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SPECIALTY SECTION

This article was submitted to
Infectious Disease Epidemiology,
a section of the journal
Frontiers in Epidemiology

RECEIVED 24 July 2022

ACCEPTED 16 September 2022

PUBLISHED 13 October 2022

CITATION

Yaro AS, Linton Y-M, Dao A, Diallo M,
Sanogo ZL, Samake D, Ousmane Y,
Kouam C, Krajacich BJ, Faiman R,
Bamou R, Woo J, Chapman JW,
Reynolds DR and Lehmann T (2022)
Diversity, composition, altitude, and
seasonality of high-altitude windborne
migrating mosquitoes in the Sahel:
Implications for disease transmission.
Front. Epidemiol. 2:1001782.
doi: 10.3389/fepid.2022.1001782

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Sanogo, Samake, Ousmane, Kouam,
Krajacich, Faiman, Bamou, Woo,
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Diversity, composition, altitude, and seasonality of high-altitude windborne migrating mosquitoes in the Sahel: Implications for disease transmission

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Recent studies have reported *Anopheles* mosquitoes captured at high-altitude (40–290 m above ground) in the Sahel. Here, we describe this migration modality across genera and species of African Culicidae and examine its implications for disease transmission and control. As well as *Anopheles*, six other genera—*Culex*, *Aedes*, *Mansonia*, *Mimomyia*, *Lutzia*, and *Eretmapodites* comprised 90% of the 2,340 mosquitoes captured at altitude. Of the 50 molecularly confirmed species ($N = 2,107$), 33 species represented by multiple specimens were conservatively considered high-altitude windborne migrants, suggesting it is a common migration modality in mosquitoes (31–47% of the known species in Mali), and especially in *Culex* (45–59%). Overall species abundance varied between 2 and 710 specimens/species (in *Ae. vittatus* and *Cx. perexiguus*, respectively). At altitude, females outnumbered males 6:1, and 93% of the females have taken at least one blood meal on a vertebrate host prior to their departure. Most taxa were more common at higher sampling altitudes, indicating that total abundance and diversity are underestimated. High-altitude flight activity was concentrated between June and November coinciding with availability of surface waters and peak disease transmission by mosquitoes. These hallmarks of windborne mosquito migration bolster their role as carriers of mosquito-borne pathogens (MBPs).

Screening 921 mosquitoes using pan-*Plasmodium* assays revealed that thoracic infection rate in these high-altitude migrants was 2.4%, providing a proof of concept that vertebrate pathogens are transported by windborne mosquitoes at altitude. Fourteen of the 33 windborne mosquito species had been reported as vectors to 25 MBPs in West Africa, which represent 32% of the MBPs known in that region and include those that inflict the heaviest burden on human and animal health, such as malaria, yellow fever, dengue, and Rift Valley fever. We highlight five arboviruses that are most likely affected by windborne mosquitoes in West Africa: Rift Valley fever, O'nyong'nyong, Ngari, Pangola, and Ndumu. We conclude that the study of windborne spread of diseases by migrating insects and the development of surveillance to map the sources, routes, and destinations of vectors and pathogens is key to understand, predict, and mitigate existing and new threats of public health.

KEYWORDS

arbovirus, disease-spread, dispersal, malaria, mosquito-borne pathogen, surveillance, Africa, one health

Introduction

High-altitude windborne migration of large insects such as locusts has been recognized over millennia. However, over the past decades, our knowledge of this behavior has greatly expanded thanks to systematic studies using aerial sampling, radar, and other methodologies (1–3). Radar studies have been key to estimate direction, speed, altitude, and magnitude of migrating insects, but they seldom provide species-level information (4). Given the enormous number of insects involved, the distances they cover, and their interactions with other organisms, windborne migration influences food security (3, 5–8), public health (9–14), and ecosystem vigor (15, 16). Although windborne flight at altitude is common in many groups of insects including Diptera (10, 16–20), most studies on mosquito movements have been focused on host location and on movement at a village scale (21–24) under the premise that dispersal of most mosquito species does not exceed 3–5 km, and long-range movements represent rare “accidental events” that are of minimal epidemiological importance (25–27). The term dispersal is commonly used in mosquito literature (25), but the definition of migration as “persistent movements unaffected by immediate cues for food, reproduction, or shelter, with a high probability of relocating the animal in a new environment” (3, 28) is apt here and will be used hereafter.

Recent studies in the Sahel revealed that *Anopheles* mosquitoes, like other windborne insect migrants, regularly engage in high-altitude windborne migration: e.g., 40–290 m above ground level (agl), between July and November (14, 29, 30). Furthermore, gravid females (i.e., females with mature eggs, following a blood meal) predominated among mosquito migrants. The number of individuals that cross a 100 km line

perpendicular to the wind at altitude was found to range between tens of thousands to millions per year, depending on mosquito species. Finally, aided by winds at altitude, these mosquitoes cover tens to hundreds of kilometers per night (assuming up to 9-h nightly flights). These findings pertain to Sahelian *Anopheles* species and were interpreted in the context of malaria transmission and control. Here, we expand the scope of our first analysis (14) and include the full collection of mosquitoes that were intercepted at altitude over the Sahel to better assess the implications of this migration modality in mosquitoes to public health and ecosystem stability in West Africa. Specifically, we identify the species intercepted at altitude, and for each species estimate their sex ratio and gonotrophic state composition, mean flight altitude, and seasonality – all parameters that impinge on risk of infection with mosquito-borne pathogens (MBPs) (31). We report on the initial screening of these mosquitoes to infection with species of *Plasmodium*, which infect a range of vertebrate hosts. Focusing on the windborne mosquito species that have previously been reported as vectors of MBPs in West Africa, we rank both the risk of the pathogens for windborne spread and the mosquito species in terms of their total contributions to overall MBP spread. Our results indicate that windborne migration of mosquitoes is a key driver shaping the epidemiology of many mosquito-borne diseases and that this newly recognized aspect of the vector and MBP ecology should be considered in predicting disease spread as well as in planning disease control strategies.

Materials and methods

The study area has been described in detail previously (32–35), as has the field and most laboratory methods used in this

study (14, 30, 36). Below is a brief description of the field and laboratory operations previously reported and a detailed description of other procedures. Aerial sampling stations were placed in four Sahelian villages with traditional mud-brick houses, surrounded by fields, set in a dry savanna, consisting of grasses, shrubs, and scattered trees: Thierola (13.6586, – 7.2147) and Siguima (14.1676, – 7.2279) were sampled between March 2013 to November 2015; Markabougou (13.9144, – 6.3438) sampled between June 2013 to June 2015, and Dallowere (13.6158, – 7.0369) sampled between July to November 2015. Sampling was not carried out in January and February. Over 90% of the rains fall in the wet season (June–October, ~550 mm annually), forming temporary puddles and pools that usually dry by November. Rainfall during the dry season (December–May) is negligible (0–30 mm) (37).

Aerial insect sampling was conducted using sticky nets (1 × 3 m panels) attached to the tethering line of 3 m diameter helium-filled balloons, with each balloon typically carrying three panels. Initially, panels were suspended at 40, 120, and 160 m agl, but after preliminary results showed higher mosquito panel densities at higher elevations (from August 2013), the typical altitude was 90, 120, and 190 m agl. When a larger balloon (3.3 m dia.) was deployed at Thierola (August–September 2015), two additional panels were added at 240 and 290 m agl. Balloons were launched ~1 h before sunset (~ 17:00) and retrieved 1 h after sunrise (~ 07:30), the following morning. To control for insects trapped near the ground as the panels were raised and lowered, comparable control panels were raised up to 40 m agl and immediately retrieved during each balloon launch and retrieval operation. Between September and November 2014, the control panels were raised to 120 m agl. Following panel retrieval, inspection for insects was conducted in a dedicated clean area. Individual insects were removed from the nets with forceps, counted, and stored in labeled vials containing 80% ethanol.

Mosquitoes were morphologically identified to genus, sex, and gonotrophic state before DNA was extracted from whole body of males and separately from abdomen and thorax/head of female mosquitoes. Molecular barcode identification of mosquito species was carried out by PCR amplification of 658 bp of the mitochondrial COI gene, amplicons were sequenced, and sequences compared with public databases: BOLD and GenBank. Targeted pathogen analysis was performed on head-thorax extractions of females to detect avian and mammalian plasmodia, using 18S RNA pan-*Plasmodium* assay (38, 39). Sequencing results will be reported elsewhere (Bamou et al.: unpublished).

Data on vectorial status of the mosquito species at altitude was extracted from an extensive literature search summarized by (40). Here, we included only mosquito species that were reported as probable or likely vectors of pathogens in West Africa based on the references listed in (Supplementary Table S3). For example, our literature records showed that *Ae. mcintoshi*

vectors at least 10 MBPs in Africa, but this species was conservatively excluded from the present analysis because no record pertained to West Africa. West Africa was defined to include the following countries: Benin, Burkina Faso, Gambia, Ghana, Guinea, Guinea Bissau, Ivory Coast, Liberia, Mali, Mauritania, Senegal, Sierra Leone, Togo (40). The evidence used by these sources to implicate vectors includes one or more of the following findings as shown in Supplementary Table S3: (a) multiple isolations of a pathogen from wild vector from whole body or midgut [Win], (b) from salivary glands/saliva [WIs], (c) from legs [WIL], (d) from Thorax and Head [Wlth], (e) detection of pathogen by circumsporozoite ELISA [WlC], (f) laboratory experiments demonstrating vectorial competence (physiological capacity to support the pathogen to the infectious stage and capacity to transmit to a new host, following blood feeding on infectious blood meal), provided the mosquito naturally feeds on relevant hosts [LF], (g) transovarial transmission, provided the mosquito naturally feeds on relevant hosts [TF], (h) synchronous appearance of vector that feeds on relevant hosts during outbreak/peak transmission [Epi]. Frequently vectors were incriminated by multiple independent studies with either the same or different criteria. The use of “primary” and “secondary” vectors is widespread in the literature, attesting for its utility (40–42). Despite being vaguely defined, primary vectors are often those that are responsible for the largest share of the transmission (in a particular region). Unless a mosquito species was explicitly indicated as primary vector for that MBP, it was conservatively assumed to be a secondary vector.

Data analysis

Heterogeneity across species in categorical traits such as sex or exposure to vertebrate blood was evaluated with contingency likelihood ratio chi-square test. If the fraction of cells with expected counts <5 were >20%, we used exact tests based on Monte Carlo simulation of 10,000 samples using Proc Freq (43).

To estimate a species' mean flight altitude given our panel heights, we used a weighted mean across panel altitudes, whereby each altitude was weighted by the ratio of the proportion of specimens collected at that altitude over the proportion of nights that panel altitude was used throughout the study. The weighed estimate accommodates variation in panel altitude over the course of the study due to systematic changes (i.e., eliminating panel heights of 40 m after the first year of the study) and opportunistic changes (i.e., adding panels at 240 and 280 m when a larger balloon and abundant helium supply were available). If mosquitoes were equally likely to be sampled across all heights, then the ratio would be 1; however, if mosquitoes were more (or less) likely to be caught at particular panel heights then the ratio would deviate accordingly.

To estimate seasonality in high-altitude flight, panel densities of each species (zeros included) were averaged for each month over data across Sahelian stations and years. The log of the monthly mean multiplied by 1,000 (to form integers) was used as a relative index of monthly flight activity accommodating the large variation between species.

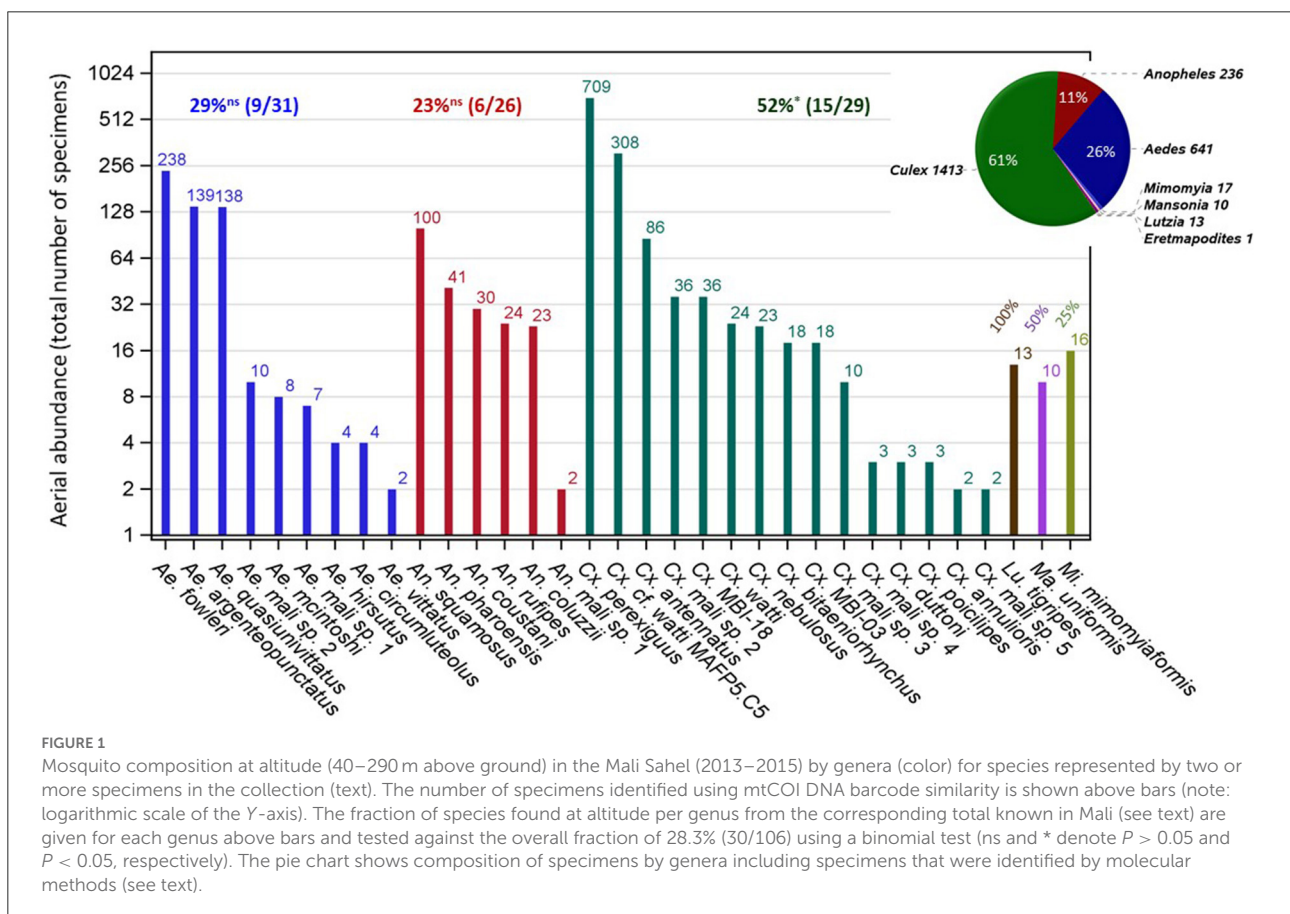
The overall risk of a given pathogen to be dispersed at altitude by a mosquito species depends on the abundance of the mosquito at altitude and on its likelihood to carry that pathogen. The logarithm of the total aerial abundance was used as a measure of windborne presence for each mosquito species, and we considered the ratio of 10:1 as the typical difference between primary and secondary vector in their rate of infection with a particular MBPs (see below for sensitivity analysis). The sum of the risk estimates for each pathogen across windborne mosquito vectors (when more than one vector species was indicated for that pathogen) was used to rank pathogens as to their overall risk of windborne spread, and a similar approach was applied to rank the relative contribution of each mosquito species to the overall windborne spread of MBPs. To address the uncertainty and natural variance in differential weights between primary and secondary vectors and assess the sensitivity of the results to error in these weights, we also used estimates that span the range of these weights. We considered a ratio of 4:1 as the minimum differential rate (in rates of infection) in keeping with

the definition of primary and secondary vectors. Because typical sample size per species in each study ranges between a few hundreds and a few tens of thousands, the maximum differential rate would be near 10,000:1 (in rates of infection). Because we aim to rank mosquito borne diseases (MBDs) in term of risk for airborne spread by mosquitoes, we applied and compared their ranking based on these three sets weights: 4:1, 10:1, and 10,000:1 (Supplementary Results and discussion) as a sensitivity analysis to address the uncertainty associated with difference between primary and secondary vectors and its effect on our results.

Results

The composition of high-altitude mosquitoes in the West African Sahel

A total of 2,340 culicines and 236 anophelines were identified by morphology among the insects collected at altitude during 617 sampling nights on 1,894 sticky nets (panels) in the Sahel of Mali. No mosquitoes were collected on 508 control panels that were raised to 40–100 m agl and immediately retrieved upon launch and retrieval of the standard panels (14, 30), showing that the samples were not contaminated by mosquitoes flying near the ground



during the very short duration launching and retrieval procedure. Based on mitochondrial COI sequence similarity, 2,331 specimens (90.5%) were molecularly assigned to seven genera: *Culex* (1,413), *Aedes* (641), *Anopheles* (236), *Mimomyia* (17), *Lutzia* (13), *Mansonia* (10), and *Eretmapodites* (1), and 2,107 mosquitoes (81.8%) were assigned to 50 species (Supplementary Table S1). Species represented by a single specimen ($N = 17$, Supplementary Table S1) probably fly at altitude in lower numbers. To exclude accidental capture, however, we confined our analyses to the species that were represented by at least two specimens in our collection (Figure 1, Supplementary Table S1), conservatively assuming that only 33 mosquito species, in six genera are high-altitude migrants. With 106 mosquito species spanning 11 genera reported from Mali (40, 42, 44), this estimate supports that at least 31% of the Malian mosquito species use this migration modality. Test of homogeneity in this fraction among the three most diverse genera in Mali ($N > 25$) revealed that *Culex* has exhibited higher than expected fraction of species in altitude (52%, $P = 0.0164$, $\chi^2_{[df=1]} = 5.7$, binomial test). *Culex* has also had the largest number of specimens/species (85, Supplementary Figure S1). Generally, the mean number of mosquitoes per species at altitude increased across genera with the total number of species found at altitude ($r = 0.99$, $P < 0.001$, $N = 6$).

Sex ratio and female gonotrophic state composition in high-altitude mosquitoes

Overall, female mosquitoes predominated at altitude across all taxa (85%, $N = 2,219$) with only five species below 75% and five species above 95% (Figure 2, Supplementary Table S2). Variation across taxa was detected ($P < 0.005$, Exact χ^2 test), however, the species showing the largest departures were all associated with small sample size. For example, *Aedes hirsutus* had the lowest female proportion (50%) and a sample size of just $N = 4$, (Supplementary Table S2). Likewise four species, including *Ae. mcintoshi* and *Ma. uniformis* had the highest proportion (100%, $N = 9$, Supplementary Table S2, Figure 2).

Gravid females predominated over all other gonotrophic states combined (Figure 2). Except unfed females, all other gonotrophic states (which must have taken at least one blood meal on a vertebrate host, unless are autogenous) are hereafter referred to as “exposed”. The overall mean proportion of exposed females was 93% ($N = 924$ representing 21 species with sample size of four or more females of known gonotrophic state). No heterogeneity was detected in exposure rate among species ($P > 0.38$, Exact χ^2 test). Accordingly, only species with small sample sizes exhibited the largest apparent departure from that value, e.g., *Aedes*

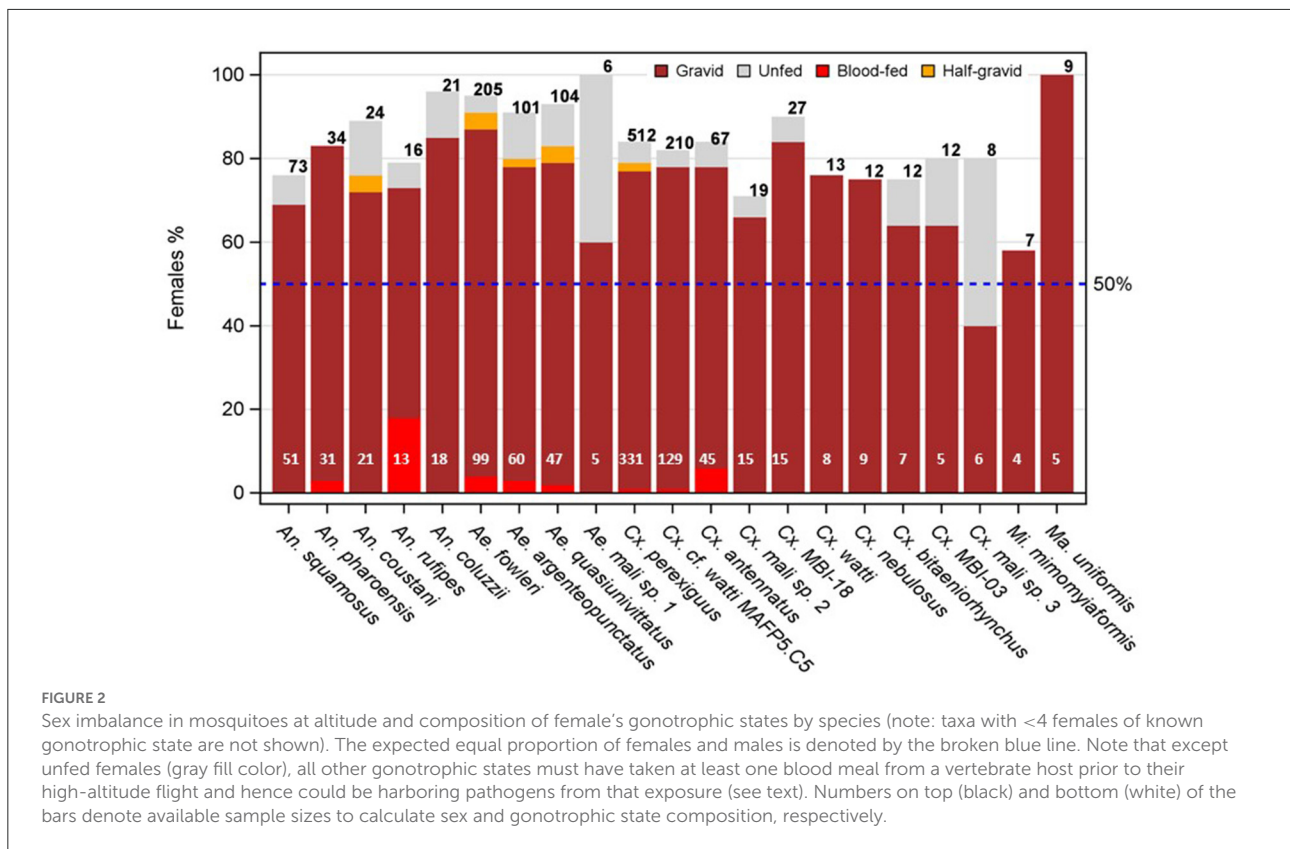
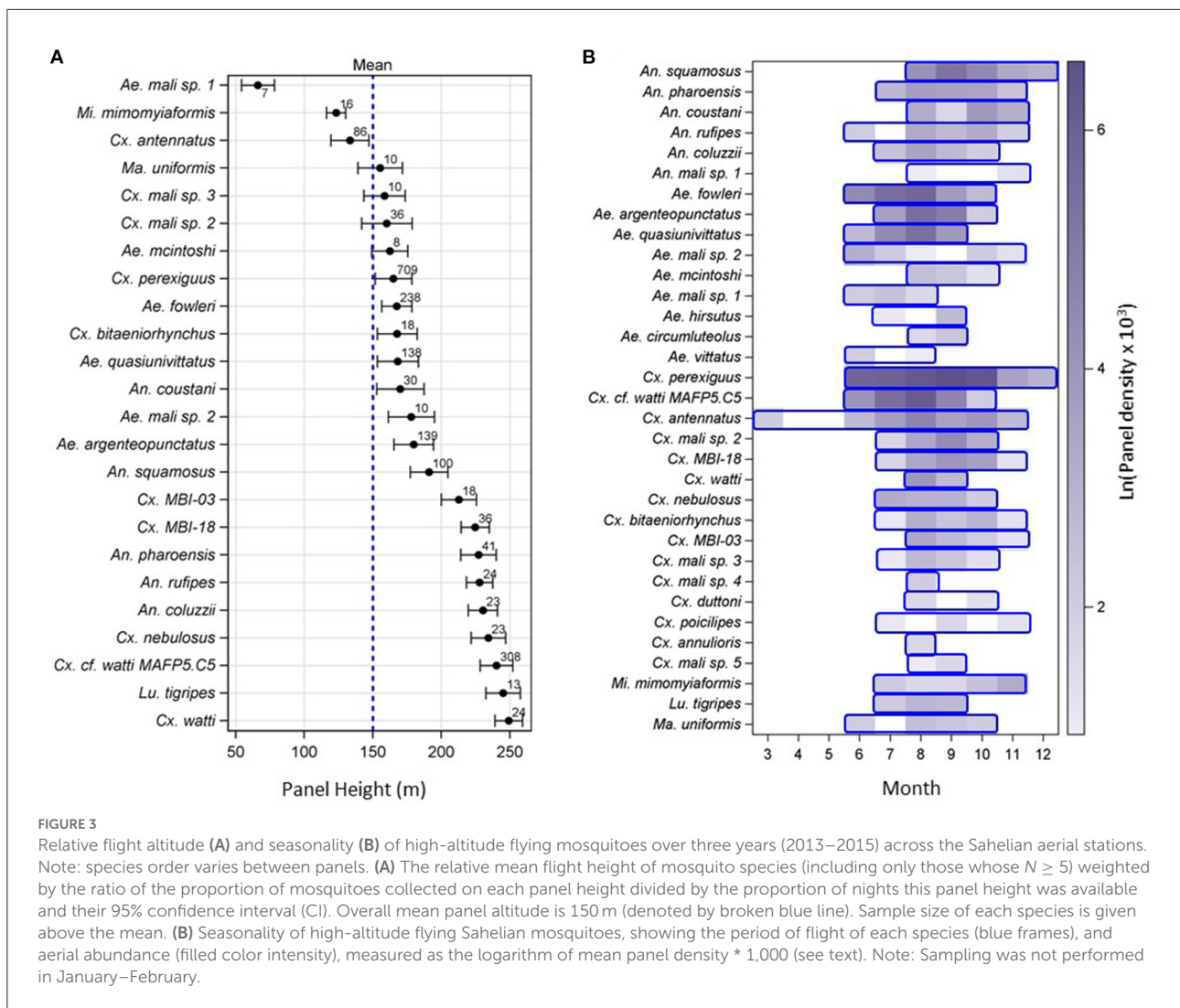


FIGURE 2
Sex imbalance in mosquitoes at altitude and composition of female’s gonotrophic states by species (note: taxa with <4 females of known gonotrophic state are not shown). The expected equal proportion of females and males is denoted by the broken blue line. Note that except unfed females (gray fill color), all other gonotrophic states must have taken at least one blood meal from a vertebrate host prior to their high-altitude flight and hence could be harboring pathogens from that exposure (see text). Numbers on top (black) and bottom (white) of the bars denote available sample sizes to calculate sex and gonotrophic state composition, respectively.



Mali sp. 1 had the lowest exposure rate (60%, $N = 5$, Figure 2, Supplementary Table S2), and five species including *An. pharoensis*, *Cx. watti*, *Cx. nebulosus*, *Mi. mimomyiaformis*, and *Ma. uniformis* had the highest proportion (100%, $4 < N < 30$, Figure 2, Supplementary Table S2).

Flight altitude and seasonality of windborne mosquitoes

Mosquitoes were intercepted on all panels set from 40 to 290 m agl. Estimating a species' typical flight altitude is constrained by the set panel heights (Materials and methods). Thus, although it cannot accurately represent the species actual flight altitude, it can be used to compare species as to their relative typical height weighted by sampling effort at each panel altitude. We estimated the species' relative mean flight altitude by weighting altitudes based on the

ratio of the proportion of mosquitoes captured at that altitude standardized to the proportion of nights that panel altitude was used (Materials and methods, Figure 3A). The weighted mean altitude profile revealed heterogeneity among species as there were non-overlapping 95% CIs. The 95% CI of five species intersected with the mean panel height, but those of three species were below and those of 17 species were above the mean panel height (Figure 3A, $P < 0.002$, Binomial test of $P = 0.5$). Thus, most culicid species tend to fly at higher altitudes than 150 m (Figure 3B). Species of *Culex* and *Anopheles* tended to fly higher than species of *Aedes* although the difference was not significant ($P = 0.071$, $F_{2/17} = 3.07$ ANOVA excluding genera with one species).

The mean number of mosquitoes per panel varied between zero (May, $N = 48$ panels) and 3.7 (August, $N = 395$ panels). Indeed, high-altitude flight was detected across most mosquito species in June–July and ended October–November

(Figure 3B). Exceptions included *Cx. antennatus* (March–November), *Cx. perexiguus* (June–December) and *An. squamosus* (July–December).

Additional sampling would have probably extended these periods, especially for low-abundance species; however, these data suggest a common period of activity across Culicidae (July–October) that coincides with the wet season.

Assessment of the risk for long-range spread of mosquito-borne pathogens by high-altitude windborne mosquitoes in the West African Sahel

Screening of 921 high-altitude mosquito females using pan-*Plasmodium* qPCR assay (39), revealed 22 positive samples. Because assays were conducted on thorax-head tissues (Materials and methods), these results indicate the development of the *Plasmodium* to the sporozoite stage, i.e., an overall infectiousness rate of 2.4%. Such results are used to incriminate mosquito vectors of *Plasmodium* spp., rather than a mosquito that was only exposed *via* infected bloodmeal but cannot transmit it further. *Plasmodium*-infected high-altitude mosquitoes included members of *Culex* (18), *Aedes* (2), and *Anopheles* (2), with *Cx. perexiguus* comprising ~50% of the infected/infectious mosquitoes. Full detail on pathogen detection in high-altitude mosquitoes will be reported elsewhere (Bamou et al.: in preparation).

To assess the risk for long-range spread of pathogens by the windborne high-altitude mosquito species, we collated literature records (40) demonstrating involvement in disease transmission by the mosquito species described above. Only species categorized as primary or secondary vectors for a mosquito-borne pathogen (MBP) in West Africa were included (Methods and Supplementary Table S3). The risk of a given pathogen to be dispersed in altitude by a particular mosquito species depends on the abundance of the mosquito at altitude and on the vector likelihood to carry the pathogen. Accordingly, our risk estimate was approximated by the product of the overall aerial abundance and a weight measuring the relative difference in infectiousness of primary and secondary vectors (1 vs. 0.1, Materials and methods). The sum of the risk estimates for each pathogen across windborne mosquito vectors (when more than one vector species was implicated) was used to rank pathogens by their overall risk of windborne spread, whereas the sum of the risk estimates across pathogens for each mosquito species was used to rank the relative contribution of each mosquito species to the overall windborne spread of mosquito-borne diseases in that region (Figure 4). To evaluate the sensitivity of the results to different weights, we have compared the rankings of the MBPs and the mosquitoes using differential weights reflecting

the range of difference between primary and secondary vectors (Materials and methods and Supplementary Figure S2).

Of the 33 high-altitude flying mosquito species (above), 14 were reported as probable vectors of a total of 25 MBPs in West Africa, including 22 arboviruses, two protozoans (*Plasmodia*) and one nematode (*Wuchereria bancrofti*, WUBNC, Figure 4, Supplementary Table S3). Four mosquito species were recognized as primary vectors for one or more West African MBPs, namely: *An. coluzzii*, *An. coustani*, *Cx. poicilipes*, and *Cx. antennatus*, and the other 10 species were categorized as secondary vectors (Figure 4). Eleven MBPs have been reportedly vectored by one secondary windborne vector species whereas 14 MBPs have been reportedly vectored by multiple vector species or by one primary windborne vector (Figure 4). The MBPs with elevated risk for windborne spread included all those of highest burden on human health: West Nile virus (WNV), malaria (PLFLC and PLVIX), Zika virus (ZIKV), chikungunya virus (CHIKV), yellow fever virus (YFV), and dengue virus (DENV), as well as Rift Valley Fever virus (RVFV) that threatens domestic animals and humans (Figure 4 and Supplementary Figure S2).

That *Ae. vittatus*, *Cx. antennatus*, *Cx. poicilipes*, *An. coluzzii*, and *Ma. uniformis* are confirmed to undertake long distance migrations is of epidemiological significance in West Africa as these taxa are reported to transmit multiple MBPs (6–8 each), whereas *An. coustani* is reported to transmit five (Figure 4). *Aedes hirsutus*, *An. pharoensis*, and *An. rufipes* are known to transmit three MBPs each, whereas the remaining six aerial mosquito species contribute only to the transmission of one or two MBPs each (Figure 4). The index of aerial abundance was not correlated with overall contribution to pathogen spread ($r = 0.009$, $P = 0.97$, $N = 14$).

Discussion

Attributes of high-altitude windborne mosquitoes

High-altitude windborne migration of mosquitoes is a dispersal modality of considerable importance in shaping the shifting landscape of mosquito and pathogen composition over space and time, including the generation of disease outbreaks and range expansion (12, 18, 19, 45, 46). It also is bound to affect gene flow between mosquito populations, mediating the spread of insecticide resistance and possibly, in the future, the spread of genetically engineered constructs (e.g., gene drive systems) designed to impact disease control or elimination (47–49). Here, we expand our first analysis that covered windborne *Anopheles* vectors of malaria (14), which consisted of a mere 10% of the mosquitoes in our aerial collection. Leveraging the full collection of mosquitoes that were

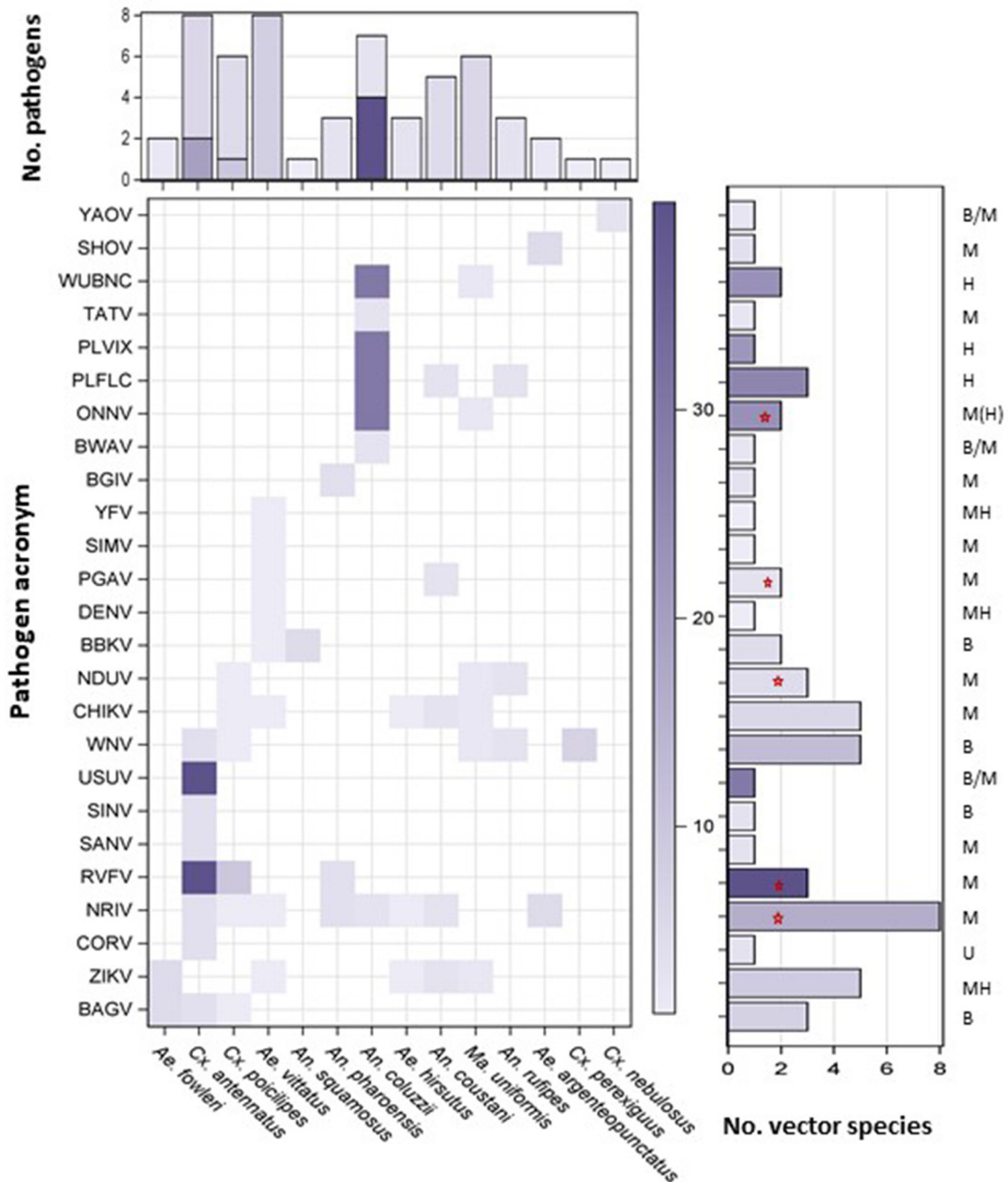


FIGURE 4

Risk of disease spread by windborne mosquitoes. A heatmap showing pathogen (Y-axis, pathogen acronym, see Supplementary Table S3) and vector combinations with weight given by the product of vector importance (primary vs. secondary) and aerial abundance (see text). Vertical bars show the number of diseases reportedly transmitted by each vector, either as primary or secondary vector with the total risk weight (color intensity, see text). Horizontal bars show the number of reported vectors of each pathogen, either as primary or secondary vector with the total risk weight (color intensity, see text) for each pathogen. Pathogens circulating between humans (H), mammals (M), birds (B), or unknown (U) natural hosts are listed above horizontal bars (M > B denotes mostly mammals but some indication of birds; B/M indicates evidence for both without clear ranking; MH indicates that wild mammals are reservoir hosts but human to human transmission by vector can sustain human infection over several years, M(H) denotes that human-to-human transmission occurs rarely). Red stars indicate MBPs of highest risk for being primarily affected by windborne mosquitoes (see text).

intercepted at high-altitude (40–290 m agl) over a systematic three-year surveillance period, we provide compelling support for windborne mosquito migration as being (i) widespread across diverse genera and species; (ii) fivefold more common in females than in males; (iii) 13 times more common in females after they took at least one blood meal than in those that had not; (iv) typically associated with flight altitudes >150 m and probably exceeding our currently highest trap (290 m); (v) coincidental with the wet season (June–November) and peaking in August–October; (vi) mediating the spread of plasmodia of vertebrates, and thus representing a proof of principle that pathogens exploit windborne mosquitoes for their own spread, and (vii) involving mosquito species that have been reported as vectors to 25 MBPs, including those that inflict the heaviest burden on human and animal health in Africa, such as malaria and Rift Valley fever. Together these findings bolster the role of high-altitude windborne mosquitoes as carriers of pathogens over large distances. Therefore, we propose that aerial sampling, coupled with near real-time holistic pathogen detection approaches, can serve as a powerful monitoring system to map regular pathogen movement, and detect irregular events—functioning as an early warning system to address spillover events, and predict the trajectory of outbreaks and epidemics on regional and continental scales.

The culicid diversity in the aerial collections was high, representing 33 confirmed migrant species and additional 17 likely migrant species of the 106 species reported from Mali (40, 42, 44) which are equivalent to 31–47% of the country's fauna. Conservatively, we excluded 17 species represented by a single specimen because some may have been accidental captures (see [Supplementary Results and discussion](#)), however, others were likely less common high-altitude flyers and would be detected through a greater sampling effort. This variation may reflect differences between source population sizes, distances from aerial stations, the fraction of windborne migrants per species, and whether they are diurnal or nocturnal flyers (our sampling was aimed at nocturnal migrants) among other factors. Considering that this 3-year collection is only from the Sahel whereas the mosquito fauna of Mali was studied for over a century by many collections using multiple methods targeting diurnal, crepuscular, and nocturnal species from five ecozones, i.e., humid Gallery Forest, wet and dry savannas, Sahel, and desert (44), we expect the actual number of high-flying species in Mali to increase as more studies address this issue. These results suggest that high-altitude flight is a frequent and widespread behavioral trope in mosquitoes, and that this migration modality appears to be part of a mosquito “blueprint,” as also suggested from high-altitude sampling of species of *Culex*, *Aedes*, *Anopheles*, and other genera from North America, Asia, Europe, and Australia (11, 17, 50, 51), and widely known in many groups of insects that track resources in space and time, e.g., locusts, Hemiptera,

moths, and other dipterans (3, 5, 11, 16, 30, 52–56). It is expected that the abundant seasonal resources available in the Sahel reward long-range migrants, resulting in higher diversity and abundance of migrants (4, 57, 58). Accordingly, the high diversity of insects over the Sahel may reflect migrant mosquitoes from large areas including neighboring ecozones several hundred km away (10, 14, 30, 45, 59–61). Total aerial abundance varied between culicid species from 2 (e.g., *Ae. vittatus*) to 710 (*Cx. perexiguus*), spanning almost three orders of magnitude (Figure 1). In each genus the total specimens of 1–2 species outnumbered all remaining specimens of 4–15 species combined. Surprisingly, some of the most common species in ground collections using human and domestic animal baits, or indoor-resting collections in human and animal dwellings (e.g., *Ae. aegypti*, *Cx. pipiens quinquefasciatus*, *An. gambiae s.l.*) are less common, rare, or altogether absent in the aerial collection. The abundance of a species at altitude primarily depends on the propensity to engage in high altitude flight and on ground density. If ground densities of these “domesticated species” are among the highest, it seems that their propensity to engage in high altitude flight is relatively low, whereas those of the dominant species (e.g., *Cx. perexiguus*, *Ae. fowleri*, *An. squamosus*) are considerably higher, suggesting that the propensity to migrate varies across mosquito species and that only a fraction of the population engages in windborne migration (partial migration).

Sex ratio at altitude showed that, across species, female mosquitoes outnumber males 5.7 to 1, with modest variation around this ratio (Figure 2) attributed to small sample size. Similarly, a 3:1 ratio of female to male mosquitoes (84:29) was reported in a high-altitude collection from South Asia (51). A high proportion of females at altitude being gravid (89.6%, Figure 2) adds weight to the deliberate evolutionary strategy underpinning this long-distance migration. This ratio likely reflects a fitness differential between migrant female and male mosquitoes (62). In species where females typically are inseminated before embarking on migratory flight, and thus, do not require mating upon arrival to destination—the fitness benefit is expected to be asymmetric between the sexes and favor female biased migration (62–64). Unlike male mosquitoes, whose fitness depends primarily on the number of virgin females they inseminate, the fitness of gravid females depends primarily on finding larval sites where predation, competition, and pathogen pressures are low and water remains available until larvae complete their development (65–71); these are also key drivers of migration in other insects with similar life-histories and habitat requirements (3, 55, 72, 73). Arrival of migrant females on the winds that also bring the monsoon rains into the Sahel (8, 14, 30, 74) ensures finding surface water for oviposition. On the other hand, males' prospects of mating with a larger number of (virgin) females are considerably lower because typically the sex ratio of offspring is 1:1, suggesting similar competition per female at the destination as around the

area of departure. Moreover, the days lost in travel, recovery of energy reserves, and finding mating swarms at destination areas where population density is low probably further depress males' prospects of mating at its destination, compared with at its site of origin. Nonetheless, the reward of migratory males is the expected larger number of offspring from the fewer successful mates they acquire due to the better conditions in larval sites at the destination. The strong female predominance, therefore, suggests that unless they are propelled to migrate by their pathogens (75–80), the reduction in the number of expected mates of windborne migrating males is large and that the female expected reproductive success is high enough to expect that most windborne females would be capable of at least one successful egg-laying cycle [with >150 eggs/cycle (81, 82)].

At least 93% of female mosquitoes collected at altitude have taken one or more blood meals on a vertebrate host prior to departure (Figure 2). Although we cannot rule out that some of our gravid mosquitoes are in fact autogenous (capable of egg development and deposition without a blood meal for their first egg batch), none of the 33 species considered as high-altitude migrants are known to be autogenous (83). Furthermore, the relatively high rate of infection with plasmodia implies high rate of blood feeding on vertebrate host. However, further assessment of autogeny in high altitude species is warranted. The variation associated with this proportion was fully attributed to sample size. Because these diverse mosquito species take blood meals from different domestic and wild mammals and bird species, they are exposed to diverse pathogens found in blood or skin of these diverse hosts. Because of the large daily influx of newly emergent mosquitoes and the time—typically 2–3 days—they take before their first blood meal, this proportion is higher than that found in most ground samples (84–88). The age of these females, however, is important for pathogen detection (29). If the majority are at their first or second gonotrophic cycle, i.e., 2–3 days after their first or second blood meal (~5–6 or 8–9 days old), then pathogen density in infected females would be still low, yielding false negative results for infected mosquitoes that would be found positive a few days later. For example, a typical ELISA or PCR assay using thoracic tissues to detect *Plasmodium* sporozoites would be negative in females before day 7–9 post infectious bloodmeal (89, 90). Thus, sensitive methods for pathogen detection should be used and pooling mosquitoes should be minimized.

Our results suggest that most mosquitoes tend to fly at higher altitudes than our average panel height (150 m, Figure 3A). These results agree with radar studies showing that nocturnal insect migrants typically reach heights >200 m above ground and often concentrate in the low-level nocturnal jet that develops above the temperature inversion layer, usually about 200–700 m agl (4, 91, 92). Therefore, our data underestimate the aerial abundance of most species as well as overall culicid diversity. Species of *Culex* and *Anopheles* appeared to fly higher

than species of *Aedes* ($P = 0.071$, $F_{2/18} = 3.07$ ANOVA, excluding genera with one species). Because panel heights varied between years and locations (Methods) and because sample size of most species is below 30 (Figure 1), we cannot evaluate the variation in flight altitude between years and locations. Implicit in our approach is the assumption that the variation in flight altitude of a given species between years and location is negligible. The variation in typical species flight altitude may be related to the expected displacement due to faster winds at higher altitudes.

As expected for Sahelian insects, high-altitude flight of mosquitoes was markedly seasonal, peaking in August–October and was seldom detected outside June–November (Figure 3B). The migration period coincides with the availability of surface water for larval sites (above) and with peak disease transmission in the Sahel and surrounding ecozones, thus, increasing the prospects for pathogen transport by windborne mosquitoes. The high-altitude migration period is probably longer than measured in low-abundance species. Indeed, the longest migration period was observed among the most abundant species, e.g., *Cx. perexiguus* (June–December), *Cx. antennatus* (March–November) and *An. squamosus* (July–December). Likewise, the correlation coefficient between the length of the migration period and log of total aerial abundance was 0.59 ($N = 30$, $P = 0.0006$) when the minimum sample size was 2, but fell to 0.4 ($N = 23$, $P > 0.06$) when the minimum total aerial abundance was 10.

The spread of pathogens by windborne mosquitoes

Finding widespread infection with plasmodial sporozoites (2.4%) in high-altitude mosquitoes across genera (*Culex*, *Aedes*, and *Anopheles*) is a proof of principle that pathogens of vertebrates are carried by windborne mosquitoes (and presumably by other blood-sucking insects) at altitude. Because finding infected (and infectious) mosquitoes using pan-*Plasmodium* assays and because these assays represent our first targeted screen for pathogens, we expect that additional pathogens including arboviruses and filarial nematodes will also be detected in high-altitude mosquitoes. Full detail on pathogen detection in high-altitude mosquitoes will be reported elsewhere (Bamou et al.: in preparation). Importantly, these results establish that infection with pathogens including the development and amplification of the pathogen in the vector's body does not prevent mosquito's high-altitude flight. Nonetheless, comparable data from aerial and ground collections are required to determine whether infection with MBPs suppresses or increases (or does not affect) high-altitude flight in mosquitoes. Unlike laboratory data which remain unresolved (75–80), empirical field data might best address this

issue. Because probability of infection increases with age, the age distribution of high-altitude and ground mosquitoes remains key to the interpretation of these infection rates. Finding thoracic infections with plasmodia indicates that at least some of the females are ≥ 7 days post infectious blood meal and thus are ≥ 9 days old.

Pathogen transport over tens and hundreds of kilometers in a single night by windborne mosquitoes might result in introduction of MBPs to a vulnerable host population and generate disjointed outbreaks (10, 14, 25, 93, 94). Such a process of disease spread is considerably faster than by movements of most terrestrial wild hosts and mosquitoes at the ground level. Mixing pathogen genotypes from different localities by windborne transport of MBPs by mosquitoes might also result in new variants. Furthermore, disease control and elimination are complicated by windborne mosquito movement because man-made barriers and boundaries (e.g., insecticide-sprayed zones) are ineffective against windborne mosquitoes. Re-introduction of pathogens into regions after successful local elimination, such as malaria in South Africa may be complicated by windborne mosquitoes (14). Hence, the contribution of windborne plasmodium to the case load of malaria in wild rodents appears small by the number of direct cases compared with local mosquito transmission, but the rare introduction of distinct genotype MBPs from distant sources into other endemic areas can have far reaching implications (95–99).

In West Africa, mosquitoes are known to transmit 79 pathogens (40). Of these, 25 MBPs (32%) are at risk for being transported and introduced into distant sites by one or more of the 14 windborne mosquito species that were reported as probable or partially implicated vectors (Figure 4 and Supplementary Table S3). Because we conservatively considered only vectors implicated in transmission of MBPs in West Africa that were intercepted at altitude ($N \geq 2$) and because the vectors of many MBPs remain unknown, especially those vectors transmitting MBPs among wild vertebrates in sylvatic cycles [40–42), the actual number of MBPs capable of windborne spread is expected to be higher. Indeed, all the species of plasmodia found to infect our high-altitude mosquitoes represent new vector-MBP records for West Africa (Bamou et al.: unpublished). This sizable proportion of windborne MBPs and the inclusion of the most burdensome (e.g., YFV, DENV, human malaria) and others posing severe threats for new outbreaks and expansion e.g., RVFV, O’Nyong-Nyong virus (ONNV), highlight the importance of elucidating the impact of windborne mosquito migration on MBD spread and control.

In MBPs circulating between humans as natural hosts i.e., *P. falciparum*, *P. vivax*, *W. bancrofti*, and dengue, the importance of spread by windborne mosquitoes is likely smaller than the role of human transport (at least nowadays). Additionally, YFV, ZIKV, CHIKV are maintained among wild mammals (*via* mosquitoes), but once transmitted to humans, these MBPs

can be transmitted between humans (*via* mosquitoes) for up to several years. Thus, further reducing the number of putative MBPs whose disease ecology is shaped by windborne mosquitoes as a key factor. O’Nyong-Nyong virus is capable of transmission between humans (100), but this appears to be a rare occurrence in relation to the maintenance of the virus in sylvatic cycles, suggesting that the ecology of ONNV is shaped by transmission among nonhuman mammalian hosts. The spread of WNV, which is transmitted among birds, and occasionally transmitted to humans as a zoonosis, is primarily shaped by migratory birds (101). However, the case of MBPs that circulate in birds is complex. Firstly, many bird species are residential and move on a distance scale quite similar to resident mammals (102, 103). Secondly, bird migration often occurs out of synchronization with the transmission season, e.g., the migration from Africa to Europe and Asia occurs in March–May, months before the main transmission season in Sahelian and adjacent ecozones (104). Thirdly, many birds fly several hundreds of kilometers a day and exit the ranges of many MBPs before transmission could take place. Seven MBPs are circulating in birds as natural hosts: BAGV, BBKV, BWAV, USUV, WNV, SINV, and YAOV (Figure 4). Because detailed information on migration of the bird host is lacking, we conservatively excluded all these MBPs from the diseases that are most affected by windborne mosquitoes. The remaining 10 MBPs, circulating among wild mammals are less likely to move large distances, and therefore windborne mosquitoes may play a major role in their spatio-temporal dynamics. These include SHOV, NRIV, SIMV, PGAV, TATV, BGIV, RVFV, NDUV, SANV, and ONNV (Figure 4). Among these 10 MBPs, five are transmitted by multiple windborne vectors and thus are ranked as the most affected by this mode of spread: RVFV (with *Cx. antennatus* as primary vector), ONNV (with *An. coluzzii* as primary vector), NRIV (with eight secondary vectors), PGAV, and NDUV (with two and three secondary vectors, respectively; Figure 4). The long-range spread of RVFV is characterized by disjointed outbreaks (105), as expected when MPB is transmitted by windborne mosquitoes.

The three factors that determine the role of windborne spread of an MBP include: (1) the abundance of the vector species at altitude, (2) the number of windborne mosquito species acting as vectors to this pathogen, and (3) whether the vector is a primary or secondary vector (Figure 4). Arguably, the epidemiological influence of a single secondary vector might be high even if it introduces a particular MBP into a new host population once every several years, as compared to a single primary vector that does that hundreds of times a year. Fourteen of the 33 high-altitude flying mosquito species identified here, were reported as vectors of one or more MBPs in West Africa (Figure 4, Supplementary Table S3). Five mosquitoes—*Ae. vittatus*, *Cx. antennatus*, *Cx. poicilipes*, *An. coluzzii*, and *Ma. uniformis*—are reportedly vectors for 6–8

MBPs each (Figure 4), and together contribute to transmission of 22 of the 25 MBPs (Figure 4), posing a disproportional risk for windborne spread of MBPs. Aside from differences in infection rate, the actual vector status of secondary vectors is typically less established than that of those considered primary vectors. Seventeen of the 25 MBPs are known to be vectored by multiple windborne mosquito species or by a single primary vector (Figure 4), thus are ranked high in terms of risk of being transported by windborne mosquitoes. They include RVFV, ZIKV, CHIKV, YFV, *P. falciparum*, *P. vivax*, O'Nyong-Nyong virus (ONNV), Ngari virus (NRIV) whose natural hosts are mammals, as well as WNV and Usutu virus (USUV) whose natural hosts are birds. Many of the MBPs that are transmitted among mammals have been characterized by distant outbreaks – an expected signature of windborne spread, e.g., RVFV, ZIKV, CHIKV, ONNV, NRIV, and YFV, however, this information is not available for many MBPs.

Conclusion and implications

Despite the low efficiency of our passive aerial sampling method and evidence that mosquito density is likely higher above the height of our highest panel, our results suggest that windborne mosquitoes at altitude represent a key driver of MBP spread at least in the Sahel. Moreover its unique features highlight it as an especially informative monitoring system from public health and food security perspectives because it can reveal the sources and destinations of migrating mosquitoes, pests, and pathogens because of the following reasons: (1) the sizable diversity of mosquito species (and genera) at altitude (which matches or surpasses that of most other sampling methods); (2) the large catchment region from which mosquitoes originate that extends tens and possibly hundreds of kilometers upwind from the sampling station; (3) the predominance of female mosquitoes that have already taken at least one blood meal; (4) the timing of mosquito migration at altitude that coincides with the rainy season when MBP transmission peaks; and (5) the fact that many high-altitude windborne mosquito species (14 of 33) have been fully or partly incriminated as vectors of a total of 25 MBPs in West Africa. These include the MBPs that inflict the heaviest burden on human and animal health (e.g., malaria, yellow fever, dengue, Rift Valley fever), affecting public health and food security. A monitoring system based on high throughput aerial sampling stations will inform about regular mosquito and MBP movement patterns including their major sources, routes, and destinations, which is key to the implementation of successful vector and disease control to reduce buildup of mosquito migrants and the likelihood of carriage of MBPs, as well as prevent arriving immigrants from successful disease transmission and establishment in main destination sites. Moreover, departures from “baseline

migration” such as the appearance of a new MBP (or blood/skin pathogen) of vertebrates will be evaluated as a potential new risk, so it could be tracked before it appears in hospitals or veterinary clinics. The prerequisites for such a monitoring system include the demonstration of widespread windborne migration of mosquitoes (and other pests) across African ecozones, infection in high-altitude mosquitoes, and ideally evidence that infection rate at altitude is similar or higher than infection of mosquitoes at ground level. Altogether these results underscore the need to investigate high-altitude windborne mosquitoes and pathogens in Africa to understand the process of disease/outbreak spread, improve disease control strategies, and advance outbreak prediction.

Data availability statement

All DNA barcode sequences used to identify the mosquitoes captured, together with their associated collection data are publicly available on the Barcode of Life Database (www.boldsystems.org) under the project code “TOMAL - High Altitude Culicidae”. The original contributions presented in the study are included in the article/Supplementary material. Further inquiries can be directed to the corresponding author.

Author contributions

This project was conceived by TL. Field methods and operations were designed with DR and JC. Fieldwork, protocol optimization, sampling data acquisition and management, and initial specimens processing was performed by AY, AD, MD, DS, ZS, and YO. Laboratory processing and molecular identification and bioinformatics analyses were done by Y-ML, BK, RB, RF, and CK. Literature search, review, and database construction was carried out by CK, JW, Y-ML, and TL. Data analysis were carried out by TL with inputs from all authors, especially Y-ML, DR, and JC. The manuscript was drafted by TL and AY. All authors have revised the manuscript, contributed in key ingredients and ideas that have shaped the work and the final paper.

Funding

This study was supported by the Division of Intramural Research, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda MD (ZIA AI001196-06) and the Bill & Melinda Gates Foundation (Grand Challenges Explorations grant (OPP1217659) awarded to TL). Y-ML was supported by the Global Emerging Infections Surveillance Branch of the Armed Forces Health Surveillance Division (AFHSD-GEIS) FY21 project P0030_21_WR and P0065_22_WR. Rothamsted Research receives grant-aided

support from the United Kingdom Biotechnology and Biological Sciences Research Council (BBSRC).

Acknowledgments

We are grateful to the residents of Thierola, Siguima, Markabougou, and Dallowere for their permission to work near their homes, and for their wonderful assistance and hospitality. We thank to Moussa Keita, Boubacar Coulibaly, and Ousmane Kone for their valuable technical assistance with field and laboratory operations. We thank Thomas Wellems, Dick Sakai, Laure Juompan, Fatoumata Bathily, Sekou F. Traore, Margie Sullivan, and Sam Moretz for vital and often creative administrative and logistic support.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fepid.2022.1001782/full#supplementary-material>

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Supplementary Materials:

Diversity, composition, altitude, and seasonality of high-altitude windborne migrating mosquitoes in the Sahel: Implications for disease transmission

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Supplementary Results and Discussion

Because no mosquito was collected on 508 control panels that were raised to 40-100 m agl and immediately retrieved during launch and retrieval of the standard panels, these mosquitoes most probably were captured at altitude rather than near the ground (Huestis et al 2019, Florio et al. 2020). Mosquitoes are not captured on the control panels not only because of their short duration but also because launch stations were set in open fields away from humans, animals, and shelters and because most mosquito species are nocturnal, while launch and retrieval occurred during the day. As it was found in over 55% of the genera, i.e., six (or seven) of the 11 genera in Mali (Lehmann *et al.*, 2021; Wilkerson *et al.*, 2021): *Culex*, *Aedes*, *Anopheles*, *Mansonia*, *Mimomiya*, *Lutzia*, and *Eretmapodites* (Table S1), this migration modality is rather common in mosquitoes. Likewise, high proportion of the species in Mali engage in high-altitude migration because depending on whether species sampled by a single specimen are included, our estimate ranges between 31% and 47%. Yet, the actual proportion is expected to be higher because: i) 473 of 2,576 mosquitoes (18.4%) were not assigned to species, and likely include several new species, ii) our aerial sampling was carried out in the Sahel, whereas other ecozones of Mali, which have distinct mosquito fauna have yet to be sampled, and iii) sampling at higher altitudes, e.g., 300-700 m agl, and during stronger winds (precluded given the helium balloons vulnerability to strong winds) would likely increase the number of specimens and the species diversity. The genus *Culex* predominated in the aerial collection both in terms of the number of mosquito specimens and the number of species (Table S1 and Fig. S1). Test of homogeneity among the three largest genera in this fraction revealed that *Culex* has exhibited higher than expected fraction of species in altitude given its total number of species in Mali (45%, $P=0.048$, $\chi^2_{[df=1]}= 3.9$, binomial test). *Culex* has also had the largest number of specimens/species (99).

To ensure we do not include accidentally caught mosquitoes, we excluded species that were represented by a single specimen even though most probably are species that are less abundant in high altitude (see

Figure S1. (Supp. Mat.). Mean number of specimen per species across genera and 95% confidence interval for $N_{\text{species/genus}} > 7$. Mean values are shown above bars.

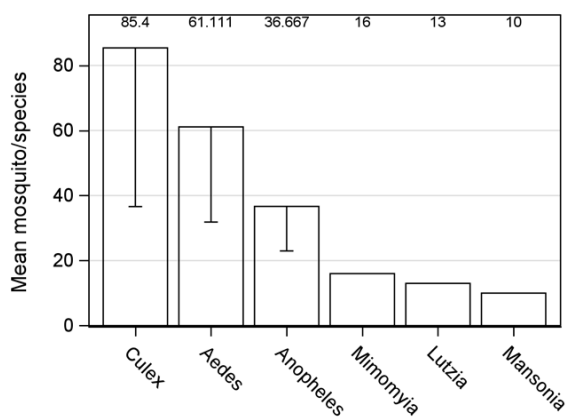


Table S1. Mosquito genera and species in high-altitude in the Sahel and their sample size

Seq.No.	Genus	Species	No. Specimens
1	<i>Aedes</i>	<i>Ae. fowleri</i>	238
2	<i>Aedes</i>	<i>Ae. argenteopunctatus</i>	139
3	<i>Aedes</i>	<i>Ae. quasiunivittatus</i>	138
4	<i>Aedes</i>	<i>Ae. mali sp. 2</i>	10
5	<i>Aedes</i>	<i>Ae. mcintoshi</i>	8
6	<i>Aedes</i>	<i>Ae. mali sp. 1</i>	7
7	<i>Aedes</i>	<i>Ae. hirsutus</i>	4
8	<i>Aedes</i>	<i>Ae. circumluteolus</i>	4
9	<i>Aedes</i>	<i>Ae. vittatus</i>	2
10	<i>Aedes</i>	<i>Ae. aegypti^a</i>	1
11	<i>Aedes</i>	<i>Ae. (Stg.) sp. 1</i>	1
12	<i>Aedes</i>	<i>Ae. ochraceus</i>	1
13	<i>Aedes</i>	<i>Ae. triseriatus</i>	1
14	<i>Aedes</i>	<i>Ae. mali sp. 3</i>	1
15	<i>Aedes</i>	<i>Ae. mali sp. 4</i>	1
16	<i>Aedes</i>	<i>Ae. mali sp. 6</i>	1
17	<i>Aedes</i>	<i>Aedes spp.</i>	84
18	<i>Anopheles</i>	<i>An. squamosus</i>	100
19	<i>Anopheles</i>	<i>An. pharoensis</i>	41
20	<i>Anopheles</i>	<i>An. coustani</i>	30
21	<i>Anopheles</i>	<i>An. rufipes</i>	24
22	<i>Anopheles</i>	<i>An. coluzzii</i>	23
23	<i>Anopheles</i>	<i>An. mali sp. 1</i>	2
24	<i>Anopheles</i>	<i>An. gambiae</i>	1
25	<i>Anopheles</i>	<i>An. sp. nr concolor</i>	1
26	<i>Anopheles</i>	<i>An. mali sp. 2</i>	1
27	<i>Anopheles</i>	<i>An. cf. coustani 1 NFL-2015</i>	1
28	<i>Anopheles</i>	<i>Anopheles spp.</i>	12

29	<i>Culex</i>	<i>Cx. perexiguus</i>	709
30	<i>Culex</i>	<i>Cx. cf. watti</i> MAFP5.C5	308
31	<i>Culex</i>	<i>Cx. antennatus</i>	86
32	<i>Culex</i>	<i>Cx. mali</i> sp. 2	36
33	<i>Culex</i>	<i>Cx. MBI-18</i>	36
34	<i>Culex</i>	<i>Cx. watti</i>	24
35	<i>Culex</i>	<i>Cx. nebulosus</i>	23
36	<i>Culex</i>	<i>Cx. bitaeniorhynchus</i>	18
37	<i>Culex</i>	<i>Cx. MBI-03</i>	18
38	<i>Culex</i>	<i>Cx. mali</i> sp. 3	10
39	<i>Culex</i>	<i>Cx. mali</i> sp. 4	3
40	<i>Culex</i>	<i>Cx. duttoni</i>	3
41	<i>Culex</i>	<i>Cx. poicilipes</i>	3
42	<i>Culex</i>	<i>Cx. annulioris</i>	2
43	<i>Culex</i>	<i>Cx. mali</i> sp. 5	2
44	<i>Culex</i>	<i>Cx. cinereus</i>	1
45	<i>Culex</i>	<i>Cx. decens</i>	1
46	<i>Culex</i>	<i>Cx. pipiens</i> ^a	1
47	<i>Culex</i>	<i>Cx. simpsoni</i>	1
48	<i>Culex</i>	<i>Culex</i> spp.	128
49	<i>Eretmapodites</i>	<i>Er. intermedius</i>	1
50	<i>Lutzia</i>	<i>Lu. tigripes</i>	13
51	<i>Mansonia</i>	<i>Ma. uniformis</i>	10
52	<i>Mimomyia</i>	<i>Mi. mimomyiaformis</i>	16
53	<i>Mimomyia</i>	<i>Mi. mediolineata</i>	1

^a It cannot entirely be rule out is that using colony specimens as positive controls may have resulted in erroneous identification. Although there is no evidence for this possibility, additional caution is needed when this species is being considered as a high-altitude migrant.

main text). Importantly, two species that appear in our aerial collection as singletons are especially important disease vectors, i.e., *Ae. aegypti*, *Ae. ochraceus*, *An. gambiae*, and *Cx. pipiens* (Table S1) (Braack *et al.*, 2018; Lehmann *et al.*, 2021; Wilkerson *et al.*, 2021). An additional concern, we cannot entirely rule out is that during the early phase of the molecular identification, certain specimens used as positive control might have resulted in possible laboratory error due to contamination. Although there is no evidence for this possibility, the inclusion of *Ae. aegypti*, *An. gambiae*, and *Cx. pipiens* from our laboratory as positive controls requires additional prudence. The *An. gambiae* specimen (Table S1) was separated and identified by another laboratory as previously described (Huestis *et al.*, 2019) prior to the processing of specimens in our own laboratory, precluding this possibility for that species identification.

Table S2. Female proportion across species ($N \geq 4$) sorted by the proportion of females in the aerial collection and the proportion of females exposed to vertebrate blood (see text).

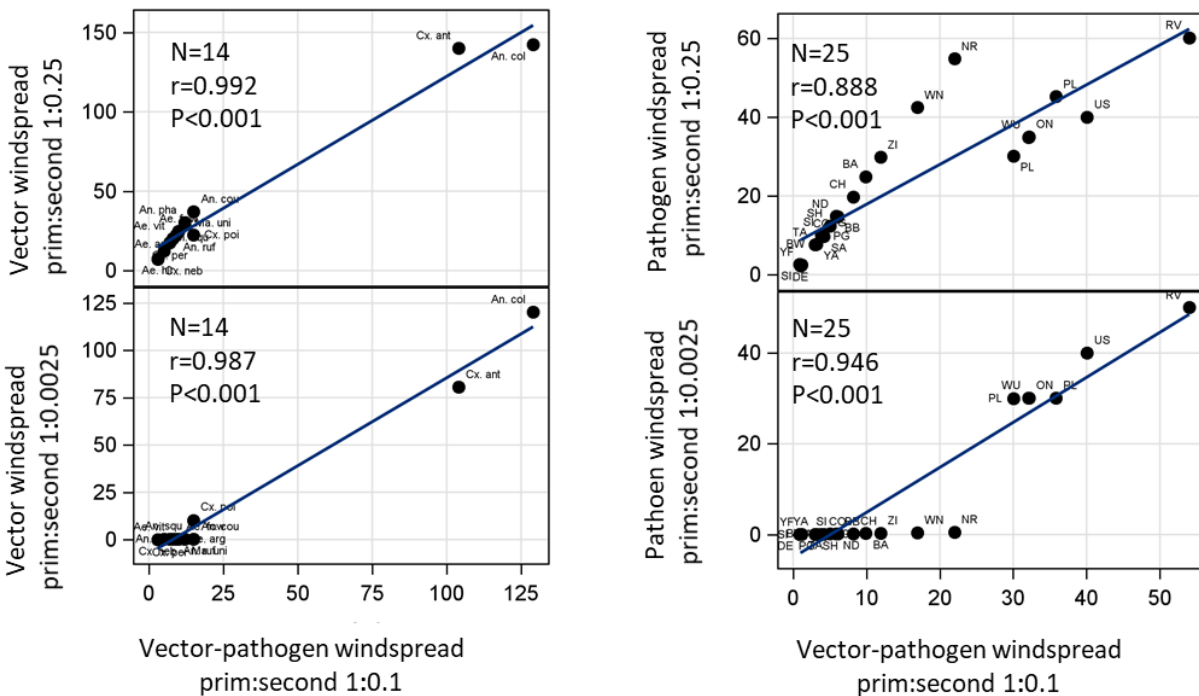
Genus	Species	Females	Female (%)	N (sex)	Exposed	Exposed (%)	N (gono.)
Aedes	Ae. hirsutus	2	50.0	4	ND	ND	1
Lutzia	Lu. tigripes	5	55.6	9	ND	ND	4
Mimomyia	Mi. mimomyiaformis	7	58.3	12	4	100	4
Aedes	Ae. mali sp. 2	5	62.5	8	ND	ND	3
Culex	Cx. mali sp. 2	19	70.4	27	14	93.3	15
Culex	Cx. nebulosus	12	75.0	16	9	100	9
Culex	Cx. bitaeniorhynchus	12	75.0	16	6	85.7	7
Anopheles	An. squamosus	73	76.0	96	46	90.2	51
Culex	Cx. watti	13	76.5	17	8	100	8
Anopheles	An. rufipes	16	80.0	20	12	92.3	13
Culex	Cx. MBI-03	12	80.0	15	4	80.1	5
Culex	Cx. mali sp. 3	8	80.0	10	3	50	6
Culex	Cx. cf. watti MAFP5.C5	210	81.7	257	122	94.6	129
Culex	Cx. antennatus	67	82.7	81	42	93.3	45
Anopheles	An. pharoensis	34	82.9	41	31	100	31
Culex	Cx. perexiguus	512	83.5	613	312	94.3	331
Anopheles	An. coustani	24	88.9	27	18	85.7	21
Culex	Cx. MBI-18	27	90.0	30	14	93.3	15
Aedes	Ae. argenteopunctatus	101	90.2	112	53	88.3	60
Aedes	Ae. quasiunivittatus	104	92.9	112	42	89.4	47
Aedes	Ae. fowleri	205	94.5	217	95	95.9	99
Anopheles	An. coluzzii	21	95.5	22	16	88.9	18
Aedes	Ae. mcintoshi	5	100.0	5	ND	ND	3
Aedes	Ae. mali sp. 1	6	100.0	6	3	60.1	5
Aedes	Ae. circumluteolus	4	100.0	4	ND	ND	2
Mansonia	Ma. uniformis	9	100.0	9	5	100	5

Sensitivity analysis was used to evaluate the effect of the uncertainty (and natural variance) in the relative difference in likelihood of transmission by primary and secondary vectors on our estimates of importance of windborne spread of different pathogens and on the relative roles of different vectors on overall windborne spread of pathogens (see Main Text). Accordingly, we compared the correlations between our best estimates of windborne spread depicted in Figure 4. with those based on high and low values that span the range of the difference between primary and secondary transmission ratios. We considered an infection ratio (approximately equivalent to transmission contribution ratio) of 4:1 as the minimum differential ratio in keeping with the definition of primary and secondary vectors because similar size difference may also be found between two primary vectors in which one contributes 75% and the other 19%, whereas additional three or more secondary vectors contribute less than the remaining 6%. On the other hand, we consider a ratio of 1:0.0025 among the highest differential rate (in their infection rates) because typical sample size per species in most studies ranges between a few hundreds and a few tens of

thousands and the infection rate of the primary vector(s) are near 1% so a ratio of 1:0.0025 implies finding mere 2-3 infected mosquitoes of a sample of 100,000, which very few studies have ever reached not to mention exceeded. Our sensitivity analysis reveals that the estimates of windborne spread based on the mid-range (1:0.1) were highly correlated with those in the extreme low ($r=0.99$, $P<0.0001$, $N=53$) and high of the range ($r=0.979$, $P<0.0001$, $N=53$). When summed over pathogens or mosquito species, high correlations persisted ($r>0.95$, $P<0.0001$, $N=14$, Fig. S1).

[Table S3. Please see separate file in Supplementary Materials.]

Figure S2. Sensitivity of estimates of the relative contribution of windborne transport by different mosquito species (a) and pathogens (b) to the uncertainty in transmission likelihood by primary and secondary vectors (see main text). The relationship between estimates based on the “mid-range” difference between primary and secondary vectors (1:0.1 X-axis), and the lowest difference (1:0.25 Y-axis, top) and highest difference (1:0.0025 Y-axis, bottom) are shown. Values close to the diagonal indicate low sensitivity to different weights or similar patterns across weights. Linear regression between estimators are shown (blue) and observations are labeled by abbreviated vector species and pathogen acronym. Pearson correlation coefficients are shown in top left corner of each panel.



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