Evolving the world’s most dangerous animal

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The devastating consequences of malaria are well known but many mysteries remain about its key protagonists, a handful of Anopheles species. New work provides a framework for solving such puzzles, by generation and analysis of whole genome assemblies for 16 Anopheles species, with genomic flexibility a key emergent theme.

The question of what makes a malaria vector is fundamental to man’s unfortunately intimate relationship with the disease. There are roughly 500 species of Anopheles, over 80% of which cannot or do not transmit malaria. Unraveling the complex interplay between morphological, physiological, life history, and behavioural differences that separate major, minor, and non-malaria vectors is a daunting task, but one which could provide fundamental information for long-term elimination. This question is at the heart of the Anopheles 16 genomes project, the first major works from which have recently been published [1,2].

Genomes and transcriptomes of Anopheles from across the world have been sequenced and assembled, including some of the most efficient malaria vectors. To facilitate comparative analyses, species were chosen to occupy progressively greater phylogenetic distances from the primary African vector species pair Anopheles gambiae s.s. and Anopheles coluzzii (formerly the S and M forms of Anopheles gambiae [3]), overall representing 100 million years of evolution. Why is this project important when the genome of A. gambiae (PEST strain) was fully sequenced and near-fully assembled over a decade ago [4]? The answer becomes clear when comparing the newly sequenced genomes to those of the existing PEST assembly. A. gambiae is a species complex of morphologically identical, relatively closely-related species. Genome sequence data from each of the five members of the A. gambiae complex in the project align well to the PEST assembly (with >85% alignment success). However, this concordance drops dramatically once sequences of the more distantly related species are considered; down to <15% for the major Central/South American malaria vectors Anopheles albimanus and Anopheles darlingi [1]. In other words, while major malaria vectors are often not closely related, a single reference genome provides a poor general template. Indeed, even for species within the A. gambiae complex, and despite gaps in sequence coverage, inevitable in newly sequenced genomes, alignment of additional sequenced specimens to their species-specific assemblies was significantly better in every case than to the fully-assembled PEST reference [2].

The first paper by Neafsey, Waterhouse et al. [1] presents an in-depth description of the methodologies involved in the project, including a very useful glossary of the many software tools applied (pages 123-4 of the supplementary materials). Assembly qualities were evaluated by searching each reference genome and paired RNA transcripts for almost 3000 evolutionary conserved single-copy orthologous genes from Drosophila, that is, those which should be present in all of the genomes. Assembly qualities generally appear impressive and surprisingly consistent, given the highly variable number and length of the sequence scaffolds. Variation in contiguity of assemblies was highlighted however when analysing cuticular genes, which comprise 2% of protein coding genes and tend to occur in clusters. All new assemblies show fewer genes per cluster than the fully assembled A. gambiae PEST genome, which is most likely a result of assembly difficulties for highly similar genes in clusters, toward the end of scaffolds [1]. This highlights an easily overlooked issue with genome sequence data now obtained relatively quickly and inexpensively. Meaningful comparative analyses require extensive work on assembly and gene annotations, and rigorous quality checks, of the kind performed by the 16 genomes team, other major sequencing consortia (http://www.malariagen.net/projects/vector/ag1000g), and the community as a whole. To this end the 16 genomes consortium are to be congratulated for a release policy which has allowed substantial deposition of data in VectorBase far in advance of publication.

Intriguing insights emerge from comparing variation across the Anopheles species sequenced to results from the Drosophila 12 genomes project [5]. For example, there was a five-fold higher rate of gene gain/loss among Anopheles than among Drosophila species, along with higher rates of intron loss, and a lack of the constraining influence of the codon usage bias that is evident in Drosophila. All hint at flexibility in Anopheles which might be a key to the multiple occurrences of adaptation to human hosts. A distinct evolutionary profile of the X chromosome is highlighted by elevated rates of gene gain and loss relative to autosomes [1] and comes to the fore in the second paper. Fontaine, Pease et al. [2] conduct an in-depth analysis of species relationships and introgression (inter-specific gene flow) within the A. gambiae complex. The true species tree within the complex has been unresolved for many years but compelling evidence now shows that the major vector Anopheles arabiensis appears closely related to A. gambiae and A. coluzzii throughout most of the genome as a result of extensive introgression. In fact the true species tree is reconstructed against a vast genomic majority rule, and
is recovered most accurately from a section of the X chromosome retardant to introgression. Presumably therein lie the key genes maintaining reproductive isolation. These results beautifully advertise the power of phylogenomics, not via obtaining an overwhelming amount of data from which an average consensus can be obtained, but rather by comparative analysis across species and different parts of the genome. Some long-held views of *A. gambiae* history also appear to have been overturned, notably the timing of speciation in relation to the expansion of human agriculture (which it now seems to pre-date) and the transfer of a huge, ecologically important chromosomal inversion from *A. arabiensis* to *A. gambiae* (the reverse now seems probable).

The ability of *Anopheles* genomes to accept and incorporate large sections of autosomes, even if highly distinct, without loss of species integrity, could be a key to successful adaptation [6]. Such flexibility may result from interplay of properties of the genomes themselves as suggested [1], and the strong selective pressures incurred by adaptation to human hosts. The importance of genetic transfer long after species split will be a familiar story to plant and microbial geneticists, but was long thought rare in animals [7]. However, results from both the *Anopheles* 16 genomes consortium and the *Heliconius* genome consortium [8] suggest that interspecific hybridization plays a major role in adaptive phenotypic change. A chilling difference is that whilst butterflies exchange genetic variants controlling their often beautiful colour patterns, *Anopheles* species may transfer adaptive variation which enhances their efficiency as malaria vectors.

### References