Salivary Polytene Chromosome Map of Anopheles darlingi, the Main Vector of Neotropical Malaria

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Abstract. A new photomap of Anopheles (Nyssorhynchus) darlingi Root, 1926, is described for a population from Guajará-Mirim, State of Rondônia, Brazil. The number of sections in the previous A. darlingi reference map was maintained and new subsections were added to the five chromosome arms. Breakage points of paracentric inversions had been previously incorporated into the photomap of this species. An additional inversion is reported, called 3Lc, totaling 14 inversions in the A. darlingi chromosome arms. The proposed photomap is potentially useful for further evolutionary studies in addition to physical and in silico chromosome mapping using A. darlingi genomic and transcriptome sequences. Furthermore, in our attempt to compare sections of the 2R chromosome arm of A. darlingi with Anopheles funestus, A. stephensi, and A. gambiae, we found great differences in the arrangement of the polyploid chromosome bands, which are consistent with the known phylogenetic divergence of these species.

INTRODUCTION

Anopheles darlingi Root, 1926, has historically been considered the primary vector of malaria and the most anthropophilic and endophilic Anopheles species in the Americas.1 In the Brazilian Amazon Basin, this mosquito is the most important vector of human malaria.2 Anopheles darlingi has been found in clear water such as lakes, ponds, and streams, and in smaller water courses such as artificial deposits and puddles.3,4 The relevance of A. darlingi as vector of malaria, its isoenzymes, behavior, genetic structure, biology, metaplace karyotype, mapping, and chromosome polymorphism were demonstrated in several studies.5–14

In South America, 17 Anopheles species are known as important vectors of malaria. Of those, more than two-thirds belong to the subgenus Nyssorhynchus, which contains 29 named species throughout the Neotropics.15 They are of primary importance for transmission of human malaria and include the A. darlingi. Several areas in Amazonia have shown significant differences in A. darlingi incidence.2 At the outer edge of the Aripuanãs locality in Rondônia State, in Brazil, the registered index of adult Anopheles species has been 30.6 mosquitoes per person per hour, and A. darlingi is abundant. This mosquito has been observed almost exclusively at the modified sites, especially around and into the houses, a fact that confirms its high degree of association with human dwellings.16

Karyotypes of the studied Anopheles species are relatively conserved. Morphological data obtained from metaphase karyotypes of six chromosomes in brain cells of A. darlingi describe a pair of metacentric (chromosome 2) and a pair of submetacentric (chromosome 3) autosomes, and sex chromosomes (female XX; male XY). The X chromosome is acrocentric in females and the Y chromosome is punctiform in males.11 This pattern is observed in the majority of the species in the Nyssorhynchus subgenus, including Anopheles strodei, A. albimanus, A. aquasalis, A. noroestensis, and A. argyritarsis.17 Morphological variation was detected in the X chromosome, which was metacentric in A. aquasalis and A. noroestensis, A. strodei, A. albimanus, acrocentric in A. argyritarsis,17,18 A. darlingi,11 and A. albitalis sensu latu,19 whereas it is metacentric in A. aquasalis and A. noroestensis,17 and submetacentric in A. nuneztovari.13 This pattern is observed also in Anopheles cruzii and Anopheles bellator of subgenus Kerteszia,20,21 A. subpictus,22 and A. gambiae species complex, serie Pyretophorus, subgenus Cellia.23

The morphological variation of the sexual chromosome pair suggest selective pressure in those anopheline species, which is apparently occurring only at a genetic level. In the larval salivary gland cells, the well-developed and well-banded polyploid chromosome X, 2 and 3 represent this diploid set.

Currently, a new photomap description of A. darlingi polyploid chromosomes in substitution of the drawn reference map2 would be highly helpful. The first study on polymorphism of inversions of A. darlingi polyploid chromosomes was reported.3 The authors described nine independent inversions and a complex array of references on a drawn map of polyploid chromosomes of populations from the regions of Manaus and Itacoatiara, in the State of Amazonas, Northern Brazil. These populations were highly polymorphic compared with a studied A. darlingi population from the Southeast of the country, in the region of Araquara, in the State of São Paulo. Subsequently, the previously mentioned inversions and described three new inversions, one on chromosome 2 (2Rd), one on chromosome 3 (3Rc), and one on chromosome X (Xb), all belonging to the A. darlingi population from the vicinities of the BR-174 highway, Manaus-Boa Vista, in the State of Amazonas were analyzed.6,7 In three populations of this same region, Tadei and Santos10 conducted a comprehensive study on frequency variation in most of the chromosome inversions. These populations were characterized as highly polymorphic, with up to 60% of individuals presenting three to five inversions, depending on what time of the year the specimens were sampled. High frequencies of inversions in the heterozygous state were also found in this same population.

The major advantage of the A. darlingi photomap is to better define chromosome regions and subsections, allowing structural analysis by means of traditional methods and computer resources. It will be a useful tool in identifying the large-scale physical mapping using probes of expressed sequence tags (EST) from the A. darlingi transcriptome (Rafael MS and others, unpublished data). The polyploid

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chromosome photomap will also offer a better understanding of the anchoring of the complete genome of *A. darlingi* for the *Drosophila* 11 Genomes.24

The objective of this work was to construct a polytene chromosome photomap for *A. darlingi* based on a population with low chromosomal polymorphism from the Northern region of the Amazon, aiming at contributing to the understanding of the *A. darlingi* genome structure and organization, and of the evolutionary diversification and chromosomal variability of this important vector of Neotropical malaria.

**MATERIALS AND METHODS**

**Sampling and identification of mosquitoes.** *Anopheles darlingi* females were sampled in August 1999, in the region of Guajará-Mirim (10°47′0″S, 65°21′0″W), in the State of Rondonia, Brazil. They were captured when resting on stable walls or biting humans in inner houses, between 6:30 pm and 9:00 pm. The Guajará-Mirim population was chosen because of its low chromosomal polymorphism, which was inferred from rearrangement patterns. Wild-captured individuals were transported in moist chambers to the Laboratory of Vectors of Malaria at the National Institute for Amazonian Research (INPA) in Manaus. Morphological identification of specimens was carried out according to the taxonomic keys.25, 26 The sampled females were confined individually in plastic cups for egg laying. Offsprings were fed with finely ground liver and fish powder, and reared to the fourth instar larvae.

**Chromosome preparation.** Salivary glands of fourth stage *A. darlingi* larvae were dissected and used to prepare chromosome slides.12, 13, 27 Preparations of well-spread polytene chromosomes on 11 slides were photographed using a light microscope (Carl Zeiss Axioplan, Ulm, Germany), for further analysis and construction of a photomap.

**Construction of the photomap and description of inversions.** The photomap of the *A. darlingi* analyzed population was constructed using carefully selected photos of different chromosomes considered representative of the banding pattern. The best images were captured with the aid of a scanner and processed using Adobe Photoshop (Adobe, 2002 7.0. v 701; Adobe Systems Inc., San Jose, CA) and Corel Draw (Corel-DRAW 10 2000, 10.410, Corel Corp., Ottawa, Canada). All chromosomes were divided into sections represented by numbers and into subsections represented by capital letters, starting with A in the distal region. Chromosomes were divided into 45 sections, but the previous name of each paracentric inversion was changed following the system.8 Thus, the diverse arrangements of heterozygous inversions in the chromosome X and autosomes 2 (chromosomal arms 2R and 2L) and 3 (chromosomal arms 3R and 3L) were identified by lower case letters, in alphabetical order according to their previous descriptions. Wherever possible, the correspondence was maintained between our descriptions and the earlier studies.5–7 The former authors studied a highly polymorphic *A. darlingi* population from the Central region of the Amazon and described nine independent inversions and a complex chromosome arrangement. Tadei and others’ studied populations from the vicinities of the BR-174 highway, between Manaus and Boa Vista, in the State of Amazonas and described two previously unknown independent inversions, increasing to 12 the number of chromosome inversions.

A comparative analysis of chromosome arm tips, except of the X chromosome, was done for the major malaria vectors *Anopheles gambiae* Patton, *Anopheles stephensi* Liston, *Anopheles funestus* Giles, and *Anopheles darlingi*. The *Anopheles gambiae* and *A. funestus* species are the major malaria vectors in Africa. *A. stephensi* is the major vector in Indo-Pakistan and Middle East, and *A. darlingi* is the primary vector of Netropical malaria. The species *Anopheles gambiae*, *A. stephensi*, and *A. funestus* belong to the subgenus *Cellia* and *A. darlingi* belongs to the subgenus *Nyssorhynchus*.17, 28, 29 Homologies of bands in the chromosome arm tips of *A. darlingi* was compared with the photomaps previously described for *A. gambiae*,30 *A. stephensi*,31 and *A. funestus*.3

**RESULTS**

A photomap of polytene chromosomes from salivary glands of *A. darlingi* was developed for a low polymorphic population
from the region of Guajará-Mirim, in the Northern Brazilian Amazon (Figure 1). The photomap is oriented so that the centromere of each chromosome is positioned on the right and the chromosome distal region is on the left. The photomap was divided into 45 sections, following the original draw reference map. Sections were numbered from the starting point of the distal part of the X chromosome (section 1) ending at the base of the distal part chromosome 3L (section 45). Correspondence of the sequences followed, to the possible extent, the division proposed in the draw reference map. Alternatively, photos were used in this work to visualize the nature of the banding pattern of chromosomes and their sizes. Thus, new subsections were included in several of the sections, so that each subsection includes approximately 3 to 4 bands of similar size.

New divisions were added, as sections 2 and 4 of the X chromosome, subdivided into three sections (Table 1). In the chromosome arm 2R, sections 6, 7, 8, 9, 10, 12, 13, and 15 were subdivided into 4, 3, 3, 3, 3, 3, 3, and 5 subsections, respectively. In the chromosome arm 2L, sections 16, 17, 19, 20, 21, 22, 23, and 24 were subdivided into 3, 3, 3, 3, 3, 3, 3, and 3 subsections, respectively. Chromosome arm 3R shows the sections 26, 27, 28, 29, 30, 33, and 34, which were subdivided into 3, 3, 3, 3, 3, 3, and 3 subsections, respectively. In the chromosome arm 3L, sections 36, 37, 38, 39, 40, 41, 42, 43, and 44 were subdivided into 4, 4, 4, 4, 4, 4, 4, and 4 subsections, respectively. Other sections in which not all chromosome arms were modified, the number of subdivisions was maintained according to the reference map. The final result was a photomap with well-distributed sections, subsections, and total size of all chromosomes together.

Table 1. New divisions for polytene chromosome photomap of Anopheles darlingi of Guajara-Mirim locality

<table>
<thead>
<tr>
<th>Chromosome X</th>
<th>Right arm (2R)</th>
<th>Left arm (2L)</th>
<th>Chromosome 2</th>
<th>Right arm (3R)</th>
<th>Left arm (3L)</th>
<th>Chromosome 3</th>
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<tbody>
<tr>
<td>Sections</td>
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</tr>
<tr>
<td>2</td>
<td>A, B, C</td>
<td>6</td>
<td>A, B, C, D</td>
<td>16</td>
<td>A, B, C, D</td>
<td>26</td>
</tr>
<tr>
<td>4</td>
<td>A, B, C</td>
<td>7</td>
<td>A, B, C</td>
<td>17</td>
<td>A, B, C</td>
<td>27</td>
</tr>
<tr>
<td></td>
<td></td>
<td>8</td>
<td>A, B, C, D</td>
<td>18</td>
<td>A, B, C, D</td>
<td>28</td>
</tr>
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<td></td>
<td></td>
<td>9</td>
<td>A, B, C, D, E</td>
<td>19</td>
<td>A, B, C, D</td>
<td>29</td>
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<td></td>
<td></td>
<td>10</td>
<td>A, B, C, D, E</td>
<td>20</td>
<td>A, B, C</td>
<td>30</td>
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<td></td>
<td></td>
<td>12</td>
<td>A, B, C, D, E</td>
<td>21</td>
<td>A, B, C, D</td>
<td>33</td>
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<td>24</td>
<td>A, B, C</td>
<td>36</td>
<td>A, B, C, D</td>
<td>38</td>
</tr>
</tbody>
</table>

The relative size of each chromosome arm in the photomap was carefully estimated from individual measurements with a digital caliper. Several nuclei were measured and only photomicrographs displaying all chromosomes together were chosen, to minimize experimental errors. It can be observed in the photomap of A. darlingi (Figure 1), the X chromosome is 10% of the total size of all chromosomes together. Chromosome 2 corresponds to 49% (2R = 28% and 2L = 21%) and chromosome 3 corresponds to 41% (3R = 18% and 3L = 23%). These results agree with Kreutzer and others who reported the approximate proportions of 10% for each of the X chromosomes, 52% for chromosome 2 (2L = 30% and 2R = 22%), and 40% for chromosome 3 (3R = 20% and 3L = 20%).

In general, all chromosomes are quite stable regarding polymorphism and presence of puffs and constrictions. However, the X chromosome was an exception in the analyzed population. Its variation in levels of distension and compression, especially in sections 1A and between 5A and 5C, is shown in Figure 2. Figure 3 shows the X chromosome pattern and the breakpoints of the heterozygous inversion Xa.

The photomap of chromosome arm 2R is shown in Figure 4, and the chromosome arms 2L is shown in Figure 5. The probable breakpoints of the inversions 2Ra (between sections 7B and 8C), 2Rb (between 9D and 10A), and 2Rc (between 11A and 12E) are indicated in Figure 4. They were first described by Kreutzer and others and later by Tadei and others, who also...
described a new reversal in the 2Rd (from 12E to 14B). Figure
5 shows the photomap of the chromosome arm 2L, probable
breakpoints of the heterozygous inversion 2La (between sections
20A and 23C), and a complex of inversions 2La + b, with
breakpoints between sections 23C and 21B. The latter authors
also described an inversion homozygote 2Lb present only in a
population from the region of Araraquara, in the State of São
Paulo.

Figures 6 and 7 show, respectively, the photomap of chromo-
some arms 3R and 3L. For the inversion in chromosome arm
3R, Figure 6 shows the probable breakpoints of the heterozy-
gous inversions 3Ra (between sections 28D and 32C) and 3Rc
(between sections 34A and 35A), and the inversions 3Rb
(between sections 35A and 35C) and 3Rd (between the sec-
tions 31A, 28B). The probable breakpoints of the heterozy-
gous inversions 3La (between sections 43C and 42C) and 3Lb
(between 40A and 37A) and 3Rd are shown in Figure 7.
A new inversion 3La in A. darlingi, with breakpoints between
sections 42B and 37D, is presented here, for the first time,
occuring in the Guajará-Mirim population. This inversion
is the largest of all the investigations already described for
A. darlingi. It was found only in heterozygous state and repre-
sents approximately 57% of the length of chromosome arm 3L.
Thus, the A. darlingi chromosomes comprise at least 14 inver-
sions, with one inversion on the X chromosome, 4 inversions in
the chromosome arm 2R, 2 in 2L, 4 in 3R, and 3 in 3L.

DISCUSSION

Autosome arm tips of A. darlingi were compared with the
species A. gambiae, A. stephensi, and A. funestus. Figure 8
shows the probable chromosome homology of the 2R arm tip
between A. darlingi and A. gambiae, and of A. stephensi com-
pared with A. gambiae and A. funestus. Figure 9 shows part
of the homology in the 2R chromosome arm tip of A. gam-
biae and 2R of A. funestus. The homology in the 2R arm tip
of both species was unclear. Therefore, we suggest another
possibility of homology between A. funestus and A. gambiae,
plus A. stephensi, as presented in Figure 8, where A. stephensi
and A. gambiae are very similar in relation to the 2R arm tip.
Figures 9–15 present the probable homology between tips of
the chromosome arms 2L, 3R, and 3L, except X chromosome
of A. darlingi that was quite divergent.

A standard polytene chromosome photomap is one of
the essential requirements for cytogenetic investigation of
Dipteran insects, including the anopheline mosquitoes. The
data described herein show that A. darlingi is no exception.
The construction of a photomap of A. darlingi polytene chro-

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**Figure 4.** Photomap and arrangements of Anopheles darlingi 2R chromosome arm. Lines over the chromosome arm indicate probable break-
points of the heterozygous inversion 2Ra (between sections 7B and 8C), found in our study and first described, in addition to inversions 2Rb
(between sections 9D and 10A) and 2Re (between 11A and 12E), and inversion 2Rd (between section 12E and 14B) plus 2Rc. Grid lines rep-
resent chromosome arm continuation.

**Figure 5.** Photomap and arrangements of the Anopheles darlingi 2L chromosome arm. Lines over the chromosome arm indicate probable break-
points of the heterozygous inversion 2La (between sections 23C and 20A), and the complex inversion 2La + b (inversion 2Lb with a break-
point between sections 23C and 21B). The inversion 2Lb was also described by the latter authors as a homozygous inversion described only in the
population from Araraquara. Grid lines represent chromosome arm continuation.
mosomes was motivated by the need to map, in silico and in situ, the ESTs generated from the genome sequencing of this species (Rafael and others, unpublished data). This was triggered by the first mapping of ribosomal genes and genes of single copy (Hsp70) in polytene chromosomes of *A. darlingi* populations of Manaus and Macapá regions, in the Brazilian Amazon.

In previous studies, photomaps were described for other species as *Anopheles funestus*, *Anopheles stephensi*, *Anopheles gambiae*, *Anopheles subpictus*, *Anopheles albimanus*, *Anopheles triannulatus*, *Anopheles cruzii*, and *Anopheles belattor*. The photomaps have been useful for identifying species with high chromosome polymorphism and monomorphic species constructing maps of low resolution. Furthermore, chromosome arm translocations have been described for the important Neotropical vectors of malaria, *A. gambiae*, *A. funestus*, and *A. stephensi*, of the subgenus Cellia. In contrast, among the important Neotropical vectors of malaria, *A. darlingi* does not show homologies in the X chromosomes, 3R or 3L similarly to *A. aquasalis*, *A. albimanus*, or *A. nuneztovari*. However, there is homology in the 2R arms of *A. darlingi*, and in *A. argyritarsis*, *A. aquasalis*, *A. albimanus*, and *A. nuneztovari*. The morphologically alike species are expected to and do share more banding pattern similarity than species with low morphological similarity.

In this study, we showed probable chromosome homology between the 2R arm tips of *A. darlingi* and *A. gambiae*. The 2L arm of *A. funestus* corresponds to the 3R arms of *A. gambiae* and *A. stephensi*. The 2L arm of *A. stephensi* corresponds to the 3L arms of *A. funestus* and *A. gambiae*. The 2L arm of *A. funestus* corresponds to the 3R arms of *A. gambiae* and *A. stephensi*. In contrast, among the important Neotropical vectors of malaria, *A. darlingi* does not show homologies in the X chromosomes, 3R or 3L similarly to *A. aquasalis*, *A. albimanus*, or *A. nuneztovari*. However, there is homology in the 2R arms of *A. darlingi*, and in *A. argyritarsis*, *A. aquasalis*, *A. albimanus*, and *A. nuneztovari*. The morphologically alike species are expected to and do share more banding pattern similarity than species with low morphological similarity.

In this study, we showed probable chromosome homology between the 2R arm tips of *A. darlingi* and *A. gambiae*, and the probable homology between 2R arms of *A. funestus* and *A. stephensi*. In these four species, the 2R chromosome arm is the longest and contrasts with the X chromosome, the smallest. Despite chromosome length, the heat shock protein gene Hsp70 was mapped on the 2R chromosome arm of *A. gambiae* and *A. darlingi*. The Hsp70 gene mapped on 12A and 14A bands of the 2R chromosome arm of *A. darlingi* populations.

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**Figure 6.** Photomap and arrangements of the *Anopheles darlingi* 3R chromosome arm. Lines over the chromosome arm indicate probable breakpoints of the heterozygous inversion 3Ra (between sections 28D and 32C) and inversion 3Rc (between sections 34A and 35A), as well as inversions 3Rb (between sections 35A and 35C) and 3Rd (between sections 28B and 31A). Grid lines represent chromosome arm continuation.

**Figure 7.** Photomap and arrangements of the *Anopheles darlingi* 3L chromosome arm. Lines over the chromosome arm indicate probable breakpoints of the heterozygous inversion 3La (between sections 43C and 42C) and inversion 3Lb (between sections 40A and 37A), besides the inversion 3Lc (between sections 42B and 37D) here described as a new inversion for the *A. darlingi* species. Grid lines represent chromosome arm continuation.
in Manaus (AM) and Macapá (AP) in the Amazonian region. The Hsp70-12A and Hsp70-14A genes were useful in analysis of their fixed inversion and chromosome evolution. *Anopheles darlingi* is very divergent from *A. gambiae*, possibly because of one little inversion only in the 2R chromosome tip of *A. darlingi*. Furthermore, it was infeasible to compare all 2R chromosome arm sections of *A. darlingi* with *A. funestus*, *A. stephensi*, and *A. gambiae*, because of the great differences in the order of the polytene chromosome bands, which seems consistent with the phylogenetic divergence of these species. Despite the probable tip homology among the chromosome arms 2L, 3R, and 3L, except the X chromosome of *A. darlingi*, they were very divergent comparing *A. funestus*, *A. stephensi*, and *A. gambiae*. In addition, phylogenetic relationships between *A. gambiae*, *A. funestus*, and *A. stephensi* have been estimated by polymorphism on ribosomal DNA (rDNA) (18S and 28S) sequence data. Accordingly, *A. funestus* and *A. gambiae* are more distant species from each other than from *A. stephensi*. In this latter mosquito, genetic studies indicated chromosomal positions for the breakpoints of the 19 paracentric inversions. In addition, polymorphic inversions are not distributed evenly across the *A. stephensi* genome. More than two-thirds of them are distributed among two arms, such as five inversions are on 2R and nine on 3L. Nonrandom distribution of polymorphic inversions has been described for other species. For instance, of eight polymorphic inversions described in *A. gambiae* s.s., seven occur on chromosome 2R and 11 of 17 polymorphic inversions involve the 2R of *A. funestus*. Similar to the previous species, the *A. darlingi* complement of polytene chromosomes consists of five chromosome arms (X, 2L, 2R, 3L, and 3R). It includes one inversion on the X chromosome, four inversions in the chromosome arm 2R, two in 2L, four in 3R, and three in 3L totaling 14 inversions. Moreover, chromosome markers and morphological traits of *A. darlingi* populations revealed significant genetic differentiation among populations from Northern and Southern Brazil, and along with its geographic distribution from Southern Mexico to Northern Argentina, respectively, but *A. darlingi* consists of a single taxon. Micросatellite data suggest little genetic structure for *A. darlingi* across the Eastern, Central, and Western Amazonian regions. The latter authors described seven microsatellite loci of *A. darlingi* populations from nine localities in Central and Western Amazonian Brazil. Genetic differentiation was low and not significant among populations separated by distances less than 152 km, with exceptions for Castanho-Puraquequara (CAS-PUR), Ramal do Brasilieirinho (CAS-RBR) localities, both in the state of Amazonas, and the Nm values suggested extensive gene flow among *A. darlingi* populations.

The importance of chromosomal inversions in ecological adaptations of anopheine mosquitoes has been well established, suggesting that the differences in chromosomal rearrangements are indicators of adaptation to different ecological habitats. The data on chromosomal polymorphism of *A. darlingi* that are related to behavioral aspects may be associated to genetic structure of this species, possibly affecting their capacity to adapt to different environments. Ecological observations of several anopheine vectors of malaria in the Brazilian
Amazon describes low density and infectivity indexes, which is caused by control methods used against adult mosquitoes. However, only a few A. darlingi individuals infected with Plasmodium vivax and Plasmodium falciparum, according to enzyme-linked immunosorbent assay (ELISA) tests are needed to maintain the malaria circulating in a population. Studied parameters consisted of their breeding sites, samples distribution, incidence, feeding preferences, hour of maximum activity of adult samples, seasonality, resting places, and the presence of Plasmodium species. They support the hypothesis that seasonality, deforestation, and associated ecological alterations may contribute to the behavior, chromosome differentiation, and are conducive to A. darlingi presence, thereby they increase malaria risk.

Another important aspect of A. darlingi chromosomal polymorphism refers to the identification of populations based on the degree of heterozygosity. Comparisons of chromosomal variability among A. darlingi populations from Northern and Southern Brazil have revealed a higher frequency of heterozygous inversions, and significant genetic differentiation in the Northern populations. Salivary polytene chromosome analysis of A. darlingi have shown high frequencies of inversions in the heterozygous state and in two random regions, one near the centromere of the X chromosome and two on chromosome 2 in A. darlingi population along the Highway BR-174, Manaus-Boa Vista. In subsequent studies, populations of the central axis of the Amazon showed a number of heterozygous inversions per individual that was almost double in the marginal populations. These inversions represent 50% reduction in the values of mean inversion per individual in the heterozygous state. In addition, this work describes a new inversion, named 3Lc, which is located in chromosome arms of the A. darlingi population from Guajará-Mirim in the State of Rondonia. These high polymorphism inversions for A. darlingi populations could be a consequence of a high degree of variability that can be seen in this species, giving it sufficient adaptive plasticity to exploit the different habitat. Furthermore, its extensive geographic distribution, and in accordance with similar morphological characteristics support independent speciation processes for this mosquito.

These aspects confirm the important heterozygosity of A. darlingi populations found in Amazon region. This region is considered the hub of inhabitation in the Brazilian Amazon, as are the vicinities of the Highway BR-174, km 137, in the State of Amazonas, Highway BR-319 from Manaus to Porto Velho and Ariquemes, in the State of Rondonia. In these areas, heterozygosity of A. darlingi populations averaged between 3.20 ± 0.22 and 4.45 ± 0.34 were found. In the Marabá and Tucurui marginal populations, in the State of Pará, and in Cruzeiro do Sul, in the State of Acre, the averages ranged from 2.07 ± 0.19 to 2.19 ± 0.15. The heterozygosity averages were low in specimens from Cachoeira Porteira and Trombetas River, in the State of Pará, and in Maruanum, in the State of Amapá. The data, however, are based on the reference maps. Errors of interpretation may occur caused by the non-existence of a photomap default, or detailed pictures of heterozygous or homozygous inversions. Such limitations justified the investment on the construction of the
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