Molecular Tools for Early Detection of Invasive Malaria Vector Anopheles stephensi Mosquitoes

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Reports of the expansion of the Asia malaria vector Anopheles stephensi mosquito into new geographic areas are increasing, which poses a threat to the elimination of urban malaria. Efficient surveillance of this vector in affected areas and early detection in new geographic areas is key to containing and controlling this species. To overcome the practical difficulties associated with the morphological identification of immature stages and adults of An. stephensi mosquitoes, we developed a species-specific PCR and a real-time PCR targeting a unique segment of the second internal transcribed spacer lacking homology to any other organism. Both PCRs can be used to identify An. stephensi mosquitoes individually or in pooled samples of mixed species, including when present in extremely low proportions (1:500). This study also reports a method for selective amplification and sequencing of partial ribosomal DNA from An. stephensi mosquitoes for their confirmation in pooled samples of mixed species.

Reports of the expansion of the malaria vector *Anopheles stephensi* mosquito (1–6) are increasing, intensifying the threat of urban malaria (7). The gradual southward expansion of this species has been recorded in India since the 1970s (1). The first occurrence of *An. stephensi* mosquito in Africa was reported as early as 1966 in a town in Egypt (2). During the past 2 decades, several reports of expansion of this species in the Horn of Africa, Sudan, Sri Lanka, and Lakshadweep (a union territory of India) have appeared (1–6). A high probability of presence within many urban cities across Africa has been predicted, which warrants prioritizing vector surveillance (8).

As a consequence of recent invasions of this vector species in several countries, the World Health

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Organization (WHO) recommended conducting active surveillance of An. stephensi mosquitoes in urban and periurban areas, in addition to routine surveillance in rural areas in the affected and surrounding geographic areas (9). However, identifying this species, a process that relies mainly on the morphologic characteristics of adult female mosquitoes, is often challenging. Sampling of adult *An*. stephensi mosquitoes from their resting habitats is difficult because they are secretive in their habits (10). WHO has recommended sampling immature An. stephensi mosquitoes from natural breeding habitats and rearing them in the laboratory until their emergence into adults to enable species identification (9). This practice is being adopted for sampling this species in regions of Africa (4,5,11), which is a time-consuming and labor-intensive process. Even freshly emerged adults have been reported to be misidentified as An. gambiae mosquitoes because of superficial resemblance (11).

Adult mosquito collection through light-trap or pyrethrum spray collections are alternative and popular methods of sampling An. stephensi mosquitoes, but identifying adults collected through such methods can be difficult because of the loss of morphologic characteristics critical for their correct identification. Therefore, developing highly specific PCR-based assays is crucial for identifying both larval and adult An. stephensi species collected by a variety of methods. Such diagnostics will be helpful to field technologists who are not familiar with the morphology of this species. Those assays can detect An. stephensi mosquitoes in a large pool of mosquitoes when their proportion is extremely low. Equally vital is developing a DNA sequencing strategy to confirm the presence of An. stephensi mosquitoes in such pools of mixed species. Such molecular tools will help detect invasions of An. stephensi mosquitoes early in new geographic areas where they are present in extremely low densities.

Methods

Mosquito Samples

We obtained adult mosquitoes or their DNA samples belonging to a total of 17 anopheline and 3 culicine species from different parts of the world either from BEI Resources (https://www.beiresources.org) or locally (Appendix Table, https://wwwnc.cdc.gov/EID/article/29/1/22-0786-App1.pdf). Field-collected *An. stephensi* mosquitoes were processed for identifying biologic forms according to the methods described by Singh et al. (10).

DNA Isolation from Individual and Pooled Samples

We isolated DNA from individual specimens as well as pooled samples of mosquitoes to standardize PCR-based assays and their validation. We used pooled samples of different sizes, each consisting of a single *An. stephensi* mosquito and the rest *An. culicifacies* mosquitoes. We also used a pool of field-collected mosquitoes consisting of a single *An. stephensi* mosquito and other species.

We selected 2 commercial kits for DNA isolation. For DNA isolation from individual mosquitoes or smaller pool sizes, we used the DNeasy Blood and Tissue kit (QIAGEN, https://www.qiagen.com), recommended for ≥25 mg of tissue. For larger pools, we used DNAzol Reagent (ThermoFisher Scientific, https://www.thermofisher.com), recommended for 25–50 mg of tissue per milliliter of reagent. In addition, we isolated DNA from individual *An. stephensi* mosquitoes by boiling method.

DNeasy Blood and Tissue Kits

We isolated DNA from individual mosquitoes of some *Anopheles, Culex*, and *Aedes* species (Appendix Table) following the manufacturer's protocol and eluted in 200 μ L elution buffer. We isolated DNA from smaller pools of III–IV instar larvae or adults, each containing a single *An. stephensi* mosquito and the rest *An. culicifacies* mosquitoes in different pool sizes (i.e., 2, 5, 10, 20, and 50). We also isolated DNA from the single leg of an *An. stephensi* mosquito, which was eluted in 50 μ L of elution buffer.

DNAzol Reagent

We used DNAzol Reagent for DNA isolation from pools (i.e., 25, 100, and 500) of adult mosquitoes and pools of 100 larvae (III–IV instar), each containing 1 An. stephensi mosquito and the rest An. culicifacies mosquitoes. We directly homogenized pools of 25 and 100 mosquitoes in 1 mL of DNAzol reagent in a microcentrifuge tube. In the case of pools of 100 mosquitoes, we transferred 250 μ L of the triturate to

a separate microcentrifuge tube to make up a volume of 1 mL with DNAzol. We ground the pools of 500 mosquitoes in liquid nitrogen and transferred $\approx\!25$ mg of triturate in a microcentrifuge tube and homogenized in 1 mL of DNAzol reagent. We centrifuged all triturates at $10,000\times g$ for 10 min and transferred 500 μL of supernatant to a fresh 1.5-mL microcentrifuge tube, which we subjected to ethanol precipitation, washing, and solubilization of DNA following the vendor's protocol. We dissolved DNA in 200 μL of 8.0 mM NaOH. We also isolated DNA from a pool of 100 mosquitoes containing a single An. stephensi mosquito and field-collected (through hand-catch method) mosquitoes belonging to An. culicifacies, An. subpictus, An. fluviatilis, and Culex quinquefasciatus.

Boiling Method

We isolated DNA from 10 individual specimens of An. stephensi mosquitoes by the boiling method described by Sharma et al. (13). We either used the DNA isolated from this method immediately after isolation or preserved it at -20 °C for later use.

Selecting Target Sites for Designing Primers and Probes

We selected ribosomal DNA (rDNA) as a target site for developing diagnostics to identify An. stephensi mosquitoes, which are present in hundreds of copies in an individual and are highly conserved in a species because of homogenization of sequence through unequal crossing over and gene conversion, a process known as concerted evolution. We selected the internal transcribed spacer 2 (ITS2) rDNA, which is conserved within a species but highly variable across taxa, for designing An. stephensi-specific primers and probes. For designing An. stephensi-specific primers and probes, we conducted a homology search of ITS2 sequences of An. stephensi (14) through a nucleotide BLAST search (https://blast.ncbi.nlm.nih.gov/Blast. cgi). We optimized the search for blastn (somewhat similar sequences) and excluded the taxon Anopheles stephensi from the search. All 297 search returns belonged to the Anopheles mosquitoes, all belonging to the Neocellia series (subgenus Cellia); however, none of the returns showed homology to the last 122 bp segment of ITS2. We considered this region unique to An. stephensi mosquitoes and exploited this region for designing highly specific An. stephensi-specific primers and probes (Figure 1). For designing universal primers and a probe, we identified highly conserved regions from 5.8S and 28S rDNA (Figure 1) based on the alignment of sequences of anophelines available in the GenBank. To verify the specificity of each

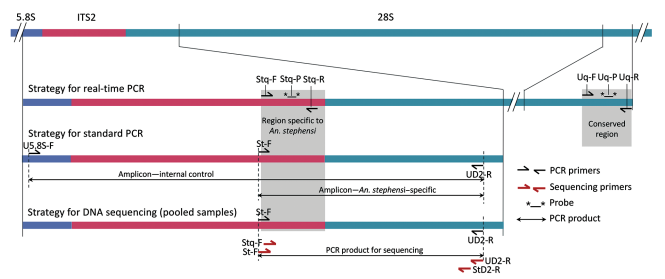


Figure 1. Schematic representation of PCR and sequencing strategies used for early detection of invasive malaria vector *Anopheles stephensi* mosquitoes

An. stephensi-specific primer and probe (Table 1), we performed a blastn search in silico, which ensured that none of the primers and probes matched rDNA sequences of any other mosquitoes.

Development of a Hydrolysis Real-Time PCR

For An. stephensi-specific hydrolysis real-time PCR, we designed 2 oligonucleotide primers, (Stq-F and Stq-R) and a hydrolysis probe (Stq-P) from the An. stephensispecific ITS2 region (Table 1; Figure 1). For internal control (IC), we designed primers (Uq-F and Uq-R) and a hydrolysis probe (Uq-P) from a region of 28S-rDNA conserved in anophelines (Table 1; Figure 1). We performed real-time PCR in 10 µL of reaction mixture containing 0.4 µM each of Uq-F, Uq-R and Stq-F; 0.5µM of Stq-R; 0.2 μM of each probe (Stq-P and Uq-P); 1X QuantiFast Multiplex PCR kit (QIAGEN); and 1 µL of template DNA in Bio-Rad CFX96 Touch Real-Time PCR Detection System (https://www.bio-rad.com). The cyclic conditions were predenaturation at 95°C for 5 min, followed by 35 cycles, each with denaturation at 95°C for 10 s and annealing/extension at 60°C for 30 s. We scored the number

of cycles required for the fluorescent signal to cross the threshold (cycle threshold [Ct] values) by using the software CFX Maestro (Bio-Rad) (Appendix Figure 1).

We evaluated PCR efficiencies of each hydrolysis probe assay by performing duplex real-time PCR assays in triplicate at 6 different concentrations, diluted serially by 10-fold. We performed the experiments on 2 samples of An. stephensi DNA with different DNA concentrations, 8.7 and 3.2 ng/ μ L (Figure 2). To determine the limit of detection (LOD), we conducted real-time PCR on the diluted DNA of An. stephensi mosquitoes with concentrations of 160 fg, 80 fg, 40 fg, and 20 fg, each with 12 replicates.

Development of Size-Diagnostic PCR

For a size-diagnostic PCR, we designed 3 primers: we designed the *An. stephensi*-specific forward primer (St-F) from the *An. stephensi*-specific region of ITS2 and designed 2 flanking universal primers from conserved 5.8S rDNA (U5.8S-F) and D2 domain of 28S rDNA (UD2-R). According to the strategy designed (Figure 1), an *An. stephensi*-specific diagnostic

Table 1. Sequences of primers and probes used in study of molecular tools for early detection of invasive malaria vector *Anopheles stephensi* mosquitoes

Name	Sequence, $5' \rightarrow 3'$	Annealing specificity	Reference
Stq-F	TCTTTCCTCGCATCCAGTTG	An. stephensi	This study
Stq-R	CGGGAGAAGCGGTGATAAAT	An. stephensi	This study
Stq-P	/56-FAM/CGTGCTAAC/ZEN/CTCACTCACCCACAC/3IABKFQ/	An. stephensi	This study
Uq-F	GAGATTCCCTCTGTCCCTATCT	Universal	This study
Uq-R	AGCTCAACAGGGTCTTCTTTC	Universal	This study
Uq-P	/5HEX/TAGCGAAAC/ZEN/CACAGCCAAGGGAA/3IABKFQ/	Universal	This study
U5.8S-F	ATCACTCGGCTCATGGATCG	Universal	(15)
St-F	CGTATCTTTCCTCGCATCCA	An. stephensi	This study
UD2-R	GCACTATCAAGCAACACGACT	Universal	This study
StD2-R	GTCTGCCACCACAGTCCT	An. stephensi	This study

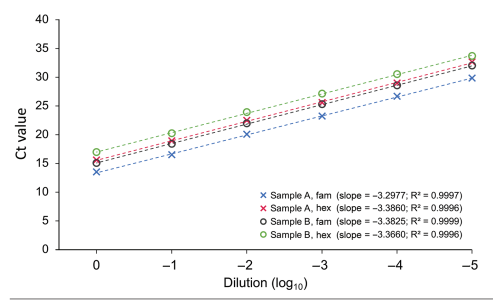


Figure 2. Standard curve showing correlation of Ct values against 10-fold serially diluted DNA samples of *Anopheles stephensi* mosquitoes (2 samples, A and B) in the duplex hydrolysis fluorescent probe assay. The slope of each line represents [–1/log₁₀ (PCR efficiency)] for a hydrolysis probe assay. R² represents correlation coefficient of a slope. Ct, cycle threshold.

amplicon of 438 bp size will be formed by the primers St-F and D2-R, and a universal amplicon of varying sizes (>600 bp), depending upon the length of ITS2 in a particular species, will be formed by the primers 5.8S and D2-R. The universal amplicon will serve as an IC to rule out PCR failure.

Because of the competitive nature of primers in multiplex-PCR, we optimized 2 protocols of PCR on the basis of the number of mosquitoes in a pool. For individual mosquitoes or smaller pools, we conducted size-diagnostic PCR assays by using Hot Start Taq 2X Master Mix (New England Biolabs, https://www.neb.com) in a 20-µL reaction mixture containing 0.5 units of taq polymerase, 1.5 mM of MgCl₂, 0.25 μM of primer St-F, 0.25 μM of primer U5.8S-F, 0.375 µM of UD2-R, and 0.50 µL of DNA template (protocol-1). In another protocol (protocol-2), we reduced the concentration of primer U5.8S-F to 0.10 µM and increased the concentration of UD2-R to 0.50 μM for larger pools of mosquitoes (25–500). We also observed that the intensity of amplicon in size-diagnostic PCR with pools of mosquitoes (>25) could be improved by the dilution of template DNA. Therefore, the template DNA of larger pools of mosquitoes (>25) was further diluted by 1/10 before using it as template DNA for size-diagnostic PCR reactions to minimize the PCR inhibitors in PCR reactions. However, such dilution was not required for real-time PCR. Optimized thermal cycling conditions were an initial denaturation step at 95°C for 30 s, 30 cycles each with a denaturation step at 95°C for 30 s, annealing at 55°C for 30 s and extension at 68°C for 45 s, and final extension at 68°C for 7 min. Five µL of PCR product

was run on 2% agarose gel and visualized in the gel documentation system.

DNA Sequencing Strategy for the Confirmation of PCR-Based Identification of *An. stephensi* Mosquitoes in Pooled Samples

We amplified *An. stephensi*-targeted amplicons from DNA isolated from pools of 100 (field-collected) and 500 mosquitoes, each pool containing a single An. stephensi mosquito, using primers St-F and UD2-R. We performed PCR by using Hot Start Taq 2X Master Mix in a 20 µL reaction mixture containing 0.25 µM of each primer. PCR conditions were similar to PCR protocol-1 but with extension time reduced to 30 s and number of cycles increased to 35. The amplified products were treated with Exo-Sap II, and sequence termination reactions were performed from both directions of strands using BigDye Terminator v3.1 Cycle Sequencing Kit (both ThermoFisher Scientific). The primers used for sequencing were the primers used for PCR amplification as well as the 2 internal primers (Stq-F and StD2-R) (Table 1; Figure 1). Both internal primers are specific to *An*. stephensi and were expected to provide a noise-free sequence by eliminating the possibility of sequencing nonspecific PCR product. The sequencing products were electrophoresed on an ABI Prism 3730xl (ThermoFisher Scientific).

Results

Real-time PCR

PCR efficiencies, as estimated based on Ct values of 6 serially diluted concentrations of DNA, were 97.5%-

101% for *An. stephensi*-diagnostic (Fam-labeled probe) PCR (Figure 2). The dynamic range of Ct valprobe) PCR and 97.4%–98.2% for IC (Hex-labeled ues for real-time PCR for *An. stephensi*-specific PCR

Table 2. Results of hydrolysis real-time PCR and size-diagnostic PCR on individual and pooled mosquitoes for early detection of invasive malaria vector. *Ann helps, tenhensi*, mosquitoes*

	DNA		Real-time PCR			Size-diagnostic	PCR
	isolation		Ct values (An.			An. stephensi-	
Specimen type	method	No.	stephensi)†	Ct values IC	No.	specific band	IC ban
Anophelines							
Neocellia series							
An. stephensi type	DNeasy	26	13.47-17.23	15.20-18.93	24	Positive	Positiv
An. stephensi intermediate	DNeasy	2	14.75-15.60	15.39-16.06	2	Positive	Positiv
An. stephensi var. mysorensis	DNeasy	1	14.54	15.01	1	Positive	Positiv
An. stephensi strain STE2	DNeasy	1	14.85-15.25	16.06-16.51	2	Positive	Positiv
An. stephensi type form	Boiling	10	15.17-17.41	15.97-18.05	-	Not done	Not do
An. stephensi type form, single leg	DNeasy	4	18.64-22.29	19.02-23.51	4	Positive	Positiv
Pyretophorus series	,						
An. gambiae	DNeasy	3	Negative	14.58-17.32	3	Negative	Positiv
An. quadrimaculatus	DNeasy	2	Negative	14.52-15.64	2	Negative	Positiv
An. merus	DNeasy	2	Negative	15.22-15.35	2	Negative	Positiv
An. subpictus form A	DNeasy	2	Negative	16.61-17.02	1	Negative	Positiv
An. subpictus form A‡	Pre-isolated§	1	33.09	18.32		Negative	Positiv
An. subpictus form B	Pre-isolated§	2	Negative	16.17–16.35	1	Negative	Positiv
An. sundaicus cytoform D	Pre-isolated§	1	Negative	14.47	1	Negative	Positiv
Albimanus series		•			•	. 10944110	
An. albimanus	DNeasy	2	Negative	16.21-16.58	1	Negative	Positiv
Anopheles series	2.1000)	_			•	. 10944110	
An. freeborni	DNeasy	2	Negative	15.73-15.81	1	Negative	Positiv
An. atroparvus	DNeasy	2	Negative	14.70–15.88	1	Negative	Positiv
Neomyzomyia series	Divodoy	_	rioganio	11.70 10.00		rtogativo	1 00111
An. dirus	DNeasy	2	Negative	17.34-18.01	1	Negative	Positiv
An. farauti	DNeasy	2	Negative	19.09–19.48	1	Negative	Positiv
Myzomyia series	2.1040)	_		10100 10110	•	. rogain o	
An. funestus	DNeasy	2	Negative	15.50-15.62	1	Negative	Positiv
An. culicifacies species A	DNeasy	1	Negative	17.41	1	Negative	Positiv
An. culicifacies species B	Pre-isolated§	1	Negative	17.48	1	Negative	Positiv
An. fluviatilis species S	Pre-isolated§	1	Negative	16.61	1	Negative	Positiv
An. fluviatilis species T	Pre-isolated§	1	Negative	17.02	1	Negative	Positiv
Culicines	1 10 100141043	•	rioganio	17.02		rtoganto	1 00111
Culex quinquefasciatus	DNeasy	3	Negative	16.58-19.20	2	Negative	Positiv
Aedes albopictus	DNeasy	2	Negative	16.65–16.68	2	Negative	Positiv
Ae. aegypti	DNeasy	2	Negative	18.96–19.20	2	Negative	Positiv
Ae. aegypti‡	Pre-isolated§	1	32.95	20.97	_	Negative	Positiv
Single <i>An. stephensi</i> individual pooled wi			02.00	20.01		rrogativo	1 001111
Adults (1/2)	DNeasy	1	17.06	18.31	1	Positive	Positiv
Adults (1/5)	DNeasy	1	15.37	16.11	1	Positive	Positiv
Adults (1/10)	DNeasy	1	15.77	15.30	1	Positive	Positiv
Adults (1/20)	DNeasy	1	15.74	13.71	1	Positive	Positiv
Adults (1/20) Adults (1/50)	DNeasy	1	15.43	11.43	1	Positive	Positiv
Adults (1/30) Adults (1/25)	DNAzol	1	14.53	12.61	1	Positive	Positiv
Adults (1/23) Adults (1/100)	DNAzol	1	16.37	11.27	1	Positive	Positiv
Adults (1/500)	DNAzol	2	20.12–23.96	11.08–11.50	2	Positive	Positiv
Larvae (1/2)	DNeasy	1	15.56	16.87	1	Positive	Positiv
Larvae (1/5)	DNeasy	1	15.49	16.65	1	Positive	Positiv
Larvae (1/10)	DNeasy	1	16.36	15.53	1	Positive	Positiv
Larvae (1/20)	DNeasy	1	16.51	15.06	1	Positive	Positiv
Larvae (1/50)	DNeasy	1	16.27	14.40	1	Positive	Positiv
Larvae (1/100)	DNAzol	1	18.14	13.16	1	Positive	Positi
Single An. stephensi individual pooled wi							
Adults (1/100)	DNAzol	1	17.61	11.70	1	Positive	Positiv

^{*}Ct, cycle threshold; IC, internal control.

[†]Ct value shown as Negative implies that no Ct value was scored within 35 cycles of reactions.

[‡]Contaminated with An. stephensi DNA.

[§]Preserved DNA isolated manually for other studies.

Denominator of numeric expression inside parenthesis indicate total number of mosquitoes in a pool.

was 13.5–32 and for IC-PCR was 15.5–33.5; the LOD was 40 fg of genomic DNA.

The real-time PCRs conducted on DNA isolated from individual An. stephensi mosquito samples, 19 different nontarget Anopheles mosquito species, and a single An. stephensi mosquito in different pool sizes were specific based on software-determined Ct value scored within 35 cycles of reactions, except in the case of 2 preisolated DNA (1 each of Aedes aegypti and An. subpictus mosquitoes) showing false positivity with late Ct values (>32) (Table 2). The 2 false-positive samples were contaminated with the An. stephensi mosquito DNA as revealed through An. stephensitargeted sequencing (Appendix). Although primers and probe for IC were designed on the basis of anophelines' 28S rDNA sequences, they also worked on all 3 nonanopheline species tested (i.e., Cx. quinquefasciatus, Ae. aegypti, and Ae. albopictus mosquitoes). The real-time PCR also successfully identified An. stephensi mosquito from DNA samples isolated by boiling method and DNA isolated from a single leg of mosquitoes. The real-time PCR was sensitive to detecting a single An. stephensi mosquito in pools of 500 mosquitoes with low Ct values (<24) (Table 2). Rising of fluorescent signal was noticed with IC probe in some experiments only in negative controls after 30 cycles but not with An. stephensi-specific probe (Appendix Figure 1).

Size-Diagnostic PCR

We performed PCR protocol-1 on DNA samples isolated from individual *An. stephensi* mosquito samples, nontarget mosquitoes, and pooled samples of mixed

species (≤50 mosquitoes), which provided desired amplicons. All An. stephensi mosquitoes were positive for An. stephensi-specific band (438 bp), and all other mosquito species were negative (Table 2; Figures 3, 4). All species exhibited amplification of an IC band of varying sizes (>600 bp). Ae. aegypti mosquitoes exhibited the smallest IC band (≈650 bp) because of the shortest length of ITS2 (200 bp). An. funestus and An. dirus mosquitoes exhibited the largest IC bands (>1 kb) because of longer ITS2 (700 bp). PCR protocol 1 successfully identified An. stephensi mosquitoes in all pooled samples of mixed species, where the An. stephensi-specific band was prominent in pools of <20 mosquitoes. We observed that the An. stephensi mosquito diagnostic band grew fainter as the concentration of An. stephensi mosquito DNA decreased in larger pools (Figure 3, panel A). Therefore, a different protocol (protocol 2) with different primer concentrations was adopted for larger pools. PCR protocol 2 successfully identified An. stephensi mosquitoes in pools of 25-500 mosquitoes and provided clearly visible *An. stephensi*-specific band in pools of ≤100 mosquitoes. On the basis of these results, we found PCR protocol-1 suitable for individual samples or smaller pools (up to 25) (Figure 3, panel A) and PCR protocol-2 suitable for larger pools (25-100 mosquitoes) (Figure 3, panels B and C).

Sequencing Results

An. stephensi-targeted DNA sequencing was successful with all 4 sequencing primers. The quality of DNA sequences generated from pooled mosquitoes was reasonably high (Appendix Figure 2). The output

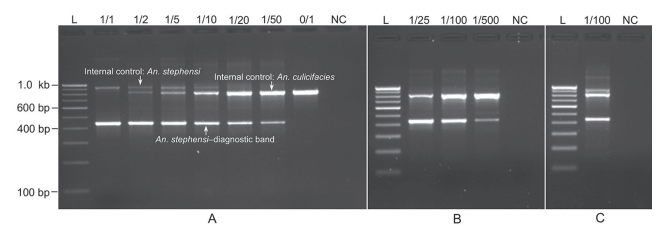


Figure 3. Gel photographs visualizing the result of PCRs specific to invasive malaria vector *Anopheles stephensi* mosquitoes. A) PCR protocol 1 on individual specimens and pools of *An. stephensi* and *An. culicifacies*. B) PCR protocol 2 on larger pools of *An. stephensi* and *An. culicifacies*. C) PCR protocol 2 on a pool of 100 mosquitoes containing a single *An. stephensi* mixed with other wild-caught anophelines (*An. culicifacies*, *An. fluviatilis*, *An. minimus*, and *An. subpictus*). The numerator of numeric expression shown on the top of each lane indicates number of *An. stephensi* in a pool, and denominator indicates size of mosquito-pool. L, 100-bp ladder; NC, negative control.

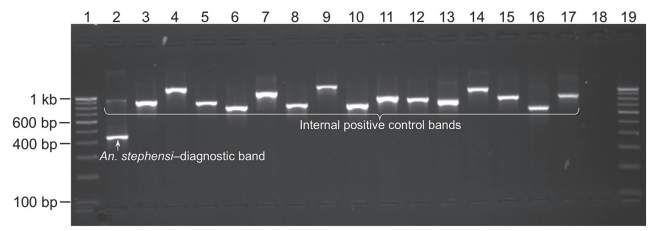


Figure 4. Gel photograph showing the result of PCR specific to invasive malaria vector *Anopheles stephensi* mosquitoes (protocol 1) on some individual mosquitoes belonging to the genus *Anopheles*, *Culex*, and *Aedes*. Lanes 1 and 19, 100-bp DNA ladder; 2, *An. stephensi*; 3, *An. gambiae*; 4, *An. dirus*; 5, *An. albimanus*; 6, *An. quadrimaculatus*; 7, *An. farauti*; 8, *An. freeborni*; 9, *An. funestus*; 10, *An. atroparvus*; 11, *An. merus*; 12, *An. fluviatilis* species T; 13, *Cx. quinquefasciatus*; 14, *An. subpictus* molecular form A; 15, *An. minimus* sensu strictu; 16, *Ae. aegypti*; 17, *Ae. albopictus*; 18, negative control.

sequences showed 100% similarity with *An. stephensi* mosquito sequences in a BLAST search. The second highest similarity was with *An. superpictus* mosquitoes, which demonstrated 94.88% similarity on the basis of only 73% coverage. The remaining 27% nucleotide sequence belonging to the ITS2 region did not show a match with any organism other than *An. stephensi* mosquitoes.

Discussion

The molecular methods developed in this study can identify and confirm *An. stephensi* mosquitoes individually or in a large pool of mixed mosquito species, which could enable screening of large numbers of samples collected through a variety of methods (such as light trap, pyrethrum spray collection, larval collection, etc.) with relatively limited effort and time. This method could help in early reporting of the presence of *An. stephensi* mosquitoes to concerned state health agencies and WHO.

In molecular diagnostics, we designed *An. ste-phensi*-specific primers and probes from a segment of ITS2 lacking homology to any other organisms for which the rDNA sequence database is available in the public domain, enabling the design of primers that are highly specific to *An. stephensi* mosquitoes and refractory to nonspecific annealing. However, this process does not preclude the possibility of any mosquito's sequence, as-yet unreported, having matching sequences with *An. stephensi*-specific primers or probes. Therefore, confirmatory DNA sequencing should be performed in new areas using the *An. stephensi*-specific primers suggested in this report.

In earlier studies, confirmation of this species was done through sequencing of ITS2 and mitochondrial DNA using universal primers, which cannot be used in pooled samples of mixed species. Moreover, Mishra et al. (14) have shown that direct DNA sequencing of ITS2 in the case of *An. stephensi* mosquitoes is not fruitful because of the presence of indel variants, which causes the collapse of a sequence starting from the indel position. The method proposed here for sequencing *An. stephensi* mosquitoes in a pooled sample was targeted to indel-free partial ITS2 (which lacks homology with any organism) and D1-D2 domains of 28S of rDNA, which are species-informative.

The real-time PCR developed in this study is for diagnostic purposes only and by no means intended for quantitative PCR (qPCR); qPCR is not reliable in the case of pooled mosquitoes belonging to different species because of interspecific variations in rDNA copy number (16) and body mass. However, the proportion of *An. stephensi* mosquitoes in a mosquito population, when present in extremely low density, can be obtained by the method used for the estimation of infection rates in hematophagous insects by estimating minimum infection rate or maximum-likelihood procedure (17) on the basis of the number of pools positive, methods frequently used for xenomonitoring.

LOD for the real-time PCR is considered a vital criterion for assessing the sensitivity of real-time PCR when the copy number of target nucleic acid is a limiting factor (e.g., detecting pathogens in an organism). However, LOD is not a limiting factor for the *An. stephensi*-specific diagnostic real-time PCR; rDNA is abundantly found in the organism because

of its high copy number. We observed in this study that LOD cannot be a limiting factor even when the proportion of target mosquitoes is 1/500 in a pooled sample or tested on DNA isolated from a single leg (Ct values <24). On the basis of our observations, we suggest a cutoff value of 30 for real-time PCR for more reliable results. Ct values above this threshold can be suspected to be DNA contamination which should be verified through DNA sequencing (Appendix). In this study, we observed false positivity for *An. stephensi* mosquitoes with late Ct values (>32) in 2 DNA samples (1 each of the *Ae. aegypti* and *An. subpictus* mosquito) because of the contamination of DNA from *An. stephensi* mosquitoes.

The diagnostic PCRs in this study were designed to identify An. stephensi mosquitoes in large pools of samples. However, pooling of a large number of samples can accumulate potential PCR inhibitors. The heme compound in the blood (18) and eye pigment in the head of an insect (19) are reported potential inhibitors. Although we did not observe inhibitory effect of pooling in the real-time PCR, we observed substantial inhibitory effect in a size-diagnostic PCR with a pool of \geq 50 mosquitoes. In this study, we experienced an improvement in the intensity of the band in size-diagnostic PCR in such pools by diluting DNA.

Although we have successfully demonstrated identifying a single An. stephensi mosquito in pools of 500 mosquitoes, using a pool of up to 100 mosquitoes that can be ground in a single microcentrifuge tube during DNA isolation without the need for grinding by mortar and pestle is recommended. Grinding by using a mortar and pestle might increase the risk for carryover contamination. For confirmation of An. stephensi mosquitoes in pooled samples through Sanger sequencing, we suggest using internal primers (Stq-F and StD2-R, both of which are specific to *An. stephensi* mosquitoes) for sequence termination reactions as a precautionary measure. This step is critical to rule out sequencing of false-positive PCR products because of nonspecific annealing with unknown nontarget species, if any.

In conclusion, the molecular tools developed in this study can be used to identify and confirm *An. ste-phensi* mosquitoes, individually or in a pool of mixed mosquito species. This process will enable health authorities to detect early invasion of the species, especially in areas where it exists with low density.

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O.P.S. designed PCR strategies, analyzed data, and wrote the first draft of the manuscript; G.S., T.K., M.P.K., and P.K.M. performed laboratory experiments; P.K.M., S.M., and N.K. contributed to the manuscript. All authors approved the final version of the manuscript.

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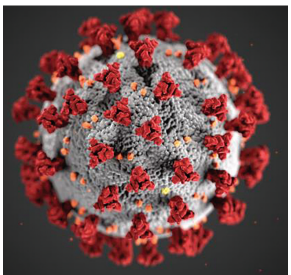
References

- Sharma SK, Hamzakoya KK. Geographical spread of Anopheles stephensi vector of urban malaria, and Aedes aegypti, vector of dengue/DHF, in the Arabian Sea Islands of Lakshadweep, India. Dengue Bulletin. 2001;25:88–91. WHO Regional Office for South-East Asia [cited 2020 Nov 24]. https://apps.who.int/ iris/handle/10665/148798
- Gad AM. Anopheles stephensi liston in Egypt, UAR. Mosq News. 1967;27:171–4.
- Faulde MK, Rueda LM, Khaireh BA. First record of the Asian malaria vector *Anopheles stephensi* and its possible role in the resurgence of malaria in Djibouti, Horn of Africa. Acta Trop. 2014;139:39–43. https://doi.org/10.1016/j.actatropica. 2014.06.016
- Carter TE, Yared S, Gebresilassie A, Bonnell V, Damodaran L, Lopez K, et al. First detection of *Anopheles stephensi* Liston, 1901 (Diptera: culicidae) in Ethiopia using molecular and morphological approaches. Acta Trop. 2018;188:180–6. https://doi.org/10.1016/j.actatropica.2018.09.001
- Balkew M, Mumba P, Dengela D, Yohannes G, Getachew D, Yared S, et al. Geographical distribution of Anopheles stephensi in eastern Ethiopia. Parasit Vectors. 2020;13:35. https://doi.org/10.1186/s13071-020-3904-y
- Gayan Dharmasiri AG, Perera AY, Harishchandra J, Herath H, Aravindan K, Jayasooriya HTR, et al. First record of *Anopheles stephensi* in Sri Lanka: a potential challenge for prevention of malaria reintroduction.

- Malar J. 2017;16:326. https://doi.org/10.1186/s12936-017-1977-7
- Takken W, Lindsay S. Increased threat of urban malaria from *Anopheles stephensi* mosquitoes, Africa. Emerg Infect Dis. 2019;25:1431–3. https://doi.org/10.3201/eid2507.190301
- Sinka ME, Pironon S, Massey NC, Longbottom J, Hemingway J, Moyes CL, et al. A new malaria vector in Africa: predicting the expansion range of *Anopheles stephensi* and identifying the urban populations at risk. Proc Natl Acad Sci U S A. 2020;117:24900–8. https://doi.org/10.1073/ pnas.2003976117
- 9. World Health Organizaton. Vector alert: *Anopheles stephensi* invasion and spread. 2019. WHO reference number: WHO/HTM/GMP/2019.09 [cited 2020 Nov 24]. https://apps.who.int/iris/rest/bitstreams/1242915/retrieve
- Christophers SR. The fauna of British India, including Ceylon and Burma. Diptera. 4. Family Culicidae, Tribe Anophelini. London: Taylor and Francis; 1933.
- 11. Ahmed A, Khogali R, Elnour MB, Nakao R, Salim B. Emergence of the invasive malaria vector *Anopheles stephensi* in Khartoum State, Central Sudan [Erratum in: Parasit Vectors. 2021;14:562]. Parasit Vectors. 2021;14:511. https://doi.org/10.1186/s13071-021-05026-4
- 12. Singh OP, Mishra S, Sharma G, Sindhania A, Kaur T, Sreehari U, et al. Evaluation of intron-1 of odorant-binding protein-1 of Anopheles stephensi as a marker for the identification of biological forms or putative sibling species. PLoS One. 2022;17:e0270760. https://doi.org/10.1371/journal.pone.0270760
- Sharma G, Lather M, Singh OP. Variations in palpal ornamentation of *Anopheles fluviatilis* species T and U (Diptera: Culicidae) and their taxonomic consequence. Indian J Exp Biol. 2020;58:64–8. https://doi.org./10.56042/ ijeb.v58i01.65321
- Mishra S, Sharma G, Das MK, Pande V, Singh OP. Intragenomic sequence variations in the second internal transcribed spacer (ITS2) ribosomal DNA of the malaria vector *Anopheles stephensi*. PLoS One. 2021;16:e0253173. https://doi.org/10.1371/journal.pone.0253173
- Manonmani A, Townson H, Adeniran T, Jambulingam P, Sahu S, Vijayakumar T. rDNA-ITS2 polymerase chain reaction assay for the sibling species of Anopheles fluviatilis. Acta Trop. 2001;78:3–9. https://doi.org/10.1016/ S0001-706X(00)00154-6
- Kumar A, Rai KS. Chromosomal localization and copy number of 18S + 28S ribosomal RNA genes in evolutionarily diverse mosquitoes (Diptera, Culicidae). Hereditas. 1990;113:277–89. https://doi.org/10.1111/j.1601-5223.1990.tb00094.x
- Walter SD, Hildreth SW, Beaty BJ. Estimation of infection rates in population of organisms using pools of variable size. Am J Epidemiol. 1980;112:124–8. https://doi.org/10.1093/ oxfordjournals.aje.a112961
- Akane A, Matsubara K, Nakamura H, Takahashi S, Kimura K. Identification of the heme compound copurified with deoxyribonucleic acid (DNA) from bloodstains, a major inhibitor of polymerase chain reaction (PCR) amplification. J Forensic Sci. 1994;39:362–72. https://doi.org/10.1520/JFS13607J
- Boncristiani H, Li J, Evans J, Pettis J, Chen Y. Scientific note on PCR inhibitors in the compound eyes of honey bees, *Apis mellifera*. Apidologie (Celle). 2011;42:457–60. https://doi.org/10.1007/s13592-011-0009-9

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