

PATHWAYS OF EXPANSION AND MULTIPLE INTRODUCTIONS ILLUSTRATED BY LARGE GENETIC DIFFERENTIATION AMONG WORLDWIDE POPULATIONS OF THE SOUTHERN HOUSE MOSQUITO

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Abstract. The southern house mosquito *Culex quinquefasciatus* is a principal vector of human lymphatic filariasis, several encephalitides (including West Nile virus), avian malaria, and poxvirus, but its importance as a vector varies considerably among regions. This species has spread with humans and is ubiquitous in tropical urban and suburban environments. This was the first mosquito to reach Hawaii and we performed a worldwide genetic survey using microsatellite loci to identify its source. Our analyses showed divergent Old World and New World genetic signatures in *Cx. quinquefasciatus* with further distinctions between east and west African, Asian, and Pacific populations that correlate with the epidemiology of human filariasis. We found that in Hawaii south Pacific mosquitoes have largely replaced the original New World introduction of *Cx. quinquefasciatus*, consistent with their reported expansion to higher elevations. We hypothesize worldwide pathways of expansion of this disease vector.

INTRODUCTION

A select group of insect vectors of disease have expanded their ranges radically in association with humans.¹ Although their introduction to new areas has sometimes heralded disease outbreaks, e.g., yellow fever epidemics in the New World after the introduction of *Aedes aegypti*,² the distribution of disease vectors does not always correlate with the distribution of the diseases they transmit.³ For example, nocturnal periodic lymphatic filariasis (*Wuchereria bancrofti*) is transmitted primarily (some say exclusively) by *Culex* (*Culex*) *quinquefasciatus* Say in urban eastern Africa and in Asia.⁴ However, although *Cx. quinquefasciatus* is omnipresent and very common across all tropical and subtropical regions of the world,⁵ the primary vectors of nocturnal filariasis in rural and western Africa are several *Anopheles* species, while in the Pacific islands the nocturnal form of *W. bancrofti* is virtually absent and diurnal/sub-periodic forms are transmitted by local *Aedes* species.⁴ Across the world *Cx. quinquefasciatus* is also a locally important vector of St. Louis encephalitis and West Nile virus, as well as of avian malaria and pox viruses.⁶

New or modified vector-mediated host/parasite interactions occur when natural or human-assisted introductions of vectors and parasites are made into new ranges.^{1,7} However, in addition to possible new vector-disease-host combinations, the vector itself may undergo local selection, genetic drift, or hybridizations that can modify their ability to transmit a disease.^{8,9} Our capacity to predict, prepare, and react to emerging arthropod-borne diseases depends not only on our understanding of emerging disease organisms, but also their vectors. As a model system we have been focusing on one of the best-known mosquito introductions, that of *Cx. quinquefasciatus* to the Hawaiian Islands where as the sole vector of avian malaria it has contributed to the endangerment of many endemic forest bird species.¹⁰ T. R. Peale, an American naturalist, found no mosquitoes when he visited the islands in 1823,¹¹ so as the first introduced mosquito, *Cx. quinquefasciatus* was conspicuous to natives, explorers, and missionaries.

The most detailed and possibly the most speculative description of the introduction is that of Reverend William Richards¹² who indicts the crew of the "Wellington" for releasing larvae of the southern house mosquito with old drinking water obtained in San Blas, Mexico, while at port in Lahaina, Maui, in 1826. Other authors provide fewer details, but because *Cx. quinquefasciatus* was then not likely present in the Pacific,¹³ the mosquito source is always the New World.^{11,14} The epidemiology of avian malaria (*Plasmodium relictum*) in the Hawaiian Islands is often cited as a classic example of co-evolution subsequent to the introduction of disease to a highly susceptible, isolated wildlife population.¹⁵ However, the role of the vectors has not been previously considered. The objective of our study was to examine the history of introductions of *Cx. quinquefasciatus* to Hawaii. While doing so we uncovered evidence of multiple introductions diagnosed by an unexpected degree of genetic differentiation worldwide.

MATERIALS AND METHODS

Samples and microsatellite genotyping. In an attempt to obtain a snapshot of the genetic signature of *Cx. quinquefasciatus* across the world, we obtained specimens from as many different locations as possible with the aid of local entomologists (PHS permit no. 99-05-0660) or from colleagues (Table 1). For an updated yet still incomplete map of the distribution of *Cx. quinquefasciatus*, see the report by Smith and Fonseca.¹⁶ We excluded samples from Shanghai, China, because they showed considerable hybridization with *Cx. pipiens pallens*, an east Asian member of the *Cx. pipiens* complex.¹⁶ To achieve representative sample sizes, we chose to combine close-by samples (within each country or geographic location). The exceptions are Hawaii, where we examined separately the current (Oahu) and historic (Maui) main entry ports because we had ample access to specimens and one of our objectives was to understand better the timing of introductions. We also decided not to combine specimens from the Midway Atoll (Hawaii) with other Hawaiian specimens because we had prior knowledge of separate introductions to this atoll,¹⁷ which was the site of extensive naval traffic during World War II. We also did not combine specimens from

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TABLE 1

Collection location, samples sizes, stage from which DNA was extracted, date of collection, and sources of *Culex quinquefasciatus* analyzed*

Location	No.	Stage	Date	Source
1 Makurdi and Zaria, Nigeria	20	A	June 1999 ¹	Light traps near houses
2 Dongola, Sudan	28	L	Jan 2003 ²	Three larval dips within city limits
3 Kisumu and Malindi, Kenya	31	A	May 1999 ³	Collected inside houses in five villages
4 Kochi, India	9	A	May 2000 ⁴	Hand collected inside medical compound
5 Depok (near Jakarta), Java, Indonesia	16	A	1999, 2003 ⁵	Adults reared from larvae collected within city limits
6 Makassar, Sulawesi, Indonesia	8	A	Aug 2003 ⁶	Adults reared from larvae collected within city limits
7 Kupang, West Timor, Indonesia	32	A	1999, 2003 ⁵	Adults reared from larvae collected within city limits
8 Okinawa, Japan	24	A/L	April 1999 ⁷	Light traps and larval dips
9 Cairns, Australia	22	A	2004 ⁸	Adults raised from > 7 egg rafts
10 Kingswood and Glandore, Australia	25	A	April 1999 ⁸	Adults raised in the lab from > 100 egg rafts
11 Auckland, New Zealand	35	L	Jan 1999 ⁹	Larval dips. Three separate sites within city limits
12 American Samoa	14	L	1999 ¹⁰	Larval dips
13 Midway, HI, USA	10	L	1998 ¹⁰	Larvae in rainwater pooled in plastic covers
14 Oahu, HI, USA	25	A	1998 ¹¹	Light traps in airport, Pearl Harbor, and Honolulu
15 Maui, HI, USA	53	A/L	1997 ¹²	Lihue, Lahaina, Airport, and cattle puddle up Haleakala
16 Chino, CA, USA	46	A	Oct 2002 ¹³	Light traps within city limits
17 Jalisco, Mexico	44	A	1998 ¹⁴	Light traps at Estacion Biologica de Chamela
18 Tapachula, Chiapas, Mexico	24	A	July 1998 ¹⁵	Light traps near Centro de Investigacion de Paludismo
19 Santa Cruz, Galapagos Islands, Ecuador	29	A	May 2004 ¹⁶	Oviposition trap next to house (27 egg rafts), Puerto Ayora
20 Manta, Ecuador	17	A	Dec 2004 ¹⁷	Light traps within city limits
21 New Orleans, LA, USA	24	L	March 1999 ¹⁸	Larval dips at sewage treatment plant
22 Archer, FL, USA	24	L	Sept 1998 ¹⁹	Collected from 5 scrap tires
23 Bermuda, UK	44	L	Dec 2004 ²⁰	Human containers (boat, tires, etc.) across the island
24 George Town, Cayman Islands, UK	24	A	June 2003 ²¹	Light traps within city limits
25 Savanna la Mar, Jamaica	22	A	Dec 2004 ¹	Adults reared from larvae collected with oviposition traps
26 Trujillo, Venezuela	30	L	Mar 2004 ²²	Larval dips into large cemetery flower pot (multiple cohorts)
27 Amapá, Macapá, Brazil	13	A	May 1997 ²³	Hand-net collection of swarming specimens inside restaurant
28 São Paulo, Brazil	42	A	Dec 2002 ²⁴	Aspirations from ferns, Alto do Pinheiro, next to houses
Total	735			

* The numbers in the left column correspond to those in Figures 1 and 2 and in the allelic frequency tables in the Supplementary Materials. A = adult; L = larva.
¹ Jan Conn, Wadsworth Center, Albany, NY; ² Colin Malcolm, Queen Mary, University of London, UK; ³ Shirley Luckhart, Virginia Tech, Blacksburg, VA and Dina M. Fonseca, National Museum of Natural History, Smithsonian Institution; ⁴ Motoyoshi Mogi, Saga Medical School, Saga, Japan; ⁵ Ichiro Miyagi, University of the Ryukyus, Okinawa, Japan; ⁶ Craig R. Williams, James Cook University, Cairns, Australia; ⁷ Jo Kent and Craig R. Williams, University of South Australia, Adelaide, Australia; ⁸ Mark Dullman, AgriQuality New Zealand, Auckland, New Zealand; ⁹ Dennis LaPointe and Carter Atkinson, U.S. Geological Survey Pacific Island Ecosystem Research Center, HI; ¹⁰ George Kitajuchi, Hawaii State Department of Health, Oahu, HI and Dina M. Fonseca; ¹¹ Michael D. Tancayo, Jr., Hawaii State Department of Health, Kahului, HI and Dina M. Fonseca; ¹² Min Lee Cheng, West Valley Mosquito and Vector Control District, Chino, CA; ¹³ Felipe Nogueira, Estacion de Biologia Chamela, Jalisco, Mexico; ¹⁴ Juan Arredondo Jimenez, Centro de Investigacion de Paludismo, Mexico; ¹⁵ Laura Kramer, Wadsworth Center, Albany, NY; ¹⁶ Ben Pagac, Army Center for Health Promotion and Preventive Medicine-North, Fort Meade, MD; ¹⁷ Mike Carroll, New Orleans Mosquito and Termite Control Board, New Orleans, LA; ¹⁸ George O'Meara, Florida Medical Entomology Laboratory, Vero Beach, FL; ¹⁹ David Kendell, Bermuda Government Pest Control, Hamilton, Bermuda and Julie L. Smith; ²⁰ William Petric, Mosquito Research and Control Unit, Cayman Islands; ²¹ Richard C. Wilkerson; ²² Anice Sallum, Universidade de São Paulo, São Paulo, Brazil.

mainland Ecuador and the Galapagos Islands so we might investigate the sources of the recent introduction of *Cx. quinquefasciatus* to the Galapagos Islands. Furthermore, we extracted DNA from five-dried museum specimens collected between 1919 and 1944 in Hawaii (Bishop Museum collections).

All specimens were field collected as adults or larvae and they were sent to us either dry or in ethanol. Larval collections were made from multiple oviposition sites in each location to prevent a few families from influencing the results. A morphologic examination, including genitalia analysis of males¹⁸ to confirm species, was performed prior to DNA extraction. In later samples, rapid polymerase chain reaction (PCR)-based assays were used to confirm species identification.¹⁶ DNA was extracted using a standard phenol/chloroform method¹⁷ and 1 μ L of the DNA was used in each PCR. To prevent contamination with male DNA, female abdomens were not used. The DNA extraction and PCR preparations involving the five dried museum specimens were performed in a separate room away from the main laboratories. We examined twelve microsatellite loci: CQ11, CQ26, CQ29, CQ41, CQ46, pGT12, pGT46, pGT51, and qGA12, qGT4, qGT8, and qGT17 (Table 2). Analyses of mosquito families have showed that all the microsatellite loci used in this study are inherited in a Mendelian fashion and are not sex-linked.^{19–21} Microsatellite loci were amplified and sized as described by Smith and others.²¹

Statistical analyses. Tests of Hardy-Weinberg equilibrium were conducted using GENEPOP 1.2,²² and allelic richness was calculated with FSTAT 2.9.3²³ using a rarefaction index (2N = 10) to account for different sample sizes. We used Cavalli-Sforza and Edwards chord distance as well as Nei's genetic distance as a measures of population differentiation,²⁴ which were implemented using the programs MSA3.0,²⁵ fol-

TABLE 2
Microsatellite loci used in this study*

Locus name	H _e	No. of alleles
CQ11 ^A	0.68	27
CQ26 ^A	0.61	14
CQ29 ^A	0.41	7
CQ41 ^A	0.64	26
CQ46	0.74	11
qGA12 ^B	0.70	19
qGT4 ^B	0.58	10
qGT8 ^B	0.68	24
qGT17 ^B	0.68	21
pGT12 ^C	0.36	8
pGT46 ^C	0.65	15
pGT51 ^C	0.71	20

* The summary statistics were calculated for all populations combined (N = 735), except for CQ11. The information for CQ11 refers only to American populations, where the locus is in Hardy-Weinberg equilibrium. For all loci, expected heterozygosity (H_e) was calculated as $1 - \sum p_i^2$, where $\sum p_i^2$ is the sum of the squared frequencies of alleles i . Subscripts after the name of each locus refer to membership to one of three multiplexes.

lowed by NEIGHBOR and CONSENSE in Phylip 3.573c²⁶ with 1,000 replicate bootstraps. Trees were assembled with TreeViewPPC.²⁷ Furthermore, we assigned individuals to clusters (strains) based on their multilocus genotypes with a maximum likelihood algorithm implemented in the program Structure 2.0.²⁸ We used 20,000 burn-in steps and 1,000,000 runs with a model of uncorrelated allele frequencies allowing admixture ($\gamma = 0.34$, calculated at $K = 1$).²⁹ In this analysis, the origin of each specimen is not disclosed but the number of clusters (K) is decided *a priori* for each run. To assess the consistency of the analysis, we performed an exhaustive comparison of 10 runs at each K scoring the similarity coefficient described by Rosenberg and others.³⁰ Formal estimation of admixture proportions (i.e., calculation of the putative contribution of introductions from different locations to the current Hawaiian populations of *Cx. quinquefasciatus*) was performed with the Markov Chain Monte Carlo method LEADMIX, a superior maximum likelihood based method that allows for more than two source populations and can incorporate admixture due to non-simultaneous migration from different source populations.³¹

RESULTS

We obtained 735 specimens from 28 locations across the world always near or inside human dwellings (Table 1). Although some collecting devices were placed in areas less impacted by humans, those did not yield *Cx. quinquefasciatus*. Of the specimens examined, 70% were adults, but less than 40% of those were collected with light traps or other methods that mostly collect females. Many larvae were reared to adults giving us access to known mixes of males and females. Furthermore, as mentioned before, the microsatellite loci used in this study are not sex-linked.

Even after we redesigned primers,²¹ two of the 12 microsatellite loci used in the analyses (CQ46 and CQ41) had significant heterozygote deficits in several populations (see Tables in Supplemental Material). Thus, they were excluded from the analyses. Neighbor-joining distance trees (Cavalli-Sforza and Edwards chord distance and Nei's genetic distance produced nearly identical trees) using the remaining 10 microsatellite loci showed two well-supported geographically structured groups (Figure 1): a Pacific group and a New World group. The samples from east Africa and those from Southeast Asian locations also clustered together, but samples from both Nigeria (West Africa) and Japan clustered with the New World populations. Because of DNA degradation, we only attempted to amplify three loci (CQ26, CQ29, and CQ41) from the museum specimens. Two specimens (both collected in the 1920s) did not yield products. The remaining three specimens (1919, 1931, 1944) yielded 11 alleles but only the specimen from 1944 had an allele unique to Pacific populations (CQ29-186, Table A3). Table A3 appears online at www.ajtmh.org.

The results of the multilocus genetic structure analysis, which combines all individual multilocus genotypes and separates them into distinct clusters analogous to the hierarchical branching of tree diagrams,³² gave similar results to the distance analysis but with a higher resolution (Figure 2). The similarity of results across 10 replicates at each number of inferred clusters (K) was high (0.9–0.99) for K from 2 to 4. At $K = 2$, the 735 specimens of *Cx. quinquefasciatus* examined cluster into two clearly defined groups: Hawaii plus Old

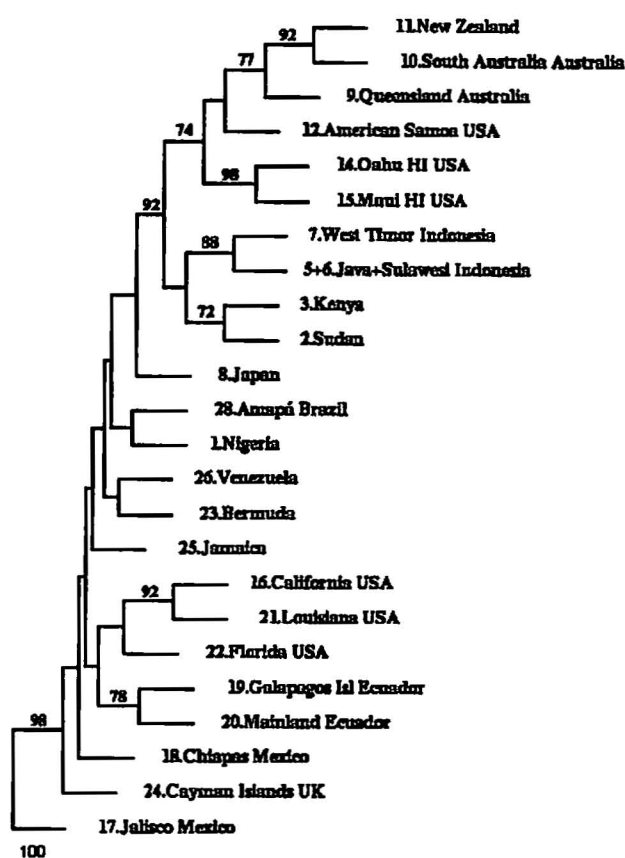


FIGURE 1. Unrooted consensus nearest-neighbor tree depicting the relationships between populations used in this study (to decrease sampling related artifacts, we excluded populations with less than 14 specimens from this grouped analysis). Numbers on branches indicate bootstrap percentages. The numbers before each name correspond to those in Figure 2. All populations group first according to geographic proximity except for Nigeria and Japan, which consistently group with the American populations. Isl = Islands.

World and New World, although similar to earlier findings in this report, Nigeria has New World ancestry. At $K = 3$, Austral and Pacific specimens separate from the Old World cluster, while Brazilian, Nigerian, and Japanese specimens have a mixed signature (Old and New World). Finally, at $K = 4$, populations along the Atlantic Coast of the Americas, as well as Japan and Nigeria, all separate into a distinct group. Throughout the analysis, Hawaiian populations cluster with Australia and New Zealand although Maui has more specimens with New World ancestry than does Oahu.

Admixture analyses showed that Hawaiian populations have an uneven mixture of alleles from New World and Old World populations (less than 20% input from New World populations), which agrees with our other analyses that assign a predominantly South Pacific ancestry to current Hawaiian populations. The concordance between the different analyses is not surprising because a few microsatellite alleles that are common in the Americas (e.g., qGA12-157) occur in Hawaiian populations but are absent from the South Pacific (Tables A1–A11), while many alleles common in the south Pacific and also common in Hawaii are conspicuously absent from the Americas (e.g., CQ26-220, CQ29-186). American popula-

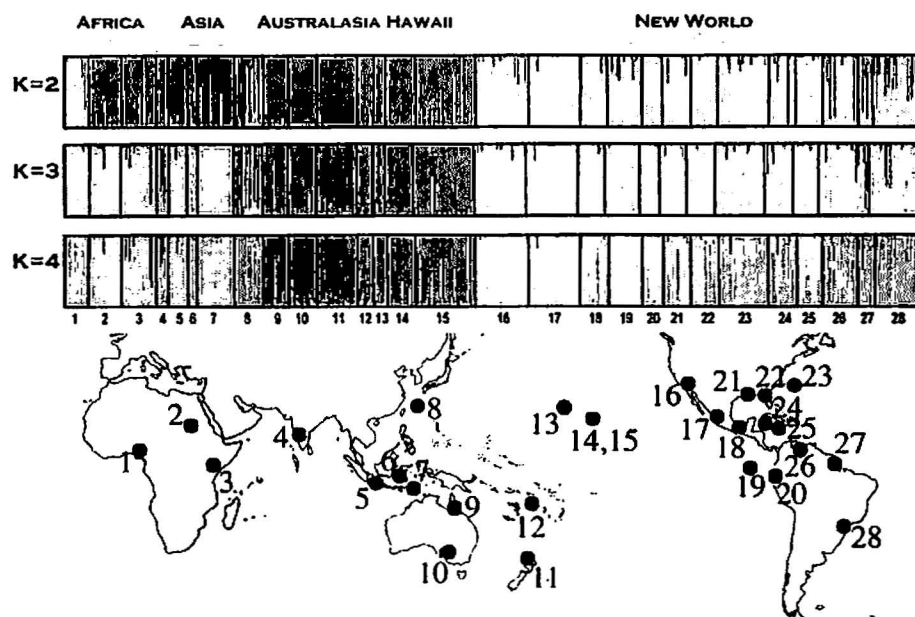


FIGURE 2. Results of a Bayesian cluster analysis of multilocus microsatellite genotypes. Each of the 735 individuals included in the analysis is represented by a thin vertical line, partitioned into colored segments that represent the individual's probability of belonging to one of each of the genetic clusters. Although the origin of each specimen is not used in the analysis, in this figure specimens were grouped by location (separated by a vertical line). The geographic locations of all samples with associated location numbers are shown in the world map and are the same as in Figure 1. Included are two populations with samples of less than 14 specimens: India ($n = 9$), Midway Island ($n = 10$), and Amapá, Brazil ($n = 13$), which were excluded from the distance analysis. Their location numbers are 4, 13, and 27, respectively.

tions, especially those on the Pacific Coast, had significantly ($P < 0.05$) lower allelic richness than Old World populations. The highest allelic richness was found in Asian and east Africa populations ($P < 0.05$) and the lowest was found in west Africa.

DISCUSSION

Although we were able to obtain specimens from multiple locations in some countries but not from others, those events are randomly distributed across the samples and do not reflect the allelic richness encountered. For example, in both Nigeria and Kenya, specimens came from a range of collection sites but those two locations in our study have the lowest and one of the highest allelic richness, respectively. The similarity between Hawaiian and South Pacific populations is unexpected (based on historical accounts) and significant because South Pacific populations of *Cx. quinquefasciatus* are adapted to cold southern hemisphere environments. In New Zealand, where winter temperatures are occasionally below their putative survival minimum, larvae are found from July through September, the southern winter months.³¹ The introduction and genetic swamping of such populations could explain the apparent recent expansion of *Cx. quinquefasciatus* to higher elevations in Hawaii.^{7,34} The fact that Maui contains many more admixed specimens argues that the non-Australasian introduction occurred when Maui was the main port of entry to Hawaii (in the 1800s), which supports the original accounts of the first introduction of mosquitoes. These analyses also indicate that most of the dozens of *Cx. quinquefasciatus* arriving monthly in Oahu in aircrafts from both Asia and the Americas¹⁵ do not reproduce, an important

observation from a control standpoint. The presence of a unique South Pacific allele only in the specimen from 1944 gives us a date by which we know South Pacific *Cx. quinquefasciatus* had already arrived in Hawaii, although the small number of specimens available does not allow us to reject the possibility they had arrived earlier.

It is not known when *Cx. quinquefasciatus* arrived in Australia. Marks argued that it arrived with or shortly after the colonial First Fleet in 1788,^{1,36} while others have attributed its arrival to the opening of Australian ports to American whalers in 1831.³⁷ Its introduction to New Zealand appears to be recent because there are suggestions that it is just starting to penetrate inland.³⁸ Supporting the idea that an Australian strain of *Cx. quinquefasciatus* could have remained localized, the introduction of *Ae. australicus* (a species native to Australia) to New Zealand is well documented and occurred only during the last 50–80 years.³⁷ The expansion of *Cx. quinquefasciatus* to the smaller Pacific islands is thought to be even more recent and linked to events during World War II.¹³ Such introductions are supposedly linked to the increased connectivity between Australia and the Pacific islands because of the intense traffic of whaling boats and later passenger airplanes and warships.³⁷

The low diversity of New World populations of *Cx. quinquefasciatus* agrees with assertions that it is a recently introduced species. So does its only recent arrival to the Galapagos Islands and the similarity between mainland Ecuador and Galapagos specimens. However, the source of New World *Cx. quinquefasciatus* is unclear. Its arrival from west Africa as claimed³⁹ is unlikely because the species was reported absent there before 1942.³⁰ The similarity between Nigerian and American populations instead indicates that the former were

introduced from the New World, which is supported by the even lower allelic richness of Nigerian specimens. The differences between Pacific and Atlantic coasts in the New World are consistent across latitude (Figure 2) and close examination of the allelic frequencies (Tables A1–A11) and the higher allelic richness in Atlantic Coast populations supports the hypothesis that the extensive boat traffic across the Caribbean and the Atlantic may have led to extensive mixing.

East African and Asian populations cluster together (Figure 2). Although the heavy human traffic across the Indian Ocean might be the reason, if, as it has been proposed, *Cx. quinquefasciatus* originated in Africa,⁴¹ African populations would have ancestral polymorphism. Instead, both the east African and Asian populations we examined had the highest allelic richness across all populations examined. The very different signatures of east and west African populations, as well as the uniqueness of the interaction between *Cx. quinquefasciatus* and its sibling species *Cx. pipiens* in Africa,⁴² hint at a complex origin and distribution of the species there and will require further sampling in the continent. Critically, our findings agree with the epidemiology of nocturnal filariasis described earlier and are supported by vector competence studies showing low susceptibility of *Cx. quinquefasciatus* from west Africa and Polynesia to *W. bancrofti*.^{43,44}

The presence of distinct strains of *Cx. quinquefasciatus* across the world was unexpected because previous studies examining loci involved in insecticide resistance concluded this species is undergoing a human-aided expansion with non-trivial levels of gene flow between populations.⁴⁵ Our results, however, show that recombination may break the connection between selected and neutral loci very quickly, maintaining the integrity of the microsatellite signature while allowing the penetration of useful insecticide resistance alleles. This result is a critical example of cryptic introgression of useful genes, which may be a common phenomenon with very broad consequences.⁴⁶

In conclusion, we found that worldwide populations of *Cx. quinquefasciatus* are significantly genetically differentiated and correlated to known *W. bancrofti* vector competence, and their pathways of expansion are remarkably similar to those inferred for *Ae. aegypti*, another vector species associated with humans.² Furthermore, we found there has been at least a second introduction of *Cx. quinquefasciatus* into Hawaii. This conclusion is significant because changes in the dynamics of avian malaria in Hawaii, especially the perceived increase in the altitudinal range of the mosquitoes^{7,34} as well as changes in parasite virulence,⁴¹ may be associated with this secondary introduction. We are currently examining differences in vector competence to avian malaria of the various genetic strains. Multiple introductions may be a fundamental evolutionary agent in invasive species⁴⁷ and in disease vectors they may impact important epidemiologic parameters.

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AA CQ29	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28
168				0.1	0.2	0.3	0.4						0.3			0	0.2	0		0	0	0			0.1			
174	0.3				0.2		0.1					0				0.1	0		0.1	0.1	0	0.3	0.2	0	0.1		0	0.2
176	0	0.2	0.6	0.3	0.1	0		0.1		0.1	0.2	0.1		0.2	0.2		0	0	0	0.1	0.1		0.1		0.1	0.4		0.5
178	0.7	0.8	0.4	0.6	0.6	0.7	0.5	0.9	0.8	0.8	0.5	0.8	0.2	0.5	0.5	0.9	0.8	0.9	0.8	0.9	0.9	0.7	0.7	0.9	0.7	0.6	1	0.3
181		0																										
186			0.1						0.2	0.1	0.2	0.1	0.6	0.3	0.3									0				
188													0	0														
Ho	0.3	0.44	0.57	0.78	0.27	0.38	0.59	0.13	0.42	0.36	0.71	0.29	0.6	0.72	0.62	0.28	0.43	0.13	0.28	0.18	0.25	0.5	0.38	0.09	0.42	0.75	0.08	0.61
He	0.44	0.36	0.54	0.6	0.45	0.59	0.6	0.19	0.34	0.34	0.61	0.43	0.58	0.64	0.61	0.25	0.38	0.12	0.27	0.17	0.23	0.44	0.45	0.17	0.44	0.49	0.08	0.64
P																							‡	‡				
A4																												
CQ41	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28
117						0.1																						
120							0.1																					
137		0.1		0.1																								
139	0.2		0											0	0													0.3
140			0																									0.1
141	0.1	0.1	0.2	0.3	0.3	0.1	0.1	0.3		0	0.1	0.5	0.2			0.1	0.8	0.2	0.7	0.8	0.2	0	0.6	0.6	0.4	0.8	0.6	0.3
143	0.2		0	0.2	0.1	0.2	0.1	0.2	0.2	0.1	0.3	0.3		0	0	0.1	0.1	0.5	0.3	0.1	0.2	0.7	0	0.2	0	0.1	0	0.1
144																												
145		0.1			0.1					0		0	0.1															
146									0.2																			
147	0.4	0.3	0.2	0.1	0.3	0.6	0.1	0.3	0.1	0.2	0.2	0	0.7	0.5	0.7	0.6	0	0			0.6	0.3	0.1	0		0.1	0.2	
148																												
149			0																									
150	0.1	0.4	0	0.1	0.1		0.1	0.1						0.1	0													0
152		0	0.1						0.5	0.6	0.4	0.1	0.2	0.2	0.2	0.1												
154					0.1		0							0	0													
155																							0.1					
156	0	0	0.2			0.1	0.1							0.1	0		0	0.2			0		0.1		0.2	0.4		
157				0.1							0.1					0					0		0.1					
158																										0.2		
159	0.1		0		0		0.1							0.1	0								0					
160				0.1																					0			
161	0.1		0		0		0.1									0												
163			0																									
165							0.1																					
167							0																					
Ho	0.35	0.63	0.71	0.78	0.79	0.63	0.65	0.83	0.79	0.56	0.71	0.64	0.4	0.58	0.38	0.24	0.23	0.55	0.34	0.24	0.17	0.5	0.45	0.45	0.41	0.41	0.46	0.63
He	0.82	0.74	0.85	0.84	0.85	0.67	0.91	0.73	0.7	0.56	0.73	0.67	0.56	0.7	0.51	0.62	0.36	0.63	0.44	0.4	0.57	0.42	0.66	0.6	0.71	0.35	0.52	0.75
P	‡						‡								‡	‡	‡			‡	‡	‡	‡	‡	‡	‡	‡	
A5																												
aG12	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28
91			0																									
111										0																		
113	0.1	0.2	0.1	0.1	0.2	0.3	0.4	0.2	0	0.1	0.1		0.1	0.1	0	0.3	0.2	0.4		0.1	0.2	0.4	0.2	0.3	0.3	0.5	0	0
115	0.8	0.1	0.4	0.5	0.6	0.4	0.4	0.4	0.2	0	0.3	0.1	0.4	0.2	0.1	0.1	0.2	0	0.1		0.1	0	0.4	0.1	0.2	0	0.4	0.3
117	0	0.5	0.4	0.2	0.2	0.3	0.2	0.2	0.3	0.7	0.4	0.3	0.2	0	0.1	0.2	0.2	0.3		0.7	0.6	0.3	0.1	0.1	0.1	0.1	0.2	0.2
119								0.2					0.1															
121																									0			
123									0.4	0.1	0.1			0	0													

TABLE A3
Continued

A3 CO29	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28
125			0	0.1					0	0	0				0.1	0		0.1	0.1		0.1	0	0		0		0	0
127	0.1			0.1	0					0	0	0.3	0.2	0.2	0.3	0.4	0.2	0.2	0.1	0.3	0.3	0.3	0.2	0.3	0.4	0.3	0.5	0.3
129		0	0			0						0.3				0.1		0.1		0	0					0	0.1	
131			0	0.1													0.2						0.1		0.2	0		
133														0										0				
135														0.1	0								0					
137											0	0.2	0.2	0.2								0.1						
139		0.2	0											0.1	0.1													
141														0	0													
143															0													
145															0													
He	0.44	0.52	0.7	0.57	0.63	0.14	0.31	0.25	0.86	0.56	0.8	0.86	0.8	1	0.9	0.58	0.73	0.92	0.34	0.44	0.67	0.75	0.77	0.7	0.86	0.83	0.69	0.93
He	0.38	0.69	0.71	0.74	0.7	0.69	0.58	0.67	0.7	0.49	0.72	0.75	0.82	0.86	0.85	0.73	0.81	0.76	0.55	0.59	0.78	0.74	0.76	0.79	0.76	0.64	0.63	0.75
P						‡	‡									‡			‡									
A6 qGT4	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28
141																												
145		0.1																			0					0		
147		0								0.1	0.2					0.1	0		0		0.1							0.2
149	0.1	0.6	0.3	0.3	0.4	0.4	0.8	0.6	0.7	0.6	0.5	0.2	0.4	0.3	0.2	0.3	0.7	0.3	0.1	0.3	0.4	0.7	0.6	0.4	0.7	0.4	0.5	0.3
151	0.8		0.2	0.2	0.3	0.4	0.2	0.1	0.3	0.2	0.1	0.6	0.6	0.7	0.7	0.6	0.2	0.6	0.6	0.4	0.5	0.3	0.4	0.4	0.1	0.5	0.3	0.3
153	0.1	0.3	0.4	0.3	0.2				0	0.1	0.1	0.2	0.1	0	0.1	0	0	0.1	0.2	0.2	0.1	0.1	0	0.2	0.2	0.1	0.1	0.1
155				0.1	0						0.1								0					0			0	0
157			0.1	0.1	0.2	0.1		0.3																				
159						0				0.1													0					0
161																												0
He	0.35	0.48	0.83	0.6	0.63	0.33	0.75	0.75	0.33	0.6	0.71	0.5	0.3	0.48	0.42	0.67	0.45	0.63	0.57	0.69	0.58	0.67	0.53	0.48	0.38	0.59	0.62	0.68
He	0.31	0.52	0.73	0.84	0.63	0.3	0.74	0.7	0.45	0.62	0.72	0.57	0.56	0.46	0.51	0.61	0.47	0.56	0.57	0.66	0.65	0.5	0.52	0.67	0.49	0.62	0.62	0.74
P																								‡				
A7 pGT12	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28
144		0																										
150	0			0.3										0.1	0.2	0.2	0.4	0	0.1	0.4	0.2	0.1	0.2	0.2	0.4		0	
152		0.1	0.2	0.2	0.1		0.1		0.5	0.3	0.3	0.1			0	0		0			0			0	0	0.1	1	0.2
154			0																									
156																												
158		0	0		0.1	0.5	0.2	0	0.2	0.1	0.3		0.1	0	0			0.2							0.1		0	
160	1	0.9	0.7	0.5	0.8	0.5	0.8	1	0.3	0.6	0.3	0.9	1	0.9	0.8	0.7	0.6	0.7	0.9	0.6	0.8	0.9	0.8	0.8	0.5	0.9	0	0.8
176																0												
He	0.05	0.26	0.39	0.44	0.63	0.38	0.19	0.09	0.81	0.32	0.66	0.29	0.1	0.24	0.43	0.36	0.5	0.61	0.17	0.53	0.46	0.21	0.4	0.21	0.62	0.17	0.08	0.31
He	0.05	0.27	0.4	0.66	0.52	0.43	0.39	0.09	0.61	0.55	0.68	0.25	0.1	0.25	0.39	0.41	0.49	0.48	0.16	0.49	0.38	0.19	0.35	0.32	0.6	0.21	0.08	0.37
P																												
A8 qGT8	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28
77											0																	
83										0	0					0				0.1								
85																				0								
87	0.6	0.1	0.3	0.3	0.1	0.3	0.3	0.3	0.4	0.5	0.2	0.3		0.2	0.2	0.8	0.5	0.6	0.4	0.6	0.6	0.8	0.5	0.4	0.5	0.5	0.3	0.3
88		0.2	0.1	0.2	0	0				0.3	0.2	0.5	0.2	0.4	0.3								0.1	0	0.1		0.3	0.2
89		0	0				0.2	0.2	0.1	0.2	0.1	0.4	0.2	0	0.1	0	0.5	0.2	0.2	0.3								
90					0.2	0.1	0.1	0.1					0.1						0									

TABLE A3
Continued

A8 qGT8	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27		
91	0.4	0.1	0.3	0.3	0.5	0.4	0.1	0.3	0	0		0		0	0	0.1		0.2	0.2	0.2	0.1	0.1	0	0.3	0.3	0.3	0.2	0.3	
93																													
95																			0										
97												0																	
98																													
99	0.1															0.1	0				0.1		0		0.1	0.1		0.1	
103																													
105						0	0.1							0.1	0						0								
107			0.1	0.2	0	0.1	0.1	0.2	0.1		0		0.4	0.3	0.4	0					0.1	0.1	0.1	0.1		0.1			
109			0		0.1	0.1	0.1							0	0								0.2		0.1		0	0	
110					0																								
111			0.2												0														
113			0.3							0	0.2												0				0.1	0.1	
115			0																										
119																													
120																						0	0	0	0.1				
122																													
Ho	0.55	0.82	0.71	0.89	0.81	0.88	0.75	0.92	0.68	0.6	0.82	0.71	0.4	0.76	0.75	0.39	0.52	0.46	0.76	0.44	0.54	0.46	0.67	0.68	0.52	0.76	0.85	0.86	
He	0.53	0.83	0.79	0.78	0.76	0.87	0.7	0.78	0.71	0.68	0.71	0.7	0.74	0.71	0.73	0.42	0.52	0.53	0.72	0.6	0.57	0.39	0.74	0.72	0.7	0.67	0.74	0.75	
P																													
A9 qGT7	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	
126										0.1																			
128		0.2	0.1	0.2		0.1		0.2	0.1		0	0	0.4			0.1													
130	0.8	0.2	0.2	0.4	0.2	0.2	0.3	0.7	0.2	0.4	0.3	0.5	0.3	0.6	0.4	0.3	0.6	0.5	0.3	0.7	0.5	0.3	0.3	0.5	0.7	0.4	0.7	0.4	
132	0.1	0.4		0.1	0.1	0.3	0.3	0.1	0.3	0.1	0.5		0.1	0.1	0.1	0.3	0.1	0.2	0.2	0.1	0.2	0.4	0.5	0.1	0.1	0.2		0.3	
134		0	0.3	0.2	0.2	0	0.1		0	0.2	0.2	0.2	0.2	0.1	0.2					0	0				0				
136		0.1	0.4	0.1	0.3				0.2																				
138					0.1														0										
140	0.2	0	0.1		0	0.1	0.1	0								0.2	0.1	0.2	0.4	0.1	0.2	0.2	0.1	0	0.1	0.1	0.1	0.3	
142					0		0.1							0	0				0	0				0		0.2	0.1		
144			0				0.1	0	0.1	0.2	0	0.3	0.2	0.2	0.2	0.1	0.2	0.1		0.1	0.1	0.1	0.1	0.3	0.1	0.2	0		
146																													
148						0																							
150						0																							
152						0																							
154							0.1																						
156						0.1																							
158					0.1	0																							
162					0																								
164						0.1																							
166						0.1																							
170											0																		
Ho	0.4	0.46	0.67	0.5	0.81	0.88	0.78	0.54	1	0.6	0.53	0.64	0.5	0.52	0.56	0.65	0.55	0.54	0.76	0.56	0.71	0.83	0.67	0.59	0.52	0.72	0.46	0.71	
He	0.41	0.74	0.74	0.81	0.85	0.88	0.84	0.47	0.8	0.75	0.65	0.67	0.79	0.63	0.71	0.74	0.58	0.65	0.68	0.55	0.71	0.73	0.69	0.59	0.48	0.75	0.53	0.7	
P																													
A10 pGT40	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	
256																0.1		0.1		0.1	0.2	0.1	0.1						
261	0.1	0.1	0.1	0.1	0.3	0.3	0.3		0.4	0.2	0.2	0		0.1	0.2	0.6	0.5	0.5	0.3	0.5	0.4	0.2			0.3	0.2	0	0.2	0.4
262		0.3		0.3		0.2	0.1	0.2	0.2	0.2	0.5	0.4	0.5	0.1			0	0						0				0	

TABLE A3
Continued

Alto pG146	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27
263	0	0.2	0.1	0.2	0	0.1	0.2		0.2	0.4	0.1	0.2	0.2	0.2	0.2	0	0	0.1			0		0.2	0	0	0.4	0.7
264	0.1	0.2	0	0.2	0.1	0	0.1		0.2	0.2	0.1	0.2	0.2	0.3	0.3	0	0	0.1			0		0.2	0	0	0.4	0.7
265	0.7	0.3	0.3			0.1		0.4	0	0.2	0.1	0.2	0.4	0.3	0.3	0.2	0.4	0.3	0.3	0.4	0.4	0.7	0.6	0.6	0.7	0.6	0.1
266	0.1	0.2	0.3	0.3	0.5	0.2	0.4	0.4			0.4	0.4	0	0	0	0.1	0	0	0.4	0.1	0	0.1	0	0	0	0	0.2
267	0.1	0.2	0.3	0.3	0.5	0.2	0.4	0.4				0.4				0											0
268	0.1	0.1														0							0				0
270	0															0											0
271																0											0
274	0															0											0
286																0											0
288						0	0	0.1																			0
290						0																					0
H ₀	0.5	0.78	0.58	0.67	0.81	1	0.55	0.77	0.73	0.84	0.58	0.71	0.6	0.64	0.77	0.59	0.52	0.65	0.66	0.8	0.58	0.46	0.63	0.46	0.43	0.83	0.38
He	0.46	0.78	0.82	0.88	0.82	0.8	0.67	0.66	0.73	0.74	0.71	0.7	0.7	0.75	0.74	0.61	0.54	0.64	0.67	0.63	0.67	0.52	0.53	0.53	0.51	0.56	0.53
<i>P</i>																											
Alto pG146	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27
86	0.4	0.3	0.1	0.1	0.7	0.3	0.3	0.2	0.1	0.1	0.2	0.2	0.4	0.2	0.4	0.3	0.3	0.5	0.3	0.4	0.2	0.4	0.2	0.4	0.3	0.5	0.3
88																											
90																											
92																											
108																											
110	0	0.3	0.1						0.1	0.1	0.1	0.2	0.2	0.2	0.1	0.1	0	0.1	0.1	0	0.1	0	0	0	0.4	0.1	0.2
112	0.1	0	0.1	0.1	0	0.1	0.1	0.3	0.2	0.2	0.2	0.2	0.2	0	0	0.1	0.3	0.1	0	0.1	0.1	0.1	0.1	0	0.1	0.2	0.1
114	0	0.2	0.1	0.1	0	0	0.1	0	0	0	0.1	0.1	0.1	0.1	0.1	0.2	0.2	0.2	0.2	0.1	0.1	0.3	0.3	0.2	0.2	0.2	0.1
116	0.2	0.2	0.2	0.3	0	0	0.1	0.1	0	0.5	0.4	0.4	0.5	0.5	0.5	0.3	0.4	0.3	0.2	0.4	0.5	0.3	0.7	0.5	0.1	0.1	0.1
118	0.3	0.2	0.1	0.4	0.1	0.3	0.2	0.5	0.4	0	0	0	0	0	0	0.3	0.1	0.1	0.2	0.2	0.4	0.5	0.7	0.5	0.1	0.1	0.2
120	0.1	0.1	0.1		0.1	0.1	0.1	0.1	0	0	0	0	0	0	0	0	0.1	0	0	0	0	0	0	0	0	0	0
122					0.1	0.1	0.1	0.1							0												
124						0.1	0.1																				
126		0				0.1	0.1		0				0.1	0	0											0.2	0
127							0.1		0.1				0.1	0	0												0
128					0	0.1			0	0	0																
130					0	0																					
134					0	0																					
136																											
140				0.1							0																
H ₀	0.7	0.70	0.74	0.67	0.75	1	0.5	0.96	1	0.72	0.77	0.64	0.5	0.64	0.66	0.8	0.8	0.71	0.76	0.76	0.71	0.83	0.36	0.42	0.68	0.67	1
He	0.75	0.79	0.83	0.79	0.77	0.88	0.47	0.7	0.77	0.67	0.79	0.76	0.62	0.65	0.61	0.78	0.71	0.63	0.77	0.7	0.73	0.7	0.51	0.58	0.7	0.71	0.82
<i>P</i>																											

* 0 = 0.001, He and H₀ in expected and observed heterogeneity, respectively. *P* is the significance level of the difference between He and H₀.
 † *P* = 0.05
 ‡ *P* = 0.01