1 <u>Ultra-conserved sequences in the genomes of highly diverse Anopheles mosquitoes</u>,

2 with implications for malaria vector control.

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ABSTRACT

18 DNA sequences that are exactly conserved over long evolutionary time scales have 19 been observed in a variety of taxa. Such sequences are likely under strong functional 20 constraint and they have been useful in the field of comparative genomics for 21 identifying genome regions with regulatory function. A potential new application for 22 these ultra-conserved elements has emerged in the development of gene drives to 23 control mosquito populations. Many gene drives work by recognising and inserting at 24 a specific target sequence in the genome, often imposing a reproductive load as a 25 consequence. They can therefore select for target sequence variants that provide

26 resistance to the drive. Focusing on highly conserved, highly constrained sequences

- 27 lowers the probability that variant, gene drive-resistant alleles can be tolerated.
- 28

29 Here we search for conserved sequences of 18bp and over in an alignment of 21 30 Anopheles genomes, spanning an evolutionary timescale of 100 million years, and 31 characterise the resulting sequences according to their location and function. Over 8000 32 ultra-conserved elements were found across the alignment, with a maximum length of 33 164 bp. Length-corrected gene ontology analysis revealed that genes containing 34 Anopheles ultra-conserved elements were over-represented in categories with structural 35 or nucleotide binding functions. Known insect transcription factor binding sites were 36 found in 48% of intergenic Anopheles ultra-conserved elements. When we looked at 37 the genome sequences of 1142 wild-caught mosquitoes we found that 15% of the 38 Anopheles ultra-conserved elements contained no polymorphisms. Our list of 39 Anopheles ultra-conserved elements should provide a valuable starting point for the 40 selection and testing of new targets for gene-drive modification in the mosquitoes that 41 transmit malaria.

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INTRODUCTION

44 DNA sequences that are highly conserved over long evolutionary timescales have been 45 identified in many organisms. Some of these sequences show complete conservation at 46 the nucleotide level and are often known as ultra-conserved elements (UCEs). 47 Originally, UCEs were defined as sequences of at least 200bp that were identical 48 between human, mouse and rat genomes (Bejerano *et al.* 2004). Subsequently the 49 search for UCEs has been extended to other vertebrates, insects and plants (e.g.

Makunin *et al.* 2013; Siepel *et al.* 2005; Baxter *et al.* 2012), and to sequences of length
50 bp or more.

52

53 There are several reasons why UCEs are of interest. First, in the field of comparative 54 genomics, UCEs are thought to represent functionally important regions. While there 55 is still some mystery around why sequences might be conserved at the nucleotide level 56 over long evolutionary timescales, it has been shown that UCEs 1) often are involved 57 in regulation of transcription of genes, especially essential genes involved in 58 development (e.g. Visel et al. 2008); 2) may have a role in chromosomal structure (e.g. 59 Chiang et al. 2008); and 3) are sometimes non-coding RNA genes (e.g. Kern et al. 60 Even UCEs in protein coding regions may have multi-functional roles 2015). 61 (Warnefors et al. 2016). Second, UCEs can act as probes to facilitate genomic 62 sequencing of non-model organisms using sequence-capture methods (Faircloth et al. 63 2012). Third, alterations in UCEs have been shown to have an association with human 64 cancers (e.g. Calin et al. 2007; Lin et al. 2012).

65

66 A new potential role for UCEs has recently emerged in the fight against malaria using 67 gene-drive mosquitoes (Kyrou et al. 2018). Anopheles mosquitoes are the vectors of 68 malaria parasites, and mosquito control has been responsible for much of the recent 69 success in reduction of malaria cases (78% of the 663 million malaria cases averted 70 globally since 2000 (Bhatt et al. 2015)). Progress in reducing malaria cases has stalled 71 (WHO 2018), probably in part due to resistance of the mosquitoes against commonly 72 used pesticides. One novel method under consideration is the development of 73 mosquitoes containing gene drives that either reduce the population size (Windbichler 74 et al. 2011; Hammond et al. 2016) or make them unable to transmit the malaria parasite

75 (Gantz et al. 2015). Both methods currently rely on nuclease-based synthetic gene drive 76 systems that introduce a desired trait at a precise genomic location, spreading it in a 77 target population at such a rate that outweighs fitness costs associated with the trait 78 (Burt 2003). The technologies include RNA-guided endonucleases (such as 79 CRISPR/Cas9) and homing endonucleases (Jinek et al. 2012; Windbichler et al. 2011). 80 These enzymes recognise and cleave a particular target size of about 18 bp. When the 81 sequence coding for these enzymes is engineered into its own target site in the genome 82 and is expressed in the germline, it creates a double-strand break in the homologous 83 chromosome. The break will usually be repaired by homology-directed repair using the 84 drive-containing chromosome as a template which results in conversion of the repaired 85 to also contain the drive element in greater than the usual 50% inheritance rate among 86 the gametes. An efficient gene drive can be inherited by almost 100% of progeny 87 (Hammond et al. 2015). Theoretical and laboratory studies have shown that changes to 88 the recognition site can result in alleles that cannot be recognised or cleaved. If these 89 alleles confer increased fitness compared to the wild type allele in the presence of the 90 gene drive they can be expected to spread and retard the spread of the gene drive 91 (Deredec et al. 2008; Unckless et al. 2017; Hammond et al. 2017). For population 92 suppression gene drives that are designed to impair essential genes the selection 93 pressure for resistance alleles to arise is high. These alleles can arise from standing 94 variation at the target site in a wild population, or may come about from the action of 95 the endonuclease. This is because non-homologous end joining can sometimes repair 96 the double-strand break, and random insertions and deletions can be introduced to the 97 target site.

99 Two of the most important vector species in sub-Saharan Africa are the close relatives 100 Anopheles gambiae and An. coluzzii, both of which are highly genetically diverse. A 101 study of 765 mosquitoes in phase 1 of Ag1000G project, which looked to sample 102 genetic diversity in the wild through the resequencing of wild caught individuals across 103 Africa (Anopheles gambiae 1000 Genomes Consortium 2017), found a polymorphism 104 on average every 2.2 bases of the accessible genome. Nucleotide diversity (π) ranged 105 from ~ 0.008 to ~ 0.015 per population sampled, and even non-degenerate sites (which 106 are expected to be strongly constrained) had an average π of ~0.0025.

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108 Proof of principle for retarding the evolution of resistance to nuclease-based gene-drive 109 by targeting an evolutionarily conserved sequence has recently been demonstrated. A 110 strain of mosquitoes with a CRISPR/Cas9 gene-drive targeting the *doublesex* gene fully 111 suppressed laboratory caged populations of An. gambiae (Kyrou et al. 2018) without 112 selecting for resistance. The CRISPR/Cas9 target sequence in this strain is an 113 intron/exon junction that is highly conserved across the An. gambiae species complex, 114 and only one rare single nucleotide polymorphism was found in the sequence in An. 115 gambiae and An. coluzzii in the Ag1000G data. Consistent with the target site being a 116 region of high functional constraint, monitoring of potential resistant mutations during 117 the cage experiment revealed that although some indels had been introduced by the 118 endonuclease, none of them showed signs of positive selection.

119

This strong constraint at the nucleotide level may exist at other loci in *An. gambiae*.
The Ag1000G project looked for conserved putative CRISPR/Cas9 target sites (18
invariant bases followed by the -NGG motif necessary for Cas9 cleavage) in the 765

123 mosquitoes of Phase 1 of the project, and found 5474 genes containing such sequences.

- 124 However, they note that more variation is likely to be found with further sampling.
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126 Here we take an approach that is likely to be more stringent in identifying functionally 127 constrained sequences by searching for regions that are ultra-conserved across the 128 whole Anopheles genus, which has a most recent common ancestor ~100 million years 129 ago (Neafsey et al. 2015). Although sequence constraint across such a long time scale 130 is not necessary for a good target (as indicated by the *doublesex* locus, which is ultra-131 conserved within the An. gambiae species complex, but shows less conservation outside 132 the complex), we are hypothesising that such highly conserved sequences will contain 133 few polymorphisms in the wild Anopheles gambiae population, and any 134 polymorphisms that do arise (either spontaneously or due to the action of the 135 endonuclease) are likely to have strong fitness costs. We also do not confine our 136 analysis to sequences compatible with any single nuclease architecture (e.g the 5'-137 NGG-3' PAM sequence required by the SpCas9 nuclease) since the range and 138 flexibility of nuclease architectures is constantly expanding, meaning that these 139 requirements may be relaxed (Anders et al. 2016; Chaterjee et al. 2018; Hu et al. 2018). 140 We extracted UCEs from an alignment of the genomes of 21 Anopheles species and 141 strains that was constructed by the Anopheles 16 genomes consortium (Neafsey et al. 142 2015). We characterise the UCEs according to their locations in the genome, and use 143 data from Drosophila orthologues to group genic UCEs according to potential 144 phenotype. We then use the Ag1000G data (765 An. gambiae and An. coluzzii) to see 145 whether these conserved elements contain any variation in natural populations of 146 potential target mosquito species.

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METHODS

149 **Data.**

150 Two sources of genomic data were used in this study: a multi-species alignment from 151 the Anopheles 16 genomes project (Neafsey et al. 2015) and variation data from phase 152 1 of the MalariaGEN An. gambiae 1000 genomes project (Anopheles gambiae 1000 153 Genomes Consortium 2017). The Anopheles 16 genomes project multi-species 154 alignment contains reference genomes from 21 Anopheles species and strains: An. 155 gambiae PEST, An. gambiae s.s., An. coluzzii, An. merus, An. arabiensis, An. 156 quadriannulatus, An. melas, An. christyi, An. epiroticus, An. minimus, An. culicifaces, 157 An. funestus, An. stephensi S1, An. stephensi I2, An. maculatus, An. farauti, An. dirus, 158 An. sinensis, An. atroparvus, An. darlingi, An. albimanus. A description of the methods 159 used to create the alignment is found in Neafsey et al. 2015. Phase 1 of the Ag1000G 160 project comprises 590 An. gambiae and 129 An. coluzzii collected from 9 countries in 161 Africa, plus 46 hybrid individuals from Guinea Bissau. A detailed description of the 162 samples and data is given in Anopheles gambiae 1000 Genomes Consortium 2017.

163

164 Identifying UCEs.

165 To identify invariant regions we used only parts of the multi-species alignment where 166 sequence data was available for all 21 strains. We used Variscan v2.03 (Vilella et al. 167 2005) to find regions of the alignment of 18bp or longer containing no variation. We 168 mapped the resulting regions back to the PEST reference genome using BWA-aln with 169 strict mapping parameters (exact parameters can be provided on request; bwa-0.7.10 170 (Li and Durbin 2010)). Sequences that mapped at multiple places in the genome were 171 included in the analysis, but flagged as 'repeat sequences' as these would not be suitable 172 for use as CRISPR targets. We used BEDTools (Quinlan and Hall 2010) to classify the

genomic location of the UCEs (such as exonic, intronic etc). The AgamP4.12
basefeatures file was used from VectorBase (Giraldo-Calderón *et al.* 2015). Genic
sequences were defined as those with an AGAP gene annotation so include exons,
UTRs and introns. UCEs that partly or wholly fell within genes were classified by us
as genic, and those outside genes were classified as intergenic.

178

For comparison, we used the same method to identify invariant sequences of 18bp or more just in the *An. gambiae* complex species (*An. gambiae PEST, An. gambiae s.s., An. coluzzii, An. merus, An. arabiensis, An. quadriannulatus, An. melas*). We also looked for conservation beyond the *Anopheles* genus by performing a BLAST with default search parameters (Altschul *et al.* 1990) search of the UCEs against *Culex quinquefasciatus* and *Aedes aegypti* reference genomes. From the BLAST results we extracted sequences of 18bp or more with no substitutions, insertions or deletions.

186

187 **Random control sequences.**

So that we could compare the location of UCEs with non-UCEs we extracted 10 randomly distributed sets of control sequences from the multi-species alignment file that were matched to give the same number of sequences with the same base-lengths. To compare variation in the Ag1000G data in UCEs and non-UCEs, we also extracted 10 sets of control sequences from the AgamP4 genome but also matching for genic and intergenic locations.

194

195 **Orthology between species**.

For UCEs that fell within genes, we compared the orthology identifiers betweenAgamP4 and *An. arabiensis* Dongola references genomes, and between *An. gambiae*

198 PEST and An. funestus FUMOZ reference genomes. We chose these species because 199 An. gambiae (and its sister species An. coluzzii), An. arabiensis and An. funestus are the 200 most important malaria vectors in sub-Saharan Africa. An. gambiae PEST is a hybrid 201 strain of An. gambiae and An. coluzzii (previously known as S and M forms of An. 202 gambiae). An. gambiae and An. arabiensis are closely related (in the same species 203 complex) and An. funestus is more distantly related. Genic UCEs were checked for 204 orthology between An. gambiae and An. arabiensis and between An. gambiae and An. 205 *funestus*. Coordinates of UCEs were extracted from the multiple-alignment file for An. 206 arabiensis and An. funestus reference genomes, and annotated with gene names from 207 the basefeatures files Anopheles-arabiensis-Dongola_BASEFEATURES_AaraD1and 208 Anopheles-funestus-FUMOZ_BASEFEATURES_AfunF1.3 (from VectorBase). 209 Orthology identifiers for each gene in each species were found from the 210 ODBMOZ2 Anophelinae database at OrthoDb.org (Kriventseva et al. 2019). 211 Orthology identifiers that match between species indicated that the genes were 212 orthologous. We could not use orthology to directly compare intergenic UCEs, so 213 instead we identified flanking genes for each intergenic UCE in the reference genome 214 of each species, and then compared the orthology identifiers for these genes as before. 215

216 Ontology analysis of genes containing UCEs.

PANTHER software (version 14.0) (Mi *et al.* 2016) was used to categorise the gene
ontology (GO-Slim) terms of the genes containing UCEs. A gene was represented in
the analysis once, regardless of how many UCEs it contained. The genes were clustered
by GO-Slim molecular function, biological process and cellular component terms.

222 Because the Panther categorisation does not take into account how much of the genome 223 is covered by each GO term, we used GOseq (Young et al. 2010) to carry out length-224 bias corrected gene ontology (GO) enrichment analysis, implemented in Galaxy (Afgan 225 et al. 2018). GOseq corrects for gene length using a Wallenius non-central hyper-226 geometric distribution. We used GO-Slim terms extracted from VectorBase (Giraldo-227 Calderón et al. 2015) for AgamP4.12 gene set. GO terms with a Benjamini-Hochberg 228 corrected false discovery rate (FDR) of less than or equal to 0.05 were considered over-229 represented. We also looked for over-representation of GO-Slim terms in the genes 230 flanking integenic UCEs. We were interested to see how our set of UCEs compared to 231 UCEs from *Drosophila* studies, so as well as our full data set, we also performed the 232 GO term analysis on a subset of genes that contained at least one UCE over 50bp long, 233 to make the data comparable.

234

235 Targets for mosquito control.

236 One form of gene drive aimed at population suppression looks to disrupt essential 237 mosquito genes and thereby impose a strong reproductive load on the population as it 238 spreads. UCEs may offer good targets for control of An. gambiae by a gene drive 239 method; if any sequence variation at these sites results in high fitness costs there would 240 be little selective advantage to a mosquito having the variant allele over the gene drive 241 allele. We searched the functional annotations of genes containing UCEs to find genes 242 that may have a suitable function to be targeted for control. Gene descriptions were 243 obtained from VectorBase (Giraldo-Calderón et al. 2015). Gene drives that confer 244 recessive female sterility are particularly potent since both sexes can transmit the drive 245 at very high rates to offspring yet only females homozygous for the drive display the 246 phenotype, which results in a drastic reduction of the population's reproductive

capacity (Burt 2003, Burt and Deredec 2008). P-sterile values were available for some
genes from (Hammond *et al.* 2016). P-sterile is a sterility index based on a logistic
regression model that correlates gene expression features in *Anopheles* with the
likelihood that mutations of the gene produce female sterile alleles in the model
dipteran *Drosophila melanogaster* (Baker *et al.* 2011).

252

To narrow down the gene list to potential vector control targets, we leveraged the large amount of phenotype data already available for *Drosophila* mutants. Where possible, *Drosophila* orthologues were identified for genes containing UCEs (in Vectorbase). We used ID converter in FlyBase (Gramates *et al.* 2017) to batch convert *Drosophila* gene identifiers into alleles associated with the genes (FBal numbers). The alleles have associated phenotype data provided by the research community; we searched for phenotypes conferring female sterility or recessive lethality.

260

261 Transcription factor binding site motifs in UCEs.

We used the 'Find Individual Motif Occurrences' (FIMO, Grant *et al.* 2011) scanning module (MEME suite 4.12.0, Bailey *et al.* 2009) to look for transcription factor binding motifs in UCEs and controls. The UCEs were scanned for known insect transcription factor binding sites using weighted matrices from the JASPER CORE collection (Insect position frequency matrices 8th release (2020), Khan *et al.* 2018). The results were filtered by q-value to account for multiple tests. A cut-off of q<0.05 was used.

268

269 Variation at UCE locations in Ag1000G data.

270 Using the final filtered variant file from phase 2 of the Ag1000G project (described at

271 https://www.malariagen.net/data/ag1000g-phase-2-ar1) we extracted single nucleotide

polymorphisms for the UCEs identified above, and for matched non-UCE regions.
Diversity statistics were calculated in scikit-allel v1.3.2 (Miles et al 2020): number of
segregating sites (s), nucleotide diversity (pi) and the neutrality test Tajima's D (Tajima
1989).

276

Data used in this study are publicly available from the *Anopheles* 16 genomes
consortium and the *Anopheles* gambiae 1000 Genomes project. Data generated in this
study are given in the supplementary tables.

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RESULTS

282 Ultra-conserved regions from the multi-species alignment.

283 Much of the MAF file does not include alignments of all 21 species and strains (see 284 Table S8 in Neafsey et al. 2015). The total number of aligned bases from which we 285 extracted the UCEs was 17,095,206 (7.4% of the AgamP4 reference genome (Suppl 286 Table 1). A total of 8338 invariant regions of 18bp or more were identified; 1675 on 287 chromosome arm 2L, 3015 on chromosome arm 2R, 1375 on chromosome arm 3L, 288 2188 on chromosome arm 3R and 85 on chromosome X (Table 1; we have also included 289 the same metrics at different evolutionary timescales for comparison). The longest UCE 290 was 164bp. Genomic coordinates of the UCEs relative to the Anopheles gambiae PEST 291 reference genome are given in Supplementary table 2. The UCEs were distributed 292 throughout the chromosomes, but were significantly under-represented on the X 293 chromosome (See Supplementary figure 1 and Supplementary Table 1). The X 294 chromosome is already under-represented in the MAF as it was less alignable than other 295 chromosomes (see Figure 2 in Neafsey et al 2015). It is well established that the X 296 chromosome shows higher differentiation between species than autosomes (due to

297 'Haldanes Rule' and the 'Large X effect') and genomic studies have reinforced this 298 observation (Presgraves 2018). However, the under-representation in the MAF is not sufficient to explain the paucity of UCEs on the X. In the Anopheles genus, the X 299 300 chromosome was observed to have undergone particularly dynamic evolution, with 301 chromosome rearrangements at a rate of 2.7 times higher than the autosomes, and a 302 significant degree of observed gene movement from X to other chromosomes relative 303 to Drosophila (Neafsey et al 2015). This dynamic evolution of the chromosome may 304 explain why it would be less likely to contain functional sequences that require 305 conservation at the nucleotide level.

306

307 Size distributions of the UCEs are shown in Supplementary figure 2. In the autosomal 308 genic UCEs there is pattern of a jump in frequency every 3 bases, indicating the 309 tendency for runs of ultra-conserved bases to neither start nor end on third codon 310 positions in coding regions. UCEs are significantly more AT-rich than random 311 sequences (64% and 54% respectively, t-test p<0.001).

312

| | 2L | 2R | 3L | 3R | Х |
|---------------------|------------|------------|------------|------------|-----------|
| Gambiae complex | | | | | |
| No. UCEs | 452,281 | 612,824 | 376,383 | 498,473 | 99,561 |
| No. Invariant bases | | | | | |
| within UCEs | 15,365,491 | 21,350,270 | 12,886,437 | 17,278,830 | 3,338,454 |
| Anopheles | | | | | |
| No. UCEs | 1,675 | 3,015 | 1,375 | 2,188 | 85 |

| within UCEs | 45,916 | 81,186 | 37,102 | 59,055 | 2,299 |
|-----------------------|--------|--------|--------|--------|-------|
| Anopheles+Aedes | | | | | |
| No. UCEs | 278 | 344 | 193 | 293 | 15 |
| No. Invariant bases | | | | | |
| within UCEs | 8,161 | 10,275 | 5,499 | 8,339 | 456 |
| Anopheles+Culex | | | | | |
| No. UCEs | 279 | 350 | 202 | 310 | 16 |
| No. invariant bases | | | | | |
| within UCEs | 8,201 | 10,184 | 5,716 | 8,691 | 503 |
| Anopheles+Aedes+Culex | | | | | |
| No. UCEs | 192 | 247 | 133 | 217 | 12 |
| No. invariant bases | | | | | |
| within UCEs | 5,995 | 7,579 | 3,989 | 6,391 | 393 |

313

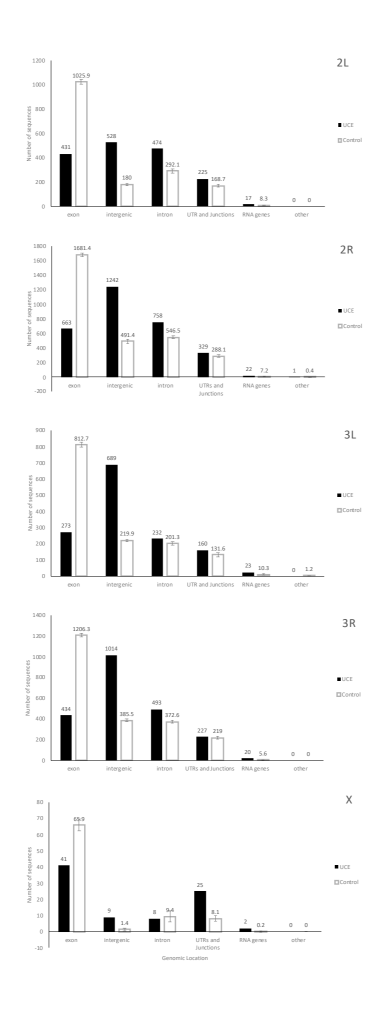
No. Invariant bases

Table 1. Number of ultra-conserved sequences of 18bp or more, and total number of
invariant sites within these sequences. *Gambiae* complex = 7 species and strains; *Anopheles* = 21 species and strains; *Culex* = *Culex quinquefasciatus* reference genome; *Aedes* = *Aedes aegypti* reference genome.

318

We annotated the UCEs in BEDtools to identify where they were found in the genome with regards to exons, introns, UTRs, intergenic regions etc (Figure 1). The 21-genome aligned parts of the MAF file from which we extracted the UCEs is not representative of the reference genome with respect to these features, so we extracted randomly

| 323 | distributed sets of 'control' sequences from the MAF, and only from sequences where |
|-----|--|
| 324 | all 21 genomes were aligned. These control sequences were matched to give the same |
| 325 | number of sequences with the same base-lengths as the UCEs, and were compared with |
| 326 | the UCE locations to see whether the UCEs were randomly distributed. The UCE |
| 327 | sequences were significantly over-represented (compared to control sequences) in |
| 328 | intergenic regions (42% vs. 15%, ANOVA, p<0.05) and in RNA genes (1% vs. 0.4%, |
| 329 | p<0.05), and less frequent in exons (22% vs. 57%, p<0.05). The MAF itself is heavily |
| 330 | skewed to including exonic sequences, as only about 7% of the An. gambiae genome |
| 331 | as a whole is exonic (Holt et al. 2002). |
| 332 | |
| 333 | |



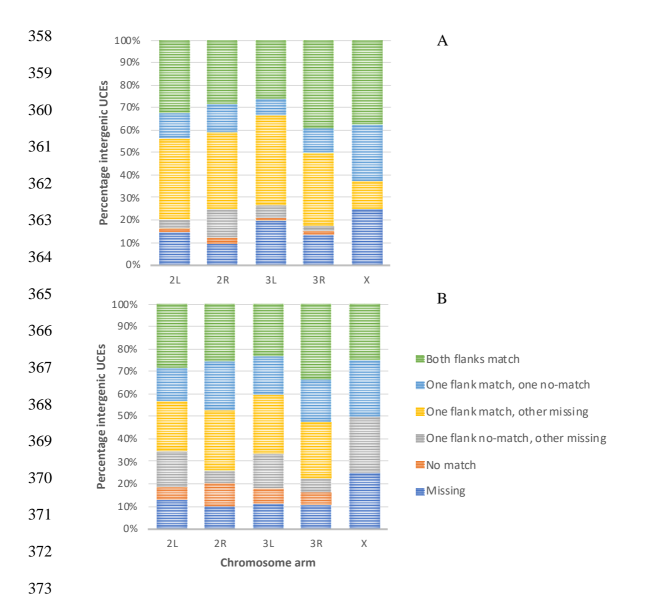
336 Figure 1. Distribution of UCE and non-UCE control sequences according to

337 genomic location. Genomic locations annotated with BEDtools. Control error bars =
338 standard deviation for 10 control data sets of sequences of matched length and number
339 to the UCEs, extracted randomly from the MAF, only from regions where data for all
340 21 genomes is present.

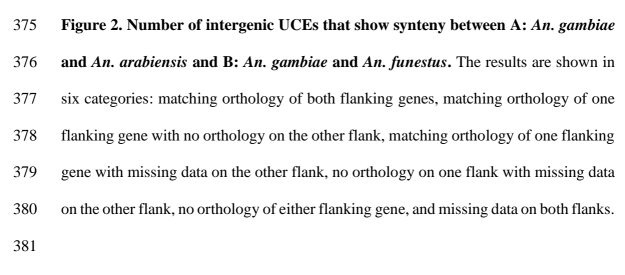
341

342 Orthology between important vector species.

343 To ensure that the UCEs were not falling in sequences that had aligned across the 21 344 genomes by chance, we checked for orthology between some species in the UCEs. For 345 UCEs that fell within genes this was done by simply by comparing orthology identifiers 346 (from OrthDB.org) between An. gambiae and An. arabiensis, and between An. gambiae 347 and An. funestus. For An. gambiae and An. arabiensis, 94% of autosomal genes 348 containing UCEs shared orthology. For An. gambiae and An. funestus, this number was 349 87%. The amount of orthology fell to 54% and 63% for genes containing UCEs on the 350 X chromosome. For UCEs that were intergenic, we looked at orthology of the flanking 351 genes. The results fell into six categories: orthology of both flanking genes, orthology 352 of one flanking gene with no orthology on the other flank, orthology of one flanking 353 gene with missing data on the other flank, no orthology on one flank with missing data 354 on the other flank, missing data on both flanks, and no orthology of either flanking 355 gene. Ignoring missing data, 92% of intergenic UCEs showed full or half synteny 356 between An. gambiae and An. arabiensis, and 77% of UCEs showed full or half synteny 357 between An. gambiae and An. funestus (Figure 2).



374



382 Functional profile analysis of the genes containing UCEs via GO term enrichment

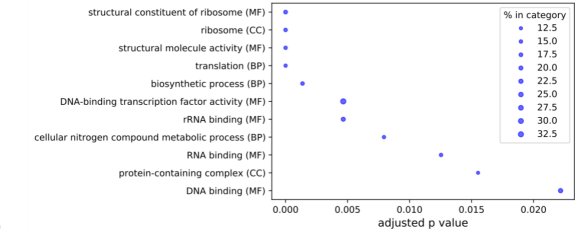
383 Of the 13,796 genes annotated in the Anopheles gambiae PEST gene set Agam4.12,

1,601 (12.9%) had at least one UCE. We clustered the genes based on GO-Slim terms

- for molecular function, biological process and cellular component (Suppl Figure 3).
- 386

Because the clustering does not take into account the amount of the genome covered
by each GO class, we carried out length-bias corrected GO term enrichment analysis.
This showed that certain functional groups were over-represented compared with the

390 whole *Anopheles* PEST reference gene set (Figure 3).



391

Figure 3. GOseq GO term enrichment analysis with length-bias correction. GO-Slim categories were extracted from the AgamP4.12 gene set. Results are shown for categories that were enriched with an FDR adjusted p-value<0.05. Bubble size is relative to the percentage of AgamP4.12 genes in a GO category that were present in the UCE gene set. MF=molecular function; BP=biological process; CC=cellular component.

398

399 In the genes containing UCEs over 50bp long, only 4 categories were over-represented:

400 transmembrane transporter activity (MF), transmembrane transport (BP), transport

401 (BP) and protein-containing complex (CC), (adjusted p values 0.0047, 0.0047, 0.0272, 402

403

404 Genes flanking intergenic UCEs were enriched for the GO-Slim categories DNA 405 binding (MF), DNA-binding transcription factor activity (MF) and anatomical structure

development (BP) (adjusted p values 4.16E-06, 1.46E-05 and 0.016 respectively).

407

406

408 Potential targets for vector control.

0.0272 respectively).

409 AGAP001189 (odorant-binding protein 10) contained the highest number of invariant 410 bases in UCEs (1215/135306). Nine genes contained UCEs longer than 100bp, of which 411 3 are annotated as being involved in ion transport. These include the voltage gated 412 sodium channel gene (VGSC, AGAP004707), which is a target for (and therefore has 413 a significant role in conferring resistance to) some of the main classes of insecticides 414 used for malaria vector control. VGSC is one of the most conserved genes we found, 415 containing 13 UCEs with a total of 507 invariant bases, of which 91% were in exons 416 and most coded for trans-membrane domains. A total of 357 genes contained 100 or 417 more invariant bases. A full list of genes containing UCEs is given in Supplementary 418 table 3.

419

420 Eleven genes containing UCEs had a p-sterile score of greater than 0.5 implying that 421 they could be good targets to affect female fertility.

422

423 Drosophila orthologues were identified for 1309 of the 1601 genes containing UCEs. 424 Allele and phenotype classes for these genes were extracted from Flybase where 425 available. For an effective population suppression gene-drive, the target would affect

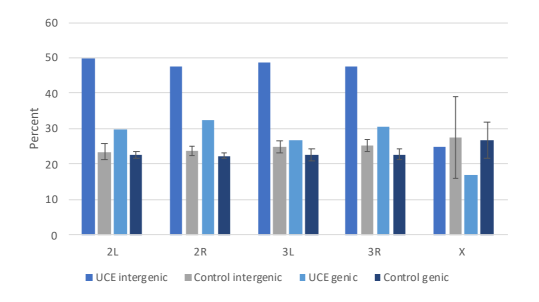
426 female fertility or impose a genetic load as a homozygote, so we extracted UCE 427 containing genes that have *Drosophila* orthologues annotated with a female sterile term 428 or a lethal recessive term (shown in Supplementary table 3). 177 genes containing 429 UCEs have *Drosophila* orthologues with an allele phenotype affecting female fertility, 430 and 367 genes have *Drosophila* orthologues with an allele conferring a lethal recessive 431 phenotype.

432

433 **Transcription factor binding motifs in UCEs.**

434 DNA binding motifs recognised by transcription factors might be expected to be 435 constrained and hence enriched for UCEs since this protein:DNA interaction is 436 sequence-specific. The FIMO search found that 38% of UCEs contained hits for insect 437 transcription factor binding sites with a q value <0.05 (48% of intergenic and 30% of 438 genic UCEs). This compared with 23% for control (non-conserved) sequences of the 439 same number and length. On the X chromosome, where data is sparse (only 8 intergenic 440 and 75 genic UCEs (Figure 4)), the numbers of transcription factors binding sites were 441 not significantly different between UCEs and controls.





444 Figure 4. Percent of UCEs and control sequences that contain at least one insect

445 **transcription factor binding motif.** Control error bars = standard deviation for 10 446 control data sets. UCEs were searched for known insect transcription factor binding 447 sites from the JASPER CORE collection (Insect position frequency matrices 8th release 448 (2020), Khan *et al.* 2018). The results were filtered by q-value to account for multiple 449 tests. A cut-off of q<0.05 was used.

450

451 Genetic variation at UCE locations in Ag1000G data.

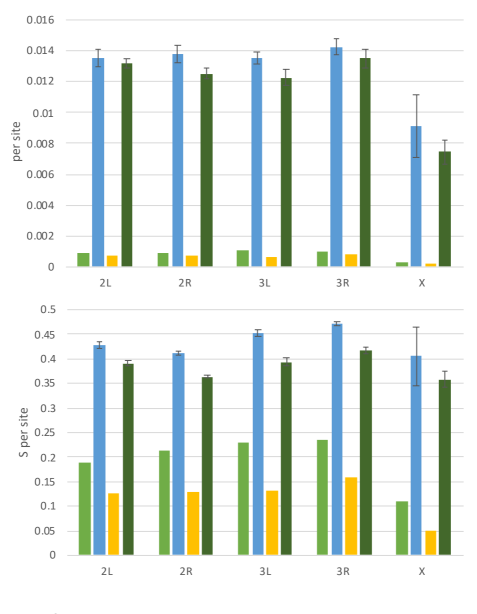
452 In order to see whether sequences that are ultra-conserved across the Anopheles genus 453 show variation in wild mosquito populations, we searched for single nucleotide 454 polymorphisms (SNPs) in the 1142 samples from phase 2 of the Ag1000G project. 455 There were significantly fewer sites containing polymorphisms in UCEs than control 456 sequences (Figure 5 middle), and those SNPs that were present were at significantly 457 lower frequency (Figure 5 top). Of the 8338 UCEs, 1213 (15%) contained no SNPs in 458 the 1142 samples (229 on 2L, 470 on 2R, 226 on 3L, 259 on 3R and 29 on X). Tajima's 459 D is significantly more negative for UCEs than controls, with the exception of X 460 chromosome intergenic sequences (Figure 5 bottom). Negative values of Tajima's D 461 are expected for sequences under purifying selection.

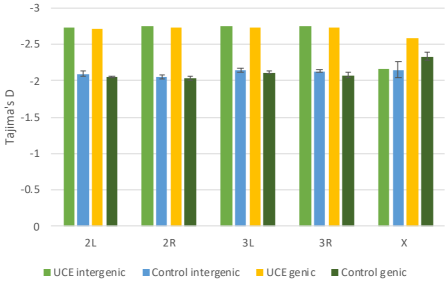
462

The Ag1000G study (*Anopheles gambiae* 1000 genomes consortium 2017), performed a search within the Phase 1 data to look for potential Cas9 targets (non-overlapping exonic invariant sequences of 21bp, ending in the 'NGG" motif). They identified 13 genes containing sequences corresponding to this motif. None of these 13 genes contained UCEs as defined by our study, so these genes also did not overlap with invariant sequences in Ag1000G data found here. We did not confine our search for

- 469 UCEs to current Cas9 target site restrictions because of the growing possibility of
- 470 relaxation of these constraints. However, for completeness we looked within our final
- 471 set of UCEs for the Cas9 motif (18bp followed by -NGG, or CC- followed by 18bp).
- 472 We found 1997 (24%) UCEs contained suitable targets for Cas9.

473





476 Figure 5. Genetic diversity per chromosome arm in 1142 Anopheles gambiae s.l.

477 samples in UCE locations. Top: nucleotide diversity (π); middle: segregating sites

478 (s); bottom: Tajima's D. Calculations were made in scikit-allel v1.3.2 (Miles et al
479 2020).

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- 481

DISCUSSION

482 Similarities and differences of Anopheles UCEs with UCEs from *Drosophila*.

483 Despite approximately 100 million years since their most recent common ancestor, we 484 identified in the Anopheles genus over 8000 sequences of 18bp or more where there 485 was no nucleotide variation across the alignment of 21 species and strains. By 486 coincidence, this is approximately the same span of evolutionary time covered in the 487 human/mouse/rat data set in which UCEs were originally identified (Bejerano et al. 488 2004). 481 UCEs of more than 200bp were observed between these genomes, but the 489 longest we found in the Anopheles genus was 164bp. This is consistent with previous 490 reports that UCEs are fewer and shorter in insects (mainly *Drosophila*) than vertebrates 491 (Makunin et al. 2013; Glazov et al. 2005). Our criteria for identifying UCEs were 492 somewhat different than those used previously. First, we only considered sequences 493 that were present in all 21 species/strains in the alignment; some of these species have 494 poorly assembled genomes, so this may have reduced the number of UCEs that we 495 uncovered. Second, we also included invariant stretches of 18bp or more, whereas 496 Drosophila studies have used cut-offs of 50bp (Glazov et al. 2005, Warnefors et al. 497 2016), 80bp (Kern et al. 2015) or 100bp (Makunin et al. 2013). Despite this we see 498 some similarities between our UCEs and UCEs found in Drosophila. UCEs are located 499 in all parts of the genome and, like Drosophila, the majority are found in intergenic 500 regions and introns. We also found that junction locations (e.g. intron-exon, exon-

501 intergenic etc) are over-represented compared to random sequences, which in 502 Drosophila has been linked to conservation of splice-sites (Glazov et al. 2005; 503 Warnefors et al. 2016). Another similarity with Drosophila is the high proportion of 504 genes with the GO terms 'binding' and 'transporter activity' (Kern et al. 2015; Glazov 505 et al. 2005). In Drosophila, ion channel/transporter genes have been shown to undergo 506 extensive RNA editing (Hanrahan et al. 2000; Hoopengardner et al. 2003; Rodriguez 507 et al. 2012) which is thought to explain the high level of conservation. This is because 508 RNA adenosine deaminases require double stranded RNA as a substrate, which means 509 that there is likely to be strong selection at the nucleotide level. The high number of 510 UCEs in Anopheles ion channel/transporter genes suggests a similar mechanism is 511 responsible for the high conservation in the Anopheles genus. However, these genes 512 are extremely long and are not over-represented in the UCE data when a length-bias 513 corrected analysis is carried out in GOseq. In the GOseq analysis, the most over-514 represented molecular functions are mostly involved in binding or structure. 515 Transcription factor binding, enzyme binding and nucleic acid binding have also been 516 shown to be associated with ultra-conservation in both invertebrates and mammals 517 (Bejerano et al. 2004; Glazov et al. 2005). A noteworthy addition to highly represented 518 GO terms in Anopheles that has not been reported in Drosophila, is the category of 519 'catalytic activity' genes, although again, these were not over-represented when gene 520 length was taken into account. When the GO term clustering was carried out on genes 521 containing UCEs of 50bp or more in length, we found that the category reduced from 522 28% to 18% suggesting that these shorter ultra-conserved regions most likely code for 523 a small number of key residues around an active site.

525 The high number of UCEs that we observe in intergenic regions and introns suggests 526 that we have found numerous unannotated locations in the Anopheles PEST reference 527 genome with putative regulatory functions. At least 70% were syntenic between An. 528 gambiae/An. arabiensis and An. gambiae/An. funestus so the location of these highly 529 conserved sequences is likely to be important. A GOseq analysis of the genes flanking these intergenic sequences showed significant over-representation of genes with DNA 530 531 binding GO terms. Sequences that are ultra-conserved at the nucleotide level across a 532 long evolutionary time have been shown to be linked to regulatory functions such as 533 cis-regulation of genes (e.g. enhancers, insulators, silencers) and RNA genes (e.g. 534 miRNA, snRNA), likely because of the sequence-specific nature of protein:nucleotide 535 or nucleotide:nucleotide interactions. 19 of the 77 miRNA genes that are annotated in 536 the Anopheles PEST genome were included in our set of UCEs (other miRNAs may 537 contain ultra-conserved regions that did not meet our criteria). We also found known 538 insect transcription binding factors in 48% of the intergenic UCEs.

539

540 **Polymorphisms in UCEs in** *Anopheles* **populations.**

541 All of the UCEs discovered from the alignment of the reference genomes of 21 542 Anopheles species were also found to be highly conserved in the sample of 1142 wild 543 caught mosquitoes sequenced in phase 1 of Ag1000G. Although the majority of UCEs 544 contained one or more polymorphisms, they were almost all rare. 1213 UCEs showed 545 no polymorphisms at all in this sample. This does not rule out the existence of 546 polymorphisms in the wild populations, but does imply that there may be strong 547 constraint at a nucleotide level that means alteration of the sequence either naturally or 548 by the action of a gene-drive may have a strong fitness cost. This would need to be 549 tested experimentally as different levels of underlying functional constraint may have

different fitness costs. For instance, deletion of certain ultra-conserved sequences in mice gave no discernible fitness cost (Ahituv *et al.* 2007), but a similar experiment in *Drosophila* showed promise, with 4 out of 11 UCEs with inserted transposons having a lethal recessive phenotype (Makunin *et al.* 2013). For a resistance-proof gene drive, selecting target sites that show high levels of conservation is a good starting point, but the targets would need to be tested under selection pressure to ensure that functional mutants do not arise.

557

558 UCEs and vector control

559 UCEs occur within many genes that could have potential for vector control. Nearly 200 560 genes have *Drosophila* orthologues with an allele phenotype affecting female fertility, 561 and over three hundred genes have Drosophila orthologues with an allele conferring a 562 lethal recessive phenotype. These phenotypes could both be used for a population 563 suppression strategy i.e. to reduce the numbers of mosquitoes to a level where malaria 564 could no longer be transmitted (Deredec et al. 2011). More investigation would be 565 needed to see whether disrupting the genes at the ultra-conserved loci gives the same 566 phenotype in Anopheles. There are also genes that confer recessive phenotypes in 567 Drosophila such as 'flightless' or 'behaviour defective' that could also be used for 568 population suppression, or for a population modification type of strategy, where instead 569 of reducing the mosquito population it is replaced by a strain that cannot transmit 570 malaria (Carballar-Lejarazú et al. 2018). Precise targeting of sequences using 571 CRISPR/Cas9 gene editing had made testing for these phenotypes feasible.

572

573 Another potential mode of mosquito genetic alteration that has not yet been explored 574 would be to target sequences involved in gene regulation. Many ultra-conserved

sequences in mammals and invertebrates are thought to be involved in regulation ofgenes important in development.

577

578 Targeting a sequence that is conserved between species means that the gene drive could 579 spread between closely related species that hybridise in the wild. For this to happen the 580 species would need to mate in the wild, produce some fertile offspring, and be able to 581 express the CRISPR enzyme using the same promoter. Three species (An. gambiae, An. 582 *coluzzii* and *An. arabiensis*) are responsible for the majority of malaria transmission in 583 some parts of sub-Saharan Africa, and are known to hybridise in nature (e.g. Weetman 584 et al. 2014, Anopheles gambiae 1000 Genomes Consortium 2017, Fontaine et al 2015). 585 For effective vector control it would be desirable to be able to reduce or alter all three 586 species with one construct. The gene drive would not spread to Anopheles species that 587 do not mate in the wild, so would not spread beyond the Anopheles gambiae species 588 complex. However, if a particular target site was proved to be effective for vector 589 control in An. gambiae, a gene drive targeting an orthologous site could be developed 590 in the laboratory for other important malaria vectors such as An. funestus.

591

592

CONCLUSION

Thousands of short genomic regions exist that are conserved across the *Anopheles* genus. These sequences show many of the same traits as ultra-conserved elements found in *Drosophila* (such as an association with gene regulation and ion channel activity). Our list of ultra-conserved elements in the *Anopheles* genus should provide a valuable starting point for the selection and testing of new targets for gene-drive modification in the mosquitoes that transmit malaria. Focussing on sequences that have

| 599 | been tightly conserved over long evolutionary time has promise for mitigating against |
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| 600 | or slowing the development of resistant alleles in the wild population. |
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| 803 | |
| 804 | |
| 805 | Authors' contributions |
| 806 | SO and AB jointly devised the study; SO performed and guided data analysis and wrote |
| 807 | the manuscript; AF carried out data analysis; SF designed and performed preliminary |
| 808 | GO term analysis; TD assisted in bioinformatics; TN and AC gave advice on analysis; |
| 809 | all authors provided editorial comments and read and approved the final manuscript. |
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| 815 | |
| 816 | |