

Small dams drive *Anopheles* abundance during the dry season in a high malaria burden area of Malawi

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Abstract

This study explores the influence of small dams on the exposure to malaria vectors during the dry season in Kasungu district, Malawi, an area recently identified as high priority for malaria interventions by the National Malaria Control Programme. Small dam impoundments provide communities with a continuous supply of water for domestic and agricultural activities across sub-Saharan Africa and are considered vital to food security and climate change resilience. However, these permanent water bodies also create ideal breeding sites for mosquitoes in typically arid landscapes. The study focuses on a specific dam impoundment and its vicinity, aiming to assess its spatial and temporal influence on indoor vector densities. From May to August 2021, CDC light traps were used to measure indoor mosquito densities for two consecutive nights per month in three communities located at increasing distances from the dam (0, ~1 and ~2 km). Simultaneously, drone imagery was captured for each community, enabling the identification of additional standing water within approximately 400 m of selected households. Larval sampling was carried out within the impoundment periphery and in additional water bodies identified in the drone imagery. Generalised linear mixed models (GLMMs) were employed to analyse the indoor *Anopheles* abundance data, estimating the effects of household structure (open/closed eaves), month, temperature and water proximity on malaria vector exposure. Throughout 685 trapping nights, a total of 1256 mosquitoes were captured, with 33% (412) being female *Anopheles*. Among these, 91% were morphologically identified as *Anopheles funestus* s.l., and 5% as *Anopheles gambiae* s.l. Catches progressively decline in each consecutive trapping month as the environment became drier. This decline was much slower in Malangano, the community next to the dam, with abundance being notably higher in June and July. Further, the majority of *An. gambiae* s.l. were caught in May, with none identified in July and August. *Anopheles*

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larvae were found both in the impoundment and other smaller water bodies such as irrigation wells in each survey month; however, the presence of these smaller water bodies did not have a significant impact on adult female mosquito catches in the GLMM. The study concludes that proximity to the dam impoundment was the primary driver of differences between survey communities with the abundance in Chikhombwe (~1 km away) and Chiponde (~2 km away) being 0.35 (95% confidence interval [CI], 0.19–0.66) and 0.28 (95% CI, 0.16–0.47) lower than Malangano, respectively, after adjusting for other factors. These findings underscore the importance of targeted interventions, such as larval source management or housing improvements, near small dams to mitigate malaria transmission risks during the dry season. Further research is needed to develop cost-effective strategies for vector control within and around these impoundments.

KEY WORDS

anthropogenic environmental change, dry season transmission, malaria vectors, small dams, vector ecology

INTRODUCTION

The seasonal fluctuation of *Anopheles* mosquito populations is inextricably linked with rainfall. During the rainy season, substantial areas of standing water are available for mosquito egg-laying and larval development. Should these habitats persist long enough to exceed the temperature-dependent time it takes for adult emergence, then lagged malaria transmission will inevitably follow in endemic areas. While there is a positive association between rainfall and mosquito density (Koenraadt et al., 2004; Patz et al., 1998), the process is highly non-linear and dependent on local topography, hydrology and rainfall patterns (e.g., distribution and severity of rainfall events) (Bombaliis, 2012) as well as the ecology of the local dominant species. Once the rains subside, however, mosquito habitat gradually succumbs to evapotranspiration. In certain areas (e.g., the Sahel), this leaves virtually no extant pools for oviposition, causing *Anopheles* populations to enter either a state of dormancy (aestivation) (Faiman et al., 2022) or undergo wind-assisted migration (Huestis et al., 2019) to maintain population viability. That said, in many parts of Africa, enough standing water persists into the dry season extending the malaria transmission season and disproportionately increasing the risk of those communities living in proximity to dry season habitat.

Natural river and lakeside habitat may provide suitable dry-season habitat for mosquito production, but most remaining water bodies are invariably tied to human practices and particularly those related to agriculture and water security. This creates a dichotomy between socio-economic development and the risk of infectious disease (Rohr et al., 2019). Permanent and semi-permanent water-holding structures such as irrigation canals, water impounded by dams, urban agriculture and water storage units, all promote *Anopheles* production in the absence of rainfall (McCann et al., 2018). Epidemiological and entomological observations surrounding individual dams repeatedly show an increased risk of malaria in those people living in proximity to the dam periphery, and these observations are supported by larger systematic data analyses across the continent. Most of the research on

dams and malaria to date has focussed on large dams (Kibret et al., 2015). These are defined as those constructions with a height of 15 m resulting in impoundments holding 3 million m³ of water. The relative contribution of earthen small (or micro) dams—only a few metres high with impoundments containing 100–500,000 m³ of water—on malaria incidence may in fact be greater. Small dam impoundments provide a more suitable habitat for *Anopheles* due to their shallower and more vegetated edges (McCann et al., 2018). Furthermore, communities tend to live in closer proximity to these structures with greater accessibility and use the water directly for livestock watering and domestic activities.

Malawi has a predominantly agriculture-based economy that depends on increasingly erratic annual rainfall patterns (Siderius et al., 2021). Small dams form part of Malawi's growing efforts to double the area of land under irrigation by 2035 (Malawi Government, 2016). They provide some security in terms of water accessibility and are found across Malawi; however, no current database of the location, size and status of these dams exists (Nkhoma, 2011). The general interplay between irrigation and malaria in Africa is well-understood but the relationship is complex and depends on the agroecosystem, socio-economic factors, local vector behaviour and ecology, and the seasonality of disease (Chan et al., 2022; Ijumba & Lindsay, 2001). Efforts to characterise this relationship in Malawi have been performed in a commercial sugar growing estate and within an irrigated rice valley (Mangani et al., 2022; Spaans et al., 2024), and research is underway to do the same within a World Bank-funded large-scale transformation programme (Jones et al., 2023). These systems are, however, much larger and concentrated on a single geographic area. Small dams are different in that their impoundments are much smaller and their impact on mosquito abundance and disease is likely to be far more focal.

The purpose of this study was to determine the relative impact of small dam impoundments on the abundance of *Anopheles* mosquitoes in surrounding villages compared with smaller and more transient water bodies. Combining entomological surveillance with a

drone-captured imagery and on-the-ground larval habitat surveys, the study focussed our efforts on Malawi's dry season given the hypothesis that small dam impoundments have a greater impact on malaria during this period of the year. The study was conducted in Kasungu district, which has recently been identified as a high priority district for malaria interventions in a recent malaria burden stratification exercise (Mategula et al., 2023). Furthermore, Kasungu reports a high percentage of its annual malaria cases during the dry season. In the year in which this study was conducted (2021), facilities in Kasungu reported a total of 431,277 confirmed malaria cases with 24% of these occurring in the dry months of June–October (Malawi Ministry of Health, 2023).

METHODS

Study area

This study was conducted in three communities in Kasungu district, Central Malawi, which were situated approximately 7 km from the district capital of Kasungu town (Figure 1). These communities fall within UNICEF's humanitarian drone testing corridor (UNICEF, n.d.) and form part of the *Maladrone* study, which aims to (1) evaluate the feasibility of using drone imagery to identify mosquito larval habitat during the dry season and (2) to use remotely sensed data on dry

season larval habitat to measure the influence of habitat proximity on the spatial distribution of malaria risk. Further details relating to aim (1) can be found in the study by Stanton et al. (2021), and further analysis using passive surveillance of malaria case data will be published elsewhere.

The three communities of focus in this study were selected based on their proximity to a small dam, which is being used by the local population to irrigate the surrounding area. The dam and its impoundment are situated next to Malangano village, covering an area of approximately seven hectares in the dry season (Figure 1). Preliminary larval surveys undertaken in the area by the study team confirmed the presence of *Anopheles* and *culex* larvae within this impoundment during both the wet and dry season (Stanton et al., 2021), indicating its likely influence on malaria transmission in the local area throughout the year. To determine the range over which the mosquito population is sustained by the impoundment during drier periods, in addition to undertaking mosquito surveys in Malangano village (total population of 165 households), two additional villages with houses situated approximately 1–1.5 km (Chiponde, 92 households) and 1.8–2.6 km (Chinkhombwe, 78 households) from the impoundment boundary were included in this study. An additional small dam was situated approximately 2 km from houses in Malangano, with no additional small dams within 3.5 km of the study communities. The nearest large dam was the Chitete dam, situated on the outskirts of Kasungu town.

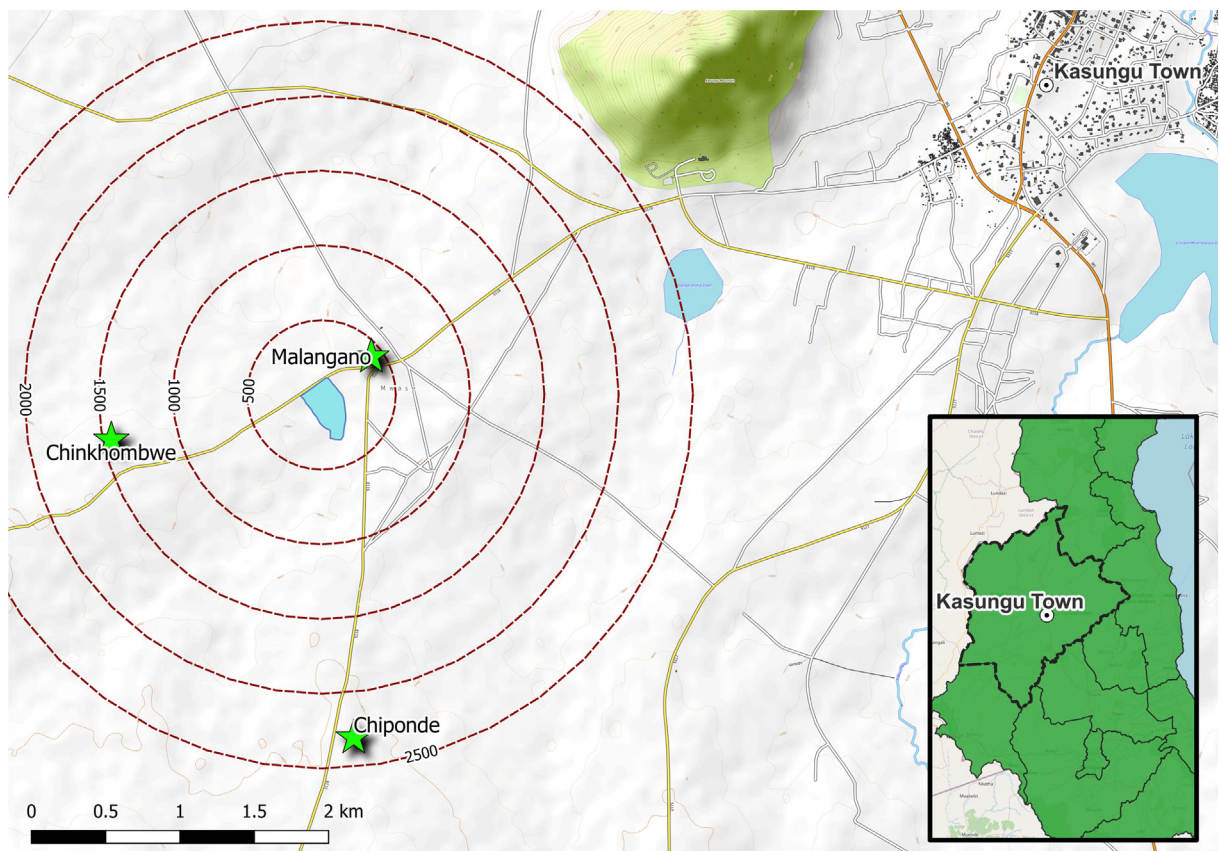


FIGURE 1 Map of the three study communities in relation to Kasungu town and surrounding dam impoundments.

Household selection

A simulation study was conducted to determine the number of households to include in an indoor mosquito sampling exercise. Simulations were generated under the assumption that there needed to be a power of 80% to detect a range of differences in mosquito abundances (number of mosquitoes/trap/night) between the two communities separated by the greatest distance (Malangano and Chiponde). Based on the resources available to conduct this study, it was assumed that the smallest feasibly detectable difference was based on a relative risk (RR) of 0.5, that is, to detect if the mosquito abundance in Chiponde was at most half of the abundance of Malangano. The selected study design comprised of the sampling of 30 households for two consecutive nights per month for 4 months, with the same households being sampled each month. Further details on the sample size simulation study are presented in Supporting Information A.

A list of households in each community was obtained from each village head, and 30 households were randomly selected using a random number generator. Consent to participate in the study was then sought from each of the selected household heads. If any household withdrew from the study during the 4-month period, a replacement household was randomly selected from the household list.

Ethical approval for this study was obtained from both the Liverpool School of Tropical Medicine Research Ethics Committee (Protocol 20.004) and the Kamuzu University of Health Sciences Research Ethics Committee (Protocol P.07/19/2745). Further, a series of stakeholder engagement events were held to engage district officials and community leaders in the study, and community engagement activities took place within all three study communities prior to any recruitment taking place.

Data collection

This study was conducted between May and August 2021, with data collection comprising four components: household surveys, indoor collections of adult mosquitoes, drone image capture and larval sampling.

Household surveys

Prior to starting the indoor mosquito collections, each household head was asked a series of questions to ascertain details about the household structure and composition including the number of people who usually slept in the house/the room where the trap was being placed, the number of bednets used in the house/room, roof materials and whether or not the eaves and windows were open or closed (Supporting Information B). Questionnaire data were captured on smartphones using OpenDataKit (ODK), with anonymised responses being uploaded and stored on Google Drive.

Indoor mosquito collections

Centers for Disease Control (CDC) light traps (CDC-LT, Model 512; John W. Hock Company, Gainesville, Florida, USA) were placed overnight within the sleeping area of selected households to capture adult mosquitoes. Traps were set approximately 1.5 m above the ground next to the foot of the bed between 6 PM and 6 AM each trapping night. Households in each community were sampled sequentially over a 2-week period, with approximately 15 houses being sampled each night. Collected mosquitoes were separated from other insects and were morphologically identified to genus level to separate the *Anopheles* from culicines. All mosquitoes were sexed as either males or females. Culicine mosquitoes were pooled together in storage bottles and no further identifications to species level were conducted. For the *Anopheles* mosquitoes, males were discarded while females were further identified as either *Anopheles gambiae* s.l., *Anopheles funestus* s.l. and other anophelines. Individual female *Anopheles* were stored in 1.5 mL Eppendorf tubes containing a desiccant (silica gel) and each individual female *Anopheles* tubes were assigned unique IDs, and abdominal status (fed/unfed, gravid). All entomological data were subsequently recorded using ODK.

Drone image capture

Drone imagery was used to identify small water bodies in the study area, which had the potential to harbour mosquito larvae. Imagery of the area surrounding each of the study villages was captured each month within the 2-week period in which indoor mosquito collections were being conducted. Image capture was supported by nationally trained drone pilots via the drone service provider Globhe (<https://www.globhe.com/>). Due to the costs involved in capturing imagery, it was necessary to limit the size of the area covered by the imagery. Images were, therefore, captured within a 400-m buffer around the edges of each community only. Images were captured by multirotor drones (DJI Mavic Air and DJI Enterprise) and the resulting images were processed into orthomosaics with a spatial resolution of approximately 3 cm. Due to the time required to upload the imagery and process the orthomosaic there was a time lag of approximately 2 weeks between the images being captured and the orthomosaic being available to the study team. Two members of the study team then independently reviewed each orthomosaic manually and each site where surface water was observed was defined as a potential larval habitat and was subsequently georeferenced and categorised. Categories included irrigation wells, irrigation channels and small ponds. These two datasets were then reviewed together and consolidated, resulting in a single dataset for potential larval habitat locations for each community each month.

Larval sampling

Larval sampling was conducted during the same 2-week window in which indoor adult mosquito collections were being undertaken. An

adaptive sampling strategy was undertaken based on sampling data collected in the previous months, and practical considerations such as accessibility and weather. During the first sampling month, sampling took place along the shoreline of small dam impoundment at Malangano and in additional sites identified from ground-based surveys. The ground-based surveys were conducted with the support of local field staff and volunteers. In subsequent months, the previous month's dataset of georeferenced potential larval habitat was used to guide the larval surveys. Sites were selected based on habitat category (irrigation wells, small ponds, pools around water pumps and dam impoundment) and accessibility. Due to time restrictions, some categories were excluded from the latter drier months' surveys after no larvae were found in any of that category's previously sampled sites in the early wetter months.

A standardised larval sampling approach was adopted for all habitat categories that were sampled. Larvae collections were done using sweep nets. Sweeping nets were dipped in the water at a 45° angle on the water surface, sweeping the entire accessible area (up to 1 m wide) at once to minimise interruptions of the larvae. Sweeping was repeated three to five times, and all contents were emptied into a white plastic bowl and counted by genera (*Anopheles* or culicines) and by stage (early stage [L1/L2] and late stage [L3/L4]). The number of pupae was also counted. Count data were recorded using ODK.

Additional data

Meteorological data were obtained for the study period to enable the variability in temperature and rainfall across the 4-month period to be accounted for in subsequent statistical models of mosquito abundance. These data were obtained from ERA5-Land, which combines ground data from meteorological stations with modelled data to produce hourly estimates at a 9 km spatial resolution (Muñoz-Sabater et al., 2021). Data were extracted for the grid cells covering the study area, and aggregated measurements of total daily rainfall and minimum and maximum daily temperature were calculated from 1 April 2021 up to the final data collection day (mid-August 2021).

Data analysis

Summaries of the number of mosquitoes captured per household per sampling night were produced for each month, disaggregated by community, sex and genera. Further summaries of the female *Anopheles* were generated, including the number of each species complex identified by abdominal stage (unfed, blood-fed, half-gravid and gravid), disaggregated by community and month.

Focusing on female *Anopheles*, a univariate analysis of the community effect was undertaken. A simple generalised linear mixed model (GLMM) that included household-level random effects and a community effect only was fitted to the daily count data for each individual sampling month. A likelihood ratio test was then performed to compare this simple model to a null model (household-level random

effects only) to assess whether significant differences between communities were present.

Summaries and plots of household characteristic data captured within household questionnaires were produced to explore the association between mosquito exposure risk, household composition (number of people who generally sleep in the room in which the net has been placed), the presence of bednets and house construction (eaves open or closed, windows open or closed, roof type). Simple GLMMs fitted to the catch data aggregated over time were fitted which included household-level random effects and single household characteristic fixed effects only. These models were compared to a base model that contained household-level random effects only via a likelihood ratio test.

Summaries of the types of potential larval habitat sites captured within the drone imagery were produced for each community by month. A 400-m buffer was then created for each surveyed household using QGIS (version 3.28), and the number of small potential larval habitats (irrigation wells, irrigation channels, ponds, etc.), that were located within the buffer was calculated, in addition to the distance between the surveyed house and the boundary of the small dam impoundment at Malangano. These variables were used to create potential larval habitat proximity categories (no habitats within 400 m, only small habitats within 400 m, Malangano's dam within 400 m). Plots of these variables against the number of female *Anopheles* catches per sampling night were generated to explore the association between mosquito exposure risk and surface water proximity, and simple GLMMs fitted to the monthly catch data were fitted, which included household-level random effects and a surface water proximity fixed effect only. These models were again compared to a base model which contained household-level random effects only.

The number of sampled potential larval habitats, summarised by genus and stage (early/late), were tabulated by month and habitat type. Maps were produced of all identified potential larval habitat surrounding each community by month, with sampled habitats differentiated according to whether larvae were present or absent.

As weather conditions including temperature and rainfall impact the immature stages of the *Anopheles* lifecycle, temporally lagged effects of ERA5-Land derived variables on indoor adult mosquito catches were anticipated (Agyekum et al., 2021; Hast et al., 2019; Ngowo et al., 2022). To explore this, the daily number of female *Anopheles* per trap per night was aggregated across all sampled houses, and the Pearson's correlation between this value and lagged daily minimum, lagged maximum temperature and lagged rainfall were calculated for lags of up to 21 days. To account for the cumulative effects of temperature and rainfall conditions, moving averages were calculated with a window size of 1 and 2 weeks, such that $\rho_{ij}^{\max(t)}, \rho_{ij}^{\min(t)}, \rho_{ij}^{\text{rain}}$ represent the correlation between weather conditions lagged by i days, averaged over $j = 1, 7, 14$ days for maximum daily temperature, minimum daily temperature and total daily rainfall, respectively. As such, conditions up to 35 days prior to the trapping date were taken into consideration. Values of i and j , which maximised the correlation, were reported, and simple univariate GLMMs were

fitted to the daily data and compared to a base model using the corresponding lagged and averaged weather variables.

More complex spatio-temporal GLMMs were fitted to the mosquito count data to further quantify the influence of household-level characteristics and the surrounding environment on indoor catches. Counts were initially assumed to follow a Poisson distribution, and the fitted GLMMs were tested for overdispersion. Three sets of models were fitted. In the initial model (Model 1), household characteristics were considered as fixed effects to explain between-household differences in catch numbers. Community name and month were included as fixed effects to crudely account for spatial and seasonal differences in catch data respectively. Random effects in Model 1 included collection date to account for differences in localised condition affecting indoor catches and household ID to account for unmeasured differences in households (Supporting Information C). To explore whether seasonal differences could be fully or partially explained by the weather conditions, Model 2 incorporated additional fixed effects using variables derived from the ERA5-Land dataset. Finally, Model 3 explored whether any further spatio-temporal variability could be explained by the presence or absence of small potential larval habitats within 400 m of the house. Akaike's Information Criterion (AIC) was calculated for each of the models under consideration within each model set, and the results of the model with the lowest AIC were reported. The marginal and conditional R^2 values, calculated using methods reported in the study by Nakagawa et al. (2017), were generated for each of the selected models to evaluate the variance explained by the fixed effects in the model and the fixed

and random effects combined. Model diagnostics were performed on the best overall model with respect to AIC, which included tests for overdispersion, heteroscedasticity, collinearity and the normality of random effects.

All data analyses were conducted in R (v4.3.0), with GLMMs fitted using the glmer function within the lme4 (version 1.1-29) package. Model diagnostics were generated using the check_model function within the performance package (version 0.9.0) (Lüdecke et al., 2021).

RESULTS

Table 1 displays the number of *Anopheles* mosquitoes caught and the mean catch per trap per night by sex in each of the three study communities for each sampling month (May–August 2021). A similar table for culicines is available in Supporting Information D. In total, 1256 mosquitoes were captured, of which 32.8% ($N = 412$) were female *Anopheles*, 58.8% ($N = 739$) were female culicines, 1.7% ($N = 21$) were male *Anopheles* and 6.7% ($N = 84$) were male culicines. Catches were consistently higher in the community bordering the dam impoundment (Malangano) across all sampling months, with the largest differences observed in June and July 2021. Figure 2 presents the mean number of female *Anopheles* and culicines per trap per day for each community over time and their respective 95% confidence intervals (CIs). Simple GLMMs fitted to the catch data for each individual month indicate that in May, at the start of the dry season, there was

TABLE 1 Summaries of *Anopheles* trap data by sex and month for houses within the three study villages.

	May			June			July			August			Total		
			Mean/ trap/ night			Mean/ trap/ night			Mean/ trap/ night			Mean/ trap/ night			
	N	Nights		N	Nights		N	Nights		N	Nights		N	Nights	
Malangano															
Female	77	49	1.54	82	60	1.37	51	60	0.85	25	60	0.42	235	229	1.02
Male	3		0.06	2		0.03	1		0.02	6		0.10	12		0.05
Total	80		1.60	84		1.40	52		0.87	31		0.52	247		1.07
Chikhombwe															
Female	46	48	0.96	16	60	0.27	9	60	0.15	15	60	0.25	86	228	0.38
Male	1		0.02	4		0.07	1		0.02	1		0.02	7		0.03
Total	47		0.98	20		0.33	10		0.17	16		0.27	93		0.41
Chiponde															
Female	62	48	1.29	17	60	0.28	6	60	0.10	6	60	0.10	91	228	0.40
Male	2		0.04	0		0.00	0		0.00	0		0.00	2		0.01
Total	64		1.33	17		0.28	6		0.10	6		0.10	93		0.41
Total															
Female	185	145	1.27	115	180	0.64	66	180	0.37	46	180	0.26	412	685	0.60
Male	6		0.04	6		0.03	2		0.01	7		0.04	21		0.03
Total	191		1.31	121		0.67	68		0.38	53		0.29	433		0.63

Note: Malangano borders the dam impoundment, Chikhombwe is approximately 1 km from the dam and Chiponde is approximately 2 km from the dam.

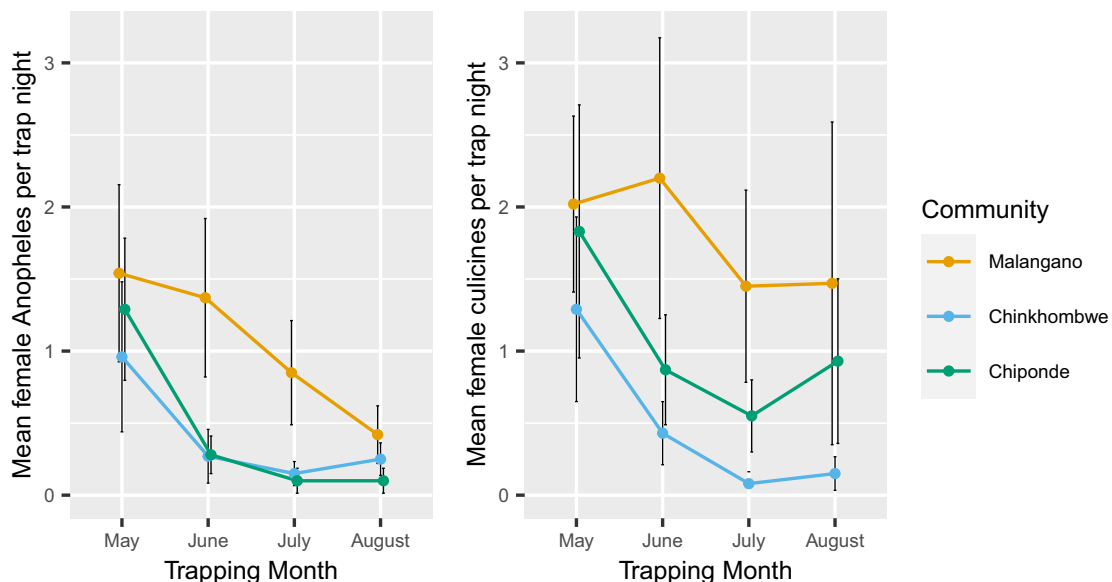


FIGURE 2 Mean number of female *Anopheles* (left) and culicine (right) mosquitoes captured in each community by month. Bars represent the 95% confidence intervals. Malangano borders the dam impoundment, Chinkhombwe is approximately 1 km from the dam, and Chiponde is approximately 2 km from the dam.

no significant difference ($p = 0.2343$) in mean female *Anopheles* catches between the three communities (1.54 [95% CI: 0.926–2.154], 0.96 [95% CI: 0.439–1.481] and 1.29 [95% CI: 0.797–1.783] per trapping day for Malangano [0 km from dam], Chinkhombwe [\sim 1 km from dam] and Chiponde [\sim 2 km from dam], respectively). While catches declined over time in all three communities, the decline was sharpest in the two communities further from the dam, whereas catches in Malangano remained significantly higher ($p < 0.0001$) in both June (1.37 [95% CI: 0.820–1.920]) and July (0.85 [95% CI: 0.489–1.211]). In August, a significant difference remained ($p = 0.0087$), although mean catches in all three communities were low. The lowest catches in August were observed in Chiponde (0.10 [95% CI: 0.013–0.187]), with Malangano and Chinkhombwe being marginally higher (0.42 [95% CI: 0.220–0.620], 0.25 [95% CI: 0.137–0.363] per trapping day for Malangano and Chinkhombwe respectively). For culicines, a similar pattern was observed for May ($p = 0.1263$), June ($p < 0.0001$) and July ($p < 0.0001$), however in August ($p = 0.0023$) catches remained relatively high in Malangano (1.47 [95% CI: 0.350–2.590]) and Chiponde, the community furthest from the dam (0.93 [95% CI: 0.359–1.501]) compared to Chinkhombwe (0.15 [95% CI: 0.033–0.267]).

Of the 412 female *Anopheles* captured during the indoor trapping exercise, 91% (373/412) were morphologically identified as *An. funestus* s.l., 4% (18/412) were *An. gambiae* s.l., and 5% (21/412) were other secondary anopheline species (unidentified) (Table 2). Most of the *An. gambiae* s.l. mosquitoes were captured within houses in Malangano in the first month of sampling (11/18), and none were captured in July or August in any of the sampled houses. A total of 52 (13%) of the 412 female *Anopheles* were blood fed (28 blood fed, 22 gravid and

TABLE 2 Females *Anopheles* captured in each community by month, summarised by morphologically identified species complex.

	May	June	July	August	Total
Malangano					
<i>An. funestus</i>	57	77	49	24	207
<i>An. gambiae</i>	11	2	0	0	13
Other	9	3	2	1	15
Total	77	82	51	25	235
Chinkhombwe					
<i>An. funestus</i>	44	15	7	14	81
<i>An. gambiae</i>	1	1	0	0	2
Other	1	0	2	1	4
Total	46	16	9	15	86
Chiponde					
<i>An. funestus</i>	60	15	6	5	86
<i>An. gambiae</i>	2	1	0	0	3
Other	0	1	0	1	2
Total	62	17	6	6	91
Total					
<i>An. funestus</i>	161	107	62	43	373
<i>An. gambiae</i>	14	4	0	0	18
Other	10	4	4	3	21
Total	185	115	66	46	412

Note: Malangano borders the dam impoundment, Chinkhombwe is approximately 1 km from the dam and Chiponde is approximately 2 km from the dam.

2 semi-gravid), with the majority of these (41) being captured in Malangano, five in Chikhombwe (~1 km from the dam) and six in Chiponde (~2 km from the dam). See Supporting Information E.

Household characteristics of the sampled houses are summarised in Table 3 by community. Communities were similar with respect to the number of residents in each household (median of 4 in Malangano and Chikhombwe, 4.5 in Chiponde), and the number of people sleeping in the room in which the CDC-LT was situated (two in Malangano and Chikhombwe and three in Chiponde). Bed net ownership was highest in Chiponde, the community furthest from the dam (all 30 houses owned at least one net), and lowest in Malangano, the community closest to the dam (21/31 owned at least one net). With regards to house construction, there were no clear differences between communities with approximately half of surveyed houses having an iron sheet roof and half having a thatched roof in all three communities. Only three houses had tiled roofs (two in Chikhombwe and one in Chiponde). The eaves were open or partially open in approximately half of surveyed houses in all three communities,

whereas most houses (76/92, 83%) had open or partially open windows.

Whilst household characteristics may not explain differences in mosquito catches between communities, their influences on within-community variability of aggregated catches of female *Anopheles* were explored (Supporting Information F). These figures, combined with the results of the simple GLMMs, indicated that there were no significant differences in females *Anopheles* per trap per sampling night for all sampling months combined and the number of people usually sleeping in the room ($p = 0.4405$), whether or not there was at least one bednet in the room ($p = 0.5321$), and whether or not windows were closed or (partially) open ($p = 0.8728$). Significantly higher catches were however observed between houses with thatched roofs compared to iron roofs ($p = 0.0221$) and between houses with (partially) open eaves compared to closed eaves (0.0171). It was noted that most houses with iron roofs had closed eaves (38/44), and thatched roofs had open eaves (39/45).

Drone imagery captured for each of the four study months clearly shows a decline in green vegetation over the period in all areas excluding the periphery of the impoundment next to Malangano village (Figure 3). Manual reviews of the imagery, independently undertaken by two team members, identified additional small water bodies in the vicinity of the three communities, primarily in the form of small irrigation wells (Figure 3), the majority of which were in the surrounds of Chiponde (28 observed in May, compared with 8 surrounding Malangano and 11 surrounding Chikhombwe). Chikhombwe, the community approximately 1 km from the dam impoundment, had far fewer potential larval habitat sites in its immediate surroundings compared with the other two communities. Most sites dried up over the observation period in Chikhombwe and Chiponde (Table 4), whereas the number of irrigation wells in Malangano was sustained as a result of several new wells being created close to the impoundment in July and August.

Figure 4 presents boxplots of female *Anopheles* catches per night by potential larval habitat proximity categories (no habitats within 400 m, only small habitats within 400 m, Malangano's dam within 400 m) by sampling month. The presence of small potential habitats (irrigation wells, irrigation channels/puddles, ponds and pooling around boreholes) near houses in Chikhombwe and Chiponde appear to have higher indoor catches than those houses with no potential habitat nearby for May and June only, with this effect diminishing in July and August. Simple GLMMs fitted to the household-level data for each month indicate a significant difference in female *Anopheles* catches in June (0.0006), July (<0.0001) and August (0.0103). Given the higher number of small habitats in the vicinity of Chiponde in comparison to Chikhombwe in May and June (Table 4), this may explain why a greater mosquito abundance was observed in Chiponde over this period despite it being situated the furthest from Malangano's dam.

To explore the potential for these small water bodies as larval habitats, preliminary larval sampling of 13 sites within the study area was undertaken in May 2021 using local field staff's knowledge of the area. In June–August, additional sampling sites were selected by

TABLE 3 Household characteristics by study community.

	Malangano	Chikhombwe	Chiponde
No. of sampled houses			
Median residents	31	31	30
≤5	1	1	1
>5	3	4	3
Total	4	4	4.5
Median no. of people sleeping in room with trap			
≤5	0	0	0
>5	2	2	2
Total	2	2	3
No. of houses with at least one net	21	25	30
Mean nets per resident	0.30	0.28	0.39
No. of houses with at least one net in room with trap	20	23	29
Mean nets per person in room with trap	0.30	0.38	0.44
Roof type			
Iron sheets	15	15	14
Thatched	16	14	15
Tiles	0	2	1
Eaves			
Closed	14	15	14
Open ^a	17	16	16
Windows			
Closed	7	5	4
Open ^a	24	26	26

Note: Malangano borders the dam impoundment, Chikhombwe is approximately 1 km from the dam and Chiponde is approximately 2 km from the dam.

^aOpen includes partially open.

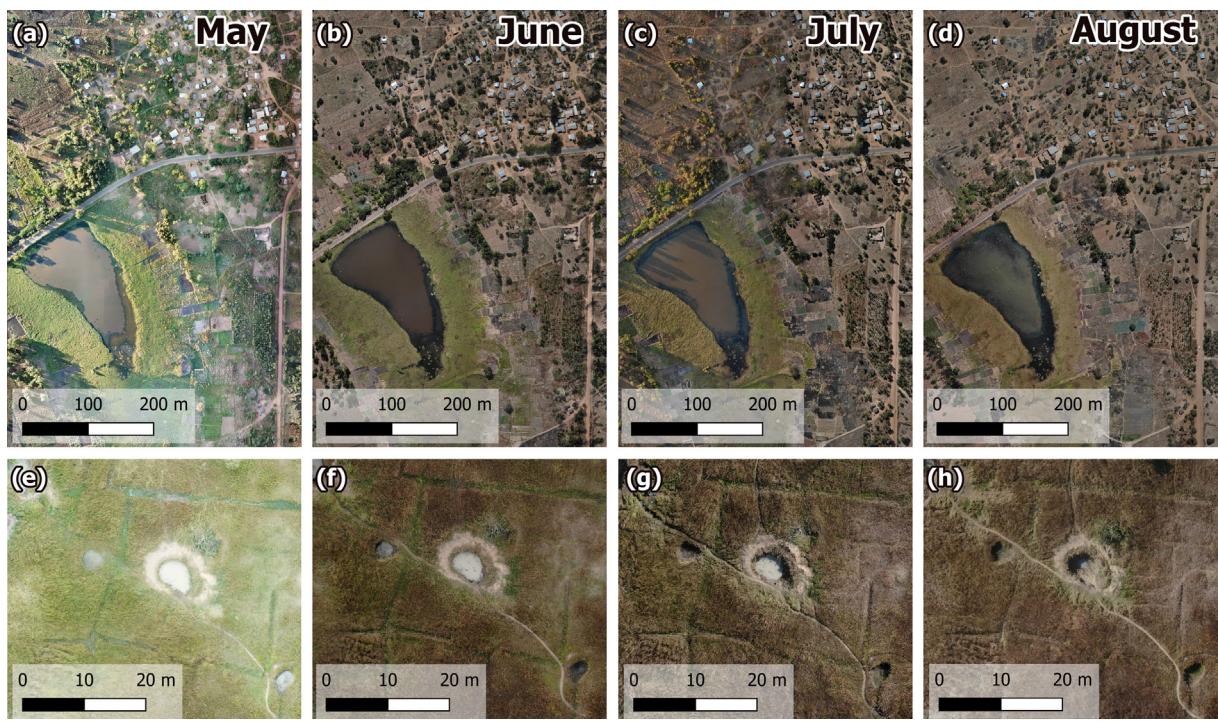


FIGURE 3 Drone imagery of the study area for May–August 2021. (a–d) display the small dam impoundment for each month, with a contemporary close-up of an irrigation well and channels shown in (e–h).

TABLE 4 Number of water bodies detected through manual examination of drone imagery within the surrounds (400 m) of households included in the study.

	May	June	July	August
Malangano				
Irrigation wells	8	5	7	7
Ponds	6	4	0	0
Pooling around water pumps	5	4	2	2
Puddles/irrigation channels	3	1	1	0
Total	22	14	10	9
Chikhombwe				
Irrigation wells	11	5	4	2
Ponds	0	0	0	0
Pooling around water pumps	0	0	0	0
Puddles/irrigation channels	4	0	0	0
Total	15	5	4	2
Chiponde				
Irrigation wells	28	21	9	4
Ponds	3	2	0	0
Pooling around water pumps	1	1	1	1
Puddles/irrigation channels	3	1	0	0
Total	32	28	10	5

Note: Malangano borders the dam impoundment, Chikhombwe is approximately 1 km from the dam and Chiponde is approximately 2 km from the dam.

combining information extracted from the drone imagery with local knowledge. A summary of sites sampled by habitat type is presented in Table 5. Due to the 1-month delay between the drone imagery and larval sampling activities, an additional 23 drone-identified sites were visited by the field team (10 in June, 6 in July and 7 in August) that had since dried up and are therefore not included in this summary.

Summaries of the results of the larval sampling by month and habitat type are presented in Table 6. Additional summaries by month and stage (early or late) are presented in Supporting Information G. Due to the nature of the sample site selection, it is not possible to use these data to draw any robust conclusions about trends over time for each habitat type. Irrigation wells, for which the most samples are available, indicate that while the proportion of sites where no larvae were observed is high (60%, 25/37), they plausibly contribute to malaria transmission in dryer months. Maps of the sampled sites for each month in relation to the sampled households are presented in Figure 5.

ERA5-Land weather data comprising of hourly temperature and rainfall were extracted for the 9 km-by-9 km grid cell containing the study area for April to August 2021 inclusive using the ncdf4 package (version 1.21) in R. Hourly data were aggregated to daily summary statistics (minimum temperature, maximum temperature and total rainfall) (Figure 6). In the month prior to the start of the indoor surveys, daily minimum temperature ranged between 13 and 18°C (median = 16°C), with temperatures lowering from 8 to 16°C (median 11°C) over the 4-month study period, rising again after the final collections had been conducted. Daily maximum temperature was less variable, ranging between 22 and 27°C (median = 24°C) in the month prior to the study and lowering slightly to 20–26°C (median = 23°C)

TABLE 5 Number of sites sampled for larvae by habitat type and month.

	May	June	July	August	Total
Irrigation wells	7	4	10	16	37
Ponds	2	7	2	0	11
Pooling around water pumps	2	2	5	0	9
Impoundment	2	1	5	7	15
Total	13	14	22	23	72

TABLE 6 Larval sampling data by habitat type and month.

Habitat	Month	N	Anopheles larvae only	Culicine larvae only	Both	No larvae
Irrigation wells	May	7	1	0	2	4
	June	4	0	1	0	3
	July	10	1	0	0	9
	August	16	0	6	1	9
Ponds	May	2	0	0	2	0
	June	7	0	3	2	2
	July	2	0	0	1	1
	August	0	0	0	0	0
Pooling around water pumps	May	2	0	1	0	1
	June	2	0	0	0	2
	July	5	0	0	0	6
	August	0	0	0	0	0
Impoundment	May	2	0	1	1	0
	June	1	0	0	1	0
	July	5	3	1	0	1
	August	7	1	1	1	4

during the study. In the month prior to the study collections starting, daily rainfall ranged between 0 and 7 mm (median = 0.4 mm), whereas during the study period there was very little rainfall (min = 0, max = 1 and median = 0), with just 31 of the 103 days (30%) between the first and last collections recording more than 0.01 mm. To account for the lagged and cumulative effects of temperature and rainfall on mosquito development, the Pearson's correlation (ρ_{ij}) between temperature lagged by $i=1, \dots, 21$ days and averaged over $j=1, 7, 14$ days against the daily female *Anopheles* catch data were generated (Figure 7). Generally, a stronger correlation was observed when averaging over the previous 14 days compared to considering the weather on a single day, with a small amount of variation across the considered time lags. A stronger correlation was observed between daily female *Anopheles* catches and minimum temperature ($\max(\rho_{ij}) = \rho_{20,14} = 0.75$) and total rainfall ($\max(\rho_{ij}) = \rho_{20,14} = 0.68$), compared with maximum temperature ($\max(\rho_{ij}) = \rho_{10,14} = 0.56$). Simple univariate GLMMs fitted to the optimal weather variables such that the weather variable was included as a linear fixed effect and the collection date was included as a random effect indicated a significant positive relationship ($p < 0.001$) between all three weather variables and daily female *Anopheles/trap/night*.

Building upon the above exploratory analysis, a series of GLMMs were fitted to the daily household-level female *Anopheles* abundance data (Table 7). Initially, community name and collection month were included as fixed effects to crudely capture spatial and temporal differences, with household-level and community-level random effects being incorporated to account for further unmeasured differences (Model 1, AIC = 1293, marginal $R^2 = 0.29$, conditional $R^2 = 0.50$). Compared with baseline (catches in Malagano in May in houses with closed eaves), it was observed after adjusting for other fixed effects, catches were significantly smaller in Chinkhombwe, located approximately 1 km from the dam (RR = 0.32 [95% CI, 0.18–0.55]) and Chiponde, located approximately 2 km from the dam (RR = 0.31 [95% CI, 0.18–0.54]). Catches declined significantly in successive months (June RR = 0.43 [95% CI, 0.28–0.66], July RR = 0.25 [95% CI, 0.16–0.40], August RR = 0.18 [95% CI, 0.11–0.30]). Catches were significantly higher in houses with (partially) open eaves compared with closed eaves (RR = 1.93 [95% CI, 1.33–2.78]).

Weather variables in the form of minimum temperature averaged over 2 weeks, lagged by 20 days, maximum temperature averaged over 2 weeks lagged by 10 days, and total rainfall summed over 2 weeks, lagged by 20 days were considered in Model 2 to

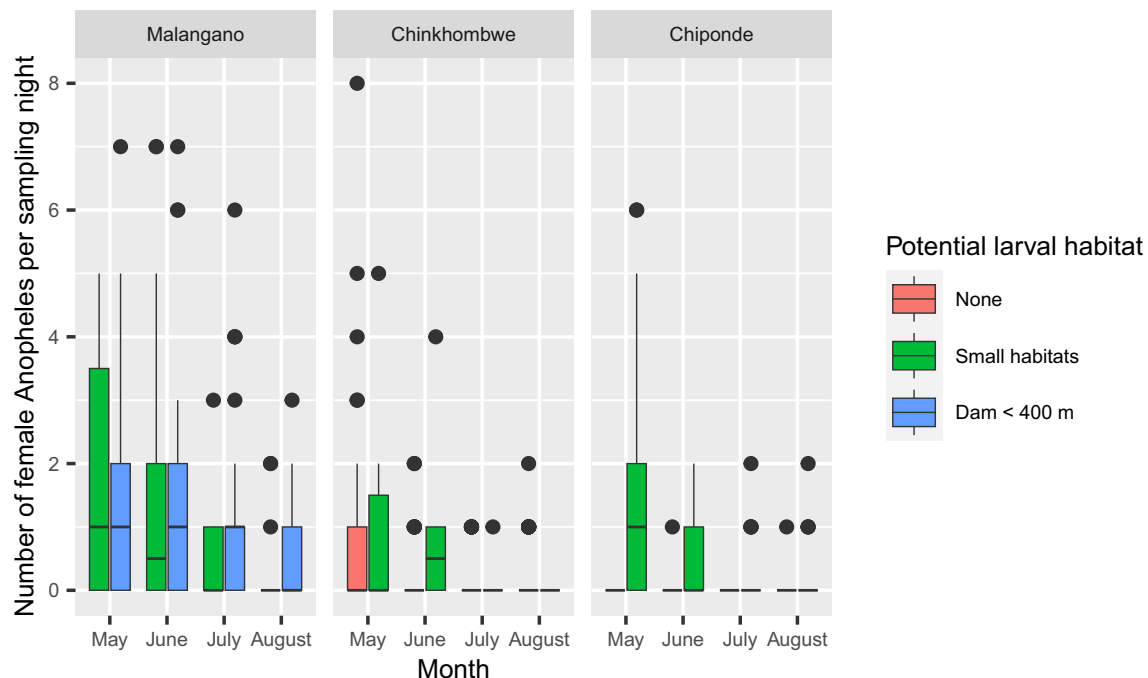


FIGURE 4 Boxplot of the number of female *Anopheles* catches per sampling night in each household by potential larval habitat proximity category (None: no habitat within 400 m of the household; Small habitats: only small habitats within 400 m of the household; Dam < 400 m: Malangano's dam within 400 m of the household) for each study month by study community. Note that the 'None' category is excluded from the Malangano panel as all houses were either within 400 m of the dam impoundment or other small habitats, whereas the 'Dam < 400 m' category is excluded from the Chinkhombwe and Chiponde panels due to these communities being located approximately 1 km and 2 km from the dam impoundment respectively.

account for temporal variability in female *Anopheles* catches. A community effect was included in the model to account for both crude spatial variability and as a proxy for proximity to the Malangano dam. Exploratory plots of weather against catches indicated that the relationship varied between the three study communities, which is likely due to the influence of the dam (Supporting Information H). An interaction between community and weather was therefore considered to account for this variability. Using AIC for model comparison, the minimum temperature variable best captures the variability in catches over time (Table 7). Model 2 had an AIC of 1280, with a marginal R^2 of 0.34 and a conditional R^2 of 0.51. The effect of minimum temperature on catches varied between communities. In Malangano (the baseline community), no significant relationship was observed ($RR = 1.11$ [95% CI, 0.87–1.43]), whereas a positive significant relationship was demonstrated in Chinkhombwe (interaction $RR = 1.42$ [95% CI, 1.05–1.92]) and Chiponde (interaction $RR = 1.97$ [95% CI, 1.43–2.73]). The inclusion of minimum temperature explains some of the variability in catches over time; however, after adjusting for minimum temperature changes, the catches were still lower than expected in July/August compared with May/June (interaction $RR = 0.55$ [95% CI, 0.37–0.83]). Note that time was grouped into 2-month periods in Model 2 as opposed to single months as it was observed that the grouping of months resulted in a marginally lower AIC and the simpler model was, therefore, favoured.

The inclusion of proximity to larval habitat was incorporated into Model 3 ($AIC = 1275$, marginal $R^2 = 0.35$ and conditional R^2 of 0.51). Due to the design of the study, proximity to the dam was already captured in the community effect. Proximity to small water bodies was included as a binary variable (small water present/absent within 400 m). The number of small water bodies within 400 m was considered; however, the inclusion of this additional information did not significantly improve the model. An interaction between the presence of water and month was included to account for the observation that catches were very low in July/August regardless of water being detected in the area. In May/June (baseline time period), a non-significant increase in RR was observed in houses with small water bodies identified within 400 m ($RR = 1.71$ [95% CI, 0.79–1.31]). This effect was however significantly reduced in July/August (interaction $RR = 0.36$ [95% CI, 0.18–0.73]).

Model diagnostics were performed on the best-fitting model (Model 3). No significant overdispersion was detected ($p = 0.494$). Further, no substantial evidence of heteroscedasticity was detected, and distributional assumptions about the random effects in the model appeared reasonable (Supporting Information I). A boxplot of observed against fitted values (Supporting Information J) indicates that the model captures the general trend in catches, differentiating reasonably well between low and high catches, but generally under-predicts larger observed values.

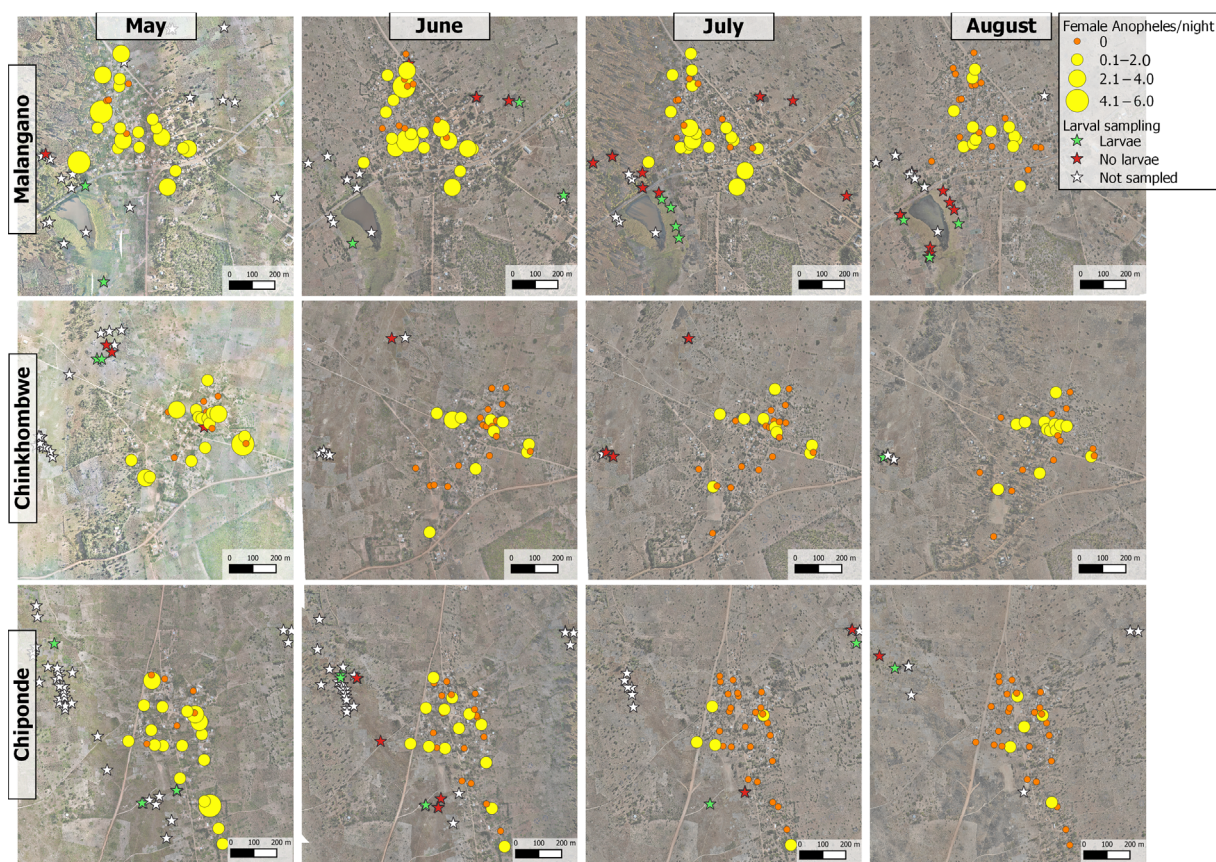


FIGURE 5 Entomological data consisting of household-level number of female *Anopheles* per trap per night and larval sampling sites by larval presence/absence for each community by month. Malangano borders the dam impoundment, Chinkhombwe is approximately 1 km from the dam, and Chiponde is approximately 2 km from the dam.

DISCUSSION

Anopheles mosquito abundance during the dry season was highly focal within our study, with the focality seemingly driven by the presence of a small dam and its impoundment. The influence of proximity to this impoundment on indoor mosquito abundance dropped much more sharply than anticipated, with significantly fewer *Anopheles* caught in villages situated just 1–2.6 km from the impoundment periphery. This extended impact on mosquito numbers, however, lasted only 2 months into the dry season with *Anopheles* abundances falling considerably thereafter. The relative contribution of smaller water bodies on *Anopheles* production in the area—identified from drone-captured imagery and on-the-ground surveys—was inconclusive but any contribution certainly diminished into the dry season. Most of the female *Anopheles* caught by indoor CDC light traps were *An. funestus*, suggesting that small dam impoundments are colonised by this predominant vector of malaria during the dry season in Kasungu.

Water harvesting structures, whether large power-generating hydrodams or smaller community-based dams, perpetuate malaria transmission in Africa (Kibret et al., 2017). Most of the epidemiological and entomological evidence gathered to date has focussed on larger dam structures (Kibret et al., 2015) but meta-analyses across the continent suggest that smaller dams may have an even greater role to

play (Kibret et al., 2021). In an analysis of malaria incidence in populations living less than 5 km from the shoreline to small and large dams in four river basins across Africa, smaller dams contributed to 77%–85% of cases (Kibret et al., 2021). Here, our primary goal was to determine how temporal changes in the local landscape surrounding a small dam in the dry season—in an area designated as ‘highest malaria burden’ by the NMCP— influences mosquito abundance and diversity. Proximity to the dam had a clear influence on indoor female *Anopheles* catch numbers in the adjacent village of Malangano. Shortly after the rains subsided in May, *Anopheles* densities were similar in all three study villages, but as the dry season continued into June and July, households in Malangano received significantly higher numbers of female *Anopheles* compared with households in Chiponde and Chinkhombwe; villages situated at maximal distances of 1.5 and 2.6 km from the dam, respectively. While small differences were observed in bed net ownership and housing structure between communities, large between-community differences were still observed after adjusting for this within the GLMM framework. Furthermore, whereas there is a positive significant relationship between lagged minimum temperature and *Anopheles* abundance observed in Chiponde and Chinkhombwe, no such relationship is observed in Malangano, indicative of an overriding effect of the dam on mosquito numbers in this village.

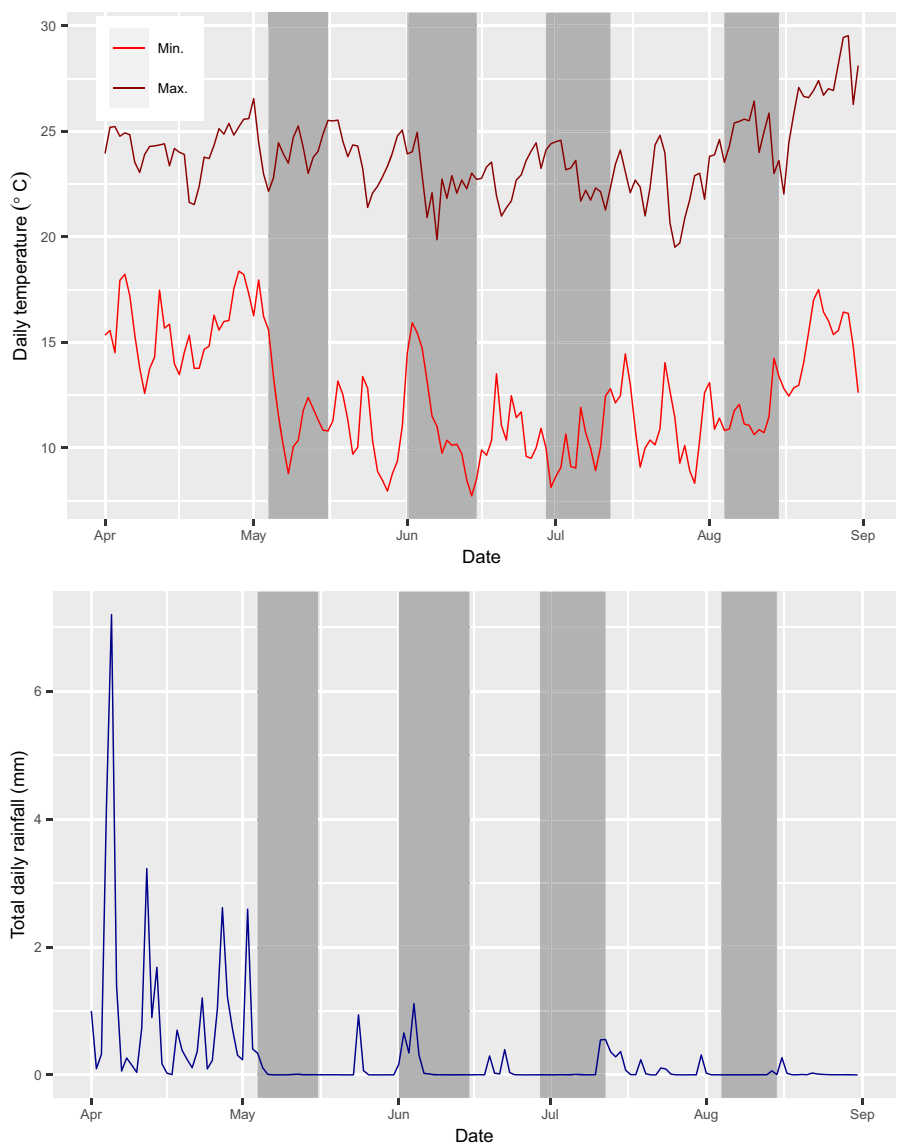


FIGURE 6 Daily minimum and maximum temperature and total rainfall for the study area, derived from ERA5-Land data. Dark grey shaded sections indicate indoor mosquito sampling periods.

The perimeter of small dam impoundments provides an excellent habitat for the larvae of *Anopheles* malaria vectors (McCann et al., 2018). Small dams are generally earthen structures relying on natural drainage with shallow edges often supporting a variety of floating and/or emergent vegetation. Furthermore, receding waters in the dry season leave exposed soil and clay that create transient pools. This creates a wider variety of mosquito habitat around the periphery compared with larger heavily constructed dams. In one of the few detailed larval ecology studies focussed on small dams (McCann et al., 2018), vegetation and soil type along the shoreline had a significant impact on *Anopheles* species abundance and richness, with associations between habitat type and species corresponding to known ecologies and population dynamics of the vector. Small dam impoundments are not homogenous habitats, and extensive heterogeneity in mosquito habitat can exist around a single dam.

Small dams are scattered across the district of Kasungu providing a vital source of water for domestic and agricultural activities. We observed nearly zero rainfall during the period of mosquito collections (May–August) and so these dam impoundments represent one of the few permanent habitats available for *Anopheles* to lay their eggs during the dry season. This is supported by drone-captured declines in green vegetation across the area and the disappearance of smaller alternative water bodies such as irrigation wells and ponds by the end of August, presumably through evapotranspiration (Figure 3). Earlier on in the dry season, these smaller water bodies do appear to influence *Anopheles* abundance in households situated within 400 m, particularly in Chiponde and Chikhombwe (Figure 4) but overall, their relative contribution remains equivocal compared to the dam impoundment.

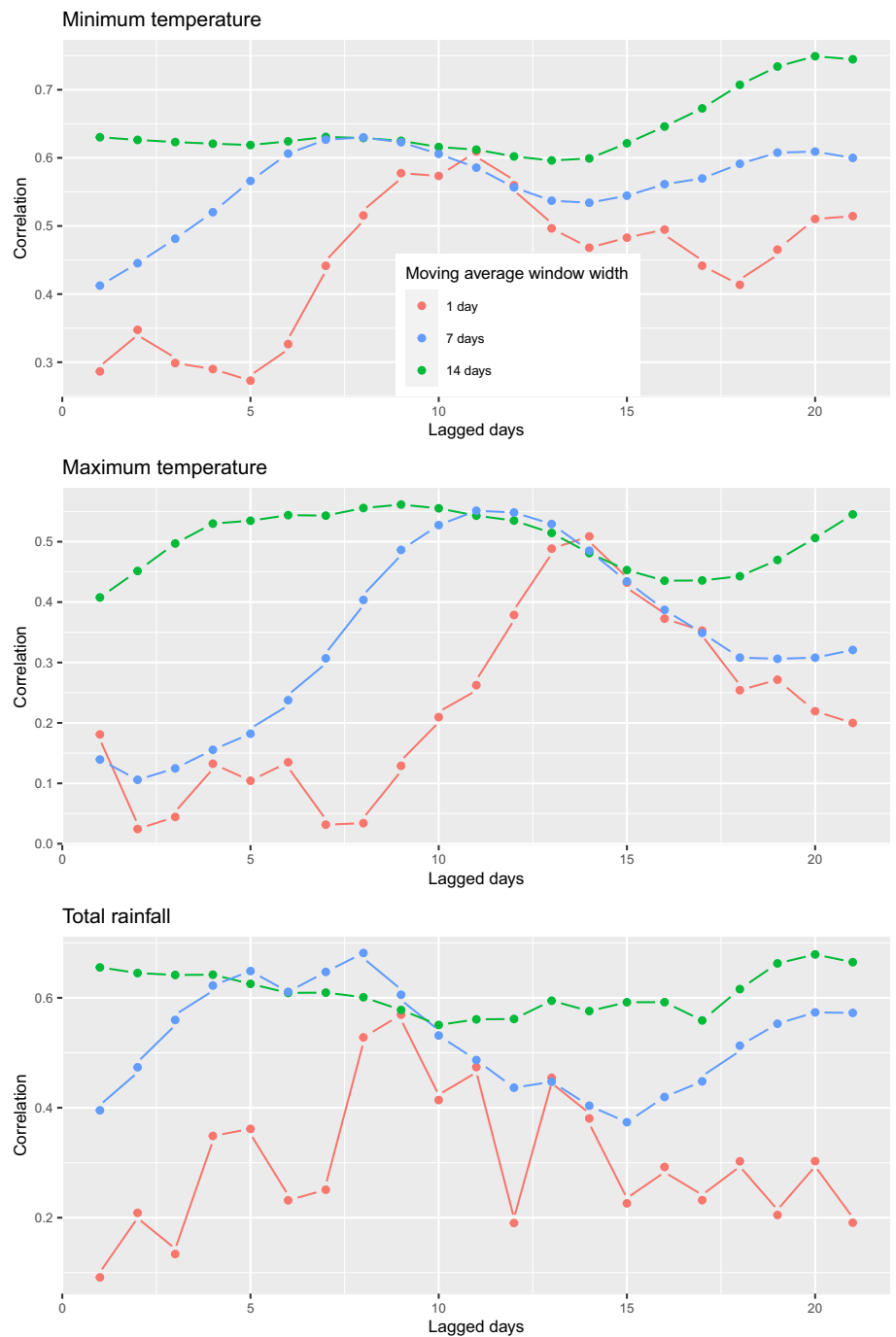


FIGURE 7 Pearson's correlation between daily number of female *Anopheles*/trap/night and moving averages (1 [red line], 7 [blue line] and 14 [green line] days) lagged by 1–21 days.

An. funestus was the primary *Anopheles* vector collected in households suggesting that small dam impoundments act as dry season habitat for this vector. The relative importance of *An. funestus* in malaria transmission in Malawi has increased over the past decade (The PMI VectorLink Project, n.d.) echoing similar species shifts across East Africa (Kaindoa et al., 2017; Ogola et al., 2018) due to its high anthropophilic behaviour and efficiency as a vector of *P. falciparum*. In national annual vector monitoring surveys using CDC-LTs during the same period of our study (2021), *An. funestus* was the predominant vector in two sites sampled within Kasungu district (The PMI

VectorLink Project, n.d.). Seasonal entomological collections in different parts of Tanzania have shown that populations and entomological rates of *An. funestus* are higher in the dry season (Kaindoa et al., 2017; Matowo et al., 2021), consistent with its preference for permanent water bodies. These breeding sites of *An. funestus* have been, however, notoriously difficult to find compared with *An. gambiae* s.l. but recent extensive ecological surveys in Tanzania show that this vector prefers to lay eggs in open semi-permanent water bodies with emerging vegetation and clear water as well as in those habitats exposed to sunlight and high temperatures (Nambunga et al., 2020)—fitting the

TABLE 7 Fixed effects for best fitting GLMM within each model set—Model 1 (AIC = 1293, marginal $R^2 = 0.29$), Model 2 (AIC = 1280, marginal $R^2 = 0.34$) and Model 3 (AIC = 1275, marginal $R^2 = 0.35$).

	Model 1			Model 2			Model 3		
	Relative risk	95% confidence intervals	p-value	Relative risk	95% confidence intervals	p-value	Relative risk	95% confidence intervals	p-value
Intercept	1.29	0.79–2.11	0.316	0.65	0.41–1.05	0.081	0.46	0.22–0.95	0.035
Eaves									
Closed	-	-	-	-	-	-	-	-	-
Open	1.93	1.33–2.78	<0.001	1.95	1.35–2.81	<0.001	1.91	1.33–2.75	<0.001
Community									
Malangano	-	-	-	-	-	-	-	-	-
Chikombwe	0.32	0.18–0.55	<0.001	0.33	0.20–0.55	<0.001	0.35	0.19–0.66	0.001
Chiponde	0.31	0.18–0.54	<0.001	0.29	0.17–0.49	<0.001	0.28	0.16–0.47	<0.001
Month									
May	-	-	-	-	-	-	-	-	-
June	0.43	0.28–0.66	<0.001						
July	0.25	0.16–0.40	<0.001	0.55	0.37–0.83	0.005	1.06	0.57–1.97	0.860
August	0.18	0.11–0.30	<0.001						
Minimum temperature (scaled)									
Lagged 20 days averaged over 2 weeks	-	-	-	1.11	0.87–1.43	0.396	1.02	0.79–1.31	0.896
Min temp: Chikombwe	-	-	-	1.42	1.05–1.92	0.023	1.79	1.27–2.51	0.001
Min temp: Chiponde	-	-	-	1.97	1.43–2.73	<0.001	2.00	1.45–2.76	<0.001
Presence of small water bodies									
Present within 400 m	-	-	-	-	-	-	1.71	0.88–3.31	0.113
Present: July/August	-	-	-	-	-	-	0.36	0.18–0.73	0.005

description of the type of small dam impoundments found across Kasungu, including the one in our study. While we did not perform any species-level larval identification, the lack of any other semi-permanent water bodies in Kasungu, strongly suggests that small dam impoundments support *An. funestus* populations in Malawi. The small percentage of *An. gambiae* s.l. were all captured by June, suggesting their presence was associated with rainfall at the end of the wet season.

The CDC-LT data suggest that the vast majority of *An. funestus* emerging as adults from the small dam impoundment flew a short distance (<500 m) to households in Malangano. Mosquito dispersal is influenced by many factors including odour plume (van Breugel et al., 2015), host availability and density (Kaindoa et al., 2016), wind patterns and the spatial configuration of households and villages (Sedda et al., 2022). The sharp decline in *An. funestus* abundance in Chiponde and Chikombwe in July and August suggests that the dam and village of Malangano are acting as a ‘hotspot’ for *An. funestus*, with the presence of human settlements next to the dam essentially limiting mosquito dispersal and concentrating vector densities. Whether this interaction between mosquito breeding site and the spatial configuration of houses intensifies or extends malaria transmission into the dry season within our setting is unknown. Human biting

rates are expected to occur closest to the breeding site (Smith et al., 2004), whereas exposure to infectious, older mosquitoes, should occur further away from the site of adult emergence (Pinda et al., 2022). Indeed, the RR of catching indoor *Anopheles* in households with open eaves (confirming previous findings in Malawi (Mburu et al., 2018)) demonstrates the importance of other factors such as house construction.

What are the practical implications of our findings? Our investigation into the impact of small dams on mosquito populations and malaria transmission aims to inform recommendations for additional vector control that complement frontline tools like insecticide-treated nets. Based on our findings, small dam impoundments provide focal habitat for the most efficient malaria vector in Malawi, and targeting these areas with larval source management (LSM) could have substantial benefits for those communities living within their vicinity. The optimal timing and coverage of LSM are determined by the local vector ecology and seasonality of malaria, balanced with the practicalities of implementation. The current World Health Organization recommendation is that LSM is most impactful in the dry season, yet a recent modelling study using the OpenMalaria platform suggests that LSM is more effective at the beginning or during the rainy season in seasonal settings (Runge et al., 2021). Investment in LSM as part of an

integrated vector management strategy remains subject to debate and is context-specific, but the small dams situated in Kasungu are worthy of exploring LSM as a viable option. A community-based approach employing a range of environmental management activities significantly reduced both larval and adult *Anopheles arabiensis* densities around a small dam in highland Ethiopia where malaria transmission is markedly seasonal (Yohannes et al., 2005).

To conclude, this study highlights the influence of human-made structures, specifically small dams, on malaria vector exposure in Central Malawi. As these small dams continue to play a vital role in food security and climate change resilience across sub-Saharan Africa, urgent measures are needed to mitigate their public health impact. The highly localised effects of dam impoundments on vector densities revealed in this study can inform the design of targeted control methods, including LSM. Future research should prioritise cost-effective strategies for implementing these targeted interventions.

AUTHOR CONTRIBUTIONS

Kennedy Zembere: Investigation; resources; project administration; supervision; writing – original draft. **Christopher M. Jones:** Conceptualization; funding acquisition; methodology; supervision; writing – original draft. **Rhosheen Mthawanji:** Investigation; resources; writing – review and editing. **Clinton Nkolokosa:** Investigation; resources; writing – review and editing. **Richard Kamwezi:** Investigation; resources; writing – review and editing. **Patrick Ken Kalonde:** Investigation; resources; writing – review and editing. **Michelle C. Stanton:** Conceptualization; data curation; formal analysis; funding acquisition; methodology; project administration; supervision; visualization; writing – original draft.

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FUNDING INFORMATION

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CONFLICT OF INTEREST STATEMENT

The authors declare no conflicts of interest.

DATA AVAILABILITY STATEMENT

The household-level entomological data that support the findings of this study are openly available in Zenodo at <https://doi.org/10.5281/zenodo.8414747>. Household information data are openly available in

Zenodo at <https://doi.org/10.5281/zenodo.10067093>. Household coordinates have been removed from this dataset due to privacy restrictions. Drone imagery relating to this study are openly available from openaerialmap.org.

ETHICS STATEMENT

This study was approved by the Kamuzu University of Health Sciences ethics committee (protocol number P.07/19/2745) and Liverpool School of Tropical Medicine Research Ethics committee (protocol number 20-004).

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

Data S1. Supporting Information.

A. Sample size simulation study.

Figure A. Results of the sample size simulations assuming one village per distance band is sampled, with each house being sampled for one night.

B. Household questionnaire.

- C. Model specification.
- D. Summaries of culicine trap data by sex and month for houses within the three study villages.
- E. Status of female anophelines by sampling month and community.
- F. Boxplots of number of female *Anopheles* per trap per night by household characteristics.
- G. Larval sampling results by genera, stage and month.
- H. Plots of abundance against averaged weather conditions. Solid line represents smoothed conditional means generated using LOESS.
- I. Model diagnostics for the best fitting model.
- J. Boxplots of observed vs fitted female *Anopheles*/trap/night for the best fitting model.

Data S2. Structured reflexivity statement for international research partnerships.

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