Diversity, resistance and vector competence of endophilic anophelines from southern Ghana

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Abstract

Background. As part of efforts to monitor the impact of vector control strategies so that they can be improved and more targeted, we collected baseline data on aspects of the bionomics of endophilic anophelines in southern Ghana.

Materials and Methods. Indoor resting anophelines were collected using mouth aspirators and pyrethroid spray catch. Anopheles females were identified to species level using morphological characteristics and sibling species were distinguished by PCR. The presence of the L1014F mutation, conferring resistance to insecticides, was determined in An. gambiae s.s. and An. coluzzii samples using TaqMan real-time PCR. Host blood meal sources were determined by PCR, and the presence of Plasmodium falciparum circumsporozoite proteins determined by ELISA.

Results. A total of 892 female Anopheles (31% An. gambiae, 41% An. coluzzii and 28% An. funestus) were collected from six villages. The L1014F mutation was almost fixed in all populations studied (allele frequencies: 0.87-1.00). Both An. gambiae s.l. and An. funestus fed mainly on humans, with a human blood index of 1, although some animal feeding was recorded in An. gambiae. P. falciparum was detected in all ecological zones and in all three major vector species, being 4.9% in An. funestus, 3.8% in An. gambiae s.s. and 1.1% in An. coluzzii.

Conclusions. These findings suggest that the three major vectors of malaria are present in all ecological zones of southern Ghana and contribute to disease transmission. The near fixation of the L1014F mutation in southern Ghana poses a great threat to vector control, thus highlighting the urgent need to implement measures to maintain the efficacy of current control tools and to develop novel control strategies.

1 Introduction

In Ghana, four dominant malaria vector species are responsible for malaria transmission: Anopheles funestus s.s., and three members of the An. gambiae complex: An. gambiae s.s., An. arabiensis and An. coluzzii [1]. An. gambiae s.l. and members of the An. funestus group are sympatric over much of their range in Africa [2]. Studies in southern Ghana have found that An. gambiae s.s., An. coluzzii and An. funestus are the dominant malaria vector species [3-6]. These species are sympatric across much of Ghana but An. gambiae s.s. predominates in the middle belt, characterised by forest type vegetation, whereas An. coluzzii predominates in the northern and coastal savannah regions [7,8]. However, this information alone is too general and cannot be used to extrapolate effective vector control strategies against these species under the diverse ecological conditions experienced in Ghana. It has been established that occurrence, population density and vector competence of An. gambiae s.l. and An. funestus vary geographically and are influenced by a number of environmental conditions, such as climate, rainfall and vegetation, as well as activities which influence their breeding [2,3,7,9,10]. It is therefore essential to understand the distribution and population dynamics of these anophelines in the various ecological settings before implementing appropriate malaria vector control interventions.

In Ghana, the national vector control programme, spearheaded by the Ghana National Malaria Control Program (NMCP), among other malaria control initiatives, prioritises use of insecticide-treated bednets (ITNs) and indoor residual spraying (IRS) as the main malaria vector control tools [1]. These strategies rely heavily on few insecticides recommended by WHO [11]. Unfortunately, there is strong evidence that mosquito populations in Ghana have developed resistance to insecticides used in malaria vector control [4-8,12]. This development calls for a shift towards integrated vector management (IVM). There
are several approaches available, including, but not limited to: insecticide-treated wall linings (ITWL), entomopathogenic fungi, larviciding, and the development and implementation of novel strategies to counter this problem. In addition, insecticide resistance management needs to be implemented to prolong effectiveness of the insecticides currently available for malaria vector control [1]. It is against this background that the studies reported in this article focused on understanding the bionomics and susceptibility levels of vector populations from southern Ghana as an essential step towards the development and implementation of effective IVM with a strong insecticide resistance management component.

2 Materials and methods

2.1 Study area

The study was carried out in six villages in southern Ghana, covering three ecological zones: forest ecological zone (FEZ), forest transition ecological zone (FTEZ) and coastal savannah ecological zone (CSEZ) (Figure 1).

In the FEZ, mosquitoes were collected from three villages: Atatam (06° 17.377’N, 001° 27.545’W), Agyenkwaso (06° 15.547’N, 001° 26.689’W) and Kusa Dinkyeae (06° 18.894’N, 001° 29.622’W), all located in the Adansi North District of the Ashanti Region (Figure 1). Atatam lies 257 m above sea level, is surrounded by mountains, from which several streams reach the village, and has 1,220 inhabitants. Agyenkwaso lies 223 m above sea level, with a population of about 1,300, while Kusa Dinkyeae has only 800 inhabitants and lies 268 m above sea level. Atatam and Agyenkwaso are proximal to each other while Kusa Dinhyeae is more isolated in the forest. A mountain separates Atatam and Kusa Dinkyeae. The main economic activity in these three villages is cocoa farming, interspersed with coffee, oil palm and subsistent farming. All three villages have been targeted by expansion of the Obuasi malaria control programme run by the AngloGold Ashanti Gold Mine because of their proximity to Obuasi municipality.

From the CSEZ, two villages, Gomoa Okyereko (05° 24.828’N, 000° 36.284’W) and Gomoa Adawukwa (05° 24.960’N 000° 36.729’), were sampled. Okyereko and Adawukwa are located in the Gomoa East District of the Central Region of Ghana, with an elevation of 19 and 22 meters above sea level, respectively. Okyereko is an irrigated rice farming village with about 1,500 inhabitants. Water from a dam about 500 m away from the village flows through canals to irrigate the rice fields, which are less than 100 m from the village. Flooded rice fields provide active breeding sites for different species of mosquitoes, including anophelines. Cement block houses with corrugated iron sheet roofing dominate, and eaves are either open or closed. Adawukwa is located about 500 m from Okyereko, and the two villages are separated by a river. About 500 m upstream the river, is a swamp where sugarcane cultivation, the main occupation of the inhabitants, takes place. There are about 1,200 people in the village, who, besides sugarcane farming, are involved in maize farming and sugarcane alcohol distillation. There is a brick and roofing tile factory in the village and, as a result, most of the houses, which are either made of brick or cement blocks, are roofed with brick tiles and often have open eaves. Apart from a few households with ITNs and the use by individuals of mosquito-repelling coils, there is no active malaria control programme running in any of the six villages. Domestic animals, such as goat, sheep, chicken, dogs, cats and pigs, are present in all villages. Cattle were not seen in any of the villages sampled.

The sixth village where mosquito sampling was carried out, Osorongma (05° 52.757’N, 000° 06.563’W), lies 94 m above sea level and is located in the FTEZ. It is a peri-urban village close to Dodowa in the Dangme West District of the Greater Accra Region of Ghana, with about 200 inhabitants, who are mainly farming cassava and mango. Houses in the village are a mix of cement block and mud houses, all with corrugated iron sheet roofing, usually with open eaves.

Figure 1. Map of Ghana showing the geographical location of study villages. 1: Osorongma, 2: Okyereko, 3: Adawukwa, 4: Kusa Dinkyeae, 5: Atatam, 6: Agyenkwaso.
Table 1. Indoor resting mosquito densities in six villages in southern Ghana.

<table>
<thead>
<tr>
<th>Village</th>
<th>Mean number of mosquitoes/room ± S.E.</th>
<th>An. Funestus group</th>
<th>Culex spp.</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atatam</td>
<td>7.20 ± 1.59b</td>
<td>7.45 ± 1.46c</td>
<td>0.00 ± 0.00i</td>
<td>14.65 ± 1.90*</td>
</tr>
<tr>
<td>Agyenkwaso</td>
<td>4.36 ± 1.15b</td>
<td>5.18 ± 1.33a</td>
<td>0.27 ± 0.14v</td>
<td>9.81 ± 1.08b</td>
</tr>
<tr>
<td>Kusa Dinkyae</td>
<td>8.63 ± 2.82a</td>
<td>1.50 ± 0.76b</td>
<td>0.00 ± 0.00i</td>
<td>10.13 ± 2.57*</td>
</tr>
<tr>
<td>Osorongma</td>
<td>10.00 ± 2.49a</td>
<td>0.00 ± 0.00b</td>
<td>0.00 ± 0.00b</td>
<td>10.00 ± 2.49a</td>
</tr>
<tr>
<td>Adawukwa</td>
<td>14.75 ± 2.60a</td>
<td>4.38 ± 2.78b</td>
<td>0.38 ± 0.38</td>
<td>19.15 ± 1.15*</td>
</tr>
<tr>
<td>Okyereko</td>
<td>12.13 ± 2.19a</td>
<td>0.13 ± 0.13b</td>
<td>0.33 ± 0.33</td>
<td>12.39 ± 2.13*</td>
</tr>
<tr>
<td>Total</td>
<td>9.18 ± 0.91b</td>
<td>3.54 ± 0.65c</td>
<td>0.11 ± 0.05</td>
<td>12.83 ± 0.90*</td>
</tr>
</tbody>
</table>

Means in the same row with different letters are significantly different at P<0.05. S.E.: standard error of mean.

2.2 Mosquito collections and field processing

Sampling was carried out between September 2011 and January 2012, spanning the minor rainy season (September - November) and the dry season (December - March). Prior to mosquito sampling, permission to gain access to each village was sought from an opinion leader, such as the chief or assemblyman. In all cases, verbal informed consent was obtained from the head of each household before mosquito collections were carried out. Mosquitoes were collected early in the morning between 6 and 10 am inside houses, using standard indoor resting collection methods (mouth aspirators and pyrethroid spray collection). Immediately after collection, mosquitoes were killed in a killing bottle containing ethyl ether, individually placed in 1.5 ml microcentrifuge tubes containing silica gel and labelled. All specimens were transported to the laboratory where they were morphologically identified and sorted into species groups using the keys by Gillies and Coetzee [13], and stored at −20°C for further processing.

2.3 Laboratory analysis

A sub-sample of 30 or all (where less than 30 mosquitoes were collected) for each round of collection from every village was selected randomly for molecular assays. The head and thorax of each specimen was dissected and placed in labelled microcentrifuge tubes for enzyme-linked immunosorbent assay (ELISA) to determine the presence of *P. falciparum* sporozoites. DNA from the rest of the specimen (legs, wings and abdomen) was extracted using standard procedures described by Collins et al. [14]. The extracted DNA was used for species identification, kdr mutation determination and blood meal analysis by polymerase chain reaction (PCR).

Specimens identified morphologically as *An. gambiae s.l.* were identified to species level using the PCR assay described by Scott et al. [15]. Those specimens identified as *An. gambiae s.l.* were further identified as *An. gambiae s.s.* or *An. coluzzii* using the method described by Favia et al. [16]. Specimens identified morphologically as *An. funestus* were identified to species-specific level using the PCR assays of Koekemoer et al. [17], and those that failed to amplify after a single repeat were subjected to the IGS-based real-time PCR protocol for identification of *An. funestus* subgroup developed by Vezenegho et al. [18].

The West African knockdown resistance mutation (L1014F) was assayed in samples identified as *An. gambiae s.s.* and *An. coluzzii* using the TaqMan real time PCR (RT-PCR) assay described by Bass et al. [19]. The presence of *P. falciparum* parasites was determined in all specimens identified to the species-specific level using the ELISA protocol of Wirtz et al. [20]. To determine the blood meal source in blood-fed mosquitoes, the multiplex PCR method developed by Kent and Norris [21] was used. The assay was used to test for the presence of human, cow, pig, goat or dog blood in the specimens. Any specimens that gave multiple blood meals were repeated for confirmation. To confirm blood meal type, PCR products for each blood type were sequenced and checked for alignment with the respective host blood sequence.

2.4 Data analysis

Mosquito species density was calculated as the number of each species divided by the total number of mosquitoes collected per room per collection date. Knockdown resistance (*kdr*) data were analysed using a Chi-square test for goodness of fit, from which the allele frequencies were generated. Human blood index (HBI) was calculated as number of specimen positive for human blood divided by the total number of specimens tested. Sporozoite rate was calculated as number of specimen positive for *P. falciparum* divided by total number tested.

Data for indoor resting density and *kdr* allele frequencies for each species were compared across villages using ANOVA and Tukey’s test used for means separation. Student’s t-test was used to compare percentage prevalence of *An. gambiae s.s.* and *An. coluzzii* in each village. All statistical analyses were performed using SPSS version 16, except for Chi-square tests, which were performed online using the Hardy-Weinberg equilibrium calculator [22].

2.5 Ethical clearance

This study was exempt from ethical approval by the Institutional Review Board of the Noguchi Memorial Institute for Medical Research, University of Ghana on the basis that it did not involve human participants.
Table 2. Relative prevalence (%) of *An. gambiae* s.s. and *An. coluzzii* in six villages in southern Ghana.

<table>
<thead>
<tr>
<th>Village</th>
<th>N</th>
<th><em>An. coluzzii</em></th>
<th><em>An. gambiae</em> s.s.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adawukwa</td>
<td>47</td>
<td>1.96 ± 1.96(^a)</td>
<td>98.04 ± 1.96(^c)</td>
</tr>
<tr>
<td>Agyenkwaso</td>
<td>22</td>
<td>23.61 ± 1.39(^a)</td>
<td>76.39 ± 1.39(^b)</td>
</tr>
<tr>
<td>Kusa Dinkyeae</td>
<td>29</td>
<td>0.00(^a)</td>
<td>100.00(^b)</td>
</tr>
<tr>
<td>Osorongma</td>
<td>43</td>
<td>6.00 ± 3.24(^a)</td>
<td>94.00 ± 3.24(^a)</td>
</tr>
<tr>
<td>Adawukwa</td>
<td>79</td>
<td>100.00(^a)</td>
<td>0.00(^b)</td>
</tr>
<tr>
<td>Okyereko</td>
<td>88</td>
<td>100.00(^a)</td>
<td>0.00(^b)</td>
</tr>
<tr>
<td>Total</td>
<td>308</td>
<td>57.71 ± 11.76(^a)</td>
<td>42.86 ± 11.69(^b)</td>
</tr>
</tbody>
</table>

Means in the same row with different letters are significantly different at P<0.05. S.E.: standard error of mean.

3 Results

3.1 Indoor resting mosquito density

A total of 925 indoor resting mosquitoes, comprising 640 (68%) female *An. gambiae* s.l., 252 (27%) female *An. funestus*, 34 (4%) male anophelines and 9 (1%) *Culex* spp. were collected from the six villages (Table 1). The indoor resting densities of *An. gambiae* s.l. were generally higher (P<0.05) than those of *An. funestus* in all villages except Adawukwa and Agyenkwaso, where there was no significant differences in their abundance (P>0.05).

From a total of 308 female *An. gambiae* s.l. processed, 176 (57.71%) were identified as *An. coluzzii* and 132 (42.86%) as *An. gambiae* s.s. (Table 2). *An. gambiae* s.s. was predominant in Adawukwa, Agyenkwaso, and Osorongma, and was the only member of the species complex collected from Kusa Dinkyeae. On the other hand, collections from Adawukwa and Okyereko, which are very close to each other, were exclusively *An. coluzzii*.

3.2 Prevalence of kdr mutations

The L1014F mutation was present in both *An. gambiae* s.s. and *An. coluzzii* (Table 3). The allele frequencies were slightly lower in *An. coluzzii*, ranging from 0.75 in the Osorongma population to 0.90 in the Agyenkwaso population. In Adawukwa and Okyereko, the allele frequencies were 0.87 and 0.86, respectively. In *An. gambiae* s.s., it was almost fixed in all the populations, being 0.97 in Atatam, 0.99 in Osorongma and 1.00 in Agyenkwaso and Kusa Dinkyeae.

Table 3. Allele distribution of the west African kdr mutation in *An. gambiae* s.l. from six villages in southern Ghana.

<table>
<thead>
<tr>
<th>Village</th>
<th><em>An. coluzzii</em></th>
<th><em>An. gambiae</em> s.s.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adawukwa</td>
<td>–(^a)</td>
<td>0.87 (79)(^a)</td>
</tr>
<tr>
<td>Agyenkwaso</td>
<td>0.97 (39)(^a)</td>
<td>1.00 (29)(^b)</td>
</tr>
<tr>
<td>Kusa Dinkyeae</td>
<td>0.75 (2)(^a)</td>
<td>0.99 (39)(^c)</td>
</tr>
<tr>
<td>Osorongma</td>
<td>0.90 (5)(^a)</td>
<td>1.00 (17)(^b)</td>
</tr>
<tr>
<td>Adawukwa</td>
<td>1.00 (17)(^c)</td>
<td>0.97 (48)(^c)</td>
</tr>
<tr>
<td>Okyereko</td>
<td>0.87 (79)(^a)</td>
<td>0.97 (39)(^c)</td>
</tr>
<tr>
<td>Okyereko</td>
<td>0.86 (88)(^a)</td>
<td>–</td>
</tr>
</tbody>
</table>

* Insufficient specimens available for analysis. Values in the same column with same superscript letters are not significantly different at P<0.05.

3.3 Blood meal source and sporozoite rates

Results for blood meal source identification are shown in Table 4. The PCR method could not identify the source of blood meal in some specimens despite physical evidence of blood in the abdomen. A total of 414 specimens were assayed, with success rates ranging from 50 to 100%. A high level of anthropophagy was recorded for the three endophilic anophelines in all villages sampled. *An. funestus* populations from the three ecological zones sampled were exclusively anthropophagic (HBI = 1), i.e. no animal feeding was observed. *An. gambiae* s.s. and *An. coluzzii* populations were similarly highly anthropophagic; however, evidence of animal blood feeding was recorded in both species. All *An. gambiae* s.s. and *An. coluzzii* specimens that fed on animals had also fed on humans, except for one *An. coluzzii* specimen from Okyereko that had exclusively fed on animal blood (cow).

*Plasmodium falciparum* infection rates were determined for 429 specimens, comprising 131 *An. gambiae* s.s., 175 *An. coluzzii* and 123 *An. funestus* (Table 5). There was evidence of infection in all the villages except Okyereko, where *P. falciparum* was not present in any of the 90 specimens processed. After pooling the data from all villages, infectivity was highest in *An. funestus*, followed by *An. gambiae* s.s. and lowest in *An. coluzzii*.

4 Discussion

4.1 Indoor resting mosquito diversity

*Anopheles gambiae* s.l. and *An. funestus* s.s. were the only anophelines collected from the villages sampled. The in-
The presence of breeding sites suitable for proliferation of \textit{An. gambiae} could be attributed to the availability of breeding sites and environmental variables, such as temperature and relative humidity [7, 25, 26]. In this study, breeding sites around villages in the CSEZ were collected indoors compared to \textit{An. funestus} in Adawukwa and Okyereko from the CSEZ. Only one \textit{An. funestus} specimen was collected from Osorongma in the FTEZ. Members of the \textit{An. gambiae} complex showed varied distribution across the study area, with collections from the FEZ and FTEZ being predominantly \textit{An. gambiae} s.s. with very few \textit{An. coluzzii}, except in Kusa Dinkyeae, where no \textit{An. coluzzii} was collected. \textit{An. gambiae} s.l. from the CSEZ were, however, exclusively \textit{An. coluzzii}. These results are similar to other findings from southern Ghana [4, 6, 7]. All collections from Obuasi in 2011 [6], a location closest to the three FEZ sampling sites, were \textit{An. gambiae} s.s. Similarly, 100% of all collections from Kumasi, a locality in the FEZ, were \textit{An. gambiae} s.s., and 98% from Osorongma in the FTEZ [4]. Results from Okyereko and Adawukwa (100% \textit{An. coluzzii} from each site) also confirm findings dating back to 2004 [4]. The relative abundance of these two species, which until recently were considered molecular forms of the same species (\textit{An. gambiae} s.s.), has been associated with breeding site characteristics. \textit{An. coluzzii} tends to be associated with flooded or irrigated sites that provide permanent breeding conditions, whereas \textit{An. gambiae} s.s. is associated with rain-dependent temporary sites [23].

The results obtained in the current study are in accordance with several reports across Africa, where the three species occur in sympatry over much of their range [2, 24]. Prevalence of anophelines in any locality is influenced by ecological factors, such as availability of preferred hosts, breeding sites and environmental variables, such as temperature and relative humidity [7, 25, 26]. In this study, mosquitoes were collected using indoor collection techniques; therefore, availability of host and resting behaviour were the main expected determinants of occurrence. As all species collected in this study are highly anthropophilic, endophagic and endophilic [2], it was expected that they would be equally represented in the collections. However, this was not the case. The differences in the distribution observed could be attributed to the availability of breeding sites around the villages where mosquitoes were sampled. Presence of breeding sites suitable for proliferation of \textit{An. gambiae} s.s. and \textit{An. funestus}, i.e. availability of temporary water puddles during the rainy season due to poorly permeable clay soils and several rivers around the villages of Atatam and Agyenkwaso, could be the underlying factors for the presence of both species in equal densities. Since there were no permanent water bodies in Kusa Dinkyeae this may explain the predominance of \textit{An. gambiae} s.s.

The breeding sites around villages in the CSEZ are different from those in the FEZ. Okyereko is an irrigated rice-cultivating community, and Adawukwa has a vast swamp and a nearby stream. The rice fields close to these two villages are ideal breeding sites for \textit{An. coluzzii} and, to a lesser extent, \textit{An. funestus}, while the edges of the stream and swamps are ideal sites for \textit{An. funestus} breeding. This probably explains the presence of both species in Okyereko and Adawukwa. The highest density of \textit{An. funestus} was recorded in Adawukwa, which is located closer to a swamp, suitable for \textit{An. funestus} breeding. Our findings agree with those by Charlwood and Edoh [26], who reported that anopheline adult density is negatively correlated with the distance between larval habitats and houses. The complete absence of \textit{An. funestus} in Osorongma could be explained by the absence of suitable breeding sites, as there are no permanent water bodies suitable for \textit{An. funestus} breeding near this village. Furthermore, according to de Souza et al. [7], the distribution of the members of the \textit{An. gambiae} complex appears to be driven by environmental factors, with \textit{An. gambiae} s.s. more predominant in the forested areas characterised by lower mean daily temperatures, whereas \textit{An. coluzzii} is predominant in the northern and coastal savannah areas, where mean daily temperatures are slightly higher.

The densities reported here may not represent the actual annual anopheline distribution in the study areas. Sampling was only limited to two seasons: the minor rainy season (September to November, 2011) and the dry season (December 2011 - January 2012). Species composition and abundance varies with time of year [6]. Yawson et al. [4] collected 58 \textit{An. funestus} out of 341 specimens from 10 collections in Osorongma during the major rainy season (June to September), whereas in the present study, only one specimen of \textit{An. funestus} was collected from the same village during the dry season. In this study, more \textit{An. gambiae} was collected during the minor rainy season than the dry season and vice versa for \textit{An. funestus} (data not shown).

### 4.2 Prevalence of the \textit{kdr} mutation

We observed high variability in the prevalence of the L1014F mutation in \textit{An. gambiae} s.s. and \textit{An. coluzzii}. The near fixation of the L1014F mutation in the populations from the areas sampled is understandable, since this mutation is widespread and predominant in West Africa [27]. High frequencies of the L1014F mutation have been reported from same location or locations close to the sampled villages across southern Ghana [4-6, 27]. Results from the current study show that the mutation is prevalent in both \textit{An. coluzzii} (90%) and \textit{An. gambiae} s.s. (98%). The L1014F mutation has shown a steady increase in \textit{An. gambiae} s.s.
Anopheles gambiae s.s., An. coluzzii and An. funestus are the major indoor resting malaria vectors in all the three ecological zones where this study was carried out, but showing variations in their distribution. An. gambiae s.s. predominates in the forested areas, whereas An. coluzzii predominates in the coastal savannah and An. funestus shows a wider distribution across southern Ghana. This distribution is highly influenced by breeding site availability and environmental factors. Thus, any research and development activity that focuses on developing control tools targeting these major malaria vectors might have to be carried out across all the ecological zones to be relevant for area-wide implementation. Similarly, to carry out any control programme across these ecological zones, the tools must target all three species.

The L1014F kdr mutation is present at very high frequencies in all three ecological zones, with its spread in the population nearing fixation in both An. gambiae s.s. and An. coluzzii. This implies that control programmes must include resistance management strategies and incorporate novel tools that are not affected by the kdr mutation. There should also be extensive resistance monitoring in the country and further research into the impact of this and other resistance mechanisms on vector control activities in Ghana.
6 Acknowledgements

The authors would like to thank the chiefs and people of all six communities where the study was carried out. Many thanks to the International Atomic Energy Agency (IAEA) for the Technical Cooperation Fellowship to M.O. through the Graduate School of Nuclear and Allied Science, University of Ghana, and to the Management and staff of the Vector Control Reference Laboratory of the National Institute for Communicable Diseases in South Africa for hosting and assisting M.O. with all laboratory work. Thanks also to the management and staff of the Biotechnology and Nuclear Agriculture Research Institute of Ghana Atomic Energy Commission for logistical support during field collections.

References


