005) also, shown to be present in the genomes of other *Anophelines* (*A. arabiensis* and *A. merus*) and in *A. aegypti* (Arensburger et al. 2011) have all been shown to be mobilized in different species.

Sequences belonging to three different *hAT* elements were identified in *A. funestus*. We used the full-length *hAT1_Afun* sequence as a query in a BlastN search in the *A. funestus* sequenced genome. Five *hAT* sequences were identified in the genome. Two pairs of intact TIRs flanking

the transposase gene were also identified. A phylogeny of the C-terminal dimerization region of the sequences identified both in the transcriptome and the genome together with reference sequences from *hAt* elements from other insect genomes showed a cluster of three pairs of transcriptome and genome sequences (*hAT1-3_Afun*) (Fig. 9). Two other genomic sequences, named *hAT4,5_Afun* presented no expressed counterpart. The p-distance among the genomic sequences corresponding to the three expressed elements is 39.18%, indicating that the three of them belong to the same family. On the other hand, the p-distances between the sequences in the cluster and *hAT4_Afun* and *hAT5_Afun* are 47.89 and 56.96%, respectively.

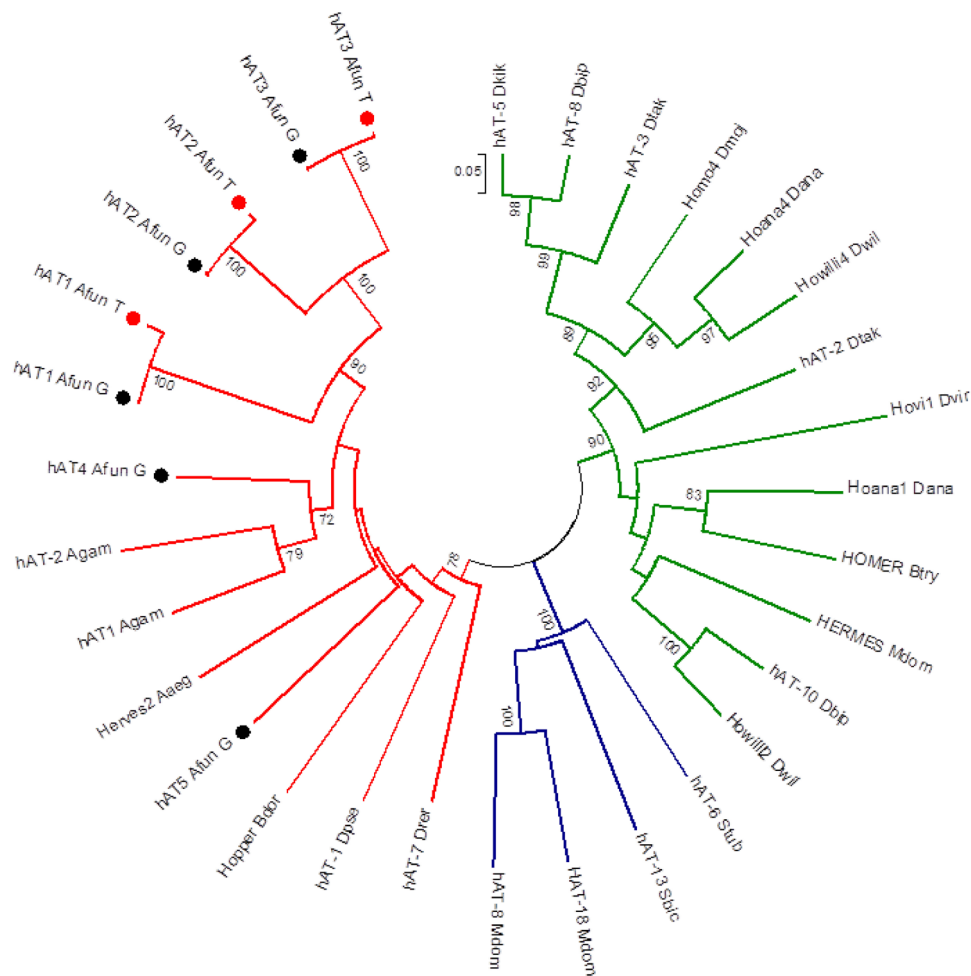


Fig. 9 Phylogenetic relationships of *hAT* sequences from *A. funestus*. Phylogenetic relationships of three sequences from the *A. funestus* transcriptome and five sequences from the genome together with 23 sequences from other insect genomes (accession numbers in Table S3). The phylogeny was inferred using the Neighbor-Joining method (Saitou and Nei 1987). The optimal tree with the sum of branch length = 6.28 is shown. The tree is drawn to scale, with branch lengths in the same units as those of the evolutionary distances used to infer the phylogenetic tree. The evolutionary distances were computed using the p-distance method and are in the units of the number

The finding of an expressed element, presenting full-length transposase gene and intact TIRs at the genomic level is suggestive of recent or ongoing active transposition of this family. The ability to transpose in diverged species from their hosts appears to be common feature of members of the *hAT* superfamily. Consequently, the *hAT* elements have great potential to serve as non-drosophilid insect gene vectors.

of amino acid differences per site. The analysis involved 31 amino acid sequences. All ambiguous positions were removed for each sequence pair. There were a total of 107 positions in the final dataset. The analyses were conducted in MEGA5 (Tamura et al. 2011). The numbers above the branches indicate the bootstrap value of a total of 1000 resamplings (only values higher than 70 are shown in the Figure). *DifferenthAT* lineages are coloured in red, blue and green. The *A. funestus* sequences are highlighted with colored dots corresponding to each lineage, in red the sequences isolated from the transcriptome (T) and in black the sequences obtained from the genome (G)

Concluding remarks

In this article we have presented data related to the identification and characterization of transposable elements in *A. funestus*, the second most important vector of malaria in Africa. We have used publicly available data of two whole transcriptome sequencing projects (Crawford et al. 2010; Gregory et al. 2011) to elucidate the extent and character of the repetitive elements being expressed in this mosquito. TEs belonging to all the classes and to most of the TE superfamilies already characterized are present and expressed in this organism. The methodology implemented resulted in the identification of TE superfamilies already identified and characterized in other sequenced genomes avoiding the identification of putative novel elements. The determination of the activity or potential activity of these elements will require further functional verification.

The identification of a vast diversity of TEs expressed in *A. funestus* suggests that there is ongoing amplification of several families in this organism. On the other hand, the lack of genomic representation of many TE's reflects the difficulties related to the correct assembling of transposable elements sequences in genome assemblies.

The data presented here is based primarily on the similarity of the transcript sequences to functional domains for known transposable elements in other species. This might be insufficient support for actual activity of these elements. It is known that the potential for TE activity on a structural level may be restricted by cell type, especially the soma and germ-line, or epigenetic microRNA regulation. And, it is well established that the transcriptional activity of transposable elements is tightly controlled, although some retrotransposons are transcribed under stress conditions such as pathogen infection, physical injuries or abiotic stresses (Grandbastien 1998).

TE-mediated mechanisms for developing resistance against insecticides in mosquitoes have been reported previously. An amplification of at least 250-fold of the esterase gene related to the overproduction and increased activity of esterase B1, involved in the resistant phenotype of *Culex* species to organophosphorus (OP) insecticides, has been identified in *Culex pipiens quinquefasciatus*. An amplicon (30 kb) in the resistant mosquitoes contained the esterase gene (2.8Kb) framed by DNA sequences homologous to middle or highly repetitive elements present in the genome of susceptible and OP-resistant mosquitoes, which were thought to be of TE origin (Mouches et al. 1990, 1991). Microbial larvicides have also commonly been used for controlling mosquitoes-borne diseases. A binary toxin from *Bacillus sphaericus* has a larvicidal property following ingestion by susceptible larvae. However, high levels of resistance has also been reported in field populations of *Culex* species

isolated from different countries where the larvicide has been used extensively (Rao et al. 1995; Yuan et al. 2000; Chevillon et al. 2001). One of the mechanisms of the larvae resistance is related to the insertion of a TE-like element in the coding region of the gene that codifies for the receptor involved in the interaction with the toxin (Darboux et al. 2007). The TE insertion modified a splicing site, creating an intron and leading to the production of a shorter receptor, unable to interact with the toxin and leading to the insect survival. Even if the relevance of TEs to insecticide resistance as a rule is not especially strong, they exemplify relations between the 'TE landscape' and the appearance of adaptive traits, which is of big importance given the fact that vector control is one of the only control measures that show broad efficacy against diseases such as malaria.

We used stringent criteria to designate putative functional transcripts, however, we cannot exclude the possibility that some of these sequences might represent read-throughs transcripts. Still, given the importance of the transposable elements and their role in many spontaneous mutations influencing evolution, and the adaptive traits in insects (for instance insecticide resistance) it is important to characterize the presence of these elements at the transcriptome level.

It is therefore not only important to understand the landscape of elements present in any given genome but also the expression of those sequences in transcriptomes. As the examples showed above indicate, TEs might play an important role in the appearance of larvicide- and insecticide-resistant phenotypes, emphasizing the significance of studies aiming at the identification and characterization of TEs in genomes and transcriptomes.

On the other hand, the use of transposable elements as tools for the introduction of desirable genes into target populations has also been pursued as a means for controlling VBD, particularly, the transformation of *A. gambiae* as a means to control the spread of malaria. Active TEs can be used in genetic engineering as transformation vectors and can be used for gene and enhancer trapping; they also can be used for genome-wide insertional mutagenesis studies (Tu and Li 2009). In this respect, we have identified a full-length putative active element (including the presence of full length TIRs in the genomic sequence) belonging to the hAT superfamily, which presents active members in other insect genomes (Herves in *Anopheles gambiae*, Hermes in *Musca domestica* and *Aedes aegypti*, and hobo in *Drosophila*) and that have been also used as driver elements. Moreover, a great diversity of active elements is present in *A. funestus*. Further functionality tests by mobility assays could be of great importance in order to determine the use of these elements as genetic tools in other species. This work contributes overall to a comprehensive understanding

of the landscape of transposable elements in this important vector for malaria.

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Compliance with ethical standards

Conflict of interest The authors declare no competing financial interests.

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