Insecticide resistance in indoor and outdoor-resting *Anopheles gambiae s.l.* in Northern Ghana

Majidah Hamid-Adiamoh (mojedamid@gmail.com)  
University of Ghana  https://orcid.org/0000-0002-0101-5445

Alfred Amambua-Ngwa  
MRC Laboratories The Gambia at LSHTM

Davis Nwakanma  
Medical Research Unit The Gambia at LSHTM

Umberto D’Alessandro  
Medical Research Council Unit The Gambia at LSHTM

Gordon A. Awandare  
West African Centre for Cell Biology of Infectious Pathogens (WACCBIP), University of Ghana

Yaw A. Afrane  
Department of Medical Microbiology, College of Health Sciences, University of Ghana

Research

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Abstract

**Background** Selection pressure from continued exposure to insecticides drives the development of insecticide resistance and changes in resting behavior of malaria vectors, which may support residual transmission in several endemic settings. There is a need to understand how resistance drives changes in resting behavior within vector species. Here, we examined the association between insecticide resistance and resting behavior of *Anopheles gambiae s.l.* in Northern Ghana.

**Methods** Adult mosquitoes were collected both indoors and outdoors from two communities using mouth aspirators and pit shelters. F₁ progenies from a subset of mosquitoes were exposed to dichloro diphenyl trichloroethane (DDT), deltamethrin, malathion and bendiocarb using WHO insecticide susceptibility tests. Insecticide resistance markers including voltage-gated sodium channel (Vgsc)-1014F, Vgsc-1014S, Vgsc-1575Y, glutathione-S-transferase epsilon 2 (GSTe2)-114T and acetylcholinesterase (Ace1)-119S, as well as blood meal sources were investigated using PCR methods. Activities of metabolic enzymes, acetylcholine esterase (AChE), non-specific β-esterases, glutathione-S-transferase (GST) and monoxygenases (oxidases) were measured from unexposed F₁ progenies using microplate assays.

**Results** Susceptibility of *An. coluzzii* to deltamethrin 24hr post-exposure was significantly higher in indoor (mortality=5%) than the outdoor (mortality=2.5%) populations (P=0.02). The mosquitoes were fully susceptible to malathion (mortality: indoor=98%, outdoor=100%). Susceptibility to DDT was significantly higher in outdoor (mortality=9%) than indoor (mortality=0%) mosquitoes (P=0.006). Mosquitoes were also found with suspected resistance to bendiocarb but mortality was not statistically different (mortality: indoor=90%, outdoor=95%. P=0.30). The frequencies of all resistance alleles were higher in F₁ outdoor (0.11-0.85) than indoor (0.04-0.65) mosquito populations, while Vgsc-1014F in F₀ *An. gambiae s.s* significantly associated with outdoor-resting behavior (P=0.01). Activities of non-specific β-esterase enzymes were significantly higher in outdoor than indoor mosquitoes (Mean enzyme activity: Outdoor= 1.70/mg protein; Indoor=1.35/mg protein. P<0.0001). AChE activity was also more elevated in outdoor (0.62/mg protein) than indoor (0.57/mg protein) mosquitoes but this was not significant (P=0.08). Human blood index (HBI) was predominantly detected in indoor (18%) than the outdoor mosquito population (3%).

**Conclusions** These findings revealed higher phenotypic resistance in indoor than outdoor-resting mosquitoes, but genotypic and metabolic resistance levels were higher in outdoor than the indoor mosquito populations. However, the overall results did not establish that there was a significant preference of resistant malaria vectors to solely rest indoors or outdoors, but varied depending on the resistant alleles present. Indication that human and animal blood meal indices were more prevalent in indoor-resting mosquitoes was also shown. Continued monitoring of changes in resting behavior within *An. gambiae s.l.* populations is recommended.

Background

Malaria control and elimination efforts rely heavily on vector control interventions, more specifically on long-lasting insecticidal nets (LLINs) and indoor residual spraying (IRS), that involve the use of insecticides [1]. The scale-up of LLIN, has contributed significantly to the decline of malaria burden observed over the last 10–15 years in sub-Saharan Africa [2]. Unfortunately, malaria vectors have developed resistance to the insecticides employed in vector control programs and indeed to almost all the classes of available insecticides [3]. Insecticide use has been associated with widespread physiological resistance and behavioral changes of malaria vectors which may contribute in maintaining residual malaria transmission [4, 5]. IRS and LLINs are meant to provoke a knock down or mortal effect on vectors upon contact, targeting their classical anthropophilic (human feeding), late night indoor biting (endophagic) and indoor resting (endophilic) behaviors [6, 7]. This applies specifically to the most efficient malaria vectors, namely Anopheles arabiensis, An. coluzzii, An. gambiae s.s. and An. funestus. Contrary to expectations, in settings where IRS and LLINs were extensively deployed, highly anthropophilic, late-indoor biting and indoor resting vectors have switched to animal feeding and outdoor human feeding following the deployment of vector control activities [8, 9]. For instance, An. gambiae s.l. populations in Bioko Island [10], Ghana [11], Senegal [12] and Tanzania [8] increased outdoor feeding behavior following extensive intervention with IRS and LLINs. Outdoor biting was also found in...
naturally endophilic An. funestus populations in Western Kenya [13]. Furthermore, vector populations have adapted to early and early-morning biting, targeting a time when humans are not protected by LLINs [14].

Intriguingly, recent studies done in areas of high IRS and LLINs coverage have shown concurrent indoor and outdoor feeding behavior within sibling species of An. gambiae s.l. from Benin [15], Ethiopia [16], Libreville [17], Tanzania [8] and Western Kenya [13]. However, there is little evidence of intra-species consistency or differences in insecticide-driven vector resting behavior. It is plausible that insecticide pressure may select for behavioral changes within species, such that resistant mosquitoes feed and survive indoors while susceptible mosquitoes adopt exophilic behavior. This can be further modulated by variation in molecular mechanisms that enable survival against insecticides.

Target site and metabolic resistance mechanisms have been shown to confer resistance to insecticides in An. gambiae s.l. [18–20]. Target site resistance involves mutation in the voltage-gated sodium channel (Vgsc) gene, mediating resistance to dichlorodiphenyltrichloroethane (DDT) and pyrethroids [21, 22], as well as acetylcholinesterase (ACE), responsible for carbamate and organophosphate resistance [23, 24]. Increased detoxifying activities of metabolic enzyme families including non-specific esterases, glutathione-S-transferases (GSTs) and monooxygenases (cytochrome P450s) were associated with resistance to the various malaria control insecticides [25, 26]. Several markers have been identified and widely used for resistance surveillance. In An. gambiae s.l., knockdown resistance (kdr) Vgsc-1014F, Vgsc-1014S, Vgsc-1575Y and glutathione-s-transferase epsilon 2 (GSTe2)-114T are markers associated with DDT and pyrethroid resistance [27, 28], whereas Ace1-119S is linked to carbamate and malathion resistance [29]. The prevalence of the resistance phenotypes and polymorphisms, as well as enzymatic activities, in association with vector behavioral patterns may help understand the effect of vector interventions and strategies to improve efficacy in specific malaria endemic populations.

In Ghana, reduction of the malaria burden has been attributed to the scale-up of IRS and LLINs [30]. However, emergence of insecticide resistance and changes in mosquito biting behavior have been documented in the local vector populations [11, 31], both of which could contribute to maintaining transmission despite high IRS and LLINs coverage [32]. This study therefore investigated the association between resting behavior of members of An. gambiae s.l. and insecticide resistance and its contribution to residual malaria transmission in Northern Ghana.

Materials And Methods

Study sites

The study was conducted in two rural communities in Northern Ghana which are 16 km apart, Kpalsogu (9.33⁰ N, 1.02⁰ W) and Libga (9.35⁰ N, 0.51⁰ W) (Fig. 1). Northern Ghana was chosen because the region continues to experience a high malaria burden than other regions in Ghana despite scaled-up malaria control interventions [33]. Kpasolgu is one of the sites for annual IRS conducted by the President Malaria Initiative (PMI) and Ghana National Malaria Control Programme (NMCP) since 2008. However, IRS started in Libga in 2008 but was discontinued from 2014. Both communities are in close proximity to dams linked to an irrigation scheme which allows uninterrupted farming activities throughout the year but also supports perennial breeding of mosquitoes [34]. Malaria transmission is seasonal in the areas without irrigation [35].

Mosquito collections and rearing in the insectary

Indoor and outdoor mosquito collections were conducted in July-November 2017 from each site every other day. Collections were done between 06:00 hr and 09:00 hr. Live indoor-resting mosquitoes were sampled using prokopack electrical aspirators [36]. Pit traps [37] were constructed outside houses to attract live outdoor-resting mosquitoes which were later collected with prokopack aspirators. Four pit traps were constructed in each village. Each trap was placed about 5 m from each compound and the houses were 50 m apart from each other. Both indoor and outdoor collections were done in 6–8 randomly selected compounds in each community. Mosquitoes were transferred into paper cups labeled as per their resting locations.
Mosquitoes were immediately transported to the insectary for morphological identification of species and abdominal status using taxonomic keys [38]. All blood fed, half-gravid and gravid F₀ female An. gambiae s.l. were kept in cages to lay eggs. They were provided with laying pads, made of filter paper on top of a wet cotton wool in a petri dish. Eggs were subsequently allowed to hatch and larvae reared to adult stage.

**Insecticide susceptibility bioassay**

Batches of 20–25 emerging F₁ adult females (2–5 days old) from 480 wild-caught F₀ females, were exposed to insecticide-impregnated papers containing 0.05% deltamethrin, 5% malathion, 0.1% bendiocarb and 4% DDT at a temperature range of 27-30°C and relative humidity of 63–67%, in a closed tube for one hour. Two batches of the same number of mosquitoes were exposed to untreated test papers as negative controls. Mosquitoes were then supplied with 10% sugar solution in a holding tube and mortality after 24 hours was recorded and scored according to WHO protocol [39]. Dead and surviving mosquitoes were separately stored in 1.5 ml Eppendorf tubes with silica gel for subsequent molecular tests for insecticide resistance mechanisms.

**Anopheles species identification**

Genotypic DNA was extracted from the legs of individual F₀ and F₁ female mosquitoes using Qiagen QIAxtractor robot. Species identification to the molecular level was carried out as previously done [40, 41]. All phenotyped F₁ and F₀ An. gambiae s.l. mosquitoes were analyzed for species identification. Primers included in the reaction were those that detect sibling species of An. gambiae complex, including An. arabiensis, An. coluzzii. An. gambiae s.s and An. melas; which are the relevant vectors of malaria in Ghana [42].

**Analyses of target site modifications**

From each resting location, 50 mosquitoes were selected per insecticide for genotyping of insecticide resistance polymorphisms in phenotyped mosquitoes. Selection was done using dplyr package in R (cran.r-project.org). Similarly, all 480 F₀ that laid eggs and the remaining wild mosquitoes that were not selected for egg laying were processed for genotypic assessment of insecticide resistance mechanisms. Single nucleotide polymorphism (SNP) markers of insecticide resistance were screened from DNA of each specimen using a TaqMan SNP genotyping probe-based assays [43]. These markers include Vgsc-1014F, Vgsc-1014S and Vgsc-1575Y for target site resistance mutations to DDT and pyrethroids in voltage-gated sodium channel [21, 22, 27]; Ace1-119S mutation, marker of resistance to carbamates and organophosphates [29] and Gste2-114T, a molecular marker of metabolic resistance to DDT [28]. Analysis of allele frequencies of kdr mutations was conducted in the F₁ generation of the An. coluzzii alone because they were the majority species encountered in the study sites.

**Metabolic enzyme activity assays**

Other subsets of emerging F₁ adult females (2–5 days old) were immediately frozen in -20°C for biochemical assays. The frozen specimens were analyzed for activities of metabolic enzymes including AChE, non-specific β-esterases, GSTs and monooxygenases (oxidases). 50 F₁ mosquitoes were analyzed from each of the study localities and they were not exposed to any insecticide prior to the assays. Microplate assay standard protocols as described [44] were followed for each enzyme, where all assays were run in triplicates and along with Kisumu strain as susceptible control population.

Briefly, individual whole adult mosquitoes (enzyme source) were homogenized in potassium phosphate (KPO₄) buffer and substrates to respective enzymes were added as well as chromogenic agents. Absorbance was measured using Varioskan Lux multimode microplate reader (Thermo Scientific) at specific wavelengths depending on the enzyme being measured. Acetylcholine esterase was measured at 414 nm in the presence of acetylthiocholine iodide (ATCh) as substrate; while β-esterases at 540 nm in the presence of β-naphthyl acetate. Monooxygenases (cytochrome P450) level was determined using 3, 3’, 5,5’-Tetramethyl-Benzidine Dihydrochloride (TMBZ) and absorbance captured at 620 nm. Lastly, glutathione-S-transferase with 1-chloro-2, 4’-dinitrobenzene (cDNB) at 340 nm. Total protein from individual mosquitoes was also analyzed to standardize the mean enzyme activity of the test samples.

**Analysis of blood meal sources**
Blood meal origins were determined from DNA extracted from the abdomens of blood-fed F₀ mosquitoes using the multiplex PCR protocol [45] modified by including primers that could amplify donkey and horse. This assay involves amplification of mitochondrial cytochrome B of An. gambiae and vertebrate hosts including cow, dog, donkey, goat, human, horse and pig from a single mosquito specimen.

**Data analysis**

Data from both study sites were pooled together as there was no significant difference in the results obtained. The level of insecticide susceptibility of mosquitoes was evaluated following WHO 2016 criteria [39]. Pearson's Chi squared test was used to determine the differences in mortality to insecticides by resistance allele and their frequencies between indoor and outdoor mosquito populations. Odds ratio was applied to determine the association between resistance phenotype and frequency of resistance alleles in F₁ mosquito populations exposed to insecticides.

Mean activities of each enzyme per mg of protein were compared between mosquitoes from the two resting locations and the reference susceptible strain, using one-way analysis of variance (ANOVA) with Holm-Sidak's multiple comparisons test. The mean enzyme activities between indoor and outdoor mosquitoes were compared using Mann-Whitney test. Human (HBI) and animal (BBI) blood indices were each calculated as total number of mosquitoes positive for human and animal DNA as a proportion of all blood fed mosquitoes expressed in percentage. All statistical analyses were performed using Stata/IC 15.0 (2017 StataCorp LP) and GraphPad Prism 8.0.1 software. P value of < 0.05 was considered significant in all data interpretations.

**Results**

**Anopheles mosquito species composition**

A total of 1,122 female An. gambiae s.l. mosquitoes were collected during the study, with majority found resting outdoors (58%, 652) than indoors (42%, 470). An. coluzzii was the predominant species, both indoors (36%, 375) and outdoors (39%, 413), followed by An. arabiensis (3%, 33) indoors and outdoors (12%, 125), and An. gambiae s.s. indoors (2%, 27) and outdoors (8%, 83). Five (5) hybrids of An. coluzzii/gambiae s.s were also identified.

Phenotypic resistance in F₁ An. coluzzii populations.

Species identification of all phenotyped samples revealed 98% of F₁ progeny were An. coluzzii both indoors and outdoors while the remaining species (An. arabiensis and An. gambiae) represented 2%. Mortality was generally higher in outdoor mosquitoes than the indoor populations. A 24-hour post-exposure mortality of 0% and 9% (95%CI: 3–12%) was observed for DDT with progeny of mosquitoes from indoor and outdoor respectively (Fig. 2) and this difference was statistically significant (Pearson $X^2 = 7.58$, df = 1, P = 0.006). Progeny of mosquitoes exposed to deltamethrin showed an overall mortality of 5% (95% CI: 1–12%) for indoor mosquitoes and 2.5% (95%CI: 8–34%) for outdoor-resting mosquitoes (Pearson $X^2 = 5.44$, df = 1, P = 0.02).

The indoor and outdoor mosquitoes exposed to bendiocarb showed suspected resistance with mortality of 90% (95%CI: 64–95%) in the indoor population and 95% (95%CI: 87–100%) in the outdoor population (Pearson $X^2 = 1.07$, df = 1, P = 0.30). Both the indoor and outdoor populations were fully susceptible to malathion, with 98% and 100% (95%CI: 87–100%) mortality for indoor and outdoor mosquitoes, respectively (Pearson $X^2 = 2.02$, df = 1, P = 0.16). There was no observed mortality (0%) in the controls for all insecticides tested.

Detection of resistance alleles in F₁ An. coluzzii populations

Resistance-associated allele frequencies were higher in outdoor-resting mosquitoes than the indoor population (Table 1). Vgsc-1014F and GSTe2-114T alleles were the most common in both phenotypically resistant and susceptible indoor and outdoor mosquitoes. In the deltamethrin-resistant mosquitoes, Vgsc-1014F frequency was 0.65 (indoor) and 0.67 (outdoor). However in the DDT-resistant mosquitoes, Vgsc-1014F frequency was 0.65 (indoor) and 0.73 (outdoor). These observed differences were
not statistically significant between the indoor and outdoor mosquito populations (Deltamethrin: Pearson $X^2 = 0.22$, df = 1, $P = 0.64$. DDT: Pearson $X^2 = 0.41$, df = 1, $P = 0.52$). The carriage of Vgsc-1014F mutation was strongly associated with resistance to deltamethrin (OR = 5.46, $P = 0.001$, 95% CI: 1.94−15.41) but not with DDT resistance (OR = 0.69, $P = 0.75$, 95% CI: 0.066−7.14). No Vgsc-1014S allele was detected in any of the mosquitoes.

Vgsc-1575Y mutation was detected mainly in the deltamethrin-resistant outdoor An. coluzzii populations (frequency = 0.27). GSTe2-114T mutation was significantly higher in outdoor-resting (0.85) mosquitoes than the indoor (0.56) DDT-resistant mosquitoes (Pearson $X^2 = 5.73$, df = 1, $P = 0.02$). This mutation was also identified in mosquitoes resistant to deltamethrin (indoor = 0.62, outdoor = 0.84).

Ace1-119S was detected in a single indoor and an outdoor An. coluzzii specimens that survived bendiocarb exposure. It was also found in a single bendiocarb-resistant outdoor mosquito. The allele was detected only in malathion-susceptible mosquitoes at frequency of 0.08 (indoor) and 0.12( outdoor) with no significant difference (Pearson $X^2 = 0.003$, df = 1, $P = 0.96$).

Table 1: Frequencies (proportions) of resistance alleles in indoor and outdoor F$_1$ An. coluzzii populations based on insecticide resistance phenotypes (dead and alive)

<table>
<thead>
<tr>
<th></th>
<th>Dead</th>
<th>Alive</th>
<th>Dead</th>
<th>Alive</th>
<th>Dead</th>
<th>Alive</th>
<th>Dead</th>
<th>Alive</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Indoor</td>
<td>0.5</td>
<td>0.65</td>
<td>0</td>
<td>0.07</td>
<td>0.5</td>
<td>0.62</td>
<td>1</td>
</tr>
<tr>
<td>Deltamethrin</td>
<td>(N = 8)</td>
<td>(N = 81)</td>
<td>(N = 8)</td>
<td>(N = 82)</td>
<td>(N = 8)</td>
<td>(N = 82)</td>
<td>(N = 8)</td>
<td>(N = 82)</td>
</tr>
<tr>
<td>Outdoor</td>
<td>1</td>
<td>0.65</td>
<td>1</td>
<td>0.27</td>
<td>1</td>
<td>0.84</td>
<td>0.75</td>
<td>0.73</td>
</tr>
<tr>
<td>DDT</td>
<td>(N = 4)</td>
<td>(N = 49)</td>
<td>(N = 0)</td>
<td>(N = 50)</td>
<td>(N = 4)</td>
<td>(N = 52)</td>
<td>(N = 55)</td>
<td>(N = 3)</td>
</tr>
<tr>
<td>Indoor</td>
<td>0</td>
<td>0.65</td>
<td>0</td>
<td>0.04</td>
<td>0</td>
<td>0.56</td>
<td>0.06</td>
<td>1</td>
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<tr>
<td>Outdoor</td>
<td>0.75</td>
<td>0.73</td>
<td>0</td>
<td>0.11</td>
<td>1</td>
<td>0.85</td>
<td>0.12</td>
<td>0</td>
</tr>
<tr>
<td>Bendiocarb</td>
<td>(N = 55)</td>
<td>(N = 3)</td>
<td>(N = 69)</td>
<td>(N = 3)</td>
<td>(N = 50)</td>
<td>(N = 52)</td>
<td>(N = 56)</td>
<td>(N = 3)</td>
</tr>
<tr>
<td>Indoor</td>
<td>0.06</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Outdoor</td>
<td>(N = 59)</td>
<td>(N = 1)</td>
<td>(N = 60)</td>
<td>(N = 1)</td>
<td>(N = 50)</td>
<td>(N = 52)</td>
<td>(N = 54)</td>
<td>(N = 1)</td>
</tr>
<tr>
<td>Malathion</td>
<td>0.08</td>
<td>0</td>
<td>0.12</td>
<td>0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Detection of resistance alleles in F$_0$ An. gambiae s.l populations

The frequency of resistance alleles between the indoor and outdoor mosquitoes varied by mosquito species. Whereas Vgsc-1014S was not detected in the F$_1$ An. coluzzii, it was observed mainly in the F$_0$ An. arabiensis resting outdoors. Vgsc-1014F mutation was significantly higher in outdoor (0.99) resting mosquitoes compared to those indoors (0.77) in An. gambiae ss. (Pearson $X^2 = 31.6$, df = 2, $P = 0.001$) (Table 2). There was an indication of association of outdoor-resting behavior with resistance in An. gambiae ss. population carrying the Vgsc-1014F mutation (OR = 0.05, $P = 0.01$, 95% CI: .005-0.419). Although, An. coluzzii was the predominant species collected both indoors and outdoors, the difference in the frequency of this mutation in indoor (0.65) and outdoor (0.70) populations was not statistically significant (Pearson $X^2 = 0.7$, df = 2, $P = 0.4$). However, the higher prevalence of the mutation in indoor (0.48) than the outdoor (0.21) An. arabiensis population was significant (Pearson $X^2 = 6.42$, df = 2, $P = 0.04$). Vgsc-1014S was mainly found in indoor An. arabiensis (0.42).
Vgsc-1575Y was detected at an almost similar level (frequencies: Indoor = 0.21, outdoor = 0.2) in An. coluzzii populations. Further, no significant difference was observed in indoor (0.30) and outdoor (0.18) An. gambiae s.s (Pearson $X^2 = 1.2$, df = 1, P = 0.27).

Ace1-119S mutation was most frequent in An. gambiae s.s, although there was no statistically significant difference in the frequencies between indoor (0.25) and outdoor (0.31) populations (Pearson $X^2 = 0.2$, df = 1, P = 0.65). The prevalence was 0.1 in indoor and outdoor An. coluzzii.

<table>
<thead>
<tr>
<th></th>
<th>Vgsc-1014F</th>
<th>Vgsc-1014S</th>
<th>Vgsc-1575Y</th>
<th>GSTe2-114T</th>
<th>Ace1-119S</th>
</tr>
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<tbody>
<tr>
<td>An. arabiensis</td>
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<td>Outdoor</td>
<td>Indoor</td>
<td>Outdoor</td>
<td>Indoor</td>
</tr>
<tr>
<td></td>
<td>0.48</td>
<td>0.21</td>
<td>0.42</td>
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<td>N = 19</td>
<td>N = 97</td>
<td>N = 15</td>
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<tr>
<td>An. coluzzii</td>
<td>Indoor</td>
<td>Outdoor</td>
<td>Indoor</td>
<td>Outdoor</td>
<td>Indoor</td>
</tr>
<tr>
<td></td>
<td>0.65</td>
<td>0.7</td>
<td>0.01</td>
<td>0.01</td>
<td>0.21</td>
</tr>
<tr>
<td></td>
<td>N = 352</td>
<td>N = 401</td>
<td>N = 125</td>
<td>N = 122</td>
<td>N = 358</td>
</tr>
<tr>
<td>An. gambiae s.s</td>
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<td>Outdoor</td>
<td>Indoor</td>
<td>Outdoor</td>
<td>Indoor</td>
</tr>
<tr>
<td></td>
<td>0.77</td>
<td>0.99</td>
<td>0</td>
<td>0.5</td>
<td>0.3</td>
</tr>
<tr>
<td></td>
<td>N = 22</td>
<td>N = 74</td>
<td>N = 5</td>
<td>N = 2</td>
<td>N = 23</td>
</tr>
<tr>
<td>An. coluzzii/gambiae s.s</td>
<td>Indoor</td>
<td>Outdoor</td>
<td>Indoor</td>
<td>Outdoor</td>
<td>Indoor</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

Metabolic enzyme activities in F$_1$ An. coluzzii populations

An overall highly significant elevated levels of AChE ($F_{2,237} = 55.93$, $P < 0.0001$) and β-esterase ($F_{2,237} = 159.0$, $P < 0.0001$) activities were observed in both indoor and outdoor mosquito populations compared to the susceptible reference strain, Kisumu (Figs. 3A-D). Conversely, in both mosquito populations, the activities of the GSTs and monooxygenases were less relative to Kisumu but this was not significant in monooxygenase activity ($F_{2,237} = 0.6589$, $P = 0.52$).

AChE activity was not significantly higher in the outdoor (0.62/mg protein) than the indoor (0.57/mg protein) population (Mann–Whitney U = 5037, Z=-1.73, P = 0.08). The elevation in enzyme activity was found (Table 3) to be 2.48 fold (indoor) and 2.7 fold (outdoor) significantly higher than in Kisumu ($P < 0.0001$). Similarly, non-specific β-esterase activity in the outdoor-resting mosquitoes (1.70/mg protein) was significantly more than the indoor mosquitoes (1.35) (Mann–Whitney U = 0.5, Z=-8.33, P < 0.0001); with 1.69 (indoor) and 2.13 (outdoor) significant fold changes ($P < 0.0001$). No significant difference was detected in the level of GST activity between the two mosquito populations (indoor: 0.01/mg protein, outdoor: 0.02/mg protein, (Mann–Whitney U = 5709, Z = -0.29, P = 0.78). Monooxygenase activities also showed a similar level in both indoor and outdoor mosquitoes (mean activity = 0.21/mg protein, (Mann–Whitney U = 4989, Z = -1.84, P = 0.07). The fold difference in monooxygenase activities in Kisumu population (1.11) was not statistically significant ($P = 0.59$).
Table 3
Mean activities of individual enzyme and the fold change in mosquito populations relative to Kisumu

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Mosquito population</th>
<th>Mean enzyme activity (95% CI)</th>
<th>Fold change</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Kisumu</td>
<td>0.23 (0.22–0.24)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AChE</td>
<td>Indoor</td>
<td>0.57 (0.54–0.60)</td>
<td>2.48</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td></td>
<td>Outdoor</td>
<td>0.62 (0.58–0.66)</td>
<td>2.7</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td></td>
<td>Kisumu</td>
<td>0.80 (0.78–0.81)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>β-esterase</td>
<td>Indoor</td>
<td>1.35 (1.31–1.39)</td>
<td>1.69</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td></td>
<td>Outdoor</td>
<td>1.70 (1.65–1.76)</td>
<td>2.13</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td></td>
<td>Kisumu</td>
<td>0.46 (0.45–0.47)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>GST</td>
<td>Indoor</td>
<td>0.01 (0.0–0.01)</td>
<td>0.02</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td></td>
<td>Outdoor</td>
<td>0.02 (0.01–0.02)</td>
<td>0.04</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td></td>
<td>Kisumu</td>
<td>0.19 (0.18–0.19)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Monooxygenase</td>
<td>Indoor</td>
<td>0.21 (0.18–0.23)</td>
<td>1.11</td>
<td>0.59</td>
</tr>
<tr>
<td></td>
<td>Outdoor</td>
<td>0.21 (0.19–0.22)</td>
<td>1.11</td>
<td>0.59</td>
</tr>
</tbody>
</table>

Host blood meal sources of wild $F_0$ mosquitoes

A total of 165 out of 214 blood-fed mosquitoes were successfully identified to have fed either on human or animal hosts. The overall vertebrate positivity rate was higher in indoor-resting mosquitoes (Table 4), predominantly in An. coluzzii, which was the most abundant species in the study sites. Overall human blood index (HBI) was 21% and again more prominent in indoor (18%) than outdoor (3%) mosquitoes. 69% of HBI was detected from indoor An. coluzzii, followed by 17% in outdoor An. coluzzii. An. arabiensis was found with only 9% (indoor) and 1% (outdoor) HBI, while 1% HBI was identified in indoor An. gambiae only.

The principal animal blood meal source was from goat, representing 36% of indoor against 8% of outdoor-resting An. coluzzii. The other animal blood sources included cows, dogs, donkeys, horses and pigs and were mainly detected in indoor An. coluzzii specimens. Fewer (4%) An. arabiensis resting indoors fed on animal blood compared to the outdoor population (10%), similar to An. gambiae s.s indoor (0.6%) and outdoor (1.2%) population. Mixed human and goat blood meal was identified from a single indoor An. coluzzii specimen. Also, mixed cow and goat meal were found in three outdoor An. coluzzii specimens and one outdoor An. gambiae s.s specimen.
Table 4
Proportion of blood meal origin of the indoor and outdoor-resting mosquito populations

<table>
<thead>
<tr>
<th></th>
<th>An. arabiensis</th>
<th>An. coluzzii</th>
<th>An. gambiae</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Proportion (n)</td>
<td>Proportion (n)</td>
<td>Proportion (n)</td>
</tr>
<tr>
<td>Human</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Indoor</td>
<td>0.02 (3)</td>
<td>0.15 (24)</td>
<td>0.01 (1)</td>
</tr>
<tr>
<td>Outdoor</td>
<td>0.01 (1)</td>
<td>6 (0.04)</td>
<td>0</td>
</tr>
<tr>
<td>Cow</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Indoor</td>
<td>0</td>
<td>0.01 (2)</td>
<td>0</td>
</tr>
<tr>
<td>Outdoor</td>
<td>0</td>
<td>0.01 (2)</td>
<td>0</td>
</tr>
<tr>
<td>Dog</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Indoor</td>
<td>0</td>
<td>0.05 (8)</td>
<td>0</td>
</tr>
<tr>
<td>Outdoor</td>
<td>0</td>
<td>0.01 (2)</td>
<td>0</td>
</tr>
<tr>
<td>Donkey</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Indoor</td>
<td>0</td>
<td>0.04 (7)</td>
<td>0</td>
</tr>
<tr>
<td>Outdoor</td>
<td>0</td>
<td>0.01 (1)</td>
<td>0</td>
</tr>
<tr>
<td>Goat</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Indoor</td>
<td>0.04 (7)</td>
<td>0.36 (59)</td>
<td>0.01 (2)</td>
</tr>
<tr>
<td>Outdoor</td>
<td>0.07 (12)</td>
<td>0.08 (13)</td>
<td>0.01 (1)</td>
</tr>
<tr>
<td>Horse</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Indoor</td>
<td>0</td>
<td>0.02 (3)</td>
<td>0</td>
</tr>
<tr>
<td>Outdoor</td>
<td>0.01 (2)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Pig</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Indoor</td>
<td>0</td>
<td>0.01 (1)</td>
<td>0</td>
</tr>
<tr>
<td>Outdoor</td>
<td>0.01 (2)</td>
<td>0.01 (1)</td>
<td>0</td>
</tr>
<tr>
<td>Mixed hosts</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Indoor (Human + goat)</td>
<td>0</td>
<td>0.01 (1)</td>
<td>0</td>
</tr>
<tr>
<td>Outdoor (Cow + goat)</td>
<td>0</td>
<td>0.02 (3)</td>
<td>0.01 (1)</td>
</tr>
</tbody>
</table>

Discussion

In this study, we found phenotypic resistance to be higher in indoor-resting than outdoor-resting An. coluzzii mosquito progeny. Higher resistance-associated alleles and elevated activities of two metabolic enzymes were however recorded in the outdoor-resting population. Overall, there was not a significant association between insecticide resistance and resting location of the mosquito populations; but there was a tendency for An. coluzzii to rest indoors when phenotypically resistant to DDT and An. gambiae ss to rest outdoors when resistance was genotypically due to Vgsc-1014F mutation. Enhanced AChE and β-esterases
activities were also prominent in outdoor-resting F₁ An. coluzzii mosquitoes. Moreover, human and animal blood meal indices were higher in indoor than the outdoor mosquito population but with no statistical significance.

A probe into insecticide-driven genetic adaptation in vector population at intra-species level was a main interest in this study. We hypothesized higher resistance levels in indoor compared to the outdoor populations, due to increased contact with insecticide which amplifies their propensity to develop resistance [46]. This was mainly evident in the F₁ An. coluzzii populations exposed to insecticides where indoor mosquitoes were less susceptible to three out of four insecticides. Phenotypic resistance was especially prominent in DDT-exposed indoor populations. In contrast, genotypic resistance and enzymatic activities were more prevalent in all outdoor F₁ mosquito populations. This may be due to selection pressure and cross-resistance from other pyrethroids used in LLINs [47].

This study showed an association between outdoor-resting behavior in F₀ An. gambiae s.s population and Vgsc-1014F mutation. High frequencies of resistance markers associated with DDT resistance, Vgsc-1575Y and GSTe2-114T, were also observed in both F₀ and F₁ indoor and outdoor An. coluzzii populations. These may result from selection pressure due to the widespread use of similar insecticides for both public health and agriculture. As noted, year-round agriculture is practiced in the study sites with pesticide control measures in place [48, 49]. This could further explain why there were no significant difference in frequencies between the indoor and outdoor populations. Notably, resistance has been previously reported at varying levels to DDT and deltamethrin across all vector species in Ghana [44, 47] and the neighboring countries including Benin [19], Burkina Faso [27] and Togo [50]. High frequencies of resistance loci may compromise the effectiveness of vector control in the study areas that could subsequently accentuate residual transmission [35].

Target site polymorphisms may not fully explain resistance in vector populations [51], thus we probed the possible metabolic mechanisms involved. We identified significant increase in the activities of AChE and β-esterases, both of which have been associated with resistance to the insecticides tested [52]. Consistently, the observed fold change in β-esterases activities which was significantly higher than the susceptible strain, may demonstrate a possible role in deltamethrin resistance in the vector populations as previously reported [53, 54]. On the other hand, the decreased levels of GSTs and monooxygenases detected may indicate that they do not contribute to the DDT and deltamethrin resistance in the study mosquito populations. Therefore, genotypic mechanism alone may be mediating the documented DDT resistance in these mosquito populations. Interestingly, an increased activity of AChE was identified despite low level of Ace1-119S and no phenotypic resistance in the carbamate and organophosphate insecticides in the study. This may likely reflect other role of this enzyme which may not be related to resistance in the vector population. Since there was no documented use of carbamate insecticide for IRS in this region expect for agricultural use [55], perhaps this resistance selection may be from agriculture use.

We found human and animal blood indices to be higher in indoor-resting mosquitoes than the outdoor population despite a higher outdoor collection. This indicates that in spite of the fact that the study areas were under high IRS and LLINs interventions, mosquitoes were able to have either fed on their host indoors or outdoors and still successfully rested indoors despite interventions; thus retained their indoor-resting behavior. Plausibly, the blood-fed endophilic population could be among the indoor-resistant populations that are capable of maintaining contact with insecticides due to their age and feeding status [56, 57]. This scenario could also expose human to infective bites and possible malaria risk thus promoting residual malaria transmission under high intervention as earlier described [4]. Due to logistical reasons, we could not undertake circumsporozoite detection.

The predominant vector species identified was An. coluzzii, which is known to be highly endophilic and anthropophilic [58], however, the results here suggested that this vector population were displaying high zoophilic behavior. The abundant presence of animals in the study areas and reduced access to human host due to intervention may have driven zoophagy and exophagy in this vector species as previously suggested [59, 60]. Further studies could explore the dynamics of this behavior and its implication on the control efforts in the study region.

**Conclusions**
This study demonstrated that An. coluzzii phenotypically resistant to DDT had a higher propensity for indoor-resting behavior, while outdoor-resting tendency was found in those phenotypically-resistant to deltamethrin. Also, An. coluzzii with increased AChE and β-esterases activity, and An. gambiae s.s with Vgsc-1014F mutation displayed outdoor-resting behavior. Mosquitoes resting indoors were found to have fed more on both human and animals than their outdoor counterparts. Our findings highlight variation in response of mosquitoes within the same species to insecticide-based interventions. We recommend continued monitoring of vector behaviors in surveillance programs to help in the control of malaria.

Data Availability Statement:

All relevant data are within the paper. No supporting Information is available.

**Abbreviations**

Ace-1  
acetylcholinesterase 1  
DDT  
dichlorodiphenyltrichloroethane  
DNA  
deoxyribonucleic Acid  
GSTe2  
glutathione-s-transferase epsilon 2  
HBI  
human blood index  
IRS  
indoor residual spraying  
Kdr  
knockdown resistance  
LLIN  
long-lasting insecticidal net  
NMCP  
National Malaria Control Programme  
PCR  
polymerase chain reaction  
PMI  
President Malaria Initiative  
SNP  
single nucleotide polymorphism  
Vgsc  
voltage-gated sodium channel  
WHO  
World Health Organization

**Declarations**

**Acknowledgements**

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Authors’ contributions

MHA designed, performed the laboratory work, analyzed data and drafted the manuscript. YAA conceived and supervised the study, analyzed data and revised the manuscript. AAN and DA supervised study and revised the manuscript. UDA and GAA revised the manuscript.

Competing Interests:

The authors have declared that no competing interests exist.

Ethical Approval

Ethical approval was obtained from the Institutional Review Board of the Noguchi Memorial Institute for Medical Research (NMIMR), University of Ghana. Verbal informed consent of the village leaders and compound/household heads were also taken before mosquitoes were collected from the study villages.

Consent for publication

Not applicable

References


**Figures**
Figure 1

Map of study sites
Figure 2

Phenotypic resistance to the four insecticides tested in indoor and outdoor mosquitoes. * P=0.02. ** P=0.006
Figure 3

Mean enzyme activities observed in individual enzymes in indoor and outdoor mosquitoes and the susceptible reference strain, Kisumu.