Multiplex Assay for Species Identification and Monitoring of Insecticide Resistance in Anopheles punctulatus Group Populations of Papua New Guinea

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Abstract. Anopheles punctulatus sibling species (An. punctulatus s.s., Anopheles koliensis, and Anopheles farauti species complex [eight cryptic species]) are principal vectors of malaria and filariasis in the Southwest Pacific. Given significant effort to reduce malaria and filariasis transmission through insecticide-treated net distribution in the region, effective strategies to monitor evolution of insecticide resistance among An. punctulatus sibling species is essential. Mutations in the voltage-gated sodium channel (VGSC) gene have been associated with knock-down resistance (kdr) to pyrethroids and DDT in malaria regions. By examining VGSC sequence polymorphism we developed a multiplex assay to differentiate wild-type versus kdr alleles and query intron-based polymorphisms that enable simultaneous species identification. A survey including mosquitoes from seven Papua New Guinea Provinces detected no kdr alleles in any An. punctulatus species. Absence of VGSC sequence introgression between species and evidence of geographic separation within species suggests that kdr must be monitored in each An. punctulatus species independently.

INTRODUCTION

Malaria vectors in the Southwest Pacific region belong to the Anopheles punctulatus group. The species group range extends from Moluccas to Vanuatu, including New Guinea and islands of the Bismarck Archipelago, the Solomon Islands, and northern Australia.1 Thirteen sibling species (An. punctulatus sensu stricto [s.s.], An. species near punctulatus, morphologically indistinguishable Anopheles farauti 1–8 [Farauti complex; former An. farauti 1, 2, 3, and 7 now An. farauti s.s., Anopheles hinesorum, Anopheles torresiensis, and Anopheles irenicus, respectively], Anopheles koliensis, Anopheles clowi, and Anopheles renellensis)1–9 are known to comprise the sibling group. Blood feeding behaviors and breeding habitats of the Punctulatus group members have been characterized as unspecialized.6 Distribution of individual sibling species is heterogeneous, displaying unique and occasional overlapping patterns of dispersal.9 In Papua New Guinea (PNG), studies have specifically implicated An. farauti s.s., An. hinesorum, An. farauti 4, An. koliensis, and An. punctulatus s.s. as the primary vectors of malaria and filariasis.7,12 As evidence accumulates to verify the reproductive independence of these sibling species, it becomes increasingly important to collect evidence regarding their species-specific susceptibility to vector control strategies and competence as malaria vectors.

The importance of malaria transmission by An. punctulatus group mosquitoes was first recognized by Heydon and others in the 1920s.13,14 Despite establishing this connection over 90 years ago, mosquito and therefore malaria control in New Guinea has been inconsistently supported and pursued. As malaria casualties during WWII consistently outnumbered those from combat, military activities in New Guinea prompted intense United States and Australian interest in malaria control from 1942 to 1945.15–17 During this time troop protection included military discipline, efforts of malaria control and survey units beginning in 1943, atabrine prophylaxis and treatment of closed environments with “bug bomb” (pyrethrum).15,17 Perhaps the most effective means for controlling mosquito populations, DDT (dichloro-diphenyl-trichloroethane), was not available to troops in the Southwest Pacific area until late 1944, when military operations in New Guinea were drawing to a close.

DDT has been of greatest relevance to mosquito population management in New Guinea because it changed World Health Organization (WHO) strategies from malaria control to eradication.18,19 During the 1940–50s DDT mixed in oil-based solvents and sprayed on larval breeding sites showed short-term effectiveness against malaria transmission.20–22 However, DDT indoor residual spraying (IRS) was associated with reduced exposure to An. punctulatus group mosquitoes in New Guinea23–26 and remained active for 6 months after application. These combined features suggested that DDT IRS would enable deployment of time-efficient “attack” phase strategies by limited numbers of disease control specialists and unskilled workers.19,25,27 These findings encouraged DDT IRS activities from the late 1950s to 1970s27–29 to cover ~50% of the population in 1972.29 Limited WHO insecticide susceptibility tests performed in the early 1970s showed high levels of DDT susceptibility among An. punctulatus group species in surveyed parts of PNG and the Solomon Islands.23,24 In 1972 the WHO Malaria Eradication Program in PNG ceased because of operational failure,23 however DDT IRS remained an important control strategy through provincial government activities.25,26 Further attempts at National Program IRS coverage in the late 1970s appeared to reduce anopheline mosquito and Plasmodium falciparum prevalence, but by 1983 insufficient resources and strained community relations brought this effort to a close.35–37

In the 1980s, the PNG Institute of Medical Research and Ministry of Health began to assess the effect of low-cost bednets. Although the studies were conducted in two different endemic sites north of the PNG Central ranges, results showed that both untreated and permethrin-treated nets (Wosera38,39 and Madang,40 respectively) were associated with protection
from malaria exposure and prevalence. Additionally, studies showed that untreated bednets were associated with reduced transmission of lymphatic filariasis on Bagabag Island north of Madang.41

Presently, the Global Fund to Fight AIDS, Tuberculosis and Malaria (GFATM) has provided support for the distribution of over 2.5 million long-lasting insecticide-treated nets (LLIN; deltamethrin) at no cost to individual families.42 Introduction of LLINs to all of PNG’s 20 Provinces represents the first attempt at a comprehensive plan for mosquito-based malaria control. Of importance, our recently published study observed no phenotypic evidence of resistance to the synthetic pyrethroid insecticides among Anopheles mosquitoes collected at five different PNG malaria-endemic sites.43

In response to intensive insecticide exposure, malaria vectors in other regions of the world have developed resistance to a number of different classes of insecticide compounds. Because several insect species have shown evidence for development of specific polymorphisms in the voltage-gated sodium channel (VGSC) after prolonged exposure to DDT and pyrethroid insecticides,46 a number of studies have focused on sequence encoding the S4–S6 region of VGSC domain II.46,47 In Anopheles mosquitoes, knockdown resistance (kdr) has been associated with an A → T mutation (L1014F)48 or a T → C mutation (L1014S)49 in VGSC. These single nucleotide polymorphisms (SNPs) have been found in African, Indian, and Asian malaria vectors,50–52 and most recently in neighboring Indonesia,53 showing the capacity of these mutations to develop independently in different Anopheles species. Another point mutation in VGSC associated with very high levels of DDT resistance, super-kdr (M918T), has been described in many insects, however this point mutation has not yet been detected in Anopheles.43

To date, Southwest Pacific malaria vectors have not been surveyed for any mutations associated with insecticide resistance. Assuming validity of An. punctulatus sibling species definitions, and the heterogenous history of insecticide application in PNG, we hypothesize that development, prevalence, and distribution of these mutations would occur independently among members of the An. punctulatus group. Here, we have developed a multiplex DNA-based assay to detect knockdown resistance (kdr) polymorphisms and simultaneously identify individual species in the An. punctulatus group by analysis of sequence polymorphisms in the VGSC gene. Results from our study make it possible to evaluate kdr mutations in a species-specific context by a single assay and provide insight into the evolution of VGSC sequence in the An. punctulatus group. Although additional factors can contribute to insecticide resistance and avoidance, population-based studies from other malaria-endemic regions and results presented here may provide important background data on VGSC polymorphism predicting selective pressure of insecticides in PNG.

MATERIALS AND METHODS

Mosquito collections. The Entomology Unit of Papua New Guinea Institute of Medical Research (PNGIMR) collected mosquitoes throughout PNG as part of surveillance studies associated with distribution of LLINs and insecticide susceptibility trials.41 Mosquitoes were collected through landing catch methods, larva collections, and Center for Disease Control light traps as previously described.54,55 Collection sites span 24 villages from seven provinces in PNG and all locations were defined by the global positioning system (Table 1). A subset of these mosquitoes (n = 62) were collected as larvae,

Table 1

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<th>Longitude</th>
<th>Distance to coast**</th>
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<th>F1</th>
<th>H</th>
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* Number of individual mosquitoes included in this study; P = Anopheles punctulatus; F1 = Anopheles farauti s.s.; H = Anopheles hinesorum; F4 = An. farauti 4; K = Anopheles kolensis.
† Villages represented by voltage-gated sodium channel (VGSC) sequence data.
‡ Locations of insecticide susceptibility trials.
** altitude and distance to coast measured in meters.
§ An. longirostris collected in these locations for genetic comparison purposes.
¶ WHP = Western Highlands Province.
and then females reared to adulthood were evaluated by WHO insecticide susceptibility assays by Keven and others.  

**Morphological species identification.** Morphological identification of mosquitoes was completed using methods previously described where members of the An. punctulatus group were classified as An. koliensis, An. farauti s.l. (sensu lato) or An. punctulatus s.l. by morphologically distinguishing characteristics. Collected larvae were allowed to mature in the laboratory to facilitate adult morphological identification. Mosquitoes morphologically identified as members of the An. punctulatus species group (n = 357) were kept in coded vials containing silica gel until DNA extraction could be completed.

**DNA extraction and ITS2 molecular species identification.** Genomic DNA was extracted from single whole mosquitoes and molecular species identification was determined by analyzing polymorphisms within the internal transcribed spacer (ITS2) region of ribosomal DNA (rDNA) using a high-throughput molecular probe method previously described.

**Amplification and sequencing of VGSC.** Polymerase chain reaction (PCR) amplification of the IIS4-IIS6 region of the VGSC gene including introns preceding and following the coding region containing the kdr allele (referred to as intron 1 and intron 2, respectively) was modified from Martinez-Torres and others molecular characterization of VGSC in An. gambiae. The PCR amplification reactions (25 μL) were preformed (67 mM Tris-HCl, pH 8.8, 6.7 mM MgSO4, 16.6 mM (NH4)2SO4, 10 mM 2-mercaptoethanol, 10 μM dATP, dGTP, dCTP, and dTTP, and 2.5 units of thermostable Taq DNA polymerase) using upstream and downstream primers (L) were conducted in a solution containing 20 mM Tris-HCl buffer, pH 7.6, 25 mM potassium acetate, 10 mM magnesium acetate, 1 mM NAD+, 10 mM dithiothreitol, 0.1% Triton X-100, 10 mM (200 pmol) of each LDR probe, 1 μL of each PCR product and 2 units of Taq DNA ligase (New England Biolabs, Beverly, MA). The LDR reactions were initially heated at 95°C for 1 min, followed by 32 thermal cycles of 95°C for 15 sec and 58°C for 2 min. The labeling and detection of LDR products was completed as previously described using Bio-Plex array reader (Bio-Rad Laboratories).

**RESULTS**

Previous analysis of a small number of mitochondrial and nuclear genes has demonstrated that significant genetic polymorphism exists within and between members of the An. punctulatus species group considered to be the most significant vectors of malaria in PNG. Additionally, some polymorphisms within species, namely An. farauti s.s. and An. koliensis, have been shown to be geographically localized. Given these previous findings, and because the VGSC gene has not yet been described, we preformed DNA sequence analysis across a segment of this gene (~1,365 bp) observed to include insecticide resistance polymorphisms in a number of insect species.

**kdr-associated mutations.** The analysis was based on comparative alignment of 90 An. punctulatus group individual’s VGSC sequences where mutations associated with knockdown (kdr) and super knockdown resistance (super-kdr) have previously been identified. This includes sequence encoding the fourth to sixth transmembrane regions of domain II; codons 908 to 1026, including two introns. The occurrence of kdr mutations was evaluated following alignment of 31 alleles of An. punctulatus s.s. (98.5% similar), 14 of An. farauti s.s. (99.4% similar), 13 of An. koliensis (98.1% similar), 12 of An. farauti 4 (99.0% similar), and 20 of An. koliensis (98.4% similar), kdr (L1014F/S), and super-kdr (M918T). Among the surveyed sequences there was no evidence of mutations associated with insecticide resistance. Additional VGSC sequence comparisons were made between An. punctulatus species group members and similar sequence fragments.
MULTIPLEX KDR DETECTION AND SPECIES IDENTIFICATION

from *An. longirostris* (GenBank accession no. HQ173903) co-endemic to PNG, *An. dirus* (DQ026439) Southeast Asia/Malaysia, and *An. gambiae* (Y13592) Africa. *An. longirostris* (GenBank accession no. HQ173903) co-endemic to PNG, *An. dirus* (DQ026439) Southeast Asia/Malaysia, and *An. gambiae* (Y13592) Africa.

**VGSC coding sequence analysis.** Further examination of the coding region (358 bp) revealed 53 variable sites where 56 unique mutations were found. Among these, 20 SNPs were observed repeatedly within and between the *An. punctulatus* sibling species; 36 SNPs were observed only once. Although this error rate (1.08 × 10⁻³; 35 of 32,310 nucleotides) is higher than standard ampliqaq error rates [2.0 × 10⁻⁵ to 2.1 × 10⁻⁴/bp] [70, 71], it was not possible to validate these latter SNPs through repeated observation in this study.

**Repeated synonymous SNPs.** Within the VGSC coding region sequence, 10 SNPs occurred in multiple *An. punctulatus* sibling species (codons 909, 910, 924, 956, 973, 987, 996, 999, 1007, 1026; all occupying the third position of each codon).
999C

Table 3). Codons 909 (G>C) and 910 (G>A) were found to harbor mutations that occurred together and separately in each species with the exception of An. farauti s.s. where the mutations were not observed together. The predominant 909-910 SNPs were observed in separate codons in multiple species (codons 947, 967, 999, 1017). At codon 947 a T>C SNP was observed in one An. hinesorum allele and one An. punctulatus s.s. allele (codon position 1) and resulted in a phenylalanine to leucine amino acid substitution; at 967 an A>G SNP was observed in one An. farauti s.s. allele (codon position 2) and one An. punctulatus s.s. allele and resulted in an asparagine to serine substitution; at 999 a T>C SNP was observed in two An. farauti 4 alleles and one An. farauti s.s. allele (codon position 2) and resulted in a valine to alanine substitution; at 1017 a T>C SNP observed in one An. hinesorum, one An. koliensis, and one An. punctulatus s.s. allele (codon position 2) resulted in a leucine to proline substitution.

### Repeated nonsynonymous SNPs

Coding sequence mutations were observed in separate codons in multiple species (codons 947, 967, 999, 1017). At codon 947 a T>C SNP was observed in one An. hinesorum allele and one An. punctulatus s.s. allele (codon position 1) and resulted in a phenylalanine to leucine amino acid substitution; at 967 an A>G SNP was observed in one An. farauti s.s. allele (codon position 2) and one An. punctulatus s.s. allele and resulted in an asparagine to serine substitution; at 999 a T>C SNP was observed in two An. farauti 4 alleles and one An. farauti s.s. allele (codon position 2) and resulted in a valine to alanine substitution; at 1017 a T>C SNP observed in one An. hinesorum, one An. koliensis, and one An. punctulatus s.s. allele (codon position 2) resulted in a leucine to proline substitution.

### VGSC sequence comparisons outside the An. punctulatus species group

To augment the analysis of sequence variation among the An. punctulatus complex sibling species evaluated here, we performed additional VGSC sequence comparisons.
with similar coding region sequence fragments for An. longirostris (358 bp), An. dirus (213 bp), and An. gambiae (348 bp). Between An. longirostris and the 90 An. punctulatus sibling species alleles we observed five synonymous mutations unique to An. longirostris (954 T>C, 979 C>T, 981 A>C, 989 G>C, and 1012 C>A). Two synonymous mutations in An. longirostris were shared with at least one An. punctulatus species group allele (995 C>T An. punctulatus s.s. [n = 1]; 996 C>T An. punctulatus s.s. [n = 31], An. farauti 4 [n = 14]). One synonymous An. longirostris mutation occurred at the An. punctulatus s.s. SNP site 981 with a unique sequence change A>C compared with A>G. Between a partial An. dirus sequence and the An. punctulatus sibling species alleles we identified eight synonymous mutations unique to An. dirus (978 G>A, 989 G>C [also observed in An. longirostris], 993 T>C, 999 C>G, 1004 T>C, 1012 C>A [also observed in AL], 1013 C>T and 1014 T>C). Six synonymous mutations were observed to be shared among one or more An. punctulatus group members (956 C>T An. hinesorum [n = 2], An. punctulatus s.s. [n = 2]; 973 T>C An. punctulatus s.s. [n = 1], An. koliensis [n = 1]; 980 T>G An. farauti 4 [n = 12]; 981 A>G An. farauti 4 [n = 12], 996 C>T An. farauti 4 [n = 12], An. punctulatus s.s. [n = 31]; and 1026 C>A An. farauti 4 [n = 2], An. hinesorum [n = 5], An. farauti s.s. [n = 6], An. koliensis [n = 8], An. punctulatus s.s. [n = 10]). Finally, between a partial An. gambiae sequence and the An. punctulatus sibling species alleles we identified 19 unique synonymous mutations in An. gambiae (940 G>A, 955 T>G, 963 A>G, 969 G>A, 974 T>C, 979 T>C [also observed in An. longirostris], 982 G>A, 989 G>A, 992 C>T, 999 C>A, 1002 T>A, 1003 T>A, 1004 C>T, 1008 C>T, 1010 A>G, 1012 C>A [also observed in An. longirostris and An. dirus], 1013 C>T [also observed in An. dirus], 1015 G>C, and 1018 T>C). Nine synonymous SNPs were observed to be shared among one or more An. punctulatus group members (956 C>T [also observed in An. dirus] An. hinesorum [n = 2], An. punctulatus s.s. [n = 2]; 962 C>T An. farauti 4 [n = 2]; 981 A>G [also observed in An. dirus] An. farauti 4 [n = 12]; 983 T>C An. punctulatus s.s. [n = 1]; 995 C>T [also observed in An. longirostris] An. punctulatus s.s. [n = 1]; 997 C>T An. farauti 4 [n = 1]; 1000 T>C An. koliensis [n = 1]; 1007 A>C An. farauti s.s. [n = 11], An. punctulatus s.s. [n = 30]; and 1026 C>A [also observed in An. dirus] An. farauti 4 [n = 2], An. hinesorum [n = 5], An. farauti s.s. [n = 6], An. koliensis [n = 8], An. punctulatus s.s. [n = 10]).

Test for evidence of selection. Because the potential exists for this gene to be under selection and since there has been a history of insecticide exposure among these species it becomes necessary to look at the nature of the mutations in the coding regions of the VGSC. Previously, we described 20 SNPs (16 synonymous and 4 nonsynonymous) occurring in the coding region analyzed, which were found in more than one sequence. Seven synonymous mutations (964 A>G An. koliensis n = 20, 980 T>C An. koliensis n = 20, 980 T>G An. farauti 4 n = 12, 981 A>G An. farauti 4 n = 12, 985 A>G An. punctulatus s.s. n = 31, 987 T>C An. punctulatus s.s. n = 31, and An. farauti 4 n = 12) were fixed within at least one An. punctulatus sibling species. The remaining 9 synonymous mutations were found to be polymorphic in one or more of the sibling species (909 G>C An. farauti 4 n = 3, An. farauti s.s. n = 7, An. hinesorum n = 13, An. koliensis n = 4, An. punctulatus s.s. n = 31, 910 G>A An. farauti 4 n = 5, An. farauti s.s. n = 3, An. hinesorum n = 6, An. koliensis n = 6, An. punctulatus s.s. n = 8, 924 G>A An. farauti s.s. n = 1, An. hinesorum n = 1, 956 C>T An. punctulatus s.s. n = 2, An. hinesorum n = 2, 962 C>T An. farauti 4 n = 2, 973 T>C An. farauti s.s. n = 1, An. farauti 4 n = 1, 999 C>T An. koliensis n = 9, An. farauti 4 n = 1, 1007 A>C An. punctulatus s.s. n = 30, An. farauti s.s. n = 11, 1026 C>A An. punctulatus s.s. n = 10, An. koliensis n = 8, An. farauti s.s. n = 6, An. hinesorum n = 5, An. farauti 4 n = 5). All 4 nonsynonymous mutations were found to be polymorphic in one or multiple sibling species (947 T>C An. punctulatus s.s. n = 1, An. hinesorum n = 1, 967 An. punctulatus s.s. n = 1, An. farauti s.s. n = 1, 999 T>C An. farauti s.s. n = 1, An. farauti 4 n = 2, 1017 C>T An. punctulatus s.s. n = 1, An. koliensis n = 1, An. hinesorum n = 1). The assumption of neutrality was tested by the McDonald-Kreitman test\(^2\) using this coding region data. The results were non-significant suggesting that there is no evidence for selection in this region of the gene among the species in this sample. Given that the focus of this analysis was only on a portion of the VGSC gene, it is possible that selection could be occurring elsewhere.

**VGSC intron sequence analysis.** Intron 1 was of variable length within (exception An. farauti s.s.) and between species; An. punctulatus s.s. ranged from 880 to 923 bp, An. farauti s.s. 928 bp, An. hinesorum 939 to 950 bp, An. farauti 4 925 to 938 bp, and An. koliensis 940 to 953 bp. Intron 2 varied in size between species; An. punctulatus s.s. 65 bp, An. farauti s.s. 65 bp, An. hinesorum 68 bp, An. farauti 4 66 bp, and An. koliensis 68 bp. Further unique species-specific insertions and deletions were also observed within introns 1 and 2. These differences are shown in Figure 1 where consensus sequences from each An. punctulatus sibling species of interest were aligned using ClustalW. When the An. punctulatus sibling species consensus sequences were compared with An. longirostris, they were found to be 77–79% similar.

We then analyzed the intronic regions for each species independently (Table 4). For intron 1 (ranging 880–953 bp) all An. farauti s.s. sequences had the greatest pairwise percent identity, 99.5%. High percentage identity is not surprising as all 14 sequences were the same length, leaving differences to be SNPs alone. Pairwise percent identity for the remaining within species comparisons were as follows: An. punctulatus s.s. (98.1%), An. hinesorum (97.8%), An. farauti 4 (97.8%).

### Table 4

<table>
<thead>
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<th>Intron</th>
<th>AP</th>
<th>AFL</th>
<th>AH</th>
<th>AF4</th>
<th>AK</th>
<th>AP Complex</th>
<th>APC + AL()*</th>
<th>% Pairwise identity</th>
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<td>899</td>
<td>899</td>
<td>521</td>
<td>409</td>
<td>409</td>
</tr>
</tbody>
</table>

\* AP = Anopheles punctulatus; AFL = Anopheles farauti s.s.; AH = Anopheles hinesorum; AF4 = An. farauti 4; AK = Anopheles koliensis.
\* Base pairs.
\* APC = An. punctulatus species complex; AL = Anopheles longirostris.
and *An. koliensis* (98.1%). Pairwise percent identity was 88.5% when comparing all *An. punctulatus* complex members. When comparing all *An. punctulatus* sibling species with *An. longirostris* the pairwise percent identity dropped slightly to 87.8%, however when comparing *An. longirostris* individually to a representative of each species, pairwise percent identities were < 69%. Intra- and inter-species complex relationships are further defined by considering the percent of identical sites within each comparison. Intra-species comparisons observed nucleotide identity over the analyzed gene fragment of 84.2% for *An. punctulatus* s.s., 97.2% for *An. farauti* s.s., 91.3% for *An. hinesorum*, 95.5% for *An. farauti* 4, and 91.3% for *An. koliensis*. With inter-species comparisons among *An. punctulatus* sibling species allelic percent identity dropped to 51.7%; 40% when *An. longirostris* was added to this sequence comparison. This further demonstrates that intronic sequences appear to have been conserved within each species and among members of the *An. punctulatus* species group. Given that this gene is possibly under selective pressure and natural selection reduces the genetic variation of organisms we would expect to see little to no variation within this gene within species.

Unlike intron 1, intron 2 was not observed to vary in size within species and is much smaller (65–68 bp). Pairwise percent identity of all the species was found to be greater than 97.7% (*An. hinesorum*) with four species (*An. punctulatus* s.s., *An. farauti* s.s., *An. farauti* 4, and *An. koliensis*) above 99%. Comparison of all *An. punctulatus* species group members showed 90.8% pairwise percent identity and when *An. longirostris* was added to the comparison this percentage dropped slightly to 90.2%. Conserved intron 2 sequences further demonstrated species similarities through analysis of sequence identity. *An. punctulatus* s.s., *An. farauti* s.s., *An. farauti* 4, and *An. koliensis* all have greater than 96.9% identical sites, and *An. hinesorum* has 89.7% identical sites. When comparing all *An. punctulatus* s.s. group members together, there were only 60.6% identical sites; overall sequence identity dropped to 43.7% when *An. longirostris* was added to these comparisons.

**Phylogenetic comparison of *An. punctulatus* sibling species using VGSC sequence.** Given that species-specific polymorphism in both the coding and intronic regions were observed in the multiple species alignment (Figure 1) in addition to the multiple SNPs shared among more than one species (Table 3), we preformed a phylogenetic comparison of the entire sequenced portion of VGSC (1,303–1,379 bp) of the 90 *An. punctulatus* species group members. *Anopheles longirostris* was used as an out-group. Using ClustalW (at default parameters), sequences were aligned and phylogenetic relationships were inferred using the Neighbor-Joining method. The optimal tree with the sum of branch length = 0.58406478 is shown. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) are shown next to the branches. Branches with bootstrap values < 80 were condensed. All positions containing alignment gaps and missing data were eliminated only in pairwise sequence comparisons (Pairwise deletion option). There were a total of 1,430 positions in the final dataset. Phylogenetic analyses were conducted in MEGA5.

![Figure 2](image-url)  
**Figure 2.** Phylogenetic relationships of 90 *Anopheles punctulatus* species complex members. Individual mosquitoes have been described, including GenBank accession numbers for all VGSC sequences. Designations for each mosquito included in this analysis are represented by an abbreviated identifier for each species (e.g., AP), collection site, and unique mosquito number. For example, “AF1_bil_02” is a unique *An. farauti* s.s. representative collected from Bibibil village. *Anopheles longirostris* (AL), a non-*An. punctulatus* species group member, was used as the out-group. The evolutionary history was inferred using the Neighbor-Joining method. The optimal tree with the sum of branch length = 0.58406478 is shown. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) are shown next to the branches. Branches with bootstrap values < 80 were condensed. All positions containing alignment gaps and missing data were eliminated only in pairwise sequence comparisons (Pairwise deletion option). There were a total of 1,430 positions in the final dataset. Phylogenetic analyses were conducted in MEGA5.
We observed that each of the An. punctulatus sibling species formed its own clade with bootstrap support greater than 97, and that no individual mosquito was found in a clade that was different from its initial species identification based on detailed comparisons of ITS2 rDNA sequence polymorphisms. Phylogenetic analyses also observed significant stratification between An. hinesorum mosquitoes collected north and south of the PNG Central Ranges. Overall, our sequence comparisons suggested that the VGSC gene sequence would be a reliable candidate for molecular species identification that could be coupled with probes investigating the kdr-associated SNPs.

**Post-PCR LDR-FMA design and validation.** Conserved regions of the amplified VGSC sequence surrounding both the location of the M918T and L1014F/S mutations afforded the opportunity to build ligase-detection (LDR) probes to monitor for the presence (heterozygous or homozygous) or absence of these mutations in a post-PCR-based LDR-FMA. Given the high bootstrap support for the ability of VGSC to differentiate members of the An. punctulatus group, we targeted species-specific regions of the amplified VGSC region in development of LDR probes to differentiate members of the An. punctulatus group. These probes, coupled with the kdr detection probes, could simultaneously monitor the presence or absence of the kdr mutations and differentiate members of the An. punctulatus species group. Locations where kdr mutation detection and species-specific classification LDR reporter probes hybridized with VGSC template PCR products are shown in the VGSC consensus alignment for An. punctulatus s.s., An. farauti s.s., An. hinesorum, An. farauti 4, and An. koliensis (Figure 1).

The specificity of the assay was demonstrated using probes identified in Table 2 to differentiate PCR products amplified from previously sequenced species-specific controls including two controls, known to harbor the sensitive (TTA) or resistant (TTT or TCA) kdr mutation at amino acid position 1014. As members of the An. punctulatus species group are known to carry Plasmodium, Wuchereria parasites as well as human blood meals, six additional controls were included containing P. falciparum, Plasmodium vivax, Plasmodium malariae, Plasmodium ovale, Wuchereria bancrofti, and human genomic DNA.

Results of control LDR-FMA tests (Figure 3) showed high specificity for kdr allele differentiation (kdr-sensitive [TTA] mean fluorescence intensity [MFI] > 23,000; kdr-resistant [TTT or TCA] < 15,000 MFI; background signals < 3,000 MFI) as each LDR-FMA probe set detected only the kdr allele present in the genomic controls and displayed. Simultaneously, each species-specific LDR-FMA probe set (An. punctulatus s.s. MFI > 21,000; An. koliensis > 23,000; An. farauti s.s. > 23,000; An. hinesorum > 10,500; An. farauti 4 > 22,000), detected only the species expected; all species probe background signals were below MFIs of 2,000. Results showed that the LDR probes designed to target the species-specific polymorphisms described here detected only the species present and no others. Fluorescent signals from the parasite and human samples were below a background of 160 MFI (data not shown), showing that the presence of parasites and human blood meal would not obscure Anopheles species identification or kdr mutation detection.

**Application of VGSC multiplex assay: analysis of mosquito collections for kdr mutation and species determination.** The VGSC multiplex assay was then used to evaluate a sample collection of 312 mosquitoes for the kdr mutation and species identification. All 312 mosquitoes assayed were determined to be homozygous for the sensitive kdr allele (TTA/TTA). Results in Table 5 summarize the comparison of species identification using the VGSC multiplex assay to a similar molecular classification method that probes differences in An. punctulatus sibling species ITS2 rDNA sequences. One discordant sample was detected among the 312 mosquitoes assayed; by VGSC LDR-FMA this mosquito was identified as belonging to the An. punctulatus species group.
as *An. punctulatus* s.s., where ITS2 LDR-FMA species were identified as *An. farauti* s.s. These results show 99.7% concordance between the two species identification methods while simultaneously identifying the *kdr* mutation status in a large sample set of field-collected mosquitoes.

Mosquitoes that were evaluated for insecticide susceptibility using WHO insecticide bioassays (as described previously in Keven and others[4]) were also analyzed in this assay. Sixty-two mosquitoes, collected from five geographically distinct areas (Table 1) were previously tested and reported to be susceptible to either 0.05% lambda-cyhalothrin-treated filter paper (n = 26, locations: Dimer, Peneng, and Lorengau) or new 55 mg/m² deltamethrin-treated LLIN (n = 36, locations: Ramu [Madjad, Naru, and Dimer]. These mosquitoes were evaluated using the *VGSC* LDR-FMA. All 62 samples gave a strong signal for the presence of the susceptible genotype (TTA; average MFI 8,182; signals for TTT and TCA probes fell within expected background [< 3,000 MFI]) showing that these samples were both phenotypically susceptible and lacked the L1014F/S mutations associated with *kdr*.

**DISCUSSION**

Here, we have characterized DNA sequence polymorphism in a segment of the *VGSC* gene (~1,350 bp) among sibling species of the *An. punctulatus* group known to be significant vectors of human malaria and filariasis in Papua New Guinea.[7,12] This region of the *VGSC* gene encodes the fourth to sixth transmembrane regions (domain II) of the *VGSC* protein,[40] and is known to harbor mutations associated with knockdown resistance to pyrethroids and DDT in many insect species (*Anopheles*,[44,52,53,55,66,74–79] *Culex*,[60,65] *Musca domestica* [House fly],[82,83] *Blattella germanica* [German cockroach]).[84,85]

In this characterization of *VGSC* genetic variation we were able to determine that the well-known *kdr*-associated mutations (L1014F; S918T)[86] were not observed among the extant mosquitoes of the *An. punctulatus* species group. Although our collections were not comprehensive across PNG, individual mosquitoes were available at multiple locations along the PNG north coast where transmission of malaria and filariasis is intense. Because many of these collection sites occurred in regions that have historically been the focus of DDT larviciding applications[22,27] and IRS programs,[25,29] it is possible that ancestral *An. punctulatus* species group populations have been exposed to positive selection that would favor the *kdr*-associated mutations. This included mosquito collection sites (Table 1) in six mainland Provinces and Manus Island Province. The greatest distance between sample collection sites within each species ranged from 85 km for *An. farauti* 4 to 883 km for *An. farauti* s.s., with all species (except for *An. farauti* 4) having collections spanning at least 580 km in distance. For this study *An. farauti* 4 collections were limited to pockets north of the Central Ranges. Significant geographical barriers also separated collection sites for *An. farauti* s.s. (island locations) and *A. hinesorum* (locations north and south of the Central Ranges). Interestingly, although our phylogenetic analysis provided evidence for significant population stratification within *A. hinesorum* collected north and south of the PNG Central Ranges, significant *VGSC* sequence variation was not observe between *An. farauti* s.s. mosquitoes collected between the PNG mainland and Manus Island.

Furthermore, the overall sequence analysis revealed a wide range of synonymous and nonsynonymous SNP and insertion/deletion polymorphisms, the latter exclusively within “so-called” introns 1 and 2. The additional coding region SNPs enabled us to perform population genetic comparisons to evaluate evidence for selective pressure on this gene sequence and sequence-based relationships within and between sibling species. McDonald-Kreitman testing,[72] based on the observation of 16 synonymous versus 4 non-synonymous SNPs in the *VGSC* coding region analyzed here indicated that there was no evidence for selection in this gene segment. However, it may be important to note that these observations have been based on mosquitoes collected over the past 5–10 years during a time when no concerted insecticide dispersal program has been under implementation in PNG. Therefore, it is not possible to rule out the potential that the *kdr*-associated mutations arose in the past and have now drifted out of current *An. punctulatus* sibling species populations.

Given results of the McDonald-Kreitman test, reservations about using *VGSC* sequences to perform phylogenetic analyses may be reduced. The Neighbor-Joining analysis reported in Figure 2 shows very strong conservation of sequence within each species cluster and no evidence for gene flow between species. Because many of the polymorphisms characterizing the different species included consistently observed insertions and deletions, our results suggested that it might be possible to develop highly specific DNA probes for species identification. Consistent with these phylogenetic results (apart from one sample), DNA probes based on *VGSC* sequence polymorphisms produced species identification results consistent with earlier ITS2 analyses for over 300 mosquitoes collected across our widely distributed sampling sites; expanded surveys will be necessary to verify that these results can be consistently reproduced throughout the Southwest Pacific region where the *An. punctulatus* sibling species are endemic. Although our resulting LDR-FMA multiplex assay did not detect any evidence of *kdr*-associated mutations from any of the PNG study sites surveyed, our analyses revealed an important feature of *VGSC* sequence evolution and an important potential challenge for mosquito control. Because our results provide evidence for independent acquisition and maintenance of *VGSC* sequence variation, *kdr*-associated mutations in this gene sequence would also be expected to emerge independently within each species. Therefore, whether *VGSC* *kdr*-associated mutations or mutations in other genes lead to insecticide resistance, assays must be developed to monitor each of the *An. punctulatus* sibling species with equal efficiency, specificity, and sensitivity. Therefore, molecular diagnostic assay developed here, offers advantages over

### Table 5

Concordance assessment of molecular ITS2 and VGSC classification methods for *Anopheles punctulatus* sibling species

<table>
<thead>
<tr>
<th>ITS2</th>
<th>AP</th>
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<th>AF1</th>
<th>AF2</th>
<th>AF4</th>
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<td>-</td>
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<td>AF4</td>
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</table>

*AP = Anopheles punctulatus s.s.; AF = Anopheles farauti s.s.; AH = Anopheles hinesorum AF4 = An. farauti 4; AK = Anopheles kolensis.*

VGSC = voltage-gated sodium channel; LDR = ligase detection reaction; FMA = fluorescent microscopy assay.
other PCR-based species identification strategies by que-
rying insecticde resistance associated and species-specific
polymorphisms in a single multiplex test. Currently, coun-
try-wide planning and distribution of LLINs is underway in the
Solomon Islands\(^{34,37}\) and Papua New Guinea.\(^{42,43}\) A recent
report describing the discovery of the VGSC L1014F
kd\(_r\) mutation in populations of Anopheles from neighboring
Indonesia suggests that An. punctulatus sibling species in this
same region\(^{35}\) may be experiencing insecticide exposure that
will select for kdr-associated mutations.

Integrated vector management strategies require a means
to continuously monitor and assess the vector species compo-
sition in an endemic region and elements of the vector’s biol-
ogy. Although kdr is a concern, it is important to understand
this phenotype within the context of additional gene sequence
and gene expression polymorphism, physiological and behav-
ioral heterogeneity. Because insecticide resistance can develop
through many different routes\(^2\) it is becoming necessary to
broaden evaluation of insecticide resistance beyond candidate
gene-based investigations as demonstrated by recent genomic
comparison studies of An. gambiae.\(^{49,90}\)

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