# Outbreaks of target-site resistance to pyrethroid insecticides in the African malaria vectors *Anopheles gambiae* and *Anopheles*coluzzii

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<sup>4</sup>https://www.malariagen.net/projects/ag1000g#people

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14 Abstract

Resistance to pyrethroid insecticides is a major concern for malaria vector control, because these are the only compounds approved for use in insecticide-treated bed-nets (ITNs). 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 1 of the Anopheles qambiae 1000 Genomes Project (Ag1000G) to provide a comprehensive account of genetic variation at the Vqsc locus in mosquito populations from 8 African countries. In addition to three known resistance variants, we describe 20 non-synonymous variants at appreciable frequency in one or more populations that are previously unknown in mosquitoes. For each variant we predict a resistance phenotype based on genetic evidence for positive selection, patterns of linkage between variants, and functional evidence from other species. We then analyse the genetic backgrounds on which resistance variants are found, to refine our understanding of the origins and spread of resistance between species and geographical locations. We identify ten distinct outbreaks of resistance, of which five appear to be localised to a single geographical location, and five have spread between two or more countries. The most successful and widespread outbreak (F1) originates in West Africa and has subsequently spread to countries in Central and Southern Africa. Our results demonstrate that the molecular basis of pyrethroid resistance in African malaria vectors is more complex than previously appreciated, and provide a foundation for the design of new genetic tools for outbreak surveillance to inform insecticide resistance management and track the further spread of resistance.

# **Introduction**

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- <sup>39</sup> Pyrethroid insecticides are currently the cornerstone of malaria prevention in Africa [1].
- 40 Pyrethroids continue to be the only approved class of insecticide for use in insecticide-
- treated bed-nets (ITNs), and are widely used in indoor residual spraying (IRS) campaigns
- as well as in agriculture. Pyrethroid resistance is, however, now widespread in malaria vec-
- 43 tor populations across Africa [2]. The World Health Organisation (WHO) has published
- 44 plans for insecticide resistance management (IRM), which highlight the need for improve-
- 45 ments in our ability to monitor resistance, and for improvements in our understanding of
- the molecular mechanisms of resistance [3].
- The voltage-gated sodium channel (VGSC) is the physiological target of pyrethroid in-
- 48 secticides, and is integral to the insect nervous system. Pyrethroid molecules bind to sites
- within the protein channel and prevent normal nerve function, causing paralysis ("knock-
- down") and then 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) [4]. In the African malaria vectors Anopheles 52 gambiae and An. coluzzii, three substitutions have been found to cause pyrethroid resis-53 tance. Two of these substitutions occur in codon 995<sup>1</sup>, with L995F prevalent in West and Central Africa [5, 6], and L995S found in Central and East Africa [7, 6]. A third variant, N1570Y, was found in Central Africa and shown to increase resistance in association with L995F [9]. However, studies in other insect species have found a variety of other Vqsc57 substitutions inducing a resistance phenotype [10, 11, 12]. To our knowledge, no studies (prior to Ag1000g [13]) in malaria vectors have analysed the full Vgsc coding sequence, thus the genetic basis of target-site resistance to pyrethroids has not been fully explored. 60 Basic information is also lacking about the history and epidemiology of pyrethroid re-61 sistance in malaria vectors. For example, it is not known when, where or how many times 62 VGSC mediated pyrethroid resistance has emerged. The paths of transmission carrying 63 resistance between mosquito populations are also not known. Previous studies have found evidence that L995F occurs on several different genetic backgrounds, suggesting multiple independent outbreaks of resistance driven by this allele [14, 15, 16]. However, these studies analysed only a small region of the VGSC gene, and therefore had limited power to 67 make inferences about the origins or spread of resistance alleles. It has also been shown 68 that the L995F allele spread from An. qambiae to An. coluzzii in West Africa [17, 18]. However, both L995F and L995S now have wide geographical distributions [6], and no attempts have been made to reconstruct the geographical spread of either allele. If insecticide resistance were a disease, standard methods of outbreak investigation could be 72 applied, and information about epidemiological origins, transmission and virulence fac-73 tors would be used to formulate an outbreak response plan. In the absence of analogous information for pyrethroid resistance, planning an effective response is clearly difficult. Here we report an in-depth analysis of the VGSC gene, using whole-genome Illumina se-76 quence data from phase 1 of the Anopheles gambiae 1000 Genomes Project (Ag1000G) [13]. 77 We investigate variation across the complete gene coding sequence, to fully characterise 78 the primary and secondary genetic factors driving target-site resistance to pyrethroids 79

<sup>&</sup>lt;sup>1</sup>Codon numbering is given here relative to transcript AGAP004707-RA as defined in the AgamP4.4 gene annotations. A mapping of codon numbers from AGAP004707-RA to *Musca domestica*, the system in which the *kdr* mutations were first discovered [8], is given in Table 1 and in @@Supplementary data.

in natural mosquito populations. We then use haplotype data from the chromosomal 80 region spanning the VGSC gene to study the genetic backgrounds carrying resistance al-81 leles. The goal of these analyses is to diagnose how many separate outbreaks of target-site pyrethroid resistance have occurred, which outbreaks are localised, and which are spreading. We also explore ways in which variation data from Ag1000G could be used to design high-throughput, low-cost genetic assays for monitoring pyrethroid resistance, with the capability to differentiate and track separate resistance outbreaks. Finally, we investigate 86 the potential of these data to reconstruct the path of transmission of resistance alleles be-87 tween mosquito populations, and to provide information on the probable source. Although the geographical and temporal sampling of mosquito populations in Ag1000G phase 1 is too sparse to support a comprehensive outbreak analysis, our aim is to investigate methods that could provide answers to these questions, given further sequencing of mosquito 91 populations.

### Results

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#### **Functional variation**

To identify variants with a potentially functional role in pyrethroid resistance, we extracted single nucleotide polymorphisms (SNPs) from the Ag1000G phase 1 data resource that 96 alter the amino acid sequence of the VGSC protein, and computed their allele frequencies among 9 populations defined by species and country of origin. Alleles that confer resistance are expected to increase in frequency under selective pressure, and we refined the list of potentially functional variant alleles to retain only those at an appreciable frequency 100 (>5%) in one or more populations (Table 1). The resulting list comprises 23 variant alleles, 101 including the known L995F, L995S and N1570Y variants, and a further 20 not previously 102 described in these species. We reported 15 of these novel alleles in our initial analysis 103 of the Ag1000G phase 1 data [13], and we extend the analyses here to incorporate two tri-allelic SNPs affecting codons 402 and 490 and a SNP altering codon 1603. 105 The two alleles in codon 995 are clearly the main drivers of resistance at this locus. 106 The L995F allele at high frequency in populations of both species from West, Central and 107 Southern Africa, and the L995S allele at high frequency among An. gambiae populations

Table 1. Non-synonymous nucleotide variation in the voltage-gated sodium channel gene. AO=Angola; BF=Burkina Faso; GN=Guinea; CM=Cameroon; GA=Gabon; UG=Uganda; KE=Kenya; GW=Guinea-Bissau; Ac=An. coluzzii; Ag=An. gambiae. All variants are at 5% frequency or above in one or more of the 9 Ag1000G phase 1 populations, with the exception of 2,400,071 G>T which is only found in the CMAg population at 0.4% frequency but is included because another mutation (2,400,071 G>A) is found at the same position causing the same amino acid substitution (M490I); and 2,431,019 T>C (F1920S) which is at 4% frequency in GAAg but also found in CMAg and linked to L995F.

Var	Population allele frequency (%)									Function			
Position <sup>1</sup>	$Ag^2$	$Md^3$	$\overline{AOAc}$	BFAc	$\mathrm{GN}Ag$	BFAg	CMAg	GAAg	UGAg	KE	GW	Domain <sup>4</sup>	Resistance phenotype <sup>5</sup>
2,390,177 G>A	R254K	R261	0	0	0	0	32	21	0	0	0	IN (I.S4-I.S5)	L995F enhancer (predicted)
2,391,228 G>C	V402L	V410	0	7	0	0	0	0	0	0	0	TM (I.S6)	I1527T enhancer (predicted)
2,391,228 G>T	V402L	V410	0	7	0	0	0	0	0	0	0	TM (I.S6)	I1527T enhancer (predicted)
2,399,997 G>C	D466H	-	0	0	0	0	7	0	0	0	0	IN (I.S6-II.S1)	L995F enhancer (predicted)
2,400,071 G>A	M490I	M508	0	0	0	0	0	0	0	18	0	IN (I.S6-II.S1)	none (predicted)
2,400,071 G>T	M490I	M508	0	0	0	0	0	0	0	0	0	IN (I.S6-II.S1)	none (predicted)
2,416,980 C>T	T791M	T810	0	1	13	14	0	0	0	0	0	TM (II.S1)	L995F enhancer (predicted)
2,422,651 T>C	L995S	L1014	0	0	0	0	15	64	100	76	0	TM (II.S6)	driver
2,422,652 A>T	L995F	L1014	86	85	100	100	53	36	0	0	0	TM (II.S6)	driver
2,424,384 C>T	A1125V	K1133	9	0	0	0	0	0	0	0	0	IN (II.S6-III.S1)	none (predicted)
2,425,077 G>A	V1254I	I1262	0	0	0	0	0	0	0	0	5	IN (II.S6-III.S1)	none (predicted)
2,429,617 T>C	I1527T	I1532	0	14	0	0	0	0	0	0	0	TM (III.S6)	driver (predicted)
2,429,745 A>T*	N1570Y	N1575	0	26	10	22	6	0	0	0	0	IN (III.S6-IV.S1)	L995F enhancer
2,429,897 A>G	E1597G	E1602	0	0	6	4	0	0	0	0	0	IN (III.S6-IV.S1)	L995F enhancer (predicted)
2,429,915 A>C	K1603T	K1608	0	5	0	0	0	0	0	0	0	TM (IV.S1)	L995F enhancer (predicted)
2,430,424 G>T	A1746S	A1751	0	0	11	13	0	0	0	0	0	TM (IV.S5)	L995F enhancer (predicted)
2,430,817 G>A	V1853I	V1858	0	0	8	5	0	0	0	0	0	IN (IV.S6-)	L995F enhancer (predicted)
2,430,863 T>C	I1868T	I1873	0	0	18	25	0	0	0	0	0	IN (IV.S6-)	L995F enhancer (predicted)
2,430,880 C>T	P1874S	P1879	0	21	0	0	0	0	0	0	0	IN (IV.S6-)	L995F enhancer (predicted)
2,430,881 C>T	P1874L	P1879	0	7	45	26	0	0	0	0	0	IN (IV.S6-)	L995F enhancer (predicted)
2,431,019 T>C	F1920S	Y1925	0	0	0	0	1	4	0	0	0	IN (IV.S6-)	L995F enhancer (predicted)
2,431,061 C>T	A1934V	A1939	0	12	0	0	0	0	0	0	0	IN (IV.S6-)	L995F enhancer (predicted)
2,431,079 T>C	I1940T	I1945	0	4	0	0	7	0	0	0	0	IN (IV.S6-)	L995F enhancer (predicted)

(Table 1)

<sup>&</sup>lt;sup>1</sup> Position relative to the AgamP3 reference sequence, chromosome arm 2L. Variants marked with an asterisk (\*) failed conservative variant filters applied genome-wide in the Ag1000G phase 1 AR3 callset, but appeared sound on manual inspection of read alignments.

<sup>&</sup>lt;sup>2</sup> Codon numbering according to *Anopheles gambiae* transcript AGAP004707-RA in geneset AgamP4.4.

<sup>&</sup>lt;sup>3</sup> Codon numbering according to *Musca domestica* EMBL accession X96668 [8].

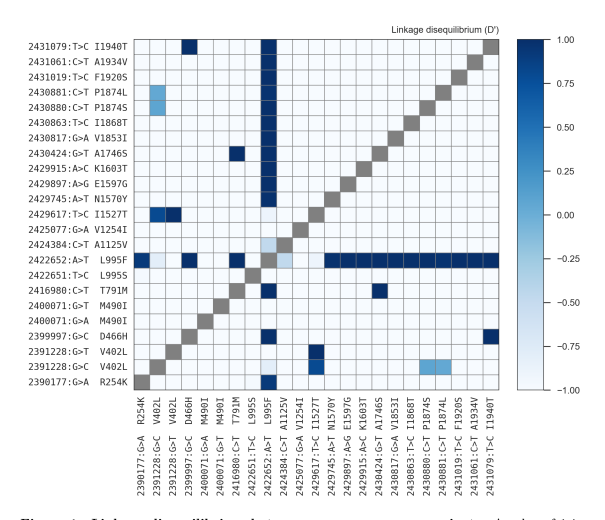
<sup>&</sup>lt;sup>4</sup> Position of the variant within the protein. IN=internal domain; TM=trans-membrane domain. The protein contains four homologous repeats (I-IV), each having six transmembrane segments (1-6). Codes in parentheses identify the specific domain, e.g., "I.S4" refers to trans-membrane segment 4 in repeat I, and "IS4-IS5" refers to the linker segment between I.S4 and I.S5.

<sup>&</sup>lt;sup>5</sup> Phenotype predictions are based on population genetic evidence and have not been confirmed experimentally.

from Central and East Africa (Table 1; [13]). All haplotypes carrying L995F or L995S have evidence for strong recent positive selection [13]. Both alleles were present in populations sampled from Cameroon and Gabon, including some individuals with a hybrid L995F/S genotype. In Cameroon these alleles were in Hardy Weinberg equilibrium ( $x^2 = 0.02$ , p > 0.05), thus there does not appear to be selection for or against carriers of both alleles; however in Gabon, they were not in equilibrium ( $x^2 = 8.96$ , p < 0.005), with an excess of heterozygotes suggesting a fitness advantage to mosquitoes carrying both alleles in this region.

The I1527T allele is present in An. coluzzii from Burkina Faso at 14% frequency, and there is evidence that haplotypes carrying this allele have been positively selected [13]. 118 Codon 1527 occurs within trans-membrane domain segment III.S6, immediately adjacent 119 to a second predicted binding pocket for pyrethroid molecules, thus it is plausible that 120 I1527T could alter insecticide binding [12]. We also found that the two variant alleles 121 affecting codon 402, both of which induce a V402L substitution, were in strong linkage with I1527T (D'  $\geq 0.8$ ; Figure 1), and almost all haplotypes carrying I1527T also carried a 123 V402L substitution. The most parsimonious explanation for this pattern of linkage is that 124 the I1527T mutation occurred first, and mutations in codon 402 subsequently arose on this 125 genetic background. Codon 402 also occurs within a trans-membrane segment (I.S6), and 126 the V402L substitution has associated with pyrethroid resistance in bedbugs [19]. Other 127 substitutions at this locus have also been associated with resistance, V402A/G in the moth 128 crop pests Helicoverpa zea [20] and V402M in Heliothis virescens, the latter of which has 129 been shown experimentally to confer resistance in *Xenopus* oocytes [21, 22]. However, 130 because V402L appears secondary to I1527T in our cohort, we classify I1527T as a putative 131 resistance driver and V402L as a putative enhancer. Because of the limited geographical 132 distribution of these alleles, we hypothesize that the I1527T+V402L combination represents a pyrethroid resistance allele that arose in West African An. coluzzii populations; however, 134 the L995F allele is at higher frequency (85%) in our Burkina Faso An. coluzzii population, 135 and is known to be increasing in frequency [23], therefore L995F may provide a stronger 136 resistance phenotype and is replacing I1527T+V402L in these populations. 137

Of the other 16 SNPs, 13 occurred almost exclusively in combination with L995F (Figure 1; [13]). These include the N1570Y allele, known to enhance pyrethroid resistance in An.



**Figure 1. Linkage disequilibrium between non-synonymous variants**. A value of 1 indicates that the two variants always occur in combination, and conversely a value of -1 indicates that the two variants never occur in combination. @TODO nuance this?

gambiae in combination with L995F [9]. These also include two variants in codon 1874 140 (P1874S, P1874L). P1874S has previously been found in a colony of the crop pest Plutella 141 xylostella with a pyrethroid resistance phenotype, but has not been shown to confer re-142 sistance experimentally [24]. 10 of these variants, including N1570Y and P1874S/L, occur 143 within internal linker domains of the protein, and so fit the model of variants that may en-144 hance or compensate for the driver phenotype by modifying channel gating behaviour [25, 145 9]. The remaining 3 variants are within trans-membrane domains, and so may enhance 146 resistance by altering or interacting with the insecticide binding sites on the VGSC [12]. 147 Because of the tight linkage between these 13 SNPs and the L995F allele, we classify all as putative L995F enhancers, although experimental work is required to confirm a resistance phenotype.

The remaining 3 variants (M490I, A1125V, V1254I) do not occur in combination with any known resistance allele, and do not appear to be associated with haplotypes under selection [13] A possible exception is the M490I allele found at 18% frequency in the Kenyan population, although the fact that this population has experienced a recent population crash makes it difficult to test for evidence of selection at this locus. All 3 variants occur in internal linker domains, and so do not fit the model of a resistance driver, although experimental work is required to rule out a resistance phenotype.

### 158 Haplotype structure

Although it is known that pyrethroid resistance is increasing in prevalence in malaria vector populations across Africa, it has not been clear whether this is being driven by the spread of resistance alleles via gene flow, or by resistance alleles emerging independently in multiple locations, or by some combination of both processes. The Ag1000G data resource provides a rich source of information about the evolutionary and demographic history of insecticide resistance in any given gene, because data are available not only for SNPs in gene coding regions, but also SNPs in introns and flanking intergenic regions, 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 analysis of the Vqsc [13], we used 1710 biallelic SNPs from within the 73.5 168 kbp Vqsc gene (@@N exonic, @@N intronic) to compute the number of SNP differences 169 between all pairs of 1530 haplotypes derived from 765 wild-caught mosquitoes. To visualise 170 these patterns, we used the pairwise genetic distances to perform hierarchical clustering 171 and found that haplotypes carrying resistance alleles were grouped into 10 distinct clus-172 ters. Five of these clusters carried the L995F allele (labelled F1-F5), and a further five 173 clusters carried L995S (labelled S1-S5). If we assume that haplotypes within each cluster 174 share a common ancestor since the introduction of insecticides, which is reasonable given 175 the high degree of similarity (clustered haplotypes were nearly identical across all 1710 176 SNPs spanning 73.5 kbp), then each of these clusters provides evidence that resistance 177 alleles have been spreading between geographical locations and species via adaptive gene 178 flow. Here we present several new analyses of these haplotype data, to confirm our initial inferences regarding gene flow, and provide further details regarding the origins and 180

movement of resistance alleles.

To provide an alternative view of the genetic similarity between haplotypes carrying 182 resistance alleles, we used haplotype data from within the Vgsc gene region to construct 183 median-joining networks (Figure 2). This analysis is similar to hierarchical clustering, however, it allows for the reconstruction and placement of intermediate haplotypes that 185 may not be observed in the data and allows much easier visualisation of the closely related 186 haplotypes at the leaves of the dendrogram branches. We constructed these networks using 187 a maximum edge distance of 2 SNP differences, to ensure that each connected component 188 in the resulting networks represents a collection of haplotypes with a recent common ancestor (analogous to the cutting of the dendrogram in [13]). For haplotypes carrying 190 both L995F and L995S alleles, the resulting networks confirm the presence of ten distinct 191 clusters, with close correspondence to the clusters we identified previously [13]. 192

The haplotype networks bring into sharp relief the explosive evolution of amino acid substitutions secondary to the L995F allele. Within the F1 network, nodes carrying non-

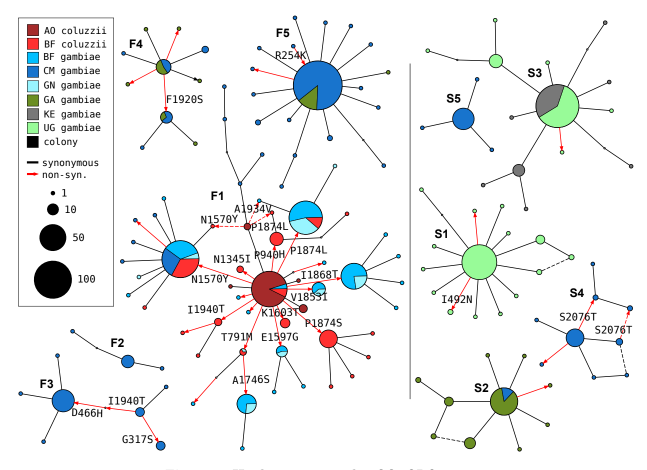


Figure 2. Haplotype networks. @@TODO caption

synonymous variants radiate out from a central node carrying only L995F (Figure 2), 195 suggesting that the central node represents the ancestral haplotype carrying L995F alone 196 which initially came under selection, and these secondary variants have arisen subsequently 197 as new mutations. Many of the nodes carrying secondary variants are large, consistent with positive selection and a functional role for these secondary variants as enhancers of the 199 L995F resistance phenotype (Table ??). The F1 network also allows us to infer multiple 200 introgression events between the two species. The central (ancestral) node comprises 201 haplotypes from both species, as do nodes carrying the N1570Y, P1874L, and T791M. This 202 structure is consistent with an initial introgression of the ancestral F1 haplotype, followed later by introgressions of haplotypes carrying secondary mutations. The contrast between 204 the haplotype networks for the L995F and L995S alleles is striking because of the near-205 total absence of non-synonymous variation within the L995S networks. As we reported 206 previously @@Where?, this difference is highly significant – the ratio of non-synonymous 207 to synonymous nucleotide diversity (@@piN/piS) is @@N times higher among haplotypes 208 carrying L995F relative to haplotypes carrying L995S (@@Test; P=@@) ([13]). Some 209 secondary non-synonymous variants are present within the L995S networks, but all are at 210 low frequency, and thus may be neutral or mildly deleterious variants that are hitch-hiking 211 on selective sweeps for the L995S allele. 212

While the haplotype clustering and network analyses provide evidence for the spread 213 of resistance alleles via adaptive gene flow, and for the secondary evolution of L995F 214 enhancer alleles, they have several limitations. These analyses only leverage information 215 about genetic distance within the Vasc gene, and for very recent events, insufficient time 216 has elapsed for informative mutations to accumulate within this relatively small genome 217 region. The fact that we observe five distinct clusters for each of the codon 995 alleles 218 suggests that each cluster is in some sense independent from the others, and thus gene flow is not required for resistance to emerge in multiple geographical locations. However, 220 the threshold for the genetic distance at which we have chosen to divide haplotypes into 221 different networks or clusters is to a certain extent arbitrary, and based on an intuitive 222 sense of how much variation could have accumulated among the descendants of a single 223 resistant ancestor since the onset of selective pressure. Finally, analyses of genetic distance within a fixed genome region can be confounded by recombination events occurring within 225

that region. For example, a recombination event within the *Vgsc* gene upstream of codon
995 could cause us to split a collection of haplotypes into two clusters, even though they
are ancestrally related within the region downstream of the recombination event. In
the next sub-sections we provide some further analyses to help clarify these ambiguities,
using haplotype sharing from the genome regions flanking the *Vgsc* gene to provide finer
resolution to diagnose recent gene flow and recombination events.

#### 232 Insecticide resistance outbreaks

To provide an aid to further interpretation of the genetic data, and relating them to the 233 challenges of insecticide resistance management, we introduce the concept of an insecticide resistance outbreak. Informally, we define a resistance outbreak by analogy with the epidemiological concept of an outbreak, as a rapid increase in the prevalence 236 of insecticide resistance among mosquitoes at a particular place and time. Note that 237 this does not imply that the overall abundance of mosquitoes is increase, just that the 238 relative frequency of resistance within mosquito populations is increasing. We also re-239 quire that all occurrences of insecticide resistance within the same outbreak are connected by a chain of transmission of resistance alleles from parent to progeny mosquitoes, and 241 thus can be traced back to a single resistant common ancestor. A resistance outbreak 242 can be **localised**, meaning that it affects a small group of mosquitoes of a single species 243 from a limited geographical area. Alternatively, a resistance outbreak may be **spreading**, 244 meaning that resistance alleles have been transmitted since the introduction of insecti-245 cides by interbreeding of mosquitoes of different species and/or originating from different geographical locations. 247

Our goal for the *Vgsc* gene can now be restated, which is to perform an insecticide resistance outbreak analysis. We would like to diagnose how many separate outbreaks have occurred, which outbreaks are localised, and which are spreading. For spreading outbreaks, we would like to reconstruct the path of transmission of resistance alleles between mosquito populations, and to provide information on the probable source. We would, of course, also like to identify the primary and secondary genetic factors that are driving each outbreak. Stated in this way, it is easier to discuss how this information is potentially relevant to insecticide resistance management, and to frame key epidemiological questions. For

example, we would like to begin to build a picture of where and when local conditions 256 have favoured the evolution of insecticide resistance, and whether those conditions are 257 relatively patchy (and hence outbreaks are mainly localised) or whether conditions are 258 consistent over broad areas (and hence can support a spreading outbreak). We would also like to know which mosquito populations are sufficiently connected to enable outbreak 260 spread, and if there is any consistent pattern to the direction of spread. This information 261 could be relevant to discussions about how resources for insecticide resistance management 262 might be targeted, what strategies are appropriate in which settings, and where and when 263 insecticide resistance management needs to be coordinated between different countries and/or at different levels of administration. 265

For clarity, we also define the concept of an insecticide resistance outbreak formally in terms of coalescent theory, as a collection of lineages (1) sharing a resistance driver allele by descent, (2) coalescing more recently than the onset of insecticide pressure, and (3) having increased in frequency because of positive selection due to insecticides. This definition is illustrated for four hypothetical outbreaks in Figure 3. Because mosquitoes

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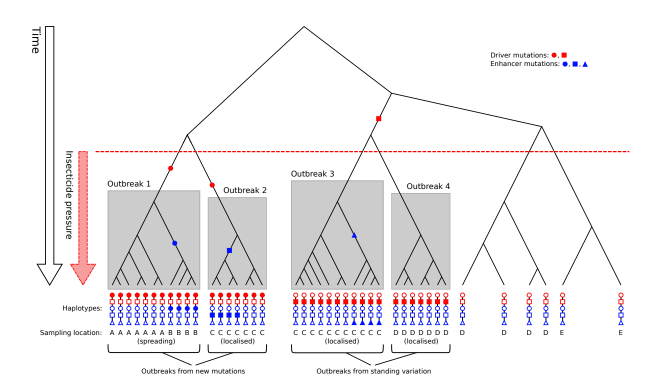


Figure 3. Illustration of insecticide resistance outbreaks. @@TODO explanation.

are sexually recombining, genealogical trees vary along the genome, and so we define 271 resistance outbreaks with respect to a specific gene locus, which for the present study 272 is codon 995 within the *Vgsc* gene. Note that separate outbreaks may be driven by 273 the same resistance allele, and this can occur if multiple mutational events occur after the introduction of insecticides (Figure 3, outbreaks 1 and 2), or if a resistance allele 275 is present in mosquito populations as standing variation prior to insecticide use (Figure 276 3, outbreaks 3 and 4). Here we are primarily concerned with whether outbreaks are 277 localised or spreading, because this has immediate epidemiological relevance. We do not 278 attempt to infer whether separate outbreaks with the same driver allele arose via standing variation or new mutations, however this is an interesting biological question to address 280 in future studies. As a technical note, there is a simple correspondance with terminology 281 conventionally used in the population genetics literature to describe selective sweeps. At 282 a given gene locus, a hard selective sweep gives rise to a single resistance outbreak, and a 283 soft selective sweep gives rise to multiple resistance outbreaks.

## Outbreak analysis from haplotype age

As described above, haplotype data from genome regions both within and flanking the Vgsc gene provide a higher resolution for reconstructing recent historical events. To leverage this information, we used a heuristic approach to estimate the time to most recent common ancestor (TMRCA) or "age" for each pair of haplotypes in our dataset, centering the analysis on Vgsc codon 995. For each pair of haplotypes, we estimated the length

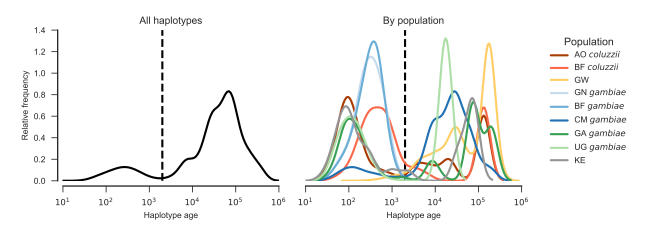


Figure 4. Haplotype age distribution. @@TODO caption.

of the region shared identical by descent (IBD), and the number of mutations that have 291 accumulated since the most recent common ancestor. We then combined these two pieces 292 of information to produce a point estimate for the haplotype age (Methods). We studied 293 the overall distribution of pairwise haplotype ages (Figure 4), and used hierarchical clustering to construct a dendrogram and visualise the overall age structure (Figure 5). We 295 caution that although the estimated ages are in units of generations, these estimates have 296 not been calibrated, and there is substantial uncertainty regarding both the mutation and 297 recombination rate parameters. The ages therefore should not be interpreted as reliable 298 absolute values, but they can be compared to each other to investigate the relative age of different events. 300

A key feature of the overall age distribution is that it is bimodal, with a minor mode of haplotypes coalescing recently, and a major mode coalescing further in the past (Figure 4). This is expected at an insecticide resistance locus experiencing one or more resistance outbreaks. Within each outbreak, all haplotypes share a very recent common ancestor, but between outbreaks and among haplotypes without any resistance allele, haplotypes are more distantly related, and the distribution of ages is influenced by mosquito population

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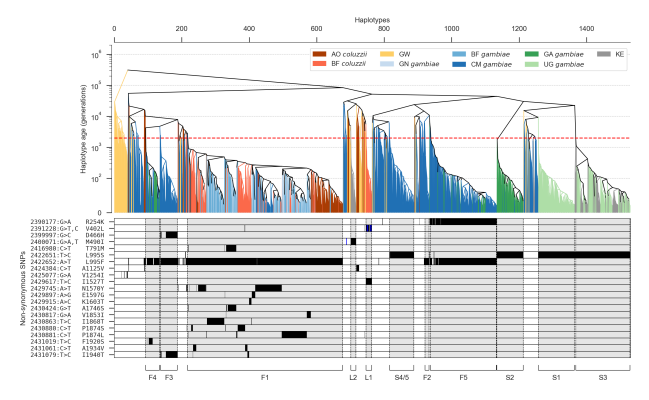


Figure 5. Clustering of haplotypes by age. @@TODO caption.

size and other demographic factors. In particular, mosquito populations generally have 307 a large effective population size (@@REF Ag1000G), and so in the absence of selection, 308 haplotypes are expected to coalesce slowly. The bimodal age distribution is not due to 309 geographical population structure, because the same bimodality is observed within several populations. We take the midpoint between these two modes as an estimate for the earliest 311 time of onset of selective pressure due to insecticides, and thus for the maximum age of 312 a resistance outbreak. To identify haplotype clusters representing putative resistance 313 outbreaks, we then cut the haplotype dendrogram at this maximum outbreak age (Figure 314 5). Comparing this to previous analyses of haplotype structure based on genetic distance, we find clusters F1-F5 and S1-S3 recapitulated with close correspondence, and S4 and 316 S5 merged into a single cluster. We label a new cluster "L@@" representing an outbreak 317 driven by the I1527T allele in combination with one or the other V402L allele. We also label 318 a cluster "L@@" capturing a set of haplotypes from Kenya carrying the M490I variant, 319 although the fact that these haplotypes all share a recent common ancestor may be a 320 reflection of the unusual demography of the Kenyan population which has experienced 321 a severe population crash (@@REF) and not be due to recent selection for insecticide 322 resistance. As in earlier analyses, clusters F1, F4, F5 and S3 all include haplotypes 323 sampled from multiple geographical locations, and thus represent spreading outbreaks. 324 Clusters F2, F3, S1, S2, S4/5 and L1 include only haplotypes from a single sampling 325 location, and thus appear to represent localised outbreaks. 326

We then studied the distribution of haplotype ages within each spreading outbreak, to 327 attempt to reconstruct information about the historical path of transmission of resistance 328 alleles between locations. To do this, we grouped the haplotypes within each spreading 329 outbreak by sampling location, and compared the distribution of haplotype ages both 330 within and between locations. To aid in interpreting these data, we define three possi-331 ble spreading scenarios, being: (1) a directional spread from one population to another; 332 (2) spread from an unsampled population into the sampled populations; and (3) a com-333 plex scenario involving multiple gene flow events. In Figure 6 we illustrate the expected 334 genealogy and haplotype age distribution under each of these scenarios. 335

The clearest result was obtained for outbreak F1 (Figure 7). Within this outbreak, haplotypes from Cameroon and Angola are significantly younger than haplotypes from

Burkina Faso and Guinea. The age distributions are consistent with an outbreak originating in West Africa and subsequently spreading towards Cameroon and separately towards
Angola. We were surprised that the age distributions for An. gambiae and An. coluzzii
from Burkina Faso are very similar, despite the fact that previous studies have shown that
introgression has occurred from An. gambiae into An. coluzzii. This may indicate that
the initial introgression event happened during the early phases of the outbreak, but is
also consistent with a complex history of multiple gene flow events between the species.

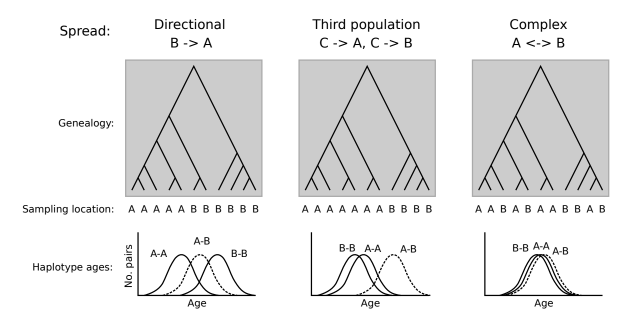


Figure 6. Inferring history of spread from haplotype ages. @@TODO explain.

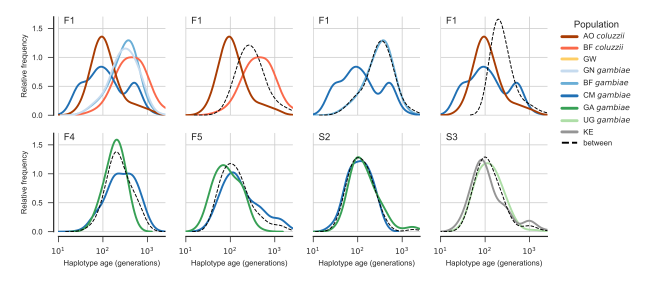


Figure 7. Haplotype age distributions within spreading outbreaks. @@TODO caption.

Outbreaks F4, F5 and S2 each involve haplotypes from both Cameroon and Gabon. 345 Interpreting the age distributions for these outbreaks is difficult, because mosquitoes from 346 Gabon were collected at a much earlier time point (2000) than mosquitoes from Cameroon 347 (20@@). If our haplotype age estimates were well-calibrated, and we also had reliable estimates for the number of mosquito generations per year, then we might be able to 349 adjust for this time difference, however we are not able to do so presently. An interesting 350 feature of these outbreaks, however, is that we would expect haplotypes from Gabon to 351 appear older due to the time of sampling, which is observed for outbreak S2 but not 352 for F4 or F5. Indeed, S2 is at a high frequency among all Gabon haplotypes and a low frequency among Cameroon haplotypes, whereas the reverse is true for F4 and F5. These 354 data suggest that F4 and F5 have spread from Cameroon towards Gabon, while S2 has 355 spread in the opposite direction. A lot can happen in mosquito populations in @@N years, 356 however, and these conclusions remain highly speculative pending further sampling from 357 both locations. 358

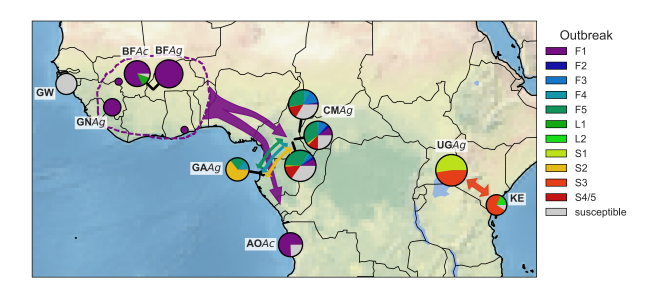
For outbreak S3 involving haplotypes from Uganda and Kenya, the age distributions 359 do not suggest any clear direction of gene flow. This could reflect multiple gene flow 360 events in either or both directions. However, another outbreak (S1) is localised in Uganda 361 and represented within the Ugandan population at roughly equal frequency with S3. If 362 transmission was occurring from Uganda towards Kenya, we might expect both outbreaks 363 to have spread to Kenya. Thus the localisation of S1 suggests S3 has spread into Uganda 364 from Kenya or another location. Again, this conclusion remains tentative and requires 365 confirmation via further sampling. 366

To summarise these conclusions in a concise way, we have depicted the distribution and spread of resistance outbreaks via the map shown in Figure 8. We have plotted haplotypes from each sampling location as a pie chart. The overall size of each pie chart represents the number of haplotypes sampled, and coloured wedges within each pie represent the frequency of each resistance outbreak within the population. Coloured arrows are used to depict our inferences regarding the transmission paths for spreading outbreaks. Our conclusions regarding direction of spread for outbreaks F4, F5, S2 and S3 are tentative, and we indicate this with a question mark. Because of the relatively sparse geographical representation within the Ag1000G phase 1 dataset, and the fact that collections were

not synchronized but span several years, we cannot be precise about the geographical 376 origins of these resistance outbreaks. Even for outbreak F1 where we have clear evidence 377 of spread from West Africa towards Central and Southern Africa, we have only sampled 378 mosquitoes from Guinea and Burkina Faso, and the true source of the outbreak may not be either of these countries. We indicate this uncertainty regarding the outbreak source 380 as a coloured area with a dashed border. This representation is imperfect, as is our 381 knowledge regarding the sources and transmission paths of these outbreaks, but we hope 382 this depiction may at least serve to stimulate further sampling, analysis and discussion, 383 with the aim of improving our knowledge of resistance outbreaks for Vgsc as well as other insecticide resistance genes. 385

# Design of genetic assays for outbreak surveillance

The insecticide resistance outbreaks we have identified here are undoubtedly ongoing, affecting many more mosquito populations than we have sampled in Ag1000G phase 1, and continuing to spread. In addition, other outbreaks may be occurring in populations that we have not sampled, or in populations we have sampled but since the sampling date. Wholegenome sequencing of individual mosquitoes clearly provides data of sufficient resolution to identify resistance outbreaks, and could also be used to provide ongoing outbreak surveil-



**Figure 8.** Geographical distribution of resistance outbreaks. @@TODO caption. @@TODO explain Clarkon and Norris points.

lance. The cost of whole-genome sequencing continues to fall, with the present cost being 393 approximately 100 GBP to obtain ~30× coverage of an individual Anopheles mosquito 394 genome with 150 bp paired-end reads. Mobile sequencing using nanopore technology is 395 also developing rapidly [26] and may be a realistic prospect for mosquito whole-genome sequencing within a few years. There is an interim period, however, during which it may be 397 more practical to develop targeted genetic assays for outbreak surveillance that could scale 398 to tens of thousands of mosquitoes at low cost. For example, both next-generation and 399 mobile sequencing platforms can be used for amplicon sequencing, where specific genome 400 regions are amplified and sequenced in highly multiplexed libraries [27, 28]. 401

To facilitate the development of targeted genetic assays for Vgsc insecticide resistance outbreak surveillance, we have produced two supplementary data tables. In Supplementary

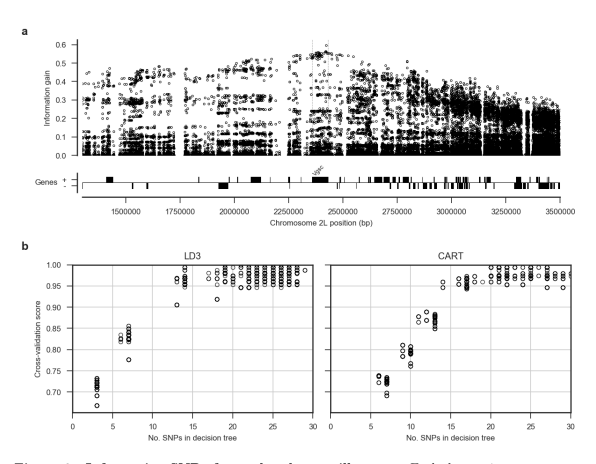


Figure 9. Informative SNPs for outbreak surveillance. a, Each data point represents a single SNP. The information gain value for each SNP provides an indication of how informative the SNP is likely to be if used as part of a genetic assay for testing whether a mosquito carries a resistance haplotype, and if so, which resistance outbreak it derives from. b, Number of SNPs required to accurately classify which outbreak a haplotype derives from. Decision trees were constructed using either the LD3 (left) or CART (right) algorithm for comparison. Accuracy was evaluated using 10-fold stratified cross-validation.

tary Table 1 we provide a list of all biallelic SNPs discovered with high confidence in this 404 study within the Vqsc gene and in the 100 kbp upstream and downstream flanking regions. 405 To aid in PCR primer design, for each SNP we provide the flanking sequence for 250 bp 406 upstream and downstream of the SNP position, including information about polymorphisms within these flanking regions. Not all SNPs are informative for detecting whether 408 an individual mosquito carries a haplotype from a resistance outbreak, and we provide 409 some summary statistics for each SNP to aid in the selection of the most informative 410 SNPs. This includes allele frequencies within each of the outbreaks identified here, as well 411 as for populations of susceptible haplotypes. We also provide the overall variance in allele frequencies, the information gain [29], and the Gini impurity [30] for each SNP. Note that 413 recombination events are more likely at increasing distances upstream and downstream 414 of the resistance variants under selection, and thus the most informative SNPs are found 415 closest to the resistance variants within the gene (Figure 9). However, SNPs with some 416 information gain are available throughout the gene and in flanking regions.

We suggest that the design of a genetic assay proceed by (1) performing an initial 418 round of filtering to remove SNPs which are not informative (e.g., low information gain); 419 (2) performing a round of primer design to remove SNPs for which primers are unlikely to 420 be successful; (3) performing a full analysis of the remaining SNPs to select a subset that 421 is sufficient to classify all outbreaks identified here, including some redundancy; (4) finalise 422 primer designs for the chosen panel of SNPs. A possible methodology for step 3 would be 423 to use an algorithm such as ID3 [29] or CART [30] to build a decision tree, although many 424 other algorithms for building classifiers are also applicable. To aid in the development of 425 a classifier, in Supplementary Table 2 we provide our classification for each of the 1530 426 haplotypes sampled here, along with the alleles carried by each haplotype for each of 427 the SNPs included in Supplementary Table 1. To test the methodology, we constructed decision trees using either LD3 or CART algorithms, and using all available SNPs from 429 within the Vasc plus 20 kbp flanking regions as input features (i.e., assuming primers could 430 be designed in all cases). Figure 9b shows the cross-validation scores obtained for trees 431 constructed allowing increasing numbers of SNPs. This analysis suggests that it should 432 be possible to construct a tree able to classify haplotypes from all 10 resistance outbreaks with >95% accuracy using 20 SNPs or less.

#### 435 Recombination

To look for evidence that haplotypes have experienced recent positive selection, we performed an analysis of extended haplotype homozygosity (EHH) decay @@REF. We defined a core region spanning Vgsc codon 995 and an additional 4 kbp of flanking sequence
(Methods). Within this core region, we found @@N distinct haplotypes at a frequency >
1% within the cohort, including core haplotypes representing each of the resistance outbreaks we identified above, and a further @@N core haplotypes not carrying any known
or putative resistance allele for comparison. @@TODO finish this

Sandbox paragraph: @@TODO integrate or remove In this section we present

Sandbox paragraph: @@TODO integrate or remove In this section we present analyses of recombination both within the *Vgsc* gene itself and on either flank. These analyses provide information about which haplotypes have experience recent selection, and an alternative view of how different haplotypes are related. They also provide in-

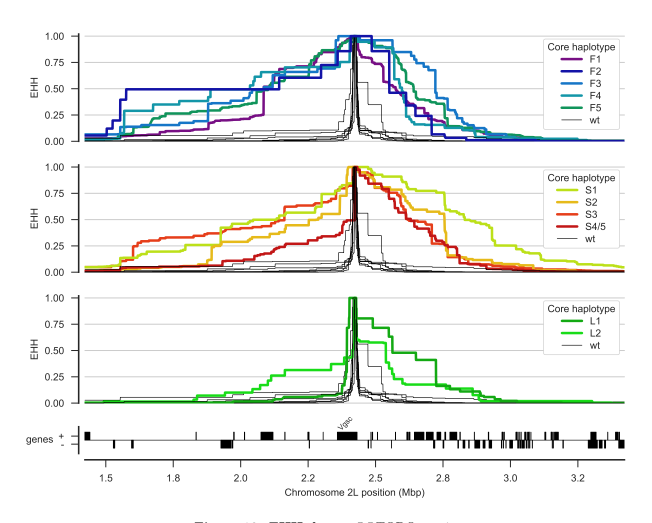


Figure 10. EHH decay. @@TODO caption

formation about where in the genome recombination events have occurred, and whether 447 these recombination events may have biased or otherwise influenced the outcome of analy-448 ses presented in other sections. EHH analysis first identifies collections of haplotypes with 449 the same alleles at a core locus. The haplotypes within each collection are then compared, and the fraction of haplotype pairs that remain identical (EHH) is computed moving both 451 up- and down-stream of the core locus. Recombination events break haplotype homozy-452 gosity, and so a slow decay of EHH indicates fewer recombination events, A collection of 453 haplotypes where EHH decays more slowly provides evidence for positive selection on the 454 core allele, Haplotypes that have risen rapidly in frequency due to selection will be younger on average, and thus the length of regions of homozygosity between pairs of haplotypes 456 These analyses provide confirmation of which haplotypes have experience recent positive 457 selection, as haplotypes that have recently increased in frequency will 458

As mentioned earlier, analyses of haplotype structure based on genetic distance within 459 the fixed window of the Vgsc gene could be affected if recombination events occurred within the gene. Our analyses of haplotype age should be less affected by recombination, 461 because they explicitly take recombination into account, estimating the positions at which 462 recombination events have occurred to interrupt regions shared IBD between pairs of 463 haplotypes. However, these analyses were based on a heuristic method for estimating 464 recombination breakpoints, and there are several potential sources of error. To study 465 the evidence for recombination within the genome region spanning the Vqsc gene, and 466 provide some additional confirmation that our inferences regarding insecticide resistance 467 outbreaks have not been affected by recombination or other sources of error, we performed 468 an additional analysis of genetic distance between haplotypes. We first constructed a 469 putative ancestral haplotype for each of the outbreaks we identified, by starting from 470 the codon 995 position and separately moving upstream and downstream, assuming the major allele at each SNP bifurcation point represents the ancestral haplotype. We then 472 computed the genetic distance  $(D_{XY})$  between each of our sampled haplotypes and each 473 of the inferred ancestral outbreak haplotypes, computing the distance in @@ overlapping 474 windows of @@ bp across a 2 Mbp region spanning the Vqsc gene. The results for outbreaks 475 F1-F5 are plotted in Figure 11, and outbreaks S1-S4/5 are shown in Figure ??. In these plots we expect that all haplotypes from a given outbreak should share very close genetic

similarity  $(D_{XY} \approx 0)$  with each other and with the ancestral haplotype for that outbreak within the Vgsc gene itself, with an increasing number of haplotypes recombining away from the ancestral outbreak haplotype as we move away from the gene in either the upstream or downstream direction. Conversely, haplotypes from one outbreak should not share any close genetic similarity  $(D_{XY} > 0)$  with the inferred ancestral haplotype from a different outbreak, either within the Vgsc gene or in flanking regions.

The results for all outbreaks are largely consistent with this expectation. For this

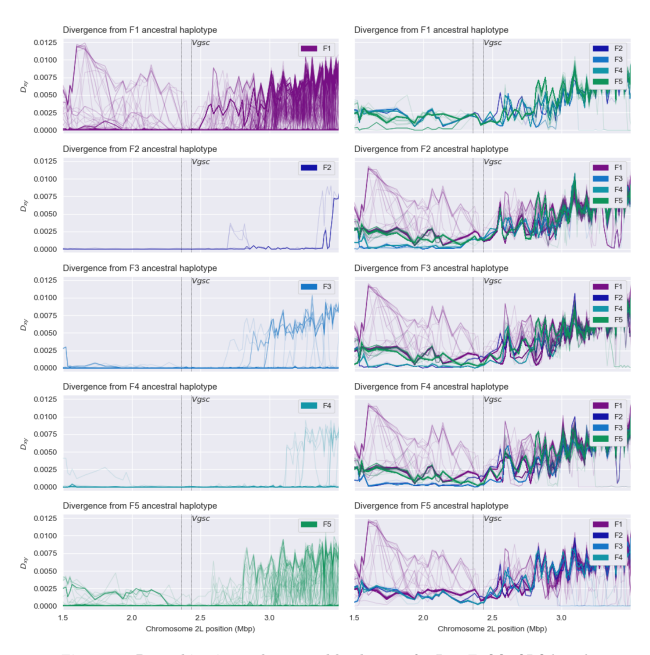


Figure 11. Recombination and ancestral haplotypes for L995F. @@TODO legend

analysis we treated S4/5 as a single outbreak, as indicated by the haplotype age analysis, and we can gain some insight into why these two were split into separate clusters in earlier analyses. All haplotypes in the S4/5 outbreak share close similarity with the ancestral haplotype on both flanks of the *Vgsc* gene, but there is a short region of within the gene where a subset of haplotypes are diverged. This region of divergence accounts for the S4/S5 split in earlier analyses. @@TODO explain @@TODO also note relatively low divergence among F2, F3, F4 on upstream flank and explain

## **Discussion**

- 493 @@TODO Discuss accessibility, have we missed any functional variation?
- @@TODO Discuss weaknesses, caveats and potential improvements to method for estimating haplotype age.
- @@TODO What are the implications for insecticide resistance management? Realistically how could this information be used?
- @@TODO What about DDT? If prior selection for DDT resistance, how might this complicate the picture? Do we see any evidence for multiple phases of selection?
- ©@TODO Speculate on why L995F but not L995S has evolved secondary variation.

# Methods

#### 502 Code

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

#### 505 Data

- We used variant call data from the phase 1 AR3 release and phased haplotype data from
- AR3.1. These data are publically downloadable via ftp from https://www.malariagen.
- $_{508}$  net. @@add ENA from paper

### 509 Data collection and processing

For detailed information on Ag1000g WGS sample collection, sequencing, variant calling, 510 quality control and phasing see [13]. In brief, An. gambiae and An. coluzzii mosquitoes 511 were collected from eight countries across Sub-Saharan Africa: Angola, Burkina Faso, 512 Cameroon, Gabon, Guinea, Guinea Bissau, Kenya and Uganda. From Angola just An. 513 coluzzii were sampled, Burkina Faso had samples of both An. qambiae and An. coluzzii 514 and all other populations consisted of purely An. gambiae except for Kenya and Guinea Bissau, where species status is uncertain [13]. Mosquitoes were individually whole genome 516 sequenced on the Illumina HiSeq 2000 platform, generating 100bp paired-end reads. Se-517 quenced reads were aligned to the [An. gambiae] AgamP3 reference genome assembly 518 [31]). Aligned bam files underwent improvement, before variants were called using GATK 519 UnifiedGenotyper. Quality control included removal of samples with mean coverage <= 520 14x and an accessibility map was employed following a similar approach to that used for 521 human data by The 1000 Genomes Project Consortium [32]). Various quality control filters 522 were applied to remove samples and SNPs with poor quality data. This process produced 523 a call set containing @@n SNPs genotyped in 765 wild-caught individual mosquitoes [13]. 524 The Ag1000g variant data was functionally annotated using the SnpEff v4.1b software 525 which allowed investigation of potential phenotype altering variants within Vqsc [33]. Nonsynonymous Vgsc variants were identified as all variants in AGAP004707, 2L:2358158-527 2431617, with a SnpEff annotation of âĂIJmissenseâĂİ and an ALT allele frequency of 528 >5% in at least one of the nine mosquito populations, with the exceptions of the multi-529 allelic SNP 2L:2400071 G>A which is shown despite only being found in An. qambiae from 530 Cameroon at 0.4% frequency, as the G>T variant at the same position which causes the same codon change (M4901), is found above 5% frequency in Kenya. F1920S is included for 532 continuity with recent An. qambiae Vqsc research [13]. A minimum ALT allele frequency 533 was employed to discriminate towards variants that may be undergoing selective sweeps 534 and against less informative low frequency alleles. 535 For ease of comparison with previous work on Vgsc, pan Insecta, in Table 1 we report 536 codon numbering for both An. gambiae and Musca domestica (the species in which the 537 gene was first discovered). The M. domestica Vqsc sequence (EMBL accession X96668 -538

[8]) was aligned with the *An. gambiae* AGAP004707-RA sequence (AgamP4.4 gene-set), using the Mega v7 software package [34]. A map of equivalent codon numbers between the two species can be download from the MalariaGEN website (@@include as supplementary data file?)- https://www.malariagen.net/sites/default/files/content/blogs/domestica\_gambiae\_map.txt.

Haplotypes for each chromosome of each sample were estimated (phased) using using 544 phase informative reads (PIRs) and SHAPEIT2 v2.r837 [35], see [13] supplementary text 545 for more details. The SHAPEIT2 algorithm is unable to phase multi-allelic positions, therefore the two multi-allelic non-synonymous SNPs within the Vgsc gene (>5% ALT frequency in at least one population), altering codons V402 and M490, were phased onto the 548 haplotypes using MVNcall v1.0 [36]. Conservative filtering had removed one of the three 549 known insecticide resistance conferring kdr variants, N1570Y [9]. After manual inspection 550 of the read alignment revealed that the SNP call could be confidently made, it was added 551 back into the data set and then also phased onto the haplotypes using MVNcall. To evaluate the linkage disequilibrium (LD) of non-synonymous Vqsc mutations with the two 553 most widespread kdr resistance mutations (L995S/F), the D1 statistic was calculated using 554 haplotypes. 555

#### Haplotype networks

Discerning the relationships between similar haplotypes can be difficult when using bifurcating trees as, inherently, the distance between the leaves at the tips (haplotypes) will be
small. As these relationships may be informative of the history of selection, we utilised
a network approach to elucidate them. We constructed haplotype networks using the
median-joining algorithm [37] as implemented in a custom Python script available from
https://github.com/malariagen/agam-vgsc-report Networks were rendered with the
graphviz library and a composite figure constructed using Inkscape.

# 564 Haplotype age

Haplotype age. @@TODO - AM -Length of shared haplotype and number of mutations between them are informative of ageâĂę -Pairwise t values were hierarchically clustered and visualised as a dendrogram using the Python library Scipy and its cluster hierarchy functions linkage method. -Cutting the dendrogram at @@generations clustered haplotypes together into haplogroupsâĂę - Naming of haplogroups with reference to Ag1000g... -dendro figure/distro figures/map - Python libraries...

#### 571 Recombination

Recombination. @@TODO - AM - Absolute divergence dxy...

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