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DATA NOTE

A chromosome-scale assembly of the major African malaria vector Anopheles funestus

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Abstract

Background: Anopheles funestus is one of the 3 most consequential and widespread vectors of human malaria in tropical Africa. However, the lack of a high-quality reference genome has hindered the association of phenotypic traits with their genetic basis in this important mosquito. **Findings:** Here we present a new high-quality A. *funestus* reference genome (AfunF3) assembled using 240× coverage of long-read single-molecule sequencing for contigging, combined with 100× coverage of short-read Hi-C data for chromosome scaffolding. The assembled contigs total 446 Mbp of sequence and contain substantial duplication due to alternative alleles present in the sequenced pool of mosquitos from the FUMOZ colony. Using alignment and depth-of-coverage information, these contigs were deduplicated to a 211 Mbp primary assembly, which is closer to the expected haploid genome size of 250 Mbp. This primary assembly consists of 1,053 contigs organized into 3 chromosome-scale scaffolds with an N50 contig size of 632 kbp and an N50 scaffold size of 93.811 Mbp, representing a 100-fold improvement in continuity versus the current reference assembly, AfunF1. **Conclusion:** This highly contiguous and complete A. *funestus* reference genome assembly will serve as an improved basis for future studies of genomic variation and organization in this important disease vector.

Keywords: Anopheles mosquito; malaria; genome assembly; DNA sequencing; Hi-C chromosome conformation capture

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Data Description

Introduction and background

Many insect genomes remain a challenge to assemble, and mosquito genomes have proven particularly difficult owing to their repeat content and structurally dynamic genomes. These issues are compounded by the fact that long-read sequencing technologies typically require >10 μ g of DNA for library construction. As a result, it is often impossible to construct a sequencing library from a single individual. Instead, it has been necessary to sequence a pool of individuals from an inbred population [1]. For species that are amenable to extensive inbreeding, this approach has led to reference-grade genomes directly from the assembler [2]. However, when inbreeding is not possible, the sequenced pool of individuals can carry population variation that fragments the resulting assembly. In this case, instead of assembling a single genome, the assembler must reconstruct some unknown number of variant haplotypes.

Motivated by the goal of genome-enabled malaria control, a large international consortium previously sequenced and assembled the genomes of 16 Anopheles species using short-read Illumina sequencing [3, 4]. Although these draft assemblies represented a crucial first step, their potential for (i) understanding and manipulating vectorial capacity traits, (ii) inferring how key vector adaptations to hosts and habitats have arisen and are maintained, and (iii) accurately defining vector breeding units and migration between them is constrained by 2 major limitations. First, many of these Anopheles assemblies are highly fragmented collections of relatively short scaffolds, causing gene annotation problems such as missing genes, missing exons, and genes split between scaffolds or sequencing gaps. Thus, one of the consequences of fragmented assemblies is that it is difficult to estimate gene copy number, which may be linked to important phenotypic traits (e.g., insecticide resistance) [5, 6]. Genes of particular interest with respect to arthropod disease vectors (e.g., cytochrome P450s and odorant/gustatory receptors) may be especially prone to annotation errors because many belong to gene families whose members are often physically clustered into tandem arrays.

A second major limitation of fragmented insect assemblies is that they are rarely scaffolded into chromosomes, owing to difficulty and lack of funding for physical or linkage mapping. Among other consequences, the unknown placement of scaffolds along chromosome arms means that their position within or outside of chromosomal inversions is difficult or impossible to determine. Many anopheline species are highly polymorphic for chromosomal inversions, which tend to occur disproportionately on particular chromosome arms [7-9]. In a heterozygote carrying 1 inverted and 1 uninverted chromosome, recombination between the reversed chromosomal segments is greatly reduced [10], creating cryptic population structure that can cause spurious associations in genome-wide association studies (GWAS) [11] and mislead recombination-based inference of selection and gene flow [12, 13]. Importantly, chromosomal inversions also directly or indirectly influence traits affecting malaria transmission intensity-anopheline biting and resting behavior [14, 15], seasonality [16], aridity tolerance [14, 17-21], ecological plasticity [22, 23], morphometric variation [24], and Plasmodium infection rates [25, 26]. Thus, correct population genomic and GWAS inferences depend upon knowing the location of a marker in the genome.

Anopheles funestus (NCBI:txid62324) is one of the 3 most important and widespread vectors of human malaria in tropical Africa [27–30], and unlike Anopheles gambiae with which it

broadly co-occurs, it is a relatively neglected species. It is considered even more highly anthropophilic and endophilic than A. gambiae and amenable to conventional indoor-based vector control such as bed nets and indoor spraying of houses with residual insecticides. Indeed, historical house spraying campaigns in eastern and southern Africa not only locally eliminated this species, but the effect was maintained for several years following the cessation of spraying, due to the apparent inability of A. funestus to recolonize some areas. Likewise, A. funestus was eliminated from a humid forest and degraded forest areas in West Africa where malaria is meso- or hypoendemic [31]. However, in the savanna environment of West Africa where malaria is holoor hyperendemic, similar historical indoor spraying campaigns failed to eliminate the species. Exophilic populations persisted, which-despite marked anthropophily-continued to feed outdoors on cattle but also entered sprayed houses to bite humans. Today, the situation is worsened by the emergence and spread of insecticide resistance in this species [29, 32-34].

Mastery over malaria will require tackling A. funestus, but it remains understudied; information on its behavior and genetics lags far behind that of A. gambiae. At least part of the reason for its neglect may be the historical lack of laboratory colonies, a problem solved with the establishment of the FU-MOZ colony and its registration with the Anopheles program of BEI Resources [35]. A. funestus shares with A. gambiae not only a broad sub-Saharan distribution and major vector status but also abundant chromosomal inversion polymorphism and shallow range-wide population structure [36]. However, there are behavioral and genetic heterogeneities relevant to malaria transmission that remain poorly understood. In West Africa, strong cytogenetic evidence points to cryptic, temporally stable assortatively mating populations co-occurring in the same villages [37-40]. These chromosomally recognized forms of A. funestus, named Kiribina and Folonzo, seem to differ in larval ecology, and-importantly-they also differ in adult behaviors affecting vectorial capacity, most notably indoor resting behavior. Mechanistic understanding of the genomic determinants of these and other epidemiologically important phenotypic and behavioral traits ultimately depends on upgrading the A. funestus reference to a chromosome-based assembly in which the unanchored scaffolds are united, ordered, and oriented on chromosome arms.

Chromosome-scale assembly of Anopheles funestus

To achieve a complete and highly contiguous assembly of the A. funestus genome (AfunF3), we first assembled contigs from long, single-molecule reads and then scaffolded these contigs into chromosome-scale scaffolds using Hi-C proximity ligation data. A similar strategy was recently used to improve the genome of Aedes aegypti [41]. An initial assembly of the long-read data alone (AfunF3 contigs) yielded a contig N50 size of 94.05 kbp (N50 such that 50% of assembled bases are in contigs of this size or greater) and extensive haplotype separation as evidenced by an inflated assembly size of 446.04 Mbp and a high rate of core gene duplications (48%) as measured by BUSCO [42]. These alternative alleles likely derive from natural variation circulating within the sequenced FUMOZ colony, as the DNA from a pool of adult mosquitoes was required for Pacific Biosciences (PacBio) library preparation. Identifying and removing duplicate contigs via an all-vs-all alignment reduced the primary assembly size to 211.75 Mbp and improved the N50 size to 631.72 kbp (Table 1).

The primary set of contigs (excluding alternative alleles) was then scaffolded using Hi-C Illumina reads to first bin the con-

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Assembly		Contigs			Scaffolds		Total assembly size	QV (accuracy)	
	No.	N50	Maximum size	No.	N50	Maximum size		Illumina	10X Genomics
AfunF1	9,880	60,925	563,645	1,392	671,960	3,832,769	225,223,604	38.93 (99.84%)	22.69 (99.46%)
AfunF3 contigs	10,245	94,259	7,564,979	9,175	238,902	99,362,816	446,039,041	29.82 (99.89%)	28.18 (99.84%)
AfunF3 primary	1,053	631,722	7,564,979	3	93,811,348	99,362,816	210,827,327	24.94 (99.64%)	25.82 (99.73%)

Table 1: Assembly statistics for the A. funestus genome

AfunF1 represents the prior reference assembly, AfunF3 contigs denotes the complete long-read assembly with all contigs included, and AfunF3 primary denotes the assembly after deduplication and scaffolding. The assembly quality value (QV) was estimated using Illumina or 10X Genomics data. QV (Illumina) is highest for the AfunF1 assembly because it is the same data used to generate that assembly, whereas QV (10X Genomics) is based on data from a single mosquito of the same FUMOZ colony. The numbers in parentheses in the QV columns denote the estimated accuracy of the assembly based on QV score.

tigs into 3 chromosomes, followed by ordering and orientation of the contigs using the Proximo method (Phase Genomics, Seattle, WA, USA). The final scaffolded assembly (AfunF3 primary) contains 210.82 Mbp of sequence and a scaffold N50 of 93.81 Mbp. The resulting scaffolds represent the entirety of the 3 A. *funestus* chromosomes: 2, 3, and X (Fig. 1).

Because single-molecule PacBio data are prone to insertion and deletion errors, all AfunF3 contigs were polished twice with Arrow [43] using the signal-level PacBio data and once with Pilon [44] using paired-end Illumina data from the same FUMOZ colony. Because Illumina-based polishing tools typically do not correct bases that appear heterozygous in the read set, we anticipated that variation in the FUMOZ colony would prevent the correction of variant bases. To help address this issue, we finally polished the assembly using 10X Genomics Illumina data obtained from an individual mosquito. As an independent test of base accuracy, we compared our new assembly (AfunF3 primary) and the prior assembly (AfunF1) to a 10X Genomics dataset from a different individual mosquito. The average Phred-scaled quality value (QV) [45] of the new assembly was estimated as QV 28 (99.84% identity) versus QV 23 (99.49% identity) for the Illuminabased AfunF1 assembly. These independent data from a single mosquito of the FUMOZ colony indicate that the new AfunF3 assembly is of comparable accuracy to the prior Illumina-based assembly and that the small differences between quality estimates could be due to genetic diversity within the colony.

We next evaluated the structural accuracy of the AfunF1 and AfunF3 assemblies by measuring their agreement with the raw PacBio reads. The intermediate assembly AfunF2 [46] was assembled before collection of all PacBio and Hi-C data and so was deemed redundant and excluded from these analyses. When compared to the raw data, the AfunF3 primary assembly had fewer called structural differences (insertions, deletions, duplications, and inversions) than AfunF1 (Table 2). Despite the substantial single-nucleotide polymorphism observed within the FUMOZ colony, no large polymorphic inversions could be identified from the combined PacBio, Hi-C, and 10X Genomics data. Comparison of the chromosome-scale AfunF3 primary assembly versus the A. gambiae reference genome (AgamP4) confirmed a known reciprocal whole-arm translocation between 2L and 3R, as well as substantial intra-chromosomal shuffling (Fig. 2). AfunF3 contigs also had fewer fragmented BUSCO core genes and a similar number of complete BUSCOs compared to AfunF1 (Table 2) but also a high rate of duplication. The AfunF3 primary scaffolds reduce duplication at the expense of lower BUSCO completeness.

To further evaluate AfunF3's suitability as an updated reference for A. *funestus*, we mapped RNA-sequencing (RNA-Seq) expression data to the assemblies and computed the number of concordant paired-end reads. A better assembly is expected to have both a higher fraction of mapped reads (completeness) as well as a higher fraction of correctly spaced and oriented pairs (structural accuracy). Both primary and complete AfunF3 assemblies have better agreement of mapped read pairs as well as a higher overall mapping rate versus the AfunF1 assembly (Table 2). The AfunF3 contigs do have a higher rate of multimapping RNA-Seq reads, but this is reduced in the primary assembly while preserving the high mapping rate. In addition to a higher mapping rate, more complete transcripts were mapped to single contigs within the long-read assemblies. The average number of complete transcripts contained per contig was 67.38 for AfunF3 primary versus 5.28 for the AfunF1 assembly. These results demonstrate the greater continuity of the updated assembly, which provides sequence-resolved reconstructions of many A. funestus intergenic regions for the first time.

Discussion

Anopheles funestus is one of the leading vectors of malaria, and understanding the organization and function of its genome is key to controlling this deadly disease. Herein we describe a chromosome-scale assembly of the A. funestus genome using multiple sequencing technologies and assembly methods. The tremendous improvement in the completeness and contiguity of its genome will provide a valuable resource for future genomic analyses and functional characterization of this important species and enable a mechanistic understanding of the genomic determinants of epidemiologically important phenotypic and behavioral traits.

Materials and Methods

Library preparation and sequencing

A gravid female mosquito of the FUMOZ colony was allowed to lay eggs, and her offspring were inbred for a single generation. From this, an isofemale line was grown and DNA extracted from the adult females for sequencing with PacBio and Hi-C. A total of 46 single-molecule real-time (SMRT) cells of PacBio RSII sequencing using the P6-C4 chemistry were run by the core facility at the Icahn School of Medicine at Mount Sinai (New York, NY), resulting in 173× coverage (assuming a 250-Mbp genome size). A previous study generated 70× coverage of the same colony using the older PacBio P5-C3 chemistry sequencing [46]. These older data were combined with the additional 173× coverage, totaling 60.95 Gb of long-read data in 10.93 million sequences (average length 5.6 kb, N50 read length 8.4 kb) and an estimated total coverage of 234×. Two Hi-C libraries were prepared and sequenced

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Figure 1: Circos plot comparing the AfunF1 assembly of A. *funestus* to the updated AfunF3 assembly. AfunF1 scaffolds (colored half of the outer ring) are ordered by majority alignment location onto AfunF3 (black half of the outer ring). Connecting lines indicate pairwise alignments between the 2 assemblies, and crossing lines indicate that part of the AfunF1 scaffold aligns to discordant regions on the AfunF3 chromosome. The first internal ring color corresponds to the AfunF1 scaffold color. The second internal ring represents the orientation of the AfunF1 scaffolds onto AfunF3, where orange is forward and green is reverse.

Table 2: Validation of A. funestus genome assemblies using BUSCO gene set completeness, agreement of the assemblies with RNA-Seq transcriptome data, and structural accuracy inferred using PacBio long-read data

	BUSCO statistics				Trar	Structural variants called with long reads					
Assembly	C/S	C/D	F	М	Alignment rate	Multi-mapped reads	Transcripts in a single contig	Deletions	Duplicati	onsversion	s Insertions
AfunF1	2,756	16	27	16	81.79	23.92	84.96	9,036	455	152	3,798
AfunF3 contigs AfunF3 primary	2,765 2,685	1,068 54	18 30	17 81	84.34 84.86	36.97 27.03	91.16 89.40	NA 571	NA 6	NA 10	NA 702

AfunF1 represents the prior reference assembly, AfunF3 contigs denotes the complete long-read assembly with all contigs included, and AfunF3 primary denotes the assembly after deduplication and scaffolding. For BUSCO categories C denotes "complete genes," S denotes "single copy genes," D denotes "duplicated genes," F denotes "fragmented genes," and M denotes "missing genes."



Figure 2: Hi-C interaction map for assembled A. *funestus* scaffolds generated using the Juicebox Hi-C visualization program [47]. Darker colors indicate a higher frequency of chromatin interaction. The plot shows clear separation of chromosome boundaries and limited off-diagonal interactions, supporting the global structure of the chromosome-scale scaffolds. Note that the light colored "cross" centered near the centromere of chromosome 3 is the repetitive rDNA locus, which could not be confidently placed using the Hi-C data alone and may require future correction using other mapping techniques (see Methods).

(one from mixed-sex larvae, the second from adult females) by Phase Genomics (Seattle, WA, USA), resulting in ${\sim}100{\times}$ coverage of Illumina Hi-C data containing ${\sim}187$ million 80-bp paired-end Illumina reads.

Assembly and scaffolding

PacBio contig assembly was performed with Canu v1.3 (Canu, RRID:SCR_015880) [48] using the following parameters: corOut-Coverage = 100 genomeSize = 250m errorRate = 0.013 batOptions = "-dg 3 -db 3 -dr 1 -ca 500 -cp 50". The resulting contigs were then polished with Arrow [43] using default parameters and the P6-C4 PacBio signal data (because Arrow does not support the older P5-C3 data). After polishing, the assembly was separated into primary and alternative contigs to remove unnecessarily duplicated alleles from the AfunF3 contigs. This was performed using 2 different approaches. First, contigs containing \geq 1 complete BUSCO gene were identified. For each BUSCO gene, if it was found contained in \geq 2 contigs, the contig with the highest alignment score was kept as the primary. Next, all contigs not containing a BUSCO gene but assembled with high coverage (>40X) were added to the primary set.

To order and orient the primary contigs along the chromosomes, Hi-C reads were aligned using Bowtie2 (Bowtie, RRID:SC R_005476) [49] and scaffolding using Proximo (Phase Genomics, Seattle, WA, USA). Scaffold gaps spanned by PacBio reads were filled using PBJelly (PBJelly, RRID:SCR_012091) [50]. This assembly was again run through Arrow to polish the sequences inserted by PBJelly and fill any remaining short gaps. The Hi-C assembled scaffolds were then aligned using NUCmer [51] to the AfunF1 contigs for validation and the alignments visualized using Circos (Circos, RRID:SCR_011798) [52] and mummerplot. This identified a mis-join of chromosomes 3R and X, which was manually corrected. Additional manual curation using mapped transcripts, fluorescence in situ hybridization (FISH) probes [46], and comparison to AfunF1 scaffolds identified a few additional inversion errors in the scaffolds, mainly on distal 2L. Visual inspection of the Hi-C data showed clear signatures of scaffolding error. These errors were corrected by manually extracting the region and placing the sequence at the correct locus, as indicated by the Hi-C interactions. After these corrections, the scaffolded chromosomes (AfunF3 primary) show good agreement with the Hi-C data (Fig. 3). The largest remaining ambiguity in the Hi-C map is the placement of the ribosomal DNA (rDNA) locus, which is placed near the centromere of chromosome 3 in the AfunF3 assembly. Given that the rDNA locus in A. gambiae is known to be on the X chromosome [53], this is possibly a mis-assembly in AfunF3 mediated by the increased proportion of repetitive transposable elements surrounding the rDNA and centromeres. However, there was insufficient long-read or Hi-C evidence to confidently place this highly repetitive locus in AfunF3, which may require correcting in future *A. funestus* assemblies.

Because diploid and population variation introduces indels in the Arrow polishing process [55], the final assemblies were also polished by Pilon using paired-end Illumina data (NCBI SRA accession numbers: SRX209628 and SRX209387) and 10X Genomics Illumina data from a single individual (NCBI SRA accession number: SRX4819916). The paired-end Illumina data were mapped using BWA-MEM [56] and the 10X Genomics data mapped using Lariat [57] in a barcode-aware manner, so as to improve the mapping quality. Consensus quality of the final assemblies was then estimated using an independent 10X Genomics dataset (NCBI SRA accession number: SRX4819903) of a different mosquito of the same FUMOZ colony. Based on the alignment of reads to the assembly, variants were called using freebayes (parameters: -C 2 - 0 - 0 - q 20 - z 0.10 - E 0 - X - u - p 2 - F 0.5),



Figure 3: Whole-genome alignment dotplot for Anopheles funestus and Anopheles gambiae genomes generated using D-GENIES [54]. A dot in the plot corresponds to a match between the corresponding genomic positions indicated on the axes. The A. gambiae reference genome is displayed on the x-axis, and the A. funestus AfunF3 primary assembly on the y-axis. A reciprocal whole-arm translocation between 2L and 3R is apparent, as well as substantial intra-chromosomal shuffling between these genomes.



Figure 4: GC content versus coverage plot for all assembled A. funestus contigs. The orange points denote the contigs classified by Kraken as A. funestus and green points denote everything else. A majority of the contigs are classified as A. funestus by Kraken, and there is no indication of extensive contamination.

and the assembly QV was estimated using called homozygous variants (i.e., positions where nearly all Illumina reads agreed with each other yet disagreed with the assembly).

Validation

To check for the presence of contamination, assembled contigs were classified using Kraken [58] using a custom database including all microbial RefSeq genomes and all available mosquito genomes. Most of the assembled sequence (96.00%) was classified as A. *funestus* or Culicidae. The remaining sequences were primarily unannotated or annotated at a higher taxonomic level (3.76%), from possible bacterial/human sources (0.24%, 32 contigs), and had slightly lower guanine-cytosine (GC) content (Fig. 4). However, none of these contigs were called contaminants by NCBI's independent contamination check and so all contigs were included in the submitted assembly to avoid excluding novel mosquito sequence missing from the prior draft assemblies.

The structural accuracy of the assemblies was evaluated by mapping raw PacBio reads and calling structural variants. PacBio reads were aligned to each assembly using NGMLR [59] with the following parameters: -t 16 -x pacbio –skip-write. Using these alignments, variants were called using Sniffles [59] with the following parameters: -t 32 -s 10 -f 0.25. Variants were then filtered to avoid capturing heterozygous population variants such that variants for which the alternate variant had \geq 45 supporting reads and the assembly variant had <10 supporting reads were called as assembly errors.

Paired-end RNA-Seq for the A. funestus FUMOZ colony was downloaded from NCBI under accession SRR826832. These reads were aligned to all assemblies using the HISAT2 aligner (HISAT2, RRID:SCR_015530) [60] and assembled into transcripts using Trinity (Trinity, RRID:SCR_013048) [61] with default parameters. The assembled transcripts were then mapped to all assemblies using GMAP (GMAP, RRID:SCR_008992) [62]. Transcripts were required to be aligned over 90% of their length to a single contig to be considered "complete" in the assembly.

Availability of supporting data and materials

Raw genomic sequence reads are available in the NCBI Sequence Read Archive under project accession PRJNA494870. This Whole Genome Shotgun project has been deposited at DDBJ/ENA/GenBank under the accession RCWQ00000000. The version described in this paper is version RCWQ01000000. Supporting data and materials are available in the *GigaScience* GigaDB database [63].

Abbreviations

bp: base pairs; BUSCO: Benchmarking Universal Single-Copy Orthologs; FISH: fluorescence in situ hybridization; GC: guaninecytosine; GWAS: genome-wide association studies; kbp: kilobase pairs; Mbp: megabase pairs; NUCmer: NUCleotide MUMmer; PacBio: Pacific Biosciences; rDNA: ribosomal DNA; RNA-Seq: RNA-sequencing; NCBI: National Center for Biotechnology Information; QV: quality value; SMRT: single-molecule real-time; SRA: Sequence Read Archive.

Competing interests

The authors declare that they have no competing interests.

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Authors' contributions

A.M.P. and N.J.B. conceived and coordinated the project. J.G., S.K., S.T.S., and A.M.P. performed the genome assembly, validation, and comparative analyses. S.R. provided the 10X Genomics data and analysis. P.H. provided FUMOZ samples for sequencing. J.G., A.M.P., and N.J.B. drafted the manuscript. All the authors have read and approved the manuscript.

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References

- 1. Kim KE, Peluso P, Babayan P, et al. Long-read, whole-genome shotgun sequence data for five model organisms. Sci Data 2014;1:140045.
- Berlin K, Koren S, Chin C-S, et al. Assembling large genomes with single-molecule sequencing and localitysensitive hashing. Nat Biotechnol 2015;33:623–30.
- Neafsey DE, Christophides GK, Collins FH, et al. The evolution of the Anopheles 16 genomes project. G3 (Bethesda) 2013;3:1191–4.
- Neafsey DE, Waterhouse RM, Abai MR, et al. Mosquito genomics. highly evolvable malaria vectors: the genomes of 16 *Anopheles* mosquitoes. Science 2015;347:1258522.
- Assogba BS, Milesi P, Djogbénou LS, et al. The ace-1 locus is amplified in all resistant Anopheles gambiae mosquitoes: fitness consequences of homogeneous and heterogeneous duplications. PLoS Biol 2016;14:e2000618.
- Weetman D, Djogbenou LS, Lucas E. Copy number variation (CNV) and insecticide resistance in mosquitoes: evolving knowledge or an evolving problem? Curr Opin Insect Sci 2018;27:82–8.
- Coluzzi M. A polytene chromosome analysis of the Anopheles gambiae species complex. Science 2002;298:1415–8.
- Pombi M, Caputo B, Simard F, et al. Chromosomal plasticity and evolutionary potential in the malaria vector Anopheles gambiae sensu stricto: insights from three decades of rare paracentric inversions. BMC Evol Biol 2008;8:309.
- Sharakhov I. A microsatellite map of the African human malaria vector Anopheles funestus. J Hered 2004;95:29–34.
- 10. Kirkpatrick M. How and why chromosome inversions evolve. PLoS Biol 2010;**8**:e1000501.
- Ma J, Amos CI. Investigation of inversion polymorphisms in the human genome using principal components analysis. PLoS One 2012;7:e40224.

- Seich Al Basatena N-K, Hoggart CJ, Coin LJ, et al. The effect of genomic inversions on estimation of population genetic parameters from SNP data. Genetics 2013;193: 243–53.
- Houle D, Márquez EJ. Linkage disequilibrium and inversiontyping of the Drosophila melanogaster genome reference panel. G3 (Bethesda) 2015;5:1695–701.
- Coluzzi M, Sabatini A, Petrarca V, Di Deco MA. Chromosomal differentiation and adaptation to human environments in the Anopheles gambiae complex. Trans R Soc Trop Med Hyg 1979;73:483–97.
- Main BJ, Lee Y, Ferguson HM, et al. The genetic basis of host preference and resting behavior in the major African malaria vector, Anopheles arabiensis. PLoS Genet 2016;12:e1006303.
- Rishikesh N, Di Deco MA, Petrarca V, et al. Seasonal variations in indoor resting Anopheles gambiae and Anopheles arabiensis in Kaduna, Nigeria. Acta Trop 1985;42:165–70.
- Ayala D, Zhang S, Chateau M, et al. Association mapping desiccation resistance within chromosomal inversions in the African malaria vector *Anopheles gambiae*. Mol Ecol 2019;28:1333–42.
- Petrarca V, Nugud AD, Elkarim Ahmed MA, et al. Cytogenetics of the Anopheles gambiae complex in Sudan, with special reference to An. arabiensis: relationships with East and West African populations. Med Vet Entomol 2000;14:149–64.
- 19. Gray EM, Rocca KAC, Costantini C, et al. Inversion 2La is associated with enhanced desiccation resistance in *Anopheles* gambiae. Malar J 2009;8:215.
- Rocca KAC, Gray EM, Costantini C, et al. 2La chromosomal inversion enhances thermal tolerance of Anopheles gambiae larvae. Malar J 2009;8:147.
- 21. Fouet C, Gray E, Besansky NJ, et al. Adaptation to aridity in the malaria mosquito *Anopheles gambiae*: chromosomal inversion polymorphism and body size influence resistance to desiccation. PLoS One 2012;7:e34841.
- 22. Ayala D, Acevedo P, Pombi M, et al. Chromosome inversions and ecological plasticity in the main African malaria mosquitoes. Evolution 2017;71:686–701.
- Cheng C, Tan JC, Hahn MW, et al. Systems genetic analysis of inversion polymorphisms in the malaria mosquito. Proc Natl Acad Sci U S A 2018;115:E7005–14.
- 24. Ayala D, Caro-Riaño H, Dujardin J-P, et al. Chromosomal and environmental determinants of morphometric variation in natural populations of the malaria vector Anopheles funestus in Cameroon. Infect Genet Evol 2011;11:940–7.
- Riehle MM, Bukhari T, Gneme A, et al. The Anopheles gambiae 2La chromosome inversion is associated with susceptibility to Plasmodium falciparum in Africa. Elife. 2017;6:e25813. http: //dx.doi.org/10.7554/elife.25813
- Petrarca V, Beier JC. Intraspecific chromosomal polymorphism in the Anopheles gambiae complex as a factor affecting malaria transmission in the Kisumu area of Kenya. Am J Trop Med Hyg 1992;46:229–37.
- 27. Gillies MT, De Meillon B. The Anophelinae of Africa South of the Sahara (Ethiopian Zoogeographical Region). South African Institute for Medical Research; 1968.
- Coetzee M, Fontenille D. Advances in the study of Anopheles funestus, a major vector of malaria in Africa. Insect Biochem Mol Biol 2004;34:599–605.
- Coetzee M, Koekemoer LL. Molecular systematics and insecticide resistance in the major African malaria vector Anopheles funestus. Annu Rev Entomol 2013;58:393–412.

- 30. Dia I, Guelbeogo MW, Ayala D. Advances and perspectives in the study of the malaria mosquito Anopheles funestus. In: Anopheles Mosquitoes - New Insights into Malaria Vectors. 2013, doi:10.5772/55389.
- 31. Zahar AR, World Health Organization. Vector Bionomics in the Epidemiology and Control of Malaria: The WHO African region & the southern WHO eastern Mediterranean region. 1985. http://www.who.int/iris/handle/10665/62183. Accessed 2 Nov 2018.
- 32. Menze BD, Riveron JM, Ibrahim SS, et al. Multiple insecticide resistance in the malaria vector *Anopheles funestus* from northern Cameroon is mediated by metabolic resistance alongside potential target site insensitivity mutations. PLoS One 2016;11:e0163261.
- 33. Riveron JM, Ibrahim SS, Mulamba C, et al. Genome-wide transcription and functional analyses reveal heterogeneous molecular mechanisms driving pyrethroids resistance in the major malaria vector Anopheles funestus across Africa. G3 (Bethesda) 2017;7:1819–32.
- 34. Ndo C, Kopya E, Donbou MA, et al. Elevated Plasmodium infection rates and high pyrethroid resistance in major malaria vectors in a forested area of Cameroon highlight challenges of malaria control. Parasit Vectors 2018;11:157.
- 35. Anopheles Program. https://www.beiresources.org/AnophelesProgram.aspx. Accessed 2 Nov 2018.
- Michel AP, Ingrasci MJ, Schemerhorn BJ, et al. Rangewide population genetic structure of the African malaria vector Anopheles funestus. Mol Ecol 2005;14:4235–48.
- Michel AP, Guelbeogo WM, Grushko O, et al. Molecular differentiation between chromosomally defined incipient species of Anopheles funestus. Insect Mol Biol 2005;14:375–87.
- Guelbeogo WM, Grushko O, Boccolini D, et al. Chromosomal evidence of incipient speciation in the Afrotropical malaria mosquito Anopheles funestus. Med Vet Entomol 2005;19:458– 69.
- Costantini C, Sagnon N, Ilboudo-Sanogo E, et al. Chromosomal and bionomic heterogeneities suggest incipient speciation in Anopheles funestus from Burkina Faso. Parassitologia 1999;41:595–611.
- Guelbeogo WM, Sagnon N 'fale, Grushko O, et al. Seasonal distribution of Anopheles funestus chromosomal forms from Burkina Faso. Malar J 2009;8:239.
- Matthews BJ, Dudchenko O, Kingan SB, et al. Improved reference genome of Aedes aegypti informs arbovirus vector control. Nature 2018;563:501–7.
- Waterhouse RM, Seppey M, Simão FA, et al. BUSCO applications from quality assessments to gene prediction and phylogenomics. Mol Biol Evol 2018;35(3):543–8.
- 43. Chin C-S, Alexander DH, Marks P, et al. Nonhybrid, finished microbial genome assemblies from long-read SMRT sequencing data. Nat Methods 2013;10:563–9.
- 44. Walker BJ, Abeel T, Shea T, et al. Pilon: an integrated tool for comprehensive microbial variant detection and genome assembly improvement. PLoS One 2014;9:e112963.

- Ewing B, Hillier L, Wendl MC, et al. Base-calling of automated sequencer traces using phred. I. Accuracy assessment. Genome Res 1998;8:175–85.
- Waterhouse RM, Aganezov S, Anselmetti Y, et al. Leveraging evolutionary relationships to improve Anopheles genome assemblies. bioRxiv 2019, doi:10.1101/434670
- Durand NC, Robinson JT, Shamim MS, et al. Juicebox provides a visualization system for Hi-C contact maps with unlimited zoom. Cell Syst 2016;3:99–101.
- Koren S, Walenz BP, Berlin K, et al. Canu: scalable and accurate long-read assembly via adaptive k-mer weighting and repeat separation. Genome Res 2017;27:722–36.
- 49. Langmead B, Salzberg SL. Fast gapped-read alignment with Bowtie 2. Nat Methods 2012;9:357–9.
- English AC, Richards S, Han Y, et al. Mind the gap: upgrading genomes with Pacific Biosciences RS long-read sequencing technology. PLoS One 2012;7:e47768.
- Kurtz S, Phillippy A, Delcher AL, et al. Versatile and open software for comparing large genomes. Genome Biol 2004;5:R12.
- Krzywinski M, Schein J, Birol I, et al. Circos: an information aesthetic for comparative genomics. Genome Res 2009;19:1639–45.
- Sharakhov IV, Sharakhova MV. Heterochromatin, histone modifications, and nuclear architecture in disease vectors. Curr Opin Insect Sci 2015;10:110–7.
- 54. Cabanettes F, Klopp C. D-GENIES: dot plot large genomes in an interactive, efficient and simple way. PeerJ 2018;6:e4958.
- 55. Koren S, Rhie A, Walenz BP, et al. De novo assembly of haplotype-resolved genomes with trio binning. Nat Biotechnol 2018, doi:10.1038/nbt.4277.
- Li H, Durbin R. Fast and accurate long-read alignment with Burrows-Wheeler transform. Bioinformatics 2010;26:589–95.
- 57. Bishara A, Liu Y, Weng Z, et al. Read clouds uncover variation in complex regions of the human genome. Genome Res 2015;**25**:1570–80.
- Wood DE, Salzberg SL. Kraken: ultrafast metagenomic sequence classification using exact alignments. Genome Biol 2014;15:R46.
- Sedlazeck FJ, Rescheneder P, Smolka M, et al. Accurate detection of complex structural variations using single-molecule sequencing. Nat Methods 2018;15:461–8.
- Kim D, Langmead B, Salzberg SL. HISAT: a fast spliced aligner with low memory requirements. Nat Methods 2015;12:357– 60.
- Grabherr MG, Haas BJ, Yassour M, et al. Full-length transcriptome assembly from RNA-Seq data without a reference genome. Nat Biotechnol 2011;29:644–52.
- 62. Wu TD, Watanabe CK. GMAP: a genomic mapping and alignment program for mRNA and EST sequences. Bioinformatics 2005;**21**:1859–75.
- 63. Ghurye J, Koren S, Small ST, et al. Supporting data for "A chromosome-scale assembly of the major African malaria vector Anopheles funestus." GigaScience Database 2019. http: //dx.doi.org/10.5524/100602.