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Author(s): T. F. SILVA-DO-NASCIMENTO, R. C. WILKERSON, R.

LOURENÇO-DE-OLIVEIRA, and F. A. MONTEIRO

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Molecular Confirmation of the Specific Status of *Anopheles halophylus* (Diptera: Culicidae) and Evidence of a New Cryptic Species within *An. triannulatus* in Central Brazil

T. F. SILVA-DO-NASCIMENTO, 1 R. C. WILKERSON, 2 R. LOURENÇO-DE-OLIVEIRA, 1 and F. A. Monteiro 3

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ABSTRACT Anopheles halophylus Silva-do-Nascimento & Lourenço-de-Oliveira was recently described using morphological and biological variants in specimens previously identified as Anopheles triannulatus (Neiva & Pinto). Because these two species occur in sympatry in central Brazil, we used allozymes to determine the extent of gene flow to confirm that they are different species. Of 11 allozyme loci analyzed, one (Mpi) was found to be diagnostic for $An.\ halophylus$ and $An.\ triannulatus$, confirming their specific status. This locus revealed a second sibling species within $An.\ triannulatus$ sensu lato. $An.\ halophylus$ and the new undescribed species were confirmed using random amplified polymorphic DNA markers that showed moderate genetic divergence among these three sympatric and closely related taxa (D = 0.145-0.428). Moreover, this marker indicates that $An.\ halophylus$ and the new species are more closely related to each other than either is to $An.\ triannulatus$.

KEY WORDS Anopheles, taxonomy, Culicidae, allozymes, RAPD

Several important malaria vectors are members of morphologically similar (sometimes indistinguishable) species complexes. The *Anopheles* subgenus *Nyssorhynchus* is one of the most studied subgenera because of the importance of its members as malaria vectors in South and Central America.

Anopheles triannulatus (Neiva & Pinto) is a polymorphic species with a wide geographic distribution, ranging from Argentina to Nicaragua (Faran and Linthicum 1981), and recently reported from the island of Trinidad, in the West Indies (Chadee and Wilkerson 2005). Although the species is essentially zoophilic, it seems to be able to transmit malaria when in high densities (Charlwood and Wilkes 1981); thus, it has been considered to be a potential vector in some areas of Latin America (Rubio-Palis 1994). In Brazil, the morphological variability of this species has resulted in five names currently in synonymy (Faran 1980) and the designation of two subspecies (Galvão and Lane 1941). This issue was a matter of controversy because although they were recognized by some researchers (Deane et al. 1947), others still considered An. triannulatus to be a single polymorphic species (Faran

Recent studies have attempted to distinguish An. triannulatus from morphologically similar species

based on multiplex polymerase chain reaction (PCR) assays (Fritz et al. 2004) as well as determine its phylogenetic position within the *Nyssorhynchus* subgenus (Marrelli et al. 2005).

Morphological studies on An. triannulatus from several localities in Brazil, Peru, and Argentina revealed the existence of two distinct sympatric forms in Salobra, central Brazil, An. triannulatus and the recently described Anopheles halophylus Silva-do-Nascimento and Lourenço-de-Oliveira, 2002. However, certain individuals seemed to share morphological characters with both An. halophylus and An. triannulatus (Silvado-Nascimento and Lourenço-de-Oliveira, 2002) and could thus represent natural hybrids between them. We used allozymes and random amplification of polymorphic DNA (RAPD) to genetically compare sympatric An. triannulatus and An. halophylus from Salobra to corroborate their status as separate species and to shed light on the identity of the specimens presenting the variant morphology.

Materials and Methods

Mosquito Sampling and Collection Site. Two field trips were carried out in August 1996 and July 1997 to collect female *An. triannulatus* and *An. halophylus* (undescribed at the time) in Salobra (20°12′40″ S, 56°29′30″ W), Mato Grosso do Sul State, Brazil. Salobra is in the Pantanal region of Central Brazil, an area of nearly 140,000 km² subjected to annual flooding in the rainy season, from November to June (Galdino and Resende 2000). Blood-fed females were collected

¹ Departamento de Entomologia, Instituto Oswaldo Cruz, Av. Brasil 4365, 21045-900 Rio de Janeiro, RJ, Brasil.

² Department of Entomology, Walter Reed Army Institute of Research, Silver Spring, MD 20910.

³ Departamento Medicina Tropical, Instituto Oswaldo Cruz, Av. Brasil 4365, 21045-900 Rio de Janeiro, RJ, Brasil.

Table 1. Gene frequencies for 11 allozyme loci for sympatric An. triannulatus, An. halophylus, and the new species from Salobra, Brazil

		Species			
Locus	Allele	An.	An.	New	
		triannulatus	halophylus	species	
Fum	100 (n)	1.000 (30)	1.000 (23)	1.000 (16)	
α - $Gpdh$	108	0.019	0.000	0.000	
	100 (n)	0.981 (27)	1.000 (41)	1.000 (16)	
Gpi	100 (n)	1.000 (41)	1.000 (51)	1.000 (22)	
Hk-1	100	1.000	1.000	0.932	
	93 (n)	0.000(24)	0.000(27)	0.068 (22)	
Hk-2	100 (n)	1.000 (26)	1.000 (27)	1.000 (22)	
Idh-1	104	0.047	0.020	0.031	
	100	0.953	0.922	0.938	
	102 (n)	0.000 (32)	0.059(51)	0.031 (16)	
Idh-2	102	0.000	0.000	0.156	
	100	0.413	0.946	0.812	
	103 (n)	0.587 (23)	0.054(37)	0.031 (16)	
Mdh-1	104	0.048	0.000	0.000	
	100 (n)	0.952 (21)	1.000 (36)	1.000 (16)	
Mdh-2	100 (n)	1.000 (29)	1.000 (45)	1.000 (16)	
Mpi	111	0.000	0.104	0.000	
	109	0.000	0.896	0.000	
	103	0.000	0.000	0.016	
	100	0.000	0.000	0.984	
	93	0.221	0.000	0.000	
	86	0.353	0.000	0.000	
	82	0.147	0.000	0.000	
	78 (n)	0.229 (34)	0.000 (48)	0.000 (31)	
Pgm	103	0.013	0.000	0.000	
· ···	100	0.900	0.898	1.000	
	97	0.000	0.082	0.000	
	95 (n)	0.087 (40)	0.020 (49)	0.000 (16)	

 $\it n$ is number of individuals analyzed. Most frequent alleles per locus per species are in bold.

from horse bait between 1800 and 2100 hours for seven consecutive days on each occasion. Mosquitoes were identified (Consoli and Lourenço-de-Oliveira 1994), individually separated into labeled vials to lay eggs, and the progeny were separately reared. For An. halophylus, however, complete larval development was only achieved when brackish water from the local breeding sites was added to the rearing containers (Silva-do-Nascimento and Lourenço-de-Oliveira 2002). A single mosquito from each F1 was used for subsequent molecular analyses. Eggs, fourth instars, and adults of each progeny were stored in 4% glutaraldehyde, 70% ethanol, or pinned, respectively, for further morphological analyses.

Allozymes. Each mosquito was homogenized in 25 μ l of lysis buffer (0.05 mM Tris-HCl, pH 8.0, 0.01 mM EDTA, 15 mM dithiothreitol, 0.01 mM ϵ -amino-n-caproic acid, and 1% Triton X-100). Two μ l of each

homogenate were loaded onto a 1% agarose gel and subjected to electrophoresis as described by Momen and Salles (1985) and stained according to Manchenko (1994). Specimens were analyzed for eight enzymes: fumarate hydratase (FUM, E.C. 4.2.1.2.), glucose phosphate isomerase (GPI, E.C. 5.3.1.9.), α-glycerophosphate dehydrogenase (α -GPDH, E.C. 1.1.1.8.), hexokinase (HK, E.C. 2.7.1.1.), isocitrate dehydrogenase (IDH, E.C. 1.1.1.42.), malate dehydrogenase (MDH, E.C. 1.1.1.37.), mannose phosphate isomerase (MPI, E.C. 5.3.1.8.), and phosphoglucomutase (PGM, E.C. 2.7.5.1.). Genotype frequencies for each locus were obtained directly from the electrophoretic phenotypes. The most frequent electromorph for each enzyme was designated arbitrarily as the "100" reference allele, and motilities for the other electromorphs were determined relative to this common electromorph.

Genepop online (version 3.4) (Raymond and Rousset 1995) was used to calculate allele frequencies and agreement with Hardy–Weinberg (H-W) equilibrium. F_{IS} was computed as in Weir and Cockerham (1984). Goodness-of-fit to H-W expectations was assessed for each locus in each species by using the exact probability test available in Genepop. Levels of genetic variability (i.e., number of alleles per locus, percentage of polymorphic loci, and observed and expected heterozygosities) were calculated for the three species with the Biosys-1 software (Swofford and Selander 1981).

RAPD-Polymerase Chain Reaction (PCR). DNA isolation followed the protocol described in Wilkerson et al. (1993). Sixty-seven decamer primers (Operon Technologies, Alameda, CA) were screened and 14 (A8, A11, A19, B3, B7, B8, B10, B11, B15, C4, C10, C16, D15, and D16) were selected on the basis of their reproducibility, polymorphism, and efficiency in the PCR amplification. The reproducibility of the amplification was tested using the same DNA sample in three different PCR reactions. These primers were used to amplify five to six specimens of each of the three putative species from Salobra. The procedure was essentially as described in Wilkerson et al. (1995), except that 40 PCR cycles were used.

RAPD band patterns were interpreted following Wilkerson et al. (1993). It was assumed that the populations were in H-W equilibrium and that bands that comigrate are homologous. RAPD markers were analyzed using the software program TFPGA version 1.3 (Miller 1997) to calculate Nei's unbiased genetic distances and characterized using the unweighted pair group method with arithmetic means (UPGMA). Re-

Table 2. Genetic variability in An. triannulatus, An. halophylus, and the new species

Species	Mean sample size/locus	Mean no. of alleles/locus	Polymorphic loci (%) ^a	Mean heterozygosity	
Species	Mean sample size/ locus	Mean no. of aneles/locus	rolymorphic loci (%)	Но	He
An. triannulatus	28.4 (2.4)	1.8 (0.3)	0.45	0.142 (0.071)	0.153 (0.074)
An. halophylus New species	39.5 (3.1) 19.0 (1.5)	1.5 (0.2) 1.6 (0.2)	0.27 0.36	0.054 (0.024) 0.061 (0.035)	0.057 (0.025) 0.055 (0.031)

Standard error values are in parentheses

 $[^]a$ A locus was considered polymorphic if the frequency of the most common allele was \leq 0.95.

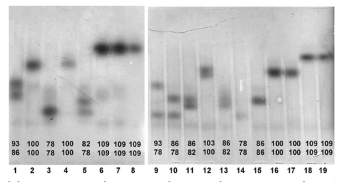


Fig. 1. Mpi agarose gel showing genotypes for An. triannulatus (samples 1, 3, 5, 9–11, and 13–15), An. halophylus (samples 6–8, 18, 19), and the new species (samples 2, 4, 12, 16, and 17).

liability of the groups in the phenogram was assessed through 1,000 bootstrap replicates of the alleles.

Results

Allozymes. In total, 123 specimens were analyzed. Allele frequencies for the 11 enzyme loci scored are given in Table 1. Six loci (Fum, α -Gpdh, Gpi, Hk-2, and Mdh-1 and -2) were monomorphic for all samples. A locus was considered monomorphic if the frequency of the most common allele was >0.95 (Tables 1 and 2).

The Mpi locus was diagnostic (sensu Avala 1983) between An. triannulatus and An. halophylus, corroborating their morphological separation. Furthermore, this locus revealed a third isolated genetic group composed of the variant individuals that share morphological characters with both An. triannulatus and An. halophylus (Fig. 1: Table 1). The analysis of the variant individuals together with either An. triannulatus or An. halophylus, would lead to strong departures from the H-W equilibrium ($F_{IS} = 0.521, \bar{P} < \hat{0}.001; F_{IS} = 0.841,$ P < 0.001, respectively) for this locus, as opposed to the lack of disequilibrium when the three species are analyzed separately ($F_{IS}=0.044,\,P=0.109,\,{\rm for}\,An.\,triannulatus,\,{\rm and}\,F_{IS}=0.340,\,P=0.059,\,{\rm for}\,An.\,halo$ *phylus*; there is no \overline{F}_{IS} value for the variant individuals as a single heterozygote was detected; Table 3). The presence of different alleles in the Mpi locus for the three putative species was observed not only when insects from the two consecutive field trips (1996 and

Table 3. Wright's fixation index $(F_{\rm IS})$ for An. triannulatus, An. halophylus, and the new species

	F _{IS}				
Locus	An. triannulatus	An. halophylus	New species		
Hexokinase-1	_	_	-0.050 ns		
Isocitrate dehydrogenase-1	-0.033 ns	-0.058 ns	-0.017 ns		
Isocitrate dehydrogenase-2	0.036 ns	-0.043 ns	-0.161 ns		
Manose-6-phosphate- isomerase	$0.044~\mathrm{ns}$	0.340 ns	_		
Malate dehydrogenase-1	-0.026 ns	_	_		
Phosphoglucomutase	$-0.085 \mathrm{\ ns}$	$-0.084~\mathrm{ns}$	_		

^{—,} a single allele was detected; ns, nonsignificant (P > 0.05).

1997) were analyzed together but also when samples from each year were analyzed separately (data not shown). Significant (although not diagnostic, as with Mpi) differences in allele frequencies among the three species were detected for the Idh-2 locus ($\chi^2=2.25$, df = 2, P < 0.002, between An. halophylus and the new species; $\chi^2=42.05$, df = 1, P < 0.0001, between An. halophylus and An. triannulatus; and $\chi^2=28.64$, df = 2, P < 0.0001, between the new species and An. triannulatus), whereas the Pgm loci presented differences only between An. halophylus and An. triannulatus ($\chi^2=11.68$; df = 3, P < 0.001).

No departures from H-W equilibrium were observed for the other four polymorphic loci (Hk-I, Idh-I and -2, and Pgm) for any of the three species ($-0.161 < F_{IS} < 0.340, P > 0.05$) (Table 3). The mean number of alleles per locus ranged from 1.5 to 1.8, the percentage of polymorphic loci from 27 to 45%, and the mean expected heterozygosity (He) from 0.055 to 0.153 (Table 2).

RAPD-PCR. Five An. triannulatus, six An. halophylus, and six individuals of variant morphology were amplified with 14 primers, producing 123 reproducible bands. Bands ranged in size from 310 to 2,300 bp. Average genetic distance (D; Nei 1978) between An. triannulatus and An. halophylus was 0.288; between An. triannulatus and the morphological variant was 0.428, and between An. halophylus and the morphological variant was 0.145, supporting the observations seen with allozymes.

A UPGMA dendrogram based on Nei's distances was constructed, revealing the existence of three well supported clusters (bootstrap values \geq 78), each containing a single species consistent with the Mpi allozyme segregation of the three species (Fig. 2).

Discussion

Allozyme and RAPD markers confirmed that there are genetic differences between *An. halophylus* and *An. triannulatus* and revealed the existence of a third sympatric species in the Triannulatus Complex. The lack of shared *Mpi* alleles and moderate RAPD-based genetic distances among the three forms collected in

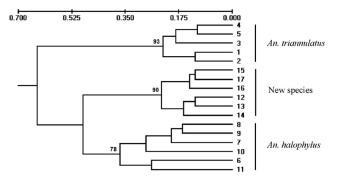


Fig. 2. UPGMA dendrogram constructed based on RAPD-derived distance matrix (Nei 1978) for *An. triannulatus*, *An. halophylus*, and the new undescribed species. Numbers represent individual mosquitoes tested using RAPD. Numbers to the left of each group are bootstrap support values.

sympatry in Salobra demonstrate the occurrence of three different biological species in that region. In addition to the *Mpi* diagnostic locus, there are significant gene frequency differences in the *Idh*-2 locus among the three species (see Results).

The existence of fixed or diagnostic alleles between sympatric populations is evidence for a barrier to gene flow among the populations and that they are different species. The observation that the diagnostic difference at the *Mpi* locus was detected in independent collections in two consecutive years reinforces that conclusion and rules out the possibility of the specimens with intermediate morphology being natural hybrids between *An. halophylus* and *An. triannulatus*. Natural hybrids would present different combinations of the *Mpi* alleles present in both parental species (78, 82, 86, 93, 109, and 111), instead of the new allelic variants observed (alleles 100 and 103; Table 1; Fig. 1).

An. triannulatus, An. halophylus, and the new species presented heterozygosity values ranging from 0.055 ± 0.031 to 0.153 ± 0.074 . These values are similar to what has been reported for other anopheline species (Manguin et al. 1999, Santos et al. 2004).

Salobra is located in the Pantanal region of central Brazil, which is an immense alluvial plain with a complex ecological system that includes seasonally flooded grasslands, gallery and dry forests, numerous river corridors, and lakes of fresh and brackish water. Salobra (Portuguese for brackish water) is at the Miranda River margins, one of the major tributaries of the Paraguay River. The Miranda River originates in the Serra da Bodoguena, where the calcareous soil contributes to the high calcium carbonate and chloride concentration in the water. During the rainy season, both fresh and brackish water habitats are present around Salobra, providing a variety of ecosystems for mosquito breeding. It was noted that the water of one breeding site of An. halophylus had a high concentration of chloride (from 2.0 to 4.0 mg/liter) that was critical for larval development. In contrast, An. triannulatus breeds solely in freshwater (Silva-do-Nascimento and Lourenço-de-Oliveira 2002). Therefore, the chemical composition of breeding site waters may be selective for species development in Salobra and elsewhere in the Paraguay basin.

Results of both allozyme and RAPD analyses of *An*. triannulatus and An. halophylus from Salobra are in accordance with previous morphological findings (Silvado-Nascimento and Lourenço-de-Oliveira 2002) and reveal the existence of a third taxon in the Triannulatus Complex. Although females of the three species are morphologically indistinguishable, An. triannulatus and An. halophylus males, larvae, and eggs may be distinguished based on several characters (Silva-do-Nascimento and Lourenço-de-Oliveira 2002). The specimens that belong to the new species were initially classified as variants because they shared morphological characters with both An. triannulatus and An. halophylus. For example, the new species presents usually long lateral arms of the median plate of the spiracular apparatus similar to An. triannulatus, as opposed to the rudimentary or completely absent arms of An. halophylus. However, with respect to the male genitalia, the new species presents broad and usually directed proximally apicolateral lobes of the claspette, similarly to An. halophylus, but different from the narrow structures that characterize An. triannulatus. In addition, we cannot associate this taxon with any of the current An. triannulatus synonyms, which led us to conclude that it represents an undescribed species (see Silva-do-Nascimento and Lourenco-de-Oliveira 2002 for a discussion). Work toward the description and diagnosis of the new species is underway.

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