Lymphatic filariasis, infection status in *Culex quinquefasciatus* and *Anopheles* species after six rounds of mass drug administration in Masasi District, Tanzania

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**Abstract**

**Background:** Lymphatic filariasis (LF) elimination program in Tanzania started in 2000 in response to the Global program for the elimination of LF by 2020. Evidence shows a persistent LF transmission despite more than a decade of mass drug administration (MDA). It is advocated that, regular monitoring should be conducted in endemic areas to evaluate the progress towards elimination and detect resurgence of the disease timely. This study was therefore designed to assess the status of *Wuchereria bancrofti* infection in *Culex quinquefasciatus* and *Anopheles* species after six rounds of MDA in Masasi District, South Eastern Tanzania.

**Methods:** Mosquitoes were collected between June and July 2019 using Center for Diseases Control (CDC) light traps and gravid traps for indoor and outdoor respectively. The collected mosquitoes were morphologically identified into respective species. Dissections and PCR were carried out to detect *W. bancrofti* infection. Questionnaire survey and checklist were used to assess vector control interventions and household environment respectively. A Poisson regression model was run to determine the effects of household environment on filarial vector density.

**Results:** Overall, 12,452 mosquitoes were collected of which 10,545 (84.7%) were filarial vectors. Of these, *Anopheles gambiae* complex, An. funestus group and *Cx. quinquefasciatus* accounted for 0.1%, 0.7% and 99.2% respectively. A total of 365 pools of *Cx. quinquefasciatus* (each with 20 mosquitoes) and 46 individual samples of *Anopheles* species were analyzed by PCR. For *Cx. quinquefasciatus* pools, 33 were positive for *W. bancrofti*, giving an infection rate of 0.5%, while the 46 samples of *Anopheles* species were all negative. All 1859 dissected mosquitoes analyzed by microscopy were also negative. Households with modern latrines had less mosquitoes than those with pit latrines (OR = 0.407, P < 0.05). Houses with unscreened windows had more mosquitoes as compared to those with screened windows (OR = 2.125, P < 0.05). More than 80% of the participants own bednets while 16.5% had no protection.

**Conclusions:** LF low transmission is still ongoing in Masasi District after six rounds of MDA and vector control interventions. The findings also suggest that molecular tools may be essential for xenomonitoring LF transmission during elimination phase.

**Keywords:** Lymphatic filariasis, *Wuchereria bancrofti*, *Culex quinquefasciatus*, *Anopheles gambiae*, *Anopheles funestus*, Mass drug administration, Infection rate
LF is caused by the filarial nematode *Wuchereria bancrofti*, transmitted by mosquitoes of the species of *Culex quinquefasciatus*, *Anopheles gambiae* s.l. and *Anopheles funestus* [3–5].

LF has been targeted as a public health problem for elimination by the World Health Assembly due to the fact that, the disease causes disability and may be irreversible if not detected and treated on time. The World Health Organization (WHO) launched the Global Programme for Elimination of Lymphatic Filariasis in 2000 with the aim of interrupting and eventually halting the transmission through repeated mass drug administration (MDA) of anthelmintics. Depending on the country situation, ivermectin (IVM) or diethylcarbamazine citrate (DEC) in combination with albendazole (ALB) were recommended [6]. The effect of IVM and DEC is to kill microfilariae [7, 8], albendazole has no effect on LF microfilaria so the addition of ALB in MDA is meant to synergize the elimination of soil transmitted helminths (STH) [9]. DEC is not recommended in some countries of sub-Saharan Africa due to the co-endemicity with onchocerciasis. Administration of DEC in patients with onchocerciasis causes serious adverse effects such as encephalopathy, confusion, stupor, or coma [10]. Therefore, in many countries of sub-Saharan Africa, IVM and ALB are used for MDA campaigns against LF. In Tanzania, MDA involves an annual single dose administration of IVM plus ALB. After five rounds of annual MDA, the prevalence of microfilaria (MF) in endemic settings was expected to fall below 1% and hence reducing the potential for new transmission by mosquitoes [11]. The success of this strategy is evidenced in Western Pacific Region including Cambodia, Cook Islands, Egypt, Maldives, Marshall Islands, Niue, Sri Lanka, Thailand, Tonga, and Vanuatu where the MF prevalence fell below 1% after several rounds of implementing MDA with DEC and ALB [12–14]. Nevertheless, the elimination target has not been met in Sub-Saharan Africa after more than a decade of MDA, but there has been remarkable progress towards LF control and elimination in the region. Recently, Togo was declared as the first country in sub-Saharan Africa to achieve LF elimination following six rounds of MDA with ivermectin and albendazole using a network of community health workers [15].

In addition to MDA which is the main strategy for LF elimination programs, there has been a growing recognition on the potential role of vector control as a supplementary component to MDA [16–18]. Some studies demonstrated that, the use of insecticide treated bed nets (ITNs) resulted in reduction in prevalence and transmission of LF [19–21]. A study conducted in Papua New Guinea revealed a significant decrease in *W. bancrofti* infection rate among *Anopheles* mosquitoes from 1.8% to 0.4% following distribution of ITNs [19]. Promoting vector control strategies in addition to MDA may have a significant contribution towards elimination of LF.

LF is widespread in many regions of Tanzania with an estimated six million people with disabilities due to the disease [22]. In response to WHO efforts on LF elimination, the National Lymphatic Filariasis Elimination program (NLFEP) began its operations in 2000, using ivermectin (150–200 μg/kg) and albendazole (400 mg) for individuals aged five years and above in selected endemic areas [22]. Since then, there have been evidence of decline in LF transmission in human populations [23–27]. Low infection and infectivity rates in mosquitoes have been demonstrated by studies in North Eastern Tanzania [26, 28] and Rufiji, South Eastern Tanzania [24].

Despite the reported decline in LF in Tanzania, the results from a recent entomological study established evidence of potential for on-going transmission of *W. bancrofti* in Mafia Island after 15 rounds of MDA [29]. It is therefore essential to conduct disease monitoring surveys both in areas undergoing MDA and after stopping MDA in order to evaluate the progress towards elimination and early detection of resurgence respectively. In addition to human blood testing for the presence of the parasites and detection of filarial antigenemia, xenomonitoring in filarial vectors has been considered as an integral component in monitoring the impact of MDA [16, 29, 30]. Xenomonitoring provides real time estimate of infection status where mosquitoes can be collected and assessed either through dissections to find the filarial larvae, or through the use of molecular methods to detect the DNA of the filarial worms [28, 29].

The implementation of MDA campaigns in Masasi district, South Eastern Tanzania, started in the year 2012, with a baseline circulating filarial antigen (CFA) prevalence of 11.7%. However, there is paucity of information on the current infection status in human and vector populations in the district. This study was therefore designed to assess the status of *Wuchereria bancrofti* infection in *Culex quinquefasciatus* and *Anopheles* species, after six rounds of MDA in Masasi District.

**Methods**

**Study site**

The study was conducted in Masasi District (10.7348° S, 38.8044° E) in Mtwara Region. It is bordered to the North by Lindi Region, to the East by the Newala District, to the South by the Ruvuma River and Mozambique, and to the West by Nanyumbu District. The population of the area is 247 993 [32], the inhabitants are mainly Makone, Makua and Wayao tribes. The average annual temperature of the area is 25.4 °C and average annual rainfall is 1024 mm with humidity of 82%.
main socio-economic activities include cashewnut farming. Masasi is an endemic district for LF; it is among the districts which are currently under MDA with ivermectin and albendazole. The implementation of LF elimination activities in the district started in 2012, with baseline CFA prevalence of 11.7% which had significantly declined to 4.7% in 2018 following six rounds of MDA.

**Study design**
This was a cross sectional study conducted in two villages namely Maparagwe and Mbuyuni which were purposely selected from 20 villages of Masasi District Council because they were considered to have high transmission. Maparagwe is in the North, 10 km from the town center, while Mbuyuni is in the South, approximately 40 km from Masasi town center (Fig. 1). For each of the two villages, 25 households were selected giving a total of 50 households. Maparagwe Village consists of five hamlets, in each hamlet, five households were randomly selected. While Mbuyuni consists of six hamlets, four households were randomly selected from each of the hamlets. The remaining one house was selected from the randomly selected hamlet in the list of the six hamlets taking into consideration the distance from the previously selected households. Prior to commencing the study; meetings were conducted with village leadership, community drug distributors (CDDs) and influential people in each respective village. The purpose of the study and methods were clarified to the community representatives. Each of the selected households was visited by the research team to seek for informed consent from the head of the household regarding participation in the study. The environmental features of the household including type of house, latrine type, presence of stagnant waters, tall grasses/bushes were recorded. A survey was also conducted to collect information on bed net ownership and use, indoor residual sprays (IRS), and mosquito repellent use as well as coverage of ITNs in the community.

**Mosquito collection procedures**
Mosquito collection was conducted from mid-June 2019 to late July 2019. Two types of traps were used in mosquito trapping, the U.S. Centers for Disease Control and Prevention (CDC) Light traps (John W. Hock Co. Gainesville, FL) were set indoor, and CDC Gravid traps (John W. Hock Co. Gainesville, FL), containing grass infusion were set in the outdoor position. Light traps are effective in collecting *Anopheles* mosquitoes while gravid traps are effective in collecting *Cx. quinquefasciatus* [28, 33]. The presence of grass infusion provide oviposition cues and potentially collect gravid mosquitoes as they approach the organic infusion in the pan below the trap. Each night, five light traps were set indoors, and five gravid traps were set outdoors at the selected houses. Two traps were set at each house, the indoor and outdoor trap were only set twice throughout the time of the study. Indoor mosquito collection was conducted in one room with an occupant(s) sleeping under mosquito nets. Light traps were set near occupied bed nets at the foot end, at approximately 1.5 m from the ground, as described previously [34, 35]. Gravid traps containing 4 L of grass infusion were placed outdoor as described in previous studies [28]. Briefly, grass infusion was prepared by soaking grass in water for 24 h in a covered plastic bucket to prevent any mosquito oviposition. Both light and gravid traps were set between 18:00 and 19:00 PM and collected early in the morning of the next day between 6:00 and 7:00 AM. Trap nets were removed from the traps and returned to the processing area for morphological identification, dissection and packaging.

**Morphological identification and packaging**
The collected mosquitoes were morphologically identified to species, based on available keys for *Anophelines* [36] and *Culicines* [37]. Briefly, mosquitoes were identified to their respective species based on common structural features including: wings, abdomen, head, thorax, legs, mouth parts and scales. After identification, mosquitoes were counted and segregated into filarial and non-filarial vectors. Individual filarial vectors were recorded in a mosquito recording sheet. Female *Cx. quinquefasciatus* mosquitoes were stored in dry clean eppendorf tubes with silica gel in a pool of 20 mosquitoes while *Anopheles* species were stored individually for molecular analysis by PCR.

**Detection of Wuchereria bancrofti in mosquitoes**

**Dissection and detection of W. bancrofti larval stages by microscopy**
A total of 1822 freshly killed female *Cx. quinquefasciatus* and 37 *Anopheles* species were dissected to assess if they harbored any stage of *W. bancrofti* larvae. Dissection standard procedures were followed, as previously described [29, 38]. Dissection results were recorded into a designed sheet. Later on, the data was entered into Microsoft excel data-base then imported into SPSS version 22 (SPSS, Inc., IL, USA) for analysis. Mosquito infection was defined as the presence of any larval stages, first stage larvae (L1), second stage larvae (L2), and/or third stage larvae (L3), while mosquito infectivity was defined as the presence of L3 larvae in any of the body segments [39].

**Dissection and detection of W. bancrofti larval stages by molecular methods**
A total of 1822 freshly killed female *Cx. quinquefasciatus* and 37 *Anopheles* species were stored in dry clean eppendorf tubes with silica gel in a pool of 20 mosquitoes while *Anopheles* species were stored individually for molecular analysis by PCR.
Detection of *W. bancrofti* by polymerase chain reaction (PCR)

**Deoxyribonucleic acid (DNA) extraction from mosquitoes**
Mosquito genomic DNA (gDNA) extraction was carried out by using the Chelex-100 Resin with modifications as described earlier [3]. Briefly, the pulsating vortex machine was used for homogenizing mosquitoes (20 in each tube) in 250 μl of 10% chelex buffer solution (C7901, Sigma, CA, USA). The Extracted gDNA was analyzed for the presence of *W. bancrofti* DNA by PCR targeting highly repetitive amino acid sequences of *W. bancrofti* DNA. Each reaction mixture consisted of 0.125 μmol/L of forward and reverse primers, 10 μl Hot StartTaqTaqPase polymerase master mix and 2 μl of DNA extract. The amplified DNA for *W. bancrofti* specimen were separated based on their fragment size by gel electrophoresis and visualized under ultra violet light as previously described [40].

Assessment of household characteristics and surrounding environment

A survey was conducted to collected information on the characteristics of households and the surrounding environment. This information included type of the house based on construction materials which were classified as, brick with iron roof, brick with grass roof and mud with grass roof. Window screening, the type of latrine, the presence of tall grasses and bushes around the house, and the presence of stagnant waters was also recorded (Additional file 1).

Assessment of vector control strategies in the study community

Pre-tested questionnaires were administered to 588 individuals in the study area to obtain information on use of vector control interventions. The questions included but were not limited to: whether they own and use bed net(s), whether the bed nets were insecticide treated or not treated. Information on other vector...
control interventions was also sought, such as use of IRS and mosquito repellent (Additional file 2).

Data analysis
All data were entered in Excel spreadsheets (Microsoft corp., Redmond, USA) and transferred to SPSS version 20.0 (SPSS, Inc., IL, and USA). The ‘infection rate’ of the dissected mosquitoes was calculated as the percentage of mosquitoes infected with any stages of W. bancrofti that is L1, L2 or L3. While the ‘infectivity rate’ was calculated as the percentage of mosquitoes infected with infective larvae (L3) as previously described [29]. For pooled mosquitoes which were analyzed by PCR technique; The Pool Screen (v.2.02) software (Department of Biostatistics and Division of Geographic Medicine, University of Alabama at Birmingham, USA) [22] was used to calculate the probability that any one of the mosquitoes is infected with any stage of W. bancrofti. Two sample t-tests for proportions were used to compare the infection rates among mosquitoes caught indoor and outdoor, and between the two villages. A P-value of less than or equal to 0.05 was considered statistically significant. Poisson regression model was run to assess the influence of household environments on vector density. Whereby, vector density was modeled as the function of house type, latrine type, and window screen, presence of tall grasses, bushes and stagnant waters around the house. The differences in bed net ownership between the two villages were compared using chi-square tests, where P ≤ 0.05 were considered as statistically significant.

Results
Mosquito populations and composition
A total of 12 452 mosquitoes were collected in the two villages during the study period, whereby, 7860 (63.1%) were collected from Maparagwe and 4592 (36.9%) from Mbuyuni village. Of the total mosquitoes collected, 1868 (15%) were collected indoor, while 10 583 (85%) were collected outdoor. Majority 10 545 (84.7%) of the collected mosquitoes were filarial vectors. The remaining 1907 (15.3%) were mosquitoes belonging to the species of Culex sinilius, Coquillettidia spp. and Aedes spp. The composition of the filarial vectors included An. gambiae complex 15 (0.1%), An. funestus group 73 (0.7%) and Cx. quinquefasciatus 10 457 (99.2%) (Table 1).

Microscopy examination
A total of 1822 female Cx. quinquefasciatus and 37 female Anopheles species from both CDC light and gravid traps were dissected and examined for infection with W. bancrofti larvae giving a total of 1859 dissected mosquitoes. None of the 1859 dissected mosquitoes were found to be infected with any of the larval stages (L1, L2 and/or L3) of W. bancrofti (Table 2).

Molecular analysis of mosquito samples
Using PCR technique, a total of 365 pools of Cx. quinquefasciatus each containing 20 mosquitoes and 46 individual Anopheles species (An. gambiae and An. funestus) were tested for infection with W. bancrofti. Of the 365 pools of Cx. quinquefasciatus, 33 were found to be infected with W.bancrofti. All 46 Anopheles samples were found to be negative. Analysis by study site indicates a significant difference in infection rate between the two study villages (two sample, t-test for proportions, P = 0.004). For both species and study villages, the probability that any one mosquito in the pool was infected with any stage of W. bancrofti parasite was estimated at 0.5% (Table 2).

Characteristics and environmental features of the sampled households
Overall, 94.0% (n = 47) of sampled households (n = 50) had pit latrines. Only, 14.0% (n = 7) of households had screened windows to prevent mosquito entry. None of them had stagnant waters around; this is because the survey was done during the dry season, between June and July. In addition to that the majority of the households 98% (n = 48) had no tall grasses and bushes around (Table 3).

Effects of household characteristics and the surrounding environment on vector density
The results showed that, houses constructed with bricks had significantly fewer mosquitoes compared to those constructed with mud regardless of the roof type (iron sheets or grass) (OR = 0.638 and OR = 0.412, respectively P < 0.05). Houses with unscreened windows had two times more mosquitoes as compared to those with screened windows, (OR = 2.125, P < 0.05). Households with modern latrines had 60% times less mosquitoes as compared to the households with pit latrines (OR = 0.407, P < 0.05). Households without tall grasses around were four times more likely to have mosquitoes compared to the households with tall grasses in the surrounding (OR = 4.320, P < 0.05) (Table 4).

Vector control interventions in the study community
Bednet use was the main vector control intervention where 80.8% (475/588) bednet ownership was recorded in the community. While only about two percent used mosquito repellent and IRS use only one percent (Fig. 2). The majority, 78.7% reported to have slept under bednets
during the last night before the day of the interview and among those who own bednets; only 52.6% had ITNs (Fig. 3). Analysis of bednet ownership by village revealed no statistically significant differences between the two villages (Chi-square = 0.673, df = 1, P = 0.412).

**Discussion**

The global programme for elimination of LF advocates annual MDA with ALB and IVM or DEC in specific endemic areas, complimented with regular monitoring of the disease status towards elimination. This study was therefore designed to assess the LF infection status in filarial vectors after six rounds of MDA in Masasi District.

The current study reports an infection rate of 0.5% in mosquitoes after six rounds of MDA which indicates a low ongoing LF transmission in the district. The obtained infection rate is above the cut-off point of 0.25%, a threshold that has been suggested for areas where *Culex* mosquitoes are the vectors [41]. This suggests a potential for persistent transmission which may be facilitated by presence of the parasite reservoirs [42]. In Tanga Region, where LF is also endemic, six rounds of MDA resulted into a decline in vector infection rate by 99.3% [43] and a follow up study reported no infection in mosquitoes after eight rounds of MDA in the same region [23]. It is therefore evident that, there is significant progress towards interrupting LF transmission in Tanzania using MDA interventions.

The infection rate obtained in this district is four times higher than the previously reported infection rate of 0.1% in Rufiji district after twelve rounds of MDA [24], which is in the same geographical region. But it is four times less than the infection rate of 1.7% reported in Mafia Island after 15 rounds of MDA [29]. The observed differences might be attributable to a number of factors, including but not limited to: the initial level of LF prevalence and density of microfilaremia, the competence and vectorial capacity of the local vectors as well as population coverage and compliance to MDA [42]. The effect of some of these factors is exemplified by the reports in Mafia Island and Masasi District, where the baseline CFA prevalence in Mafia Island was 49%, while that of Masasi District was 11.7%. The observed differences in the levels of infection rates were 1.7% vs 0.5% respectively; which may be inherent in the significantly different levels of infection status at the initiation of the MDA program.

### Table 1  Mosquito populations and composition from two villages in Masasi District

| Mosquito taxa                  | Maparagwe Village | Mbuyuni Village | Total collection, n (%) |
|-------------------------------|-------------------|-----------------|-------------------------|
|                               | Indoor, n (%)     | Outdoor, n (%)  | Indoor, n (%)           | Outdoor, n (%)  |                     |
| **Anopheles gambiae complex** | 15 (1.1)          | 0 (0.0)         | 0 (0.0)                 | 0 (0.0)         | 15 (0.1)           |
| **An. funestus group**        | 63 (4.4)          | 0 (0.0)         | 10 (2.2)                | 0 (0.0)         | 73 (0.6)           |
| **Culex quinquefasciatus**    | 1199 (84.3)       | 5484 (85.2)     | 347 (77.6)              | 3427 (82.7)     | 10 457 (84)        |
| **Other mosquitoes**          | 145 (10.2)        | 954 (14.8)      | 90 (20.1)               | 718 (17.3)      | 1907 (15.3)        |
| **Total by trap/village**     | 1422              | 6438            | 447                     | 4145            | 12 452             |

*Other mosquito species include Culex sinilius, Coquilettidia spp. and Aedes spp.*

### Table 2  Vector infection rate by methods of analysis and study site

| Analysis method | Mosquito species | No. analysed | No. infected (%) | 95% CI       | P-value |
|-----------------|------------------|--------------|------------------|-------------|---------|
| **Microscope**  | Anopheles spp.   | 37           | 0                | 0.0–0.095   |         |
|                 | Culex quinquefasciatus | 1822      | 0                | 0.0–0.001   |         |
|                 | All species      | 1859         | 0                | 0.0–0.001   |         |
| **PCR**         | Anopheles spp.   | 46†          | 0                | 0.0–0.077   |         |
|                 | Culex quinquefasciatus | 365†⁺⁺      | 33 (0.5)†        | 0.44–0.46   | 0.1     |
|                 | All species      | 411          | 33 (0.5)†        | 0.45–0.46   |         |
| **Study site**  | Maparagwe        | 280          | 15 (0.3)‡        | 0.256–0.36  |         |
|                 | Mbuyuni          | 131          | 18 (0.7)‡        | 0.61–0.77   | 0.004*  |
|                 | All villages     | 411          | 33 (0.5)†        | 0.45–0.46   |         |

*CI confidence interval
* † † † Pool of 20 mosquitoes; † Individual/Single mosquito; † Infection rate (Pool Screen, V2.0.2; Likelihood ratio method); * Two sample test for proportion to compare mosquito infection rates between villages
addition the number of MDA rounds, the coverage and community compliance to the MDA programme would to a great extent influence the decline in parasite prevalence, microfilaria density and hence the transmission dynamics and vector infectivity levels [42].

The findings of this study indicate that, out of 1859 microscopically dissected mosquito samples, no single mosquito was found to be infected with any larval stages of *W. bancrofti*, while analysis by PCR revealed an infection rate of 0.5% (Table 2). These observations emphasize the superiority of molecular techniques for xenomonitoring in settings with low LF transmission [30, 31]. The findings of this study are corroborated by the findings of a similar study in Mafia Island, Tanzania which reported infection rates of 0.3% by microscopy technique vs 1.7% by PCR technique [29]. Given the existing low transmission levels in most endemic communities during the implementation of the elimination program, molecular based techniques may be an effective tool for xenomonitoring.

**Table 3 Characteristics and environmental features of the samples households**

| Features                  | Status/observation | Number of households |
|---------------------------|--------------------|----------------------|
| House type                |                    |                      |
| Bricks/grass roof         | Yes                | 16                   |
|                           | No                 | 34                   |
| Bricks/iron roof          | Yes                | 32                   |
|                           | No                 | 18                   |
| Mud/grass roof            | Yes                | 2                    |
|                           | No                 | 48                   |
| Window                    |                    |                      |
| Not screened              | 43                 |
| Screened*                 | 7                  |
| Latrine type              |                    |                      |
| Modern                    | Yes                | 3                    |
|                           | No                 | 47                   |
| Pit latrine               | Yes                | 47                   |
|                           | No                 | 3                    |
| Uses bednets              | Yes                | 48                   |
|                           | No                 | 2                    |
| Uses bednets + IRS        | Yes                | 2                    |
|                           | No                 | 48                   |
| Presence of tall grasses  |                    |                      |
| Around the house          | Yes                | 48                   |
|                           | No                 | 2                    |
| Presence of bushes around house | No | 48   |
| Presence of stagnant water| Yes                | 0                    |
|                           | No                 | 50                   |

Table 4 Effects of household characteristics and surrounding environment on vector density (*n* = 50)

| Variable                              | Category   | OR          | 95% CI        | *P*-value |
|---------------------------------------|------------|-------------|---------------|-----------|
| House type                            | Bricks/grass roof | 0.412       | 0.385–0.440   | 0.000     |
|                                       | Bricks/iron roof | 0.644       | 0.606–0.684   | 0.000     |
|                                       | Mud/wood*     | 1           |               |           |
| Window                                | Not screened | 2.125       | 1.970–2.293   | 0.000     |
|                                       | Screened*     | 1           |               |           |
| Type of latrine                       | Modern latrine | 0.407       | 0.362–0.457   | 0.000     |
|                                       | Pit latrine*  | 1           |               |           |
| Presence of tall grasses around the house | No  | 4.320       | 3.461–5.391   | 0.000     |
|                                       | Yes*          | 1           |               |           |

*P* ≤ 0.05 is significant
OR odds ratio, CI confidence interval
*Reference category

The current findings also indicate that, *Cx. quinquefasciatus* is a dominant vector of *W. bancrofti* in Masasi District, which accounted for 99.2% of the filarial vectors sampled. More mosquitoes were caught outdoor using CDC gravid traps compared to the indoor with CDC light traps. It is hypothesized that, the difference in trap performance may be due to the organic grass infusion attractant added to the CDC gravid traps.

The dominance of *Cx. quinquefasciatus* has been reported in a number of earlier studies [29, 44, 45] and has been considered as a dominant vector in urban areas [46]. Interestingly, in recent years studies have shown that *Cx. quinquefasciatus* is now a dominant vector both in rural and urban settings [47]. A study in North Eastern Tanzania, demonstrated a major shift in vector species composition; from predominantly *Anopheles* in the pre-MDA period to almost exclusively *Culicines* after six rounds of MDA [43]. This shift in vector composition has been linked to multiple factors including; environmental and climate changes as well as documented evidence that IVM may have effects on *Anopheles* species.

The current study demonstrated low population densities of *An. gambiae* and *An. funestus*. It is possible that the observed low densities of *Anopheles* species from this study may be linked to the effects of ivermectin which is administered annually in the study area. Increased mortality and decreased fecundity of *Anopheles* species taking blood meals on ivermectin treated individuals has been demonstrated in a number of clinical trials [48, 49], field reports [50, 51] and laboratory studies [52–54].

The current study findings also indicate that, some household characteristics including pit latrine and unscreened windows have influence on density of filarial vectors. Households with pit latrines had significantly...
higher counts of mosquitoes collected compared to households with modern latrines (Table 4). This observation is not surprising because pit latrines are known to be breeding habitats for *Culex* mosquitoes as they prefer to breed in polluted waters [45]. This view is corroborated by findings from an intervention study in which pit latrines and septic tanks, were treated with polystyrene beads in Dar es salaam region and the outcome was a significant reduction in densities of adult *Cx. quinquefasciatus* [55]. Unscreened windows were also associated with increased indoor mosquito densities which present increased biting rates to the house occupants and hence increased risk of LF infection. These findings coupled with the fact that *Cx. quinquefasciatus* seems to expand its horizons to both urban and rural areas; may suggest that, vector control focusing on environmental improvement may be an important factor in the LF elimination program in some of the endemic areas.

The National Malaria Control Programme in Tanzania has significantly contributed towards vector control, through its campaigns on mosquito net distribution to vulnerable groups (pregnant women and children). Thus high coverage of bednets in both villages was recorded in this study. The use of ITNs has been linked to the decline in populations of *Anopheles* species [56, 57]. Very low densities of *Anopheles* mosquitoes were observed in this study, which could be linked to both the use of ITNs and the IVM effects on these mosquito populations. Although nearly half of the bednet owners used un-treated bednets, it offers physical protection by preventing human-mosquito contact and hence reducing the risk of infection [58]. The use of ITNs may therefore be of value to both malaria and LF control programs and should be advocated by both programs, even though vector control is not advocated in the LF elimination strategy.

The lethal effects of IVM on *Anopheles* mosquitoes have been demonstrated [48, 50–52, 54, 59]. However, IVM does not seem to have lethal effects on *Culex* species [53]. Laboratory studies using adult *Cx. quinquefasciatus* fed on blood meals from volunteers treated with ivermectin revealed no effects on its survival, egg laying capacity and development of larvae [53]. Though, mortality of *Cx. quinquefasciatus* was reported when fed on chicken treated with 2000 µg/kg of ivermectin (about ten times the therapeutic dosage) [60]. In addition, *Cx. quinquefasciatus* are relatively tolerant to insecticides used for ITNs and IRS interventions [61, 62]. It may therefore be important to develop effective vector control
tools against this mosquito species in order to maximize the impact of MDA.

The main limitation of the findings of this study is that, mosquito collection covered only 50 households, which may not be sufficient to represent the vector dynamics and infection status in the district. Furthermore, data collection was done during the dry season between June and July, which may have impacted the density of *Anopheles* mosquitoes. Earlier studies have reported that there is seasonal variation in the density of *Anopheles* mosquitoes with a decline during the dry season due to shortage of breeding habitats [63. Therefore, a longitudinal entomological survey is needed to assess, effects of seasonality on vector density, vector species, transmission dynamics and the potential risk factors.

**Conclusions**

LF low transmission is still ongoing in Masasi District after six rounds of MDA and vector control interventions which are in place. There also seem to be a shift in filarial vector transmission and population dynamics in the rural setting with *Cx. quinquefasciatus* dominating over the *Anopheles* species. The findings also suggest that molecular tools may be essential for xenomonitoring in assessment of LF transmission during the elimination phase. Based on the findings of this study, it may be reasonable to suggest that, in order to halt LF transmission as per global LF elimination initiative, an integrated strategy is essential. Integrated vector control strategies including use of ITNs, environmental improvement and modernization of latrines to limit vector breeding habitats for *Cx. quinquefasciatus* may be an important addition to the MDA strategy.

**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1186/s40249-021-00808-5.

**Additional file 1:** Checklist on characteristics of the household and the surrounding environment.

**Additional file 2:** Questionnaires for community lymphatic filariasis screening in Masasi District council.

**Abbreviations**

ALB: Albendazole; CI: Confidence interval; CDC: Center for disease control; CDDs: Community drug distributors; CFA: Circulating Filarial Antigen; DEC: Diethylcarbamizine; df: Degree of freedom; ITNs: Insecticide treated nets; IRS: Indoor residual spray; IVM: Ivermectin; LF: Lymphatic filariasis; LT: Light trap;
MDA: Mass drug administration; OR: Odd ratio; PCR: Polymerase chain reaction; SPSS: Statistical package for Social Sciences.

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Authors’ contributions

EL, DG and OM conceived the study, DG: Supervised data collection, EL: Performed data collection, Data analysis and wrote the draft of manuscript. DG& OM: critically reviewed the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Ethics and consent to participate

The ethical approval to conduct the study was obtained from the Research and publications Committee of Muhimbili University of Health and Allied sciences, Tanzania. Before starting data collection; meetings were conducted with the district and respective village and ten cell leaders, community drug distributors and influential people in the village to clarify the aim of the study and publications Committee of Muhimbili University of Health and Allied Sciences through Sida funding. The funders had no role in study design, data collection, analysis, and decision to publish, or preparation of the manuscript.

Consent for publication

Not applicable

Competing interests

The authors declare that they have no competing interests.

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