Insecticide resistance status in *Anopheles gambiae* (s.l.) in coastal Kenya

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**Abstract**

**Background:** The rapid and widespread evolution of insecticide resistance has emerged as one of the major challenges facing malaria control programs in sub-Saharan Africa. Understanding the insecticide resistance status of mosquito populations and the underlying mechanisms of insecticide resistance can inform the development of effective and site-specific strategies for resistance prevention and management. The aim of this study was to investigate the insecticide resistance status of *Anopheles gambiae* (s.l.) mosquitoes from coastal Kenya.

**Methods:** *Anopheles gambiae* (s.l.) larvae sampled from eight study sites were reared to adulthood in the insectary, and 3- to 5-day-old non-blood-fed females were tested for susceptibility to permethrin, deltamethrin, dichlorodiphenyltrichloroethane (DDT), fenitrothion and bendiocarb using the standard World Health Organization protocol. PCR amplification of rDNA intergenic spacers was used to identify sibling species of the *An. gambiae* complex. The *An. gambiae* (s.l.) females were further genotyped for the presence of the L1014S and L1014F knockdown resistance (kdr) mutations by real-time PCR.

**Results:** *Anopheles arabiensis* was the dominant species, accounting for 95.2\% of the total collection, followed by *An. gambiae* (s.s.), accounting for 4.8\%. *Anopheles gambiae* (s.l.) mosquitoes were resistant to deltamethrin, permethrin and fenitrothion but not to bendiocarb and DDT. The L1014S kdr point mutation was detected only in *An. gambiae* (s.s.), at a low allelic frequency of 3.33\%, and the 1014F kdr mutation was not detected in either *An. gambiae* (s.s.) or *An. arabiensis*.

**Conclusion:** The findings of this study demonstrate phenotypic resistance to pyrethroids and organophosphates and a low level of the L1014S kdr point mutation that may partly be responsible for resistance to pyrethroids. This knowledge may inform the development of insecticide resistance management strategies along the Kenyan Coast.

**Keywords:** *Anopheles*, Insecticide resistance, Kdr, Sodium channel, Coastal Kenya
vectors to these insecticide classes has become a major global threat to their long-term use in the fight against malaria [3–12].

In Kenya, IRS is reserved for small-scale sporadic spraying during epidemics, and this management strategy has only been implemented in the western region of the country [13, 14]. The main insecticide used to supplement deltamethrin (a pyrethroid ester insecticide) in IRS is pirimiphos-methyl (Actellic 300CS; Syngenta Group, Basel, Switzerland), an organophosphate [15]. These insecticides are also used to control agricultural and urban pests, thereby exposing the mosquitoes to persistent selection pressure that eventually results in the selection for insecticide resistance.

Knockdown resistance (kdr) mutations involving the substitution of leucine by serine (L1014S) or phenylalanine (L1014F) at amino acid position 1014 is the main mechanism of resistance to pyrethroids in malaria vectors [16], although metabolic enzymes have also been known to play a role in resistance to pyrethroids [17, 18]. Earlier studies reported that the L1014F mutation was restricted to East Africa while the L1014S mutation was widely distributed in West and Central Africa [19]. However, recent studies have shown that both the L1014S and L1014F kdr mutations co-exist in East and West Africa [20, 21]. Conversely, resistance to carbasates and organophosphates results from overexpression of non-specific esterase enzymes and/or alteration of acetylcholinesterase (AChE) due to a single glycine to serine amino acid substitution at position 119 of the AChE gene [22, 23].

Defining the prevalence of insecticide resistance in malaria vectors and the underlying mechanism(s) of resistance in different ecological settings is necessary for the development of rational strategies for insecticide use and resistance management. The objective of this study was to investigate the resistance status of An. gambiae (s.s.) and elucidate the kdr mutation in coastal Kenya. Historically, An. gambiae (s.s.) and An. funestus were the main vectors of malaria in coastal Kenya, but the widespread use of LLINs has led to a major shift in malaria vectors in favor of An. arabiensis, an exophilic mosquito species that has had less contact with LLINs [24–28].

Recent reports in Kilifi county [25] and the neighboring Kwale county [26, 27] have revealed a low frequency of both the kdr allele and pyrethroid phenotypic resistance in this region. In addition, malaria is endemic in Kilifi county, with 91% of households having reported possessing at least one LLIN and 51% having universal ITN coverage achieving the proposed target of ≤2 people per LLIN [29]. The findings of this study will provide critical knowledge that may contribute to improved strategies for malaria control in Africa.

Methods
Study area
This study was conducted in Kilifi county, located along the north coast of Kenya. The county is situated north and northeast of Mombasa county, which also has a coastline on the Indian Ocean, with Taita Taveta county to the west, Tana River county to the east and Kwale county to the southwest. Kilifi county covers an approximate area of 12,246 km², and the population at risk of malaria and other mosquito-borne diseases is approximately 1.5 million [30]. The study sites have been described in detail elsewhere [24, 31]. Anopheles gambiae (s.l.) and An. funestus (s.l.) are the key malaria vectors in the area. They occur throughout the year, with peak occurrence during the rainy season [24]. Coastal Kenya usually experiences two rainy seasons each year: short rains falling from October to November and long rains from April to July. The mean annual rainfall ranges between 300 mm in the hinterland and 1300 mm in the coastal belt. The mean daily temperature varies from 21 °C to 30 °C [32].

Larval mosquito sampling was carried out in eight randomly selected study sites (Fig. 1), of which seven were in Kilifi subcounty, namely Ngombeni (3.73208°S, 39.76491°E), Mbogolo (3.69806°S, 39.81706°E), Jaribu (3.62054°S, 39.73544°E), Kidutani (3.88209°S, 39.71819°E), Shibe (3.55840°S, 39.77921°E), Mapawa (3.80452°S, 39.73641°E) and Mangororo (3.56366°S, 39.74507°E), and one, Burangi (3.09828°S, 40.04817°E), was in Malindi subcounty.

Selection of the study sites was based on high coverage with LLINs, and abundance and ease of accessibility to mosquito breeding sites.

Larval mosquito collection, transportation and rearing
Larval mosquito collections were done in June (during long rains), August (dry season) and November (short rains) 2013 and in July 2014 (immediately after the long rains). The collections were spread across the various seasons to maximize the collection of the different malaria vectors in the study sites. Larvae were sampled from stagnant water bodies selected in each study site using the standard dipping technique [33]. In each study site, six larval habitats were sampled once per week. The habitats included sand pits, ponds, roadside ditches, marshes, shallow wells and river banks. Anopheline larvae were transferred into Whirl-pak® bags (Thermo Fisher Scientific, Waltham, MA, USA) and transported to the insectary at Kenya Medical Research Institute (KEMRI), Center for Geographic Medicine Research, Coast Kilifi.

Tetramin® (Tetra Werke, Melle, Germany) baby fish food was used as the larval rearing diet. The insectary was maintained at a constant temperature of 25–27 °C and relative humidity of 74–82%. The larval pans were
monitored daily, and the collected pupae transferred into plastic cups in emergence cages. Newly emerged adults (F0) were identified using morphological characteristics [34]. The adults were then kept as same-age cohorts for insecticide susceptibility testing.

**World Health Organization susceptibility bioassay tests**

Non-blood-fed female *An. gambiae* (s.l.) aged between 3 and 5 days and reared from field-collected larvae were used in the susceptibility bioassays. Each insecticide susceptibility test was performed using 25 mosquitoes in four replicates and two controls. The insecticides used and their concentrations included deltamethrin (0.05%), permethrin (0.75%), dichlorodiphenyltrichloroethane (DDT, 4%), fenitrothion (1%) and bendiocarb (0.1%), prepared at Universiti Sains Malaysia [35]. The negative control included 25 mosquitoes collected from each study site and exposed to untreated filter papers. In addition, laboratory-raised, susceptible *An. gambiae* (s.s) Kisumu strain were exposed to insecticide-treated filter papers as a measure of quality assurance. The knockdown time (KDT) of each insecticide was recorded every 10 min for 1 h. Mosquitoes were then moved to a recovery tube and provided with 10% sucrose. Final mortality was recorded after 24 h.

After recording mortality at 24 h post-exposure, those mosquitoes still alive were killed by freezing and together with a 20% randomly selected subset of the dead samples per study site stored individually in labeled Eppendorf tubes (Eppendorf Co., Hamburg, Germany). The specimens were then preserved at − 80 °C for later molecular analysis.
DNA extraction and species identification
Genomic DNA was extracted from the whole body of the *An. gambiae* (s.l.) using an ethanol precipitation method [36]. Conventional PCR amplification of ribosomal DNA intergenic spacers was used to differentiate the sibling species of the *An. gambiae* complex [37].

Detection of kdr genotype mutations
Knockdown resistance was tested using 192 mosquitoes (59 alive and 133 dead) after performing the pyrethroid insecticide susceptibility tests; 20 *An. gambiae* (s.s.) Kisumu strain were used as controls. Both the L1014S (leucine to serine substitution) kdr allele originally detected in East Africa [16] and the L1014F (leucine to phenylalanine substitution) kdr allele from West Africa [19] were tested. Real-time PCR was used to determine the genotype at amino acid position 1014 of the voltage-gated sodium channel, following the methods of Bass et al. [38] as modified by Mathias et al. [39]. Each PCR reaction was conducted in a 10-µl volume containing 5.0 µl 2× TaqMan mix (TaqMan® Gene expression Master Mix [Thermo Fisher Scientific], 0.5 µl of forward primer, 0.5 µl reverse primer [for either kdr-east or kdr-west], 0.4 µl of each probe, 1.0 µl of template DNA and 2.6 µl of PCR grade water [ddH₂O]). Samples were genotyped for the wild-type (susceptible) allele using the probe 5′-GACACTGAAATTTCT-3′ and for the L1014S kdr allele using the probe 5′-ACGACTGAAATTTCC-3′. The primer sequences used for this study were 5′-CAT TTTTCTTGGCACCATTGTAATG-3′ (forward) and 5′-CGATCTTTGGCACCATTTGTTGA-3′ (reverse). Controls loaded in the last four wells of the 96-well PCR plate included FAM (fluorescent label)-positive control, HEX (fluorescent label)-negative control, non-template control (NTC) and buffer. The PCR was carried out in a Stratogene® MxPro 3000 Real-Time PCR system (Stratagene, Agilent Technologies, Inc., La Jolla, CA, USA) at the following temperature profile: an initial denaturation step at 95 °C, 10 min; followed by 92 °C (for kdr-east) or 95 °C (for kdr-west)/15 s, 60 °C/1 min, for 40 cycles; with a final extension step at 72 °C for 10 s.

Data analysis
Resistance was obtained using the World Health Organization criteria, with mortality rates of 98–100% indicating susceptibility, 90–97% suggesting possible resistance that needs further investigation and ≤90% indicating resistance, respectively [35]. KDT₅₀ and KDT₉₅ (time [in minutes] to knock down 50 and 95% of mosquitoes, respectively) were estimated using probit analysis [40]. A post-hoc comparison test with the Bonferroni correction test was performed to compare the proportions of each mosquito species among the study sites. The scatter-plots were retrieved to assess the kdr mutation and the frequency of the resistance allele determined from endpoint fluorescence using the MxPro software (Agilent Technologies, Santa Clara, CA, USA). Final data were captured in Microsoft Excel 2010 (Microsoft Corp., Redmond, WA, USA) and analyzed using the R statistical package, version 3.3.2 (R Foundation for Statistical Computing, Vienna, Austria).

Results

**Sample size and species composition**
A total of 3683 *An. gambiae* (s.l.) adult females were collected as larvae and used for the susceptibility bioassays from the eight villages: Jaribuni (500), Shibe (500), Kidutani (475), Ng’ombeni (490), Mapawa (475), Mboogolo (455), Burangi (500) and Mangororo (288). PCR analyses showed that *An. arabiensis* (91.49%) and *An. gambiae* (s.s.) (5.02%) were the only sibling species of *An. gambiae* (s.l.). No amplification occurred in the remaining 3.49% of samples (Table 1). Mosquitoes from Mangororo, Shibe and Mboogolo were predominantly *An. arabiensis* (Mangororo: 100%; Shibe: 96.8%; Mboogolo: 99.01%). A post hoc comparison test with Bonferroni correction revealed that the proportions of *An. gambiae* (s.s) and *An. arabiensis* species among the study sites were significant (Z = −3.31, P = 0.001). No *An. funestus* were collected in this study.

**Insecticide susceptibility bioassays**
*Anopheles gambiae* (s.s) Kisumu strain were susceptible to all of the insecticides tested (deltamethrin, permethrin, DDT, bendiocarb and fenitrothion) with 100% mortality. The negative control did not show any mortality; therefore, Abbott’s formula was inapplicable to correct for the natural causes of mortality in this study.

| Study site | n | *An. gambiae* (s.s) | *An. arabiensis* | Not amplified |
|------------|---|--------------------|------------------|--------------|
| Janbuni    | 172 | 12 (6.98%) | 148 (86.04%) | 12 (6.98%) |
| Kidutani   | 124 | 6 (4.84%) | 114 (91.94%) | 4 (0.00%) |
| Mboogolo   | 101 | 0 (0.00%) | 100 (99.01%) | 1 (0.99%) |
| Ng’ombeni  | 123 | 18 (14.63%) | 102 (82.93%) | 3 (2.44%) |
| Shibe      | 124 | 0 (0.00%) | 120 (96.77%) | 4 (3.23%) |
| Mapawa     | 108 | 5 (4.63%) | 95 (87.96%) | 8 (7.41%) |
| Burangi    | 163 | 8 (4.91%) | 153 (93.86%) | 2 (1.23%) |
| Mangororo  | 60  | 0 (0.00%) | 60 (100%) | 0 (0.00%) |
| Total      | 975 | 49 (5.02%) | 892 (91.49%) | 34 (3.49%) |

n, Number of *An. gambiae* (s.l.) processed for species identification.
Figure 2 shows the bioassay susceptibility status of *An. gambiae* (s.l.) to the different insecticides used in this study. Deltamethrin resistance in *An. gambiae* (s.l.) was only recorded in one of the eight study sites (Burangi: $n = 100$, mortality 45.5%). Coincidentally, permethrin resistance was also reported in the same study site ($n = 100$, mortality 48%). However, full susceptibility to both permethrin and deltamethrin in *An. gambiae* (s.l.) was indicated in all of the other study sites (mortality 100%). The *An. gambiae* (s.l.) populations tested indicated full susceptibility to DDT ($n = 750$, mortality range 99–100%) in the seven study sites tested.

Adult mosquitoes were resistant to fenitrothion in Jaribuni ($n = 100$, mortality 88%) and showed possible resistance at Ngombeni, Mapawa and Mbogolo with a test mortality range of 94–97%. However, mosquitoes from Burangi, Kidutani and Shibe indicated full susceptibility to fenitrothion. The mosquitoes tested were highly susceptible to bendiocarb in six of the eight study sites, i.e. Jaribuni, Shibe, Kidutani, Mangororo, Burangi and Mapawa, with test mortality of 100%. However, the possibility of resistance that requires further investigation was recorded against bendiocarb in Mbogolo ($n = 75$, mortality 93%) and Ng’ombeni ($n = 90$, mortality 97%).

**Knockdown time (KDT$_{50}$) at 95% CI and KDT$_{50}$ ratio**

Most of the knockdown resistance ratios (KDT$_{50}$ ratio) for the test population to that of the control [Kisumu strain] were within the *An. gambiae* (s.s.) Kisumu strain susceptible range of 1.0 to 1.7 and varied among study sites, as indicated in Table 2. The KDT$_{50}$ ratio for deltamethrin could not be determined in Burangi due to the high resistance levels. However, in the other study sites, the KDT$_{50}$ ratio was 1.1–1.9 fold compared to Kisumu strain. The mosquito population from Burangi also recorded resistance to permethrin with a KDT$_{50}$ ratio of 3.3. The resistance ratio indicated potential resistance to permethrin in Kidutani (KDT$_{50}$ ratio: 2.0) which requires further investigation. The KDT$_{50}$ ratio was 1.0–1.7-fold in the remaining six study sites: Jaribuni, Shibe, Ngombeni, Mapawa, Mbogolo and Mangororo.

The KDT$_{50}$ ratio for DDT, fenitrothion and bendiocarb compared to that for the Kisumu strain ranged from 1.1 to 2.2-fold in all the study sites. The mosquitoes were susceptible to bendiocarb in all the study sites (KDT$_{50}$ ratio 1.0–1.6). Possible resistance was recorded in Jaribuni for both DDT (KDT$_{50}$ ratio 2.2) and fenitrothion (KDT$_{50}$ ratio 2.1). The KDT$_{50}$ ratio for both DDT and fenitrothion was 2.2 in both Mbogolo and Mapawa, indicating possible resistance.
Table 2 Knockdown resistance ratio of mosquito populations (An. gambiae [s.l]) to different insecticides

| Insecticide        | Population | KDT50 (in min) | 95% CI  | KDT50 ratio |
|--------------------|------------|----------------|---------|-------------|
| Deltamethrin       | Jaribuni   | 18             | 40.9–55.2 | 1.1         |
|                    | Shibe      | 16             | 43.9–58.1 | 0.9         |
|                    | Kidutani   | 18             | 42.9–57.3 | 1.1         |
|                    | Ngombeni   | 23             | 41.9–56.2 | 1.4         |
|                    | Mapawa     | 32             | 40.8–61.1 | 1.9         |
|                    | Mbogolo    | 25             | 39.8–60.2 | 1.5         |
|                    | Burangi    | –              | –        | –           |
|                    | Mangororo  | 30             | 39.8–60.2 | 1.8         |
| Permethrin         | Jaribuni   | 18             | 39.9–60.2 | 1.0         |
|                    | Shibe      | 21.5           | 37.9–58.2 | 1.2         |
|                    | Kidutani   | 35.5           | 38.9–59.2 | 2.0         |
|                    | Ngombeni   | 30             | 40.8–61.1 | 1.7         |
|                    | Mapawa     | 20             | 39.8–60.2 | 1.1         |
|                    | Mbogolo    | 27             | 38.9–59.2 | 1.5         |
|                    | Burangi    | 60             | 37.9–58.2 | 3.3         |
|                    | Mangororo  | 30             | 39.8–60.2 | 1.7         |
| Dichlorodiphenyl-  | Jaribuni   | 30.5           | 40.9–55.2 | 2.1         |
| trichloroethane    | Shibe      | 15             | 37.9–58.2 | 1.0         |
| (DDT)              | Kidutani   | 18             | 39.8–60.2 | 1.2         |
|                    | Ngombeni   | 27.3           | 39.8–60.2 | 1.9         |
|                    | Mapawa     | 31.5           | 39.8–60.2 | 2.2         |
|                    | Mbogolo    | 25             | 39.8–60.2 | 1.7         |
|                    | Burangi    | 24.4           | 36.9–57.2 | 1.7         |
| Bendiocarb         | Jaribuni   | 19.2           | 43.9–58.1 | 1.0         |
|                    | Shibe      | 19.3           | 39.8–60.2 | 1.0         |
|                    | Kidutani   | 22             | 39.8–60.2 | 1.2         |
|                    | Ngombeni   | 26.9           | 39.8–60.2 | 1.4         |
|                    | Mapawa     | 30             | 39.8–60.2 | 1.6         |
|                    | Mbogolo    | 23             | 37.9–58.2 | 1.2         |
|                    | Burangi    | 24             | 37.9–58.2 | 1.3         |
|                    | Mangororo  | 30             | 40.8–61.1 | 1.6         |
| Fenitrothion       | Jaribuni   | 46             | 37.9–58.1 | 2.1         |
|                    | Shibe      | 23.5           | 39.8–60.9 | 1.1         |
|                    | Kidutani   | 40             | 42.8–63.1 | 1.9         |
|                    | Ngombeni   | 31             | 38.9–59.2 | 1.5         |
|                    | Mapawa     | 21.5           | 37.9–58.2 | 1.0         |
|                    | Mbogolo    | 46.3           | 37.9–58.2 | 2.2         |
|                    | Burangi    | 36             | 36.9–57.2 | 1.7         |

Cl, Confidence interval; KDT50, time taken for 50% of the test population to be knocked down; KDT50 ratio, KDT50 of the test population to that of the control (Kisumu strain)

*Lower limit estimation not possible due to a large g value [40]

Knockdown resistance status

A total of 192 randomly selected samples comprising of An. arabiensis, An. gambiae (s.s.) and Kisumu strain were genotyped for the kdr mutation. Neither L1014F nor L1014S kdr allelic genes were observed in An. arabiensis species. The kdr allelic frequency in An. gambiae, An. arabiensis and An. gambiae (s.s.) populations is shown in Table 3. Upon exposure to pyrethroids for 24 h, a total of 15 An. gambiae, An. arabiensis and An. gambiae (s.s.) were identified with the resistant phenotype, while 34 had the susceptible phenotype. L1014S allelic genes were identified in only four An. gambiae, An. arabiensis, of which two were homozygous (RR) and two were heterozygous (RS) for the trait. Only one of the 15 phenotypically resistant mosquitoes was observed to be genotypically resistant; this mosquito was heterozygous for L1014S characteristic. The remaining An. gambiae, An. arabiensis and An. gambiae (s.s.) mosquitoes genotyped were recessive (SS).

Discussion

This study has shown that Anopheles arabiensis is the most dominant species in the eight study sites sampled in Kilifi county. An earlier study reported that Anopheles gambiae (s.s.) was the dominant subspecies along the Kenyan coast and other regions of the country [24]. However, a shift in malaria vector composition has recently been documented that coincides with the scaling up of vector control interventions, especially the ongoing widespread use of insecticide-treated nets (ITNs) along the Kenyan coast [31]. The increase in populations of An. arabiensis and declining populations of An. gambiae (s.s.) may be a result of their contrasting ecological behavior. Anopheles gambiae (s.s.) has anthropophilic, endophagic and endophilic tendencies [41, 42]. This means relatively more hours spent indoors and, consequently, longer contact hours with LLINs, which may have contributed to its population decrease. In contrast, An. arabiensis is both exophilic and zoophilic [43–45], and this limits its contact with LLINs and the insecticides used for IRS [46]. The observed shift in species composition in favor of An. arabiensis calls for an integrated vector management
(IVM) approach targeting both indoor and outdoor control of malaria vectors.

Mosquito populations from one of the eight study sites (Burangi) were resistant to both deltamethrin and permethrin and had high median KDT compared with the other study sites. These findings suggest that resistant genes are localized and that vector control management strategies should be established to prevent their spread and preserve the efficacy of the current vector control tools. Earlier studies along the Kenyan coast and neighboring Tanzania reported the presence of pyrethroid resistance characterized by a high median KDT and mortality levels ranging from 62.38 to 93% following pyrethroid exposure [25, 28, 47]. In some West African countries, such as Ivory Coast and Burkina Faso, pyrethroid resistance has been found to be much more intense, with high KDT ratios and mortality rates of ≤ 40% reported following exposure to pyrethroids [48–52]. High KDT values in the test populations indicate the presence of the kdr mechanism of resistance [53]. The high resistance levels might be attributed to selection for insecticide resistance due to increased use of LLINs in vector control programs [35, 54, 55]. Based on KDT values, we noted potential resistance to DDT in Jaribuni and Mapawa villages that might partly be linked to cross-resistance from pyrethroids [56] or the presence of recessive genes in the mosquito population [25]. These two insecticide classes (pyrethroids and organochlorines) have the same mode of action and thus pose a threat of cross-resistance.

Mosquito samples from Jaribuni were resistant to fenitrothion based on both mortality rate and the KDT ratio. In addition, potential resistance to both fenitrothion and bendiocarb was recorded in Mbogolo and Ngombeni. However, susceptibility to both fenitrothion and bendiocarb was detected in all other study sites. This resistance may be due to selection pressure resulting from the contamination of the larval habitats with carbamates and organophosphates used in agriculture [57] as well as exposure of adult mosquitoes to insecticides during either sugar-feeding or outdoor resting. Agricultural systems provide ideal habitats for mosquito breeding [27]. Earlier studies conducted in Kenya reported the use of fenitrothion and bendiocarb in agricultural settings and linked them to resistance [58, 59]. Fenitrothion (organophosphate) and bendiocarb (carbamate) have been proposed for IRS use to control pyrethroid-resistant mosquitoes [60]. Therefore, the evolution of resistance to these insecticides threatens their use in IRS and as substitute to pyrethroids in public health systems.

The occurrence of phenotypic resistance to pyrethroids (deltamethrin and permethrin) may indicate the presence of target site insensitivity. Our findings that no kdr alleles were detected in An. arabiensis are consistent with results from previous studies in western and coastal Kenya [28, 39], suggesting other underlying resistance mechanisms as the cause of phenotypic resistance. However, the L1014S kdr allele was detected in An. gambiae (s.s.) with an allelic frequency of 3.33% in the resistant test population. The low allelic frequency of the L1014S kdr gene is consistent with results reported in other studies in the neighboring Kwale county [26, 28]. In contrast, other studies have indicated a high frequency and wide distribution of the L1014S mutation in An. arabiensis in western Kenya [61, 62]. These variations in the frequency of the resistance allele may be either due to movement of the mutant genes from their original selection pressure region or DNA deletion in a given genome in a mosquito population [39, 63, 64]. Although target site resistance had been detected earlier in the neighboring Kwale county along the coastal Kenya, in practice this may not mean that the mutant genes originated from there. However, it is evident that the percentage allelic frequency is clearly increasing. Therefore, further studies are needed to ascertain the origin of the mutant genes and facilitate development of strategies to mitigate their rapid spread.

The low kdr allele frequency compared to the high phenotypic resistance observed in pyrethroids is indicative of other underlying resistance mechanisms. Indeed, metabolic-based resistance most likely plays a bigger role in insecticide resistance than target site resistance [65]. Several studies have documented the contribution of metabolic resistance enzymes in malaria vectors to pyrethroids, organophosphates and carbamates [4, 26, 62, 63, 66–68]. Knockdown resistance is considered to be a weak form of resistance compared to metabolic-based resistance. Therefore, vector control failure is likely when kdr occurs along with metabolic resistance [69]. Nevertheless, the increasing kdr allele frequency coupled with metabolic-based resistance reported in the neighboring Kwale county [28] focuses attention on the risk of subsequent failure of current vector control interventions in coastal Kenya. Therefore, further investigation on the evolution of insecticide resistance in An. gambiae (s.l.) in Kilifi county is vital.

Conclusions
This study reported phenotypic resistance to selected pyrethroids and organophosphate insecticides in malaria vectors along the Kenyan coast. The occurrence of the L1014S allele in An. gambiae (s.s.) at a low frequency was also documented. These findings highlight the need for regular entomological surveillance and monitoring of insecticide resistance and for investment in new vector control strategies that can supplement or even replace the use of synthetic insecticides. This effort may inform development
and implementation of an IVM approach targeting all mosquito life stages with diverse vector control interventions to limit the spread of insecticide resistance and preserve the efficacy of existing insecticides.

**Abbreviations**

AchE: Acetylcholinesterase; ddH2O: PCR grade water; DDT: Dichlorodiphenyltrichloroethane; IRS: Indoor residual spraying; ITN: Insecticide-treated net; IVM: Integrated vector management; KDT: Knockdown time; kdr: Knockdown resistance; LLINs: Long-lasting insecticide-treated nets; NTC: Non-template control.

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**Authors’ contributions**

DNM, CMM, JMM and BK participated in the conception and design of the study. DNM carried out the field activities and bioassay tests. CMM, BK and JMM coordinated and supervised the study. EJM and DNM sequenced the samples. DNM analyzed the data and drafted the first manuscript. All authors critically revised and contributed to drafting the manuscript. All authors read and approved the final manuscript.

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**Availability of data and materials**

All data generated and supporting the conclusions of this manuscript have been included in the article. Raw data and materials are available from the corresponding author upon request.

**Declarations**

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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