MOLECULAR IDENTIFICATION OF Aedes bahamensis (DIPTERA: CULICIDAE)

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Aedes (Howardina) bahamensis (Howardina bahamensis of Reinert et al. 2004) is an exotic species first detected in the United States in 1986 from eggs and adult females collected in Miami-Dade and Broward counties, Florida (Berlin 1969; Pafume et al. 1988; O’Meara et al. 1989). In the U.S., immature Ae. bahamensis are chiefly found in artificial containers (e.g., abandoned tires, cemetery vases, etc.) and exotic tank bromeliads (O’Meara et al. 1995a). Although Ae. bahamensis readily feeds on humans, it is not a major pest or disease vector. However, the immature stages are frequently found in habitats that are also known to harbor 2 medically important mosquito species: Ae. albopictus (Skuse) and Ae. aegypti (L.) (O’Meara et al. 1995b; Lounibos et al. 2010). Morphological differences between these 3 species provide useful characters that readily distinguish the egg, larval, and adult life stages (Linley 1989; Darsie & Ward 2005). Nevertheless, a molecular method to identify Ae. bahamensis would be useful to confirm the identity of damaged adult specimens collected in fan-based traps and would allow rapid identification of the species at any life stage. Here we report the rDNA second internal transcribed spacer (ITS2) sequence and a species-specific PCR primer for the identification of Ae. bahamensis.

Specimens of Aedes bahamensis were obtained from colony material maintained by GFO at the Florida Medical Entomology Laboratory (Vero Beach, Florida) and field samples collected by BDB. Field collected specimens were identified using key characters described by Darsie (1992). Total DNA was obtained from adult specimens (8 colony and 14 field) using the Qiagen DNeasy kit (Qiagen, Valencia, California) or the DNAzol reagent (Molecular Research Center, Inc., Cincinnati, Ohio) per the manufacturers’ instructions. The resulting extractions were PCR amplified in 50 μL reactions using the Invitrogen PCR Supermix (Invitrogen, Carsbad, CA). Each reaction mixture contained 3 μLs of DNA template (4-35 ng/μL), 1 μL of each forward and reverse primer (200 nM final concentration), and 45 μL of the PCR supermix. Amplification cycling conditions were 94 °C for 5 min followed by 35 cycles of 94 °C for 1 min, 54 °C for 30 s and 72 °C for 1 min. The CP-P1A/P1B primer pair was used to amplify the complete ITS2 (Fig. 1) (Wesson et al. 1992). A negative control (H2O en lieu of DNA template) was included in each run. PCR amplicons were visualized on a 1.5% agarose gel. There were no obvious intraspecific amplicon size polymorphisms. The PCR products were gel purified with the Qiaquick Gel Extraction kit (Qiagen) and subsequently cloned into the pCR 2.1 TOPO vector (Invitrogen). Purified plasmids were obtained using the Promega Wizard Plus SV miniprep kit (Promega, Madison, WI) and then sequenced (n = 10) using the Applied Biosystems (Carlsbad, CA) Big Dye Terminator V3.0 chemistry by the Davis Sequencing Facility, University of California (Davis, California). The sequences were verified as ITS2 after evaluating the results of an NCBI BLAST query, secondary structure analysis, and the identification of specific sequence motifs known to exist on the ITS2 of mosquitoes (Coleman 2007). Novel ITS2 sequences, partial 5.8S, and partial 28S sequences for Aedes bahamensis were annotated and representative samples were submitted to the NCBI GenBank (Accession numbers: JN020552, JN020553) (Keller et al. 2009).

The CP-P1A/P1B primer pair produces a 380 base pair (bp) amplicon for Ae. bahamensis that differs from the amplicons of Ae. albopictus (600 bp), Ae. aegypti (360 bp), and Ae. triseriatus (Say)
(385 bp) using the same primer pairs (Fig. 2). These four species are sympatric in southern Florida where the distribution of *Ae. bahamensis* remains limited to Miami-Dade and Broward counties. Because the CP-P1A/P1B size polymorphisms may not be readily distinguished between *Ae. bahamensis* and *Ae. aegypti* or *Ae. triseriatus*, a species-specific reverse primer was designed. In order to identify a suitable region to design an *Ae. bahamensis* specific primer, a multiple sequence alignment was created using Culicidae rDNA ITS2 sequences obtained from GenBank and the novel sequences obtained in this study. A species-specific primer (*Aebah1*: 5′-aacatagccaggtgtatg3′) was then designed using Primer3 to produce a 300 bp amplicon when used with the CP-P1A forward primer (Fig. 2) (Rozen & Skaletsky 2000). The PCR amplification cycling conditions for this primer pair are identical to the CP-P1A/P1B conditions reported above. The CP-P1A/Aebah1 primer pair will not amplify the sympatric container-inhabiting *Aedes* (i.e., *Ae. albopictus*, *Ae. aegypti*, or *Ae. triseriatus*) (Fig. 2) or other container inhabiting *Aedes* mosquitoes [e.g., *Ae. hensersoni* (Cockerell), *Ae. atropalpus* (Coquillett) and *Ae. japonicus* (Theobald)] found in some areas of the southeastern United States (data not shown).

To our knowledge, the work described here represents the first PCR based method for the rapid molecular identification of *Aedes bahamensis*. Such assays are particularly useful to confirm the identity of a specimen when key morphological characters become damaged. PCR based assays may be used on genomic DNA obtained from an organism at any life stage. Similarly, species-specific PCR assays may be used to validate the integrity of pooled mosquito specimens and confirm the presence of heterospecies contamination (Gerhardt et al. 2001).

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**SUMMARY**

Here we report novel rDNA ITS2 sequences for *Aedes bahamensis* and a species-specific PCR primer that produces a diagnostic 300 bp PCR product. This technique may be used to confirm the identity of damaged or degraded specimens that may not be readily identifiable using morphological characteristics.

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