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Radical remodeling of the Y chromosome in a recent radiation of malaria mosquitoes

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Y chromosomes control essential male functions in many species, including sex determination and fertility. However, because of obstacles posed by repeat-rich heterochromatin, knowledge of Y chromosome sequences is limited to a handful of model organisms, constraining our understanding of Y biology across the tree of life. Here, we leverage long single-molecule sequencing to determine the content and structure of the nonrecombining Y chromosome of the primary African malaria mosquito, *Anopheles gambiae*. We find that the *An. gambiae* Y consists almost entirely of a few massively amplified, tandemly arrayed repeats, some of which can recombine with similar repeats on the X chromosome. Sex-specific genome resequencing in a recent species radiation, the *An. gambiae* complex, revealed rapid sequence turnover within *An. gambiae* and among species. Exploiting 52 sex-specific *An. gambiae* RNA-Seq datasets representing all developmental stages, we identified a small repertoire of Y-linked genes that lack X gametologs and are not Y-linked in any other species except *An. gambiae*, with the notable exception of YG2, a candidate male-determining gene. YG2 is the only gene conserved and exclusive to the Y in all species examined, yet sequence similarity to YG2 is not detectable in the genome of a more distant mosquito relative, suggesting rapid evolution of Y chromosome genes in this highly dynamic genus of malaria vectors. The extensive characterization of the *An. gambiae* Y provides a long-awaited foundation for studying male mosquito biology, and will inform novel mosquito control strategies based on the manipulation of Y chromosomes.

Anopheles gambiae | PacBio | RNA-Seq | tandem repetitive DNA | Y-chromosome

Sex chromosomes carry a master switch gene responsible for sex determination (1). They are thought to derive from an ordinary pair of autosomes, and have multiple independent origins across the tree of life (2, 3). In animals with morphologically distinct (heterogametic) sex chromosomes, the Y has ceased crossing over with the X across some or all of its length and the nonrecombining region is transmitted clonally by males (4, 5). The absence of recombination initiates progressive genetic decay—gene loss and accumulation of repetitive sequences—but there is increasing recognition that even relatively old and otherwise highly degenerate Y chromosomes retain functional importance not only for sexual reproduction but for their contributions to global gene regulation, affecting health and survival (6–10). Notwithstanding these critical roles, the Y chromosome remains one of the most recalcitrant and poorly characterized portions of any genome

more than a decade into the postgenomic era, with current knowledge resting largely on only two animal groups: mammals and *Drosophila* (2, 11).

Mosquitoes in the genus *Anopheles* are the exclusive vectors of human malaria, a disease that claimed nearly 600,000 lives globally in 2013, the majority in sub-Saharan Africa (12). Although 15 y of

Significance

Interest in male mosquitoes has been motivated by the potential to develop novel vector control strategies, exploiting the fact that males do not feed on blood or transmit diseases, such as malaria. However, genetic studies of male *Anopheles* mosquitoes have been impeded by the lack of molecular characterization of the Y chromosome. Here we show that the *Anopheles gambiae* Y chromosome contains a very small repertoire of genes, with massively amplified tandem arrays of a small number of satellites and transposable elements constituting the vast majority of the sequence. These genes and repeats evolve rapidly, bringing about remodeling of the Y, even among closely related species. Our study provides a long-awaited foundation for studying mosquito Y chromosome biology and evolution.

Author contributions: S.J.E., I.V.S., Z.J.T., and N.J.B. conceived the project; N.J.B. coordinated the project; A.B.H., N.H.B., J.H., D.R., A.M.P., Z.J.T., and N.J.B. performed genome and BAC sequencing; S.K. and A.M.P. performed PacBio read correction and assembly; P.-A.P., C.C., O.S.A., A. Cagnetti, A. Crisanti, T.D., E.F., R.G., T.N., and N.W. performed RNA-Seq, RT-PCR, and gene validation; A.B.H., P.-A.P., C.C., L.A., X.J., A. Steele, S.Z., M.W.H., S.J.E., and Z.J.T. performed computational analysis of Y linkage; A. Sharma, M.V.S., V.A.T., and I.V.S. performed cytogenetics and FISH; C.C. and M.W.H. performed phylogeny reconstruction and simulations; and A.B.H., P.-A.P., A.M.P., Z.J.T., and N.J.B. wrote the paper with input from the other authors.

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intensified vector control efforts (mainly insecticide-impregnated bed nets) have successfully averted an estimated 663 million clinical cases of malaria (13), further progress toward elimination in the most malarious regions will depend upon the development of novel methods of vector control complementary to existing approaches (14). One method currently under development entails genetic modification of the mosquito to bias the population sex ratio toward males (which do not bite), with the goal of local population reduction or elimination (15–17). Modeling has shown that the most efficient means toward this end is the engineering of a driving Y chromosome (18). A molecular-level understanding of the *Anopheles* Y chromosome is important to inform and optimize such a strategy.

Historical cytogenetic studies established that the *Anopheles* Y chromosome is entirely heterochromatic, but also suggested that, contrary to *Drosophila* and in common with mammals, it bears partial homology to the X chromosome and plays a male-determining role (19–24). However, efforts to characterize Y chromosome sequences in *Anopheles* have been thwarted by a lack of directed resources and effective tools. The unsurpassed medical importance of the African malaria mosquito *Anopheles gambiae* motivated its early selection for whole-genome sequencing (25), making it second only to the model organism *Drosophila melanogaster* as a fully sequenced insect genome. However, because of formidable obstacles to assembling repeat-dense Y chromosome sequences (26), efforts to assemble or even to assign Y chromosome sequences in the framework of the *An. gambiae* genome project were largely unsuccessful, despite separate sequencing of males and females (27), leaving its content and organization obscure. Here, we leverage the increased length and reduced bias of single-molecule sequencing (28), along with Illumina-based sex-specific transcriptional profiling and whole-genome sequencing, to identify an extensive dataset of Y chromosome sequences and explore their organization and evolution in the young species radiation known as the *An. gambiae* complex (29), which contains some of the most important vectors of human malaria. We find massive remodeling of the repeat-dense Y chromosome and remarkably few genes, none of which have counterparts on the X. Only the *YG2* gene—the earliest to be expressed in 3-h male embryos—is conserved and exclusive to the Y across species in the complex, and thus is a possible male-determining factor. However, no sequence similarity to *YG2* can be detected in the genome of an Asian malaria vector from the same subgenus (30), underscoring the rapid evolution of Y-linked genes in the evolutionarily dynamic genus *Anopheles* (31). Contrary to the species branching order, the *YG2* gene tree from the *An. gambiae* complex supported the grouping of two major malaria vectors with a history of substantial autosomal introgression (32), consistent with the hypothesis that even the Y chromosome may have crossed species boundaries. Although a Y chromosome assembly awaits further technological developments, the compilation and comparative analysis of Y chromosome sequences in the *An. gambiae* complex substantially advances our understanding of the composition, organization, and evolution of the *Anopheles* Y chromosome and lays the groundwork for exploiting the Y chromosome to control disease transmission.

Results

Identification of *An. gambiae* Y Chromosome Sequences. Gross cytological estimates suggest that the *An. gambiae* Y chromosome constitutes ~10% of the 264-Mb *An. gambiae* genome (27, 33), yet a mere 0.18 Mb of unordered sequences have previously been assigned to the Y in the PEST reference genome assembly [<https://www.vectorbase.org> (34)]. Similarly, a recent *Anopheles stephensi* genome project identified only 57 short unordered Y sequences spanning ~50 kb from genomic reads, and 11 contigs spanning ~200 kb from BAC clones that were assigned to the Y chromosome (35). To overcome this impediment, we developed a strategy

based on long-read, PacBio single molecule real-time sequencing (36). Template genomic DNA was extracted from male siblings of a single-pair mating that inherited the same paternal *An. gambiae* Y chromosome and was sequenced to 70× autosomal (35× heterosomal) coverage with PacBio single molecule real-time sequencing (*SI Appendix, Text S1.1*). Consensus-based error correction with PBeR (37) resulted in 40× autosomal (20× heterosomal) coverage of PacBio-corrected reads with an N50 size of 2,799 bp (*SI Appendix, Text S1.2*). A whole-genome assembly of the entire PacBio dataset was performed with Celera Assembler (38, 39), resulting in a 294-Mb assembly with an N50 contig size of 101,465 bp (*SI Appendix, Text S1.2*). However, the moderate coverage and average raw read length of 2,479 bp proved insufficient for de novo reconstruction of the Y chromosome. Initial analysis of this assembly revealed highly fragmented heterosomal contigs and evidence that autosomal and heterosomal sequence had been incorrectly joined. Because of known limitations in assembling heterochromatic sequence and concerns about potential mis-assemblies, we decided to focus exclusively on individual (unassembled) PacBio-corrected reads from genomic DNA for all subsequent Y chromosome analysis.

As a complementary strategy to investigate the organization of large (100 kb) contiguous pieces of the *Anopheles* Y chromosome, PacBio sequencing of individual *An. gambiae* BAC clones was performed (*SI Appendix, Text S1.3*). These BACs were deemed potentially Y-linked based on initial computational analysis of available BAC-end sequences (*SI Appendix, Text S1.3*). Directed, high-coverage PacBio sequencing, ranging from 300× to 2,000× per BAC, yielded sufficient information to completely assemble each BAC without ambiguity. Successful BAC assembly was a result of localization of the repeat structure as well as the tremendous sequencing depth attained. Because PacBio read lengths are exponentially distributed, such deep sequencing increases the probability of obtaining some very long reads (e.g., >20 kb), which were necessary for the assembly of these highly repetitive sequences. Comprehensive computational analysis supported three of four assembled BACs as originating from the Y chromosome (*SI Appendix, Text S1.3, and Figs. S1 and S2*).

To identify presumptive Y-linked sequences among the unassembled genomic PacBio reads, we implemented two recent computational approaches (*SI Appendix, Text S2*) that exploit short-read (Illumina) genomic sequencing from sex-specific DNA pools (*SI Appendix, Text S1.4, and Table S1*). The Y chromosome genome scan (YGS) approach was designed to operate on scaffolds from a genome assembly derived from mixed sexes or males; after identification and masking of identical repeats, scaffolds are classified as Y-linked if they have few or no kmer-length matches to female Illumina sequences (40). As applied to *An. gambiae* male PacBio-corrected reads, which were treated as “scaffolds,” YGS failed to unambiguously classify Y-linked sequences, apparently because of the extremely small fraction of Y chromosome sequence that is exclusive to the Y in *An. gambiae* as opposed to highly enriched there (see below and *SI Appendix, Table S8*). A second approach, the chromosome quotient (CQ) method (30), infers Y-linkage based on the female-to-male ratio of sequence alignments to a reference, in this case *An. gambiae* female-to-male Illumina sequences aligned to PacBio reads. At a conservative threshold value (CQ ≤ 0.2) imposed across the length of a PacBio read, the CQ method classified 79,475 unassembled reads (246 Mb) as presumptive Y chromosome sequences, which populate a database that we denote Ydb (*SI Appendix, Text S2.2, and Tables S4–S7, and Dataset S1*). Although the rate of false-positives in Ydb should be low (30) (*SI Appendix, Text S2.1*), the conservative CQ threshold necessarily means that Ydb is incomplete with respect to possible Y chromosome sequences that share extended sequence identity with other chromosomes, as would be expected for pseudoautosomal regions or sequences recently acquired from elsewhere in the genome. However, it is likely that Ydb represents much of the

nonrecombining (male-limited) Y chromosome (NRY) (*SI Appendix, Table S4*). Greater than 94% of sequence classes comprising Ydb were validated as Y-linked in *An. gambiae* (Fig. 1 and *SI Appendix, Text S3*, and *Table S9*) through genomic PCR and physical mapping by FISH of representative sequences to mitotic chromosomes of male *An. gambiae* larvae (*SI Appendix, Text S4*).

The *An. gambiae* Y Contains Massively Amplified Satellites and Retrotransposons. We conducted a detailed computational assessment of Y chromosome repeat content based on analysis of Ydb. Our inferences should be minimally affected by redundant and overlapping reads, as they are based on proportional content (relative abundance), and PacBio coverage has limited bias (28, 38). Initially, both PacBio Ydb reads and assembled BAC sequences were screened for interspersed repeats and low-complexity DNA with RepeatMasker 4.0.3 (41), using the *An. gambiae* PEST RepeatMasker library augmented with previously characterized Y chromosome satellite and retrotransposon sequences (42, 43) (*SI Appendix, Text S3*). Anticipating that the *An. gambiae* Y chromosome contains previously unknown repeats, or repeats whose structures differ from those represented in the reference repeat library, we characterized both annotated and unclassified output from RepeatMasker through iterative clustering and consensus building of sequences in Ydb and the Y-linked BACs. This strategy ultimately revealed that ~98% of bases in Ydb belong to a very few repetitive sequence classes, amplified extensively (Fig. 2A and *SI Appendix, Text S3*, and *Table S9*).

Satellite DNA accounts for ~49% of all bases in Ydb (*SI Appendix, Table S9*). However, only six different satellite monomers were identified and two—AgY477 and AgY373—predominate, comprising 93% of all satellite DNA sequence in Ydb. Moreover, the satellite sequences are found as long tandem arrays in Ydb, largely devoid of transposable elements (TEs). These data suggest that satellite DNA is an abundant and homogeneous constituent of the NRY, a major Y chromosome sequence feature that we refer to as the satellite amplified region (SAR) (Fig. 2A). The absence of other repetitive sequence classes interspersed within the SAR suggests limited genetic exchange between satellites and other repeats, but we find evidence of recombination and higher-order repeat structures among satellites within the SAR. Different satellite monomers that share extensive sequence similarity (AgY477 and AgY373; AgY280 and AgY53D) frequently co-occur on the same PacBio read, often as interspersed or even chimeric monomers, indicative of sister chromatid or intrachromatid exchange (Fig. 2C, and *SI Appendix, Text S3.1*, and *Figs. S5–S8*).

Another 43.5% of bases in Ydb are TE-related, of which only eight distinct TE types, mainly retrotransposons, were identified (*SI Appendix, Text 3.2*, and *Table S9*). Remarkably, one particular element alone—a 6.9-kb Ty3/Gypsy LTR retrotransposon that we designate *zanzibar*—comprises almost 27% of bases in Ydb as a whole, and accounts for more than 61% of the bases classified as TEs (Fig. 2 and *SI Appendix, Table S9*). Unlike the typical interspersed arrangement of TEs in genomic euchromatin, *zanzibar* is arranged on the Y in massively amplified head-to-tail tandem arrays, in which one gag/pol region followed by one LTR is repeated in succession like beads on a string: (gag/pol+LTR)_n. From its abundance and organization, we infer that this *zanzibar* amplified region (ZAR) (Fig. 2A and B)—like the SAR—is another prominent organizational feature of the NRY. *Zanzibar* monomers (gag/pol+LTR) in the array may carry insertions of a variety of other TEs or TE fragments. Remarkably, every copy of *zanzibar* carrying a particular TE type (e.g., *mtanga*) contains precisely the same TE sequence inserted into precisely the same *zanzibar* sites (Fig. 2 and *SI Appendix, Text S3.2*), as though replica insertions in different *zanzibar* copies are not the result of independent transposition events. Taken together, these data—the precise tandem organization of the ZAR and the peculiar clonal nature of insertions—strongly suggest that *zanzibar* retrotransposons no longer function in the manner expected of autonomous transposable elements. Instead, *zanzibar* sequences appear to have been the substrate for illegitimate recombination and megabase-spanning tandem amplifications on the Y chromosome, analogous to the process described for the genesis of centromeric repeats in maize (44). Despite its evident origin as an autonomous TE, the present ZAR structure most closely resembles satellite sequence and thus may reflect a general pattern of sequence amplification and evolution on the *Anopheles* Y chromosome.

Of the remaining ~7.5% of Ydb bases, we were able to classify ~5.5% as repetitive, but the last ~2% could not be clustered (*SI Appendix, Table S9*). This small unclustered fraction contains a heterogeneous mixture of less abundant types of repetitive sequences, degenerate copies of repeats categorized above, and Y chromosome genes, many of which also appear to be multicopy (see below and *SI Appendix, Text S7*, and *Fig. S11*). Only four Ydb reads were classified by a metagenomic analysis as originating from another organism (*SI Appendix, Text S2.2*), suggesting that nearly all of Ydb is legitimate *An. gambiae* sequence.

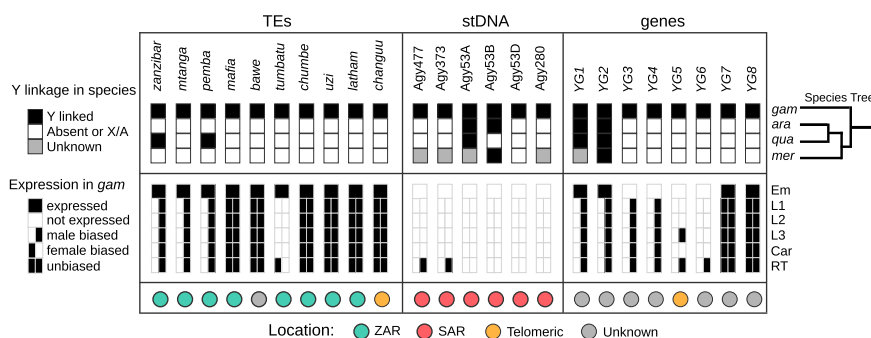


Fig. 1. Summary of major Y chromosome loci, showing rapid turnover of the Y chromosome content and expression patterns in the *An. gambiae* species complex. (Top) Black boxes indicate Y-linkage, white boxes indicate either total absence from the species or absence from its Y chromosome, and gray boxes indicate unknown status with regard to Y-linkage. Typically, sequences indicated by gray showed CQ or RCQ values of ~1, suggesting that they are either on both sex chromosomes or on autosomes. Details are provided in *SI Appendix, Tables S6, S7*, and *S15*. At right, the species branching order provides an evolutionary context of the changes in Y chromosome content within the past 2 My. Only YG2 is conserved and exclusively on the Y chromosome in all four species of the *An. gambiae* complex. (Middle) Sex-specific transcription in *An. gambiae* was assessed at different developmental stages and tissues, except for embryos (Em). (Bottom) The organization of the Y chromosome loci in *An. gambiae*, if known. *ara*, *An. arabiensis*; Car, adult carcass; Em, embryo; *gam*, *An. gambiae*; L1–L3, first- to third-instar larvae; *mer*, *An. merus*; RT, adult reproductive tissues; stDNA, satellite DNA; *qua*, *An. quadriannulatus*.

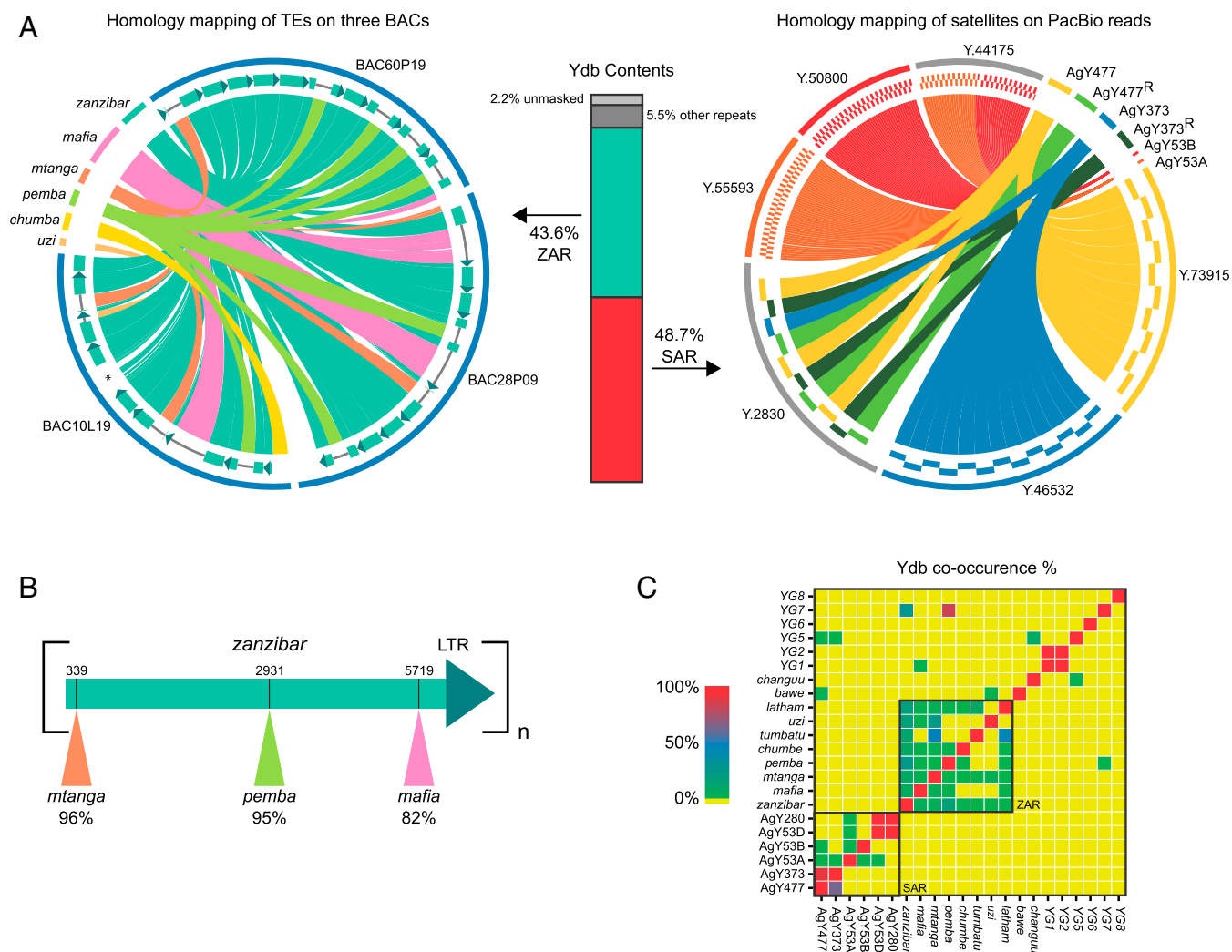


Fig. 2. The NRY of *An. gambiae* mainly consists of massively amplified tandem arrays of a small number of satellites and TEs. (A) Two major regions of the *An. gambiae* Y, the ZAR and SAR, represent 92.3% of the sequences in Ydb (vertical bar plot). Ydb reflects the content of NRY in *An. gambiae*. Percentages were calculated by masking Ydb using annotated Y chromosome loci. The *Left* circos plot, created by homology mapping of TEs on three Y chromosome BAC clones, shows the organization of the ZAR in the three BACs. As seen in these BACs, and as independently confirmed in PacBio reads, the ZAR consists of head-to-tail tandem arrays of *zanzibar*, which sometimes have other transposons inserted. The arrays of *zanzibar* units inside each BAC are shown schematically directly inside the BAC ideograms (blue semicircular lines) enclosing the circos plot. The dark green arrows of each *zanzibar* unit (shown enlarged in *B*) represent the single LTR; lines breaking *zanzibar* units indicate insertions of other TEs. A few small insertions (~200 bp) into *zanzibar* are too small to be visible in this plot. The asterisk in BAC10L19 corresponds to an atypical *zanzibar* unit that could result from recombination or misassembly. The circos plot at *Right*, constructed by homology mapping of satellite monomers on PacBio reads from Ydb, shows the organization of the SAR. Shown are representative examples of the occurrence of homo-monomeric tandem arrays (Y73915, Y46532, Y55593), junctions between homo-monomeric tandem arrays (Y44175), and recombinant arrays (Y2830). The recombinant arrays are interspersed with recombinant and nonrecombinant versions of AgY477 and AgY373 satellites (*SI Appendix, Fig. S5*). (B) Schematic of a single *zanzibar* unit, consisting of a gag/pol domain and a single LTR; each unit is organized in a head-to-tail tandem array (see *Left* circos plot in A). Shown by colored triangles are the canonical insertion sites of three other transposons (*mtanga*, *pemba*, *mafia*) into different *zanzibar* units. Percentages indicate the fraction of Ydb PacBio reads observed to carry TE insertions into *zanzibar* units at the precise insertion site illustrated (coordinates shown above the gag/pol domain). For example, we observed 243 of 256 (95%) PacBio reads in which *pemba* was inserted into *zanzibar* at position 2931. This phenomenon was independently confirmed in whole-genome sequencing Illumina reads. (C) Co-occurrence matrix of Y chromosome loci in PacBio reads from Ydb. These results show that satellite sequences co-occur (in the SAR), as do TEs (in the ZAR), but that the ZAR and SAR regions are largely independent.

Extensive Structural Dynamism of the Y Chromosome in a Young Species Radiation. Cytological observations conducted in the 1970s revealed striking differences in sex chromosome heterochromatin among populations and between species in this complex (45, 46). Not only did the staining intensity and pattern vary, but also length of the Y chromosome, ranging from less than half the length of the X in one *An. gambiae* population to almost the same length as the X in others (45). However, a mechanistic understanding of the phenomenon was lacking. Our finding that ~98% of the bases in Ydb constitute a highly repetitive sequence

organized into tandem arrays suggests that the cytological observations may have their basis in rapid expansion and contraction of tandem repeats on the Y chromosome, through unequal crossover and a variety of other mechanisms (42, 43, 47–49). We applied computational and cytogenetic methods to assess the nature and degree of Y chromosome remodeling within *An. gambiae* and among sibling species during the relatively brief (2 My) evolutionary history of the species complex.

Intraspecific variation in the SAR and ZAR of *An. gambiae* was assessed computationally among three laboratory colonies

and among 85 male and female mosquitoes sampled from a natural population in Cameroon, available through MalariaGEN's *An. gambiae* 1000 Genomes Consortium (Ag1000G) phase 1 AR3 data release (2015) (www.malariagen.net/data/ag1000g-phase1-AR3) (SI Appendix, Text S1.4–S1.5, and Tables S1 and S2). The sample set from Cameroon is one of the few available that contains both sexes. From the *An. gambiae* Pimperena colony—our reference—and two additional colonies (G3 and Asembo), we generated Illumina sequences from sex-specific genomic DNA pools, and aligned them to the consensus sequences of *An. gambiae* monomer repeat units compiled from Ydb. Alignments were performed twice, using either a strict read-mapping protocol (for CQ calculations) or a less-stringent mapping protocol used to produce a metric analogous to CQ, termed “relaxed CQ” (RCQ) (SI Appendix, Text S5, and Tables S6 and S7). The presence/absence and relative abundance of each major repeat was estimated from the number of mapped reads; male-bias was estimated from the female-to-male ratio of sequences reflected by CQ and RCQ (SI Appendix, Tables S6 and S7). Using similar strategies, we also interrogated individually sequenced wild-caught *An. gambiae* mosquitoes of both sexes (40 males, 45 females) from Cameroon (SI Appendix, Figs. S11–S13, and Tables S12–S14).

Overall, the pattern of Y-linkage as inferred by CQ and RCQ was qualitatively similar among *An. gambiae* samples. However, the corresponding copy number of Y-linked sequences was much more labile (Fig. 3 and SI Appendix, Text S5, Figs. S11–S13, and Tables S6, S7, and S13–S15). The most dramatic copy number variation of any SAR or ZAR component was displayed by satellite sequences AgY53D and AgY280 in male samples (Fig. 3). Although numbers of read alignments are not precise reflections of copy number, they can convey a rough approximation of relative abundance if copy number varies across orders of magnitude. Indeed, counts of male alignments to AgY53D and AgY280 spanned four to five orders of magnitude from the Asembo to the Pimperena colony (SI Appendix, Tables S6 and S7), and even within the natural population from Cameroon, normalized alignment counts among individual males spanned three orders of magnitude (SI Appendix, Table S13), suggesting major expansions or contractions in array length. Copy number variation of this magnitude on the Y chromosome would be expected to affect its length. We estimated the combined effect on chromosome length of copy number variation of all SAR and ZAR components among the male *An. gambiae* sampled from Cameroon. Taking these data together, we find that copy number variation can account for the gain or loss of up to ~30 Mb of the Y chromosome. Broadly consistent with this in silico estimate, our cytogenetic length estimates also varied by as much as 22 Mb among individual males from the *An. gambiae* Pimperena colony (from ~25.9 Mb to ~47.8 Mb) (SI Appendix, Text S4.2, and Table S11), in conformity with prior studies (50).

The sibling species complex to which *An. gambiae* belongs radiated rapidly and recently, within the last 2 My (32). To examine the extent of structural divergence of the Y chromosome between species over this relatively short time frame, we generated Illumina sequences from sex-specific pools of three additional members of the complex (*Anopheles arabiensis*, *Anopheles quadriannulatus*, *Anopheles merus*) (SI Appendix, Table S1), and assessed male bias and relative abundance as described above for *An. gambiae* (SI Appendix, Text S5.2, and Tables S6, S7, and S15), and by FISH (Fig. 4). Rapid and extensive remodeling of the Y chromosome between species is evidenced by dramatic examples of turnover in both the SAR and ZAR. Satellite AgY477, heavily male-biased and a major component of the Y in *An. gambiae*, is abundant but not strongly sex-biased in *An. merus*, and is not detected in *An. arabiensis* or *An. quadriannulatus* of either sex (Fig. 1 and SI Appendix, Text S5.2, and Table S15). Similarly, *zanzibar* is neither strongly sex-biased nor abundant in *An. arabiensis* and *An. merus*, yet in *An.*

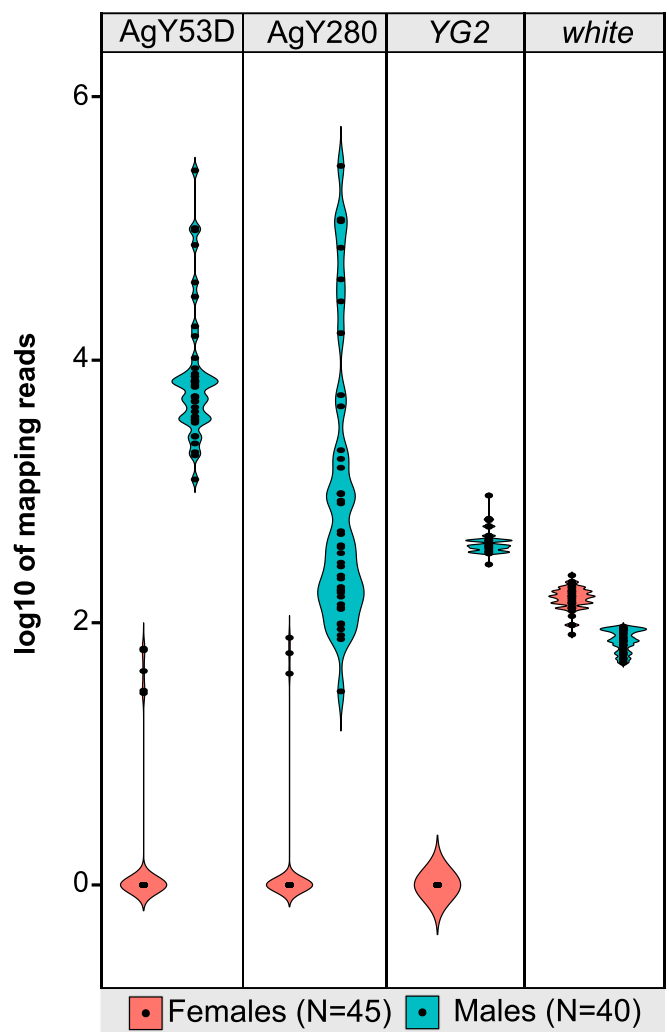


Fig. 3. Satellites AgY53D and AgY280 show extensive structural dynamism in males from a natural population of *An. gambiae*. Shown are violin plots of the \log_{10} numbers of normalized read alignments from Illumina genomic sequence derived from 40 individual male (blue) and 45 individual female (pink) mosquitoes from Cameroon, mapped to satellite monomers of AgY53D and AgY280. For comparison are similar plots of reads mapping to the presumptive male-determining gene, YG2, and to the single-copy X-linked gene, *white*. Numbers of read alignments to the satellite monomers varies drastically between individuals, in contrast to YG2 and *white*, suggesting large within-population differences in satellite abundance on the Y. Mapping reads were normalized to library size and locus length.

quadriannulatus (not a sister species of *An. gambiae*) this retrotransposon has an *An. gambiae*-like pattern of expanded tandem arrays on the Y chromosome (Figs. 1 and 4, and SI Appendix, Text S5.2, and Table S15).

The *Anopheles* Y Recombines with the X Chromosome. Meiotic pairing, chiasma formation, and crossing-over between the sex chromosomes have been reported for three anopheline species in two different subgenera (19, 21, 24), but to our knowledge, similar observations have not been reported in *An. gambiae*. Indeed, the apparent stability of X- and Y-linked translocations in *An. gambiae* (51) suggest that legitimate crossing-over between the X and Y chromosomes does not occur. However, our data are difficult to explain without some form of X–Y genetic exchange since the evolution of the sex chromosomes from an ancestral pair of autosomes, an event that must predate anopheline diversification

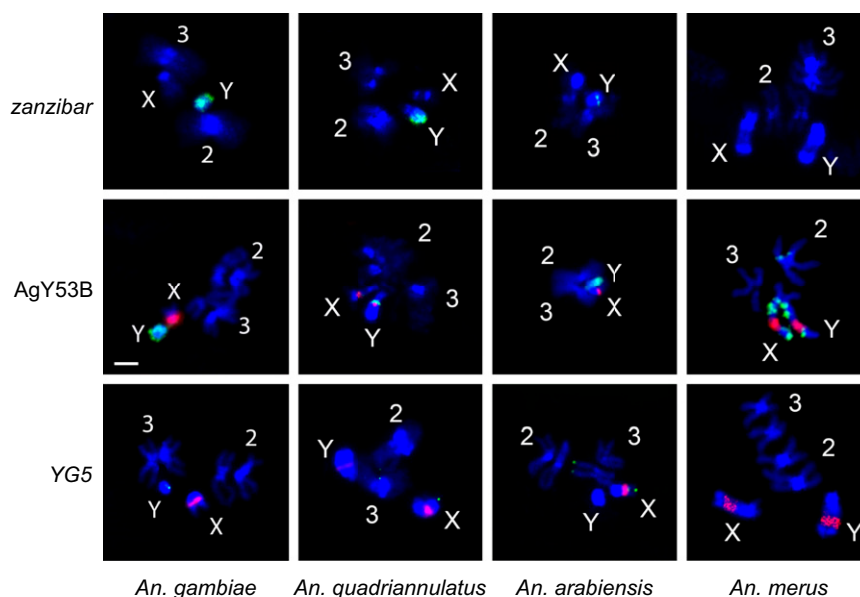


Fig. 4. Physical mapping supports structural dynamism of Y chromosome sequences in the *An. gambiae* complex. FISH of retrotransposon *zanzibar*, satellite AgY53B, and gene YG5 (green signals) was performed on chromosomes of male *An. gambiae* Kisumu (*zanzibar*, YG5), *An. gambiae* Asembo (AgY53B), *An. quadriannulatus* SANGWE, *An. arabiensis* Dongola, and *An. merus* MAF. Chromosomes were obtained from imaginal discs except for *An. merus* chromosomes hybridized to AgY53B, which were obtained from testes. The 18S rDNA probe (red signal) was used in all experiments except with *zanzibar*. Chromosomes were counterstained with DAPI (blue). (Scale bar, 2 μ m; applies to all images.)

~100 Mya. First, there is a very high degree of sequence similarity between Y-associated and X-associated repetitive DNA in *An. gambiae*. Our initial indication of this was the near-complete failure of the YGS computational method to identify Y-associated sequences in *An. gambiae*, presumably arising from the fact that such sequences are only rarely exclusive to the Y chromosome, as opposed to highly enriched there. Strongly supporting empirical evidence comes from physical (FISH) mapping of individual components of the SAR (e.g., AgY53B) (Fig. 4 and *SI Appendix*, Fig. S10), or of fluorescently labeled sequences derived from the entire microdissected Y chromosome (Fig. 5A), which reveals extensive cross-hybridization of Y repeats with X chromosome heterochromatin because of sequence similarity between satellite monomers (*SI Appendix*, Text 4.1.2). Detailed sequence analysis indicates that satellite monomers normally abundant only on the Y (AgY373, AgY477) share ~93% pairwise sequence similarity with a satellite monomer from the X chromosome [AgX367 (42)] (*SI Appendix*, Fig. S6). Moreover, we found the footprints of recombination on individual *An. gambiae* PacBio reads. These contained AgX367 monomers, together with AgY477 and AgY373 monomers, including AgX367-AgY373 and AgX367-AgY477 recombinants (Fig. 5C), confirming previous evidence of recombination based on PCR amplicon sequencing (42). A second line of evidence for occasional X–Y genetic exchange emerged from individually sequenced male and female *An. gambiae* from Cameroon (Fig. 5B and *SI Appendix*, Text S6, and Fig. S11). In a small subset of females, normalized counts of Illumina reads mapping to consensus satellite monomers normally abundant only on the Y chromosome (e.g., AgY53A) were almost two orders of magnitude larger than the median for all females (*SI Appendix*, Tables S12 and S14). Because this female subset lacked correspondingly high copies of other Y chromosome sequences, contamination by male genomic DNA (whether in the laboratory or via sperm stored in the spermatheca) is an unlikely alternative explanation. Although intrachromosomal genetic exchange on the X may contribute to the strongly bimodal pattern of satellite abundance observed in Cameroon

females, this phenomenon must have its basis in periodic X–Y genetic exchange.

Rapid Turnover of the Small Y Chromosome Genic Repertoire. Only three Y-linked genes had been identified previously in *An. gambiae* (30), designated *gYG1* to *gYG3* (hereafter, *YG1* to *YG3*). Aiming for comprehensive gene discovery on the *An. gambiae* Y chromosome, we performed extensive transcriptional profiling of developmentally staged *An. gambiae* embryos (nine time points), sexed larvae (three time points), and adults (whole and dissected males and females) through mRNA sequencing (RNA-Seq)—52 datasets in total (*SI Appendix*, Text S1.6, and Table S3)—and integrated complementary approaches to gene finding (*SI Appendix*, Text S7). Gene candidates bearing significant similarity to known TEs or bacterial sequences were discounted. Arising from the combined approaches were eight presumptive genes (*YG1*–8), including the three previously identified (Fig. 1 and *SI Appendix*, Text S7).

To be considered valid Y genes, we required that they exhibit male-biased or male-specific expression from RNA-Seq as well as male-specific amplification by genomic PCR, conditions met by *YG1*–5. With the exception of *YG4*, this validated set was further confirmed by male-specific RT-PCR. Moreover, we were able to physically localize *YG5* to the Y chromosome by FISH (a homolog was also detected on chromosome 3) (Fig. 4 and *SI Appendix*, Text S7, and Fig. S19). Because we could not identify male-specific SNPs distinguishing *YG6*–8 from their autosomal homologs, these genes could not be validated despite exclusive expression of *YG6* in male accessory glands (*SI Appendix*, Fig. S20), and elevated numbers of normalized read alignments to *YG8* from individually sequenced males versus females in our population sample from Cameroon (*SI Appendix*, Fig. S11, and Tables S12–S14).

Beyond the strikingly small total number of genes identified as Y-linked in *An. gambiae* following this intensive search, it is noteworthy that gene number varies even between strains. Both *YG3* and *YG4* are Y-linked exclusively in the G3 strain of *An. gambiae*, not in Asembo or Pimperena (*SI Appendix*, Tables S6 and S7).

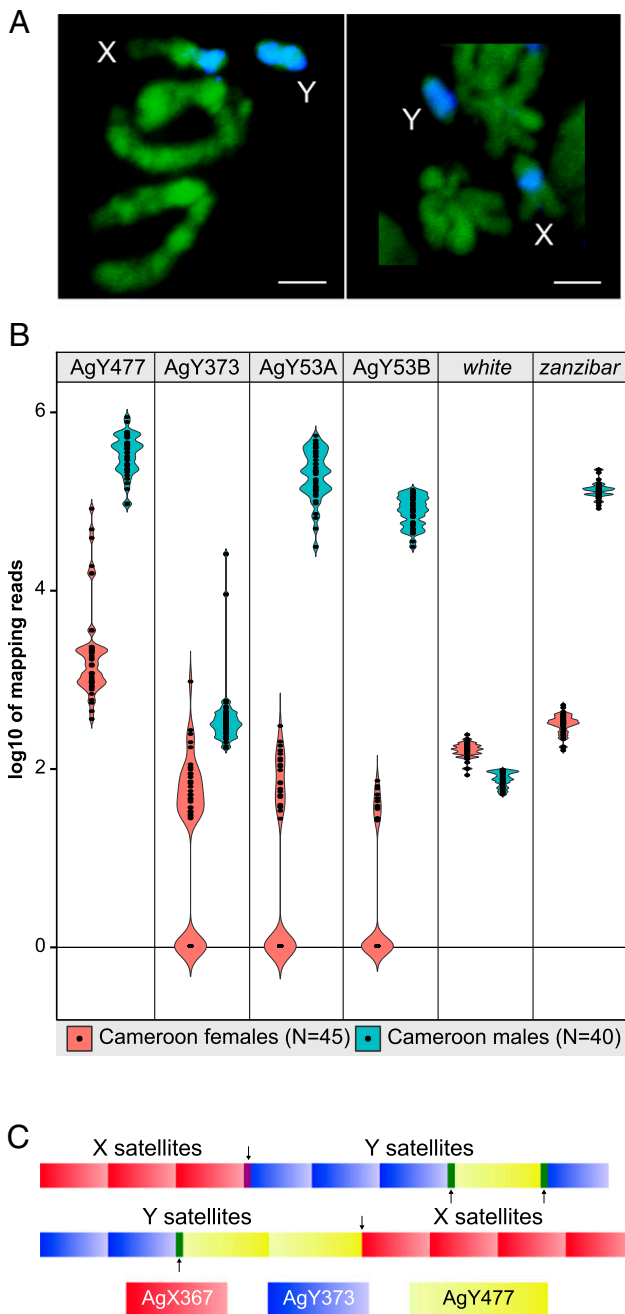


Fig. 5. The *An. gambiae* X and Y chromosomes are not genetically isolated. (A) Painting of prometaphase (Left) and metaphase (Right) chromosomes from male larval imaginal discs of the *An. gambiae* Pimperena strain with a probe generated from microdissected Y chromosomes (labeled blue by the WGA3 kit with dNTP-Cy3). Chromosomes are counterstained with YOYO-1 (green). (Scale bars, 2 μ m.) (B) Violin plots showing the \log_{10} number of normalized read alignments from 40 individual *An. gambiae* males (blue) and 45 females (pink) from the Cameroon population, to satellite AgY477, AgY373, AgY53A, and AgY53B monomers, compared with numbers of reads from these sources aligning to the *white* gene (single-copy and X-linked) and *zanzibar* (heavily Y-biased). Mapping reads were normalized to library size and locus length. (C) Two examples of single PacBio reads (pacbio_7224704_1 and pacbio_5551309_1) where predominantly X-linked (AgY367; shown in red) and predominantly Y-linked (AgY373, shown in blue; AgY477, shown in yellow) satellites occur in the same PacBio read. Black arrows indicate the junction between the predominantly X-linked and predominantly Y-linked satellites. The purple and orange boxes indicate inferred recombination, between AgX367-AgY373 and AgX367-AgY477, respectively. Green boxes indicate recombination between AgY477-AgY373.

None of these genes have recognizable gametologs on the X, yet all have partial or complete homologs on the autosomes (*SI Appendix, Text S7, and Fig. S14*), suggesting that they have been gained on the Y chromosome since its divergence from the X (see below).

To screen for candidate Y-linked genes in three *An. gambiae* sibling species (*An. arabiensis*, *An. merus*, and *An. quadriannulatus*), we used the male and female Illumina sequences from each species in conjunction with corresponding genome assemblies, mixed-sex transcript sets, and de novo RNA-Seq assemblies (31), as well as *An. gambiae* genomic resources (*SI Appendix, Text S7*). After merging the results of these approaches, we made two surprising observations. First, among all of the Y-linked genes identified in *An. gambiae*, only one—*YG2*—was computationally detected and confirmed (by male-specific genomic PCR) as Y-linked in each of the other three very closely related species (Fig. 1). None of the other *An. gambiae* Y genes, with the sole exception of *YG1*, was Y-linked in any other species examined. For *YG1*, Y-linkage was validated in *An. arabiensis* and *An. quadriannulatus*, but in *An. merus* the ratio of female-to-male alignments was inconclusive and male-specific genomic PCR amplification was not possible because of highly similar sequence elsewhere in the genome (Fig. 1 and *SI Appendix, Text S7, Fig. S14, and Tables S6 and S7*). Thus, *YG2* is the only gene both conserved on, and exclusive to, the Y chromosome in all four species examined. As it is the earliest to be expressed, at 3 h of male embryonic development (*YG1* is not expressed until 4 h), *YG2* is a possible male-determining gene. Surprisingly, *YG2* is not a single-copy gene. The first suggestion that this might be the case was hinted by the number of read alignments to *YG2* from individual males in the Cameroon sample (Fig. 3). However, we have more definitive evidence for multiple, nearly identical copies of *YG2* in *An. gambiae*. Four distinct haplotypes were sampled repeatedly in Ydb PacBio reads (which derived from the same paternal Y chromosome) (*SI Appendix, Text S7.1.2, and Fig. S17*). Variant positions among *YG2* copies were not only validated by sequencing of genomic PCR amplicons from individual male *An. gambiae* derived from natural populations, but also through RNA-Seq data, which further indicates that multiple *YG2* copies are expressed (*SI Appendix, Table S17*).

With the exception of *YG2*, the near-complete absence of conserved Y-linkage between *An. gambiae* genes and corresponding genes in the sibling species was reinforced by the converse result, our second surprising observation: all genes identified as Y-linked in any one sibling species could not be assigned to the Y chromosome in any of the other species (*SI Appendix, Text S7.2, Tables S6 and S7*). In *An. quadriannulatus* we found three novel Y-linked candidate genes (*SI Appendix, Text S7, and Table S20*). In *An. arabiensis*, no genes other than *YG1* and *YG2* were detected on the Y chromosome. In *An. merus*, we found evidence supporting the duplication of a multigene segment from chromosome 3R onto the Y since it split from other *An. gambiae* complex lineages (*SI Appendix, Text S7, Table S19*). Among seven sequential genes on 3R in this segment (corresponding to AGAP009631 to -37 in *An. gambiae*), the first three (AGAP009631 to -33) and last two (AGAP009636 and -37) have detectable copies on the Y chromosome in *An. merus* (based initially on the relative number of normalized read alignments in females and males, later validated by male-specific PCR). Our data are consistent with the two intervening genes (corresponding to AGAP009634 and -35 on 3R) having been lost from the Y, and the five flanking genes becoming amplified, although further experimental evidence will be required to confidently reconstruct these events.

Possible Y Chromosome Introgression Between Hybridizing Malaria Vectors. The *YG2* gene potentially encodes a short, 56-aa peptide whose possible role in determining maleness is under

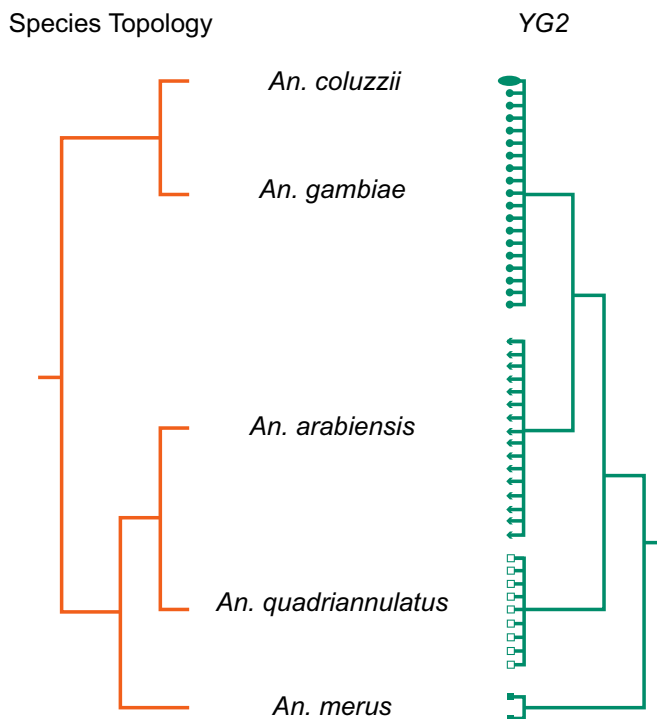


Fig. 6. Phylogeny inferred from a candidate male-determining gene on the Y chromosome, YG2, differs from the species branching order. Species topology (32) of five members of the *An. gambiae* complex (red branches) compared with a maximum-likelihood phylogeny inferred from a Y chromosome-specific region of the YG2 gene (green branches) sequenced from male *Anopheles coluzzii* (ellipse), *An. gambiae* (filled circles), *An. arabiensis* (triangles), *An. quadriannulatus* (open squares), and *An. merus* (filled squares). Samples were drawn from colonies and natural populations (see text). The YG2 tree was rooted at the midpoint; all nodes are supported by $\geq 95\%$ bootstrap replicates. The topological disagreement involves *An. arabiensis*; in the species topology *An. arabiensis* is sister to *An. quadriannulatus*, whereas the YG2 topology indicates a sister group relationship of *An. arabiensis* and *An. gambiae* + *An. coluzzii*.

investigation. In the Asian malaria vector *An. stephensi*, a Y-linked gene (*Guy1*) implicated in male determination also encodes a 56-aa sequence whose predicted secondary structure resembles that of the putative YG2 peptide (52) even though primary sequence similarity is not detectable (30) over the relatively short evolutionary span since these lineages separated, ~ 30 Mya (53). The fact that YG2 expression is detected in early embryos before any other *An. gambiae* Y-linked gene, taken together with its uniquely conserved Y chromosome location in all four *An. gambiae* complex species—in the face of otherwise rampant structural dynamism and genic turnover on the Y—is consistent with a primary role in male determination in this group. For this reason, we predicted that a gene tree reconstructed from YG2 would reflect the known species branching order (32). Although sibling species in the *An. gambiae* complex are not completely reproductively isolated, contemporary interspecific gene flow is possible only through female F1 hybrids; because their brothers are sterile, the Y chromosome cannot introgress (54). Contrary to this expectation, a YG2 tree built from sequences derived from population samples of the four species considered in this study supported *An. gambiae* and *An. arabiensis* as most closely related (Fig. 6 and *SI Appendix, Text S8*), an arrangement previously shown to be the result of massive historical introgression between these two species that involved most of the autosomes and the proximal ~ 10 Mb of the X chromosome (32). The simplest explanation for a gene tree disagreeing with the species

tree in a rapid radiation, such as the *An. gambiae* complex, is incomplete lineage sorting. We performed coalescent simulations to assess the likelihood that the grouping of *An. gambiae* with *An. arabiensis* in the YG2 tree is because of incomplete lineage sorting alone. In 62 of 1,000 simulations under the species tree, we recovered *An. gambiae* and *An. arabiensis* as sister lineages (i.e., $P = 0.062$) (*SI Appendix, Text S8*), indicating that nonintrogressing lineages could produce the observed tree a small fraction of the time. Although we cannot formally reject the null hypothesis at the 0.05 significance level, these results certainly do not rule out Y chromosome introgression. Introgression of the Y chromosome between species is conventionally viewed as unlikely (55), but it is important to consider that the pair of malaria vectors in question have historically exchanged the vast majority of the rest of their genomes, including part of the X chromosome (32). In this context, introgression of the Y chromosome is possible if not likely, as long as the introgression event(s) predated the development of male F1 hybrid sterility barriers between this species pair.

Discussion

From studies of mammals (6, 7, 11, 56, 57) and *Drosophila* (58–60), it is known that the Y chromosomes in both groups have lost most of their ancestral gene repertoire and have acquired copious amounts of repetitive and ampliconic/palindromic DNA. In the ~ 250 My since *Drosophila* and *Anopheles* last shared a common Dipteran ancestor, there has been parallel evolution of heteromorphic sex chromosomes from the same ancestral linkage group (61), implying that the *Anopheles* Y must have undergone a similar fate of massive ancestral gene loss and genomic degradation. In one main characteristic—its male determining role—the *Anopheles* Y resembles the mammalian Y more than it does *Drosophila*, in which XO flies are (sterile) males and the scant Y chromosome genes are crucial only for male fertility (2). However, in other respects, the *Anopheles* and *Drosophila* Y are much more similar. More than one-third of the human Y chromosome and 99.9% of the mouse Y is euchromatic (56, 62), whereas *Drosophila* and *Anopheles* Y chromosomes are entirely heterochromatic. Although relatively few in number, some ancestral X–Y gene pairs have been conserved throughout mammalian evolution because of their vital role as dosage-sensitive regulators of global gene expression (6, 7). Crucially, although cases are known in which a mammalian Y chromosome has acquired autosomal genes (e.g., ref. 63), most extant mammalian Y-linked genes have an X-linked gametolog. In contrast, what little gene content exists on the Y in *Drosophila* or *Anopheles* is not only poorly conserved between species, but there are no recognizable ancestral gametologs; all known Y-linked genes in *Drosophila* seem to have an autosomal origin (58). The recent DNA-based duplication of a gene from chromosome 3R to the Y chromosome in *D. melanogaster* following its split from *Drosophila simulans* ~ 4 Mya (64) mirrors our finding of a similar event in *An. merus* since the radiation of the *An. gambiae* complex, < 2 Mya. We conclude that the most salient factor uniting *Anopheles* and *Drosophila* Y chromosomes may be the continuous gain of genes and functions from the autosomes (64), in contrast to the conservation of remaining ancestral gametologs seen on the mammalian Y. However, the *Anopheles* NRY appears to stand apart from both *Drosophila* and mammalian Y chromosomes in the relative paucity of male-specific content.

One of our main findings was rapid turnover in quantity and type of repetitive DNA on the Y chromosome within and between species in the *An. gambiae* complex. It is known that both satellite and ampliconic DNA regions are prone to rapid divergence in length, structure, and sequence, as a result of unequal sister chromatid exchange between out-of-register repeat units and other mechanisms (49). On the Y chromosome such regions may be subject to accelerated rates of divergence compared with the rest of the genome. Between humans and chimps whose lineages diverged

~5 Mya, orthologous satellite arrays in the X centromere are collinear and share 93% sequence identity, whereas collinearity declines and sequence conservation drops to 78% between orthologous satellites in Y centromeres (65). Additional evidence of rapid length, structure, and sequence evolution of satellites and ampliconic structures on the Y chromosome has been reported in mice species 1–2 My diverged and mice subspecies separated by only ~900,000 y (65), between *D. melanogaster* and *D. simulans* that split ~4 Mya (60, 66), and among human males worldwide (67). Despite such pervasive remodeling of the *Anopheles* Y chromosome over short evolutionary distances, a transgene randomly inserted onto the Y chromosome of an *An. gambiae* strain in 2014 is transcriptionally active and has been stably integrated ever since, establishing that the Y chromosome is amenable to the molecular manipulation required for Y-linked genetic vector control strategies (68).

The high level of satellite DNA polymorphism within species could have important phenotypic consequences for fitness-related traits (8, 9). Moreover, the dramatic degree of satellite DNA turnover on the Y between closely related species has been implicated in hybrid incompatibility in *Drosophila* (69–71). Intriguingly, two genes known to cause hybrid incompatibility between *D. melanogaster* and *D. simulans* (*Hmr* and *Lhr*) function within species to repress transcripts from satellite DNAs and TEs (71). These species differ drastically in satellite DNA content; *D. simulans* contains fourfold less satellite DNA overall (5% versus 20% of the genome), and is particularly depauperate of the two most abundant satellite types in *D. melanogaster* (72). In *An. gambiae*, we found that AgY477 and AgY373 are the most abundant satellites on the NRY, and they are both expressed exclusively in adult male reproductive tissues; these satellite sequences are absent or altered in the other sibling species investigated (Fig. 1). Whether hybridization leads to misregulation of satellite DNA remains to be explored in the *An. gambiae* complex.

Laborious single-haplotype iterative mapping and sequencing has previously revealed the structure of mammalian Y chromosomes (11). In contrast, single-molecule sequencing now provides individual reads tens of kilobases in length, promising a resource-efficient alternative for characterizing Y chromosomes. Here, we were able to determine the content and structural characteristics of the heterochromatic *An. gambiae* NRY using this approach. Single-molecule sequencing reads were able to reveal complex repeat structures from whole-genome data and completely assemble heterochromatic BACs without manual finishing. However, the complete reconstruction of heterochromatic Y chromosomes remains a challenging open problem, as a recent PacBio assembly of *D. melanogaster* failed to completely assemble the Y chromosome (73), but it did successfully resolve the complex regions Mst77Y (74) and FDY (64). The minimum read length, accuracy, and

coverage required for the successful assembly of heterochromatin from whole-genome data are currently unknown and will vary by species, but as demonstrated here, *An. gambiae* Y chromosome BACs can be successfully reconstructed using long sequences collected from deep, directed single-molecule sequencing. These results suggest that continued single-molecule read length and throughput improvements may soon enable the complete reconstruction of Y chromosomes from whole-genome data alone.

Materials and Methods

Please see *S1 Appendix* and *Datasets S1–S3* for detailed information about: (i) genomic and transcriptomic datasets; (ii) identification of Y chromosome sequences from PacBio whole-genome sequencing; (iii) repetitive DNA content of the *An. gambiae* Y; (iv) FISH and size estimation of the Y chromosome; (v) copy number variation on the Y chromosome among individual *An. gambiae* mosquitoes and across the *An. gambiae* complex; (vi) Y chromosome recombination; (vii) genic repertoire of the Y chromosome; and (viii) phylogeny reconstruction and coalescent simulations.

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