**Abstract**

**Background:** The Thailand-Myanmar borderland is an area endemic for malaria where transmission is low, seasonal and unstable. The epidemiology has been described but there is relatively few data on the entomological determinants of malaria transmission.

**Methods:** Entomological investigations were conducted during 24 months in four villages located in Kayin state, on the Myanmar side of the Thailand-Myanmar border. *Anopheles* mosquitoes were identified by morphology, and molecular assays were used in order to discriminate between closely related sibling species of malaria vectors. *Plasmodium* infection rate was determined using quantitative real-time PCR.

**Results:** The diversity of *Anopheles* mosquitoes was very high and multiple species were identified as malaria vectors. The intensity of human-vector contact (mean human-biting rate= 369 bites/person/month) compensates for the low infection rate in naturally infected populations of malaria vectors (mean sporozoite index= 0.04 and 0.17 % for *P. falciparum* and *P. vivax* respectively), yielding intermediary level of transmission intensity (mean entomological inoculation rate= 0.13 and 0.64 infective bites/person/month for *P. falciparum* and *P. vivax*, respectively). Only 36% of the infected...
mosquitoes were collected indoors between 09:00 pm and 05:00 am, suggesting that mosquito bed-nets would fail to prevent most of the infective bites in the study area.

**Conclusion:** This study provided a unique opportunity to describe the entomology of malaria in low transmission settings of Southeast Asia. Our data are important in the context of malaria elimination in the Greater Mekong Subregion.

**Keywords**
Anopheles, human biting rate, sporozoite index, entomological inoculation rate, parasite load, residual transmission, Plasmodium juxtanucleare, zoophagy, exophagy, hypnozoite reservoir

This article is included in the Mahidol Oxford Tropical Medicine Research Unit (MORU) gateway.
Amendments from Version 3

In the version 4 of the manuscript, a sentence on the indoor/outdoor distribution of malaria vectors was rephrased in order to improve its readability by the reader. A type “underestimated” was changed to “overestimated”. In the Shift in vector-control intervention section, “16S ribosomal RNA genes” was changed to “mammalian mt16S ribosomal RNA genes”.

See referee reports

Introduction

Artemisinin resistance in Plasmodium falciparum has emerged and spread in the Greater Mekong Sub-region (GMS), leading to the failure of several artemisinin-based combination therapies (ACTs). Multi-drug resistant parasites spreading from Western Cambodia are responsible for a recent resurgence of the disease across the eastern part of the GMS. Meanwhile in Myanmar (western GMS), the incidence of clinical malaria cases has declined. In this area, dihydroartemisinin-piperaquine and artemether-lumefantrine remain effective against P. falciparum. It is therefore urgent to eliminate falciparum malaria in Myanmar, the main gateway to India and Bangladesh, before these two ACTs also fall to resistance.

Entomological aspects of malaria transmission are important in the context of elimination as they largely determine intervention design and outcome. For example, the interest of treating asymptomatic infections with mass drug administration or mass screening and treatment obviously depends on the contribution of asymptomatic carriers to the transmission. Moreover, the efficacy of vector-control interventions such as mass distribution of long-lasting insecticide-impregnated mosquito bed-nets (LLINs) and indoor residual spraying (IRS) campaigns is greatly influenced by the ecology of malaria vectors.

The transmission of P. falciparum is low, seasonal and unstable in Kayin state (Eastern Myanmar). Some entomological surveys were conducted previously, most of them on the Thai side of the Thailand-Myanmar border, where the transmission of P. falciparum is now interrupted. In this area, the primary vectors are Anopheles minimus (s.s.) (Minimus Complex, Funestus Group), An. maculatus (s.s.), An. sawadwongporn (Maculatus Group), An. dirus (s.s.) and An. baimaii (Dirus Complex, Leucophyurus Group), Anopheles pseudowillmori (Maculatus Group), An. aconitus (s.s.) (Funestus Group) and some members in the Annularis and Barbirostris Groups also play a secondary role in the transmission. Numerous aspects of malaria vector ecology and biology have not been documented and the characteristics of the entomological indices are not known precisely.

Large-scale deployment of community-wide access to early diagnosis and treatments has paved the way for the elimination of falciparum malaria in Kayin state. Mass antimalarial drug administration campaigns were used as an accelerator to elimination in villages with submicroscopic reservoirs of asymptomatic malaria infections. The objective of this study was to describe the entomological determinants of malaria transmission in Kayin state in order to guide policy making for malaria elimination.

Methods

Study villages

Four villages located in Kayin state were included in the study, namely HKT (16° 85’ N, 98° 47’ E), KNH (17° 18’ N, 98° 24’ E), TPN (17° 14’ N, 98° 29’ E) and TOT (16° 36’ N, 98° 57’ E) (Figure 1). The demographics and baseline malaria epidemiology in the study villages were described previously. Briefly, the number of households at baseline was 160, 81, 138, and 69 in HKT, KNH, TOT and TPN respectively. The census population at baseline was 908, 349, 745 and 375 at HKT, KNH, TOT and TPN respectively. Residents were mainly farmers and forest workers. The prevalence of submicroscopic malaria at the beginning of the study ranged between 4 and 22% and between 19 and 33% for P. falciparum and P. vivax respectively.

Intervention

Community-wide access to early diagnosis and treatments was deployed in all villages for the entire period of the study. Mass antimalarial drug administration campaigns with dihydroartemisinin-piperaquine and a single low dose of primaquine were repeated at monthly intervals for 3 months from 12 June to 24 August 2013 in KNH, from 27 May to 07 August 2013 in TOT, from 28 January to 29 March 2014 in TPN and from 01 April to 10 June 2014 in HKT. LLINs were also distributed to all villagers. The intervention and its impact on the parasitological and entomological indices of malaria were described into more details in Chaumeau et al. and Landier et al.

Entomological surveys

Villages were surveyed monthly from 2013 to 2015 for a total of 21 surveys per village. Entomological surveys consisted of five consecutive nights of collection from 06:00 pm to 06:00 am in five houses per village and on one cow, as described previously. There were two exceptions to the initial study design: the eleventh survey in TPN was cancelled and mosquitoes were collected for only two nights during the second survey in HKT. In each house, five traditional houses were randomly selected for mosquito sampling using the human-landing catch (HLC) collection method. In each house, one mosquito collector sat indoors and one mosquito collector sat outdoors, yielding a total of 50 person-nights of collection per survey (25 person-nights indoors and 25 person-nights outdoors). Collectors were asked to collect every mosquito landing on their uncovered legs for 50 min per hour and allowed to rest for 10 min per hour. A cow-bait trap (CBT) was also set-up yielding an additional 5 cow-nights of collection. One cow was isolated from the herd and a 1m-wide mosquito net was fenced around the animal, 30cm above the ground level. One collector was asked to collect mosquitoes resting on the net for 50 min per hour and allowed to rest for 10 min. Mosquitoes were shipped at the Shoklo Malaria Research Unit (Mae Sot, Thailand) at the end of each night of collection.
<table>
<thead>
<tr>
<th>Group</th>
<th>Subgroup</th>
<th>Complex</th>
<th>Species</th>
<th>Vectorial status on the TMB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annularis</td>
<td></td>
<td></td>
<td>An. annularis (s.l.)</td>
<td>secondary vector</td>
</tr>
<tr>
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</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>An. pampanai</td>
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<td></td>
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<td>Culicifacies</td>
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</tr>
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<td></td>
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<td>Dirus</td>
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<tr>
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<td></td>
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<td></td>
<td>An. nemophilus</td>
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<td>An. maculatus (s.s.)</td>
<td>primary vector</td>
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<tr>
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<td>An. dravidicus</td>
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<tr>
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<td>An. sawadwongporni</td>
<td>primary vector</td>
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</tr>
<tr>
<td>Tesselatus</td>
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<td></td>
<td>An. tessellatus</td>
<td>non vector ‡</td>
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</table>

† Plasmodium vivax ‡ Some species in these Groups are efficient malaria vectors elsewhere, although they were never found infected with human malaria parasites in the Thailand-Myanmar border area (e.g. An. culicifacies A, An. sinensis, An. subpictus (s.s.), An. splendidus and An. tessellatus)14,21,22.

Laboratory procedures for the processing of entomological samples

Mosquitoes were immediately identified at the genus level by morphology and Anopheles specimens were stored individually at -20°C in 1.5 mL plastic tubes containing silica gel. Anopheles mosquitoes were identified at the Group or Complex level using the key developed by Rattanarithikul et al.23. Deoxyribonucleic acid (DNA) was extracted from head/thorax using a cetyltrimethyl ammonium bromide-based method described previously24. Sibling species to the Funestus, Maculatus and Leucosphyrus Groups were discriminated using allele-specific polymerase chain reaction (AS-PCR) assays adapted from Garros et al. and Walton et al.25–27. In case AS-PCR yielded a negative result, identification at the species level was performed by sequencing the internal transcribed spacer 2 (ITS2) DNA marker using universal primers described by Beebe and Saul28. DNA extracts were screened for the presence of Plasmodium sporozoites using a quantitative real-time PCR (qPCR) assay targeting 18S rRNA genes adapted from Mangold et al.29. Specificity of the signal was confirmed in all positive samples by amplifying COX genes with primers described by Cunha et al.30. In case the confirmation assay yielded a negative result, the PCR product of the screening assay was sequenced (BioSample accessions: SAMN09845988, SAMN09845989, SAMN09845990, SAMN09845991, SAMN09845992). The presence of Plasmodium oocysts in the abdomen of sporozoites-positive specimens was assessed using the same procedures. Abdomen were preserved at -20°C in 1.5 mL plastic tubes containing silica gel during the time between the screening of head/thorax for sporozoites and the processing of abdomen from sporozoite-positive specimens. The validation of the qPCR assays used for Plasmodium detection in this study was published elsewhere24. Detailed laboratory procedures are presented in the Supplementary File 2.

Data analysis

HBR and CBR were defined as the number of mosquitoes collected divided by the number of person-nights or cow-nights respectively. Results were expressed as a number of bites/person/month or bites/cow/month. Sporozoite index (SI)
was defined as the number of mosquitoes positive in qPCR Plasmodium divided by the total number of mosquitoes analyzed. Results were expressed as a percentage. Entomological inoculation rate (EIR) was defined as the number of specimens positive in qPCR Plasmodium divided by the corresponding number of person-nights of collection. The number of person-nights used for the calculation of EIR was multiplied by the proportion of collected mosquitoes that were analyzed with qPCR Plasmodium in order to take into account that not all mosquitoes were analyzed with qPCR. Results were expressed as a number of infective bites/person/month. The exophagy index (EI) was defined as the outdoor HBR divided by the sum of the indoor and outdoor HBRs. The cow-biting index (CBI) was defined as CBR divided by the sum of the HBR and the CBR. Confidence Intervals (CIs) for Poisson counts (HBR, CBR and EIR) were estimated using the exact method of the poisson.conf.int() function in the epitools package version 0.5-10 of R software version 3.3. Binomial CIs were estimated for proportions (SI, EI and CBI) using the exact method of the binom.confint() function in the binom package version 1.1-1 of R software version 3.3. Species-specific proportions were used to compute HBR, CBR, EI and CBI estimates at the species level if ≥30 samples in the corresponding Anopheles Group were genotyped for each value of the grouping variable. The grouping variables were the collection time (06:00 pm to 06:00 am), the collection method (HLC or CBT) or the location of the mosquito collector (indoors or outdoors). For example, the number of An. maculatus (s.l.) collected by HLC indoors and outdoors was 2240 and 5041 respectively. The proportion of An. maculatus (s.s.) in the indoor and outdoor populations was 0.34 (133/392) and 0.37 (404/1084) respectively. The predicted number of An. maculatus (s.s.) collected by HLC indoors and outdoors was 762 and 1865 respectively. The EI estimate for An. maculatus (s.s.) was therefore 0.71 (1865/2627).

**Ethics approval**

The protocol for mosquito collection and analysis has been approved by the Oxford Tropical Research Ethics Committee (1015–13, dated 29 Apr 2013) and by the Ethics Review Committee for Research Involving Human Research Subjects, Health Science Group, Chulalongkorn University (COA 154/2014). All participants provided their written consent to participate in this study. This consent procedure was approved by the ethics committees.

**Results**

**Anopheles diversity**

In total, 129228 Anopheles mosquitoes were collected during 4120 person-nights and 412 cow-nights of collection (63217 by HLC and 66011 by CBT). We report the occurrence of 10 Groups of Anopheles on the basis of morphological identification: Barbirostris, Hycanmus (Anopheles Subgenus, Myzorynchus Series), Annularis, Jamesii, Maculatus (Cellia Subgenus, Neocellia Series), Funestus (Cellia Subgenus, Myzomyia Series), Kochi, Leucosphyrus, Tessellatus (Cellia Subgenus, Neomyzomyia Series) and Subpictus (Cellia Subgenus, Pyretophorus Series).
Anopheles karwari (*Cellia* Subgenus, *Neocellia* Series) was also detected at a low frequency. Less than 5% (6010/129228) of the specimens could not be identified at the Group level because they were damaged (missing legs or wings).

*A priori* malaria vectors in the Annularis, Barbirostris, Funestus, Leucosphyrus and Maculatus Groups (*i.e.* *Anopheles* taxa that were previously reported to be infected with human malaria parasites in the Thailand-Myanmar border area) accounted for 84–96% and for 40–70% of the *Anopheles* mosquitoes collected by HLC and CBT collection methods respectively (Figure 2). The abundance of malaria vectors was significantly different from one collection site to another in a given village, suggesting that local ecological factors are important drivers of exposure to malaria vectors (Supplementary File 1, Figures S1–S4). The proportion of collection-night during which a given *Anopheles* taxa was collected varied between 3% and 75% in the Subpictus and Funestus Groups respectively (Supplementary File 1, Figures S5–S6). The Funestus Group was the most abundant taxa during both the rainy and dry seasons (June to November and December to May, respectively). The Maculatus and Leucosphyrus Groups were mainly collected during the rainy season. The abundance of Annularis and Barbirostris Groups peaked during the transition period between the rainy and dry seasons, when the abundance of other groups decreased (Supplementary File 1, Figures S7–S10).

Anopheles minimus (s.s.) was by far the most abundant species among the Funestus Group (Table 2). Its sibling *An. harrisoni* and closely related members from the Aconitus and Culicifacies

![Figure 2](image-url)

**Figure 2.** Biodiversity of the *Anopheles* mosquitoes according to the collection method and study village. **A**) Proportion of malaria vectors in *Anopheles* populations collected by human-landing catch (HLC) and cow bait trap (CBT) collection methods according to the village. **B**) Relative proportions of *sensu lato* malaria vectors collected by HLC according to the village. **C**) Relative proportions *sensu lato* malaria vectors collected by CBT according to the village. † *A priori* malaria vectors refers to *Anopheles* taxa that were previously reported to be infected with human malaria parasites in the Thailand-Myanmar border area, *i.e.* the Funestus, Maculatus, Leucosphyrus, Barbirostris and Annularis Groups.

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Subgroups represented <0.5% and 13% of the specimens from the Funestus populations collected by HLC and CBT, respectively. Aconitus and Culicifacies Subgroups represented up to 12% of the Funestus specimens in TPN (Supplementary File 1, Table S2).

Anopheles maculatus (s.s.), An. sawadwongporni and An. pseudowillmori were the most abundant species in the Maculatus Group with proportion varying from 15 to 50%, 40 to 81% and 3.6 to 9% according to the village. Three species from the Dirus Complex accounted for >99% of the specimens in the Leucosphyrus Group, namely An. dirus (s.s.), An. baimaii and An. cracens. The proportion of An. dirus (s.s.) and An. baimaii in study villages varied from 60 to 97% and from 0 to 39%, respectively. The proportion of An. culicifacies B, An. maculatus (s.s.) and An. cracens increased during the dry season whereas that of An. minimus (s.s.), An. pseudowillmori and An. baimaii increased during the rainy season (Supplementary File 1, Table S3).

### Malaria vectors

The contribution of six Groups of Anopheles to malaria transmission was determined by processing 56872 samples collected by HLC in qPCR Plasmodium. Human malaria parasites P. falciparum and P. vivax were detected in 106 Anopheles specimens that belonged to four Groups of Anopheles species (Funestus, Maculatus, Leucosphyrus and Barbirostris) (Supplementary File 1, Table S4). Both P. falciparum and P. vivax were detected in the Funestus, Leucosphyrus and Maculatus Groups, whereas only P. vivax was detected in the Barbirostris Group. One specimen of the Funestus Group was co-infected with both species.

### Table 2. Molecular identification of sibling species among the Funestus, Maculatus and Leucosphyrus Groups.

<table>
<thead>
<tr>
<th>Group</th>
<th>Collection method (% of collected specimen analyzed by PCR)</th>
<th>Species</th>
<th>n/N</th>
<th>Relative proportion estimate</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Funestus</td>
<td>HLC (3294/42283=8%)</td>
<td>An. minimus (s.s.)</td>
<td>3277/3294</td>
<td>99.5</td>
<td>99.2-99.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>An. aconitus (s.s.)</td>
<td>7/3294</td>
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<td>0.1-0.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>An. culicifacies B</td>
<td>6/3294</td>
<td>0.2</td>
<td>0.1-0.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>An. harrisoni</td>
<td>2/3294</td>
<td>0.1</td>
<td>0-0.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>An. varuna</td>
<td>2/3294</td>
<td>0.1</td>
<td>0-0.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>An. pampanai</td>
<td>0/3294</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>CBT (1543/15728=10%)</td>
<td>An. minimus (s.s.)</td>
<td>1342/1543</td>
<td>87</td>
<td>85.2-88.6</td>
</tr>
<tr>
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<td></td>
<td>An. culicifacies B</td>
<td>90/1543</td>
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<td>4.7-7.1</td>
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<td>0.3-1.1</td>
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<td>0-0.4</td>
</tr>
<tr>
<td></td>
<td>CBT (1491/5239=28%)</td>
<td>An. minimus (s.s.)</td>
<td>975/1491</td>
<td>65.4</td>
<td>62.9-67.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>An. maculatus (s.s.)</td>
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<td>27.1-31.8</td>
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<td></td>
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<td>An. pseudowillmori</td>
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<td>Leucosphyrus</td>
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<td>HLC (856/1144=77%)</td>
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<td>CBT (29/46=57%)</td>
<td>An. dirus (s.s.)</td>
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<td>An. baimaii</td>
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</tr>
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<td></td>
<td></td>
<td>An. cracens</td>
<td>5/29</td>
<td>17.2</td>
<td>5.8-35.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>An. balabacensis</td>
<td>0/29</td>
<td>0</td>
<td>-</td>
</tr>
</tbody>
</table>
with both *P. falciparum* and *P. vivax*. The Funestus Group was by far the most important taxa contributing to malaria transmission ( Pf-EIR= 0.1 and Pv-EIR= 0.6 infective bites/person/month) followed by the Maculatus, Leucosphyrus and Barbirostris Groups (Table 3). Due to the relatively low sample size analyzed in the Annularis and Subpictus Groups (747 and 126 respectively), it was not possible to rule out their contribution to malaria transmission.

**Plasmodium**-infected mosquitoes were identified at the species level using molecular assays. Six formally named species were incriminated in malaria transmission: *An. minimus* (s.s.), *An. aconitus* (s.s.) (Funestus Group), *An. maculatus* (s.s.), *An. sawadwongporni* (Maculatus Group), *An. dirus* (s.s.) and *An. baimaii* (Leucosphyrus Group). *Plasmodium* sporozoites were detected in all species whereas *P. falciparum* sporozoites were detected only in *An. minimus* (s.s.), *An. maculatus* (s.s.), *An. sawadwongporni* and *An. dirus* (s.s.). Molecular identification at the species level was not possible for 6/106 positive samples because there was no DNA remaining.

Interestingly, the avian malaria *P. juxtanucleare* was detected in five specimens of the Funestus, Maculatus and Leucosphyrus Groups collected by HLC (two *An. minimus* (s.s.), one *An. baimaii* and two undetermined species). In addition, 16% (3308/21013) of the specimens from the Funestus, Maculatus and Leucosphyrus Groups collected in the CBTs were screened for *Plasmodium* sporozoites. *Plasmodium vivax* sporozoites were detected in two *An. minimus* (s.s.).

Quantitation of the sporozoite load was possible in 106/108 of the *P. falciparum* and *P. vivax* positive samples. Overall, 63% (67/106) of the infected specimens carried less than 100 sporozoites (Figure 3). The geometric mean was 41 (95%CI= [14; 98]) and 162 (95%CI= [94; 167]) sporozoites /infected mosquito for *P. falciparum* and *P. vivax* respectively (Supplementary File 1, Table S5). *Anopheles maculatus* (s.l.) appeared to be infected with lower parasite loads compared to other anopheline species. The range of sporozoite loads was 6 to 9234 sporozoites for *P. falciparum* and 4 to 517500 sporozoites for *P. vivax*. Moreover, 81/108 abdomens from sporozoite-positive samples were screened for *Plasmodium* oocysts (remaining abdomens were lost or moldy). *Plasmodium* oocysts were detected in only 57% (46/81) of the sporozoite-positive specimens. Thirty-two out of the 35 sporozoite-positive oocysts-negative specimens carried less than 500 sporozoites in their salivary glands, suggesting that these specimens were infected with a low number of oocysts.

**Blood-seeking behaviour of Anopheles mosquitoes**

Taking into account the whole dataset, the mean HBR of *Anopheles* mosquitoes was 460 bites/person/month (95%CI= [457; 463]) and the mean CBR was 4807 bites/cow/month (95%CI= [4770; 4843]). Mean HBR of primary malaria vectors varied from 2 to 306 bites/person/month in *An. dirus* (s.s.) and *An. minimus* (s.s.) respectively. Some primary malaria vectors were also frequently collected on CBT: *Anopheles minimus* (s.s.), *An. sawadwongporni* and *An. maculatus* (s.s.). had a mean CBR of 996, 249 and 112 bites/cow/month respectively (Supplementary File 1, Figure S11). The data on secondary vectors (e.g. Barbirostris and Annularis Groups) are more difficult to interpret because they probably represent a mix of several sibling species.

Leucosphyrus members were the most anthropophagic and endophagic species (mean CBI=0.16 and 0.44 and mean EI= 0.45 and 0.37 in *An. baimaii* and *An. dirus* (s.s.) respectively). Other malaria vectors in the Funestus, Maculatus and Barbirostris Groups were more zoophagic and exophagic (mean CBI=0.75-0.95 and mean EI=0.60-0.75). All remaining species were strongly zoophagic and exophagic (mean CBI= 0.83-1.00 and mean EI= 0.63-0.83) (Figure 4). Beyond zoophagy, malaria vectors appeared opportunistic in the choice of their blood meal source. Indeed, we have shown that *Anopheles* specimens collected by HLC can carry the avian malaria parasite *P. juxtanucleare*, and that *Anopheles* specimens collected on CBT can carry the human malaria parasite *P. vivax* (i.e. *Anopheles* mosquitoes collected on a given host type can carry *Plasmodium* sporozoites acquired from a previous blood meal taken on a different host type). These findings clearly demonstrate that malaria vectors can feed alternatively on different blood sources during their life span.

*Anopheles* mosquitoes exhibited an outdoor and/or early biting pattern (Supplementary File 1, Figures S12 and S13). Some species were already active at 06:00 pm and/or at 06:00 am, suggesting that exposure to malaria vectors stretches out of the standard collection time. The proportion of malaria vectors (both *Plasmodium*-infected and uninfected specimens) collected indoors between 09:00 pm and 05:00 am was 29% (range= 15–48% according to the species). Conversely, 64% of the *Plasmodium*-infected specimens were collected either out of doors, or indoors before 09:00 pm and after 05:00 am (Figure 5).

**Entomological indices of malaria transmission**

Only the Funestus, Maculatus and Leucosphyrus Groups were taken into account for the analysis of the entomological indices. Overall, mean HBR was 369 bites/person/month (95%CI= [366; 372]) and compensated for the low infection rate in these naturally infected populations of malaria vectors. Mean Pf-SI was 0.04% (95%CI= [0.02; 0.06]) and mean Pv-SI was 0.17% (95%CI= [0.14; 0.21]), yielding mean Pf-EIR of 0.13 (95%CI= [0.08; 0.21]) and mean Pv-EIR of 0.64 (95%CI= [0.51; 0.79]) infective bites/person/month (Table 4). The transmission of *P. falciparum* was highly seasonal: the rainy season was associated with a 10-fold increase in Pf-EIR. In contrast, mean Pf-EIR was 0.52 (95%CI= [0.34; 0.75]) and 0.73 (95%CI= [0.55; 0.94]) infective bites/person/month during the rainy season and the dry season respectively (Table 5). Only 6% (1/18) of the mosquitoes infected with *P. falciparum* sporozoites were detected during the dry season whereas 32% (28/87) of the mosquitoes infected with *P. vivax* sporozoites were detected during the dry season (two-sided Fisher’s exact test, p-value= 0.0218).

Average values of entomological indices concealed a high heterogeneity. When data were aggregated at the village and survey levels, mean HBR ranged from 13 to 2611 bites/person/month, mean Pf-EIR ranged from 0.00 to 3.05 infective bites/person/month and mean Pv-EIR ranged from 0.00 to 9.75 infective
Table 3. Entomological indices of malaria transmission presented per Group of *Anopheles*.

<table>
<thead>
<tr>
<th>Group</th>
<th>HBR (b/p/m)</th>
<th>Pf-SI (%)</th>
<th>Pf-EIR (ib/p/m)</th>
<th>Pv-SI (%)</th>
<th>Pv-EIR (ib/p/m)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>Estimate</td>
<td>95%CI</td>
<td>n/N</td>
<td>Estimate 95%CI</td>
</tr>
<tr>
<td>Funestus</td>
<td>42283/4120</td>
<td>307.9</td>
<td>305-310.8</td>
<td>13/41797</td>
<td>0.03   0.02-0.05</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>77/41797 0.18 0.15-0.05</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>13/4073 0.1 0.05-0.16</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>77/4073 0.57 0.45-0.71</td>
</tr>
<tr>
<td>Maculatus</td>
<td>7281/4120</td>
<td>53</td>
<td>51.8-54.2</td>
<td>4/7178</td>
<td>0.06   0.02-0.14</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>7/7178 0.1 0.04-0.14</td>
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<td></td>
<td>4/4062 0.03 0.01-0.08</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>7/4062 0.05 0.02-0.11</td>
</tr>
<tr>
<td>Leucosphyrus</td>
<td>1144/4120</td>
<td>8.3</td>
<td>7.9-8.8</td>
<td>1/1107</td>
<td>0.09   0-0.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3/1107 0.27 0.06-0.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1/3987 0.01 0-0.04</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3/3987 0.02 0-0.07</td>
</tr>
<tr>
<td>Barbirostris</td>
<td>6098/4120</td>
<td>44.4</td>
<td>43.3-45.5</td>
<td>0/5917</td>
<td>0       0-0.06</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2/5917 0.03 0-0.06</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0/3998 0 0-0.03</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2/3998 0.02 0-0.05</td>
</tr>
<tr>
<td>Annularis</td>
<td>772/4120</td>
<td>5.6</td>
<td>5.2-6</td>
<td>0/747</td>
<td>0       0-0.49</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0/747 0 0-0.49</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0/3987 0 0-0.03</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0/3987 0 0-0.03</td>
</tr>
<tr>
<td>Subpictus</td>
<td>144/4120</td>
<td>1</td>
<td>0.9-1.2</td>
<td>0/126</td>
<td>0       0-2.89</td>
</tr>
<tr>
<td></td>
<td></td>
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<td></td>
<td></td>
<td>0/126 0 0-2.89</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0/3605 0 0-0.03</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0/3605 0 0-0.03</td>
</tr>
</tbody>
</table>

*b/p/m*: bites/person/month; *CI*: confidence interval; *HBR*: human-biting rate; *ib/p/m*: infective bites/person/month; *n/N*: value of the numerator and denominator in the calculation of the corresponding indice as per the definition given in the Methods section; *Pf-EIR*: *P. falciparum* entomological inoculation rate; *Pv-EIR*: *P. vivax* entomological inoculation rate; *Pf-SI*: *P. falciparum* sporozoite index; *Pv-SI*: *P. vivax* sporozoite index.
Figure 3. Frequency distribution of the sporozoite load in naturally infected malaria vectors.

Figure 4. Exophagy index (EI) and zoophagy index (CBI) of Anopheles mosquitoes. A) EI and CBI are presented at the Group level. B) EI and CBI are presented at the species level. Vertical red lines show the mean value of the indice for the Anopheles spp. mosquitoes.

Figure 5. Distribution of infective bites according to the collection time and location. : proportion of the Plasmodium-infected mosquitoes collected indoors between 09:00 pm and 05:00 am.

The lowest HBR measured on a single collector and during a single night of collection was 0 bites/night and the highest was 289 bites/night. When taking into account the entire follow-up, mean HBR measured on single collectors ranged from 66 to 1253 bites/person/month, mean Pi-EIR ranged from 0 to 0.86 infective bites/person/month and mean Pv-EIR ranged from 0 to 4.92 infective bites/person/month (100–105 collection nights/mosquito collector). The cumulative HBR and EIR measured in the cohort of mosquito collectors followed a logarithmic distribution: 20% of the collectors received 50% of the bites and 20% of the collectors received 50% of the infective bites. In contrast, 30% of the collectors did not receive any infective bites during the study. Interestingly, the cumulative HBR followed a linear trend when paired to EIR, suggesting that the heterogeneity in
### Table 4. Entomological indices presented per study village.

<table>
<thead>
<tr>
<th>Village</th>
<th>HBR (bites/person/month)</th>
<th>Pf-SI (%)</th>
<th>Pf-EIR (infective bites/person/month)</th>
<th>Pv-SI (%)</th>
<th>Pv-EIR (infective bites/person/month)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N Estimate 95%CI</td>
<td>n/N Estimate 95%CI</td>
<td>n/N Estimate 95%CI</td>
<td>n/N Estimate 95%CI</td>
<td>n/N Estimate 95%CI</td>
</tr>
<tr>
<td>HKT</td>
<td>10736/1020 316 310-322</td>
<td>5/10625 0.05 0.02-0.11</td>
<td>39/10625 0.37 0.26-0.5</td>
<td>5/1009.5 0.15 0.05-0.35</td>
<td>39/1009.5 1.16 0.82-1.58</td>
</tr>
<tr>
<td>KNH</td>
<td>9536/1050 272 267-278</td>
<td>4/9414 0.04 0.01-0.11</td>
<td>27/9414 0.29 0.19-0.42</td>
<td>4/1036.6 0.12 0.03-0.3</td>
<td>27/1036.6 0.78 0.51-1.14</td>
</tr>
<tr>
<td>TOT</td>
<td>24293/1050 694 685-703</td>
<td>6/24083 0.02 0.01-0.05</td>
<td>15/24083 0.06 0.03-0.1</td>
<td>6/1040.9 0.17 0.06-0.38</td>
<td>15/1040.9 0.43 0.24-0.71</td>
</tr>
<tr>
<td>TPN</td>
<td>6143/1000 184 180-189</td>
<td>3/5960 0.05 0.01-0.15</td>
<td>6/5960 0.1 0.04-0.22</td>
<td>3/970.2 0.09 0.02-0.27</td>
<td>6/970.2 0.19 0.07-0.4</td>
</tr>
<tr>
<td>All villages</td>
<td>50708/4120 369 366-372</td>
<td>18/50082 0.04 0.02-0.06</td>
<td>87/50082 0.17 0.14-0.21</td>
<td>18/4069.1 0.13 0.08-0.21</td>
<td>87/4069.1 0.64 0.51-0.79</td>
</tr>
</tbody>
</table>

* b/p/m: bites/person/month; CI: confidence interval; HBR: human-biting rate; ib/p/m: infective bites/person/month; n/N: value of the numerator and denominator in the calculation of the corresponding indice as per the definition given in the Methods section; Pf-EIR: P. falciparum entomological inoculation rate; Pv-EIR: P. vivax entomological inoculation rate; Pf-SI: P. falciparum sporozoite index; Pv-SI: P. vivax sporozoite index.

### Table 5. Entomological indices presented per season.

<table>
<thead>
<tr>
<th>Season</th>
<th>HBR (bites/person/month)</th>
<th>Pf-SI (%)</th>
<th>Pf-EIR (infective bites/person/month)</th>
<th>Pv-SI (%)</th>
<th>Pv-EIR (infective bites/person/month)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N Estimate 95%CI</td>
<td>n/N Estimate 95%CI</td>
<td>n/N Estimate 95%CI</td>
<td>n/N Estimate 95%CI</td>
<td>n/N Estimate 95%CI</td>
</tr>
<tr>
<td>Dry</td>
<td>39259/2470 477 472-482</td>
<td>17/38777 0.04 0.03-0.07</td>
<td>59/38777 0.15 0.12-0.2</td>
<td>17/2439.7 0.21 0.12-0.33</td>
<td>59/2439.7 0.73 0.55-0.94</td>
</tr>
<tr>
<td>Rainy</td>
<td>11449/1650 208 204-212</td>
<td>1/11305 0.01 0-0.05</td>
<td>28/11305 0.25 0.16-0.36</td>
<td>1/1629.2 0.02 0-0.1</td>
<td>28/1629.2 0.52 0.34-0.75</td>
</tr>
<tr>
<td>All seasons</td>
<td>50708/4120 369 366-372</td>
<td>18/50082 0.04 0.02-0.06</td>
<td>87/50082 0.17 0.14-0.21</td>
<td>18/4069.1 0.13 0.08-0.21</td>
<td>87/4069.1 0.64 0.51-0.79</td>
</tr>
</tbody>
</table>

* December to May
* June to December

* b/p/m: bites/person/month; CI: confidence interval; HBR: human-biting rate; ib/p/m: infective bites/person/month; n/N: value of the numerator and denominator in the calculation of the corresponding indice as per the definition given in the Methods section; Pf-EIR: P. falciparum entomological inoculation rate; Pv-EIR: P. vivax entomological inoculation rate; Pf-SI: P. falciparum sporozoite index; Pv-SI: P. vivax sporozoite index.
the distribution of infective bites was not explained by the mean of exposure to malaria vectors (Figure 6).

Discussion

This study was a unique opportunity to document some entomological aspects of malaria transmission in low transmission settings of Southeast Asia. Our data are important in the context of malaria elimination locally in Kayin state, but also elsewhere in the Greater Mekong Subregion where malaria displays a similar transmission pattern.

The dynamics of entomological indices in an area of low, seasonal and unstable *P. falciparum* transmission setting

Our results confirm previous observations that primary malaria vectors in this area are *An. minimus* (s.s.), *An. maculatus* (s.s.), *An. sawadwongporni*, *An. dirus* (s.s.) and *An. baimaii*, and that several other species also play a secondary role in the transmission. Infection rate is low in naturally infected populations of malaria vectors and balanced by the high biting-rate of malaria vectors\(^1,1^2,1^5\), yielding mean entomological inoculation rate of 1.6 and 7.7 infective bites/person/year for *P. falciparum* and *P. vivax* respectively. These EIR estimates were made in the context of *falciparum* malaria elimination (community wide access to early diagnosis and treatment, and mass drug administration). Therefore, baseline intensity of *falciparum* malaria transmission in hotspot villages in Kayin state is likely to be higher than that reported in this study\(^1^5\). In contrast, *Plasmodium vivax* EIR estimates were higher than previously reported in the same area\(^1^1\). This implies that new infections (primo-infections and re-infections) are more common than previously thought and probably frequently asymptomatic\(^3^2\).

The transmission of *P. falciparum* was seasonal whereas infective bites of *P. vivax* occurred during both the dry and rainy seasons. The seasonality in *P. falciparum* transmission was only partially explained by the increase in malaria vector abundance during the

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**Figure 6.** Heterogeneous distribution of bites and infective bites among the cohort of mosquito collectors over the entire period of the study. **A** Cumulative distribution of the bites of malaria vectors among the cohort of mosquito collectors. **B** Paired cumulative distribution of bites of malaria vectors and infective bites among the cohort of mosquito collectors.
rainy season when compared to the dry season. The longevity of malaria vectors was most likely to be the main factor driving the seasonality of *P. falciparum* transmission. During the dry season, the life expectancy of malaria vectors was probably too short for *P. falciparum* to complete its sporogonic cycle in the mosquito. During the rainy season, the longevity of malaria vectors may have increased and malaria vectors lived long enough for *P. falciparum* sporozoites to appear in the salivary glands ([3],[4]). For *P. vivax*, the duration of the sporogonic cycle was compatible with sporozoite detection throughout the year as this parasite develops faster than any other species in its mosquito vectors ([5],[6]). Relapsing vivax malaria parasites, vector competency and temperature are also potential factors that explain seasonal trends in malaria transmission.

Although 5% of the mosquitoes infected with *Plasmodium* sporozoites had >10,000 sporozoites in their salivary glands, the sporozoite load measured in these naturally infected malaria vectors was in average very low (60% of the infected specimens carried less than 100 sporozoites). This is consistent with previous attempts to quantify *P. falciparum* and *P. vivax* sporozoites in low transmission settings ([7],[8]) and contrasts with the high sporozoite loads detected in Africa ([9],[10]). Considering a geometric mean of 162 sporozoites / *P. vivax* infected mosquito, a mean *P. vivax* EIR of 7.7 infective bites / person / year and that infected mosquitoes would transmit all their sporozoites, we estimated that each person would receive 1200 *P. vivax* sporozoites per year in average. Taking into account that not all sporozoites may be inoculated and that not all sporozoites may transform into hypnozoites, the reservoir of hypnozoite is likely to be small in most individuals exposed to the disease in this transmission setting.

The distribution of infective bites among the human population was highly heterogeneous. This pattern was not explained by the mean of exposure to malaria vectors as paired cumulative HBR and EIR did not follow the same trend. The study villages were hotspots of malaria transmission defined by the high prevalence of asymptomatic infection ([11],[12]). This implies a substantial degree of premunition in asymptomatic carriers, i.e. the development of a protective immunity that maintains parasitaemia at sub-patent levels. The broad spectrum of EIRs measured in hotspot villages may explain why some people develop such a protective immunity and manage to control the infection.

**Residual malaria transmission**

The two broadly scalable vector-control interventions recommended by the World Health Organization for the control of malaria vectors are mass distribution of LLINs or, where appropriate, IRS campaigns ([13]). By definition, LLINs target malaria vectors seeking human blood, indoors and at a time when people are sleeping under mosquito nets. In order IRS to be effective, malaria vectors targeted by the intervention must also rest indoors, before or after a blood meal. However, this stereotypical blood seeking behavior applies only to a minority of the dominant malaria vectors worldwide ([14]). Several behavioral traits drive the refractoriness of malaria vectors to LLINs and IRS including (i) their ability to take blood meals from animals (zoophagy and opportunistic blood type selection), (ii) their tendency to rest and/or feed outdoors (exophily and exophagy) and (iii) their ability to feed before dawn and after dusk, at a time when people are not protected by LLINs or IRS intervention ([15]).

As previously reported, LLINs only have a limited efficacy in preventing human-vector contact and disease transmission in the Thailand-Myanmar border area. Somboon et al. evaluated the impact of mosquito bed nets impregnated with lambda-cyhalothrin using entomological endpoints in very similar transmission settings (Karen villages located on the Thai side of the border) ([16]). The authors reported that mosquito bed nets can prevent 36–78% of the human-vector contact according to the *Anopheles* species. Universal coverage with LLINs failed to reduce the abundance and longevity of malaria vectors, suggesting that this intervention had only a limited impact on the vectorial capacity. The impact of permethrin-impregnated mosquito bed nets was also evaluated in pregnant women and children living in refugee camps using epidemiological endpoints. The use of mosquito bed nets during pregnancy was associated with a significant reduction in the incidence of severe anaemia but not of malaria ([17]). At a time when EIR was higher, the use of mosquito bed nets in children was associated with a significant decrease of *falciparum* malaria incidence but no effect was observed on *P. vivax* ([18]). More recently, Smithuis et al. failed to observe an impact of LLINs among a cohort of 175 children followed for 10 months in Western Myanmar ([19]). This negative result was explained by the early and outdoor biting pattern of malaria vectors ([20]).

In this study, only 36% of the *Plasmodium*-infected mosquitoes were collected indoors between 09:00 pm and 05:00 am (when and where people are expected to sleep under a bed-net). This proportion might have been overestimated as malaria vectors were already active at 06:00 pm and/or at 06:00 am, suggesting that the exposure stretched out of the collection time. Accurate quantitation of the part of the transmission that LLINs fail to prevent would require collecting additional data on population movements and sleeping habits of people living in this area. Moreover, we have clearly demonstrated an opportunistic blood type selection in some vectors, i.e. that a given specimen is able to feed on several blood sources during successive gonotrophic cycles. This opportunism also appears as an important factor to explain why universal coverage with mosquito bed nets fails to affect the dynamic of anopheine populations and decrease vectorial capacity in the area ([21]). Consequently, the paradigm of residual transmission as experienced in high transmission settings of Africa does not apply to the Thailand-Myanmar border area and a drastic shift in vector-control interventions is required.

**Shift in vector-control intervention**

The design of effective intervention for the control of malaria vectors in Southeast Asia should take into account malaria vector ecology and transmission dynamics. In this study, we have shown that multiple vectors have different and complementary blood-seeking behaviours, making their control particularly difficult.

Veterinary approaches such as the injection of livestock with a slow-release formulation of endectocides ([22]), or the use of insecticide-treated mosquito nets fenced around cattle ([23]) may be an interesting strategy to decrease the vectorial capacity of some
zoophilic and/or zoophagic malaria vectors (ex: An. minimus, An. maculatus and An. sawadwongporn). We have shown that malaria vectors can readily feed on a wide variety of blood types including human, cattle, pigs and birds. However, the diversity of blood sources and the relative proportion of blood meals taken on a given source remain to determine. In this regard, targeted sequencing of mammalian mt 16S ribosomal RNA genes detected in DNA extracts from blood-fed specimens is a promising tool for the determination of blood-meal sources in wild populations of malaria vectors.

The nature of Anopheles resting habitats is another important aspect of malaria vector ecology that can be targeted by residual insecticide spraying intervention. Resting habitats have been identified both indoors (ex: roof, wall, ceilings of houses and animal barns) and outdoors (ex: tree holes, rodent holes, dense bushes). However, most mosquito species rest exclusively out of doors in natural settings, and only a relatively few species rest in man-made shelters. The size and importance of the exophilic population that commonly rest inside houses are typically overlooked because the sampling of outdoor-resting population is more difficult than the sampling of indoor-resting population. In Southeast Asia, most of the life cycle of Anopheles mosquitoes is likely to take place out of doors and malaria vectors are rarely found resting indoors. Therefore, the scope of residual insecticide spraying for the control of malaria vectors may be extended to outdoor applications.

Insecticide resistance may also represent an additional threat to the control of malaria vectors in this area. We have previously detected relatively low levels of resistances to deltamethrin and permethrin in populations of An. minimus (s.l.) and An. maculatus (s.l.) collected in the villages of the present study. Further investigations are needed in order to document the extent of pyrethroid resistance elsewhere in Kayin state and to evaluate the potential effectiveness of alternative class of insecticides such as carbamate (ex. bendiocarb), organophosphate (ex. malathion) or insect growth inhibitors (ex. pyriproxyfen).

Conclusion
This study highlights the importance of entomology in the context of malaria elimination in Kayin state. A drastic shift in vector-control strategy is required in order to address early and outdoor malaria transmission, and the modalities of vector-control should be retuned to address problematic specific to malaria elimination.

Data availability
The data is available upon request to the Mahidol Oxford Tropical Medicine Research Unit Data Access Committee (Supplementary File 3; http://www.tropmedres.ac/data-sharing) and following the Mahidol Oxford Tropical Medicine Research Unit data access policy (http://www.tropmedres.ac/_asset/file/data-sharing-policy-v1-0.pdf).

Grant information
This work was supported by the Wellcome Trust [101148]; and the Bill and Melinda Gates Foundation [GH OPP 1081420]. Victor Chaumeau received a PhD scholarship by the Centre Hospitalier Universitaire de Montpellier (CHU Montpellier).

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Acknowledgments
We thank all study participants and communities from Htoo Pyin Nyar, Tar Au Ta, Ka Nu Hta and Htee Kaw Taw villages for their participation and support to the study. SMRU is part of the Mahidol Oxford University Research Unit, supported by the Wellcome Trust of Great Britain.

Supplementary material
Supplementary File 1. Supplemental data.
Click here to access the data

Supplementary File 2. Laboratory procedures for the processing of entomological samples.
Click here to access the data

Supplementary File 3. Request form for Mahidol Oxford Tropical Medicine Research Unit Data Access Committee.
Click here to access the data
References


43. https://www.who.int/malaria/areas/vector_control/core_methods/en/


Catherine Bourgouin

I approve the current version after fixing three minor points:

- Page 8: "The proportion of malaria vectors collected indoors between 09:00 pm and 05:00 am was 29% (range= 15–48% according to the species). Conversely, 64% of the infected specimens were collected out of doors, before 09:00 pm and/or after 05:00 am (Figure 5)". Based on the data in Figure 5, 29% should be 36%; the last sentence should be rephrased as for example: "Conversely, 64% of the infected specimens were collected either out of doors, or before 09:00 pm and after 05:00 am."

- Page 13: "In this study, only 36% of the Plasmodium-infected mosquitoes were collected indoors between 09:00 pm and 05:00 am (when and where people are expected to sleep under a bed-net). This proportion might have been underestimated as malaria vectors were already active at 06:00 pm and/or at 06:00 am, suggesting that the exposure stretched out of the collection time". Underestimated, should actually be "overestimated".

- Page 14, end of the 2nd paragraph: "In this regard, targeted sequencing of 16S ribosomal RNA genes detected in DNA extracts from blood-fed specimens is a promising tool for the determination of blood-meal sources in wild populations of malaria vectors". Change "16S ribosomal RNA genes" to "mammalian mt16S ribosomal RNA genes", as "16S ribosomal RNA genes" alone can be confusing as related to bacterial genomes.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Molecular entomologist, malaria transmission.

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.
Page 8: "The proportion of malaria vectors collected indoors between 09:00 pm and 05:00 am was 29% (range= 15–48% according to the species). Conversely, 64% of the infected specimens were collected out of doors, before 09:00 pm and/or after 05:00 am (Figure 5)". Based on the data in Figure 5, 29% should be 36%; the last sentence should be rephrased as for instance: "Conversely, 64% of the infected specimens were collected either out of doors, or before 09:00 pm and after 05:00 am.

Page 8, 29% refers to the proportion of malaria vectors collected indoors between 09:00 pm and 05:00 am independently of their infection status, whereas 64% refers only to the proportion of *Plasmodium*-infected specimen collected outdoors, or indoors early in the evening or early in the morning. Therefore, we do not wish to change the number cited in the main text. The sentence has been reworded in the revised manuscript as per reviewer suggestion.

Page 13: "In this study, only 36% of the *Plasmodium*-infected mosquitoes were collected indoors between 09:00 pm and 05:00 am (when and where people are expected to sleep under a bed-net). This proportion might have been underestimated as malaria vectors were already active at 06:00 pm and/or at 06:00 am, suggesting that the exposure stretched out of the collection time". Underestimated, should actually be "overestimated".

In the revised manuscript, typo page 13 "underestimated" was changed to "overestimated" as per reviewer's suggestion.

Page 14, end of the 2nd paragraph: "In this regard, targeted sequencing of 16S ribosomal RNA genes detected in DNA extracts from blood-fed specimens is a promising tool for the determination of blood-meal sources in wild populations of malaria vectors". Change "16S ribosomal RNA genes" to "mammalian mt16S ribosomal RNA genes", as "16S ribosomal RNA genes" alone can be confusing as related to bacterial genomes.

Page 14 “16S ribosomal RNA genes” was changed to “mammalian mt16S ribosomal RNA genes” as per reviewer’s suggestion.

**Competing Interests:** I have no competing interest to disclose.
Page 10, paragraph starting with “average values”. I do not follow the reasoning from “The lowest HBR measured on a single collector...” until the end of this paragraph.

What exactly do you mean by “mean HBR measured on single collectors”? Usually when using HCL, more than one collector is involved per class (indoor versus outdoor) located in different houses to avoid bias linked to the person or the surrounding of the house. We therefore combined data from several collectors within a village over a define time scale, as here month.

It may be just a matter of presenting, arguing the data presented in this paragraph. I feel it may need some clarification.

The discussion section:

I feel this could be largely improved by including conclusion of the different mosquito species biting humans and which one appears the most relevant local vector, if any, based on Plasmodium carriage. This is clearly presented in the sup figures (S7-S10).

Similarly, the huge work presented in sup data S11-S12, could argue toward “shift intervention control” and “residual malaria transmission”. Except in the second § on page 14, was a mosquito species ever mentioned.

On page 13, the 1st § discussed more the longevity of the mosquitoes for being unable to transmit Pf, than the abundance component. Is the shorter life cycle of P. vivax a dogma that could explain its persistence in mosquitoes, rather than the presence of the best competent mosquito species the year around? And relapse in people? Interesting old data, but more recent that the Boyd book, on P. vivax and mosquitoes could be found as for instance “Am. J. Trop. Med. Hyg., 37(2), 1987, pp. 241 -245”¹. In addition, duration of Plasmodium sporogonic development is highly dependent on the T° whether the parasite species is falciparum or vivax.

Additionally, discussing whether the MDA had some impact or not on parasite prevalence in mosquitoes could be interesting, if the sample sizes are enough informative.

Overall, I found some parts of the discussion too long and to some extent not linked enough to the data, ie the § on residual malaria.

Additional important comments:

- Not noticed in my former review, the ITS2 (laboratory procedure & sup data) is not a mitochondrial marker, but a rDNA marker.
- Tables 3 to 6: The “N” does not correspond to the same class between HBR, SI and EIR, please add proper legend.
- Page 8, last §: just after or in the title, it would be wise to add the analysis is done at the village scale. You could also make a brief comment on whether there is or not significant differences for the Pf/Pf EIR between villages.

Additional minor comments:

- When using “falciparum malaria”; it is a standard to not use italics “falciparum”, same for vivax malaria
- Page 3 : The last 2 sentences of the last “intro” § are somehow redundant.
- Page 3 “study villages”: could you give the range for how high is the submicroscopic malaria prevalence?
- Figure 1 legend: consider using plural for “village, house and site”
- M&M:
  - qrtPCR: for molecular biologist it should be only qPCR, as qrtPCR might be confusing with Reverse Transcription and quantitative PCR (RT-qPCR)
  - to avoid going to the publication you cited, it would be simple to add “Mangold, targeting the Plasmodium 18S” and Cunha, Plasmodium mitochondrial targets, as indeed mentioned in the sup file 2.
In the tables and in some additional places in the text: should not that be conventional to write numbers as 42,283 instead of 42283, as an example?

In different places, I think that, as an example,: “malaria vectorS ecology” should be “malaria vector ecology”.

Few typos: page 3 last 2 sentences, 2nd §: indoorresidual &theecology, Page 4 theShoklo, “internal transcribED spacer”.

Page 12 first §: “20% of the collectors received approximatively 50% of the bites and of the infective bites” => “20% of the collectors received approximatively 50% of the bites AS WELL AS 50% of the infective bites”. Correct?

Sup file 2: description of the PCR procedures for species identification: As the basic PCR is similar in terms of primer concentration, DNA template and final volume, I suggest to simplify and used a single table for the primers used adding a column with the specific temperature for primer annealing.

References

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Molecular entomologist, malaria transmission

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 26 Apr 2019

Victor Chaumeau, Centre hospitalier universitaire de Montpellier, Montpellier, France

We thank to the reviewer for her additional comments on the manuscript. Response to point specific comments are given below:

- Page 10, paragraph starting with “average values”. I do not follow the reasoning from “The lowest HBR measured on a single collector...” until the end of this paragraph. What exactly do you mean by “mean HBR measured on single collectors” ? Usually when using HCL, more than one collector is involved per class (indoor versus outdoor) located in different houses to avoid bias linked to the person or the surrounding of the house. We therefore combined data from several collectors within a village over a define time scale, as here month. It may be just a matter of presenting, arguing the data presented in this paragraph. I feel it may need some clarification.

The value of the entomological indices are longitudinal and spatially hierarchized. Therefore it is possible to describe the data at different level of aggregation ranging from hourly measurements on single individuals (mosquito collectors) to average values obtained by pooling the entire dataset. Mean values collated at the village and survey level are indeed useful to describe the data independently of bias linked to the person, the surrounding of the house or the night of collection, were reported in the first lines of the paragraph. Further insights into the characteristics of the entomological indices were be obtained by looking at different level of aggregation. For example, the range of HBR collated at the village and survey level was 13 to 2611 bites/person/month.
whereas the range of non-collated HBR estimates (number of mosquito collected during one night by a single mosquito collector) was 0 to 289 bites/person/night (8670 bites/person/month), further documenting the high abundance of malaria mosquitoes sometime observed in the area of the study. By looking at data collated at the collector level for the entire period of the study, we also observed that some mosquito collectors never captured Plasmodium-infected mosquitoes whereas 20% of the collectors received 50% of the infective bites, which would not have been shown by analyzing only aggregated data.

- **The discussion section:** I feel this could be largely improved by including conclusion of the different mosquito species biting humans and which one appears the most relevant local vector, if any, based on Plasmodium carriage. This is clearly presented in the sup figures (S7-S10).

We believe that the dynamic of malaria mosquitoes and their capacity to transmit malaria is extremely complex in the area of the study and that designating one species as the main malaria vector would be misleading. One characteristics of the study area is the high diversity of malaria vectors, which pullulate in different environments and at different time of the year. Although most of the Plasmodium-infected malaria mosquitoes were identified as An. minimus s.s. in the present study, we do not think that this species is de facto the main local malaria vectors. For example, An. dirus is an important vector in and around the forest and its contribution may have been underestimated in the present study.

- **Similarly, the huge work presented in sup data S11-S12, could argue toward “shift intervention control” and “residual malaria transmission”.

The work presented in supplemental data Figures S10-11 is discussed in the paragraph “Residual malaria transmission” and “shift in vector control intervention”. We do not wish to add additional details in the discussion of the revised manuscript.

- **Except in the second § on page 14, was a mosquito species ever mentioned.**

Names of anopheles species were added at the very beginning of the discussion in the paragraph on the dynamics of entomological indices.

- **On page 13, the 1st § discussed more the longevity of the mosquitoes for being unable to transmit Pf, than the abundance component. Is the shorter life cycle of P. vivax a dogma that could explain its persistence in mosquitoes, rather than the presence of the best competent mosquito species the year around? And relapse in people? Interesting old data, but more recent that the Boyd book, on P.vivax and mosquitoes could be found as for instance “Am. J. Trop. Med. Hyg., 37(2), 1987, pp. 241 -245”1. In addition, duration of Plasmodium sporogonic development is highly dependent on the T° whether the parasite species is falciparum or vivax.

The reviewer is right to mention that not only the longevity of malaria vectors can explain seasonal trends in malaria transmission. A sentence has been added at the end of the paragraph and the suggested bibliography was cited in the main text.

- **Additionally, discussing whether the MDA had some impact or not on parasite prevalence in mosquitoes could be interesting, if the sample sizes are enough informative.**

The impact of MDA on the entomological indices has been published separately in Chaumeau et al., 2019 in the JID and we do not wish to add more details in the current manuscript.

- **Overall, I found some parts of the discussion too long and to some extent not linked enough to the data, ie the § on residual malaria.**

We believe that the paragraph on residual malaria transmission is important given the objective of the study. Direct link to the data are made in the paragraph on residual malaria transmission (eg.
night biting pattern and corresponding distribution of infective bites indoors and outdoors over the course of the night, zoophagy, opportunistic blood-source selection). We wish to keep the paragraph on residual malaria transmission unchanged.

- Not noticed in my former review, the ITS2 (laboratory procedure & sup data) is not a mitochondrial marker, but a rDNA marker.
  
  This mistake was corrected in the revised version of the manuscript.

- Tables 3 to 6: The “N” does not correspond to the same class between HBR, SI and EIR, please add proper legend.
  
  Proper legend has been added in the Tables presenting entomological indices.

- Page 8, last §: just after or in the title, it would be wise to add the analysis is done at the village scale.
  
  The data on the entomological indices are described at different level of aggregation (eg. average values on the entire data set, mean collated by village and by survey, non-collated estimates measured on individual mosquito collectors). We do not wish to change the title of this paragraph.

- You could also make a brief comment on whether there is or not significant differences for the Pf/Pf EIR between villages.
  
  A detailed analysis of Pf- and Pv-EIRs in the context of malaria elimination has been published separately in Chaumeau et al. JID 2019. We do not wish to add more details in the current manuscript.

- When using “falciparum malaria”; it is a standard to not use italics “falciparum”, same for vivax malaria
  
  “P. falciparum malaria” p.13 was changed to falciparum malaria as per reviewer’s suggestion.

- Page 3: The last 2 sentences of the last “intro” § are somehow redundant.
  
  One sentence was removed as per reviewer’s suggestion.

- Page 3 “study villages”: could you give the range for how high is the submicroscopic malaria prevalence?
  
  The range of submicroscopic malaria prevalence at the beginning of the study was cited in the main text. Additional information can be found in Landier et al. WOR 2017.

- Figure 1 legend: consider using plural for “village, house and site”
  
  Figure 1 legend has been modified as per reviewer’s suggestion.

- qrtPCR: for molecular biologist it should be only qPCR, as qrtPCR might be confusing with Reverse Transcription and quantitative PCR (RT-qPCR)
  
  qrtPCR was changed to qPCR in revised version of the manuscript.

- to avoid going to the publication you cited, it would be simple to add “Mangold, targeting the Plasmodium 18S” and Cunha, Plasmodium mitochondrial targets, as indeed mentioned in the sup file 2.
  
  Names of PCR DNA targets were added in the main text as per reviewer’s suggestion.

- In the tables and in some additional places in the text: should not that be conventional to write numbers as 42,283 instead of 42283, as an example?
  
  We do not wish to change number format in the revised manuscript.

- In different places, I think that, as an example,: “malaria vectorS ecology” should be “malaria vector ecology”.
  
  Typos were corrected in the revised manuscript.

- Few typos: page 3 last 2 sentences, 2nd §: indoorresidual &theecology, Page 4 theShoklo, “internal transcribED spacer”.
  
  Typos were corrected in the revised manuscript.

- Page 12 first §: “20% of the collectors received approximatively 50% of the bites and of the infective bites" => "20% of the collectors received approximatively 50% of the bites AS WELL AS 50% of the infective bites”. Correct?
The sentence was modified to improve its readability by the reader.

- **Sup file 2: description of the PCR procedures for species identification:** As the basic PCR is similar in terms of primer concentration, DNA template and final volume, I suggest to simplify and used a single table for the primers used adding a column with the specific temperature for primer annealing.

We wish to keep the current outline of the supplementary file S2.

**Competing Interests:** Authors have no competing interest to disclose.

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**Reviewer Report 15 February 2019**

https://doi.org/10.21956/wellcomeopenres.16466.r34776

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Lisa J. Reimer
Department of Vector Biology, Liverpool School of Tropical Medicine, Liverpool, UK

No further comments.

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

**Reviewer Expertise:** vector biology, malaria and filariasis transmission dynamics

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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**Version 1**

**Reviewer Report 10 December 2018**

https://doi.org/10.21956/wellcomeopenres.16087.r34267

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Lisa J. Reimer
Department of Vector Biology, Liverpool School of Tropical Medicine, Liverpool, UK

This is a very comprehensive study which was completed to a high standard. The knowledge generated is essential to achieve malaria elimination in the sub-region. My two essential recommendations for revision include 1) a deeper description on vector ecology in this region and a summary of which control and
prevention activities are currently in place. This background can draw from the work that was described previously in reference 17. The end of the introduction states “Numerous aspects of malaria vectors ecology and biology have not been documented and the characteristics of the entomological indices are not known precisely” but the reader would appreciate a full description of the current gaps in our knowledge. 2) There is a significant amount of data that has been collected and analysed, but this could be organised in a way that is much more accessible. For example a table that summarizes key attributes per vector group (if it is not always available by species) such as mean and range monthly biting rates, perhaps median biting times, EI, ZI, infection prevalence.

Further specific comments are below:

1. This is described as a pilot study on targeted malaria elimination, however this is such a comprehensive entomological survey that it diminishes the significance of the work to call it a pilot.
2. Typo under entomological surveys: “mosquito sampling the using human landing catch”
3. It would be more intuitive to presented sporozoite prevalence rather than out of 1000 mosquitoes
4. Figure 4b and 5b – how is the HBR, EI and ZI calculated at the species level when a smaller proportion of samples were ID’d molecularly? Is it a proportion of those that were morphologically identified?
5. Figure 5 – what are the dashed lines for? It might be more useful to use the line to show a community mean index
6. Typo under “residual malaria transmission”: “35% of infection mosquitoes collected indoors between 9pm and 5pm”
7. Rephrase the final sentence “in order to address specific problematic in the context of malaria elimination”

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?
Partly

Are the conclusions drawn adequately supported by the results?
Yes

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

**Reviewer Expertise:** vector biology, malaria and filariasis transmission dynamics

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.
We are thankful to the reviewer for her useful feedback on the manuscript. Part of the introduction was rewritten in order to provide the reader with more background information on malaria vectors, transmission settings and on-going elimination efforts in the area of the study. A full description of the current gaps in our knowledge on vector ecology and biology would be difficult to present in the format of this research article. We instead referred the reader to the review of Sinka et al. that summarize key gaps for each dominant malaria vectors in the Asia-Pacific region. We tried to improve the presentation of the results by keeping only the key tables and figures in the main text and by adding more supplementary figures and tables in the supplementary data. Unfortunately, it was not possible to summarize in one table the statistics of all key indices, together with the raw numbers and confidence intervals. Answer to specific comments are below:

- This is described as a pilot study on targeted malaria elimination, however this is such a comprehensive entomological survey that it diminishes the significance of the work to call it a pilot.
  
  The wording pilot has been removed from the revised manuscript.

- Typo under entomological surveys: "mosquito sampling the using human landing catch"
  
  The typo has been corrected.

- It would be more intuitive to presented sporozoite prevalence rather than out of 1000 mosquito.
  
  Sporozoite prevalence was expressed as percentages rather than out of 1000 mosquitoes.

- Figure 4b and 5b – how is the HBR, EI and ZI calculated at the species level when a smaller proportion of samples were ID’d molecularly? Is it a proportion of those that were morphologically identified?
  
  Yes. Species-specific proportions were used to compute HBR, CBR, EI and CBI estimates at the species level if ≥30 samples in the corresponding Anopheles Group were genotyped for each value of the grouping variable. The grouping variables were the collection time (06:00 pm to 06:00 am), the collection method (HCL or CBT) or the location of the mosquito collector (indoors or outdoors). For example, the number of An. maculatus (s.l.) collected by HLC indoors and outdoors was 2240 and 5041 respectively. The proportion of An. maculatus (s.s.) in the indoor and outdoor populations was 0.34 (133/392) and 0.37 (404/1084) respectively. The predicted number of An. maculatus (s.s.) collected by HLC indoors and outdoors was 762 and 1865 respectively. The EI estimate for An. maculatus (s.s.) was therefore 0.71 (1865/2627). More detail were added in the methods section.

- Figure 5 – what are the dashed lines for? It might be more useful to use the line to show a community mean index
  
  Vertical lines have been modified to address reviewer’s suggestion. The solid red line represents the mean index for Anopheles spp. mosquitoes, corresponding CIs were too narrow to be shown.

- Typo under “residual malaria transmission”: “35% of infection mosquitoes collected indoors between 9pm and 5pm”
  
  The typo has been corrected.

- Rephase the final sentence “in order to address specific problematic in the context of malaria elimination”
  
  The final sentence has been changed to “the modalities of vector-control should be retuned to address problematic specific to malaria elimination”.

**Competing Interests:** No competing interests were disclosed.
The objective of the work presented by Chaumeau et al was aimed at further describing the entomological determinants of malaria transmission at the Myanmar border with Thailand, in order to guide the policy makers for malaria elimination in this region of low transmission.

The investigation involved a longitudinal survey in four villages, using mosquito collection by Human Landing Catch as well as Cow Landing Catch over 21 months. The manuscript reports on a tremendous amount of work for both collecting mosquitoes, mosquito identification and Plasmodium detection in those mosquitoes by PCR. The data presented provided sound results in line with the objective.

Nevertheless, several parts would benefit from clarification or simplification, while other information, for instance on Asian malaria vectors, would be valuable for readers who are not specialists in malaria transmission in Asia. See specific comments below.

Additionally, I was expecting some deeper analysis or comments on possible differences among the 4 villages investigated and whether the ecology of the villages and the mosquito larvae could be included in malaria control recommendation.

For the global analysis of the entomological indexes for malaria, I have found difficulties in following the calculation of HBR, SI and EIR, when related to the comparison of the 4 villages and the impact of the season (Table 4 and Table 5).

The overall discussion could be better focused 1) on the global data, 2) the specificity or not of the transmission in the 4 villages to finish with stronger arguments for recommendation than the current discussion that sounds rather dogmatic on the specificity of transmission in Asia by mostly exophagic mosquitoes….

Detailed comments:

**Title:** “Entomological determinants of malaria transmission in Kayin state, Eastern Myanmar: A 24-month longitudinal study in four villages”

The Methods mention a 21 months survey, that indeed might cover 2 years. As the survey in the 4 villages are not fully superimposed, it might be wise to describe the overlap for a better comprehensive value of the data.

The “Kayin” state is barely mentioned in the report, bringing confusion when done ie: appearing only on page 13 beside the title and the abstract with “Therefore, baseline intensity of malaria transmission in hotspot villages from the Kayin state is likely to be higher than that reported in this study”
Lastly, if there is no real discussion on the difference similarity among the 4 villages, why attracting with the “4 villages” in the title?

Abstract: Some revision could be done if the authors agree on the following comments:

Methods

Study sites: Providing the global description of the villages, as done in the J. Vector ecology Paper (ref14), would save time, rather than going to the published document as well as possibly useful for the discussion. Was the MDA still active during the whole entomological survey?

Entomological surveys: Were the house randomly chosen? Were the preliminary data (ref14) included in the analysis? Consider simplification: instead “6 sites” use 5 houses plus one cow collection. Verify the accuracy of the sentence for mosquito collection under the net covering the cow.

Data analysis: Expressing SI / 1000 mosquito can be confusing. I would keep with the classical SI expressed as %. The following sentence “Entomological inoculation rate (EIR) was defined as the number of positive in qrtPCR Plasmodium divided by the corresponding number of person-nights of collection, adjusted over the proportion of collected mosquitoes that were analyzed by qrtPCR Plasmodium” is fully unclear to me. What do you mean by “adjusted over the proportion of collected mosquitoes that were analyzed by qrtPCR Plasmodium”? Again, why not keeping with the classical definition of EIR: HBR * SI%/100.

The CI calculation might need approval from an external statistician.

Results:

Biodiversity of the Anopheles entomo-fauna

From my understanding of the Methods, the total person-nights should be 4200 and total cow-nights 420. Did I miss something?

“Potential malaria vectors”: could you provide your definition of those? A species table as sup data could be useful for non-specialist of Asian mosquito vectors. In figure 2 “other species” what does this include?

Fig2: why not commenting the comparative results for HLC and CBC and among the villages. Homogenize between fig 2 A, Legend, Table 1 and main text : CBC or CBT.

“Results of the molecular identification are presented in the Table 1.”

Malaria vectors

Table 2: Correct Mean Pf-EIR for Funestus : 0.1, not 0.01. Also in this table and followings, could not really follow the calculation from HBR and SI to the EIR, from the values of the former indexes, possibly as the mean values were calculated across the different months and/or villages. This might need to be specify at some point.

It would be also informative to provide the raw data as supp file for the identification of the species for the Plasmodium positive samples.

Sporozoite quantification: It is indeed interesting to be able to quantitate the sporozoite load for each positive mosquito. I would suggest to specify in the sup file that the calculated LOD of 6 Pf sporo or 4 Pv sporo are indeed per mosquito, according to your method.
For this load quantification data, I am not sure that any correlation with oocyst detection can be made. There is, to my knowledge, no method to determine in field collected mosquitoes that the oocysts one detects are the one providing the detected sporozoites in head-thoraces. They might come from a secondary infection. I nevertheless agree that in very low transmission area the probability for a mosquito to feed twice on a gametocyte carrier is rather very low. In addition, please indicate the oocyst detection method, PCR? And how the midgut were preserved?

Table 3 could be placed as supp data.

**Host-seeking behaviour of Anopheles mosquitoes**

First sentence: “overall”, be more precise as taking into account all mosquitoes captured by HLC… Nevertheless, again I cannot obtain the same numbers for both HBR and CBR, though very closed.

Paragraph on Zoophagy : I am not sure that a zoophagic index can indeed be calculated as the surface of a cow exceed the surface of the human skin for HLC provided by 10 persons , plus volume of air and odors and containment of the cow. This is my opinion on this, but preferentially zoophagic/anthropophagic is OK.

Outdoor and early biting paragraph: Figure 6 and 7: revised the labeling for Fig 6, no distinction in the grey colours. Although there is a tendency for early biting it does not stand for all species ie minimus. Globally there is far too much figures for this section. Relocating some to the sup data will save space for the next section, that is at the center of the question: who is transmitting and when?

**Entomological indices of malaria transmission**

As said above, this might be the most important analysis. However, I still have some difficulties with the calculation in Table 4 and 5, At what level were the mean calculated?

Because you have all these nice data as monthly collection (see sup file 1) , why not comparing HBR per species (or even group) and EIR for each village, over time. This will clearly show who and when most transmission occurs and easier to visualize as graphs rather than tables as Table 4 and 5.

**Discussion:**

In my general comment I already mentioned that the discussion could benefit from a better focus. I would add on the argument Pf and Pv transmission and season (second paragraph) that one needs to keep in mind, that beside having mosquitoes, gametocytes are also needed and Pv carriers who are not totally cured are excellent providers of gametocytes the year around. It is why it is also important that in the Methods section is included whether MDA was on or not and how Pv carriers were treated.

Also this sentence might be strongest :” In this study, only 35% of the mosquitoes infected with Plasmodium were collected indoors between 09:00 pm and 05:00 pm, because of outdoor and early biters", (page 14)., THE TIMING When people do not sleep under a net. The 35% appears as 36% in figure 8 : correct?

Lastly: the last sentence of this paragraph is unclear : “Consequently, the paradigm of residual transmission as experienced in high transmission settings of Africa does not apply to the Thailand-Myanmar border area and a drastic shift in vector-control interventions is required."
Could you explain what is the paradigm?

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

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?  
Partly

If applicable, is the statistical analysis and its interpretation appropriate?  
I cannot comment. A qualified statistician is required.

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

Are the conclusions drawn adequately supported by the results?  
Partly

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

**Reviewer Expertise:** molecular entomologist, malaria transmission

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.

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**Author Response 27 Jan 2019**

**Victor Chaumeau,** Centre hospitalier universitaire de Montpellier, Montpellier, France

We are thankful to the reviewer for her useful feed-back on the manuscript. The manuscript has been edited as per reviewer’s suggestions in order to improve the clarity of this report and to provide the reader with more background information on Southeast Asian malaria vectors. We agree with the reviewer that a more detailed explanatory analysis on the heterogeneity of malaria transmission in the four villages would have been valuable. Additional figures were added in the Supplementary File 1 in order to better show the differences in malaria transmission in the four villages. However, in the absence of appropriate datasets, the mechanisms by which local ecological factors and larval habitat interact with malaria transmission cannot be explained. Noteworthy, the primary objective of this paper was not to point out specific differences in particular study villages, but instead to identify key aspects of malaria transmission that are relevant to the Kayin state and other areas in Southeast Asia. Raw numbers used for the calculation of the entomological indices estimates were added in the corresponding tables, additional references on the method used for the calculation of entomological inoculation rate were added in the Methods section. Regarding reviewer’s suggestion on the discussion, we wish to keep the outline of the manuscript version 1. Although a comparison of the entomological indices in different geographic areas and across the endemicity spectrum would be of great interest, a review of the global data seems beyond the scope of the current paper as it would represent a
of the global data seems beyond the scope of the current paper as it would represent a tremendous amount of work and could be the object of a full review article. In addition, the specificities of malaria transmission in the 4 villages (and Southeast Asian malaria transmission settings) were discussed into details in the first version of the manuscript. The point that were discussed included the low infection rates in malaria vectors and high biting rates, the seasonality in *Plasmodium falciparum* transmission, the heterogeneity of the entomological indices, the low parasite loads in infected vectors, the high diversity of malaria mosquitoes, their outdoor and early biting pattern, their opportunistic host type selection and ability to feed on livestock's. Finally, clear recommendations for malaria vector-control in this region were made in a specific paragraph using our results as a rational: we suggested to evaluate the use of outdoor spraying of insecticides on daytime resting habitats to tackle exophilic mosquitoes and veterinary approaches to tackle zoophagic malaria vectors. Regarding reviewer's inquiry about larval source management, it is difficult for us to discuss this intervention because we did not collect data on larval stages. Existing bibliography and our data on adults suggest that larval habitats are diverse, fragmented and poorly accessible, which probably limits the feasibility of larval source management in Kayin state. Regarding the use of “village ecology” as a vector-control tool, it is unclear to us what set of interventions the reviewer was referring to. What may appear to be “dogmatic” for the reviewer (i.e. the specificity of transmission in Asia by mostly exophagic mosquitoes) may be considered by others as one of the main factor explaining the differences in mosquito bed-nets and indoor residual spraying outcomes in Southeast Asia when compared to Africa. In the current context of malaria elimination, we therefore believe that this notion should be at the center of the discussion.

Point-to-point answers to specific comments are given below:

- The Methods mention a 21 months survey, that indeed might cover 2 years. As the survey in the 4 villages are not fully superimposed, it might be wise to describe the overlap for a better comprehensive value of the data.

Reviewer's suggestion has been addressed by presenting the collection schedule in the Table S1 of the revised manuscript.

- The “Kayin” state is barely mentioned in the report, bringing confusion when done ie: appearing only on page 13 beside the title and the abstract with “Therefore, baseline intensity of malaria transmission in hotspot villages from the Kayin state is likely to be higher than that reported in this study”

Reviewer's concern was addressed by changing “Thailand-Myanmar border” to Kayin state were possible and by mentioning more often “Kayin state” in the main text.

- Lastly, if there is no real discussion on the difference similarity among the 4 villages, why attracting with the “4 villages” in the title?

We agree with the reviewer that a more detailed explanatory analysis on the heterogeneity of malaria transmission in the four villages would have been valuable. However, in the absence of appropriate datasets, this analysis is not possible. Documenting differences is particular villages was not the only objective of including 4 villages in the study design. Extensive longitudinal follow-up in four villages has also allowed us to identify key aspects of malaria entomology that would not have been accurately describe with less sites or time-point collections. Therefore, we think it is important to mention the duration of the follow-up and the number of villages in the title.

- Abstract: Some revision could be done if the authors agree on the following comments:

The abstract was modified as per reviewer’s suggestions.

- Providing the global description of the villages, as done in the J. Vector ecology Paper (ref14), would save time, rather than going to the published document as well as possibly useful for the discussion.

Reviewer's comment was addressed by adding more details in the paragraph “Study villages” of the Methods section.

- Was the MDA still active during the whole entomological survey?
The reviewer is right to inquire about the timing of mass antimalarial drug administration and to stress that the elimination intervention has been poorly described in the manuscript. As specific paragraph describing the intervention has been added to the Methods section and additional references were provided to the reader.

- Were the house randomly chosen?
  Yes, the houses were randomly selected and this has been stated in the revised manuscript.
- Were the preliminary data (ref14) included in the analysis?
  Yes, the preliminary data were included in the analysis.
- Consider simplification: instead “6 sites” use 5 houses plus one cow collection.
  The phrasing of this sentence has been simplified as per reviewer’s suggestion.
- Verify the accuracy of the sentence for mosquito collection under the net covering the cow.
  The accuracy of the sentence has been verified.
- Expressing SI / 1000 mosquito can be confusing. I would keep with the classical SI expressed as %.
  We agree with the reviewer. SI values were expressed as percentages in the revised manuscript.
- The following sentence “Entomological inoculation rate (EIR) was defined as the number of positive in qrtPCR Plasmodium divided by the corresponding number of person-nights of collection, adjusted over the proportion of collected mosquitoes that were analyzed by qrtPCR Plasmodium” is fully unclear to me. What do you mean by “adjusted over the proportion of collected mosquitoes that were analyzed by qrtPCR Plasmodium”? Again, why not keeping with the classical definition of EIR: HBR * SI%/100.
  This definition of EIR is based on the work of Drakeley et al. 2003 (An estimation of the entomological inoculation rate for Ifakara: a semi-urban area in a region of intense malaria transmission in Tanzania), also reviewed by Tusting et al. 2014 (Measuring Changes in Plasmodium falciparum Transmission: Precision, Accuracy and Costs of Metrics). When EIR is analysed as the product of SI per HBR (equation 1), computing confidence intervals (CIs) becomes difficult since HBR and SI are not independent of each other; CIs can be computed more easily when analyzing EIR as a biting rate (taking into account only the sporozoite-infected mosquitoes) (equation 2). Adjusting the number of person-nights of collection over the proportion of collected mosquitoes that were analyzed by qrtPCR allow to take into account that not all mosquitoes were screened for Plasmodium infection (equation 3). Of note, this adjustment makes the calculation equivalent to the “classical” definition of EIR (equation 4) and has little impact on the estimate value as 99% (50082/50708) of the collected specimens were analysed by PCR Plasmodium.

  (1) EIR=SI ×EIR
  (2) EIR=(Nb. sporozoite positive mosquitoes)/(Nb.person.nights of collection)
  (3) EIR=(Nb. sporozoite positive mosquitoes)/(Nb.person.nights of collection×(Nb.of mosquitoes tested by PCR “/” Nb.of collected specimens))
  (4) EIR=(Nb. sporozoite positive mosquitoes)/(Nb.of mosquitoes tested by PCR ) × (Nb.of collected specimens)/(Nb.person.nights of collection)

- The CI calculation might need approval from an external statistician.
  The method used for the CI calculation in this paper was also reviewed in Landier et al. WOR, 2017 and Chaumeau et al. JID, 2018.
- From my understanding of the Methods, the total person-nights should be 4200 and total cow-nights 420. Did I miss something?
  We are thankful to the reviewer for pointing this ambiguity. Exceptions to the initial study design have been explicitly mentioned in the corresponding paragraph of the Methods section. The number of person-nights and cow-nights for each survey is also presented in the Table S1 of the revised manuscript.
- “Potential malaria vectors”: could you provide your definition of those?
We are thankful to the reviewer for pointing this unclear wording, it has been removed from the revised manuscript.

- A species table as sup data could be useful for non-specialist of Asian mosquito vectors. Reviewer’s suggestion was addressed by adding a species table in the Introduction.
- In figure 2 “other species” what does this include?
  The legend of Figure 2 has been reworded. “Potential malaria vectors” has been changed to “a priori malaria vectors”, which refers to groups of Anopheles species that have been previously found infected with human malaria parasite in other studies (Funestus, Maculatus, Leucosphyrus, Barbirostris and Annularis Groups). “Other species” has been changed to “other groups” and includes all the other groups that have been reported in the paragraph “Anopheles diversity”. More details were also added in the legend note.
- Fig2: why not commenting the comparative results for HLC and CBC and among the villages.
  Ranges were cited in the main text to point village differences in the species distribution at the group level. Moreover, differences in species-specific distribution between villages assessed by molecular identification were commented into details in the main text and are reported in the Supplementary File 1, Table S2.
- Homogenize between fig 2 A, Legend, Table 1 and main text : CBC or CBT.

Figure and table captions, and in-text citations were harmonized to CBT in the revised manuscript.

- Table 2: Correct Mean Pf-EIR for Funestus : 0.1, not 0.01.

Mean Pf-EIR for Funestus Group has been corrected.

- Also in this table and followings, could not really follow the calculation from HBR and SI to the EIR, from the values of the former indexes, possibly as the mean values were calculated across the different months and/or villages. This might need to be specify at some point.

The raw number used for the calculation was provided in all tables. Unless specified otherwise, the calculation was made using the entire dataset.

- It would be also informative to provide the raw data as supp file for the identification of the species for the Plasmodium positive samples.

Reviewer’s suggestion was addressed by presenting the raw data of identification of the species for Plasmodium positive samples in the Supplementary File 1, Table S4.

- I would suggest to specify in the sup file that the calculated LOD of 6 P. f. spor or 4 P. v. spor are indeed per mosquito, according to your method.

Reviewer’s suggestion has been added to the revised Supplementary File 2.

- For this load quantification data, I am not sure that any correlation with oocyst detection can be made. There is, to my knowledge, no method to determine in field collected mosquitoes that the oocysts one detects are the one providing the detected sporozoites in head-thoraxes. They might come from a secondary infection. I nevertheless agree that in very low transmission area the probability for a mosquito to feed twice on a gametocyte carrier is rather very low.

We are thankful to the reviewer for pointing that the correlation between oocysts and sporozoite densities is not straightforward, especially in wild mosquitoes, and that it may yield to spurious interpretation. The correlation between oocysts and sporozoite densities has been removed from the revised manuscript.

- In addition, please indicate the oocyst detection method, PCR? And how the midgut were preserved?

A statement on oocysts detection method (PCR) and midgut preservation has been added in the Methods section of the revised manuscript in order to address reviewer’s comment.

- Table 3 could be placed as supp data.

Table 3 has been moved to supplemental data as per reviewer suggestion.
First sentence: “overall”, be more precise as taking into account all mosquitoes captured by HLC…

We are thankful to the reviewer for pointing this inaccurate wording. The word “overall” has been changed to “taking into account the whole dataset”.

Nevertheless, again I cannot obtain the same numbers for both HBR and CBR, though very closed.

We hope that the above explanations will help the reader in understanding the calculation used for the entomological indices.

I am not sure that a zoophagic index can indeed be calculated as the surface of a cow exceed the surface of the human skin for HLC provided by 10 persons, plus volume of air and odors and containment of the cow. This is my opinion on this, but preferentially zoophagic/anthropophagic is OK.

We are thankful to the reviewer for mentioning that estimating zoophagy accurately would require a different approach than that used in our study, although comparing the human- and cow-biting rates as an index can be useful to describe the capacity of some specie to feed on bovines. The wording “zoophagic index” has been changed to “cow-biting index” (CBI) to address reviewer’s comment.

Figure 6 and 7: revised the labeling for Fig 6, no distinction in the grey colours.

Distinction in the grey colours is shown in the Key box in the figure (figure S12 of the revised manuscript).

Although there is a tendency for early biting it does not stand for all species ie minimus. “Anopheles mosquitoes exhibited an outdoor and early biting pattern” has been changed to “Anopheles mosquitoes exhibited an outdoor and/or early biting pattern” in order to address reviewer’s comment. Although the biting rate of An. minimus did not peak during the early evening/night, An. minimus biting rate between 5-6 pm and 5-6 am was higher than that of any other Anopheles species. Moreover, the high biting rates observed at 5-6 pm and 5-6 am suggest that this species was active before and after the standard collection time, which would meet the definition of early biting.

Globally there is far too much figures for this section. Relocating some to the sup data will save space for the next section, that is at the center of the question: who is transmitting and when?

Some figure from this section were relocated in the supplementary file as per reviewer's suggestions.

As said above, this might be the most important analysis. However, I still have some difficulties with the calculation in Table 4 and 5. At what level were the mean calculated?

The mean was calculated using the entire dataset for the corresponding grouping variable (village in the table 4 and season in the table 5) using the definition provided in the Method section. Raw data used for the calculation of entomological indices estimates were added to all tables to help the reader understanding the details of the calculation.

Because you have all these nice data as monthly collection (see sup file 1), why not comparing HBR per species (or even group) and EIR for each village, over time. This will clearly show who and when most transmission occurs and easier to visualize as graphs rather than tables as Table 4 and 5.

The figures suggested by the reviewer were added in the Supplementary File 1, Figures S7-S10. We think that presenting mean values of entomological indices in tables 4 and 5 is important to understand malaria transmission in this area and compare with other settings, which would maybe not appear as clearly in the figures presenting the longitudinal data.

In my general comment I already mentioned that the discussion could benefit from a better focus. I would add on the argument Pf and Pv transmission and season (second paragraph)
that one needs to keep in mind, that beside having mosquitoes, gametocytes are also
needed and Pv carriers who are not totally cured are excellent providers of gametocytes the
year around. It is why it is also important that in the Methods section is included whether
MDA was on or not and how Pv carriers were treated.

More details on the elimination intervention were added in the Intervention paragraph, Methods
section and more reference were added in the revised version of the manuscript. We think that
discussing the the dynamic of gametocyte carriage may be beyond the scope of this article. For
example, Pf carriers are also likely to be excellent providers of gametocytes the year around if not
treated. A more detailed analysis of the relationship between parasitological and entomological
indices was published in the Journal of Infectious Diseases by Chaumeau et al. in 2018.

- Also this sentence might be strongest :” In this study, only 35% of the mosquitoes infected
  with Plasmodium were collected indoors between 09:00 pm and 05:00 pm, because of
  outdoor and early biters", (page 14),, THE TIMING When people do not sleep under a net.
The typo 05:00pm has been changed to 05:00am. The window 09:00pm to 05:00am indoors is
expected to cover the timing when people do sleep under a net. The stronger wording suggested
by the reviewer was added to the revised version of the manuscript.
- The 35% appears as 36% in figure 8 : correct?
The in-text citation has been corrected.
- Lastly: the last sentence of this paragraph is unclear : “Consequently, the paradigm of
  residual transmission as experienced in high transmission settings of Africa does not apply
to the Thailand-Myanmar border area and a drastic shift in vector-control interventions is
required.” Could you explain what is the paradigm?
The paradigm is that – based on observations made in Sub-Saharan Africa - universal coverage
with LLINs would prevent most of the infective bites, and conversely that the part of the
transmission that would not be covered by LLINs would be only a residue (i.e. a small part) of
transmission at baseline. Some control or elimination programs indeed still rely mainly, and
sometimes solely, on mass distribution of LLINs. The data presented in this study (as well as other
references cited in the discussion) clearly demonstrate that LLINs have only a marginal impact on
disease prevention in the GMS rather than preventing the main part of the transmission. Therefore,
we do not think that is would be accurate to refer to the part of the transmission that is not
prevented by LLINs as “residual transmission” in the transmission settings described in this study.

Competing Interests: No competing interests were disclosed.