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Osvaldo Marinotti  
*University of California Irvine*

Adam R. Wespiser  
*University of Massachusetts Medical School*

Daniel R. Caffrey  
*University of Massachusetts Medical School*

*See next page for additional authors*

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The Genome of *Anopheles darlingi*, the main neotropical malaria vector

Osvaldo Marinotti¹,*, Gustavo C. Cerqueira², Luiz Gonzaga Paula de Almeida³, Maria Inês Tiraboschi Ferro⁴, Elgion Lucio da Silva Loreto⁵, Arnaldo Zaha⁶, Santuza M. R. Teixeira⁷, Adam R. Wespirer⁸, Alexandre Almeida e Silva⁹, Aline Daiane Schlindwein¹⁰, Ana Carolina Landim Pacheco¹¹,¹², Artur Luiz da Costa da Silva¹³, Brenton R. Graveley¹⁴, Brian P. Walenz¹⁵, Bruna de Araujo Lima¹⁶, Carlos Alexandre Gomes Ribeiro¹⁷, Carlos Gustavo Nunes-Silva¹⁸, Carlos Roberto de Carvalho¹⁹, Célia Maria de Almeida Soares²⁰, Claudia Beatriz Afonso de Menezes²¹, Cleverston Matiollii²², Daniel Caffrey⁸, Demetrius Antonio M. Araújo²³, Diana Magalhães de Oliveira¹¹, Douglas Golenbock⁸, Edmundo Carlos Grisard¹⁰, Fabiana Fantinatti-Garboggini²¹, Fabiola Marques de Carvalho³, Fernando Gomes Barcellos²⁴, Francisco Prosdocimi²⁵, Gemma May¹³, Gilson Martins de Azevedo Junior²⁶, Giselle Moura Guimarães²⁶, Gustavo Henrique Goldman²⁷,²⁸, Itácio Q. M. Padilha²¹, Jacqueline da Silva Batista²⁹, Jesus Aparecido Ferro⁴, José M. C. Ribeiro³⁰, Juliana Lopes Rangel Fietto³¹, Karina Maia Dabbas⁴, Louise Cerdeira³, Lucymara Fassarella Agnez-Lima³², Marcelo Brocchi¹⁵, Marcos Oliveira de Carvalho³³, Marcus de Melo Teixeira³⁴, Maria de Mascena Diniz Maia³⁵, Maria Helena S. Goldman³⁶, Maria Paula Cruz Schneider³⁷, Maria Sueli Soares Felipe³⁴,³⁸, Mariangela Hungria³⁹, Marisa Fabiana Nicolás³, Maristela Pereira¹⁹, Martín Alejandro Montes³⁵, Maurício E. Cantão³⁴,⁰¹, Michel Vincentz⁴¹, Miriam Silva Rafael⁴², Neal Silverman⁶, Patrícia Hermes Stoco¹⁰, Rangel Celso Souza³, Renato Vicentini⁴³, Ricardo Tostes Gazzinelli⁴⁴, Rogério de Oliveira Neves¹⁷, Rosane Silva⁴⁵, Spartaco Astolfi-Filho¹⁷, Talles Eduardo Ferreira Maciel³¹, Turán P. Úrmenyí⁴⁵, Wanderli Pedro Tadei⁴², Erney Plessmann Camargo⁴⁶ and Ana Tereza Ribeiro de Vasconcelos³,⁰

¹Department of Molecular Biology and Biochemistry, University of California Irvine, Irvine, CA 92697, USA, ²Institute of Technology, Broad Institute of Harvard and Massachusetts, Cambridge, MA 02141, USA, ³Laboratório de Bioinformática do Laboratório Nacional de Computação Científica, Petrópolis, RJ 25651-075, Brasil, ⁴Departamento de Tecnologia, Faculdade de Ciências Agrárias e Veterinárias de Jaboticabal, UNESP - Universidade Estadual Paulista, SP 14884-900, Brasil, ⁵Departamento de Biologia, Universidade Federal de Santa Maria, Santa Maria, RS 97105-900, Brasil, ⁶Departamento de Biologia Molecular e Biotecnologia, Centro de Biotecnologia, Universidade Federal do Rio Grande do Sul, Porto Alegre, RS 91501-970, Brasil, ⁷Departamento de Bioquímica e Imunologia, Universidade Federal de Minas Gerais, Belo Horizonte, MG 31270901, Brasil, ⁸Department of Medicine, University of Massachusetts Medical School, Worcester, MA 01655, USA, ⁹Laboratório de Entomologia Médica IPEPATRO/FIOCRUZ, Porto Velho, RO 76812-245, Brasil, ¹⁰Departamento de Microbiologia, Imunologia e Parasitologia, Universidade Federal de Santa Catarina, Florianópolis, SC 88040-900, Brasil, ¹¹Centro de Ciências da Saúde, Universidade Estadual do Ceará, Fortaleza, CE 62042-280, Brasil, ¹²Departamento de Ciências Biológicas, Campus Senador Helvídio Nunes de

*To whom correspondence should be addressed. Tel: +55 24 2233 6065; Fax: +55 24 2233 6124; Email: atrv@lncc.br
Correspondence may also be address to Osvaldo Marinotti. Tel: +1 949 824 3210; Fax: +1 949 824 2814; Email: omarinot@uci.edu

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This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/3.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.
Anopheles darlingi is the principal neotropical malaria vector, responsible for more than a million cases of malaria per year on the American continent. Anopheles darlingi diverged from the African and Asian malaria vectors ~100 million years ago (mya) and successfully adapted to the New World environment. Here we present an annotated reference A. darlingi genome, sequenced from a wild population of males and females collected in the Brazilian Amazon. A total of 10481 predicted protein-coding genes were annotated, 72% of which have their closest counterpart in Anopheles gambiae and 21% have highest similarity with other mosquito species. In spite of a long period of divergent evolution, conserved gene synteny was observed between A. darlingi and A. gambiae. More than 10 million single nucleotide polymorphisms and short indels with potential use as genetic markers were identified. Transposable elements correspond to 2.3% of the A. darlingi genome. Genes associated with hematophagy, immunity and insecticide resistance, directly involved in
vector–human and vector–parasite interactions, were identified and discussed. This study represents the first effort to sequence the genome of a neotropical malaria vector, and opens a new window through which we can contemplate the evolutionary history of anopheline mosquitoes. It also provides valuable information that may lead to novel strategies to reduce malaria transmission on the South American continent. The A. darlingi genome is accessible at www.labinfo.lncc.br/index.php/anopheles-darlingi.

INTRODUCTION

Anopheles darlingi is the principal neotropical malaria vector (1–6), sustaining the transmission of more than a million malaria cases per year on the American continent [(7), World Health Organization Malaria Report 2011]. Anopheles darlingi has a wide geographic distribution that reaches from Southern Mexico to Northern Argentina and from East of the Andes chain to the coast of the Atlantic Ocean. Although this species has been subjected to extensive study, little is known about the molecular aspects of its biology. The A. darlingi genome presented here fills this gap in the knowledge about its genes, transcripts and proteins that determine the biological characteristics of this important malaria vector.

In spite of the availability of published genomes for three other mosquito species [Anopheles gambiae (8), Aedes aegypti (9), Culex quinquefasciatus (10)], the medical and epidemiological significance of A. darlingi and its phylogenetic position support the importance of this study. Anopheles (Nyssorhynchus) darlingi and A. (Cellia) gambiae are considered to have diverged ~100 mya (11) (Figure 1), suggesting that their most recent common ancestor lived before the geological split of western Gondwana (~95 mya). This estimation is supported by the absence of the Cellia species in the New World and Nyssorhynchus in the Afro-Eurasian continents. The most ancient human colonization of the American continent is still a matter of discussion and is estimated to have occurred 30 000–10 000 years ago, indicating that A. darlingi and its ancestral species evolved in an environment devoid of humans or human ancestral species for several million years. Furthermore, European colonialists transferred Plasmodium falciparum and Plasmodium vivax, the most prevalent malaria parasites, to the American continent in post-Colombian times (17,18). Therefore, interactions between neotropical malaria vectors and humans, and malaria parasites, are relatively recent. The evolutionary history of A. darlingi thus allows tackling basic and unanswered questions about vector–parasite and vector–human host interactions as well as about malaria parasite development within its vectors and the mosquito immune responses to the developing parasite.

MATERIALS AND METHODS

Genome

Gravid A. darlingi female mosquitoes were captured from Coari, Amazonas State, Brazil, and their progeny (F1) was reared at the insectary of the Laboratory of Malaria and Dengue Vectors, Instituto Nacional de Pesquisas da Amazônia, Manaus, Brazil. Larvae were fed powdered fish food (Tetramin®), and pupae were transferred to plastic cups filled with distilled water. Total DNA was extracted from 1884 recently emerged adults (F1, <24 h after emergence), males and females, and was used for sequencing. High-coverage whole-genome data sets were generated by 454 Life Sciences (Roche) technology using single fragment end and paired-end reads. The reads were assembled using Celera Assembler 6.1. Because the sequenced DNA was sampled from a large number of field-captured individuals, the assembly was performed with a relaxed error tolerance of 16%, except during unigig construction where it was 12%. K-mer size overlap generation was also relaxed to 16 bases.

Transcriptome

The transcriptome of adult A. darlingi was derived from two mosquito populations that were captured 524 km apart from each other (Coari, Amazonas State and Porto Velho, Rondonia State, Brazil). The extracted RNA was sequenced using two next-generation sequencing platforms: 454 Life Sciences (Roche) and Illumina (Solexa sequencing). Transcripts were reconstructed using mapping first strategy, Genomic Short-read Nucleotide Alignment Program and Scripture and the assembly first strategy, Velvet/Oases. Reconstructed transcripts were used as supporting evidence on the annotation of the genome (PASA - Program to Assemble Spliced Alignments). Additional details on genomic DNA and RNA extraction, sample preparation, sequencing, assembly and annotation are given in Supplementary Method SA.

RESULTS AND DISCUSSION

Genome size, genome and transcriptome sequencing, assembly and annotation

Five and a half billion base pairs of information were generated, resulting in an assembled A. darlingi genome.
that spans 173.9 Mb (Tables 1 and 2) (see Supplementary Tables SA1 and SA2). The size of the A. darlingi haploid genome was determined by cytometric analysis to be ~201 Mb (2C = 0.41 pg) (see Supplementary Method SB and Supplementary Figure SB1), which is ~30% smaller than the genome of A. gambiae [278 Mb, (8)] and three to six times smaller than the genome of culicinae mosquitoes C. quinquefasciatus [579 Mb, (10)] and A. aegypti [1379 Mb, (9)] but larger than the Drosophila melanogaster genome [176 Mb, (19)]. The difference between the cytometrically determined genome size and the sum of all of the contigs and scaffolds is most likely the result of unassembled centromeres, telomeres and other portions of the genome that are rich in repetitive DNA sequences. In fact, 18.66 percent of the reads were not included in the final assembly. Assuming a uniform coverage of 20x and a read average length of 248 bp, the unassembled reads correspond to 32.71 Mb, which accounts for an estimated total genome length of 206.6 Mb, a value that is similar to the value obtained by cytometry. Although the A. darlingi genome is smaller than that of A. gambiae, the sums of the lengths of all of the protein coding sequences in each of the two genomes are similar (18.2 and 19.3 Mb, respectively), which indicates a more compact genome in A. darlingi mosquitoes (see Supplementary Tables SA3 and SA4). Anopheles darlingi has shorter intergenic and intronic sequences and fewer transposable elements (TEs; these elements constitute only 2.3% of the genome; see details below). Nevertheless, A. darlingi genes display a larger average number of exons per gene (4.6) than A. gambiae (4.4) (see Supplementary Table SA5).

DNA sequences of bacterial origin were obtained along with the A. darlingi genome. For example, the complete genome of Aeromonas hydrophila was assembled during an initial analysis of the 454 reads. DNA sequences of bacterial origin were labeled as contaminants and were screened out during the assembly process. Even after applying the bacterial DNA filter, the assembled A. darlingi genome includes genes of apparent bacterial origin. The majority of these are present in small contigs (mostly <10 kb) that do not contain evident mosquito DNA, which suggests that they derive from environmental contaminations or additional microorganisms that are associated with A. darlingi. Some scaffolds apparently contain sequences of both prokaryotes and eukaryotes. Further analyses are necessary to determine the legitimacy of these assembled scaffolds and the possibility of horizontal gene transfer events that may have contributed to shaping the A. darlingi genome.

Two similar mitochondrial genomes were previously described for this species, corresponding to the Southern and Northern genotypes, which originated from Manaus, Brazil and Central Cayo District, Belize, respectively. The typical 37 genes in animal mtDNA, comprising 13 protein-encoding genes, two rRNA genes (12S rRNA and 16S rRNA), 22 tRNA genes and a control region, are found in the complete A. darlingi mitochondrial genome (11). Here, we describe a third mitochondrial genome for this species, from mosquitoes captured in Coari, Brazil, which is more similar to the Southern genotype (see Supplementary Data SC and Supplementary Figure SC1). For the first time, we report the complete A. darlingi nuclear ribosomal RNA cistron (AD11084), complementing previously published, partial rRNA sequences (20,21). Sets of 359 nuclear encoded tRNAs and 44 homologs of A. gambiae pre-microRNAs (miRNAs) were identified. miRNA precursor candidates conserved in the genomes of A. darlingi and A. gambiae, which might play important roles in the posttranscriptional

Table 1. Assembly statistics of A. darlingi reference genome

<table>
<thead>
<tr>
<th>Feature</th>
<th>Statistics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total number of good sequence reads</td>
<td>16777488</td>
</tr>
<tr>
<td>Sequence reads in assembly</td>
<td>14139351</td>
</tr>
<tr>
<td>Total number of scaffolds</td>
<td>8233</td>
</tr>
<tr>
<td>Total length of scaffolds</td>
<td>173918288</td>
</tr>
<tr>
<td>Total number of contigs</td>
<td>13857</td>
</tr>
<tr>
<td>Combined bases in contigs</td>
<td>172639290</td>
</tr>
<tr>
<td>Combined length of gaps</td>
<td>1278998</td>
</tr>
<tr>
<td>Sequencing coverage</td>
<td>20</td>
</tr>
<tr>
<td>N50 scaffold length</td>
<td>81222</td>
</tr>
<tr>
<td>N50 contig length</td>
<td>37754</td>
</tr>
<tr>
<td>Longest scaffold (number of contigs)</td>
<td>1087588 (10)</td>
</tr>
<tr>
<td>Shortest scaffold (number of contigs)</td>
<td>473 (1)</td>
</tr>
</tbody>
</table>

Table 2. General characteristics of the A. darlingi genome

<table>
<thead>
<tr>
<th>Genome feature</th>
<th>A. darlingi</th>
<th>A. gambiae(^a)</th>
<th>A. aegypti(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genome size (Mb)</td>
<td>173.92</td>
<td>278.25</td>
<td>1379</td>
</tr>
<tr>
<td>Percent of G+C (%)</td>
<td>48.15</td>
<td>40.9</td>
<td>38.2</td>
</tr>
<tr>
<td>Protein coding length (Mb) and (% genome length)</td>
<td>18.2 (10.4)</td>
<td>64.92 (23.3)</td>
<td>224.9 (16.3)</td>
</tr>
<tr>
<td>Total number of exons</td>
<td>47990(^b)</td>
<td>56210</td>
<td>66827</td>
</tr>
<tr>
<td>Number of protein-coding genes</td>
<td>10457b</td>
<td>12670</td>
<td>15704</td>
</tr>
<tr>
<td>Percent genes with introns (%)</td>
<td>91.59</td>
<td>93.6</td>
<td>90.1</td>
</tr>
<tr>
<td>Average number of exons per gene</td>
<td>4.6</td>
<td>4.4</td>
<td>4.0</td>
</tr>
<tr>
<td>Average gene length (bp)</td>
<td>1735</td>
<td>5124(^c)</td>
<td>14587(^c)</td>
</tr>
<tr>
<td>Total tRNAs</td>
<td>346</td>
<td>450</td>
<td>995</td>
</tr>
</tbody>
</table>

\(^a\)Statistics were derived from genome updates for A. gambiae AgamP3 (Vectorbase, version 66.3) and A. aegypti AaegL1 (Vectorbase, version 66.1).

\(^b\)Includes 13 mitochondrial genes.

\(^c\)Includes introns but not untranslated regions.
regulation of gene expression in these species, were described in a separate publication (22).

Synteny

In spite of ~100 million years of evolutionary divergence between *A. darlingi* and *A. gambiae*, the gene synteny between their genomes is relatively well conserved. Translocation events have occurred but were mostly restricted to large intra-chromosomal rearrangements (Figure 2). The synteny between *A. darlingi* and *D. melanogaster* presents a different scenario: each one of the 12 largest *A. darlingi* scaffolds have orthologous genes scattered through different *D. melanogaster* chromosomes, which suggests a low degree of synteny (Figure 2B).

Systematic synteny evaluation between *A. darlingi* and *A. gambiae* identified 1027 synteny clusters (Figure 3A), comprising 6312 syntenic genes or ~60% of all *A. darlingi* protein-coding genes (Figure 2B). Apart from giving an idea on how much large-scale rearrangements have been important in the divergence of these species, this analysis will help in future efforts for gene identification on the basis of conserved synteny. Similar analyses between *A. darlingi* and other dipterans, i.e. *A. aegypti*, *C. quinquefasciatus* and *D. melanogaster*, identified 848, 835 and 244 synteny clusters (Figure 3A) and 3680, 3684 and 488 syntenic genes (Figure 3B), respectively. The higher degree of synteny between *A. darlingi* and *A. aegypti* or *C. quinquefasciatus* in comparison with the values obtained by *A. darlingi*–*D. melanogaster* evaluation reflects the estimated divergence time among those species (Figure 1) and suggests that most of the interchromosomal rearrangements have taken place after the split of lineages that lead to Drosophilidae and Culicidae.

The median number of genes per synteny cluster was not significantly different among all of the pairwise synteny evaluations (Figure 3C). This observation is owing to the draft nature of the *A. darlingi* genome, which has a significant number of unclosed genome gaps; these gaps lead to premature ends of the synteny clusters. From all of the identified synteny clusters between *A. darlingi* and *A. gambiae*, 87% occur near scaffold ends (Figure 3), suggesting that those clusters will be extended further when the genome sequence gaps are mended. A detailed *A. darlingi* cytogenetic map has been described (23–28) (see Supplementary Figure SD1). It is expected that mapping of particular genes or clones on chromosomes, together with the described synteny clusters, will support a more complete and precise assembly of the *A. darlingi* genome.

**Polymorphism within and between two populations**

A database with >10 million single-nucleotide variants (SNVs) and short indels with potential use as genetic
markers was created (Table 3) (see Supplementary Method SA). Differently from most of the previous studies of sequence polymorphisms in mosquitoes, that analyzed individuals pooled from established colonies in which much of the natural diversity is lost, the A. darlingi data presented here was generated from wild caught mosquitoes. The sequencing of the 278 Mb of the A. gambiae genome revealed 445 thousand single-nucleotide polymorphisms (SNPs), with an average heterozygosity at the nucleotide level of 1.6 per kb (9). The average frequency of nucleotide variation was reported to be 7 and 12 SNPs per kb for Anopheles funestus and A.aegypti (29,30), respectively. An SNP frequency of 17 per kb was recently reported for selected gene fragments of field-captured Anopheles arabiensis (31).

Because laboratory autonomous colonies of A. darlingi are not available, the DNA and RNA sequenced in this project were extracted from >1884 individuals (F1 progeny of field-captured gravid females). While the high degree of polymorphism found in A. darlingi reads posed a challenge for genome assembly, the data acquired permitted a better representation of the sequence polymorphisms in two natural populations of this malaria vector. The distribution of SNVs is not homogeneous throughout the genome, and average values as high as 50 SNVs per kb in intergenic and intronic sequences were observed, with lower values in protein coding genes, including untranslated regions (UTRs) (40 SNVs per kb), and even lower values (26 SNVs per kb) in protein coding DNA sequences (CDSs). A total of 792 472 SNVs were uniquely found in the Coari data set, while 654 619 were identified only in the samples collected in Porto Velho. The SNVs identified in this study, though requiring validation, serve as the basis for high-throughput genotyping analysis and future population genetic and association mapping efforts.

Table 3. Number and density of SNVs per genomic feature

<table>
<thead>
<tr>
<th>Genomic feature</th>
<th>Genome (454)a</th>
<th>Transcriptome (454 + Illumina)b</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gene</td>
<td>1 643 685 (39.7 per kb)</td>
<td>819 427 (19.8 per kb)</td>
</tr>
<tr>
<td>Exon</td>
<td>488 652 (26.2 per kb)</td>
<td>494 539 (26.6 per kb)</td>
</tr>
<tr>
<td>Intron</td>
<td>1 155 083 (50.7 per kb)</td>
<td>324 926 (14.2 per kb)</td>
</tr>
<tr>
<td>CDS</td>
<td>475 903 (26.1 per kb)</td>
<td>481 588 (26.37 per kb)</td>
</tr>
<tr>
<td>Intergenic</td>
<td>6 811 677 (50.0 per kb)</td>
<td>835 447 (6.1 per kb)</td>
</tr>
<tr>
<td>Promoter</td>
<td>360 607 (41.8 per kb)</td>
<td>153 431 (17.8 per kb)</td>
</tr>
</tbody>
</table>

aGenome data from mosquitoes collected in Coari.
bTranscriptome data from Porto Velho and Coari were combined. Samples were sequenced by either 454 Life Science (454) or Illumina technologies.

Promoter = 2 kb upstream from transcript 5’-end.

Figure 3. Synteny clusters statistics. (A) Distribution of the number of genes per synteny cluster when considering A. darlingi (Ad) versus either A. gambiae (Ag), A. aegypti (Aa), C. quinquefasciatus (Cq) or D. melanogaster (Dm). Data points represent synteny clusters with more than three protein-coding genes on each pairwise comparison. The points were scattered in each column for the purpose of facilitating visualization. Red horizontal lines indicate the media values of the distribution. (B) The total number of synteny genes between each pair of species. (C) Number of synteny clusters identified on each comparison. The whole extent of the bars indicates the total number of clusters that were identified in each analysis, which was further divided into clusters located internally on scaffolds or chromosomes versus those near chromosomes or scaffold ends. Species names were abbreviated, as in panel A.

Transposable elements

TEs correspond to 2.3% of the A. darlingi genome (Table 4) (see Supplementary Data and Method SE and SF). The set of Class I and II TEs superfamilies is as diverse in A. darlingi as in the genomes of other mosquitoes; however, the number of TE copies is smaller in A. darlingi. In A. gambiae, TEs encompass 17% of the genome (9), and among the genomes of the
Drosophila species so far analyzed, TE compositions vary from 2.7 to 23% (32).

Some of the TEs found in the A. darlingi genome showed multiple identical copies and intact transposase Open reading frames (ORFs), suggesting that they are active elements. Among the putatively active TEs are the following: gypsy-like from long terminal repeats (LTR) elements order; jockey-like, Chicken repeat 1 (CR1) and retrotransposable element (RTE) families from non-LTR order; and mariner-like and Helitrons from DNA class II elements (see Supplementary Data SE and Supplementary Figures SE1 and Supplementary Table SF). Multiple applications of active TEs have been contemplated for advancing the understanding of mosquito biology as well as for genetic-based vector control strategies. 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 (33).

Protein coding genes

A total of 10 481 protein-coding genes were predicted in the A. darlingi genome. For checking the completeness of the A. darlingi gene set, the core eukaryotic gene-mapping approach (CEGMA) (34) that assess genome completeness and gene structure prediction was applied. CEGMA analysis includes a set of core genes that are supposed to be highly conserved and single-copy genes present in all eukaryotes. The integral sequences of 235 out of 248 highly conserved eukaryotic genes (94.76%) were identified in the A. darlingi genome. Other eight highly conserved genes were found as partial loci. Despite these results indicating the efficiency of the gene prediction tools used, additional A. darlingi protein coding genes are expected to be identified as future sequencing and assembling efforts will close the present gaps between scaffolds and contigs. From the A. darlingi protein coding genes, 72.3% have the closest counterpart in the A. gambiae genome and 21.3% have a gene that has the highest similarity within the genomes of other mosquitoes (A. aegypti or C. quinquefasciatus) (35). A comparative analysis of the functional categories of the genes comprising the A. darlingi and A. gambiae genomes showed that, in general, functional categories were equally represented (Figure 4). Genes associated with hematophagy

Table 4. Transposable contents in mosquito genomes

<table>
<thead>
<tr>
<th>TE class—Order</th>
<th>A. gambiae Copy number</th>
<th>A. gambiae % of genome</th>
<th>A. darlingi Copy number</th>
<th>A. darlingi % of genome</th>
<th>A. aegypti Copy number</th>
<th>A. aegypti % of genome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Class I—LTR</td>
<td>4348</td>
<td>6.2</td>
<td>241</td>
<td>0.19</td>
<td>28905</td>
<td>10.51</td>
</tr>
<tr>
<td>Class I—Non LTR</td>
<td>392</td>
<td>1.07</td>
<td>200</td>
<td>0.9</td>
<td>61938</td>
<td>14.37</td>
</tr>
<tr>
<td>Class I—SINEs</td>
<td>2389</td>
<td>3.77</td>
<td>4610</td>
<td>0.51</td>
<td>101838</td>
<td>1.88</td>
</tr>
<tr>
<td>Class I—DNN transposons</td>
<td>835</td>
<td>1.1</td>
<td>395</td>
<td>0.02</td>
<td>12930</td>
<td>3.04</td>
</tr>
<tr>
<td>Class I—Helitrons</td>
<td>5</td>
<td>0.2</td>
<td>19</td>
<td>0.02</td>
<td>244</td>
<td>1.04</td>
</tr>
<tr>
<td>Class I—MITEs</td>
<td>3399</td>
<td>5.07</td>
<td>6635</td>
<td>0.66</td>
<td>419955</td>
<td>15.8</td>
</tr>
<tr>
<td>Total</td>
<td>11 368</td>
<td>17.41</td>
<td>12 119</td>
<td>2.29</td>
<td>625810</td>
<td>46.64</td>
</tr>
</tbody>
</table>

TEs were classified as proposed by (33): Class I retrotransposon, with LTR, retroposons without LTR or SINEs (short interspersed nuclear elements). Class II were classified as DNA transposons, helitrons and MITEs (miniature inverted-repeat TEs). A. gambiae and A. aegypti data from (8,9,34).

Figure 4. Distribution and functional categories of protein-coding genes predicted in Anopheles species. The best matches distribution of all (10 481) of the A. darlingi predicted protein coding genes in the KEGG database, by organisms; and the comparison of the molecular functions of the products of the predicted protein coding genes between A. darlingi and A. gambiae.
(encoding components of mosquito olfaction and saliva), immunity and insecticide resistance are directly involved in vector-human and vector-parasite interactions and efforts to curb malaria transmission. Some of these genes, identified in the A. darlingi genome, will be further discussed.

**Olfaction**

The chemosensory system plays essential roles in food source or host location, mate choice, predator avoidance, oviposition site selection and toxic-compound avoidance (35). Molecular components of insect chemosensory systems include at least three different types of chemosensory receptors: the odorant (OR), the gustatory (GR) and the ionotropic (IR) receptors (36). Two other types of proteins, i.e. the odorant-binding proteins (OBPs) and chemosensory proteins (CSPs), are involved in the perireceptor events of the chemosensory system (36,37).

**Odorant receptors**

In A. gambiae, a family of 79 putative odorant receptor (AgOR) genes have been identified (38,39), including AgamGPRor7, now named AgamOrco (40), and the ortholog of D. melanogaster DmelOr83b, which serves as a coreceptor in all OR multimeric complexes (41). In the A. darlingi genome, we have identified 18 genes that encode putative ORs, including a gene encoding AdarOrco (GPROR7) (see Supplementary Table SH1). It appears that the number of OR paralogs is reduced in A. darlingi; OR3, which in A. gambiae is part of a group of 25 paralogs, is represented in A. darlingi by seven paralogous genes; OR33 is represented by six paralogs in A. gambiae and by four in A. darlingi. Six ORs (OR8, OR23, OR34, OR39, OR42 and OR58) are represented by single genes in A. darlingi. OR23 and OR42 are, respectively, represented by 15 and 14 paralogs in A. gambiae.

**Gustatory receptors**

Sixty-one genes encoding putative GR have been identified in the A. gambiae genome. In the A. darlingi genome, 17 GR genes were identified (see Supplementary Table SH1), three of them (AD01104, AD08863 and AD09819) as partial sequences. Among them, four genes (AD07140/GPRGR14, AD08836/GPRGR15, AD08857/GPRGR17 and AD08840/GPRGR20) encode receptors that were described as candidate sugar receptors in A. gambiae (42). The proteins encoded by the genes AD09907, AD01029 and AD09985 correspond to the receptors GPRGR22, GPRGR23 and GPRGR24, respectively, and show a high conservation (71–93%) when compared with homologous sequences in A. gambiae, A. aegypti and C. quinquefasciatus. The corresponding orthologs of GPRGR22 and GPRGR24 in D. melanogaster (DmGr21a and DmGr63A) function as a heterodimeric receptor for carbon dioxide (43,44).

**Variant ionotropic glutamate receptors**

These receptors function as chemosensory receptors in D. melanogaster (45) and A. gambiae (46,47). In A. gambiae, a family of 46 variant ionotropic glutamate receptors was identified (47). In A. darlingi, we found 14 sequences related to variant ionotropic glutamate receptors (see Supplementary Table SH1).

**Odorant binding proteins**

A total of 69 genes encoding OBP were described in A. gambiae; many of them possibly originated from recent events of gene duplications. We have found 33 OBP encoding genes (see Supplementary Table SH1) in the present A. darlingi genome assembly. The reduced number of OBP genes suggests that duplication events were not as frequent in this species. Alternatively, the missing genes may be located in unassembled portions of the genome. In fact, besides the OBP genes annotated, TBLASTN searches identified sequences that likely correspond to truncated OBP-like genes. Sequences with similarity to 10 A. gambiae OBP s could not be identified in any of the A. darlingi contigs.

The genes AD02966 (OBP34), AD00512 (OBP37), AD01405 (OBP44) and AD01406 form part of a group of paralogs that in A. gambiae is composed of 16 genes. However, in other cases, the number of related sequences is similar in both species, i.e. AD04156 (OBP10), AD03416 (OBP18), AD07879 (OBP21), AD07746 (OBP25), AD03881 (OBP26), AD03880 (OBP28), AD06986 (OBP23) and AD03882 (AGAP012322), which in A. gambiae is also represented by eight sequences. The amino acid sequences of OBP34 (AD02966) and OBP37 (AD00512) are highly similar, with only three amino acid changes. In A. gambiae, OBP 34 and 37 present identical amino acid sequences (48).

**Chemosensory proteins**

Belonging to a class of soluble proteins that are found in the sensillum lymph of insect antennae, CSP exhibit binding activity toward odorants (49). CSP encoding genes have been identified in several insects, and among the mosquitoes, 21 genes were described in C. quinquefasciatus (50) and 8 in A. gambiae (51). Six of the CSP genes (AgamCSP1 to AgamCSP6) described in A. gambiae are part of a group of paralogs. In the A. darlingi genome, we identified four CSP genes (see Supplementary Table SH1), and all presented similarity to representatives of this paralogous group.

**Salivary proteins**

The salivary gland (SG) is the only organ of A. darlingi that has been submitted to a tissue-specific transcriptome analysis (52,53). A total of 2371 clones from an adult female A. darlingi SG cDNA library were sequenced and assembled, allowing the identification of 183 protein sequences, 114 of which code for putatively secreted salivary proteins. A comparative analysis of SG transcriptomes of A. darlingi and A. gambiae reveals a significant divergence of salivary proteins. On average, salivary proteins are only 53% identical, while housekeeping proteins are 86% identical between the two species. A. darlingi proteins were found that match culicine but not anopheline proteins, indicating a loss or rapid evolution of these proteins in the old world Cellia subgenus. Additionally, several
well-represented salivary protein families in old-world anophelines are not expressed in *A. darlingi*.

**Circadian rhythm**

Rhythmic cycles of *Anopheles* mosquitoes command biting activity, mating swarms, nocturnal flight activity and egg laying; however, little work has been performed to elucidate the molecular basis for these daily rhythms (54). Throughout its geographical distribution, *A. darlingi* exhibits distinct patterns of biting behavior. One, two or three daily peaks of biting activity have been observed in different studied sites (55–58). The molecular basis for these differences in behavior is unknown. Here, we describe the *A. darlingi* circadian cycle-associated genes timeless, cycle, clock, timeout and period (see Supplementary Method and Data SI and Supplementary Figures SI1 and SI2). The identification of these genes will permit assessment of their expression levels and rhythmicity among the diverse *A. darlingi* populations.

**Insecticide resistance**

Resistance to insecticides is a major threat to sustained reductions in malaria vector populations and malaria incidence. To date, there has been only a single report of insecticide resistance in natural *A. darlingi* populations. A population from Colombia was found to be resistant to both dichlorodiphenyltrichloroethane (DDT) and lambda-cyhalothrin (59). However, a number of studies reporting insecticide resistance in the African malaria vector *A. gambiae* as well as other vector mosquitoes should caution against complacency (60–64). The changing pattern of land use in the Amazonian region, resulting in increased urbanization and agricultural initiatives, and the associated escalation in insecticide use are expected to strengthen selection for insecticide resistance in *A. darlingi*.

**Metabolic detoxification**

Three gene families that are primarily involved in insecticide metabolism have been described: the cytochrome P450s (P450s), the carboxy/cholinesterases (CCEs) and the glutathione-S-transferases (GSTs) (65). Metabolic resistance is usually a result of overexpression or allelic variation in members of detoxifying enzyme families. We identified 89 P450s, 20 CCEs and 30 GSTs genes in *A. darlingi* (see Supplementary Table SJ1). GSTs are the most conserved among the three superfamilies (66), and this conservation permitted the identification of putative orthologs between *A. darlingi* and *A. gambiae* that had a sequence identity that was >70%. Four classes of cytoolic GSTs were identified: the most conserved theta (five genes), zeta (one gene), the insect-specific delta (three genes) and epsilon (six genes) classes. Only members from the Delta and Epsilon classes have been implicated in insecticide resistance. Among the epsilon members in *A. darlingi*, GSTe2 (AdGSTe2, AD08205) is highly conserved among culicines (*A. gambiae*, *A. aegypti* and *C. quinquefasciatus*) and metabolizes DDT in *A. gambiae* and *A. aegypti* (67,68). Several AdGST genes remained unclassified, with no obvious orthologs in the *A. gambiae* genome, and thus, they might represent novel GSTs.

The CCEs and P450s appear to have undergone a slight expansion in *A. gambiae* in comparison with *A. darlingi*. It is possible, considering the redundancy in these families, that different family members are co-opted for functions in insecticide resistance in different mosquito populations, such as P450s and some GSTs that have increased mRNA accumulation in some, but not all, *A. gambiae* insecticide-resistant populations (60–64). Additionally, genes encoding a superoxide dismutase (AY745234) and a peroxiredoxin (XP_308081.2) also presented increased mRNA accumulation in these populations.

**Target-site insensitivity**

Decreased target site sensitivity to pyrethroids and DDT in *A. gambiae* has been described as being associated with two alternative substitutions at a single codon in the sodium channel gene (L1014F or L1014S) and is referred to as knockdown resistance, or kdr (69–72). A comparison of the voltage-gated sodium channel (VGSC) gene sequence across different insect species showed that it is highly conserved, but different numbers of exons are observed among species (73). In *A. gambiae*, 33 exons have been identified, which can synthesize different mRNAs through alternative splicing. Two putative VGSC genes were identified in the *A. darlingi* genome [AD07884 (2e-75; 98% identity) and AD00168 (3e-38; 45% identity)]. Primers based on the *A. gambiae* sodium channel sequence had previously failed to amplify the *A. darlingi* ortholog (59,69). The now available *A. darlingi* VGSC sequences permit the development of specific diagnostic tools for detecting kdr resistance in this species.

Target-site resistance to carbamates and, to a lesser extent, organophosphates (OP) in culicines result from a mutation in the acetylcholinesterase gene (*ace-1*). This gene is absent in *Drosophila*, possibly because of a secondary loss, and OP resistance in this organism arises from mutations in the *ace-2* gene, which is ubiquitous in insects. The putative *A. darlingi* ace-1 homolog is AD03377 (4e-38; 98% identity when compared with *Anopheles albimanus*) (74). In *A. gambiae*, a second copy of *ace-1* (ace-1D) has been described, and its high frequency and distribution in countries of West Africa points to an association with resistance (75). The availability of *A. darlingi* ace-1, VGSC and other detoxifying gene sequences allow the development of specific diagnostic tools for detecting incipient insecticide resistance in this species. This is especially important in epidemiological vigilance because evolutionary forces acting on *A. darlingi*, when facing continuous and increasing exposure to insecticides, could lead to widespread insecticide resistance.

**Immunity-related genes**

The mosquito immune system plays a critical role in limiting the spread of malaria and other vector-borne diseases. We analyzed sequences related to the three major immune response systems in Diptera, Toll, immune deficiency (IMD) and thioester proteins (TEPs) (see Supplementary Table SK1) because these genes and
their associated signaling pathways are known to limit the spread of malaria parasites in anophelines. Identifying the *A. darlingi* orthologous genes relative to each component of the *D. melanogaster* and/or *A. gambiae* pathways is challenging, especially where multigenic families such as Toll receptors or the TEPs are involved. In contrast, one to one orthologs of most of the signaling molecules were more easily identified. Although the *A. darlingi* immune system appears to be organized similar to those of other Diptera, exact orthologs of many of the important receptors have not yet been established. The presently assigned putative homologous functions must be asserted by actual bench experiments to gain a full appreciation of *A. darlingi* immunity.

**Toll pathway**

We identified four *A. darlingi* genes that are related to the Toll ligand known as spätzle (SPZ), when six SPZs were found in both *A. gambiae* and *D. melanogaster* (see Supplementary Figure SK1) (76). Two of these genes are possible orthologs of the SPZ1 group, which include *Drosophila* spätzle, the ligand for Toll. The other two are orthologous to SPZ3 or SPZ6. *Drosophila melanogaster* has nine Tolls; only Toll and Toll7 have established immune functions, while the functions of the *A. gambiae* Tolls are still largely undefined. Clear orthologs to the fruit fly genes could not be identified for most of the seven *A. darlingi* Tolls that were identified, although a Toll7 ortholog was assigned. Conversely, 1:1 orthologs were found for nearly all of the known signaling molecules in the Toll pathway, including MyD88, Tube, Pelle, TRAF6 and the NF-κB/IκB orthologs Rel1/Cactus.

**Peptidoglycan recognition proteins and the Immune deficiency pathway**

Eight peptidoglycan recognition proteins (PGRPs) were identified in the *A. darlingi* genome, three of which are likely to be catalytic type 2 amidases. PGRP-LC, a well-established receptor for DAP-type peptidoglycan and activation of the IMD pathway in fruit flies, appears to have two orthologs in *A. darlingi*. Additional orthologs of known PGRPs were identified, although only peptidoglycan recognition protein-LB 5′-untranslated region (PGRP-LB) has an established function, which is involved in degradation of PGN, a non-catalytic PGRP. Ten additional IMD pathway members (including the negative regulator CASPAR and essential signaling components such as IMD, (Fas-associated death domain containing protein - FADD and Death-related ced-3/ Nedd2-like protein - DREDD) FADD and DREDD) were found on a 1:1 orthology basis.

**Thioester proteins**

TEPs play a role in Diptera that is similar to the role of complement in humans: they directly opsonize bacteria and parasites, which leads to death and melanization. Ten possible TEPs were identified in *A. darlingi*. The *A. gambiae* TEP1 gene product has been proposed as a key regulator of malaria infection. A definite ortholog of TEP1 was not identified in *A. darlingi*, although several of the *A. darlingi* TEPs are in the subfamily in which TEP1 is included.

**Antimicrobial peptides**

*Drosophila melanogaster* has, at a minimum, seven families of antimicrobial peptides. Similar to other mosquito species, most of these antimicrobial peptides were not readily apparent in the *A. darlingi* genome. However, genes encoding two well-known classes of antimicrobial peptides that are found in the genome of other mosquitoes were identified in *A. darlingi*: one member of the Defensin family and three Cecropins.

**CONCLUSIONS**

Malaria was once epidemic in most areas in Central and South America (7,77,78). Economic development and the associated environmental changes that have occurred during the 20th century have drastically reduced malaria transmission in subtropical areas. However, malaria is still a major public health problem in the Amazon basin, where >500 thousand malaria cases occur every year. Because *A. darlingi* is the main malaria vector in the Amazon, and also for its interesting phylogenetic position, the Brazilian National Council for Research included this species among those selected as priorities for having their genomes sequenced (79). Here, we present the *A. darlingi* genome as a valuable platform for basic and applied sciences.

Laboratory colonization of *A. darlingi* has proven to be difficult, and presently there are no available autonomous colonies of this species. Nonetheless, large numbers of wild *A. darlingi* mosquitoes are easily captured in the Amazon, and raising the progeny of captured gravid females has allowed the sequencing of the mosquitoes genome and transcriptome, which complements studies of *A. darlingi* biology, behavior, physiology, genetics, biochemistry and insecticide resistance (4,11,23,22,53,80–85). The successful colonization of other neotropical anopheline species (86,87) and older reports of *A. darlingi* that were successfully adapted to breed in laboratory conditions (88–90) indicate that colonizing *A. darlingi* is an attainable task. The availability of this genome will promote efforts to establish an autonomous viable free-mating laboratory *A. darlingi* colony.

As the first neotropical Anopheles species of the subgenus *Nyssorhynchus* with its genome sequenced and annotated, the data presented here open a new window from which we can contemplate the evolutionary history of these mosquitoes. Comparative evolutionary genomics is one of the most rapidly advancing disciplines in the biological sciences and offers the opportunity to study evolutionary changes among organisms, to identify genes that are conserved among species, and to study the genes that give each organism its own specific characteristics (91). Questions that are related to malaria vectorial capacity, anthropophily and hematophagy among anophelines can now be addressed from the perspectives of two distinctly related members of the Anopheles genus that diverged ~100 mya and evolved in two distinct
environments (11). *Anopheles darlingi* orthologs of genes associated with insecticide resistance have been identified, allowing a more targeted examination of insecticide resistance status in populations of this vector species (60). A catalog of *A. darlingi* immunity-related genes will help in studies of vector–parasite interactions and will promote research to understand the determinants of vectorial capacity and competence (92). Finally, we identified 349 *A. darlingi* predicted genes that encode products with no hit in the Kyoto Encyclopedia of Genes and Genomes (KEGG) database (see Supplementary Table SL1), thus potentially related to adaptations to the New World environment. This study and other recently published and ongoing efforts to sequence the genomes and transcriptomes of malaria vectors (93,94) (vectorbase.org) will provide a needed and more complete understanding of malaria vector biology.

It is our hope that this report provides valuable information that will lead to novel strategies to reduce the rate of malaria transmission on the South American continent.

**ACCESSION NUMBERS**

The sequence of *A. darlingi* has been deposited in the DDBJ/EMBL/GenBank database under the following accession number: ADMH00000000. The version described in this paper is the second version, ADMH02000000.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online: Supplementary Tables SA1-SA5, SF, SJ1, SK1, SH1, SL1, Supplementary Figures SB1, SC1, SD1, SE1, SI1, S12, SK1, Supplementary Methods SA, SB, SE, SI, Supplementary Data SC, SE, SG, SI and Supplementary References [96–104].

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