Metazoan ribotoxin genes acquired by Horizontal Gene Transfer

Walter J. Lapadula¹*, Paula L. Marcet², María L. Mascotti¹, María V. Sánchez Puerta³, Maximiliano Juri Ayub¹*

1. Instituto Multidisciplinario de Investigaciones Biológicas de San Luis, IMIBIO-SL-CONICET and Facultad de Química, Bioquímica y Farmacia, Universidad Nacional de San Luis, San Luis Argentina.
2. Centers for Disease Control and Prevention, Division of Parasitic Diseases and Malaria, Atlanta, USA.
3. Instituto de Ciencias Básicas, IBAM-CONICET and Facultad de Ciencias Agrarias, Universidad Nacional de Cuyo, Mendoza, Argentina.

*Corresponding authors: mjuriayub@hotmail.com, wlapadula@gmail.com
Abstract

Ribosome inactivating proteins (RIPs) are RNA N-glycosidases that depurinate a specific adenine residue in the conserved sarcin/ricin loop of 28S rRNA. These enzymes are widely distributed among plants and their presence has also been confirmed in several bacterial species. Recently, we reported for the first time in silico evidence of RIP encoding genes in metazoans, in two closely related species of insects: *Aedes aegypti* and *Culex quinquefasciatus*. Here, we have experimentally confirmed the presence of these genes in mosquitoes and attempted to unveil their evolutionary history. A detailed study was conducted, including evaluation of taxonomic distribution, phylogenetic inferences and microsynteny analyses, indicating that the culicine RIP genes derived from a single Horizontal Gene Transfer (HGT) event, probably from a Cyanobacterial donor species. Moreover, evolutionary analyses show that, after transference, these genes evolved under purifying selection, strongly suggesting that they play functional roles in these organisms. In this work we confirm the presence of RIP genes in Culicinae species, and show solid evidence supporting the hypothesis that these genes are derived from a single prokaryotic transferred gene through HGT. In addition, clear evidence of purifying selection pressure has been recorded, supporting the hypothesis that these genes are functional within this subfamily.
Introduction

Ribosome inactivating proteins (RIPs, EC 3.2.2.22) irreversibly modify ribosomes through the depurination of an adenine residue in the conserved alpha-sarcin/ricin loop of 28S rRNA (1-4). This modification prevents the binding of elongation factor 2 to the ribosome, arresting protein synthesis (5, 6). The occurrence of RIP genes has been experimentally confirmed in a wide range of plant taxa, as well as in several species of gram positive and negative bacteria (7-9). Additionally, the exponential increase of information in databases has suggested the presence of these toxin genes in Fungi, Cyanobacteria and Metazoan lineages (10-12). Although several RIPs have been extensively studied at the biochemical level, their biological roles remain open to speculation. In some cases, it seems reasonable to predict their functions. For instance, the high toxicity of ricin supports an antifeedant role, whereas shiga and shiga-like toxins are strong virulence factors for their harboring bacteria. Antiviral and other defense activities have been postulated for other plant RIPs, but no concluding evidence has been obtained. Recently, the RIP of the symbiotic *Spiroplama in Drosophila neotestacea* was shown playing a defensive role in preventing a virulent nematode from infecting this insect (13).

In a previous work, we have described that phylogeny of RIP genes show incongruence with those of species. Most of these inconsistencies can be explained by gene duplication, loss and/or lineage sorting (11). Another mechanism leading to phylogeny incongruence is horizontal gene transfer (HGT); namely the non-genealogical transmission of DNA among organisms. HGT is accepted as an important force driving prokaryotic genome evolution (14, 15). In contrast, its impact on genomes from multicellular eukaryotes, in particular animals, is largely controversial (16, 17). To be maintained permanently in animal species, heritable changes (*i.e.* the transferred gene) must be incorporated into germline cells and transmitted to the offspring. Nevertheless, in the particular case of herbivore arthropods, HGT has been postulated to play a role in the adaptation to phytophagy, including the efficient assimilation and detoxification of plant produced metabolites (18).
Detection of *bona fide* HGT derived genes is not trivial, and careful data revision is required for its corroboration. Many cases of putative foreign genes have been shown, after further revision, to result from artifacts or misinterpretations, such as contamination of genomic data, incomplete sampling of sequences and/or taxa, incorrect phylogenetic inferences and hidden paralogy. Two emblematic cases illustrating these issues are the initial conclusion that the human genome contained a high percent of bacterial derived genes (19), and the recent claim that tardigrade genomes contain significant amounts of foreign DNA (20). In both cases, subsequent sounder analyses demonstrated that contamination or incomplete sampling better explained the available data (21, 22). Consequently, careful case-by-case analyses of HGT candidates are required for their efficient detection. To do so, independent evidences and alternative evolutionary scenarios should be taken into account.

Based on the previous finding of *in silico* evidence of the presence of RIP genes in two closely related species of mosquitoes, we aim to confirm the presence and determine the location of RIP genes in species of the Culicinae subfamily. Moreover, we show solid evidence supporting the hypothesis that these genes derive from a single prokaryotic transferred gene.

**Materials and Methods**

**PCR experiments**

PCR experiments were conducted to confirm the presence of RIP encoding sequences in selected organisms. Individuals of *C. quinquefasciatus* strain JHB were obtained from the MR4 colony (Malaria Research and Reference Reagent Resource Center) of the Center for Diseases Control and Prevention (CDC). Genomic DNA from these specimens was extracted employing the protocol previously described by Collins (23). Genomic DNA from wild specimens of *C. molestus, C. pipiens* and *C. torrentium* was kindly provided by Dr. Stefanie Becker (Bernhard Nocht Institute for Tropical Medicine, Hamburg) (24). Primer sets were designed to amplify the full-length RIP encoding sequence of *C. quinquefasciatus* and also to identify the predicted neighbor gene (S1 Table). High fidelity Phusion DNA polymerase (New England Biolabs) was
used. PCR conditions were: initial denaturation 30 s at 98°C, followed by 35 cycles of
denaturation (10s at 98°C), annealing (30s at Ta°C see S1 Table) and extension (30s at 72°C),
final extension of 10 min at 72°C. PCR products were cloned into the pGEMT-easy vector
(Promega) following standard methods, and sequenced. The obtained sequences are available
with the following GenBank codes: KX674699, KX674697, KX644696 and KX674698.

Multiple sequence Alignments (MSAs) and phylogenetic inferences

We used the amino acids sequences of previously reported RIPs (11) as probes for
iterative BLAST searches. All collected sequences were employed to construct MSAs using
MAFFT 7 on-line server (http://mafft.cbrc.jp/alignment/server/). MSAs were manually edited to
remove gaps. Phylogenetic trees were obtained employing Bayesian inference using Mr. Bayes
3.2 software (25). A mixed amino acid substitution model was set up, 4 gamma categories and a
proportion of invariable sites were considered. The analyses were concluded after 1,000,000
generations when the split frequency was < 0.02. FigTree 1.4.2 software was used to visualize
and edit the obtained trees.

Genomic context analyses

The whole contigs containing RIPs of *C. quinquefasciatus* (DS232037), *Aedes aegypti*
(NW001810221) and *Anopheles gambiae* (chromosome 3L) were subjected to BLASTx
searches on different protein databases to identify individual genes upstream and downstream
from the RIP open reading frame (ORF). The retrieved sequences were subjected to reciprocal
BLASTp and tBLASTn searches in order to determine putative orthologous sequences. The
obtained orthologs were confirmed using the comparative genomic tool available at VectorBase
website (https://www.vectorbase.org/).

Evolutionary analyses
Nucleotide encoding RIP sequences found in mosquitoes were aligned by codons, using PAL2NAL algorithm (26). Then, the synonymous and non-synonymous substitution rates were estimated employing SLAC, FEL and REL tests available in the Datamonkey Package (27, 28). Finally sequence logo diagram was constructed employing the codon alignment using the Weblogo software (29) and codons under purifying selection were highlighted.
Results

_Culex_ spp genomes harbor RIP encoding genes

Recently, we have found _in silico_ evidence for the presence of RIP genes in two closely related species of metazoans: _Aedes aegypti_ and _Culex quinquefasciatus_ (11). These intriguing findings led us to design experimental strategies to confirm their presence by ruling out possible database artifacts (_i.e._ contamination). For this purpose, genomic DNA was obtained from a pool of four mosquitoes of _C. quinquefasciatus_ strain JHB. Then, two independent PCR experiments were designed to demonstrate the presence of the RIP gene, and to confirm its physical linkage to the predicted neighbor gene, which is an intron-containing metazoan-derived gene (XM_001850822). Fig 1 shows that both PCR products presented the expected size. Also, further cleavage of both amplicons with EcoRI yielded the predicted patterns, confirming their identity.

![Diagram](attachment:image.jpg)

**Fig 1:** Experimental confirmation of the presence and location of RIP gene in _C. quinquefasciatus_ JHB genome. (A) Schematic representation of a fragment of the contig AAWU01015132 depicting the RIP gene (RIPcu) and its closest neighbor gene (XM_001850822). Expected amplicons and relevant EcoRI restriction sites are also depicted. (B) RIP ORF was amplified by PCR and the product was
analyzed by gel electrophoresis before (lane 1) and after EcoRI treatment (lane 2). (C) A genomic fragment of 1,882 bp linking the RIP gene with its neighbor gene was amplified and electrophoresed before (lane 1) and after EcoRI treatment (lane 2).

Once the presence and location of the RIP sequence in *C. quinquefasciatus* genome were experimentally confirmed, we successfully amplified the full length RIP coding sequences (~1300 bp) of the closely related species *C. pipiens, C. molestus,* and *C. torrentium* (S1 Fig). PCR products were cloned and sequenced, and the obtained sequences were aligned. As expected, nucleotide sequences showed high similarity (93-97% of identity at nucleotides and amino acids levels) in relation to the reported sequence in the *C. quinquefasciatus* genome database (S2 Fig). The sequence obtained from the *C. quinquefasciatus* JHB MR4-CDC repository revealed an in-frame, three-nucleotide (ACC) insertion (encoding an additional Thr residue). Interestingly, this sequence also harbors a ten-nucleotide frame-shifting deletion (nt 542-551) generating a premature stop codon (S2 Fig). By direct sequencing of PCR products from six individual specimens of *C. quinquefasciatus* JHB from the MR4-CDC colony, we confirmed that all these individuals were homozygous for the deletion, strongly suggesting that this null mutation was fixed in this colony (S3 Fig).

**Culicine RIP genes are monophyletic and syntenic**

We have previously shown that RIP genes from particular lineages (e.g. monocots or dicots, bacteria and fungi) are not monophyletic. Moreover, many RIP clades include sequences belonging to largely distant taxa (11, 30). Based on this evidence, we postulated that the evolutionary history of RIP genes is consistent with the existence of several ancient paralogues, followed by multiple lineage-specific gene duplications and losses (11). However, metazoan RIP genes are particularly interesting because they are restricted to closely related insects of the subfamily Culicinae. As can be seen in Fig 2, RIPS from *Culex quinquefasciatus* and *Aedes aegypti* form a well supported clade [posterior probability (PP): 1]. Monophyly of culicine RIPS, along with their apparent narrow taxonomic distribution, suggest that these genes are derived from a rather recent, single ancestral sequence. In order to test this, microsynteny analyses were
carried out using scaffolds from *Aedes aegypti, Culex quinquefasciatus* and *Anopheles gambiae* (the closest relative species lacking RIP genes). As expected, partially conserved synteny blocks were identified in the three species (Fig 3 and S2 Table).
Fig 2: Phclogenetic tree of RIPs protein family. The tree was constructed employing Bayesian Inference. Numbers above the branches indicate the posterior probabilities (PP) values. Names of species
are indicated for each sequence in the tree. Lineages are indicated by different color codes as follows: green (Plant), blue (Fungi), red (Metazoan), orange (Bacteria) and magenta (Cyanobacteria). The clade of dipterous RIPs is emphasized with red branches.

Fig 3: Schematic representation of the genomic context between *Culex quinquefasciatus* (DS232037), *Aedes aegypti* (NW001810221) and *Anopheles gambiae* (chromosome 3L). Greys shadows link conserved syntenic ORFs. RIP genes are represented with orange arrows. Additional information about each gene is available in Table S2.

**Culicine RIP genes are derived from a unique HGT event**

Recent sequence homology searches further confirmed that metazoan RIP genes are restricted to the Culicinae subfamily. In addition to the above described genes, we found *in silico* seven RIP genes in *Aedes albopictus* (GenBank: KXJ78156, KXJ78155, KXJ78158, KXJ73132, KXJ78157, KXJ73133 and KXJ73764), which are clearly syntenic with *Aedes aegypti* RIPs (Data File S1) and two transcriptomic sequences from another Culicine mosquito; *Armigeres subalbatus*, partially covering the ORFs (GenBank: EU212208, EU211398). In light of these findings, two alternative hypotheses were postulated:

i) These genes have been vertically inherited from the metazoan cenancestor, and were purged from other metazoan genomes by a number of independent gene loss events.

ii) These genes are derived from a unique HGT event which took place in the common ancestor of the *Culex* and *Aedes* species.

The minimal number of independent gene loss events required was determined in order to evaluate the plausibility of the vertical transmission hypothesis (see Data File S2 for details). Following a conservative approach, at least 15 gene losses should be postulated, from Bilateria
to Culicidae, to explain the narrow taxonomic distribution of RIP genes in Metazoa. The alternative HGT hypothesis involves a single gene loss event in the lineage to extant metazoans followed by a single recent HGT event in the ancestor of *Culex* and *Aedes*, yielding a more parsimonious evolutionary scenario.

Based on the strong evidence supporting that culicinae RIPs are derived from an HGT event, a search for possible donors was conducted. The phylogeny of RIPs shows that culicine RIPs form a well supported monophyletic group (PP: 1) embedded within bacterial sequences (Fig 2), suggesting a prokaryotic origin of these sequences. The lack of introns in culicine RIPs also supports this idea. Moreover, BLASTp and tBLASTn searches using mosquito RIPs as queries yielded sequences belonging to *Tolypothrix bouteillei* (Cyanobacteria), *Calothrix parietina* (Cyanobacteria) and *Spiroplasma* spp (Tenericutes). In order to perform a more robust study we focused on culicine, including the recently reported sequences from *A. albopictus*, and on bacterial RIPs. Phylogenetic analysis showed that culicine RIPs form a monophyletic group (PP: 0.96) with sequences from *Spiroplasma* spp and Cyanobacteria species (Fig 4). Thus, these organisms—or others closely related—could have been potential donors.
Fig 4: Phylogenetic relationships among metazoan and bacterial RIPs. Numbers above branches indicate Posterior Probabilities (PP) values. Light blue, green and yellow backgrounds indicate *Spiroplasma*, *Cyanobacteria* and mosquitoes RIPs, respectively.

Horizontally acquired RIP genes evolve under purifying selection pressure

Except for a few well characterized potent toxins (*e.g.* ricin, shiga and shiga-like), the physiological role of most RIPs remains unknown (31). The defensive role demonstrated for *Spiroplasma* RIP genes in *Drosophila neotestacea* (13) invites us to postulate a biological function for HGT acquired RIP genes in mosquitoes. Therefore, we searched for signs of selective pressure on HGT derived sequences as reliable evidence of functionality in insects. To do this, all RIP encoding sequences from *C. quinquefasciatus*, *C. molestus*, *C. pipiens*, *C. torrentium*, *A. aegypti* and *A. albopictus* were aligned (Dataset S1). Interestingly, the observed INDELs were always multiple of three nucleotides, strongly suggesting that frame-shifting mutations were actively purged by selection pressure. Fig 5 shows a Logo representation of the MSA, where a higher conservation degree of first and second bases of each codon seems to be
the rule. Moreover, an integrative analysis of synonymous vs non-synonymous substitutions employing three different methods (SLAC, FEL and REL) showed purifying (negative) selection for 64 codons, including most of the amino acids forming the active site (Fig 5 and S3 Table).

Fig 5: Logo representation of the MSA used to analyze the synonymous vs non-synonymous substitution rates. Codons forming the active site are indicated by colored nucleotides (A and T: yellow, G and C: blue). Codons under significant purifying selection determined by the three tests (SLAC, FEL and REL) are underlined in red color. Codons detected to be under significant purifying selection by two out of the three tests, are underlined in green color.
Discussion

Ribosome inactivating proteins form a very interesting protein family displaying a patchy taxonomic distribution. As it was mentioned above, in a previous report we have found *in silico* evidence of the presence of RIP genes in two closely related species of mosquitos (11). Due to the uniqueness of this finding, in this work we have experimentally confirmed the presence and location of RIP genes in the *C. quinquefasciatus* genome, as well as in other *Culex* species. Moreover, new exhaustive searches by BLAST and HMMER on metazoan databases, revealed the presence of additional homologous genes in *A. albopictus* and *Armigeres sulhabatus*, confirming that the RIP gene family is taxonomically restricted to the Culicinae subfamily (Data File S1 and S2).

Currently, there is not a “gold standard” methodology for automatic and reliable detection of HGT. Moreover, several reports claiming the presence of foreign genes have been undermined after more thorough analyses due to artifacts or misinterpretations (19, 32-34). Therefore, careful integration of information derived from taxonomic distribution, phylogenetic inferences and biological information is needed to detect *bona fide* horizontally acquired genes. The evidence obtained in this work shows that the most plausible origin of culicine RIPs is a single HGT event to the cenancestor of *Culex* and *Aedes* genera. This model is supported by the monophyly of metazoan RIPs and their very narrow taxonomic distribution, the gathering in a clade along with prokaryotic sequences and the shared genomic context among Culicinae species.

According to phylogenetic inferences (Figs 2 and 4), the donor of the RIP gene was, most likely, a prokaryotic organism. An obvious donor candidate for insects is *Wolbachia* spp, since several HGT events between these bacteria and arthropods have been clearly documented (35-37). It is expected that animal genomes are marginally affected by HGT because of the separation of the germline from somatic cells. This barrier to HGT, known as Weismann barrier is not present in the case of bacteria infecting germline cells, as *Wolbacchia* spp, which is consistent with the relatively high number of *Wolbacchia* to insect HGT events (17). However,
no RIPS encoding sequences can be found in any of the *Wolbachia* spp databases (including 27 fully sequenced genomes). Interestingly, homology searches using mosquitoes RIPS as probe yielded significant similarity to several *Spiroplasma* genes (Fig 5). The facts that *Spiroplasma* species lack cell walls and that they are frequent endosymbionts of arthropods make them logical donor candidates. However, considering *Spiroplasma* spp as donor involves two major drawbacks. *Spiroplasma* spp coding sequences harbor very low GC content (around 23%), whereas RIPS from Diptera range from 44.8% to 55.6% (Table 1). Secondly, *Spiroplasma* spp and some species of *Millicutes* use a non-universal UGA tryptophan codon. This variation in the genetic code is presumed to have occurred in the early divergence of these genera (dating 250 mya approximately) (38), and the Culicinae subfamily has diverged more recently (between 51 to 204 mya) (39, 40), the transferred genes containing the non-universal UGA tryptophan codons would be read as a stop. Although CG content of a transferred functional gene could be gradually modified by amelioration, the reversion of several nonsense codons to Trp does not seem plausible. Therefore, these pieces of evidence lead us to reject the hypothesis that Diptera RIP genes are derived from *Spiroplasma*.

The phylogenetically closest sequences to culicine RIPS belong to Cyanobacteria species (*Tolypothrix bouteillei* and *Calothrix parietina*). Cyanobacteria constitute a significant fraction of the microbiota at breeding sites of mosquitoes. Remarkably, Cyanobacteria species account for 40% of the bacterial midgut content of larval and pupal stages in *An. gambiae* (41). Moreover, *Calothrix* sp has been detected in the midgut bacterial flora of *An. stephensi* in larval stage (42). In addition, GC content of Cyanobacteria genomes is closer to the Diptera RIPS (Table 1). Altogether; the presented phylogenetic inferences, the shared ecological niches between these bacteria and insects, and the colonization of mosquitoes in their early developmental stages (41) strongly suggest that Cyanobacteria are the most plausible donor species of RIP genes through HGT to the Culicinae subfamily. In line with Huang (17), we postulate that Weisman barrier is, if not absent, markedly weakened in the egg, pupa and larva.
stages of mosquitoes. Consequently, these early developmental stages of mosquitoes could be particularly prone to the acquisition of heritable foreign genes by environmental bacteria.

An important question about the HGT derived genes is their fate. In other words, the issue at stake is whether "foreign" genes will impact on fitness, and if so, to what extent these genes will be affected by natural selection and/or genetic drift. Probably, mosquitoes RIP genes display a defensive role which has helped them to be fixed by natural selection. However, the fact that individuals from the *C. quinquefasciatus* MR4 colony are homozygous for a null mutation of RIP gene shows that this gene is not essential for viability under laboratory conditions. On the other hand, all mosquitoes’ sequences have several codons under purifying selection, which is also reflected by the occurrence of the majority of changes at the third position of codons (Fig 5). According to the nearly-neutral evolutionary theory, slightly deleterious mutations can be fixed in populations with low effective size (as in the case of laboratory colonies). On the contrary, these mutations are efficiently purged from larger populations, such as natural populations (43). This seems to be the case of culicine RIPs, since null mutant *C. quinquefasciatus* are viable in captivity, while clear evidence of selection pressure on coding sequences in wild specimens has been found.

**Table 1. GC content analysis.** The GC content of RIP genes was calculated using DNA/RNA GC Content Calculator ([http://www.endmemo.com/bio/gc.php](http://www.endmemo.com/bio/gc.php)). The average of GC content in coding region was obtained from the codon usage database ([http://www.kazusa.or.jp/codon/](http://www.kazusa.or.jp/codon/)).

| Organism              | RIP name | mRNA GenBank  | GC % | average in coding GC |
|-----------------------|----------|---------------|------|----------------------|
| *Culex quinquefasciatus* | RIPcu     | XM_001850821.1 | 53.7 | 51.07%               |
| *Culex pipiens*       | RIPpi     | KX674697       | 55.6 | 52.48%               |
| *Aedes aegypti*       | RIPaeI    | XM_001650114.1 | 50.7 | 50.65%               |
|                       | RIPaeI LIKE |             | 44.8 |                       |
|                       | RIPaeII   | XM_001658803.1 | 39.8 |                       |
| *Aedes albopictus*    | RIPalII   |               | 50.9 | 53.2%                |
|                       | RIPalIII  |               | 50.2 |                       |
|                       | RIPalIV   |               | 46.5 |                       |
|                       | RIPalV    |               | 46.9 |                       |
|                       | RIPalVI   |               | 45.5 |                       |
|                       | RIPalVI   |               | 46.8 |                       |
|                       |           |               | 47.2 |                       |
| Spiroplasma poulsonii   | 26.3 | 24.90% |
|-------------------------|------|--------|
|                         | 25.3 |        |
| Tolypothrix sp           | WP_038092084.1 | 38.1 | 41.73% |
| Calothrix sp             | WP_015197802.1 | 35.8 | 45.97% |
| Ancestral RIP Aedes      |       | 51.1   |
| Ancestral RIP Culex      |       | 54.5   |
| Ancestral RIP metazoan   |       | 55.4   |

**Conclusions**

We have found solid evidence suggesting that culicine RIP genes derive from a single ancestral sequence acquired by HGT from a prokaryotic organism, probably a Cyanobacterium. Moreover, we have also demonstrated that these genes have evolved, after transference, under purifying selection, implying a functional role in the host organism.

**Acknowledgements**

The DNA samples of *C. molestus*, *C. pipiens* and *C. torrentium* were kindly gifted by Dr. Stefanie Becker. The authors acknowledge the Malaria Research and Reference Reagent Resource Center (MR4, CDC) for providing the following mosquito strains through BEI Resources, NIAID, NIH: *Culex quinquefasciatus*, strain JHB, NR-43025. The authors are also grateful to Dr. Jimena Juri Ayub for her technical support. We would also like to thank GAECI for their services. The findings and conclusions in this manuscript are those of the authors and do not necessarily represent the views of the Centers for Disease Control and Prevention (CDC).
References

1. Endo Y, Mitsui K, Motizuki M, Tsurugi K. The mechanism of action of ricin and related toxic lectins on eukaryotic ribosomes. The site and the characteristics of the modification in 28S ribosomal RNA caused by the toxins. The Journal of biological chemistry. 1987;262(12):5908-12. Epub 1987/04/25.

2. Endo Y, Tsurugi K. RNA N-glycosidase activity of ricin A-chain. Mechanism of action of the toxic lectin ricin on eukaryotic ribosomes. The Journal of biological chemistry. 1987;262(17):8128-30. Epub 1987/06/15.

3. Hudak KA, Dinman JD, Tumer NE. Pokeweed antiviral protein accesses ribosomes by binding to L3. The Journal of biological chemistry. 1999;274(6):3859-64. Epub 1999/01/28.

4. Rajamohan F, Ozer Z, Mao C, Uckun FM. Active center cleft residues of pokeweed antiviral protein mediate its high-affinity binding to the ribosomal protein L3. Biochemistry. 2001;40(31):9104-14. Epub 2001/08/02.

5. Nilsson L, Nygard O. The mechanism of the protein-synthesis elongation cycle in eukaryotes. Effect of ricin on the ribosomal interaction with elongation factors. European journal of biochemistry / FEBS. 1986;161(1):111-7. Epub 1986/11/17.

6. Sperti S, Montanaro L, Mattioli A, Stirpe F. Inhibition by ricin of protein synthesis in vitro: 60 S ribosomal subunit as the target of the toxin. The Biochemical journal. 1973;136(3):813-5. Epub 1973/11/01.

7. Gribé T, Ferreras JM, Arias FJ, Stirpe F. Description, distribution, activity and phylogenetic relationship of ribosome-inactivating proteins in plants, fungi and bacteria. Mini reviews in medicinal chemistry. 2004;4(5):461-76.

8. Reyes AG, Geukens N, Gutschoven P, De Graeve S, De Mot R, Mejia A, et al. The Streptomyces coelicolor genome encodes a type I ribosome-inactivating protein. Microbiology. 2010;156(Pt 10):3021-30. Epub 2010/07/03.

9. Sandvig K. Shiga toxins. Toxicon. 2001;39(11):1629-35.

10. Di Maro A, Citores L, Russo R, Iglesias R, Ferreras JM. Sequence comparison and phylogenetic analysis by the Maximum Likelihood method of ribosome-inactivating proteins from angiosperms. Plant molecular biology. 2014;85(6):575-88. Epub 2014/06/02.

11. Lapadula WJ, Sanchez Puerta MV, Juri Ayub M. Revising the taxonomic distribution, origin and evolution of ribosome inactivating protein genes. PloS one. 2013;8(9):e72825. Epub 2013/09/17.

12. Peumans WJ, Van Damme EJ. Evolution of plant ribosome-inactivating proteins. Toxic plant proteins: Springer; 2010. p. 1-26.

13. Hamilton PT, Peng F, Boulanger MJ, Perlman SJ. A ribosome-inactivating protein in a Drosophila defensive symbiont. Proceedings of the National Academy of Sciences of the United States of America. 2016;113(2):350-5. Epub 2015/12/30.

14. Gogarten JP, Doolittle WF, Lawrence JG. Prokaryotic evolution in light of gene transfer. Mol Biol Evol. 2002;19(12):2226-38. Epub 2002/11/26.

15. Treangen TJ, Rocha EP. Horizontal transfer, not duplication, drives the expansion of protein families in prokaryotes. PLoS Genet. 2011;7(1):e1001284.

16. Ku C, Nelson-Sathi S, Roettger M, Sousa FL, Lockhart PJ, Bryant D, et al. Endosymbiotic origin and differential loss of eukaryotic genes. Nature. 2015;524(7566):427-32. Epub 2015/08/20.

17. Huang J. Horizontal gene transfer in eukaryotes: the weak-link model. BioEssays : news and reviews in molecular, cellular and developmental biology. 2013;35(10):868-75. Epub 2013/09/17.
18. Wybouw N, Pauchet Y, Heckel DG, Van Leeuwen T. Horizontal Gene Transfer Contributes to the Evolution of Arthropod Herbivory. Genome biology and evolution. 2016;8(6):1785-801. Epub 2016/06/17.
19. Lander ES, Linton LM, Birren B, Nusbaum C, Zody MC, Baldwin J, et al. Initial sequencing and analysis of the human genome. Nature. 2001;409(6822):860-921.
20. Boothby TC, Tenlen JR, Smith FW, Wang JR, Patanella KA, Nishimura EO, et al. Evidence for extensive horizontal gene transfer from the draft genome of a tardigrade. Proceedings of the National Academy of Sciences. 2015;112(52):15976-81.
21. Salzberg SL, White O, Peterson J, Eisen JA. Microbial genes in the human genome: lateral transfer or gene loss? Science. 2001;292(5523):1903-6.
22. Richards TA, Monier A. A tale of two tardigrades. Proceedings of the National Academy of Sciences. 2016;113(18):4892-4.
23. Collins FH, Mendez MA, Rasmussen MO, Mehaffey PC, Besansky NJ, Finnerty V. A ribosomal RNA gene probe differentiates member species of the Anopheles gambiae complex. The American journal of tropical medicine and hygiene. 1987;37(1):37-41.
24. Rudolf M, Czajka C, Borstler J, Melaun C, Jost H, von Thien H, et al. First nationwide surveillance of Culex pipiens complex and Culex torrentium mosquitoes demonstrated the presence of Culex pipiens biotype pipiens/molestus hybrids in Germany. PloS one. 2013;8(9):e71832. Epub 2013/09/17.
25. Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Höhna S, et al. MrBayes 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space. Systematic biology. 2012;61(3):539-42.
26. Suyama M, Torrents D, Bork P. PAL2NAL: robust conversion of protein sequence alignments into the corresponding codon alignments. Nucleic acids research. 2006;34(suppl 2):W609-W12.
27. Delport W, Poon AF, Frost SD, Pond SLK. Datamonkey 2010: a suite of phylogenetic analysis tools for evolutionary biology. Bioinformatics. 2010;26(19):2455-7.
28. Pond SLK, Frost SD. Not so different after all: a comparison of methods for detecting amino acid sites under selection. Molecular biology and evolution. 2005;22(5):1208-22.
29. Crooks GE, Hon G, Chandonia J-M, Brenner SE. WebLogo: a sequence logo generator. Genome research. 2004;14(6):1188-90.
30. Lapadula WJ, Sanchez-Puerta MV, Ayub MJ. Convergent evolution led ribosome inactivating proteins to interact with ribosomal stalk. Toxicon : official journal of the International Society on Toxinology. 2012;59(3):427-32. Epub 2012/01/17.
31. Puri M, Kaur I, Perugini MA, Gupta RC. Ribosome-inactivating proteins: current status and biomedical applications. Drug discovery today. 2012;17(13-14):774-83. Epub 2012/04/10.
32. Koutsovoulos G, Kumar S, Laetsch DR, Stevens L, Daub J, Conlon C, et al. No evidence for extensive horizontal gene transfer in the genome of the tardigrade Hypsibius dujardini. Proceedings of the National Academy of Sciences. 2016;113(18):5053-8.
33. Kurland CG, Canback B, Berg OG. Horizontal gene transfer: a critical view. Proceedings of the National Academy of Sciences. 2003;100(17):9658-62.
34. Laurence M, Hatzis C, Brash DE. Common contaminants in next-generation sequencing that hinder discovery of low-abundance microbes. PloS one. 2014;9(5):e97876.
35. Dunning Hotopp JC, Clark ME, Oliveira DC, Foster JM, Fischer P, Munoz Torres MC, et al. Widespread lateral gene transfer from intracellular bacteria to multicellular eukaryotes. Science. 2007;317(5845):1753-6. Epub 2007/09/01.
36. Klasson L, Kambris Z, Cook PE, Walker T, Sinkins SP. Horizontal gene transfer between Wolbachia and the mosquito Aedes aegypti. Bmc Genomics. 2009;10(1):1.
37. Woolfit M, Iturbe-Ormaetxe I, McGraw EA, O'Neill SL. An ancient horizontal gene transfer between mosquito and the endosymbiotic bacterium Wolbachia pipientis. Molecular biology and evolution. 2009;26(2):367-74.
38. Razin S, Yoge D, Naot Y. Molecular biology and pathogenicity of mycoplasmas. Microbiology and Molecular Biology Reviews. 1998;62(4):1094-156.
39. Reidenbach KR, Cook S, Bertone MA, Harbach RE, Wiegmann BM, Besansky NJ. Phylogenetic analysis and temporal diversification of mosquitoes (Diptera: Culicidae) based on nuclear genes and morphology. BMC evolutionary biology. 2009;9:298. Epub 2009/12/24.
40. Foley DH, Bryan JH, Yeates D, Saul A. Evolution and systematics of Anopheles: insights from a molecular phylogeny of Australasian mosquitoes. Molecular phylogenetics and evolution. 1998;9(2):262-75. Epub 1998/05/01.
41. Wang Y, Gilbreath III TM, Kukutla P, Yan G, Xu J. Dynamic gut microbiome across life history of the malaria mosquito Anopheles gambiae in Kenya. PloS one. 2011;6(9):e24767.
42. Minard G, Mavingui P, Moro CV. Diversity and function of bacterial microbiota in the mosquito holobiont. Parasites & vectors. 2013;6:146. Epub 2013/05/22.
43. Ohta T. The nearly neutral theory of molecular evolution. Annual Review of Ecology and Systematics. 1992;23:263-86.
## Supplementary Information

| Information | Page nº |
|-------------|---------|
| S1 Table    | 2       |
| S1 Fig      | 3       |
| S2 Fig      | 4       |
| S3 Fig      | 5       |
| S2 Table    | 6       |
| S3 Table    | 7       |
S1 Table: Primer sequences and annealing temperatures (Ta) are indicated in second and third column respectively.

| Seq. Names  | Primer sequences          | Ta °C |
|-------------|---------------------------|-------|
| Fw RIPcu    | ATGCAGTTCAACCGGC          | 64 °C |
| Rev RIPcu   | TTAGAAGCAATTCTTTGCGAAAAATTC | 64°C  |
| Fw intergen | GCAGAGGATTGAGAAGAAAGG     | 60°C  |
| Rev intergen| CCGTTTATGGAGAACTGGAGAAG | 60°C  |
S1 Fig: PCR amplification of RIP genes in species belonging to the *Culex* genera. *C. pipiens* (C.p, lane 1), *C. molestus* (C.m, lanes 2 and 3) and *C. torrentium* (C.t, lanes 4 and 5). Lane 6: 1Kb molecular weight size marker, lane 7: negative control.
S2 Fig: MSA of partial RIPv genes found in *Culex* ssp. Gaps are indicated in green color.

| RIPv | C. quinquefasciatus | Culex molesus | Culex pipiens | Culex torrentium |
|------|---------------------|---------------|---------------|-----------------|
| RIPv | VYGAGTCGA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA | VYGAGTCGA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA | VYGAGTCGA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA | VYGAGTCGA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA ACGGGCAAA |

| RIPv | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC |

| RIPv | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC |

| RIPv | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC |

| RIPv | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC |

| RIPv | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC |

| RIPv | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC |

| RIPv | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC |

| RIPv | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC |

| RIPv | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC | KGCCCATAT CTAATGAGG TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC TCTATCGTAC |
S3 Fig: Electropherograms of the region including the deletion of ten nucleotides in *C. quinquefasciatus* JHB MR4 colony. In all cases the region was clearly resolved. Top: sequence of *C. quinquefasciatus* JHB MR4 colony displaying the ten nucleotides deletion. Bottom: Chromatogram of *C. molestus* RIP sequence where no deletion is observed.
S2 Table: Orthologs genes between *C. quinquefasciatus*, *A. aegypti* and *An. gambiae*. The GenBank and Vector Base codes are indicated for all protein sequences. The code of each protein used in Fig 3 is shown in the last column.

| C. quinquefasciatus | A. aegypti | A. gambiae |
|---------------------|------------|------------|
| GenBank             | Vector Base code | Gen Bank   | Vector Base code | e Value | GenBank | Vector Base code | e Value | Code |
| XP_001850870        | CPU009208   | Gen Bank   | XP_001650164    | AAE005004 | 1 e-93  | A           |
| XP_001850871        | CPU009209   | Gen Bank   | XP_001650164    |         |         | B           |
| XP_001850872        | CPU009210   | Gen Bank   | XP_001650165    | AAE004999 | 5 e-65  | C           |
| XP_001850873        | CPU009211   | Gen Bank   | XP_001650165    |         |         | D           |
| XP_001850874        | CPU009212   | Gen Bank   | XP_001650165    |         |         | E           |
| XP_001850875        | CPU009213   | Gen Bank   | XP_001650165    |         |         | F           |
| XP_001850876        | CPU009214   | Gen Bank   | XP_001650165    |         |         | G           |
| XP_001850877        | CPU009215   | Gen Bank   | XP_001650165    |         |         | H           |
| XP_001850878        | CPU009216   | Gen Bank   | XP_001650165    |         |         | I           |
| XP_001850879        | CPU009217   | Gen Bank   | XP_001650165    |         |         | J           |
| XP_001850880        | CPU009218   | Gen Bank   | XP_001650165    |         |         | K           |
S3 Table: Integrative analyses of substitution rates by SLAC, FEL and REL tests. Those codons under non-neutral selection for each test are presented in violet color. In the consensus column, those codons indicated with blue and dash white inside the box display dN > dS have significant difference. These results were estimated using a p-value < 0.01 for SLAC and FEL tests, and a Bayes factor cutoff= 50 for REL test. Active site codons are indicated in yellow.

| Code | SLAC dN | SLAC dS | FEL dN | FEL dS | REL dN | REL dS | Consensus |
|------|---------|---------|--------|--------|--------|--------|-----------|
| 1    | 0.000   | 0.000   | 1.000  | -0.930 | 0.544  | 0.460  |           |
| 9    | -2.072  | 0.043   | -2.006 | 0.003  | -1.461 | 48.762 |           |
| 23   | 1.815   | 0.005   | 1.460  | 0.000  | -1.114 | 7.382  | 0.030     |
| 30   | -1.216  | 0.039   | -489.770 | 0.000  | -1.102 | 477.649 |           |
| 33   | 0.605   | 0.180   | 0.172  | 0.054  | -0.836 | 718.730 |           |
| 34   | 0.405   | 0.247   | -0.120 | 0.078  | -0.764 | 219.770 |           |
| 37   | -1.235  | 0.003   | -0.005 | 0.000  | -0.322 | 1114.200 |           |
| 46   | -1.557  | 0.072   | -0.890 | 0.007  | -0.856 | 86.225  |           |
| 48   | -1.517  | 0.052   | -1.227 | 0.007  | -0.941 | 10.976  |           |
| 49   | 1.485   | 0.012   | 1.065  | 0.000  | -1.178 | 1108.660 |           |
| 53   | -1.539  | 0.012   | -0.408 | 0.002  | -0.885 | 473.736 |           |
| 55   | -0.902  | 0.112   | -0.546 | 0.036  | -0.895 | 162.204 |           |
| 59   | -0.900  | 0.037   | -0.639 | 0.003  | -0.867 | 306.780 |           |
| 61   | -0.300  | 0.103   | 0.073  | 0.121  | -0.575 | 655.687 |           |
| 62   | -1.401  | 0.051   | -1.330 | 0.009  | -1.028 | 393.040 |           |
| 66   | -0.520  | 0.215   | -0.256 | 0.019  | -0.848 | 600.612 |           |
| 68   | -1.549  | 0.013   | -1.407 | 0.019  | -1.099 | 202.748 |           |
| 79   | -1.153  | 0.015   | -0.526 | 0.031  | -0.884 | 775.776 |           |
| 82   | -1.787  | 0.005   | -2.016 | 0.000  | -1.146 | 4880.320 |           |
| 84   | -0.605  | 0.128   | -0.121 | 0.008  | -0.827 | 1114.793 |           |
| 85   | -1.134  | 0.018   | -0.442 | 0.002  | -0.990 | 345.605 |           |
| 87   | -0.900  | 0.042   | -0.240 | 0.009  | -0.816 | 248.194 |           |
| 91   | -1.258  | 0.047   | -0.976 | 0.004  | -0.913 | 355.885 |           |
| 97   | 1.688   | 0.016   | 3.395  | 0.000  | -1.157 | 361.017 |           |
| 120  | -1.170  | 0.076   | -1.137 | 0.007  | -0.928 | 84.057  |           |
| 131  | -0.984  | 0.110   | -5.106 | 0.000  | -2.881 | 519.780 |           |
| 135  | -1.207  | 0.038   | -0.600 | 0.000  | -0.871 | 688.034 |           |
| 137  | -1.707  | 0.099   | -0.043 | 0.001  | -0.918 | 976.519 |           |
| 140  | -1.628  | 0.109   | -0.549 | 0.000  | -0.907 | 2725.530 |           |
| 151  | -1.743  | 0.036   | -1.153 | 0.006  | -0.903 | 83.679  |           |
| 153  | -1.377  | 0.083   | -2.202 | 0.001  | -1.264 | 772.423 |           |
| 155  | -1.314  | 0.033   | -1.182 | 0.002  | -0.998 | 214.374 |           |
| 157  | -1.451  | 0.020   | -0.945 | 0.001  | -0.948 | 575.293 |           |
| 158  | -1.065  | 0.025   | -0.661 | 0.001  | -0.890 | 555.446 |           |
| 159  | -0.836  | 0.050   | -0.136 | 0.007  | -0.839 | 219.591 |           |
| 159  | -1.675  | 0.005   | -1.405 | 0.000  | -1.112 | 202.302 |           |
| 166  | -0.585  | 0.171   | -0.198 | 0.046  | -0.816 | 400.256 |           |
| 169  | -0.502  | 0.215   | -0.204 | 0.009  | -0.839 | 429.773 |           |
| 173  | -1.604  | 0.070   | -1.644 | 0.070  | -1.208 | 30.940  |           |
| 178  | -0.786  | 0.117   | -0.678 | 0.003  | -0.888 | 73.948  |           |
| 180  | -1.199  | 0.012   | -0.002 | 0.000  | -0.901 | 493.560 |           |
| 183  | -0.557  | 0.179   | -0.199 | 0.004  | -0.791 | 488.112 |           |
| 185  | -1.757  | 0.056   | -1.525 | 0.002  | -1.103 | 654.990 |           |
| 187  | -1.075  | 0.039   | -0.333 | 0.009  | -0.848 | 573.723 |           |
| 188  | -1.199  | 0.012   | -0.155 | 0.000  | -0.916 | 2066.590 |           |
| 196  | -1.419  | 0.005   | -0.777 | 0.000  | -0.928 | 521.305 |           |
| 197  | -1.176  | 0.029   | -0.417 | 0.005  | -0.880 | 146.650 |           |
| 192  | -0.049  | 0.070   | -0.702 | 0.003  | -0.880 | 546.206 |           |
| 193  | -0.349  | 0.018   | -0.798 | 0.005  | -0.871 | 349.318 |           |