Evidence for an Overwintering Population of *Aedes aegypti* in Capitol Hill Neighborhood, Washington, DC

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Abstract. *Aedes aegypti* is an invasive, highly anthropophilic mosquito and a major vector for dengue and chikungunya. Population persistence in the continental United States is reportedly limited to southward of the average 10°C winter isotherm, which in the east, bisects Alabama, Mississippi, Georgia, and South Carolina. We report on summer collections and genotypic analyses of *Ae. aegypti* collected in the Capitol Hill neighborhood in Washington, DC (WDC). Analysis of a 441-bp fragment of the mitochondrial *cytochrome oxidase I* gene sequence identified the same two haplotype sequences during 2011–2014, and placed these within two discrete groups known to be derived from lineages resident in the Americas. Analysis of 10 microsatellite loci for specimens collected during 2011–2014 revealed no evidence for introgression of new alleles across years. Overall, our data support a conclusion that this represents a resident WDC population, likely maintained during winter months in a subterranean habitat that facilitates year-round survival.

*Aedes aegypti* is the primary vector of dengue, the most common arboviral disease, across most tropical and subtropical regions of the world. It is also a major vector of chikungunya, a rapidly emerging virus that is causing significant disease throughout the Caribbean islands and much of the Central and South America. In the continental United States, autochthonous transmission of dengue has been observed in Florida since 2009. No licensed vaccines or therapeutic drugs are available for preventing or treating either disease. *Aedes aegypti* is a highly anthropophilic invasive species that is well adapted for wide-spread colonization of urban environments, as it readily breeds in water in natural and man-made containers around and within human dwellings. Although *Ae. aegypti* has likely maintained an intermittent presence in the United States for ∼375 years, its distribution has been assumed to be limited by cold temperatures to boundaries roughly southward of the average 10°C winter isotherm. However, two potentially synergistic phenomena may be at play, which promote sustained northward range expansion: 1) the influence of climate changes that will allow for expansion into areas not previously colonized and 2) this species readily adapts to breeding in subterranean structures that provide a suitable and protective environment from adverse climatic conditions. Although *Ae. aegypti* collections have been reported from Virginia and Maryland, they typically comprised relatively small numbers and have been considered to be seasonal introductions, and restricted from overwintering by cold-season extremes, as no life stage is tolerant of even short-term (e.g., overnight) temperatures below 0°C. Here, we report on summer-season collections and genotypic analyses of *Ae. aegypti* larvae and adults across four contiguous years in the Capitol Hill neighborhood in Washington, DC (WDC), and conclude that this is likely an overwintering population maintained in as yet undefined subterranean habitats.

After the initial serendipitous discovery of a single adult female *Ae. aegypti* on October 22, 2011, in the Capitol Hill neighborhood, multiple subsequent surveys were performed to collect *Ae. aegypti* larvae from container habitats within 200 m of the original collection site at the square fountain (Figure 1A). Larvae were collected from a bird bath, square fountain, trash can, potted plant saucer, and a small, grated stormwater basin in an alleyway (Figure 1B and C). Collection dates (sample sizes) were October 22 and 25, 2011 (6 and 20, respectively); September 26, 2012 (17); July 13, August 25, October 19, and November 22, 2013 (6, 1, 8, and 1, respectively); and August 18 and September 14, 2014 (49 and 7, respectively). A total of 115 *Ae. aegypti* larvae were identified using standard taxonomic keys. Adults were occasionally reared from larvae, killed by freezing, and dried by placing them in open glass vials. Larvae were killed with hot water then stored in 2.5-mL Eppendorf tubes containing 70% ethyl alcohol. Samples were shipped to University of Notre Dame for genetic analysis.

Ethanol-preserved larval and dried adult samples were subjected to DNA extraction as previously described, and DNA was suspended in final volumes of 200 μL for larvae and 1,000 μL for adults containing 0.02 M NaOH and 0.036 M Tris-HCl, pH 8.0. A 710-bp amplicon from the mitochondrial *cytochrome oxidase I* (*mtCOI*) gene was obtained using LCO1490 and HC02199 primers in 25 μL PCR reactions containing 1 μL of genomic DNA and was subjected to thermocycling conditions as previously reported. Amplicons were purified (Qiagen QIAquick PCR Purification Kit) and direct sequenced with the same primers using BigDye on a 3730xl DNA Analyzer (Applied Biosystems). Multiplexes of primers for 10 microsatellite markers with fluorochrome-labeled forward primers (Integrated DNA Technologies) were prepared in 25 μL PCR reactions, amplified, and subjected to fragment analysis as previously described.

Fifty-nine *mtCOI* sequences were obtained and aligned in Clustal Omega with 20 *Ae. aegypti* *mtCOI* sequences from various geographic regions of the world downloaded from VectorBase. All sequences were trimmed to a 441-bp consensus and used to construct a neighbor-joining phylogenetic tree in MEGA6 using a *mtCOI* sequence from *Aedes albopictus* (GenBank accession no. KC690960) as an outgroup. For microsatellite analysis, samples were divided by year of collection (YOC) and tested for Hardy–Weinberg equilibrium (HWE) using Arlequin v3.5. The Bayesian clustering method in the STRUCTURE program v2.3.4 was used to assign individuals to genetic clusters. We implemented
the Evanno method\textsuperscript{15} to determine the most likely number of clusters (K) after five independent runs of STRUCTURE for K = 1 to K = 5 on individuals from four YOC, using 100,000 burn-in steps followed by 500,000 iterations.

We identified two mtCOI haplotypes from a total of 59 sequences obtained from \textit{Ae. aegypti} individuals collected during 2011–2014. The two haplotypes were shared across all four YOC, and were often identified in larvae collected from the same breeding container such as the birdbath (Figure 1B). Our sequence data were submitted to GenBank under accession numbers KM362421 and KM362422. The phylogenetic tree based on a combined analysis of the WDC sequences with 20 mtCOI haplotype sequences from various geographic regions placed the WDC haplotypes within two distinct phylogenetic groups (Figure 2A), reflective of the two African origin ancestral clades associated with extant global \textit{Ae. aegypti} populations.\textsuperscript{16} The most common WDC haplotype A showed close relationships with \textit{Ae. aegypti} populations from South America and Asia, whereas WDC haplotype B showed close relationships with \textit{Ae. aegypti} populations from the Caribbean, South America, and sub-Saharan Africa. This finding is completely consistent with results obtained from multiple studies on \textit{Ae. aegypti} populations throughout the Caribbean and Central and South America and with data from multiple mitochondrial genes.\textsuperscript{17–19} This and the fact that the two haplotype sequences are identical to the haplotype sequences reported for \textit{Ae. aegypti} populations in Brazil and Guyana support the conclusion that the WDC \textit{Ae. aegypti} population is derived from lineages resident in the Americas, and were likely transported into WDC via human-mediated travel. These data also support the expected outcome of a persistent low genetic diversity founder population; for example, in contrast, analyses of \textit{Ae. aegypti} mtND4 and mtND5 haplotypes across southern Florida in 2006 revealed 68 different haplotypes among 362 individuals examined.\textsuperscript{19}

We determined genotypes at 10 microsatellite loci for 70 individuals from the WDC population collected in 2011 (N = 22), 2012 (N = 12), 2013 (N = 8), and 2014 (N = 28). These loci fit HWE expectations with only two exceptions across all years, with the total number of alleles observed per locus across all years ranging from two to 10 (Figure 3). Examination of individual allele presence and frequency across years clearly reflect no evidence of new allele introgression.
across all 10 loci, further supporting our conclusion that this represents an overwintering population in WDC. Bayesian cluster analysis using STRUCTURE identified the likely number of clusters as $K = 2$, but failed to detect any evidence for clustering across YOC as the WDC annual populations represented similar mixtures of the identified genetic clusters (Figure 2B).

Taken together, our mtCOI haplotype and microsatellite marker data confirm consistent genetic identities among *Ae. aegypti* WDC populations across all YOC, with no evidence of introgression by novel genotypes. The most parsimonious conclusion is that this represents a small continuous resident WDC population that reflects founder effects since the original introduction during or prior to our initial sampling.
in 2011. This raises interesting questions about how the population has persisted despite the obvious extreme winter seasonal conditions, with an average January isotherm for WDC of only 2.2°C.20 A historical review of personal conditions, with an average January isotherm for WDC in 2011. This raises interesting questions about how the population has persisted despite the obvious extreme winter sea-

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FIGURE 3. Allele frequencies observed for microsatellite loci among individuals collected in Washington, DC, during 2011–2014. Numbers in brackets indicate number of alleles per locus.

