Indoor and outdoor malaria transmission in two ecological settings in rural Mali: Implications for vector control

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Abstract

Background: Implementation and upscale of effective malaria vector control strategies necessitates understanding the multi-factorial aspects of transmission patterns. The primary aims of this study are to determine the vector composition, biting rates, trophic preference, and the overall importance of distinguishing outdoor versus indoor malaria transmission through a study at two communities in rural Mali.

Methods: Mosquito collection was carried out between July 2012 and June 2016 at two rural Mali communities (Dangassa and Koïla Bamanan) using pyrethrum spray-catch and human landing catch approaches at both indoor and outdoor locations. Species of Anopheles (An.) gambiae complex were identified by polymerase chain reaction (PCR). Enzyme-Linked Immuno-Sorbent Assay (ELISA) were used to determine the origin of mosquito blood meals and presence of Plasmodium (P) falciparum sporozoite infections.

Results: A total of 11,237 An. gambiae (s.l.) were collected during the study period (5,239 and 5,998 from the Dangassa and Koïla Bamanan sites, respectively). Of the 679 identified by PCR in Dangassa, An. coluzzii was the predominant species with 91.4% of the catch followed by An. gambiae (8.0%) and An. arabiensis (0.6%). At the same time in Koïla Bamanan, of 623 An. gambiae s.l., An. coluzzii accounted for 99% of the catch, An. arabiensis 0.8% and An. gambiae 0.2%. Human Blood Index (HBI) measures were significantly higher in Dangassa (79.4%; 95% Bayesian credible interval (BCI) [77.4, 81.4]) than in Koïla Bamanan (15.9%; 95% BCI [14.7, 17.1]). The human biting rates were higher during the second half of the night at both sites. In Dangassa, the sporozoite rate was comparable between outdoor and indoor mosquito collections. For outdoor collections, the sporozoite positive rate was 3.6% (95% BCI [2.1-4.3]) and indoor collections were 3.1% (95% BCI [2.4-5.0]). In Koïla Bamanan, the sporozoite rate was higher indoors at 4.3% (95% BCI [2.7-6.3]) compared with outdoors at 2.4% (95% BCI [1.1-4.2]). In Dangassa, corrected entomological inoculation rates (cEIRs) using HBI were for indoor locations were 13.74 infective bites/person/month (ib/p/m; 95% BCI: 9.21—19.14) and were 18.66 [12.55—25.81] ib/p/m, outdoors locations. For Koïla Bamanan,cEIRs were 1.57 [2.34 —2.72] ib/p/m and 0.94 [0.43—1.64] ib/p/m for indoor and outdoor locations, respectively. EIRs were significantly higher at the Dangassa site than the Koïla Bamanan site.

Conclusion: The findings in this work may indicate the occurrence of active, outdoor residual malaria transmission is comparable to indoor transmission in some geographic settings. The high outdoor transmission patterns observed here highlight the need for additional strategies to combat outdoor malaria transmission to complement traditional indoor preventive approaches such as long-lasting insecticidal nets (LLINs) and indoor residual spraying (IRS) which typically focus on resting mosquitoes.

Background
Malaria transmission is heterogeneous and heavily dependent on the local climatic and ecological conditions. The intensity of transmission varies according to geography and seasonality. Variations are commonly observed at both the village and household levels [1, 2]. Vector control relies mainly on large-scale indoor residual spraying (IRS) and bed net distribution, both being recommended by the World Health Organization [3]. With proper implementation, these approaches have been shown to be highly effective in reducing human-mosquito contact and the burden of malaria [4]. In fact, IRS played a central role in the ultimate eradication of malaria in Europe in the 1950s [5] and recent elimination in some African countries [6]. Distribution of long lasting insecticidal nets (LLINs) was a driver for the decline in malaria incidence between 2000 and 2015, accounting for an estimated 68% of the 663 million clinical cases averted over this time period [7]. *Anopheles gambiae* s.l. (*An. coluzzii, An. gambiae* and *An. arabiensis*) and *An. funestus* groups are the predominant malaria vectors in Mali. *An. gambiae* and *An. coluzzii*, two of the four major malaria vectors in Africa, have been described as being mainly endophagic [8–13]. There is increasing evidence of declines in both endophagic and endophilic behavior in the *An. gambiae* s.l. populations after the introduction of IRS and LLINs [14]. Because they both target indoor biting and resting mosquitoes, their use over time and across space might have led to behavioral changes in vector populations [15–18]. The reaction of the mosquito populations to these indoor-use control strategies has jointly led to declining endophilic behavioral activity and increased outdoor biting rates. This phenomenon can be exacerbated in tropical areas where, due to high temperatures, human populations tend to remain outdoors for longer periods of time and are bitten before sheltering indoors. Therefore, it appears that current control strategies focusing on indoor-based methods may not be enough to eliminate malaria transmission in most endemic countries [19, 20]. Hence it is necessary to understand the importance, ecology and dynamics of outdoor transmission to develop and implement appropriate outdoor control methods. The objectives of this study were therefore to investigate potential changes in vector species composition, feeding behavior and contribution to indoor and outdoor malaria transmission in two ecologically different situated in rural parts of Mali.

**Methods**

**Study sites**

Data was obtained from longitudinal mosquito population surveys conducted at two sites in Mali, the villages of Dangassa and Koïla Bamanan located in two distinct ecological settings (Fig. 1). At each study village, longitudinal surveys were carried out during 12-day field visits at the beginning of the transmission season (June-July) and at the end of the transmission season (October) each year from 2012 to 2016.

Dangassa (8°12’37.253”W and 12°8’46.279” N) is located in the health district of Ouéléssébougou, 75 km away south of Bamako. Between the fringe of the village and the river Niger, lays a plain of about 1 km that floods in the rainy season. It is used for rice cultivation in the rainy season and vegetables gardening in the dry season. This is a Sudanian eco-geographical zone with a rainy season lasting from June to
October and a dry season from November to May. The annual mean rainfall from 2012 to 2016 was 947.92 mm. The dominant winds are the monsoon (rainy season) and the harmattan (dry season).

Koïla Bamanan (5°45′32.188″W and 13°38′24.026″N) in the health district of Dioro, is located about 385 km south-east of Bamako. This is a Sahelian eco-geographical zone with a rainy season from July to October with about 250–500 mm of rain per year and a dry season from November to May. Koïla Bamanan is located in an irrigated area where controlled submersion rice cultivation is practiced. The dominant winds are the monsoon (rainy season) and the harmattan (dry season).

According to the 2012 Demographic and Health Survey (DHS) in Mali, the prevalence of malaria among children under five years old was 52% based on microscopy and 47% based on rapid diagnostic tests (RDTs)[21]. *Plasmodium (P.) falciparum* parasites accounted for 85–90% of malaria infections, while other parasite types included *P. malariae* (10–14%) and *P. ovale* (1%). *Anopheles gambiae* s.l. (*An. coluzzii, An. gambiae, An. arabiensis*) and *An. funestus* group are currently the main vectors in Mali.

**Mosquito Collection**

Two collection methods were used for adult mosquitoes sampling: Pyrethrum spray catch (PSC) and human landing catch (HLC). In total, 12 different surveys in each village over five years (2012–2016) were performed.

During each survey, three rounds of HLC were set up with two collection points. Two volunteers (one indoor and one outdoor) collected mosquitoes at each of the two points making a total collection effort of 12 man-nights per survey. The collection sessions occurred from 18:00 h to 06:00 h indoors and outdoors in two randomly selected rooms. Collected mosquitoes were kept in separate cups, labeled by collection night, location and collection and time point. The volunteers rotated between indoor and outdoor positions after every hour to account for any biases due to variability in the attractiveness of individuals to mosquitoes [22, 23]. Volunteers were selected among young adult males at least 18 years old from the study site communities. The volunteers were trained prior to data collection periods and were informed about the study risks before agreeing to participate. Volunteers were offered treatment if they presented to health centers with malaria symptoms during the study. Informed consent was sought from room owners or users for the mosquito collections as the collections involved overnight stays. The collected specimens on human baits were morphologically identified. All these operations were supervised by two experienced entomologists.

During each survey, three sessions of PSC were conducted in the rooms to collect indoor resting mosquitoes from 07:00 h to 09:00 h according to the WHO standard protocol [24]. After each session, the collected mosquitoes were sorted according to their abdominal status (unfed, fed, half-gravid, and gravid) and dissected in two portions (head-thorax and abdomen) for the laboratory process. Each mosquito was kept in a labeled 1.5 ml eppendorf tube containing 80% ethanol. Samples were stored at –20 °C freezer in the laboratory until used for further processing.

**Laboratory Analysis Of Mosquitoes**
The species composition of *An. gambiae complex* was determined by PCR \[25, 26\]. The blood meal sources of freshly fed and half gravid *Anopheles* mosquitoes were analyzed by a direct enzyme-linked immunosorbent assay (ELISA) using human antibodies \[27\]. Head and thorax portions of the preserved *Anopheles* mosquito specimens were tested for *P. falciparum* circumsporozoite proteins (CSPs) using the ELISA technique \[28\].

**Data Management And Analysis**

Data were entered in StudyTRAX database management system (version v3.2.0802, StudyTRAX, Macon, GA). The Win BUGS application (version 1.4.1) was used to analyze the data. Pearson's chi-square tests were used to compare proportions (species composition). The mosquito density, the human biting rates, the human blood index, the *Plasmodium falciparum* infection rates and the entomological inoculation rates were estimated using formulas in Table 1. Bayesian credible intervals (BCI) were also used to compare the proportions. All analyses were carried out with a 5% type I error threshold.

| Entomological parameters                          | Formula                                                                 |
|--------------------------------------------------|--------------------------------------------------------------------------|
| Densities of female mosquitoes per room          | Total number of collected mosquitoes/total number of rooms                |
| Human blood index                                | (Total number of positive in ELISA for human blood/total number of tested mosquitoes)x100 |
| Human biting rates (HLC)                         | Total number of collected mosquitoes/(total number of volunteers x number of rounds) |
| Human biting rates (PSC)                         | Total number of freshly fed mosquitoes/ number of residents that have slept in the rooms the previous night |
| Sporozoite infection rate in mosquitoes          | (Total positive in ELISA/ total tested)x100                               |
| Entomological Inoculation Rates (HLC)            | HBR x (Total positive in ELISA/total tested)                            |

**Ethics Approval And Consent To Participate**

This study was approved by the Institutional Review Board of Tulane University (11-255609) and the Ethics Committee of the Faculty of Medicine and Faculty of Pharmacy (FMPOS) at the University of Sciences, Techniques and Technologies of Bamako (USTTB) in Mali (2011/77/FMPOS). In each of the villages, community consent had been obtained prior to the occurrence of any study activities. Individual informed consent forms were obtained for PSCs and HLCs (for both room owners and data collectors) before starting mosquito collections.

**Results**
Anopheles mosquito species composition and abundance

During the study period, a total of 11237 An. gambiae s.l. females were collected by PSC and HLC approaches. Among these specimens, 46.6% (5239/11237) were collected in Dangassa and 53.4% (5998/11237) in Koïla Bamanan. Based on logistics and practical considerations, a sub-sample was randomly selected across surveys to conduct molecular species identification. Of the 679 identified by PCR in Dangassa, An. coluzzii was the predominant species with a prevalence of 91.4% followed by An. gambiae (8.0%) and An. arabiensis (< 1%). In Koïla Bamanan, 623 An. gambiae s.l. were identified by the same methods and An. coluzzii represented 99%, An. arabiensis 0.8% and An. gambiae 0.2% of the catches. Overall, the An. coluzzii frequency was higher in Koïla Bamanan (99%) than in Dangassa (91.4%) ($\chi^2 = 38.99, P < .001$). No significant difference was observed between the frequencies of An. arabiensis ($\chi^2 = 0.001, P = 0.970$) and An. gambiae ($\chi^2 = 0.081, P = 0.770$) at both sites.

Density And Human Blood Index

In terms of abundance, Fig. 2 shows the monthly mean densities of An. gambiae s.l. in both localities using the PSC method. The mean density over the entire study period was 12.8 An. gambiae s.l. per room/per day [95% BCI: 12.5—13.2] in Koïla Bamanan and was significantly higher in Dangassa (6.4 per room/day [95% BCI : 6.2—6.7]). The highest densities were observed in April and August of 2014 for both villages. However, the densities of Koïla Bamanan were considerably higher than those of Dangassa. The lowest densities were observed in April 2016 for both study sites.

Over the 12 surveys, the mean HBIs of An. gambiae s.l. for Dangassa and Koïla Bamanan were 79.4% and 15.9%, respectively (Fig. 2). In Dangassa, the highest HBI was observed in August 2014 (96.8%) and the lowest HBI (52.1%) in April 2015, while in Koïla Bamanan, the lowest (6.7%) was observed in October 2012 and the highest (46.0%) in April 2015. HBIs were significantly higher in Dangassa than in Koïla Bamanan ($\chi^2 = 1913.6, P < 0.001$) during the entire study period.

Monthly human biting rates (MHBRs), sporozoite infection rates (SIR) and entomological inoculation rates (EIRs)

Figure 3 shows the human biting rates and the entomological inoculation rates from PSC in Dangassa and Koïla Bamanan respectively. The average monthly human biting rate (MHBR) was 2.3 times higher in Koïla Bamanan (69.8 bites/person/month) than in Dangassa (30 b/p/m). Monthly variations in MHBRs were observed at both sites. The highest MHBRs were recorded in April and August 2014. However, they were higher in Koïla Bamanan (266.6 and 249.6 b/p/m) than in Dangassa (65.9 and 105.5 b/p/m) (Fig. 3).

There was no significant difference in overall (over the entire study period) sporozoite infection rates (SIR) between the two study sites (infection rates were 2.9%, [95% BCI: 2.3—3.6 for Dangassa versus 2.6% [95% BCI: 2.2—3.1] for Koïla Bamanan; Table 2). Monthly and annual variations were observed at both sites. In Dangassa, the highest SIRs were observed in October (8.9%) in both 2012 and October 2014...
While the peak of the infection rates were observed in October in Dangassa, in Koïla Bamanan there was a second peak that was observed for April though the two peaks were seen for April 2014 (7.1%) and 2016 (78.5%).

Table 2
Sporozoite rates of *Anopheles gambiae* s.l. mosquitoes collected by spray-catch from Dangassa and Koïla Bamanan from July 2012 to June 2016.

| Survey  | Dangassa | Koïla Bamanan |
|---------|----------|---------------|
|         | #tested  | IR (%)        | BCI     | #tested  | IR (%) | BCI     |
| Jul 2012 | 226      | 2.2           | [0.7—4.5] | 656      | 0.6    | [0.2—1.3] |
| Oct 2012 | 181      | 8.9           | [5.2—13.4] | 351      | 1.7    | [0.6—3.3] |
| Nov 2012 | 224      | 4.5           | [2.2—7.5]  | 96       | 0.1    | [0.0—1.1] |
| Apr 2013 | 81       | 0.1           | [0.0—1.3]  | 269      | 0.0    | [0.0—0.4] |
| Apr 2014 | 171      | 2.9           | [1.0—5.9]  | 875      | 7.1    | [5.5—8.9] |
| Aug 2014 | 596      | 1.5           | [0.7—2.6]  | 1078     | 1.4    | [0.8—2.2] |
| Oct 2014 | 169      | 5.9           | [2.9—9.9]  | 16       | 6.4    | [0.2—22.1] |
| Apr 2015 | 337      | 1.8           | [0.7—3.5]  | 183      | 2.8    | [0.9—5.6] |
| Jun 2015 | 141      | 2.9           | [0.8—6.2]  | 572      | 3.9    | [2.4—5.6] |
| Nov 2015 | 121      | 0.1           | [0.0—0.9]  | 47       | 2.2    | [0.1—7.9] |
| Apr 2016 | 46       | 2.3           | [0.1—8.0]  | 14       | 78.5   | [54.5—94.9] |
| Jun 2016 | 369      | 3.3           | [1.7—5.3]  | 963      | 0.7    | [0.3—1.4] |
| TOTAL   | 2662     | 2.9           | [2.3—3.6]  | 5120     | 2.6    | [2.2—3.1] |

Given the low human blood index observed in Koïla Bamanan, the EIR was corrected by multiplying it by the human blood index observed at each site. The mean EIR was significantly higher in Dangassa (0.70 infective bites/person/month = ib/p/m) than in Koïla Bamanan (0.29 ib/p/m; Fig. 3). In Dangassa, the highest monthly EIRs were observed during the months of October 2012 and 2014, (2.75 and 1.81 ib/p/m). In Koïla Bamanan, the highest EIRs were recorded for April 2014 and June 2015 (3.33 and 0.59 ib/p/m, respectively). As with the sporozoite infection rates, EIRs also were subject to monthly and annual variations.

**Outdoor vs indoor biting activities of An. gambiae s.l. in Dangassa and Koïla Bamanan**

Data for comparing the location of biting activities (indoor versus outdoor locations) were generated using mosquitoes collected by human landing catches. HBRs showed a wide patterns of monthly
variation for both indoor and outdoor locations in both villages (Fig. 4).

Figure 4 shows HBR per hour and by season. HBR was higher during the second part of the night indoors and outdoors regardless of site and season. We observed a peak of HBR outdoors between 02:00 h and 03:00 h, and then two peaks respectively between 01:00 h - 02:00 h and 04:00 h -05:00 h during the rainy season at Dangassa (Fig. 4A). In the dry season, the peak was observed between 02:00 h and 04:00 h, but it was lower than in the wet season (Fig. 4B). At Koïla Bamanan, two peaks were observed indoor during the rainy season, where the first occurred between 23:00 h and 00:00 h, and the second occurred between 01:00 h and 02:00 h (Fig. 4C). In the dry season, a single peak was observed between 23:00 h and 00:00 h indoor and outdoor at Koïla Bamanan (Fig. 4D).

In Dangassa, the highest MHBRs were observed in July, October and November 2012, and in April and August 2014 (Fig. 5). The average HBR over the study period was significantly higher outdoors (648.9 b/p/m [BCI: 642.6—655.2]) than indoors (560.3 b/p/m [BCI: 554.7—565.9]) suggesting an exophagic coefficient of 1.2 (648.9/560.3).

In Koïla Bamanan, the highest MHBRs were observed in October 2012, April and August 2014 and in June 2016 indoor and outdoor (Fig. 5). As observed in Dangassa, in Koïla Bamanan the average HBR over the study period was significantly higher for outdoor locations (242.4 b/p/m [BCI: 238.1—246.7]) than indoor locations (228.9 b/p/m [BCI: 225.1—232.6]) with an exophagic coefficient of 1.1 (242.4 / 228.9). HBRs were significantly higher in Dangassa than in Koïla Bamanan, both indoors and outdoors.

In Dangassa, the sporozoite infection rate in An. gambiae s.l. was 3.1% [BCI: 2.1—4.3] and 3.6% [BCI: 2.4—5.0] for indoor and outdoor locations, respectively. No significant differences were observed between these two rates. The sporozoite infection rates were subject to wide month-to-month variability. In Koïla Bamanan, the average sporozoite infection rate was 4.3% [95% BCI: 2.7—6.3] and 2.4% [95% BCI: 1.1—4.2] for indoor and outdoor locations, respectively. The sporozoite infection rate indoors was 1.8 times higher than outdoors, though no significant difference was observed between the two. Similar monthly variations were observed both indoors and outdoors for Dangassa (Table 3),
Table 3
Sporozoite rates of *Anopheles gambiae* s.l. mosquitoes collected by human landing catch indoors and outdoors in Dangassa and Koïla Bamanan from July 2012 to June 2016.

| Surveys | Dangassa | | | Koïla Bamanan | | |
|---------|----------|---------|----------|----------------|---------|---------|
|         | Indoor   | Outdoor | Indoor   | Outdoor         |         |         |
|         | #tested  | IR (%)  | #tested  | IR (%)          | #tested | IR (%)  |
| Jul 2012| 232      | 2.2     | 88       | 1.2             | 16      | 6.5     |
|         |          | [0.7—4.4] |        | [0.0—4.3]      |         | [0.2—22.1] |
| Oct 2012| 165      | 3.7     | 180      | 5.0             | 81      | 2.5     |
|         |          | [1.4—7.0] |        | [2.3—8.6]      |         | [0.3—6.8] |
| Nov 2012| 75       | 4.0     | 74       | 1.4             | 22      | 0.5     |
|         |          | [0.9—9.5] |        | [0.0—5.1]      |         | [0.0—4.4] |
| Apr 2013| 7        | 1.4     | 12       | 0.8             | 4       | 2.3     |
|         |          | [0.0—13.0] |      | [0.0—7.8]      |         | [0.0—22.1] |
| Apr 2014| 158      | 3.8     | 77       | 3.9             | 65      | 4.7     |
|         |          | [1.4—7.3] |        | [0.8—9.2]      |         | [1.0—10.9] |
| Aug 2014| 124      | 1.6     | 131      | 0.1             | 160     | 5.0     |
|         |          | [0.2—4.5] |        | [0.0—0.8]      |         | [2.2—8.9] |
| Oct 2014| 5        | 1.8     | 10       | 20.1            | 2       | 4.3     |
|         |          | [0.0—17.9] |      | [2.9—48.4]     |         | [0.0—41.9] |
| Apr 2015| 41       | 0.3     | 59       | 6.8             | 5       | 1.9     |
|         |          | [0.0—2.4] |        | [1.9—14.4]     |         | [0.0—18.0] |
| Jun 2015| 37       | 5.5     | 65       | 9.3             | 20      | 20.1    |
|         |          | [0.7—14.7] |      | [3.5—17.3]     |         | [6.1—39.6] |
| Nov 2015| 18       | 0.6     | 0        | 0               | 1       | 7.9     |
|         |          | [0.0—5.3] |        |                 |         | [0.0—72.7] |
| Surveys | Dangassa | | | | Koïla Bamanan | | |
|---|---|---|---|---|---|---|---|
| | Indoor | Outdoor | Indoor | Outdoor | Indoor | Outdoor |
| | #tested | IR (%) | #tested | IR (%) | #tested | IR (%) | #tested |
| Apr 2016 | 2 | 4.3 | [0.0—41.8] | 6 | 1.6 | [0.0—15.1] | 0 | 0.0 | 0 | 0.0 |
| Jun 2016 | 45 | 8.9 | [2.6—18.7] | 101 | 3.0 | [0.6—7.1] | 89 | 2.3 | [0.3—6.2] | 30 | 0.4 | [0.0—3.3] |
| TOTAL | 909 | 3.1 | [2.1—4.3] | 803 | 3.6 | [2.4—5.0] | 465 | 4.3 | [2.7—6.3] | 371 | 2.4 | [1.1—4.2] |

EIRs showed monthly variations (Fig. 5). In Dangassa, the EIRs were estimated to be 13.74 [BCI: 9.21—19.14] and 18.66 [BCI: 12.55—25.81] ib/p/m, for indoor and outdoor locations, respectively. The EIR was 1.4 times higher outdoors than indoors, though the difference was not significant. The highest rates were observed in July and October 2012. In Koïla Bamanan, EIRs were 1.57 [BCI: 2.34 —2.72] and 0.94 [BCI: 0.43—1.64] ib/p/m for indoor and outdoor locations, respectively. The highest rates were observed for August 2014 and June 2015. In contrast to Dangassa, the EIR was 1.7 times higher indoors than outdoors. However this difference was not significant.

**Discussion**

Vector control remains a critical approach for malaria control and eliminations efforts. However, its success depends on periodic assessment of local malaria vector behavior. This study analyzed entomological parameters to better understand malaria transmission in two different ecological settings of Mali: Dangassa in the Sudan savanna region along the Niger River and Koïla Bamanan in the Sahelian region that conducts double rice cropping a year. In both areas, we observed the predominance of *An. coluzzii* species. Previous studies conducted in Mali reported the predominance of this species in the Sudanese river savanna, flood prone and irrigated rice farming areas [29–32]. The authors explained their findings by the presence of favorable larval breeding sites to the development of this species such as ladder like brick pits, water puddles, ruts, flood prone plains and irrigated rice farming which were also common in our study sites. The highest density rates observed for Koïla Bamanan (twice) compared to Dangassa are most likely due to the rice cultivation which offers numerous suitable larval breeding habitats, especially during the earlier development stage of rice [33–35].
The outdoor human biting rates (HBR) were significantly higher than indoor biting rates. This could be due to the side effect of ongoing control measures like LLINs in these places as *An. gambiae* s.l., which were known to be more endophagic than exophagic [36]. Indeed, several studies have shown changing its biting and resting behavior from indoors to outdoors when LLINS were widely used in an area. Such an observation was also made in Equatorial Guinea [32, 37–40]. However, recent data from Benin reported higher HBRs indoors than outdoors [36].

The human blood index (HBI) was significantly higher in Dangassa than in the irrigated rice farming area of Koïla Bamanan. This observation has now become common in Mali as rice farming has been associated with lower HBI as demonstrated by many studies [33, 41]. The same observation has been reported in Senegal [42]. Explanation of this situation has been attributed to personal protection practices as the highest densities of *An. gambiae* s.l. are known to occur in rice growing areas. Thus, mosquitoes are diverged to alternative hosts present in the area such as bovines, sheep, and goats as it is in Koïla Bamanan (unpublished observations by our study team). In addition, from 2008 to 2014, Koïla Bamanan was part of the Millennium Villages’ Project where LLINs were freely distributed and replaced every three years. These practices significantly improved ownership and usage of LLINs which may have led to a behavioral change in trophic preference in the malaria vector. Many studies across Africa showed that anthropophagic and endophilic individuals could become zoophagic and exophilic from the intensification in the use of LLINs [15, 18, 43–45].

This study revealed a higher indoor than outdoor EIR at Koïla Bamanan. The inverse relationship was observed in Dangassa where outdoor EIR was higher than indoor EIR. These findings may be due to the higher outdoor HBR observed in Dangassa than in Koïla Bamanan. The higher outdoor HBR and EIR than indoors in Dangassa could be explained by prior findings that *An. gambiae* s.l. is known to be anthropophagic and endophilic but is becoming more exophilic and humans are staying outside houses longer and are bitten before going under mosquito nets. Other environmental factors, especially elevated ambient temperature, cause people to sleep with no protection against mosquitoes’ biting [36] or outdoors. In fact, several studies reported a high outdoor transmission rate [32, 37, 46]. This finding could be attributed to the use of LLINS distributed during prenatal screenings and mothers having completed the vaccination cycles of their babies who receive treated mosquito nets for free (before 2014). Also, the universal coverage in LLINS from 2014 could itself cause mosquito behavioral changes.

**Conclusion**

*An. gambiae* s.l. was more anthropophilic in Dangassa than in Koïla Bamanan. Both sites had comparable EIR outdoors and comparable indoor transmission which was also high at both sites. These findings have important implications for the epidemiology and strategies for control of malaria in the study area. Additional control strategies are needed to complement ongoing interventions to better address the issue of outdoor transmission and reduce indoor and outdoor resting vectors using the tool package offered by integrated vector management (IVM).
Declarations

Authors’ contributions

MK, MC NS and SD designed the study and monitored its implementation. MK and NS analyzed and interpreted the results. MK, IS, MT, SID, DK conducted field work and contributed in laboratory analysis. MK wrote the manuscript which was critically revised by MC, NS, SFT, SD, ABS, JGS, DJK and MD. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Availability of data and materials

The data used and/or analyzed in this study are available from the corresponding author on reasonable request.

Consent for publication

Not applicable.

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Figures

Figure 1
Map showing the study sites

Figure 2
Monthly mean density (MMD) and human blood index (HBI) of An. gambiae s.l. in human dwellings in Dangassa & Koïla Bamanan from 2012 to 2016

Figure 3
Monthly human biting rate (MHBR) and entomological inoculation rate (EIR) of Anopheles gambiae s.l. mosquitoes collected by spray-catch from Dangassa and Koïla Bamanan form July 2012 to June 2016.
Figure 4

Hourly biting activity of An. gambiae s.l in Dangassa (A and B) and Koïla Bamanan (C and D) during study period by season. Rainy season covers June to October and dry season covers November to May.
Figure 5

Monthly Human Biting Rate (MHBR) and Entomological Inoculation Rate (EIR) of Anopheles gambiae s.l. mosquitoes collected by Human Landing Catch (HLC) at Dangassa and Koïla Bamanan indoors and outdoors from July 2012 to June 2016