A cross-sectional observational study investigating the association between sedges (swamp grasses, Cyperaceae) and the prevalence of immature malaria vectors in aquatic habitats along the shore of Lake Victoria, western Kenya

Getachew E. Bokore, Paul Ouma, Patrick O. Onyango, Tullu Bukhari, Ulrike Fillinger

Abstract

Background: Strategies that involve manipulations of the odour-orientation of gravid malaria vectors could lead to novel attract-and-kill interventions. Recent work has highlighted the potential involvement of graminoid plants in luring vectors to oviposition sites. This study aimed to analyse the association between water-indicating graminoid plants (Cyperaceae, sedges), other abiotic and biotic factors and the presence and abundance of early instar Anopheles larvae in aquatic habitats as a proxy indicator for oviposition.

Methods: A cross-sectional survey of 110 aquatic habitats along the shores of Lake Victoria was done during the rainy season. Habitats were sampled for mosquito larvae using the sweep-net method and habitat characteristics recorded.

Results: Anopheles arabiensis was the dominant species identified from aquatic habitats. Larvae of the secondary malaria vectors such as Anopheles coustani, An. rufipes and An. maculipalpis were found only in habitats covered with graminoids, whereas An. arabiensis, An. ziemanni and An. pharoensis were found in both habitats with and without graminoid plants. The hypothesis that sedges might be positively associated with the presence and abundance of early instar Anopheles larvae could not be confirmed. The dominant graminoid plants in the habitats were Panicum repens, Cynodon dactylon in the Poaceae family and Cyperus rotundus in the Cyperaceae family. All of these habitats supported abundant immature vector populations. The presence of early instar larvae was significantly and positively associated with
swamp habitat types (OR=22, 95% CI=6-86, \(P<0.001\)) and abundance of late *Anopheles* larvae (OR=359, CI=33-3941, \(P<0.001\)), and negatively associated with the presence of tadpoles (OR=0.1, CI=0.001-0.5, \(P=0.008\)).

**Conclusions:** Early instar malaria vectors were abundant in habitats densely vegetated with graminoid plants in the study area but no specific preference could be detected for any species or family. In search for oviposition cues, it might be useful to screen for chemical volatiles released from all dominant plant species.

**Keywords**
Anopheles, oviposition, larval ecology, malaria, vector control, vegetation, graminoid plants

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Background

Malaria, despite increased control efforts, is still among the leading human diseases in Africa. In 2018, 213 million people were infected and 380,000 died\(^1\). The majority of people in sub-Saharan Africa live in poverty and in areas with suitable conditions for the proliferation of malaria vectors\(^2\). The major malaria vectors are in the *Anopheles gambiae* and *An. funestus* species complexes, but a number of less efficient, so-called secondary vectors also contribute to malaria transmission\(^3\).

With growing physiological and behavioural resistance of malaria vectors to insecticides\(^4\)–\(^6\), research efforts are geared towards additional, non-insecticidal vector control strategies\(^7\)–\(^9\). Manipulations of the odour-orientation of adult vectors could lead to novel attract-and-kill interventions\(^10\)–\(^12\). The gravid female searching for a suitable oviposition site is a desirable target for control. This strategy is specifically important as a single gravid mosquito may lay between 50 to 150 eggs\(^7\), hence killing a single gravid mosquito affects the growth of the population. Understanding the cues for habitat selection is of paramount importance for the development of such a tool. Recent work has highlighted the potential involvement of graminoid plants in luring vectors to oviposition sites\(^14\). It has, for example, been shown that *Anopheles* mosquitoes respond to volatile chemical compounds that emanate from rice plants\(^15\).

Malaria vector mosquitoes lay their eggs in standing water and grass-like (graminoid) plants that often dominate wetlands associated with high *Anopheles* larval densities\(^16\)–\(^18\). Some graminoid plants, similar to lowland rice (*Oryza sativa*), are well adapted to wetlands. Most species in the sedge family, also known as swamp grasses, (Cyperaceae) are wetland indicators. One sedge species, *Cyperus rotundus*, was recently associated with the discovery of the oviposition attractant cedrol\(^19\) but its connection to the sesquiterpene compound was not clearly understood.

We considered it plausible to hypothesize that there might be an association between chemical cues released by water-indicating plants that are used by gravid malaria vectors in search of suitable oviposition sites. Therefore, we implemented this study driven by the hypothesis that sedges (Cyperaceae) are associated with the presence and abundance of early instar *Anopheles* larvae, as a proxy indicator for oviposition, in western Kenya.

Swamp habitats are very common along lakeshores and serve as permanent or semi-permanent breeding sites for malaria vectors\(^20\). Studies support that the abundance of *Anopheles* larvae are associated with habitats surrounded by grass-like plants\(^14\). In the current study, we investigated: (1) the distribution of graminoid plants associated with aquatic habitats along the shores of Lake Victoria in western Kenya, (2) the association of the graminoid plants with the occurrence and abundance of *Anopheles* larvae, and (3) the association of abiotic and biotic factors in aquatic habitats with the occurrence and abundance of *Anopheles* larvae.

Methods

**Study area**

This study was conducted on Rusinga Island (0°21’ and 0°26’ south, 34°13’ and 34°07’ east) along the shore of Lake Victoria in Homa Bay County, western Kenya\(^1\) (Figure 1). The area is endemic for malaria and the estimated prevalence of malaria in the population of Rusinga Island is around 10%\(^23\)–\(^25\). Rusinga Island is only around 100 metres away from the mainland and connected via a bridge. The island has an area of 44 km\(^2\) with altitude ranging from 1100 m to 1300 m above sea level and a population size of about 25,000\(^34\). The daily average temperatures of Rusinga Island range from 16°C to 34°C and peak in dry seasons\(^35\). The area experiences bimodal rainy seasons with long rains from March to June and short rains from November to December. Malaria transmission peaks following the end of the long rainy season in June/July\(^25\). The field survey was implemented between May and June 2018, towards the end of the long rainy season.

Habitat surveys were done along stretches of 700 m long and 300 m wide (clusters of approx. 0.2 km\(^2\)). A total of 13 sampling clusters were selected around the lake shores of Rusinga Island (Figure 1B). The areas were selected with the help of Google Earth, aiming at a homogeneous distribution around the island. Inaccessible areas with steep rocks at the shoreline were excluded. Within each sampling cluster all aquatic habitats’ locations were recorded using a smartphone, a unique identifier allocated and sampled as outlined below.

**Aquadic habitat surveys**

The aquatic habitat types were categorized as either swamp, puddle, fishpond, drainage/trench or artificial pit. The perimeter of every habitat was estimated, always by the same field worker for uniformity, by walking in large steps around the habitat. Water turbidity was measured using a turbidity meter (TRB 355IR, WTW Germany) and water pH and temperature were measured using a portable multi-parameter probe (Multi, WTW Germany). These parameters have been shown to be associated with larvae based on previous work done in the same study area.
Every aquatic habitat was inspected for the presence of larvae using the sweep-net method as described by Ndenga and others. The sweep-net (40 cm × 15 cm × 30 cm) was made from fine cotton cloth with a 150 cm long handle. It was chosen for sampling due to its better efficiency in sampling the diverse aquatic fauna including freshly hatched mosquito larvae and mosquito pupae than the standard dipper. A dipper was used for sampling when the habitat was too small to be sampled by a sweep-net. Sampling of mosquito larvae using either sampling tools was randomly done at different points of the habitats since different species of malaria vectors prefer different conditions and vegetation. The duration of sweeping was dependent on the perimeter of the habitat. About 10 minutes were taken to sweep habitats with perimeters exceeding 10 metres, while 5 minutes were taken to sweep habitats <10 m in perimeter. All sweeps were emptied into white trays and mosquito immature stages were counted separately for the two encountered genera, Anopheles and Culex. Culex larvae possess a siphon on the posterior part of their abdomen for breathing whereas Anopheles larvae have no siphon and rest horizontal to the water body. The larvae were grouped as early (1st and 2nd instar) and late (3rd and 4th instar) instars based on their body size. In addition, macroinvertebrates sampled from a habitat were grouped as Odonata (dragonfly and damselfly larvae), Coleoptera (water beetle larvae and adults), Heteroptera (Notonectidae, Naucoridae and Nepidae), fish and tadpoles. All late instar Anopheles larvae and mosquito pupae were transferred to water bottles (1 L) containing habitat water and taken to the International Centre of Insect Physiology and Ecology-Thomas Odhiambo Campus (icipe-TOC) for rearing to adults. Rearing of the field collected larvae was done in 1 L plastic containers. Larvae were fed with a pinch of ground dry cat food (Nestlé Purina Pet-Care Company, Nairobi, Kenya) once daily. The emerged adults were killed in a -20°C refrigerator, sorted by genera and all Anopheles adults stored in Eppendorf tubes (1.5 ml) at -71°C until they were identified morphologically using printed keys and molecularly using polymerase chain reaction (PCR) followed by gel-electrophoresis. Randomly selected mosquito samples were used for molecular identification. Polymerase chain reaction was implemented for the amplification of the ribosomal internal transcribed spacer (ITS2) gene using primers. Positive controls of *An. gambiae* s.s. and *An. arabiensis* (from the insectary) were analyzed with the samples from the field. Extraction of DNA was done for each mosquito separately using Tissue Kit (Quagen, GmbH Hilden, Germany). The PCR was prepared by mixing PCR mix of 2 µl of 5XHot Firepol Blended Master Mix (Ready to Load), primers (0.5 µM each), DNA template (2 µl) and nuclease-free water (5 µl). The thermal recycling conditions involved initial denaturation for 5 min at 95°C, after which 30 cycles of denaturation followed for 30 s at 94°C, annealing for 30 s at 50°C, extension for 30 s at 72°C and final extension for 5 min at 72°C. We used a Kyratec Thermal Cycler (SC300T-R2, Australia) for the thermal reactions. Agarose gel-electrophoresis (2.0%) stained with 2 µl ethidium bromide against a 100 bp DNA ladder (Bioline, A Maridian Life Science Company, UK) and a positive control was conducted to identify the species.

Vegetation coverage, vegetation types and the dominant vegetation type were recorded separately for habitat edge and water surface. Habitat edge was defined as the area along the waterline, approximately 10 cm inside and/or outside the water. Vegetation coverage was estimated visually, always by the same field worker, as the proportion of the habitats covered with vegetation and categorized as (1) 1–25%...
(2) 25–50% (3) 50–75% (4) 75–100%. Graminoid plants across the edge and inside water were recorded as Poaceae, Cyperaceae, Juncaceae, Typhaceae. The graminoid plants were identified to family using the morphology of their leaves (two or three-ranked; open or closed sheaths), and their stem type (three-sided or round; hollow or solid) using Revueltas. Furthermore, herbaceous (not woody and non-graminoid plants) were collectively recorded as forbs. The presence of water plants and algae in the aquatic habitats was also recorded. The percent coverage of water plants and algae on the water surface was visually determined as above. For each habitat, the dominant type of vegetation was identified and recorded. Full specimens of all dominant graminoid plants found in the aquatic habitats were collected and planted at icipe-TOC for further identification.

**Data analysis**

Generalised estimating equations (GEE) with Poisson distribution fitted to a log function and exchangeable correlation matrix were used to test for associations between biological and environmental factors and the abundance of early instar Anopheles larvae. The cluster ID in which habitats were located was included in the model as repeated measurement. A GEE model was also used to analyse associations between factors and the presence of early instar Anopheles larvae. Here we included the presence of early instar Anopheles larvae as dependent variable in the model with binomial distribution fitted to a logit function and exchangeable correlation matrix to analyse its association with biotic and abiotic factors of the habitats (independent variables). The presence and abundance of early instar Anopheles larvae (rather than eggs which are difficult to identify from field samples) were used as dependent variable as a proxy for oviposition. This is based on recent work confirming that early instar density correlates with the abundance of females selecting a habitat for oviposition. The statistical outputs were reported as incidence rate ratios (RR) for the abundance of the first instar larvae and odds ratios (OR) for the presence with their 95% confidence intervals (95% CI). R statistical software version 3.5.1 was used for the analyses.

**Ethics statement**

This field survey was largely descriptive and observational and had no human study participants. Habitat surveys on privately owned lands were made after seeking consent from the landowners and were implemented in their presence.

**Results**

**Aquatic habitat types**

A total of 110 aquatic habitats were identified during the survey. As expected, given the targeted areas within 300 metres of the lake shore, the most prevalent aquatic habitat types were swamps (65.5%, n=72) defined as permanent or semi-permanent water-logged sections of land with tall graminoid vegetation and/or floating plants. The water sources of these were largely groundwater supplemented by rainwater. Other habitats (see Figure 2B, Figure 3C, 3D and 3E) included ponds formerly used for breeding fish but abandoned at the survey time (11%, n=12), rainfed puddles (9%, n=10), drainages (9%, n=10) and artificial pits (5.5%, n=6). Given that all non-swamp habitats were few in number, they were pooled for statistical analysis and the swamp habitats used as the reference group (Figure 3). Early instar Anopheles larvae were found frequently during the survey in the habitat types: artificial pits (n=6, 100%), drainages (n=9, 90%), ponds (n=5, 42%), puddles (n=7, 70%) and swamps (n=61, 85%). The majority of these habitat types were characterized by possessing graminoid plants: graminoid plants dominated the vegetation in 100% of the swamps, in 83% of the ponds, in 80% of the puddles, in 70% of the drainages, and in 50% of the artificial pits.

**Association between graminoid plants and the presence and abundance of Anopheles larvae**

All the swamp habitats were bordered by graminoid plants along the water edges and had a high surface coverage. Similarly, 84% (32/38) of non-swamp habitats had graminoids along their edges and 76% (29/38) had graminoids at their surfaces. Unexpectedly, swamp grasses were not the most frequently found graminoid plants in the survey. Representatives of the Cyperaceae family were found only in 39% of the aquatic habitats sampled. Among the Poaceae family, torpedo grass (Panicum repens) and Bermuda grass (Cynodon dactylon) were the dominant species (Figure 4).

Of the surveyed habitats, 42 (38%) were found covered by P. repens along their edges and 47 (43%) of the habitats at their surfaces. Cynodon dactylon was found covering the habitats both along the edges in 35 (32%) habitats and surfaces in 25 (23%) habitats (Table 1). Overall, graminoid plants dominated in 96 habitats whilst forbs dominated only in five habitats during the survey. Nine habitats had no vegetations at their surface and five of them were colonized by early instar Anopheles larvae. We found water plants in 26 (24%) out of the 110 habitats and most (n=20, 77%) of them in swamp habitats. Filamentous algae were recorded in 21 habitats. Contrary to our hypothesis, there was no significant association between the presence or abundance of early instar Anopheles larvae and the dominant graminoid plant present in a habitat (Table 1).

**Anopheles species composition**

A total of 14,145 early and late instar Anopheles larvae and 402 pupae were collected. Out of those, 4,650 emerged into adults and were morphologically identified (Table 2). Anopheles gambiae s.l. represented 96% of all Anopheles specimen collected. Molecular identification was done for a random sample of 10% of the An. gambiae s.l. (n=480) and revealed 100% An. arabiensis.

Anopheles coustani, An. rufipes and An. maculipalpis were found only in aquatic habitats covered with graminoid plants, whereas An. arabiensis, An. ziemanni and An. pharoensis were found in both habitats with and without graminoid plants. All six species of Anopheles mosquitoes were recorded in swamp habitats. All of these Anopheles species were found in aquatic habitats with dense graminoid vegetation (50–100%) (Table 3). However, only three species of Anopheles mosquitoes (An. arabiensis, An. ziemanni, and An. pharoensis) were collected in habitats sparsely (1–25%) covered by graminoid plants.
Figure 2. Examples of habitat types. (A) Swamp, (B) Fishpond, (C) Puddle, (D) Drainage, (E) Artificial pit.

Figure 3. Bar graph showing percentage of habitats containing graminoid plants and being colonised by early instar Anopheles larvae.

Habitats with graminoids  Habitats colonised by early instar Anopheles larvae
Figure 4. The most dominant graminoid plants identified during the survey. (A) *Panicum repens* (Poaceae), (B) *Cynodon dactylon* (Poaceae) and (C) *Cyperus rotundus* (Cyperaceae).

Table 1. Association between dominant graminoid plants, and the presence and abundance of *Anopheles* early instar larvae.

| Factor                      | No. habitats | Mean (95% CI) of *Anopheles* early instar larvae | Presence of *Anopheles* early instar larvae | Abundance of *Anopheles* early instar larvae |
|-----------------------------|--------------|-------------------------------------------------|---------------------------------------------|----------------------------------------------|
|                             |              | OR (95% CI) P value                              | RR (95% CI) P value                         |                                              |
| *Cyperus rotundus* (Cyperaceae)* | 14           | 57 (22.19-149) 1                                 | 1                                           | 1                                           |
| *Cynodon dactylon* (Poaceae) | 25           | 99 (48-205) 1.1 (0.7-1.7) 0.762                   | 1.7 (0.6-5.5) 0.35                          |                                              |
| *Panicum repens* (Poaceae)   | 47           | 84 (48-146) 1.2 (0.9-1.7) 0.305                   | 1.5 (0.5-4.2) 0.99                          |                                              |
| Others (Poaceae)             | 10           | 58 (33-101) 1.4 (0.99-2) 0.057                    | 1.01 (0.3-3) 0.48                           |                                              |

*Selected as reference based on initial hypothesis and earlier association of *Cyperus rotundus* with oviposition. OR= odds ratio, RR= rate ratio, CI= confidence interval.

Table 2. Species composition of *Anopheles* collected from habitats along the lake shore of Rusinga Island.

| *Anopheles* spp | Number of mosquitoes | Percent composition |
|-----------------|----------------------|---------------------|
| *An. Arabiensis* | 4481                 | 96.24               |
| *An. coustani*   | 22                   | 0.47                |
| *An. maculipalpis* | 2                  | 0.04                |
| *An. pharoensis* | 67                   | 1.44                |
| *An. rufipes*    | 27                   | 0.58                |
| *An. ziemanni*   | 57                   | 1.22                |

* Molecular identification of a random sample of 10% of the *An. gambiae* s.l. revealed 100% *An. arabiensis.*
Aquatic habitats populated with *Panicum repens* and forbs at their edges had all the six *Anopheles* species identified (Table 4a). *Megaloprotachne albescens* was found dominant in six out of 110 habitats surveyed but was found to have all the six different species of *Anopheles* (Table 4b). *Anopheles arabiensis*, *An. coustani*, and *An. pharoensis* were coexisting with all the graminoid types and forbs found dominating along the surfaces of the habitats.

**Table 3.** Mean number ± 95% CI of different mosquito species in swamp and non-swamp habitats, habitats with and without Cyperaceae and graminoids coverage levels.

| Factor                  | Variable          | Mean (95% CI) of *Anopheles* mosquitoes identified from adults emerged from collected immature stages |
|-------------------------|-------------------|-----------------------------------------------------------------------------------------------------|
|                         |                   | *An. gambiae* | *An. ziemanni* | *An. coustani* | *An. pharoensis* | *An. maculipalpis* | *An. rufipes* |
| Habitat type            |                   |              |                |                |                 |                   |              |
| Non-swamp               |                   | 30 (17-46)   | 0.3 (0.1-0.9)  | 0.3 (0.1-0.8)  | 0.4 (0.2-0.9)   | 0                  | 0.5 (0.1-5)   |
| Swamp                   |                   | 45 (29-69)   | 0.6 (0.3-1.4)  | 0.1 (0.1-0.3)  | 0.7 (0.4-1.4)   | 0.03 (0-0.1)      | 0.1 (0-0.6)   |
| Graminoids              | Cyperaceae        | 41 (26-65)   | 0.5 (0.2-1.1)  | 0.2 (0.1-0.5)  | 0.4 (0.2-0.9)   | 0.03 (0-0.1)      | 0.4 (0-1.2)   |
|                         | Forbs             | 38 (22-66)   | 0.6 (0.2-1.8)  | 0.1 (0-1.4)    | 0.9 (0.4-1.9)   | 0                  | 0.02 (0-0.4)  |
|                         |                   | 48 (20-116)  | 0.1 (0.1-0.7)  | 0              | 0.13 (0.1-0.7)  | 0                  | 0              |
|                         |                   | 69 (22-212)  | 0.2 (0-2)      | 0.7 (0.2-3)    | 0.7 (0.14-3.4)  | 0                  | 0.10 (0-10)   |
|                         |                   | 46 (19-112)  | 0.1 (0.0-0.7)  | 0.1 (0-0.7)    | 0.2 (0.04-0.9)  | 0.1 (0-0.5)       | 0.3 (0-8)     |
|                         |                   | 31 (20-50)   | 0.9 (0.4-1.9)  | 0.2 (0.1-0.5)  | 0.9 (0.5-1.7)   | 0.02 (0-0.1)      | 0.4 (0.1-2)   |
| Graminoids coverage (%) | 1-25%             | 48 (20-116)  | 0.1 (0.1-0.7)  | 0              | 0.13 (0.1-0.7)  | 0                  | 0              |
|                         | 25-50%            | 69 (22-212)  | 0.2 (0-2)      | 0.7 (0.2-3)    | 0.7 (0.14-3.4)  | 0                  | 0.10 (0-10)   |
|                         | 50-75%            | 46 (19-112)  | 0.1 (0-0.7)    | 0.1 (0-0.7)    | 0.2 (0.04-0.9)  | 0.1 (0-0.5)       | 0.3 (0-8)     |
|                         | 75-100%           | 31 (20-50)   | 0.9 (0.4-1.9)  | 0.2 (0.1-0.5)  | 0.9 (0.5-1.7)   | 0.02 (0-0.1)      | 0.4 (0.1-2)   |

Association between aquatic habitat biotic and abiotic factors and *Anopheles* larvae presence and abundance

The presence of early instar *Anopheles* larvae in habitats was significantly and positively associated with swamp-type habitats (OR=22, 95%CI=6-86, P<0.001), presence of late instar *Anopheles* larvae (OR=359, CI=33-3941, P<0.001), and presence of *Culex* larvae (OR=17, 95%CI=3-107, P=0.002) (Table 5). In habitats containing pupae the odds of finding early instar...
**Table 5.** Output of multivariate analysis of the presence or abundance of early instar *Anopheles* larvae as outcome, and biotic and abiotic factors as explanatory variables.

| Factor               | Category | Number of habitats | Larval presence/absence | Larval abundance |
|----------------------|----------|--------------------|-------------------------|------------------|
|                      |          |                    | OR (95 % CI)            | P value          |
|                      |          |                    | RR (95 % CI)            | P value          |
| **Abiotic factors**  |          |                    | **Number of habitats**  | **OR (95 % CI)** |
|                      |          |                    | **Larval presence/absence** | **P value**  |
|                      |          |                    | **Larval abundance**     | **RR (95 % CI)** |
|                      |          |                    | **P value**              | **P value**  |
| Habitat type         | Non-swamp| 38                 | 1                       | 1                |
| Swamp                |          | 72                 | 22 (6-86)               | <0.001           |
|                      |          |                    | 1 (0.6-2)               | 0.625            |
| Perimeter (m)        | <50      | 84                 | 1                       | 1                |
| ≥50                  |          | 26                 | 0.3 (0.04-2.3)          | 0.249            |
|                      |          |                    | 0.9 (0.4-2)             | 0.754            |
| Turbidity (NTU)      | <200     | 90                 | 1                       | 1                |
| ≥200                 |          | 20                 | 1 (0.1-17)              | 0.780            |
|                      |          |                    | 2 (0.9-4)               | 0.099            |
| **Biotic factors**   |          |                    | **Anopheles late instar** | **OR (95 % CI)** |
|                      |          |                    | 359 (33-3941)           | <0.001           |
|                      |          |                    | 0.9 (0.4-2)             | 0.839            |
|                      |          |                    | **Culex larvae**        | **OR (95 % CI)** |
|                      |          |                    | 17 (3-107)              | **0.002**        |
|                      |          |                    | 2 (0.6-6)               | 0.244            |
|                      |          |                    | **Pupae**               | **OR (95 % CI)** |
|                      |          |                    | 0.08 (0.01-0.42)        | **0.003**        |
|                      |          |                    | 1 (0.7-2)               | 0.437            |
|                      |          |                    | **Odonata**             | **OR (95 % CI)** |
|                      |          |                    | 2 (0.3-11)              | **0.518**        |
|                      |          |                    | 0.5 (0.3-0.9)           | **0.019**        |
|                      |          |                    | **Coleoptera**          | **OR (95 % CI)** |
|                      |          |                    | 0.3 (0.03-3)            | 0.274            |
|                      |          |                    | 0.4 (0.2-0.8)           | **0.004**        |
|                      |          |                    | **Fishes**              | **OR (95 % CI)** |
|                      |          |                    | 0.4 (0.05-2)            | 0.288            |
|                      |          |                    | 0.6 (0.2-2)             | 0.336            |
|                      |          |                    | **Tadpoles**            | **OR (95 % CI)** |
|                      |          |                    | 0.1 (0.01-0.5)          | **0.008**        |
|                      |          |                    | 0.5 (0.2-0.9)           | **0.030**        |

**OR**= odds ratio, **RR**= rate ratio, **CI**= confidence interval.

*Anopheles* larvae was lower (OR=0.08, CI=0.01-0.42, P=0.003) than habitats without pupae. Notably, the majority of habitats with *Anopheles* larvae were also well colonised by other invertebrates, many of which are considered predators of mosquitoes, such as Odonata, Notonecta, and Coleoptera larvae. However, the presence of early instar *Anopheles* was only significantly and negatively associated with presence of tadpoles (OR=0.09, 0.01-0.53, P=0.003). Correspondingly, the abundance of early instar *Anopheles* larvae significantly decreased with the presence of tadpoles (RR=0.5, CI=0.2-0.9, P=0.03). Similarly, larval abundance was negatively associated with presence of Odonata (RR=0.5, CI=0.3-0.9, P=0.019) and presence of Coleoptera (RR=0.4, CI=0.2-0.8, P=0.004). There was no significant association between the presence and abundance of early instar *Anopheles* larvae and habitat size, habitat depth, distance to the nearest house, water pH, water turbidity, biofilm, debris, algae, and water plants.

**Discussion**

The work presented here was done with the aim of identifying graminoid plants for further behavioural and chemical ecology studies due to their association with habitats used by gravid malaria vectors for egg-laying. However, the presence of early instar *Anopheles* larvae in the majority of the surveyed habitats and the presence and high coverage of various graminoid plants did not allow us to analyse any statistically significant association. All the habitats surveyed provided excellent oviposition sites and favourable conditions for the development of immature stages based on the high and consistent number of early instar larvae as a proxy for oviposition and the associated high abundance of late instar larvae as an indicator for survival. The study, as implemented, did not allow us to infer specific plant-based factors with oviposition. Generally, the association between graminoid plants and *Anopheles* breeding sites as well as the presence and increased densities of *Anopheles* larvae in both temporary and permanent aquatic habitats have been shown before[16,17]. Recent studies have also shown promising odour-blends of volatile organic compounds identified from domesticated grasses such as rice and pollens of maize and sugarcane. These compounds, which include limonene, α-pinene, p-cymene, nonanal, benzaldehyde, sulcatone, β-caryophyllene, decanal, and 3-carene, have been reported to elicit a response in gravid *An. arabiensis*[15,39,40]. It has been suggested that vegetation can protect mosquito immature stages from being washed off by running water[41] and from predation[42,43].

Our study has several limitations that might be responsible for the negative results. The timing of the survey towards the end of the rainy season meant that all potential habitats were flooded and vegetation thrived. Habitats for oviposition were...
not a limiting factor and likely easy to identify without major cues for orientation. This might have been different if the survey had been implemented during the dry season. Furthermore, this survey was limited to locations close to the lake shores, biasing the study towards swampy habitats. Potentially a more rigorous evaluation of the plant coverage using standard methods such as a quadrant frame which might have provided more detailed information on plant numbers, could have revealed more associations. However, given the high colonisation during the rainy season such method would be better applied during drier seasons. Lastly, due to high water levels during the peak rainy season, a number of habitats with swamp graminoids of the families Cyperaceae, Typhaceae, and Juncaceae were impossible to access, hence could not be sampled. This might also explain why only very few secondary malaria vector species and no Anopheles funestus were sampled, even though An. funestus is the major vector in houses in the study area44-46.

Not many strong associations were found with early Anopheles larvae presence or abundance and other observed factors that would allow conclusions on oviposition preferences. However, the odds of finding early instars increased when late instar Anopheles larvae were present as opposed to when they were absent, potentially indicating that Anopheles arabiensis females oviposit in habitats containing late instar conspecific larvae as an indicator of suitable development conditions. This contrasts with experimental studies on An. coluzzii47,48, where it has been suggested that late instar conspecific larvae repel gravid females potentially due to the risk of cannibalism49.

The presence and abundance of early instar Anopheles larvae were negatively associated with the presence of tadpoles. It was previously shown that rainwater conditioned with tadpoles repelled gravid An. gambiae from oviposition in the laboratory50. Mature tadpoles can prey on larvae51 and might compete for resources in aquatic habitats52. Our field survey indicated that early instar larvae were cohabiting with predatory invertebrates in most habitats. Whilst Anopheles larvae might be reduced by these organisms, as suggested by the negative association between Anopheles density and the presence of Odonata and Coleoptera, gravid females nevertheless did not avoid these habitats for oviposition. This finding agrees with studies elsewhere that have shown strong associations between the presence of anopheline larvae and high invertebrate diversity53.

Six species of Anopheles mosquitoes were identified from the samples which have all been reported in previous studies in western Kenya44,46,55. Anopheles arabiensis was the predominant malaria vector from both vegetated and non-vegetated aquatic habitats in the study area during the peak rainy season. Anopheles arabiensis has historically been the predominant vector species on Rusinga Island56. In recent years however, An. funestus predominates indoor vector collections44,46,57, but larvae were not found during our survey. Breeding sites preferred by this mosquito species were inaccessible by the field team due to the large volumes of water in the lake after the long rains; An. funestus prefers breeding habitats that are covered by tall vegetations46,58,59.

Conclusions
Our results did not support the hypothesis and nullified the aim of the research to identify graminoid plant species positively associated with malaria vector oviposition. Our results did not support our initial hypothesis and did not allow us to identify any association between Anopheles larvae and specific graminoid plants. However, Panicum repens, Cynodon dactylon, and Cyperus rotundus were the predominant graminoid plants found in the aquatic habitats. The habitats covered by these vegetation were abundantly colonized by early instar Anopheles larvae even though no specific preference for any of these could be detected, likely due to study limitations. We recommend further studies on the identification of oviposition cues from graminoid plants during the dry seasons when habitats are limited and water-levels low enough to provide access to most of them. Furthermore, it might be warranted to implement bioassays in the laboratory with the here identified grass-like plants, which will allow more standardised comparisons and sufficient replication.

Data availability
Underlying data
Harvard Dataverse: Association between graminoids and the prevalence of immature malaria. https://doi.org/10.7910/DVN/NAT0YY60.

This project contains the following underlying data:
- Bokore et al. 2020_F1000Research_All_Collected_Data.csv
- Bokore et al. 2020_F1000Research_Data_used_for_final_analysis.csv
- Bokore et al. 2020_F1000Research_Variable_Codes.csv

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

Acknowledgements
We are grateful to the residents of Rusinga Island for allowing us to collect the field data in their farms. The administrative support of Ibrahim Kiche is highly appreciated.
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Eric Ochomo
Centre for Global Health Research, Kenya Medical Research Institute (KEMRI), Kisumu, Kenya

I think the authors have done a good job of revising the manuscript and it is in a indexable format.

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 1

Reviewer Report 14 September 2020

https://doi.org/10.5256/f1000research.28336.r70204

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Eric Ochomo
Centre for Global Health Research, Kenya Medical Research Institute (KEMRI), Kisumu, Kenya

This is an important article that attempts to associate the presence of sedges and the presence of immature malaria vectors along the shores on Lake Victoria on the Rusianga Island. The article is quite informative and can generate a tool that can be instrumental in the fight against malaria. I agree with most aspects of it but would like to suggest some improvements that can make it even look better. Below are some of the areas I noted that needs adjustment.

1. Exploitation of the oviposition cues can be very important in implementing the attract and
kill mosquito control technique. The authors should identify and discuss more chemical
cues that would potentially attract gravid females to lay eggs in the graminoid other than
cedrul present in *Cyperus rotundus* in the other members of Cyperaceae family identified in
this article.

2. The method used to estimate the perimeter of individual habitats is subject to errors incase
two different people are involved since one person's step cannot be exactly be the same to
another ones. I suggest a verifiable method ought to have been used in evaluating habitat
sizes.

3. The author indicates that he performed test for abiotic factors like turbidity, pH and
temperature only. I think that the parameters were not the only abiotic factors that would
influence the distribution of immature stages of malaria vectors. Abiotic factors like
Dissolved oxygen (DO), salinity atmospheric pressure could also influence the abundance
and distribution of mosquito larvae in the water and ought to have been evaluated.

4. In establishing the coverage of the various graminoid plants in the larval habitats by visually
assigning percentages in a look and see manner, I think this is subject to error too in
reporting the coverage of each Cyperaceae member. They are supposed to use a more
objective method of estimating the abundance of the sedges.

5. There is a contradiction in reporting the result for species found in the habitats densely
covered with graminoid vegetation and those that are found in habitats which are sparsely
covered by graminoid vegetation and I am not sure if this was an error.

6. The article could use a thorough review for grammatical errors.

**Is the work clearly and accurately presented and does it cite the current literature?**

Yes

**Is the study design appropriate and is the work technically sound?**

Partly

**Are sufficient details of methods and analysis provided to allow replication by others?**

Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**

Yes

**Are all the source data underlying the results available to ensure full reproducibility?**

Yes

**Are the conclusions drawn adequately supported by the results?**

Yes

**Competing Interests:** No competing interests were disclosed.
Reviewer Expertise: Medical Entomology, epidemiology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 29 Sep 2020

Getachew Bokore, International Centre of Insect Physiology and Ecology (icipe), P.O. Box 30772–00100, Nairobi, Kenya

We appreciate the constructive comments of the reviewer. Below we provide a point by point response.

1. Exploitation of the oviposition cues can be very important in implementing the attract and kill mosquito control technique. The authors should identify and discuss more chemical cues that would potentially attract gravid females to lay eggs in the graminoid other than cedrol present in Cyperus rotundus in the other members of Cyperaceae family identified in this article.

We included some discussion on this as suggested.

2. The method used to estimate the perimeter of individual habitats is subject to errors incase two different people are involved since one person's step cannot be exactly be the same to another ones. I suggest a verifiable method ought to have been used in evaluating habitat sizes.

We clarified in the methods that the perimeter was estimated always by the same person. The perimeter was a relative estimate rather than a precise measure which we did not consider necessary in context of our study.

3. The author indicates that he performed test for abiotic factors like turbidity, pH and temperature only. I think that the parameters were not the only abiotic factors that would influence the distribution of immature stages of malaria vectors. Abiotic factors like Dissolved oxygen (DO), salinity atmospheric pressure could also influence the abundance and distribution of mosquito larvae in the water and ought to have been evaluated.

We included a justification for the selection of the measures in the method section. All habitats surveyed were very similar in their characteristics, hence no major variation was expected; however, it cannot be excluded and we aimed to interpret the work carefully within its discussed limitations.

4. In establishing the coverage of the various graminoid plants in the larval habitats by visually assigning percentages in a look and see manner, I think this is subject to error too in reporting the coverage of each Cyperaceae member. They are supposed to use a more objective method of estimating the abundance of the sedges.
Indeed for an ecological mapping of plant cover, there are various methods available for sampling, for example, the quadrant method and similar approaches. We explored and piloted some of these methods prior to the survey, however did not find them very informative or feasible given the nature of habitats. The most common way to measure cover is the visual estimation method. Visual estimation is popular because it is fast, requires no specialized equipment, and can be adapted to plants of various growth forms. Again, we have clarified that the estimation was done by a single person for relative comparability across sites. In the light of our findings, we would not expect that a different method would have led to a different conclusion.

5. There is a contradiction in reporting the result for species found in the habitats densely covered with graminoid vegetation and those that are found in habitats which are sparsely covered by graminoid vegetation and I am not sure if this was an error.

We were not able to locate the contradiction, possibly this was a misunderstanding?

6. The article could use a thorough review for grammatical errors.

We have gone through the article and corrected the English for errors.

**Competing Interests:** No competing interests were disclosed.

**Reviewer Report 08 September 2020**

https://doi.org/10.5256/f1000research.28336.r70203

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Sisay Dugassa

Vector Biology and Control Research Unit, Aklilu Lemma Institute of Pathobiology, Addis Ababa University, Addis Ababa, Ethiopia

Presentation of the work is very clear with a well designed methodology. Moreover, the analyses and interpretation of the results were properly presented. Importantly, the authors analyzed the association of early instar larvae of *Anopheles* with graminoid plants and reported that “*Anopheles coustani*, *An. rufipes* and *An. maculipalpis* were found only in aquatic habitats covered with graminoid plants, whereas *An. arabiensis*, *An. ziemanni* and *An. pharoensis* were found in both habitats with and without graminoid plants”. Moreover, they analyzed the correlation between the dominant graminoid plant species and early instar larvae and indicted the correlation is not species dependent. Such results are very important for future work in this area of research. However, there might be less importance of graminoid plants for some *Anopheles* species such as *An. arabiensis*, *An. ziemanni* and *An. pharoensis*. Therefore, authors should clearly indicate the
potential importance of other factors than the plant species for the availability and density of the larvae (at least for *An. arabiensis*, *An. ziemanni* and *An. pharoensis*) in their conclusion section.

**Is the work clearly and accurately presented and does it cite the current literature?**
Yes

**Is the study design appropriate and is the work technically sound?**
Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**
Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**
Yes

**Are all the source data underlying the results available to ensure full reproducibility?**
Yes

**Are the conclusions drawn adequately supported by the results?**
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Medical Entomologist

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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