

Ultra-conserved sequences in the genomes of highly diverse *Anopheles* mosquitoes, with implications for malaria vector control

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Abstract

DNA sequences that are exactly conserved over long evolutionary time scales have been observed in a variety of taxa. Such sequences are likely under strong functional constraint and they have been useful in the field of comparative genomics for identifying genome regions with regulatory function. A potential new application for these ultra-conserved elements (UCEs) has emerged in the development of gene drives to control mosquito populations. Many gene drives work by recognizing and inserting at a specific target sequence in the genome, often imposing a reproductive load as a consequence. They can therefore select for target sequence variants that provide resistance to the drive. Focusing on highly conserved, highly constrained sequences lowers the probability that variant, gene drive-resistant alleles can be tolerated. Here, we search for conserved sequences of 18 bp and over in an alignment of 21 *Anopheles* genomes, spanning an evolutionary timescale of 100 million years, and characterize the resulting sequences according to their location and function. Over 8000 UCEs were found across the alignment, with a maximum length of 164 bp. Length-corrected gene ontology analysis revealed that genes containing *Anopheles* UCEs were found in 48% of intergenic *Anopheles* UCEs. When we looked at the genome sequences of 1142 wild-caught mosquitoes, we found that 15% of the *Anopheles* UCEs contained no polymorphisms. Our list of *Anopheles* UCEs should provide a valuable starting point for the selection and testing of new targets for gene-drive modification in the mosquitoes that transmit malaria.

Keywords: Anopheles; gene drive; conserved; malaria

Introduction

DNA sequences that are highly conserved over long evolutionary timescales have been identified in many organisms. Some of these sequences show complete conservation at the nucleotide level and are often known as ultra-conserved elements (UCEs). Originally, UCEs were defined as sequences of at least 200 bp that were identical between human, mouse, and rat genomes (Bejerano *et al.* 2004). Subsequently, the search for UCEs has been extended to other vertebrates, insects, and plants (*e.g.* Siepel *et al.* 2005; Baxter *et al.* 2012; Makunin *et al.* 2013; Quattrini *et al.* 2018), and to sequences of length 50 bp or more.

There are several reasons why UCEs are of interest. First, in the field of comparative genomics, UCEs are thought to represent functionally important regions. While there is still some mystery around why sequences might be conserved at the nucleotide level over long evolutionary timescales, it has been shown that UCEs: (1) often are involved in the regulation of transcription of genes, especially essential genes involved in development (*e.g.* (Visel *et al.* 2008); (2) may have a role in chromosomal structure (*e.g.* Chiang

et al. 2008); and (3) are sometimes non-coding RNA genes (e.g. Kern et al. 2015). Even UCEs in protein-coding regions may have multi-functional roles (Warnefors et al. 2016). Second, UCEs can act as probes to facilitate genomic sequencing of non-model organisms using sequence-capture methods (Faircloth et al. 2012). Third, alterations in UCEs have been shown to have an association with human cancers (e.g. Calin et al. 2007; Lin et al. 2012).

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

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malaria parasite (Gantz et al. 2015). Both methods currently rely on nuclease-based synthetic gene drive systems that introduce a desired trait at a precise genomic location, spreading it in a target population at such a rate that outweighs fitness costs associated with the trait (Burt 2003). The technologies include RNA-guided endonucleases (such as CRISPR/Cas9) and homing endonucleases (Windbichler et al. 2011; Jinek et al. 2012). These enzymes recognize and cleave a particular target size of about 18 bp. When the sequence coding for these enzymes is engineered into its own target site in the genome and is expressed in the germline, it creates a double-strand break in the homologous chromosome. The break will usually be repaired by homology-directed repair using the drive-containing chromosome as a template which results in conversion of the repaired chromosome to also contain the drive element in greater than the usual 50% inheritance rate among the gametes. An efficient gene drive can be inherited by almost 100% of progeny (Hammond et al. 2016). Theoretical and laboratory studies have shown that changes to the recognition site can result in alleles that cannot be recognized or cleaved. If these alleles confer increased fitness compared with the wild-type allele in the presence of the gene drive they can be expected to spread and retard the spread of the gene drive (Deredec et al. 2008; Hammond et al. 2017; Unckless et al. 2017). For population suppression gene drives that are designed to impair essential genes, the selection pressure for resistance alleles to arise is high. These alleles can arise from standing variation at the target site in a wild population or may come about from the action of the endonuclease. This is because non-homologous end joining can sometimes repair the double-strand break, and random insertions and deletions can be introduced to the target site.

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

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

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 mosquitoes of Phase 1 of the project, and found 5474 genes containing such sequences. However, they note that more variation is likely to be found with further sampling.

Here, we take an approach that is likely to be more stringent in identifying functionally constrained sequences by searching for regions that are ultra-conserved across the whole Anopheles genus, which has a most recent common ancestor ~100 million years ago (Neafsey et al. 2015). Although sequence constraint across such a long-time scale is not necessary for a good target (as indicated by the doublesex locus, which is ultra-conserved within the An. gambiae species complex, but shows less conservation outside the complex), we are hypothesizing that such highly conserved sequences will contain few polymorphisms in the wild Anopheles gambiae population, and any polymorphisms that do arise (either spontaneously or due to the action of the endonuclease) are likely to have strong fitness costs. We also do not confine our analysis to sequences compatible with any single nuclease architecture (e.g. the 5'-NGG-3' PAM sequence required by the SpCas9 nuclease) since the range and flexibility of nuclease architectures is constantly expanding, meaning that these requirements may be relaxed (Anders et al. 2016; Chatterjee et al. 2018; Hu et al. 2018). We extracted UCEs from an alignment of the genomes of 21 Anopheles species and strains that were constructed by the Anopheles 16 genomes consortium (Neafsey et al. 2015). We used data from Drosophila orthologues to group genic UCEs according to potential phenotype. We then use the Ag1000G data (1142 An. gambiae and An. coluzzii) to see whether these conserved elements contain any variation in natural populations of potential target mosquito species.

The main aim of our study was to identify potential targets for vector control, but as these are the first UCEs to be identified from an alignment of the *Anopheles* genus, we also characterized the UCEs according to their locations in the genome, and performed functional classification analyses to see how they compare with UCEs identified in other taxa.

Materials and methods Data

Two sources of genomic data were used in this study: a multispecies alignment file (MAF) from the Anopheles 16 genomes project (Neafsey et al. 2015) and variation data from phase 2 of the MalariaGEN An. gambiae 1000 genomes project (Anopheles gambiae 1000 Genomes Consortium et al. 2017). The Anopheles 16 genomes project multi-species alignment contains reference genomes from 21 Anopheles species and strains: An. gambiae PEST, An. gambiae s.s., An. coluzzii, An. merus, An. arabiensis, An. quadriannulatus, An. melas, An. christyi, An. epiroticus, An. minimus, An. culicifaces, An. funestus, An. stephensi S1, An. stephensi I2, An. maculatus, An. farauti, An. dirus, An. sinensis, An. atroparvus, An. darlingi, and An. albimanus. A description of the methods used to create the alignment is found in Neafsey et al. (2015). Phase 2 of the Ag1000G project comprises 1142 An. gambiae, An. coluzzii and hybrids, collected from 13 countries in Africa (The Anopheles gambiae 1000 Genomes Consortium (2017): Ag1000G phase 2 AR1 data release).

Identifying UCEs

To identify invariant regions, we used only parts of the multi-species alignment where sequence data were available for all 21 strains. We used Variscan v2.03 (Vilella *et al.* 2005) to find regions of the alignment of 18 bp or longer containing no variation. We mapped the resulting regions back to the PEST reference genome using BWA-aln with strict mapping parameters (zero edit distance, no gap opening allowed; bwa-0.7.10 (Li and Durbin 2010)). Sequences that mapped at multiple places in the genome were included in the analysis but flagged as "repeat sequences" as these would not be suitable for use as CRISPR targets. A recent bioinformatics resource has been published that provides an automated alternative to these methods (Kranjc *et al.* 2021).

We used BEDTools (Quinlan and Hall 2010) to classify the genomic location of the UCEs (such as exonic, intronic, etc). The AgamP4.12 base features file was used from VectorBase (Giraldo-Calderón et al. 2014). 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. Results are presented per chromosome arm; *Anopheles* chromosomes contain fixed and polymorphic inversions that can impact evolutionary influences, so treating the autosomes as a single unit would not be appropriate.

For comparison, we used the same method to identify invariant sequences of 18 bp or more just in the An. gambiae complex species (An. gambiae PEST, An. gambiae s.s., An. coluzzii, An. merus, An. arabiensis, An. quadriannulatus, and An. melas). We also looked to see whether the Anopheles UCEs were conserved at an older evolutionary scale in Culex quinquefasciatus and Aedes aegypti. The simplest way to achieve this was to use blastn with default parameters (Altschul et al. 1990) in VectorBase to search for similar sequences in the Aedes and Culex reference genomes (AaegL5.0 and CulPip1.0). Because many of our UCEs were short (18 bp) and may have random hits in the similarity search, we extended the sequences with 50 bp in either side from the An. gambiae PEST reference genome. The similarity results from blastn were filtered manually to extract DNA sequences of 18bp or more that were completely invariant, i.e., included no substitutions or indels, within the Anopheles UCE sequences.

Random control sequences

So that we could compare the location of UCEs with non-UCEs, we used custom Python scripts to extract 10 independent randomly distributed sets of control sequences from the MAF (only from locations where aligned sequences for all 21 species were present) 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 independent sets of control sequences from the AgamP4 genome but also matching for genic and intergenic locations. The custom scripts can be found on GitHub (https://github.com/soloughlinhub?tab=repositories).

Orthology between species

For UCEs that fell within genes, we compared the orthology identifiers between AgamP4 and An. arabiensis Dongola reference genomes, and between An. gambiae PEST and An. funestus FUMOZ reference genomes. We chose these species because An. qambiae (and its sister species An. coluzzii), An. arabiensis, and An. funestus are the most important malaria vectors in sub-Saharan Africa. An. gambiae PEST is a hybrid strain of An. gambiae and An. coluzzii (previously known as S and M forms of An. gambiae). An. gambiae and An. arabiensis are closely related (in the same species complex) and An. funestus is more distantly related. Genic UCEs were checked for orthology between An. gambiae and An. arabiensis and between An. gambiae and An. funestus. Coordinates of UCEs were extracted from the multiple-alignment file for An. arabiensis and An. funestus reference genomes, and annotated with gene names from the base features files Anopheles-arabiensis-Dongola_ BASEFEATURES_AaraD1and Anopheles-funestus-FUMOZ_BASE

FEATURES_AfunF1.3 (from VectorBase). Orthology identifiers for each gene in each species were found from the ODBMOZ2_Anophelinae database at OrthoDb.org (Kriventseva et al. 2019). Orthology identifiers that match between species indicated that the genes were orthologous. We could not use orthology to directly compare intergenic UCEs, so instead, we identified flanking genes for each intergenic UCE in the reference genome of each species and then compared the orthology identifiers for these genes as before.

Ontology analysis of genes containing UCEs

PANTHER software (version 14.0) (Mi et al. 2017) was used to categorize 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. We performed functional classification by GO-Slim molecular function, biological process, and cellular component terms.

Because the Panther functional classification tool does not take into account how much of the genome is covered by each GO term, we used GOseq (Young et al. 2010) to carry out lengthbias corrected gene ontology (GO) enrichment analysis, implemented in Galaxy (Afgan et al. 2018). GOseq corrects for gene length using a Wallenius non-central hypergeometric distribution. We used GO-Slim terms extracted from VectorBase (Giraldo-Calderón et al. 2014) for AgamP4.12 gene set. GO terms with a Benjamini-Hochberg corrected false discovery rate (FDR) of ≤ 0.05 were considered over-represented. We also looked for over-representation of GO-Slim terms in the genes flanking intergenic UCEs. We were interested to see how our set of UCEs compared with UCEs from Drosophila studies, so as well as our full data set, we also performed the GO term analysis on a subset of genes that contained at least one UCE over 50 bp long, to make the data comparable.

Targets for mosquito control

One form of gene drive aimed at population suppression looks to disrupt essential mosquito genes and thereby impose a strong reproductive load on the population as it spreads. UCEs may offer good targets for control of An. gambiae by a gene drive method; if any sequence variation at these sites results in high fitness costs, there would be little selective advantage to a mosquito having the variant allele over the gene drive allele. We searched the functional annotations of genes containing UCEs to find genes that may have a suitable function to be targeted for control. Gene descriptions were obtained from VectorBase (Giraldo-Calderón et al. 2014). Gene drives that confer recessive female sterility are particularly potent since both sexes can transmit the drive at very high rates to offspring yet only females homozygous for the drive display the phenotype, which results in a drastic reduction of the population's reproductive capacity (Burt 2003, Burt and Deredec, 2018). P-sterile values were available for some genes (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).

To narrow down the gene list to potential vector control targets, we leveraged a large amount of phenotype data already available for *Drosophila* mutants. Where possible, *Drosophila* orthologues were identified for genes containing UCEs (in Vectorbase). We used an 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

Table 1 Number of ultra-conserved sequences of 18 bp or more, and total number of invariant sites within these sequences

	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
No. Invariant bases 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

Numbers are displayed per chromosome arm, relative to AgamP4 reference genome. Gambiae complex, 7 species and strains (An. gambiae PEST, An. gambiae s.s. An. coluzzii, An. merus, An. arabiensis, An. quadriannulatus, An. melas); Anopheles, 21 species and strains; Culex, Culex, Quinquefasciatus reference genome; Aedes, Aedes aegypti reference genome.

phenotype data provided by the research community; we searched for phenotypes conferring female sterility or recessive lethality.

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. 2017). The results were filtered by q-value to account for multiple tests. A cut-off of q < 0.05 was used.

Variation at UCE locations in Ag1000G data

Using the final filtered variant file from phase 2 of the Ag1000G project (The Anopheles gambiae 1000 Genomes Consortium (2017): Ag1000G phase 2 AR1 data release) we extracted single nucleotide polymorphisms for the UCEs identified above, and for matched non-UCE regions. Diversity statistics were calculated in scikitallel v1.3.2 (Miles *et al.* 2020): number of segregating sites (s), nucleotide diversity (pi), and the neutrality test Tajima's D (Tajima 1989).

Data availability

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 S2 and S3, deposited along with Supplementary Table S1 and Supplementary figures are available at figshare. Custom scripts used in the data analysis can be found at https://github.com/soloughlin-hub?tab=repositories.

Supplementary material is available at https://doi.org/10. 25387/g3.14179985.

Results

Ultra-conserved regions from the multi-species alignment

Much of the MAF file does not include alignments of all 21 species and strains (Table S8 in Neafsey *et al.* 2015). The total number of aligned bases from which we extracted the UCEs was 17,095,206 (7.4%) of the AgamP4 reference genome (Supplementary Table S1). A total of 8338 invariant regions of 18 bp or more were identified; 1675 on chromosome arm 2L, 3015 on chromosome arm 2 R, 1375 on chromosome arm 3 L, 2188 on chromosome arm 3 R, and 85 on chromosome X (Table 1; we have also included the same metrics at different evolutionary timescales for comparison). The longest UCE was 164 bp. Genomic coordinates of the UCEs relative to the Anopheles gambiae PEST reference genome are given in Supplementary Table S2. The UCEs were distributed throughout the chromosomes, but were under-represented on the X chromosome (0.24% of MAF compared with 1.38% in autosomes; Supplementary Figure S1 and Table S1). The X chromosome is already under-represented in the MAF as it was less alignable than other chromosomes (Figure 2 in Neafsey et al. 2015). It is well established that the X chromosome shows higher differentiation between species than autosomes (due to "Haldanes Rule" and the "Large X effect") and genomic studies have reinforced this 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 chromosome was observed to have undergone particularly dynamic evolution, with chromosome rearrangements at a rate of 2.7 times higher than the autosomes, and a significant degree of observed gene movement from X to other chromosomes relative to Drosophila (Neafsey et al. 2015). This dynamic evolution of the chromosome may explain why it would be less likely to contain functional sequences that require conservation at the nucleotide level

Size distributions of the UCEs are shown in Supplementary Figure S2. In the autosomal genic UCEs, there is a pattern of a jump in frequency every three bases, indicating the tendency for runs of ultra-conserved bases to neither start nor end on third co-don positions in coding regions. As has been seen in some previous studies (*e.g.* Walter *et al.* 2005; Chiang *et al.* 2008), UCEs are significantly more AT-rich than random control sequences (64% and 54%, respectively, t-test P < 0.001).

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 a representative of the reference genome with respect to these features, so we extracted randomly distributed sets of "control" sequences from the MAF, and only from sequences where all 21 genomes were aligned. These control sequences were matched to give the



Figure 1 Distribution of UCE and non-UCE control sequences according to genomic location. Genomic locations annotated with BEDtools. Black bars: UCEs; Clear bars: Control sequences. Control error bars: standard deviation for 10 control data sets of sequences of matched length and number to the UCEs, extracted randomly from the MAF, only from regions where sequence for all 21 genomes is present.

same number of sequences with the same base-lengths as the UCEs, and were compared with the UCE locations to see whether the UCEs were randomly distributed. The UCE sequences were significantly over-represented (compared with control sequences) in intergenic regions (42% vs 15%, t-test, P < 0.05) and in RNA genes (1% vs 0.4%, t-test, P < 0.05), and less frequent in exons (22% vs 57%, t-test, P < 0.05). The MAF itself is heavily skewed toward exonic sequences, as only about 7% of the An. gambiae genome as a whole is exonic (Holt et al. 2002).

Orthology between important vector species

The algorithm that was used to create the sequence alignments in the MAF file results in short blocks of sequences, and is agnostic to genomic location, so to ensure that the location of our UCEs is not random, we checked for orthology between some species in the UCEs. For UCEs that fell within genes, this was done simply by comparing orthology identifiers (from OrthDB.org) between *An. gambiae* and *An. arabiensis*, and between *An. gambiae* and *An. funestus*. For *An. gambiae* and *An. arabiensis*, 94% of autosomal genes containing UCEs shared orthology. For *An. gambiae* and *An.* funestus, this number was 87%. The proportion of UCE-containing genes with orthology between species was lower on the X chromosome (54% for An. gambiae/An. arabiensis and 63% for An. gambiae/An. funestus). For UCEs that were intergenic, we looked at the orthology of the flanking genes. The results fell into six categories: orthology of both flanking genes, orthology of one flanking gene with no orthology on the other flank, orthology of one flanking gene with missing data on the other flank, no orthology on one flank with missing data on the other flank, missing data on both flanks, and no orthology of either flanking gene. Ignoring missing data, 92% of intergenic UCEs showed full or half orthology between An. gambiae and An. arabiensis, and 77% of UCEs showed full or half orthology between An. gambiae and An. gambiae and An. funestus (Figure 2). Matching orthology implies that the location of the UCEs is the same in each species with regards to shared synteny blocks.

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

Of the 13,796 genes annotated in the Anopheles gambiae PEST gene set Agam4.12, 1601 (12.9%) had at least one UCE. We performed



Figure 2 Number of intergenic UCEs that show synteny between (A) An. gambiae and An. arabiensis and (B) An. gambiae and An. funestus. The results are shown in six categories: matching orthology of both flanking genes, matching orthology of one flanking gene with no orthology on the other flank, matching orthology of one flanking gene with missing data on the other flank, no orthology on one flank with missing data on the other flank, no orthology of either flanking gene, and missing data on both flanks.

functional classification of the genes based on GO-Slim terms for molecular function, biological process, and cellular component (Supplementary Figure S3).

Because the functional classification tool 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 whole *Anopheles* PEST reference gene set (Figure 3).

In the genes containing UCEs over 50 bp long, only four categories were over-represented: transmembrane transporter activity (MF), transmembrane transport (BP), transport (BP), and proteincontaining complex (CC) (adjusted P values 0.0047, 0.0047, 0.0272, and 0.0272, respectively). Genes flanking intergenic UCEs were enriched for the GO-Slim categories DNA binding (MF), DNAbinding transcription factor activity (MF), and anatomical structure development (BP) (adjusted P values 4.16E-06, 1.46E-05, and 0.016, respectively).

Potential targets for vector control

AGAP001189 (odorant-binding protein 10) contained the highest number of invariant bases in UCEs (1215 of 135,306). Nine genes contained UCEs longer than 100 bp, of which three are annotated



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 < 0.05. MF, molecular function; BP, biological process; CC, cellular component.

as being involved in ion transport. These include the voltagegated sodium channel gene (VGSC, AGAP004707), which is a target for (and therefore has a significant role in conferring resistance to) some of the main classes of insecticides used for malaria vector control. VGSC is one of the most conserved genes we found, containing 13 UCEs with a total of 507 invariant bases, of which 91% were in exons and most coded for trans-membrane domains. A total of 357 genes contained 100 or more invariant bases. A full list of genes containing UCEs is given in Supplementary Table S3.

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

Drosophila orthologues were identified for 1309 of the 1601 genes containing UCEs. Allele and phenotype classes for these genes were extracted from Flybase where available. For an effective population suppression gene-drive, the target would affect female fertility or impose a genetic load as a homozygote, so we extracted UCE containing genes that have Drosophila orthologues annotated with a female sterile term or a lethal recessive term (shown in Supplementary Table S3). In total, 177 genes containing UCEs have Drosophila orthologues with an allele phenotype affecting female fertility, and 367 genes have Drosophila orthologues with an allele conferring a lethal recessive phenotype.

Transcription factor binding motifs in UCEs

DNA binding motifs recognized by transcription factors might be expected to be constrained and hence enriched for UCEs since this protein: DNA interaction is sequence-specific. The FIMO search found that 38% of UCEs contained hits for insect transcription factor binding sites with a q<0.05 (48% of intergenic and 30% of genic UCEs). For intergenic UCEs, this was significantly higher than control (non-conserved sequences) (48% in UCEs compared with 24% for control sequences of the same number and length, t-test across chromosome arms, P<0.005). Within genes, the difference between UCEs and controls was not significant (30% vs 23%, t-test across chromosome arms ns). This trend did not hold true for the X chromosome, where data are sparse (only 8 intergenic and 75 genic UCEs). Figure 4 shows the percentage of UCEs and control sequences containing transcription factor binding motifs broken down by chromosome arm.

Genetic variation at UCE locations in Ag1000G data

To see whether sequences are ultra-conserved across the *Anopheles* genus show variation in wild mosquito populations, we searched for single nucleotide polymorphisms (SNPs) in the 1142 samples from phase 2 of the Ag1000G project. Significance



Figure 4 Percentage of UCEs and control sequences that contain at least one insect transcription factor binding motif. Control error bars: standard deviation for 10 control data sets. UCEs were searched for known insect transcription factor binding sites from the JASPER CORE collection (Insect position frequency matrices 8th release (2020), Khan *et al.* 2017). The results were filtered by *q*-value to account for multiple tests. A cut-off of *q* < 0.05 was used.

was compared between UCEs and control sequences using a t-test across all chromosomes. There were significantly fewer sites containing polymorphisms in UCEs than control sequences (P < 0.0001, Figure 5, middle), and those SNPs that were present were at a significantly lower frequency (P < 0.0001, Figure 5, top). Of the 8338 UCEs, 1213 (15%) contained no SNPs in the 1142 samples (229 on 2L, 470 on 2R, 226 on 3L, 259 on 3R, and 29 on X). Tajima's D is significantly different and more negative for UCEs than controls, with the exception of X chromosome intergenic sequences (P < 0.005, Figure 5, bottom). Negative values of Tajima's D are expected for sequences under purifying selection.

The Ag1000G study (Anopheles gambiae 1000 genomes consortium et al. 2017) performed a search within the Phase 1 data to look for potential Cas9 targets (non-overlapping exonic invariant sequences of 21 bp, ending in the "NGG" motif) within An. gambiae and An. coluzzii. They identified 13 genes containing sequences matching these criteria. However, none of these sequences



Figure 5 Genetic diversity per chromosome arm in 1,142 Anopheles gambiae s.l. samples in UCE locations. Top: nucleotide diversity (π); middle: segregating sites (s); bottom: Tajima's D. Calculations were made in scikit-allel v1.3.2 (Miles *et al.* 2020). Results are shown per chromosome arm, divided into genic (within an annotated AGAP-) and intergenic regions. Control sequences were extracted randomly from the AgamP4 reference genome and matched to UCE sequences for length, number, and genic or intergenic location. Control error bars: standard deviation for 10 control data sets.

corresponded to UCEs fitting our more stringent definition of being conserved across the wider *Anopheles* genus. We did not confine our search for UCEs to current Cas9 target site restrictions because of the growing possibility of relaxation of these constraints as the ability to re-engineer Cas9 tolerance progresses (Walton *et al.* 2020). However, for completeness, we looked within our final set of UCEs for the Cas9 motif (18 bp followed by -NGG, or CCN- followed by 18 bp). We found 1997 (24%) UCEs contained suitable targets for Cas9.

Discussion

Similarities and differences of Anopheles UCEs with UCEs from Drosophila

Despite approximately 100 million years since their most recent common ancestor, we identified in the Anopheles genus over 8000 sequences of 18 bp or more where there was no nucleotide variation across the alignment of 21 species and strains. By coincidence, this is approximately the same span of evolutionary time covered in the human/mouse/rat data set in which UCEs were originally identified (Bejerano et al. 2004). Approximately 481 UCEs of more than 200 bp were observed between these genomes, but the longest we found in the Anopheles genus was 164 bp. This is consistent with previous reports that UCEs are fewer and shorter in insects (mainly Drosophila) than vertebrates (Glazov et al. 2005; Makunin et al. 2013). Our criteria for identifying UCEs were somewhat different than those used previously. First, we only considered sequences that were present in all 21 species/ strains in the alignment; some of these species have poorly assembled genomes, so this may have reduced the number of UCEs that we uncovered. Second, we also included invariant stretches of 18 bp or more, whereas Drosophila studies have used cut-offs of 50 bp (Glazov et al. 2005, Warnefors et al. 2016), 80 bp (Kern et al. 2015) or 100 bp (Makunin et al. 2013). Despite this, we see some similarities between our UCEs and UCEs found in Drosophila. UCEs are located in all parts of the genome and, like Drosophila, the majority are found in intergenic regions and introns. We also found that junction locations (e.g. intron-exon, exon-intergenic, etc) are over-represented compared with random sequences, which in Drosophila has been linked to the conservation of splicesites (Glazov et al. 2005; Warnefors et al. 2016). Another similarity with Drosophila is the high proportion of genes with the GO terms "binding" and "transporter activity" (Glazov et al. 2005; Kern et al. 2015). In Drosophila, ion channel/transporter genes have been shown to undergo extensive RNA editing (Hanrahan et al. 2000; Hoopengardner et al. 2003; Rodriguez et al. 2012) which is thought to explain the high level of conservation. This is because RNA adenosine deaminases require double-stranded RNA as a substrate, which means that there is likely to be strong selection at the nucleotide level. The high number of UCEs in Anopheles ion channel/transporter genes suggests that a similar mechanism is responsible for the high conservation in the Anopheles genus. However, these genes are extremely long and are not over-represented in the UCE data when a length-bias corrected analysis is carried out in GOseq. In the GOseq analysis, the most over-represented molecular functions are mostly involved in binding or structure. Transcription factor binding, enzyme binding, and nucleic acid binding have also been shown to be associated with ultra-conservation in both invertebrates and mammals (Bejerano et al. 2004; Glazov et al. 2005). A noteworthy addition to highly represented GO terms in Anopheles that has not been reported in Drosophila, is the category of "catalytic activity" genes, although again, these were not over-represented when gene length was taken into account. When the GO term functional classification was carried out on genes containing UCEs of 50 bp or more in length, we found that the category reduced from 28% to 18% suggesting that these shorter ultra-conserved regions most likely code for a small number of key residues around an active site.

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

Polymorphisms in UCEs in Anopheles populations

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

UCEs and vector control

UCEs occur within many genes that could have the potential for vector control. Nearly 200 genes have *Drosophila* orthologues with an allele phenotype affecting female fertility, and over three hundred genes have *Drosophila* orthologues with an allele conferring a lethal recessive phenotype. These phenotypes could both be used for a population suppression strategy, *i.e.*, to reduce the numbers of mosquitoes to a level where malaria could no longer be transmitted (Deredec *et al.* 2011). More investigation would be needed to see whether disrupting the genes at the ultra-conserved loci gives the same phenotype in *Anopheles*. There are also genes that confer recessive phenotypes in *Drosophila* such as "flightless" or "behaviour defective" that could also be used for population suppression, or for a population modification type of strategy, where instead of reducing the mosquito population it is

replaced by a strain that cannot transmit malaria (Carballar-Lejarazú and James, 2017). Precise targeting of sequences using CRISPR/Cas9 gene editing had made testing for these phenotypes feasible.

Another potential source of targets for genetic control approaches that has not yet been explored would be to target sequences involved in gene regulation. Many ultra-conserved sequences in mammals and invertebrates are thought to be involved in the regulation of genes important in development (Bejerano et al. 2004; Boffelli et al. 2004; Sandelin et al. 2004; Glazov et al. 2005).

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

There may be some circumstances, for example, for phased testing of a gene drive's efficacy and safety, where it is desirable to target a sequence that is unique to a particular population. For this, it would be interesting to explore conserved sites that show polymorphisms within species, a prospect that is being explored for mosquito and rodent control (Oh *et al.* 2021; Willis and Burt 2021).

Conclusion

Thousands of short genomic regions exist that are conserved across the *Anopheles* genus. These sequences show many of the same traits as UCEs found in *Drosophila* (such as an association with gene regulation and ion channel activity). Our list of UCEs 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. Focusing on sequences that have remained highly conserved over a long evolutionary time has promise for mitigating against or slowing the development of resistant alleles in the wild population.

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Literature cited

- Afgan E, Baker D, Batut B, van den Beek M, Bouvier D, *et al.* 2018. The Galaxy platform for accessible, reproducible and collaborative biomedical analyses: 2018 update. Nucleic Acids Res. 46:537–544. 10.1093/nar/gky379
- Ahituv N, Zhu Y, Visel A, Holt A, Afzal V, et al. 2007. Deletion of ultraconserved elements yields viable mice. PLoS Biol. 5:e234.
- Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. 1990. Basic local alignment search tool. J Mol Biol. 215:403–410.
- Anders C, Bargsten K, Jinek M. 2016. Structural plasticity of PAM recognition by engineered variants of the RNA-guided endonuclease Cas9. Mol Cell. 61:895–902.
- Anopheles gambiae 1000 Genomes Consortium, Data analysis group, Partner working group, Sample collections—Angola, Burkina Faso, et al. 2017. Genetic diversity of the African malaria vector Anopheles gambiae. Nature. 552:96–100.
- Bailey TL, Boden M, Buske FA, Frith M, Grant CE, et al. 2009. MEME SUITE: Tools for motif discovery and searching. Nucleic Acids Res. 37:202–208.
- Baker DA, Nolan T, Fischer B, Pinder A, Crisanti A, et al. 2011. A comprehensive gene expression atlas of sex- and tissue-specificity in the malaria vector, *Anopheles gambiae*. BMC Genomics. 12:296.
- Baxter L, Jironkin A, Hickman R, Moore J, Barrington C, et al. 2012. Conserved noncoding sequences highlight shared components of regulatory networks in dicotyledonous plants. Plant Cell. 24: 3949–3965.
- Bejerano G, Pheasant M, Makunin I, Stephen S, Kent WJ, et al. 2004. Ultraconserved elements in the human genome. Science. 304: 1321–1325.
- Bhatt S, Weiss DJ, Cameron E, Bisanzio D, Mappin B, et al. 2015. The effect of malaria control on Plasmodium falciparum in Africa between 2000 and 2015. Nature. 526:207–211.
- Boffelli D, Nobrega MA, Rubin EM. 2004. Comparative genomics at the vertebrate extremes. Nat Rev Genet. 5:456–465.
- Burt A, Deredec A. 2018. Self-limiting population genetic control with sex-linked genome editors. Proc R Soc B. 285:20180776 10.1098/rspb.2018.0776
- Burt A. 2003. Site-specific selfish genes as tools for the control and genetic engineering of natural populations. Proc Biol Sci. 270: 921–928.
- Calin GA, Liu C, Ferracin M, Hyslop T, Spizzo R, et al. 2007. Ultraconserved regions encoding ncRNAs are altered in human leukemias and carcinomas. Cancer Cell. 12:215–229.
- Carballar-Lejarazú R, James AA. 2017. Population modification of Anopheline species to control malaria transmission. Pathog Glob Health. 111:424–435.
- Chatterjee P, Jakimo N, Jacobson JM. 2018. Minimal PAM specificity of a highly similar SpCas9 ortholog. Sci Adv. 4:eaau0766
- Chiang CW, Derti A, Schwartz D, Chou MF, Hirschhorn JN, et al. 2008. Ultraconserved elements: Analyses of dosage sensitivity, motifs and boundaries. Genetics. 180:2277–2293.

- Danecek P, Auton A, Abecasis G, Albers CA, Banks E, 1000 Genomes Project Analysis Group, *et al.* 2011. The variant call format and VCFtools. Bioinformatics. 27:2156–2158.
- Deredec A, Burt A, Godfray HC. 2008. The population genetics of using homing endonuclease genes in vector and pest management. Genetics. 179:2013–2026. https://doi.org/10.1534/genetics.108. 089037
- Deredec A, Godfray HC, Burt A. 2011. Requirements for effective malaria control with homing endonuclease genes. Proc Natl Acad Sci U S A. 108:874–880.
- Faircloth BC, McCormack JE, Crawford NG, Harvey MG, Brumfield RT, et al. 2012. Ultraconserved elements anchor thousands of genetic markers spanning multiple evolutionary timescales. Syst Biol. 61:717–726.
- Fontaine MC, Pease JB, Steele A, Waterhouse RM, Neafsey DE, *et al.* 2015. Mosquito genomics. Extensive introgression in a malaria vector species complex revealed by phylogenomics. Science. 347: 1258524. https://doi.org/10.1126/science.1258524
- Gantz VM, Jasinskiene N, Tatarenkova O, Fazekas A, Macias VM, *et al.* 2015. Highly efficient Cas9-mediated gene drive for population modification of the malaria vector mosquito *Anopheles stephensi*. Proc Natl Acad Sci U S A. 112:6736–6743.
- Giraldo-Calderón GI, Emrich SJ, MacCallum RM, Maslen G, Dialynas GE, et al. 2014. VectorBase: An updated bioinformatics resource for invertebrate vectors and other organisms related with human diseases. Nucleic Acids Res. 43(Database issue):707–713.
- Glazov EA, Pheasant M, McGraw EA, Bejerano G, Mattick JS. 2005. Ultraconserved elements in insect genomes: A highly conserved intronic sequence implicated in the control of homothorax mRNA splicing. Genome Res. 15:800–808.
- Gramates LS, Marygold SJ, Santos GD, Urbano J-M, Antonazzo G, the FlyBase Consortium, *et al.* 2017. FlyBase at 25: Looking to the future. Nucleic Acids Res. 45:D663–D671.
- Grant CE, Bailey TL, Noble WS. 2011. FIMO: Scanning for occurrences of a given motif. Bioinformatics. 27:1017–1018.
- Hammond A, Galizi R, Kyrou K, Simoni A, Siniscalchi C, C, et al. 2016. A CRISPR-Cas9 gene drive system targeting female reproduction in the malaria mosquito vector *Anopheles gambiae*. Nat Biotechnol. 34:78–83.
- Hammond A, Kyrou K, Bruttini M, North A, Galizi R, *et al.* 2017. The creation and selection of mutations resistant to a gene drive over multiple generations in the malaria mosquito. PLoS Genet. 13: e1007039
- Hanrahan CJ, Palladino MJ, Ganetzky B, Reenan RA. 2000. RNA editing of the *Drosophila* para Na(+) channel transcript. Evolutionary conservation and developmental regulation. Genetics. 155: 1149–1160.
- Holt R A, Subramanian G M, Halpern A, Sutton G G, Charlab R, *et al.* 2002. The genome sequence of the malaria mosquito Anopheles gambiae. Science. 298:129–149.
- Hoopengardner B, Bhalla T, Staber C, Reenan R. 2003. Nervous system targets of RNA editing identified by comparative genomics. Science. 301:832–836.
- Hu JH, Miller SM, Geurts MH, Tang W, Chen L, et al. 2018. Evolved Cas9 variants with broad PAM compatibility and high DNA specificity. Nature. 556:57–63.
- Jinek M, Chylinski K, Fonfara I, Hauer M, Doudna JA, et al. 2012. A programmable dual-RNA-guided DNA endonuclease in adaptive bacterial immunity. Science. 337:816–821.
- Kern AD, Barbash DA, Mell JC, Hupalo D, Jensen A. 2015. Highly constrained intergenic *Drosophila* ultraconserved elements are candidate ncRNAs. Genome Biol Evol. 7:689–698.

- Khan A, Fornes O, Stigliani A, Gheorghe M, Castro-Mondragon JA, et al. 2017. JASPAR 2018: Update of the open-access database of transcription factor binding profiles and its web framework. Nucleic Acids Res. 46:260–D266.
- Kranjc N, Crisanti A, Nolan T, Bernardini F. 2021. Anopheles gambiae genome conservation as a resource for rational gene drive target site selection. Insects. 12:97.https://doi.org/10.3390/insects 12020097
- Kriventseva EV, Kuznetsov D, Tegenfeldt F, Manni M, Dias R, *et al.* 2019. OrthoDB v10: Sampling the diversity of animal, plant, fungal, protist, bacterial and viral genomes for evolutionary and functional annotations of orthologs. Nucleic Acids Res. 47: D807–D811.
- Kyrou K, Hammond A, Galizi R, Kranjc N, Burt A, et al. 2018. A CRISPR–Cas9 gene drive targeting doublesex causes complete population suppression in caged *Anopheles gambiae* mosquitoes. Nat Biotechnol. 36:1062–1066. (
- Li H, Durbin R. 2010. Fast and accurate long-read alignment with Burrows-Wheeler Transform. Bioinformatics. 26:589–595. [PMID: 20080505]
- Lin M, Eng C, Hawk ET, Huang MS, Lin J, et al. 2012. Identification of polymorphisms in ultraconserved elements associated with clinical outcomes in locally advanced colorectal adenocarcinoma. Cancer. 118:6188–6198.
- Makunin IV, Shloma VV, Stephen SJ, Pheasant M, Belyakin SN. 2013. Comparison of ultra-conserved elements in drosophilids and vertebrates. PLoS One. e82362.8:
- Mi H, Huang X, Muruganujan A, Tang H, Mills C, et al. 2017. PANTHER version 11: Expanded annotation data from Gene Ontology and Reactome pathways, and data analysis tool enhancements. Nucleic Acids Res. 45:D183–D189.
- Miles A, pyup.io bot R, Murillo P, Ralph N, Harding R, et al. 2020. cggh/scikit-allel: v1.3.2 (Version v1.3.2). Zenodo. http://doi.org/ 10.5281/zenodo.3976233
- Neafsey DE, Waterhouse RM, Abai MR, Aganezov MA, Alekseyev MA, et al. 2015. Highly evolvable malaria vectors: The genomes of 16 Anopheles mosquitoes. Science. 347:1258522.
- Oh KP, Shiels AB, Shiels L, Blondel DV, Campbell KJ, et al. 2021. Population genomics of invasive rodents on islands: Genetic consequences of colonization and prospects for localized synthetic gene drive. Evol Appl. https://doi.org/10.1111/eva.13210
- Presgraves DC. 2018. Evaluating genomic signatures of "the large X-effect" during complex speciation. Mol Ecol. 27:3822–3830. https://doi.org/10.1111/mec.14777
- Quattrini AM, Faircloth BC, Dueñas LF, Bridge TCL, Brugler MR, et al. 2018. Universal target-enrichment baits for anthozoan (Cnidaria) phylogenomics: New approaches to long-standing problems. Mol Ecol Resour. 18:281–295. https://doi.org/10.1111/ 1755-0998.12736
- Quinlan AR, Hall IM. 2010. BEDTools: A flexible suite of utilities for comparing genomic features. Bioinformatics. 26:841–842.
- Rodriguez J, Menet JS, Rosbash M. 2012. Nascent-seq indicates widespread cotranscriptional RNA editing in *Drosophila*. Mol Cell. 47: 27–37.
- Sandelin A, Bailey P, Bruce S, Engstrom PG, Klos JM, et al. 2004. Arrays of ultraconserved non-coding regions span the loci of key developmental genes in vertebrate genomes. BMC Genomics. 5:99.
- Siepel A, Bejerano G, Pedersen JS, Hinrichs AS, Hou M, et al. 2005. Evolutionarily conserved elements in vertebrate, insect, worm, and yeast genomes. Genome Res. 15:1034–1050.
- Tajima F. 1989. Statistical method for testing the neutral mutation hypothesis by DNA polymorphism. Genetics. 123:585–595.

- The Anopheles gambiae 1000 Genomes Consortium. 2017. Ag1000G phase 2 AR1 data release. MalariaGEN. http://www.malariagen. net/data/ag1000g-phase2-ar1.
- Unckless RL, Clark AG, Messer PW. 2017. Evolution of resistance against CRISPR/Cas9 gene drive. Genetics. 205:827–841.
- Vilella AJ, Blanco-Garcia A, Hutter S, Rozas J. 2005. VariScan: Analysis of evolutionary patterns from large-scale DNA sequence polymorphism data. Bioinformatics. 21:2791–2793.
- Visel A, Prabhakar S, Akiyama J A, Shoukry M, Lewis K D, et al. 2008. Ultraconservation identifies a small subset of extremely constrained developmental enhancers. Nat Genet. 40:158–160. 10.1038/ng.2007.55 18176564
- Walter K, Abnizova I, Elgar G, Gilks WR. 2005. Striking nucleotide frequency pattern at the borders of highly conserved vertebrate non-coding sequences. Trends Genet. 21:436–440.
- Walton RT, Christie KA, Whittaker MN, Kleinstiver BP. 2020. Unconstrained genome targeting with near-PAMless engineered CRISPR-Cas9 variants. Science. 368:eaba8853–296. https//doi.org/ 10.1126/science.aba8853.

- Warnefors M, Hartmann B, Thomsen S, Alonso CR. 2016. Combinatorial gene regulatory functions underlie ultraconserved elements in Drosophila. Mol Biol Evol. 33:2294–2306.
- Weetman D, Steen K, Rippon EJ, Mawejje HD, Donnelly MJ, et al. 2014. Contemporary gene flow between wild An. gambiae s.s. and An. arabiensis. Parasit Vectors. 7:345.
- WHO. 2018. World malaria report 2018. Geneva: World Health Organization. https://www.who.int/malaria/publications/worldmalaria-report-2018/en/
- Willis K, Burt A. 2021. Double drives and private alleles for localised population genetic control. BioRxiv Preprint. https://doi.org/10. 1101/2021.01.08.425856
- Windbichler N, Menichelli M, Papathanos PA, Thyme SB, Li H, et al. 2011. A synthetic homing endonuclease-based gene drive system in the human malaria mosquito. Nature. 473:212–215.
- Young MDMJ, Wakefield, GK, Oshlack Smyth A 2010. Gene ontology analysis for RNA-seq: Accounting for selection bias. Genome Biol. 11:R14.

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