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Insecticide resistance in *Anopheles arabiensis* (Diptera: Culicidae) from villages in central, northern and south west Ethiopia and detection of *kdr* mutation

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Abstract

Background: Anopheles arabiensis is the major vector of malaria in Ethiopia. Malaria vector control in Ethiopia is based on selective indoor residual spraying using DDT, distribution of long lasting insecticide treated nets and environmental management of larval breeding habitats. DDT and pyrethroid insecticides are neurotoxins and have a similar mode of action on the sodium ion channel of insects. It was therefore necessary to verify the insecticide susceptibility status of *An. arabiensis*, to better understand the status of cross-resistance between DDT and the pyrethroids in this species as well as to detect a resistant gene.

Methods: Standard WHO insecticide susceptibility tests were conducted on adults reared from larval and pupal collections from breeding sites at three villages namely: Sodere in the Rift Valley, Gorgora in the north and Ghibe River Valley in the south west of Ethiopia. The occurrence of cross-resistance between pyrethroids and DDT was determined using a DDT selected laboratory colony originally collected from Gorgora. Phenotypically characterized mosquitoes were tested for the presence of knockdown resistance (*kdr*) alleles using the standard polymerase chain reaction assay.

Results: All *An. gambiae* s.l. specimens assayed by PCR were identified as *An. arabiensis*. The knockdown and mortality results showed *An. arabiensis* resistance to DDT in all villages, resistance to deltamethrin and permethrin in the Ghibe River Valley and permethrin resistance in Gorgora. Bioassay susceptibility tests also indicated the presence of cross-resistance between DDT and permethrin, but not between DDT and deltamethrin. The knockdown resistance (*kdr*) mutation of leucine to phenylalanine in the sodium ion channel gene was detected in populations from Gorgora and the Ghibe River Valley.

Conclusion: Since *An. arabiensis* shows high levels of resistance to DDT in all villages tested and varying pyrethroid resistance in Gorgora and the Ghibe River valley, precautionary measures should be taken in future vector control operations. Moreover, the status of resistance in other locations in Ethiopia and the spread of resistant gene (s) should be investigated.

Background

Malaria vector control in Ethiopia is targeted mainly against *Anopheles arabiensis* as this species is responsible for transmitting *Plasmodium falciparum* and *P. vivax* [1]. Vector control measures including selective indoor residual spraying of dichlorodiphenyltrichloroethane (DDT), distribution of long lasting insecticide treated mosquito nets (LLINs) and source reduction of larval habitats are currently implemented by the Federal Ministry of Health in collaboration with international and non-governmental organizations. The history of utilization of DDT in the country dates back to the mid 1950s with small scale trials followed by wide and extensive application during the malaria eradication period. Thus, DDT has been in use



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for more than four decades. The emergence of DDT resistance in *An. arabiensis* has been reported from different localities in Ethiopia [2,3] and the highest levels of resistance were recorded from ArbaMinch in the South and Gambella in the West [2]. DDT resistance has also been demonstrated in populations of *An. pharoensis* in two localities [4,5], one of three secondary malaria vectors of Ethiopia (the other two being *An. funestus* and *An. nili*)[1].

DDT and pyrethroid insecticides are insect neurotoxins that interfere with ion flow regulation across the sodium ion channels. Ion channel modification via an amino-acid substitution leads to reduced target site sensitivity known as knockdown resistance (*kdr*). The substitution of leucine at position 1014 for either a phenylalanine or a serine has been reported in *An. gambiae* s.s. from West and East Africa, respectively [6,7]. These mutations have also been reported in *An. arabiensis* [8,9].

In view of the prolonged utilization of DDT for vector control, the application of different agricultural pesticides in agrodevelopment areas and the planned up-scaling of pyrethroid treated LLINs distributionace in the areasurrounding areas is found in Metehara. ArbaMinch is Metehara is one of the, it was necessary to evaluate the insecticide susceptibility status and genetic basis of insecticide resistance and cross-resistance in *An. arabiensis* from three selected localities in Ethiopia.

Methods

Study localities

The study was conducted at three localities namely Ghibe River Valley in the west (hereafter referred to as Ghibe), Sodere hot springs in the Rift Valley and Gorgora in the North West where DDT spraying has been practiced for a long time. Ghibe (8°14'N, 37°35'E, altitude 1,040-1,080 m) is situated 185 kms south-west of Addis Ababa. It has a long history of state owned farming in which vegetables, maize and citrus fruits are produced. The International Livestock Research Institute (ILRI) used to conduct research studies on cattle trypanosomiasis and Glossina in the same locality as well as in Tolay (Upper Ghibe River Valley). To control tsetse flies, the ILRI introduced deltamethrin (pyrethroid) treated cattle as well as targets impregnated with the same insecticide [10]. Currently, the Institute provides technical support to local farmers at both study sites for the application of cyfluthrin (pyrethroid) on cattle (Dr. Woudyalew Mulat, personal communication). Sodere hot springs (8° 24'N and 39° 23'E, altitude 1,360 m) is located in the middle course of the Rift Valley about 125 kms east of Addis Ababa. Gorgora (12° 12'N and 37° 16' E, altitude 1,800 m) is located at the northern tip of Lake Tana, about 800 kms north of Addis Ababa. Three private agricultural developments are underway near Gorgora.

Insecticide susceptibility tests

Insecticide susceptibility tests were carried out at different times between November 2005 and April 2007. Mosquito larvae and pupae were collected from various breeding sites and reared to adults. Two to three days old, non blood fed adult females morphologically identified as *An. gambiae* s. l were exposed to discriminating dosages of 4% DDT, 0.05% deltamethrin and 0.75% permethrin using WHO papers (obtained from Vector Control Research Unit, School of Biological Sciences, Universiti Sains Malaysia) and bioassay tubes for 1 hour [11]. Number of mosquitoes knocked down during exposure was recorded at intervals of 5 minutes and the proportions of survivors and dead mosquitoes were recorded 24 hours post exposure. The same number of mosquitoes was exposed to insecticide free papers as controls.

Samples of dead and surviving mosquitoes were preserved in 95% ethanol and kept in a freezer (-21°C) for subsequent molecular species identification and *kdr* analysis. The quality of impregnated papers was checked on a laboratory colony of *An. arabiensis* which is susceptible (100% mortality) to DDT, permethrin and deltamethrin. The colony is maintained at the Aklilu Lemma Institute of Pathobiology, Addis Ababa University, since September 2001 from original collection of mosquitoes from Debrezeit, 45 kilometers east of Addis Ababa.

Species identification

A polymerase chain reaction (PCR) assay was carried out to differentiate *An. arabiensis* from its *An. gambiae* complex sibling species according to the method of Scott *et al.* [12].

Knockdown resistance gene detection

For detection of *kdr* mutations, genomic DNA was isolated from 70 mosquitoes according to the method of Collins *et al.* [13]. Two separate PCR reactions were run, one to detect alleles of the leucine-phenylalanine substitution, the other, to detect wild-type susceptible alleles following the methods described in Matambo *et al.* [14] and Abdalla *et al.* [15]. The occurrence of *kdr* was confirmed by direct sequencing of the 293 base pair fragment of the sodium channel gene amplified using Agd1 and Agd2 primers [6]. Amplicons were submitted to Inqaba Biotec Company, Pretoria, South Africa for sequencing in both directions.

Selection for resistance, colony maintenance and cross resistance evaluation

An. arabiensis strain originating from a collection of wild females from Gorgora was selected for resistance to DDT. Selections were carried out in the insectary at the Aklilu Lemma Institute of Pathobiology by exposing male and female adults over generation to 4% DDT impregnated papers. Full resistance to DDT (zero mortality) was

achieved in the 6th generation. Subsequently insecticide susceptibility tests using 0.75% permethrin and 0.5% deltamethrin impregnated papers were carried out on the 8th generation according to the standard WHO method [11]. Tests included 5 replicates of 20 unfed, 2-3 day old adult females per replicate. Controls included exposure of females from the same cohort to insecticide free papers.

Data analysis

Percentage mortalities 24 h post exposure was used to assess the status of susceptibility/resistance to insecticides. The KT_{50} and KT_{90} (time to knockdown 50% and 90% mosquitoes, respectively) values were calculated for each insecticide using probit analysis in SPSS 15.0 for windows. Fisher's exact test was applied to observe associations between genotypes and phenotypes. Chi-square statistics was used to analyze allelic and genotype frequencies.

Results

Mosquito species identification

300 *An. gambiae* s.l. samples were assayed by PCR. All were identified as *An. arabiensis*.

Insecticide susceptibility test results

Exposure to DDT induced significantly reduced mortality in *An. arabiensis* from all study localities, implicating resistance according to established criteria [11]. The highest resistance was recorded from Ghibe followed by Gorgora and Sodere (Table 1). Mortality due to permethrin (85%) and deltamethrin (80.6%) exposure in samples from Gorgora is strongly suggestive of pyrethroid resistance. The KT_{50} and KT_{90} values are also elevated. *Anopheles arabiensis* resistance to pyrethroids has only progressed to a high level in Ghibe.

kdr gene detection

Thirty two *An. arabiensis* specimens from Gorgora and 38 from Ghibe were examined for the occurrence of 1014 F *kdr* mutation [6]. The frequency of genotypes of tested mosquitoes on each insecticide is separately depicted in Table 2. The allelic and genotypic frequencies are in Hardy-Weinberg equilibrium (Gorgora $X^2 = 0.02$, p = 0.99; Ghibe $X^2 = 2.62$, p > 0.27) (Table 3). The genotype frequency of *An. arabiensis* from Gorgora is statistically different from the population of Ghibe (Fisher's exact test, p < 0.05).

The occurrence of *kdr* mutations were further confirmed by sequencing the 293 base-pair fragment of the sodium channel gene of 14 randomly chosen *An. arabiensis* from Gorgora and Ghibe. The sequences revealed similar results (with only one exception from Ghibe) with that of PCR confirming the presence of 1014F *kdr* gene (Table 3).

Cross-resistance tests

Cross-resistance tests on F_8 DDT resistant *An. arabiensis* from Gorgora against permethrin revealed lower susceptibility with 52% mortality (Table 4). A similar proportion was knocked down at the end of 80 minutes. In contrast, deltamethrin caused 98% mortality. The KT₅₀ was 29.3 minutes which was slightly higher than the KT₅₀ of the non-selected vector population (25.3 minutes).

Discussion

The insecticide susceptibility data indicate high levels of resistance to DDT in *An. arabiensis* at all localities, although varying level of mortality and knockdown were observed. The highest level of resistance as shown by the survival rates to DDT was recorded in Ghibe (96%) followed by Gorgora (50.5%) and Sodere (21.2%). Resistance

Locality	Insecticide	Number tested	Percent mortality	KT50 in minutes (95% confidence interval)	KT90 in minutes (95% confidence interval)
Gorgora	DDT	198	49.5	*	*
	Permethrin	140	85.0	19.6 (17.5-21.6)	28.7(26.2-32.5)
	Deltamethrin	160	80.6	25.3 (23.2-27.5)	35.9(33.1-40.0)
Ghibe	DDT	160	3.8	**	**
	Permethrin	60	31.7	***	
	Deltamethrin	160	31.3	37.6 (34.7-40.5)	56.6(52.8-61.7)
Sodere	DDT	118	78.8	****	****
	Permethrin	80	100	17.8 (15.9-19.7)	25.8(23.4-25.9)
	Deltamethrin	120	99.2	21.9 (19.9-23.9)	31.0(28.6-34.8)

*60% were knockeddown at the end of 80 m, ** only 4 mosquitoes were knocked down at the end of 80 minutes, ***41.7% were knockeddown at the end of 80 m**** 77.1% were knockeddown at the end of 80 m.

Locality	Insecticide	Phenotypes	Number	Genotype (%)		
				SS	RS	RR
Gorgora	DDT	Susceptible	2	2 (100)	0	0
		Resistant	12	1 (8.3)	8 (66.7)	3 (25.0)
	Permethrin	Susceptible	7	5 (71.4)	1 (14.3)	1 (14.3)
		Resistant	11	4 (36.4)	7 (63.6)	0
Ghibe	DDT	Susceptible	5	0	4 (80.0)	1 (20.0)
		Resistant	9	0	6 (66.7)	3 (33.3)
	Permethrin	Susceptible	7	0	3 (42.9)	4 (57.1)
		Resistant	3	0	0	3 (100)
	Deltamethrin	Susceptible	6	0	2 (33.3)	4 (66.7)
		Resistant	8	0	2 (25.0)	6 (75.0)

Table 2: Genotypic frequency of kdr from bioassayed An. arabiensis from Gorgora and Ghibe, Ethiopia

SS = homozygote susceptible, RS = heterozygote resistant, = RR homozygous resistant

to DDT is higher than previously reported in *An. arabiensis* from Gambela (76%) in the West and ArbaMinch (60%) in the South [2]. The results obtained from Gorgora are comparable to those of the eastern localities (Metehara and MelkaWorer) where 45-50% resistance was reported [3]. Ameneshewa [16] noted 30% resistance from Gergedi which is close to Sodere.

Anopheles arabiensis from Ghibe shows high resistance (about 70%) to permethrin and deltamethrin as well. The Gorgora population also shows signs of resistance to pyrethroids (15%-20%) based on final mortality data. The time to 50% knockdown at Gorgora was higher (KT_{50} = 19.6 minutes for permethrin and 25.3 minutes for deltamethrin) than that of a susceptible laboratory colony (KT₅₀s of 14.6 and 18.3 minutes for permethrin and deltamethrin, respectively: data not shown) as well as of a susceptible reference strain of An. gambiae [17,18]. The pyrethroid resistance levels in Gorgora at the present are unlikely to pose an epidemiological threat but routine periodic monitoring to determine levels of resistance are needed. Anopheles arabiensis from Sodere are susceptible to the two pyrethroids and yet, their knockdown times are similar to those recorded in Gorgora. In neighboring Sudan, An. arabiensis showed comparable levels of resistance to permethrin (final mortalities varying between 10-55% in Gezira and 6-22% in Sennar) [15]. However, populations of this species proved susceptible to pyrethroids at three localities in the eastern part of the country [9]. Variable levels of resistance to DDT and pyrethroids in *An. arabiensis* populations in neighboring countries raises concerns because of unrestricted gene flow between populations through migration.

The West African kdr mutation was detected in An. arabiensis from Ghibe and Gorgora. Resistance to pyrethroids and DDT in An. gambiae associates closely with kdr [6,7,19,20]. However, a similar correlation of kdr with the phenotypic expression of resistance could neither be established in a DDT-selected An. arabiensis laboratory strain from Sudan [14] nor in natural populations [9,15]. Matambo et al. [14] and Abdalla et al. [15] observed discrepancies in their PCR data compared to sequenced samples. The findings from Sudan samples also showed that *kdr* is found in phenotypically insecticide susceptible mosquitoes, not being restricted to resistant individuals alone. The data from Gorgora and Ghibe show the same phenomenon whereby kdr homozygotes were recorded in samples characterized as phenotypically susceptible to insecticides. Brooke [21] argues that kdr might not be the

Table 3: Comparison of genotype determination between PCR and sequencing of Sodium channel gene of *An. arabiensis* from Gorgora and Ghibe River Valley, Ethiopia.

Locality	Number sequenced	PCR results	Sequence results
Gorgora	7	3 RS	Same as PCR
		4 SS	
Ghibe	7	4 RR	5 RR
		3 RS	2 RS

Insecticide	Number tested	% mortality	KT ₅₀ (minutes)	% mortality in base	KT ₅₀ (minutes) in base
Permethrin	100	52.0*	No. KD	85.0	19.6 (17.5-21.6)
Deltamethrin	100	98.0	29.3 (27.0-31.5)	80.6	25.3 (23.2-27.5)

Table 4: Bioassay test results of base and DDT selected F₈ An. arabiensis from Gorgora.

No. KD = only 56% were knocked down * Control mortality was less than 5%.

only mechanism that confers resistance to DDT and pyrethroids and suggests the potential involvement of enzyme detoxification. Target site insensitivity coupled with enzyme detoxification has been described in *An. gambiae* and *Culex quinquefasciatus* from Benin [22].

The insecticide resistance in An. arabiensis from Ghibe and Gorgora and reduced level from Sodere may be attributable to differential insecticide selection pressure. Applications of insecticide against agricultural pests and disease vectors including anophelines and tsetse flies are practiced intensively in Ghibe and, to a lesser extent, in Gorgora. Documented information is lacking on the types of insecticides in use at these agricultural localities except that deltamethrin and cyflumethrin pour-ons are used on cattle in Ghibe [10]. In Gorgora, different types of insecticides have been in use for a number of years to control agricultural pests (Ato Teklu, personal communication), although it was difficult to establish whether DDT has been used. Use of agricultural pesticides in Ghibe may account for the increased resistance of An. arabiensis to deltamethrin and permethrin. Pyrethroids have not been used for malaria vector control in these areas until the recent distribution of LLINs which started in 2005. Agricultural pesticides may exert selective pressure on malaria vectors by leaching into breeding habitats so that the immature stages are continuously exposed. In West Africa, DDT application on farms during the early WHO malaria vector control era and extensive use of pyrethroids on cotton farms has been attributed to increased resistance of An. gambiae to pyrethroids as well as increased frequency of the kdr genotype in many countries [20,22,23]. In urban areas, insecticides for households and the expansion of urban agriculture are suggested to cause resistance in malaria vectors in West Africa [24].

Cross-resistance between DDT and permethrin was shown in a DDT-resistant laboratory colony of *An. arabiensis* originating from Gorgora possibly caused by *kdr*. By contrast, cross resistance between DDT and deltamethrin was not recorded in *An. arabiensis* samples from Sudan [14]. In this case the influence of *kdr* might be less important because the test population carrying the *kdr* allele showed a high level of susceptibility to deltamethrin, suggesting the existence of an independent DDT resistance mechanism. The absence of cross-resistance between DDT and deltamethrin in Gorgora DDT-selected *An. arabiensis* is encouraging in terms of LLINs utilization, although the moderate deltamethrin resistance recorded requires careful monitoring and consideration.

The potential impact of resistance on LNs use and indoor spraying with DDT in Ghibe is unknown and needs to be established. Permethrin resistance in *An. arabiensis* from Sodere is lower than in Ghibe, and regular monitoring should identify any changes in the level of resistance in association with the continued use of LLINs. In Kenya, lower susceptibility of *An. gambiae* to permethrin was indicated in villages where permethrin impregnated mosquito nets were implemented as compared to villages without nets in a period of one year and the reduction was linked to increased detoxification caused by elevated cytochrome P450 monooxygenase activity together with kdr-w[25]. Similarly, another study by Stump *et al.* [26] confirmed an increased frequency of *kdr* in villages with nets.

In conclusion, since the country relies on DDT for indoor residual sprays, the development of resistant populations of *An. arabiensis* implies careful consideration and monitoring of ongoing vector control program. There is also a need to monitor pyrethroid susceptibility in wider areas covering a number of vector populations.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

MB conducted the field and laboratory study and drafted the manuscript, IE and MI involved in project design and conducted the study, LLK and BDB facilitated the molecular laboratory study and commented on the manuscript, HE contributed to project design, facilitated the supply of WHO bioassay materials and reviewed the draft manuscript, AA and TGM facilitated the laboratory and field study and participated in drafting the manuscript. All co-authors have read the manuscript.

Acknowledgements

The technical assistance of staff of ALIPB, IEND and VCRU is greatly acknowledged.

This study obtained financial assistance from the MIM/TDR research project (ID No. TDA 40048A) which aimed to study the epidemiology of malaria in Ethiopia and Sudan and was carried out by AHRI, IEND and ALIPB. AHRI is also acknowledged for facilitating field work and some international travel expenses.

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Received: 29 January 2010 Accepted: 25 April 2010 Published: 25 April 2010

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doi: 10.1186/1756-3305-3-40

Cite this article as: Balkew *et al.*, Insecticide resistance in Anopheles arabiensis (Diptera: Culicidae) from villages in central, northern and south west Ethiopia and detection of kdr mutation *Parasites & Vectors* 2010, **3**:40

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