The genetic architecture of target-site

resistance to pyrethroid insecticides in the

African malaria vectors Anopheles gambiae

and Anopheles coluzzii

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Genomes Consortium⁵

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16 Abstract

Resistance to pyrethroid insecticides is a major concern for malaria vector control because these are the compounds used in almost all insecticide-treated bed-nets (ITNs), and are also widely used for indoor residual spraying (IRS). Pyrethroids target the voltage-gated sodium channel (VGSC), an essential component of the mosquito nervous system, but substitutions in the amino acid sequence can disrupt the activity of these insecticides, inducing a resistance phenotype. Here we use Illumina whole-genome sequence data from phase 2 of the *Anopheles gambiae* 1000 Genomes Project (Ag1000G) to provide a comprehensive account of genetic variation in the

Vgsc gene in mosquito populations from 13 African countries. In addition to the three known kdr resistance alleles, we describe 20 non-synonymous nucleotide substitutions (at appreciable frequency in one or more populations) that are previously unknown in Anopheles mosquitoes and we mapped these variants on a molecular model of the protein. Thirteen of these novel alleles were found to occur almost exclusively on haplotypes carrying the known L995F resistance allele (L1014F in Musca domesticus codon numbering), and may enhance or compensate for the L995F resistance phenotype. A novel mutation I1527T, which is adjacent to a predicted pyrethroid binding site, was found in tight linkage with either of two alleles causing a V402L substitution, similar to a combination of substitutions found to cause pyrethroid resistance in several other insect species. We analyse the genetic backgrounds on which non-synonymous alleles are found, to determine which alleles have experienced recent positive selection, and to refine our understanding of the spread of resistance between species and geographical locations. We describe ten distinct kdr carrying haplotype groups with evidence of recent positive selection, five of which carry the known L995F resistance allele, five of which carry the known L995S resistance allele. Five of these groups are localised to a single geographical location, and five comprise haplotypes from different countries, in one case separated by over 3000 km, providing new information about the geographical distribution and spread of resistance. Two "non-kdr" haplotype groups with evidence of recent selection were also detected, one of which carries the novel I1527T allele, and one of which carries a novel M490I allele. We also find evidence for multiple introgression events transmitting resistance alleles between An. qambiae and An. coluzzii. Markers are identified that could be used to design high-throughput, low-cost genetic assays for improved surveillance of pyrethroid resistance in the field. Our results demonstrate that the molecular basis of target-site pyrethroid resistance in malaria vectors is more complex than previously appreciated, and provide a foundation for the development of new genetic tools to track the spread insecticide resistance and improve the design of strategies for insecticide resistance management.

Introduction

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Pyrethroid insecticides have been the cornerstone of malaria prevention in Africa for almost two decades [1]. Pyrethroids are currently used in all insecticide-treated bed-nets (ITNs), and are widely used in indoor residual spraying (IRS) campaigns as well as in agriculture. Resistance to these insecticides is now widespread in malaria vector populations across
Africa [2]. The World Health Organization (WHO) has published plans for insecticide
resistance management (IRM) that emphasise the need for improvements in both our
knowledge of the molecular mechanisms of resistance and our ability to monitor them in
natural populations [3, 4].

The voltage-gated sodium channel (VGSC) is the physiological target of pyrethroid insecticides, and is integral to the insect nervous system. The sodium channel protein consists of four homologous domains (DI-IV) each of which comprises six transmembrane segments (S1-S6) connected by intracellular and extracellular loops [5]. Pyrethroid molecules
bind to this protein, stabilise the ion-conducting active state and thus disrupt normal
nervous system function, producing paralysis ("knock-down") and death. However, amino
acid substitutions at key positions within the protein alter the interaction with insecticide
molecules, increasing the dose of insecticide required for knock-down (target-site resistance), and leading to this type of resistance to also be known as knock-down resistance
or kdr [6, 5].

In the African malaria vectors Anopheles gambiae and An. coluzzii, three substitutions
have been found to cause pyrethroid resistance. Two of these substitutions occur in codon
995¹, with L995F prevalent in West and Central Africa [7, 8], and L995S found in Central
and East Africa [9, 8]. A third substitution, N1570Y, has been found in West and Central
Africa and shown to increase resistance in association with L995F [11]. However, studies in
other insect species have found a variety of other Vgsc substitutions inducing a resistance
phenotype [12, 13, 5]. To our knowledge, no studies in malaria vectors have analysed
genetic variation across the full Vgsc coding sequence, thus the molecular basis of targetsite resistance to pyrethroids has not been fully explored.

Basic information is also lacking about the spread of pyrethroid resistance in malaria vectors [3]. For example, it is not clear when, where or how many times pyrethroid target-site resistance has emerged. Geographical paths of transmission, carrying resistance alleles between mosquito populations, are also not known. Previous studies have found evidence that L995F occurs on several different genetic backgrounds, suggesting multiple

¹Codon numbering is given here relative to transcript AGAP004707-RD as defined in the AgamP4.12 geneset annotations. A mapping of codon numbers from AGAP004707-RD to *Musca domestica*, the system in which *kdr* mutations were first described [10], is given in Table 1.

independent outbreaks of resistance driven by this allele [14, 15, 16, 17]. However, these studies analysed only small gene regions in a limited number of mosquito populations, and 87 therefore had limited resolution to make inferences about relationships between haplotypes carrying this allele. It has also been shown that the L995F allele spread from An. gambiae to An. coluzzii in West Africa [18, 19, 20, 21]. However, both L995F and L995S now have wide geographical distributions [8], and to our knowledge no attempts have been made to 91 infer or track the geographical spread of either allele across Africa. 92 Here we report an in-depth analysis of genetic variation in the Vgsc gene, using whole-93 genome Illumina sequence data from phase 2 of the Anopheles gambiae 1000 Genomes Project (Ag1000G) [22]. The Ag1000G phase 2 resource includes data on nucleotide variation in 1,142 wild-caught mosquitoes sampled from 13 countries, with representation of West, Central, Southern and East Africa, and of both An. qambiae and An. coluzzii. We investigate variation across the complete gene coding sequence, and report population genetic data for both known and novel non-synonymous nucleotide substitutions. We then use haplotype data from the chromosomal region spanning the Vqsc gene to study 100 the genetic backgrounds carrying resistance alleles, infer the geographical spread of resis-101 tance between mosquito populations, and provide evidence for recent positive selection. 102 Finally, we explore ways in which variation data from Ag1000G can be used to design 103 high-throughput, low-cost genetic assays for surveillance of pyrethroid resistance, with 104 the capability to differentiate and track resistance outbreaks. 105

6 Results

Vgsc non-synonymous nucleotide variation

To identify variants with a potentially functional role in pyrethroid resistance, we extracted single nucleotide polymorphisms (SNPs) that alter the amino acid sequence of
the VGSC protein, from the Ag1000G phase 2 data resource [22]. We then computed
their allele frequencies among 16 mosquito populations defined by species and country of
origin. Alleles that confer resistance are expected to increase in frequency under selective
pressure, therefore we filtered the list of potentially functional variant alleles to retain
only those at or above 5% frequency in one or more populations (Table 1). The resulting

list comprises 23 variant alleles, including the known L995F, L995S and N1570Y resistance alleles, and a further 20 alleles which prior to Ag1000G had not previously been described in anopheline mosquitoes. We reported 12 of these novel alleles in our overall analysis of the 765 samples in the Ag1000G phase 1 data resource [23], and we extend the analyses here to incorporate SNPs which alter codon 531, 697, 1507, 1603 and two tri-allelic SNPs affecting codons 402 and 490 in the 1,142 phase 2 samples.

The 23 non-synonymous variants were located on a transmembrane topology map and on 121 a 3-dimensional homology model of the *Vgsc* protein. (Figure 1). The substitutions were 122 found to be distributed throughout the channel, in all of the four internally homologous domains (DI-DIV), in S1, S5 and S6 membrane-spanning segments, in two of the intracel-124 lular loops connecting domains and in the C-terminal tail. The S5 and S6 segments that 125 form the central ion-conducting pore of the channel carry six of the eight segment substi-126 tutions, including V402 and L995 which have been shown to produce insecticide resistance 127 phenotypes [6, 5, 7, 8, 9]. Two substitutions are located on the DIII-DIV linker including the resistance conferring N1570 [11]. A further six substitutions are found concentrated 129 in the protein's carboxyl tail (C-terminus), including two alternative substitutions at the 130 resistance associated P1874 residue [24]. The DIII-DIV linker and the C-terminus seg-131 ment interact in the closed-state channel and substitutions are found throughout this 132 intracellular subdomain. Finally, there are four novel substitutions located on the DI-DII 133 intracellular linker but this region is missing from the model as it was not resolved in the 134 cockroach Na_vPaS structure used as the model template [25]. 135

The two known resistance alleles affecting codon 995 had the highest overall allele fre-136 quencies within the Ag1000G phase 1 cohort (Table 1). The L995F allele was at high 137 frequency in populations of both species from West, Central and Southern Africa. The 138 L995S allele was at high frequency among An. qambiae populations from Central and East Africa. Both of these alleles were present in An. qambiae populations sampled from 140 Cameroon and Gabon. This included individuals with a heterozygous L995F/S genotype 141 (50/297 individuals in Cameroon, 41/69 in Gabon). We calculated empirical p-values for 142 these heterozygous genotype counts using the Dirichlet distribution and 1,000,000 Monte 143 Carlo simulations. In Cameroon p=0.410 of simulations found higher proportions of heterozygous genotypes, however in Gabon this dropped to p=0.005, hinting there may be a

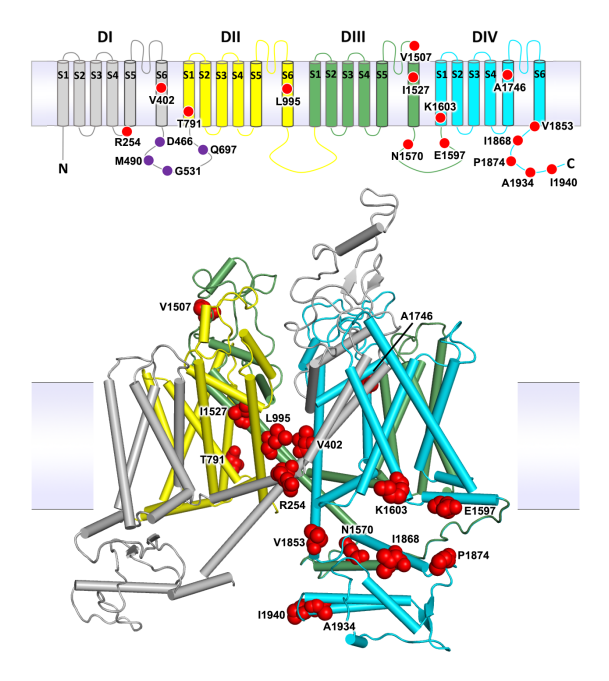


Figure 1. Voltage-gated sodium channel protein structure and non-synonymous variation. The An. gambiae voltage-gated sodium channel (AGAP004707-RD AgamP4.12) is shown as a transmembrane topology map (top) and as a homology model (bottom) in cartoon format coloured by domain. Variant positions are shown as red circles in the topology map and as red space-fill in the 3D model. Purple circles in the map show amino acids absent from the model due to the lack of modelled structure in this region.

Table 1. Non-synonymous nucleotide variation in the voltage-gated sodium channel gene. AO=Angola; GH=Ghana; BF=Burkina Faso; CI=Côte d'Ivoire; GN=Guinea; GW=Guinea-Bissau; GM=Gambia; CM=Cameroon; GA=Gabon; UG=Uganda; GQ=Bioko; FR=Mayotte; KE=Kenya; Ac=An. coluzzii; Ag=An. gambiae. Species status of specimens from Guinea-Bissau, Gambia and Kenya is uncertain [22]. All variants are at 5% frequency or above in one or more of the 16 Ag1000G phase 2 populations, with the exception of 2,400,071 G>T which is only found in the CMAg population at 0.3% frequency but is included because another mutation is found at the same position (2,400,071 G>A) at >5% frequency and which causes the same amino acid substitution (M4901).

Variant				Population allele frequency (%)															
Position ¹	Ag^2	Md^3	Domain ⁴	AOAc	$\mathrm{GH}Ac$	BFAc	CIAc	GNAc	GW	GM	CMAg	$\mathrm{GH}Ag$	BFAg	GNAg	GAAg	UGAg	GQAg	FRAg	KE
2,390,177 G>A	R254K	R261	IL45	0.0	0.009	0.0	0.0	0.0	0.0	0.0	0.313	0.0	0.0	0.0	0.203	0.0	0.0	0.0	0.0
2,391,228 G>C	V402L	V410	IS6	0.0	0.127	0.073	0.085	0.125	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
2,391,228 G>T	V402L	V410	IS6	0.0	0.045	0.06	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
2,399,997 G>C	D466H	-	LI/II	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.069	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
2,400,071 G>A	M490I	M508	LI/II	0.0	0.0	0.0	0.0	0.0	0.0	0.031	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.188
2,400,071 G>T	M490I	M508	LI/II	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.003	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
2,402,466 G>T	G531V	G549	LI/II	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.007	0.0	0.056	0.0	0.0
2,407,967 A>C	Q697P	Q724	LI/II	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.056	0.0	0.0
2,416,980 C>T	T791M	T810	IIS1	0.0	0.009	0.02	0.0	0.0	0.0	0.0	0.0	0.292	0.147	0.112	0.0	0.0	0.0	0.0	0.0
2,422,651 T>C	L995S	L1014	IIS6	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.157	0.0	0.0	0.0	0.674	1.0	0.0	0.0	0.76
2,422,652 A>T	L995F	L1014	IIS6	0.84	0.818	0.853	0.915	0.875	0.0	0.0	0.525	1.0	1.0	1.0	0.326	0.0	0.0	0.0	0.0
2,429,556 G>A	V1507I	-	IIIL56	0.0	0.0	0.0	0.0	0.125	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
2,429,617 T>C	I1527T	I1532	IIIS6	0.0	0.173	0.133	0.085	0.125	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
2,429,745 A>T	N1570Y	N1575	LIII/IV	0.0	0.0	0.267	0.0	0.0	0.0	0.0	0.057	0.167	0.207	0.088	0.0	0.0	0.0	0.0	0.0
2,429,897 A>G	E1597G	E1602	LIII/IV	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.065	0.062	0.0	0.0	0.0	0.0	0.0
2,429,915 A>C	K1603T	K1608	IVS1	0.0	0.055	0.047	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
2,430,424 G>T	A1746S	A1751	IVS5	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.292	0.141	0.1	0.0	0.0	0.0	0.0	0.0
2,430,817 G>A	V1853I	V1858	COOH	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.542	0.049	0.062	0.0	0.0	0.0	0.0	0.0
2,430,863 T>C	I1868T	I1873	COOH	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.261	0.2	0.0	0.0	0.0	0.0	0.0
2,430,880 C>T	P1874S	P1879	COOH	0.0	0.027	0.207	0.345	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
2,430,881 C>T	P1874L	P1879	COOH	0.0	0.0	0.073	0.007	0.25	0.0	0.0	0.0	0.0	0.234	0.475	0.0	0.0	0.0	0.0	0.0
2,431,061 C>T	A1934V	A1939	COOH	0.0	0.018	0.107	0.465	0.5	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
2,431,079 T>C	I1940T	I1945	COOH	0.0	0.118	0.04	0.0	0.0	0.0	0.0	0.067	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

¹ Position relative to the AgamP3 reference sequence, chromosome arm 2L.

² Codon numbering according to *Anopheles gambiae* transcript AGAP004707-RD in geneset AgamP4.12.

 $^{^3}$ Codon numbering according to $\it Musca~domestica~EMBL~accession~X96668~[10].$

⁴ Location of the variant within the protein structure. Transmembrane segments are named according to domain number (in Roman numerals) followed by 'S' then the number of the segment; e.g., 'IIS6' means domain two, transmembrane segment six. Internal linkers between segments within the same domain are named according to domain (in Roman numerals) followed by 'L' then the numbers of the linked segments; e.g., 'IL45' means domain one, linker between transmembrane segments four and five. Internal linkers between domains are named 'L' followed by the linked domains; e.g., 'LI/II' means the linker between domains one and two. 'COOH' means the internal carboxyl tail.

146 fitness advantage for mosquitoes carrying both alleles in some circumstances.

The N1570Y allele was present in Guinea, Burkina Faso (both species) and Cameroon. 147 This allele has been shown to substantially increase pyrethroid resistance when it occurs 148 in combination with L995F, both in association tests of phenotyped field samples [11] and functional tests using Xenopus oocytes [26]. To study the patterns of association 150 among non-synonymous variants, we used haplotypes from the Ag1000G phase 2 resource 151 to compute the normalised coefficient of linkage disequilibrium (D') between all pairs of 152 variant alleles (Figure 2). As expected, we found N1570Y in almost perfect linkage with 153 L995F. Of the 20 novel non-synonymous alleles, 13 also occurred almost exclusively in combination with L995F (Figure 2). These included two variants in codon 1874 (P1874S, 155 P1874L), one of which (P1874S) has previously been associated with pyrethroid resistance 156 in the crop pest moth Plutella xylostella [24]. 157

The abundance of high-frequency non-synonymous variants occurring in combination 158 with L995F is striking for two reasons. First, Vgsc is a highly conserved gene, expected to be under strong functional constraint and therefore purifying selection, so any non-160 synonymous variants are expected to be rare [12]. Second, in contrast with L995F, we did 161 not observe any high-frequency non-synonymous variants occurring in combination with 162 L995S. This contrast was clear when data on all variants within the gene were considered: 163 for haplotypes carrying the L995 allele, the ratio of non-synonymous to synonymous nu-164 cleotide diversity (π_N/π_S) was 20.04 times higher than haplotypes carrying the wild-type 165 allele, but for those carrying L995S (π_N/π_S) was 0.5 times lower than haplotypes carrying 166 the wild-type allele. These results may indicate that L995F has substantially altered the 167 selective regime for other amino acid positions within the protein, perhaps through relax-168 ation of purifying selection. Secondary substitutions have occurred and risen in frequency, 169 suggesting that they are providing some selective advantage in the presence of insecticide pressure. 171

A novel allele, I1527T, was present in An. coluzzii from Burkina Faso at 13% frequency. Codon 1527 occurs within trans-membrane segment IIIS6, immediately adjacent to residues within a predicted binding site for pyrethroid molecules, thus it is plausible that I1527T could alter pyrethroid binding [27, 5]. We also found that the two variant alleles affecting codon 402, both of which induce a V402L substitution, were in strong linkage with I1527T ($D' \geq 0.8$; Figure 2), and almost all haplotypes carrying I1527T also carried a V402L substitution. Substitutions in codon 402 have been found in a number of other

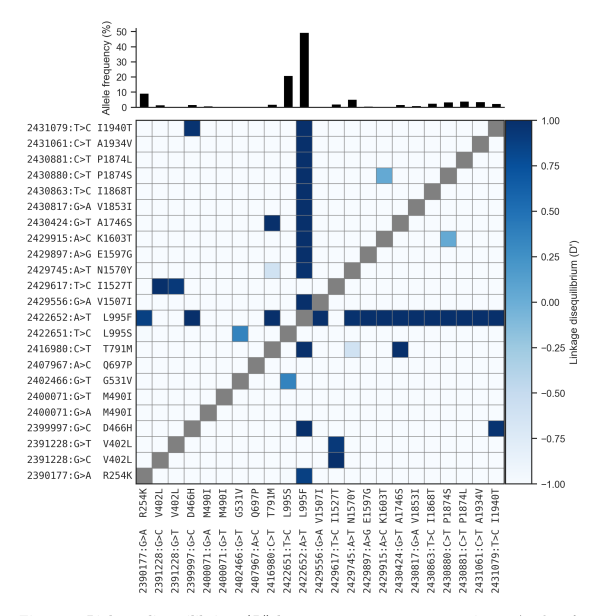


Figure 2. Linkage disequilibrium (D') between non-synonymous variants. A value of 1 indicates that two alleles are in perfect linkage, meaning that one of the alleles is only ever found in combination with the other. Conversely, a value of -1 indicates that two alleles are never found in combination with each other. The bar plot at the top shows the frequency of each allele within the Ag1000G phase 2 cohort. See Table 1 for population allele frequencies.

insect species and shown experimentally to confer pyrethroid resistance [5]. Because of the limited geographical distribution of these alleles, we hypothesize that the I1527T+V402L combination could represent a pyrethroid resistance allele that arose in West African An. coluzzii populations prior to L995F. However, the L995F allele is at higher frequency (85%) in our Burkina Faso An. coluzzii population, and is known to be increasing in frequency [28]. Therefore, L995F may provide a stronger resistance phenotype or be less deleteri-

ous in the absence of pyrethroids [29], and could be replacing I1527T+V402L. The four remaining novel alleles, Q697P, G531V and two separate nucleotide substitutions causing M490I, did not occur in combination with any known resistance allele and were private to a single population (except a single haplotype carrying G531V from Bioko) (Table 1), and to our knowledge, none have previously been found in other species [13, 5].

The Ag1000G data resource provides a rich source of information about the spread of

190 Genetic backgrounds carrying resistance alleles

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insecticide resistance alleles in any given gene, because data are not only available for 192 SNPs in protein coding regions, but also SNPs in introns, flanking intergenic regions, 193 and in neighbouring genes. These additional variants can be used to analyse the genetic backgrounds (haplotypes) on which resistance alleles are found. In our initial report of 195 the Ag1000G phase 1 resource [23], we used 1710 biallelic SNPs from within the 73.5 kbp 196 Vgsc gene (1607 intronic, 103 exonic) to compute the number of SNP differences between 197 all pairs of 1530 haplotypes derived from 765 wild-caught mosquitoes. We then used 198 pairwise genetic distances to perform hierarchical clustering, and found that haplotypes carrying resistance alleles in codon 995 were grouped into 10 distinct clusters, each with 200 near-identical haplotypes. Five of these clusters contained haplotypes carrying the L995F 201 allele (labelled F1-F5), and a further five clusters contained haplotypes carrying L995S 202 (labelled S1-S5). 203 To further investigate genetic backgrounds carrying resistance alleles, we used the 204 Ag1000G phase 2 haplotype data from the Vqsc gene (2,284 haplotypes from 1,142 mosquitoes 205 [22]), to construct median-joining networks [30] (Figure 3). The network analysis improves 206 on hierarchical clustering by allowing for the reconstruction and placement of intermedi-207 ate haplotypes that may not be observed in the data. It also allows for non-hierarchical 208 relationships between haplotypes, which may arise if recombination events have occurred 209 between haplotypes. We constructed the network up to a maximum edge distance of 2 SNP 210 differences, to ensure that each connected component captures a group of closely-related 211 haplotypes. The resulting network contained 5 groups containing haplotypes carrying 212 L995F, and a further 5 groups carrying L995S, in close correspondence with previous results from hierarchical clustering (96.8% overall concordance in assignment of haplotypes

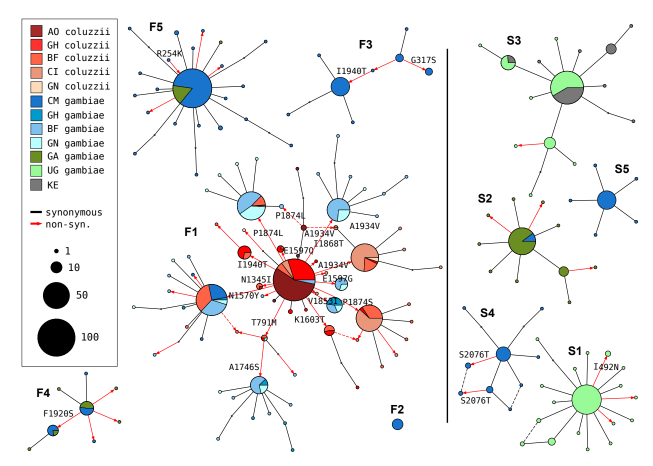


Figure 3. Haplotype networks. Median joining network for haplotypes carrying L995F (labelled F1-F5) or L995S variants (S1-S5) with a maximum edge distance of two SNPs. Labelling of network components is via concordance with hierarchical clusters discovered in [23]. Node size is relative to the number of haplotypes contained and node colour represents the proportion of haplotypes from mosquito populations/species - AO=Angola; GH=Ghana, BF=Burkina Faso; CI=Côte d'Ivoire; GN=Guinea; CM=Cameroon; GA=Gabon; UG=Uganda; KE=Kenya. Non-synonymous edges are highlighted in red and those leading to non-singleton nodes are labelled with the codon change, arrow head indicates direction of change away from the reference allele. Network components with fewer than three haplotypes are not shown.

to groups).

The haplotype network brings into sharp relief the explosive radiation of amino acid substitutions secondary to the L995F allele (Figure 3). Within the F1 group, nodes carrying
non-synonymous variants radiate out from a central node carrying only L995F, suggesting that the central node represents the ancestral haplotype carrying just L995F which
initially came under selection, and these secondary variants have arisen subsequently as
new mutations. In F1 alone, 30 network edges (shown as red arrows - Figure 3) lead to
non-synonymous nodes. Many of the nodes carrying secondary variants are large, consistent with positive selection and a functional role for these secondary variants as modifiers

of the L995F resistance phenotype. The F1 network also allows us to infer multiple intro-224 gression events between the two species. The central (putatively ancestral) node contains 225 haplotypes from individuals of both species, as do nodes carrying the N1570Y, P1874L and 226 T791M variants. This structure is consistent with an initial introgression of the ancestral F1 haplotype, followed later by introgressions of haplotypes carrying secondary mutations. 228 The haplotype network also illustrates the constrasting levels of non-synonymous varia-229 tion between L995F and L995S. Within all of the L995S groups, only eight edges lead to 230 non-synonymous nodes and all these nodes are small (low frequency variants), thus may 231 be neutral or mildly deleterious variants that are hitch-hiking on selective sweeps for the L995S allele. 233

The F1 group contains haplotypes from mosquitoes of both species, and from mosquitoes 234 sampled in six different countries (Angola, Burkina Faso, Cameroon, Côte d'Ivoire, Ghana, 235 Guinea) (Figure 4). The F4, F5 and S2 groups each contain haplotypes from both 236 Cameroon and Gabon. The S3 group contains haplotypes from both Uganda and Kenya. 237 The haplotypes within each of these five groups (F1, F4, F5, S2, S3) were nearly identi-238 cal across the entire span of the Vgsc gene $(\pi < 4.5 \times 10^{-5} \ bp^{-1})$. In contrast, diversity 239 among wild-type haplotypes was two orders of magnitude greater (Cameroon An. gambiae 240 $\pi = 1.4 \times 10^{-3} \ bp^{-1}$; Guinea-Bissau $\pi = 5.7 \times 10^{-3} \ bp^{-1}$). Thus it is reasonable to assume 241 that each of these five groups contains descendants of an ancestral haplotype that carried 242 a resistance allele and has risen in frequency due to selection for insecticide resistance. Given this assumption, these groups each provide evidence for adaptive gene flow between 244 mosquito populations separated by considerable geographical distances. 245

A limitation of both the hierarchical clustering and network analyses is that they rely 246 on genetic distances within a fixed genomic window from the start to the end of the 247 Vasc gene. Anopheles mosquitoes undergo homologous recombination during meiosis in 248 both males and females, and any recombination events that occurred within this genomic 249 window could affect the way that haplotypes are grouped together in clusters or network 250 components. In particular, recombination events could occur during the geographical 251 spread of a resistance allele, altering the genetic background upstream and/or downstream 252 of the allele itself. An analysis based on a fixed genomic window might then fail to infer gene flow between two mosquito populations, because haplotypes with and without a 254

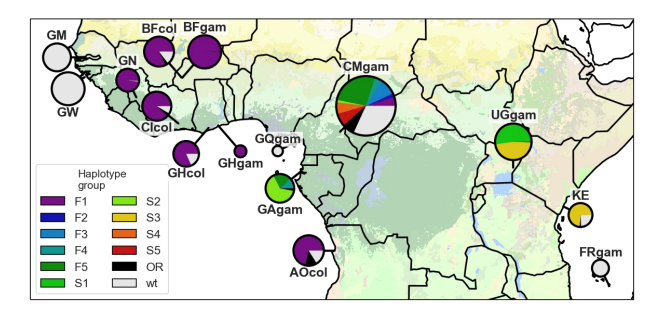


Figure 4. Map of haplotype frequencies. Each pie shows the frequency of different haplotype groups within one of the populations sampled. The size of the pie is proportional to the number of haplotypes sampled. The size of each wedge within the pie is proportional to the frequency of a haplotype group within the population. Haplotypes in groups F1-5 carry the L995F kdr allele. Haplotypes in group other resistant (OR) carry either L995F or L995S but did not cluster within any of the haplotype groups. Wild-type (wt) haplotypes do not carry any known resistance alleles.

recombination event could be grouped separately, despite the fact that they share a recent 255 common ancestor. To investigate the possibility that recombination events may have 256 affected our grouping of haplotypes carrying resistance alleles, we performed a moving 257 window analysis of haplotype homozygosity, spanning Vgsc and up to a megabase upstream 258 and downstream of the gene (Supplementary Figures S1, S2). This analysis supported a 259 refinement of our initial grouping of haplotypes carrying resistance alleles. All haplotypes 260 within groups S4 and S5 were effectively identical on both the upstream and downstream 261 flanks of the gene, but there was a region of divergence within the *Vgsc* gene itself that separated them in the fixed window analyses (Supplementary Figure S2). The 13.8 kbp 263 region of divergence occurred upstream of codon 995 and contained 6 SNPs that were fixed 264 differences between S4 and S5. A possible explanation for this short region of divergence 265 is that a gene conversion event has occurred within the gene, bringing a segment from 266 a different genetic background onto the original genetic background on which the L995S resistance mutation occurred. 268

269 Positive selection for resistance alleles

To investigate evidence for positive selection on non-synonymous alleles, we performed 270 an analysis of extended haplotype homozygosity (EHH) [31]. Haplotypes under recent 271 positive selection will have increased rapidly in frequency, thus have had less time to be 272 broken down by recombination, and should on average have longer regions of haplotype 273 homozygosity relative to wild-type haplotypes. We defined a core region spanning Vgsc codon 995 and an additional 6 kbp of flanking sequence, which was the minimum required 275 to differentiate the haplotype groups identified via clustering and network analyses. Within 276 this core region, we found 18 distinct haplotypes at a frequency above 1% within the cohort. 277 These included core haplotypes corresponding to each of the 10 haplotype groups carrying 278 L995F or L995S alleles identified above, as well as a core haplotype carrying I1527T which we labelled L1 (due to it carrying the the wild-type leucine codon at position 995). We also 280 found a core haplotype corresponding to a group of haplotypes from Kenya carrying an 281 M490I allele, which we labelled as L2. All other core haplotypes we labelled as wild-type 282 (wt). We then computed EHH decay for each core haplotype up to a megabase upstream 283 and downstream of the core locus (Figure 5). 284 As expected, haplotypes carrying the L995F and L995S resistance alleles all experience 285 a dramatically slower decay of EHH relative to wild-type haplotypes, supporting positive 286 selection. Previous studies have found evidence for different rates of EHH decay between 287 L995F and L995S haplotypes, suggesting differences in the timing and/or strength of selec-288 tion [16]. However, we found no systematic difference in the length of shared haplotypes 289 when comparing F1-5 (carrying L995F) against S1-5 (carrying L995S) (Supplementary 290 Figure S3). There were, however, some differences between core haplotypes carrying the same allele. For example, shared haplotypes were significantly longer for S1 (median 1.006 292 cM, 95% CI [0.986 - 1.040]) versus other core haplotypes carrying L995S (e.g., S2 median 293 0.593 cM, 95% CI [0.589 - 0.623]; Supplementary Figure S3). Longer shared haplotypes in-294 dicate a more recent common ancestor, and thus some of these core haplotypes may have 295 experienced more recent and/or more intense selection than others. The L1 haplotype carrying I1527T+V402L exhibited a slow decay of EHH on the downstream flank of the 297

gene, similar to haplotypes carrying L995F and L995S, indicating that this combination

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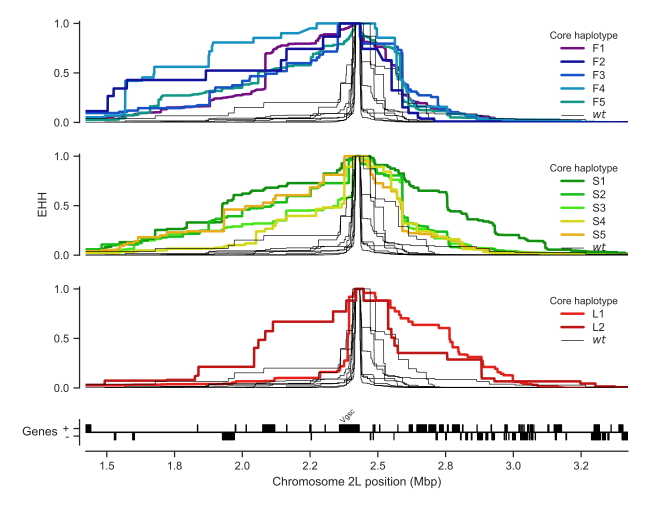


Figure 5. Evidence for positive selection on haplotypes carrying known or putative resistance alleles. Each panel plots the decay of extended haplotype homozygosity (EHH) for a set of core haplotypes centred on Vgsc codon 995. Core haplotypes F1-F5 carry the L995F allele; S1-S5 carry the L995S allele; L1 carries the I1527T allele; L2 carries the M490I allele. Wild-type (wt) haplotypes do not carry known or putative resistance alleles. A slower decay of EHH relative to wild-type haplotypes implies positive selection (each panel plots the same collection of wild-type haplotypes).

of alleles has experienced positive selection. EHH decay on the upstream gene flank was faster, being similar to wild-type haplotypes, however there were two separate nucleotide 300 substitutions encoding V402L within this group of haplotypes, and a faster EHH decay 301 on this flank is consistent with recombination events bringing V402L alleles from differ-302 ent genetic backgrounds together with an ancestral haplotype carrying I1527T. The L2 303 haplotype carrying M490I exhibited EHH decay on both flanks comparable to haplotypes 304 carrying known resistance alleles. This could indicate evidence for selection on the M490I 305 allele, however these haplotypes are derived from a Kenyan mosquito population where 306 there is evidence for a severe recent bottleneck [23], and there were not enough wild-type 307

haplotypes from Kenya with which to compare, thus this signal may also be due to the extreme demographic history of this population.

310 Discussion

311 Cross-resistance between pyrethroids and DDT

The VGSC protein is the physiological target of both pyrethroid insecticides and DDT [6]. 312 The L995F and L995S alleles are known to increase resistance to both of these insecticide 313 classes [7, 9]. By 2012, over half of African households owned at least one pyrethroid 314 impregnated ITN and nearly two thirds of IRS programmes were using pyrethroids [2]. 315 Pyrethroids were also introduced into agriculture in Africa prior to the scale-up of public health vector control programmes, and continue to be used on a variety of crops such as 317 cotton [32]. DDT was used in Africa for several pilot IRS projects carried out during the 318 first global campaign to eradicate malaria, during the 1950s and 1960s [12]. DDT is still 319 approved for IRS use by WHO and remains in use in some locations, however within the 320 last two decades pyrethroid use has been far more common and widespread. DDT was also 321 used in agriculture from the 1940s, and although agricultural usage has greatly diminished 322 since the 1970s, some usage remains [33]. In this study we reported evidence of positive 323 selection on the L995F and L995S alleles, as well as the I1527T+V402L combination and 324 possibly M490I. We also found 14 other non-synonymous substitutions that have arisen in 325 association with L995F and appear to be positively selected. Given that pyrethroids have 326 dominated public health insecticide use for two decades, it is reasonable to assume that the 327 selection pressure on these alleles is primarily due to pyrethroids rather than DDT. It has 328 previously been suggested that L995S may have been initially selected by DDT usage [16]. 329 However, we did not find any systematic difference in the extent of haplotype homozygosity 330 between these two alleles, suggesting that both alleles have been under selection over a 331 similar time frame. We did find some significant differences in haplotype homozygosity between different genetic backgrounds carrying resistance alleles, suggesting differences 333 in the timing and/or strength of selection these may have experienced. However, there 334 have been differences in the scale-up of pyrethroid-based interventions in different regions, 335 and this could in turn generate heterogeneities in selection pressures. Nevertheless, it is 336

possible that some if not all of the alleles we have reported provide some level of crossresistance to DDT as well as pyrethroids, and we cannot exclude the possibility that
earlier DDT usage may have contributed at least in part to their selection. The differing
of resistance profiles to the two types of pyrethroids (type I, e.g., permethrin; and type
II, e.g., deltamethrin) [34], will also affect the selection landscape. Further sampling and
analysis will be required to investigate the timing of different selection events and relate
these to historical patterns of insecticide use in different regions.

Resistance phenotypes for novel non-synonymous variants

The non-synonymous variants are distributed throughout the channel protein but can be considered in terms of three clusters: (i) the transmembrane domain, (ii) the DI-II intracellular linker and (iii) the DIII-DIV/C-terminal subdomain. The pyrethroid binding site is located in the transmembrane domain between the IIS4-S5 linker and the IIS5, 348 IIS6 and IIIS6 helices [35]. The I1527T substitution that we discovered in An. coluzzii 349 mosquitoes from Burkina Faso occurs in segment IIIS6 and is immediately adjacent to 350 two pyrethroid-sensing residues in this binding site [5]. It is thus plausible that pyrethroid binding could be altered by this substitution. The I1527T substitution (M. domestica 352 codon 1532) has been found in Aedes albopictus [36], and substitutions in the nearby 353 codon 1529 (M. domestica codon 1534) have been reported in Aedes albopictus and in 354 Aedes aegypti where it was found to be associated with pyrethroid resistance [5, 37, 38]. 355 We found the I1527T allele in tight linkage with two alleles causing a V402L substitution 356 (M. domestica codon 410). Substitutions in codon 402 have been found in multiple insect 357 species and are by themselves sufficient to confer pyrethroid resistance [5]. The fact that 358 we find I1527T and V402L in such tight mutual association is intriguing because haplotypes 359 carrying V402L alone should also have been positively selected and thus be present in one 360 or more populations. 361 The V402 residue is located towards the middle of the IS6 helix. The L995F and L995S 362 substitutions occur at a similar position on the IIS6 helix. It was proposed these S6 sub-363 stitutions confer resistance by allosterically modifying formation of the pyrethroid binding 364 site [35]. More recently the L995 kdr residue was speculated to form part of a second 365 pyrethroid binding site in the insect channel termed 'PyR2' [27, 39]. A major functional

effect of the L995F substitution is enhanced closed-state inactivation [40]. This contributes 367 to kdr resistance by reducing the number of channels that undergo activation, which is 368 the functional state that pyrethroids bind to with highest affinity [40]. Fast inactivation 369 involves movement of the DIV domain to form a receptor for the DIII-DIV linker fast inactivation particle containing the 'MFM' sequence motif (equivalent to the 'IFM' motif in 371 mammals) [41, 5]. Recent eukaryotic sodium channel structures reveal that the DIII-DIV 372 linker is in complex with the C-terminal segment in the closed-state conformation but the 373 DIII-DIV linker appears to dissociate and bind in close proximity in the DIV S6 helix upon transition to the inactivated state [25, 42]. It seems that binding of the DIII-DIV linker pushes the DIV S6 helix forward to occlude the pore and produce the inactivated state 376 [42]. We suggest that substitutions located on the DIII-DIV linker and C-terminal tail may 377 perturb the conformation of this subdomain when it assembles in the closed-state channel 378 and may subsequently affect capture or release of the DIII-DIV linker from this complex. 379 The expected functional outcome would be altered channel inactivation, although whether 380 inactivation is enhanced or diminished and if this compensates for a deleterious effect of 381 L995F on channel function awaits elucidation. The N1570Y substitution on the DIII-DIV 382 linker has been functionally characterised but inactivation kinetics in the mutant channel 383 were found unaltered [26]. Pyrethroid sensitivity was also unaffected by N1570Y although 384 resistance was greatly enhanced in the N1570Y + L995F double mutant [26]. 385

The final cluster of novel variants is located on the DI-DII intracellular linker. This 386 segment includes the novel M490I substitution that was found on the Kenyan L2 haplotypic 387 background potentially under selection. M490I did not occur in association with L995F or 388 any other non-synonymous substitutions. Although we were unable to model this region, 389 we speculate that the DI-DII linker passes under the DII S4-S5 linker and these regions 390 may interact, as was found in a bacterial sodium channel structure [43]. The structural 391 effects of DI-DII substitutions may be altered interactions with the DII S4-S5 linker, the 392 movement of which is critical for formation of the pyrethroid binding site [35, 44]. Overall, 393 there are a number of potential mechanisms by which a pyrethroid resistance phenotype 394 may arise and topology modelling reveals how many of the non-synonymous variants we 395 discover may be involved, though clearly much remains to be unravelled regarding the molecular biology of pyrethroid resistance in this channel.

Design of genetic assays for surveillance of pyrethroid resistance

Entomological surveillance teams in Africa regularly genotype mosquitoes for resistance al-399 leles in Vqsc codon 995, and use those results as an indicator for the presence of pyrethroid 400 resistance alongside results from insecticide resistance bioassays. They typically do not, 401 however, sequence the gene or genotype any other polymorphisms within the gene. Thus, 402 if there are other polymorphisms within the gene that cause or significantly enhance pyrethroid resistance, these will not be detected. Also, if a codon 995 resistance allele is observed, there is no way to know whether the allele is on a genetic background that 405 has also been observed in other mosquito populations, and thus no way to investigate 406 whether resistance alleles are emerging locally or being imported from elsewhere. Whole-407 genome sequencing of individual mosquitoes clearly provides data of sufficient resolution to 408 answer these questions, and could be used to provide ongoing resistance surveillance. The 409 cost of whole-genome sequencing continues to fall, with the present cost being approxi-410 mately 50 GBP to obtain ~30× coverage of an individual Anopheles mosquito genome with 411 150 bp paired-end reads. However, to achieve substantial spatial and temporal coverage 412 of mosquito populations, it is currently cheaper and more practical to develop targeted 413 genetic assays for resistance outbreak surveillance. Technologies such as amplicon sequencing [45] are already being trialled on mosquitoes [46], these could scale to tens of 415 thousands of samples at low cost and could be implemented using existing platforms in 416 national molecular biology facilities. 417

To facilitate the development of targeted genetic assays for surveillance of Vqsc-mediated 418 pyrethroid resistance, we have produced several supplementary data tables. In Supple-419 mentary Table 1 we list all 82 non-synonymous variants found within the Vgsc gene in this study, with population allele frequencies. In Supplementary Table 2 we list 756 biallelic 421 SNPs, within the Vqsc gene and up to 10 kbp upstream or downstream, that are poten-422 tially informative regarding which haplotype group a resistance haplotype belongs to, and 423 thus could be used for tracking the spread of resistance. This table includes the allele 424 frequency within each of the 12 haplotype groups defined here, to aid in identifying SNPs that are highly differentiated between two or more haplotype groups. We also provide 426 Supplementary Table 3 which lists all 10,244 SNPs found within the Vqsc gene and up to 427

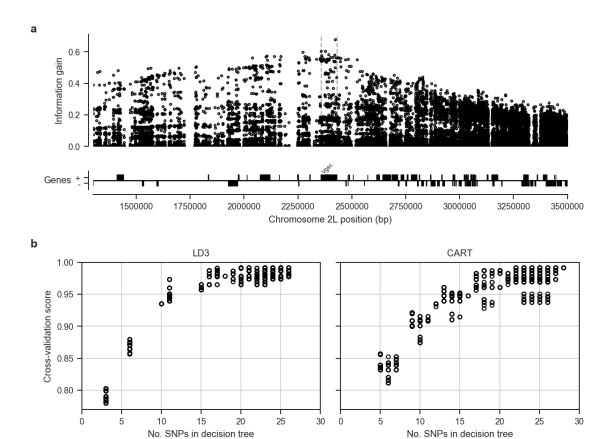


Figure 6. Informative SNPs for haplotype surveillance. a, Each data point represents a single SNP. The information gain value for each SNP provides an indication of how informative the SNP is likely to be if used as part of a genetic assay for testing whether a mosquito carries a resistance haplotype, and if so, which haplotype group it belongs to. b, Number of SNPs required to accurately predict which group a resistance haplotype belongs to. Each data point represents a single decision tree. Decision trees were constructed using either the LD3 (left) or CART (right) algorithm for comparison. Accuracy was evaluated using 10-fold stratified cross-validation.

10 kbp upstream or downstream, which might need to be taken into account as flanking 428 variation when searching for PCR primers to amplify a SNP of interest. To provide some 429 indication for how many SNPs would need to be assayed in order to track the spread of 430 resistance, we used haplotype data from this study to construct decision trees that could 431 classify which of the 12 groups a given haplotype belongs to (Figure 6). This analysis 432 suggested that it should be possible to construct a decision tree able to classify haplo-433 types with >95% accuracy by using 20 SNPs or less. In practice, more SNPs would be 434 needed, to provide some redundancy, and also to type non-synonymous polymorphisms in 435 addition to identifying the genetic background. However, it is still likely to be well within 436 the number of SNPs that could be assayed in a single multiplex via amplicon sequencing. 437 Thus it should be feasible to produce low-cost, high-throughput genetic assays for tracking 438

the spread of pyrethroid resistance. If combined with a limited amount of whole-genome sequencing at sentinel sites, this should also allow the identification of newly emerging resistance outbreaks.

442 Methods

443 Code

All scripts and Jupyter Notebooks used to generate analyses, figures and tables are available from the GitHub repository https://github.com/malariagen/ag1000g-phase2-vgsc-report.

446 Data

We used variant calls and phased haplotype data from the Ag1000G Phase 2 AR1 data release (https://www.malariagen.net/data/ag1000g-phase-2-ar1). Variant calls from
Ag1000G Phase 2 are also available from the European Nucleotide Archive (ENA; http:
//www.ebi.ac.uk/ena) under study PRJEB36277.

Data collection and processing

For detailed information on Ag1000G WGS sample collection, sequencing, variant call-452 ing, quality control and phasing, see [23, 22]. In brief, An. gambiae and An. coluzzii 453 mosquitoes were collected from 33 sites in 13 countries across Sub-Saharan Africa: Angola, Bioko, Burkina Faso, Cameroon, Côte d'Ivoire, Gabon, The Gambia, Ghana, Guinea, 455 Guinea Bissau, Kenya, Mayotte and Uganda. From Angola and Côte d'Ivoire just An. 456 coluzzii were sampled, Burkina Faso, Ghana and Guinea had samples of both An. gambiae 457 and An. coluzzii and all other populations consisted of purely An. gambiae, except for 458 The Gambia, Guinea Bissau and Kenya where species status is uncertain [22]. Mosquitoes were individually whole genome sequenced on the Illumina HiSeq 2000 platform, gener-460 ating 100bp paired-end reads. Sequence reads were aligned to the An. qambiae AgamP3 461 reference genome assembly [47]. Aligned bam files underwent improvement, before variants 462 were called using GATK UnifiedGenotyper. Quality control included removal of samples 463 with mean coverage $\leq 14x$ and filtering of variants with attributes that were correlated 464 with Mendelian error in genetic crosses. 465

The Ag1000G variant data was functionally annotated using the SnpEff v4.1b software 466 [48]. Non-synonymous Vqsc variants were identified as all variants in AgamP4.12 transcript 467 AGAP004707-RD with a SnpEff annotation of "missense". The Vgsc gene is known to 468 exhibit alternative splicing [6], however at the time of writing the An. gambiae gene annotations did not include the alternative transcripts reported by Davies et al. We wrote 470 a Python script to check for the presence of variants that are synonymous according to 471 transcript AGAP004707-RD but non-synonymous according to one of the other transcripts 472 present in the gene annotations or in the set reported by Davies et al. Supplementary Table 473 1 includes the predicted effect for all SNPs that are non-synonymous in one or more of these transcripts. None of the variants that are non-synonymous in a transcript other 475 than AGAP004707-RD were found to be above 5% frequency in any population. 476 For ease of comparison with previous work on Vgsc, pan Insecta, in Table 1 and Supple-477 mentary Table 1 we report codon numbering for both An. gambiae and Musca domestica 478 (the species in which the gene was first discovered). The M. domestica Vgsc sequence (EMBL accession X96668 [10]) was aligned with the An. qambiae AGAP004707-RD se-480 quence (AgamP4.12 gene-set) using the Mega v7 software package [49]. A map of equiva-481 lent codon numbers between the two species for the entire gene can be download from the 482 MalariaGEN website (https://www.malariagen.net/sites/default/files/content/ 483 blogs/domestica_gambiae_map.txt). 484 Haplotypes for each chromosome of each sample were estimated (phased) using using 485 phase informative reads (PIRs) and SHAPEIT2 v2.r837 [50], see [23] supplementary text 486 for more details. The SHAPEIT2 algorithm is unable to phase multi-allelic positions, 487 therefore the two multi-allelic non-synonymous SNPs within the Vasc gene, altering codons 488 V402 and M490, were phased onto the biallelic haplotype scaffold using MVNcall v1.0 [51]. 489 Lewontin's D' [52] was used to compute the linkage disequilibrium (LD) between all pairs

492 Haplotype networks

491

of non-synonymous Vgsc mutations.

Haplotype networks were constructed using the median-joining algorithm [30] as implemented in a Python module available from https://github.com/malariagen/ag1000g-phase2-vgsc-repo
Haplotypes carrying either L995F or L995S mutations were analysed with a maximum edge

distance of two SNPs. Networks were rendered with the Graphviz library and a composite figure constructed using Inkscape. Non-synonymous edges were highlighted using the SnpEff annotations [48].

499 Positive selection

Core haplotypes were defined on a 6,078 bp region spanning Vqsc codon 995, from chromosome arm 2L position 2,420,443 and ending at position 2,426,521. This region was 501 chosen as it was the smallest region sufficient to differentiate between the ten genetic 502 backgrounds carrying either of the known resistance alleles L995F or L995S. Extended 503 haplotype homozygosity (EHH) was computed for all core haplotypes as described in [31] 504 using scikit-allel version 1.1.9 [53], excluding non-synonymous and singleton SNPs. Analyses of haplotype homozygosity in moving windows (Supplementary Figs. S1, S2) and pair-506 wise haplotype sharing (Supplementary Figure S3) were performed using custom Python 507 code available from https://github.com/malariagen/ag1000g-phase2-vgsc-report. 508

Design of genetic assays for surveillance of pyrethroid resistance

To explore the feasibility of indentifying a small subset of SNPs that would be sufficient 510 to identify each of the genetic backgrounds carrying known or putative resistance alleles, we started with an input data set of all SNPs within the Vasc gene or in the flanking 512 regions 20 kbp upstream and downstream of the gene. Each of the 2,284 haplotypes in 513 the Ag1000G Phase 2 cohort was labelled according to which core haplotype it carried, 514 combining all core haplotypes not carrying known or putative resistance alleles together as 515 a single "wild-type" group. Decision tree classifiers were then constructed using scikit-learn version 0.19.0 [54] for a range of maximum depths, repeating the tree construction process 10 times for each maximum depth with a different initial random state. The classification 518 accuracy of each tree was evaluated using stratified 5-fold cross-validation. 519

Homology modelling

A homology model of the *An. gambiae* voltage-gated sodium channel (AGAP004707-RD AgamP4.12) was generated using the 3.8 Å resolution structure of the *Periplaneta americana* sodium channel Na_vPaS structure (PDB code 5X0M) [25]. Sequences were aligned

using Clustal Omega [55]. 50 starting models were generated using MODELLER [56].

The internal scoring function of MODELLER was used to select 10 models, which were
visually inspected and submitted to the VADAR webserver [57] to assess stereochemistry
in order to select the best final model. Figures were produced using PyMOL (DeLano
Scientific, San Carlos, CA, USA).

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Supplementary figures

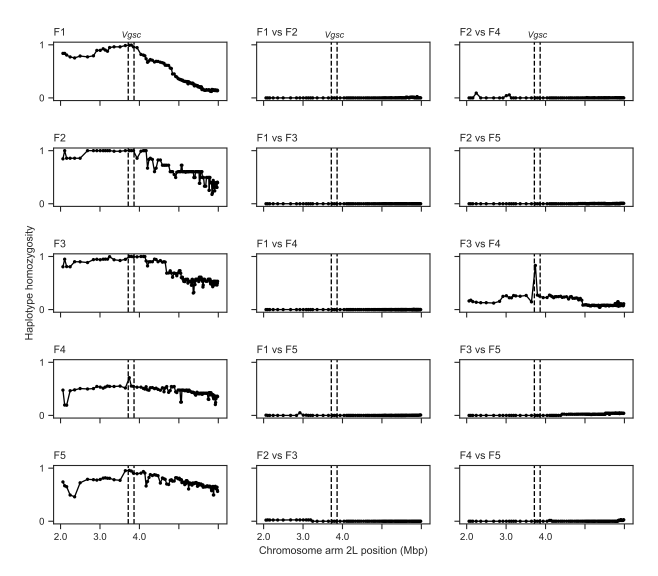


Figure S1. Windowed analysis of haplotype homozygosity for genetic backgrounds carrying the L995F allele. Each sub-plot shows the fraction of haplotype pairs that are identical within half-overlapping moving windows of 1000 SNPs. Each sub-plot in the left-hand column shows homozygosity for haplotype pairs within one of the haplotype groups identified by the network analysis. Sub-plots in the central and right-hand columns show homozygosity for haplotype pairs between two haplotype groups. If two haplotype groups are truly unrelated, haplotype homozygosity between them should be close to zero across the whole genome region. Dashed vertical lines show the location of the *Vgsc* gene.

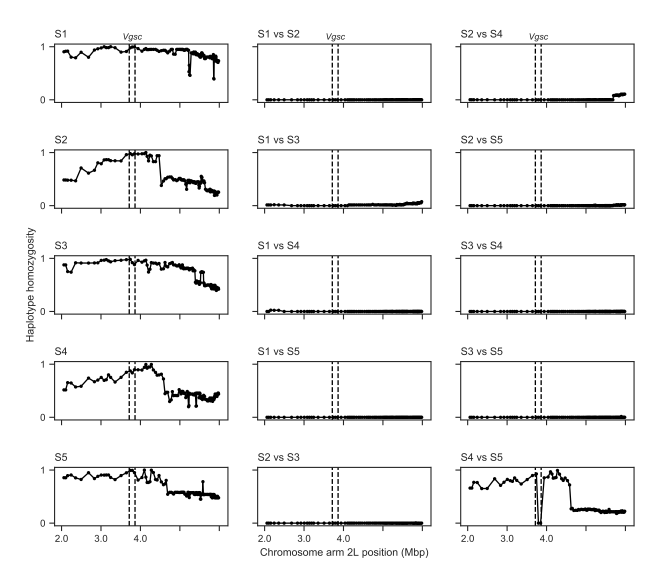


Figure S2. Windowed analysis of haplotype homozygosity for genetic backgrounds carrying the L995S allele. See Supplementary Figure S1 for explanation. Haplotype homozygosity is high between groups S4 and S5 on both flanks of the gene, indicating that haplotypes from both groups are in fact closely related.

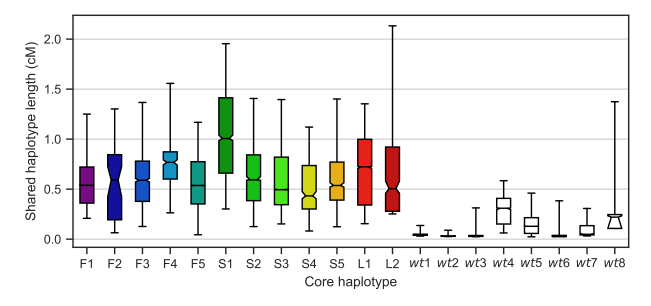


Figure S3. Shared haplotype length. Each bar shows the distribution of shared haplotype lengths between all pairs of haplotypes with the same core haplotype. For each pair of haplotypes, the shared haplotype length is computed as the region extending upstream and downstream from the core locus (*Vgsc* codon 995) over which haplotypes are identical at all non-singleton variants. The *Vgsc* gene sits on the border of pericentromeric heterochromatin and euchromatin, and we assume different recombination rates in upstream and downstream regions. The shared haplotype length is expressed in centiMorgans (cM) assuming a constant recombination rate of 2.0 cM/Mb on the downstream (euchromatin) flank and 0.6 cM/Mb on the upstream (heterochromatin) flank. Bars show the inter-quartile range, fliers show the 5-95th percentiles, horizontal black line shows the median, notch in bar shows the 95% bootstrap confidence interval for the median. Haplotypes F1-5 each carry the L995F resistance allele. Haplotypes S1-5 each carry the L995S resistance allele. Haplotype L2 carries the M490I allele. Wild-type (*wt*) haplotypes do not carry any known or putative resistance alleles.