

Single mosquito metatranscriptomics identifies vectors, emerging pathogens and reservoirs in one assay

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Abstract

Mosquitoes are major infectious disease-carrying vectors. Assessment of current and future risks associated with the mosquito population requires knowledge of the full repertoire of pathogens they carry, including novel viruses, as well as their blood meal sources. Unbiased metatranscriptomic sequencing of individual mosquitoes offers a straightforward, rapid and quantitative means to acquire this information. Here, we profile 148 diverse wild-caught mosquitoes collected in California and detect sequences from eukaryotes, prokaryotes, 24 known and 46 novel viral species. Importantly, sequencing individuals greatly enhanced the value of the biological information obtained. It allowed us to a) speciate host mosquito, b) compute the prevalence of each microbe and recognize a high frequency of viral co-infections, c) associate animal pathogens with specific blood meal sources, and d) apply simple co-occurrence methods to recover previously undetected components of highly prevalent segmented viruses. In the context of emerging diseases, where knowledge about vectors, pathogens, and reservoirs is lacking, the approaches described here can provide actionable information for public health surveillance and intervention decisions. **(167 words)**

Introduction

Mosquitoes are known to carry more than 20 different eukaryotic, prokaryotic, and viral agents that are pathogenic to humans (WHO, 2017). Infections by these mosquito-borne pathogens account for over half a million human deaths per year, millions of disability-adjusted life years (GBD 2017 Causes of Death Collaborators, 2018; GBD 2017 DALYs and HALE Collaborators, 2018; GBD 2017 Disease and Injury Incidence and Prevalence Collaborators, 2018), and periodic die-offs of economically important domesticated animals (Nonito Pagès and Lee W. Cohnstaedt, 2018). Moreover, recent studies of global patterns of urbanization and warming, as well as the possibility of mosquito transport via long-range atmospheric wind patterns point to an increasing probability of a global expansion of mosquito habitat and a potential concomitant rise in mosquito-borne diseases within the next 2-3 decades (Huestis et al., 2019; Kraemer et al., 2019). While mosquito control has played a major role in eliminating transmission of these diseases in many parts of the world, costs and resources associated with basic control measures, combined with emerging pesticide resistance, pose a growing challenge in maintaining these gains (Wilson et al., 2020).

Female mosquitoes take up blood meals from humans and diverse animals in their environment and serve as a major source of trans-species introductions of infectious microbes. For well-studied mosquito-borne human pathogens such as West Nile virus, an understanding of the transmission dynamics between animal reservoir, mosquito vector, and human hosts has been essential for public health monitoring and intervention (Hofmeister, 2011). In contrast, transmission dynamics are less clear

for emerging microbes with pathogenic potential. Metagenomic sequencing of individual mosquitoes provides a means to comprehensively identify mosquito species, the pathogens they carry and the animal hosts that define a transmission cycle with a single assay.

We also lack a comprehensive understanding of the composition of the endogenous mosquito microbiota, which has been suggested to impact the acquisition, maintenance, and transmission of pathogenic mosquito-borne microbes. For example, *Wolbachia*, a highly prevalent bacterial endosymbiont of insects (Werren et al., 2008) has been shown to inhibit replication of various mosquito-borne, human-pathogenic viruses when introduced into susceptible mosquitoes (Moreira et al., 2009). These observations have led to the development of *Wolbachia*-based mosquito control programs for *Aedes aegypti* mosquitoes, which vector yellow fever virus, dengue virus, Zika virus, and chikungunya virus. Experimental releases of *Aedes aegypti* mosquitoes transinfected with *Wolbachia* have resulted in a significant reduction in the incidence of dengue virus infections in local human populations. Laboratory-based studies have identified additional endogenous mosquito microbes, such as midgut bacteria and several insect-specific flaviviruses. Greater knowledge of these endogenous microbes could inform their potential use in interfering with mosquito acquisition of and competence to transmit pathogenic *Plasmodium* species and human flaviviruses, respectively. Quantitative analysis of the composition of endogenous microbes and the viruses in individual mosquitoes would be needed to establish a role for these agents in naturally occurring infections and/or transmission of known human pathogens.

Several recent, unbiased metagenomic analyses of batches of mosquito pools collected around the world have begun to address these issues (Atoni et al., 2018; Fauver et al., 2016; Frey et al., 2016; Li et al., 2015; Moreira et al., 2009; Pettersson et al., 2019; Sadeghi et al., 2018; Shi et al., 2019, 2015, 2017, 2016; Xia et al., 2018; Xiao et al., 2018a, 2018b) (reviewed in (Atoni et al., 2019; Xiao et al., 2018b)). These studies, which have primarily focused on analysis of viruses, have expanded our understanding of the breadth of viral diversity present in mosquito populations worldwide. However, they have not provided key epidemiologic information needed to direct intervention. This includes the measurement of viral prevalence within mosquito populations, their potential reservoir sources, or the impact that additional bacterial and eukaryotic microbes carried by mosquitoes might have on virus carriage, transmission, and pathogenesis.

Single mosquito analyses are required to link blood meal sources, endogenous microbes, and co-occurring pathogens. A handful of small-scale studies have demonstrated that it is possible to identify divergent viruses and evidence of other microbes in single mosquitoes via metagenomic next-generation sequencing (Bigot et al., 2018; Chandler et al., 2015; Shi et al., 2019). Here, we analyzed the metatranscriptomes of 148 individual mosquitoes collected in California, USA. We characterized the composition of their co-infecting microbes, quantified the prevalence and load of detectable viruses and selected bacterial and eukaryotic microbes, and identified blood meal sources and their associated pathogens. Crucially, sequencing a large number of individuals allowed for simple co-occurrence analyses that extended the sensitivity to detect missing or as-yet unidentified viral genome segments with no recognizable homology to previously described sequences. Our findings demonstrate how large-scale single mosquito metatranscriptomics can define both the mosquito's complex microbiota, including mosquito-borne pathogens, and its blood meal sources, thus contributing critical epidemiological information needed to control transmission.

Results

Mosquito host speciation by comparative whole transcriptome analysis

Adult *Aedes*, *Culex*, and *Culiseta* mosquito species circulating in California in late fall of 2017 were collected to acquire a diverse and representative set of 148 mosquitoes for metatranscriptomic next generation sequencing (mNGS) analysis. We targeted collections across a variety of habitats within 5 geographically distinct counties in Northern and Southern California (Supplemental Figure 1). Visual mosquito species identification was performed at the time of collection (Materials and Methods, results are summarized in Table S1). Primarily female mosquitoes were included to enrich for blood-feeding members of the population responsible for transmission of animal and human diseases. Total RNA extracted from each mosquito was used as the input template for mNGS to capture both polyadenylated and non-polyadenylated host, viral, prokaryotic, and eukaryotic RNAs (Materials and Methods; per mosquito sequence yields are summarized in Table S2).

Given the important role of accurate identification of mosquito species for understanding geospatial mosquito circulation and vector-pathogen interactions, and the potential for human error in visual inspection, we investigated if single mosquito mNGS could provide a complementary, unbiased molecular method for identifying mosquito species. Because complete genome sequences were not available for all mosquito species identified visually in this set, we applied a reference-free, kmer-based approach (Harris, 2018) to compute pairwise genetic distances between the complete metatranscriptomes acquired for each of the 148 mosquitoes. Samples were grouped using hierarchical clustering and the most common visually identified species within each group was taken as a consensus species call for that group (Figure 1 and see Supplemental Figure 2 for detailed alignment of visual calls with the clustered genetic distance matrix, and Supplemental Data File 1 for underlying snp distance data). These molecular groupings of mosquito genera and species agreed the visual calls for 95% of the specimens ($n=140/147$, one sample had no visual identification). The discordant calls occurred in 2 contexts reported to present challenges to morphology-based speciation: 1) within the *Culex* genus in which genetic hybridization among species members has been documented and reported to confound accurate morphological speciation in California (Cornel et al., 2003, 2003; Kothera et al., 2012; McAbee et al., 2008); and 2) between samples belonging to the *Culex* and *Culiseta* genera that share some overlap in morphology, and require detection of features (perspiracular bristles and subcostal wing vein bristles) that can be lost or damaged during trapping and handling (Darsie and Ward, 2016). Thus, we used the transcriptome-based species calls for this study. There is additional within-species structure visible on the hierarchical clustering tree that, for *Aedes aegypti* and *Culex erythrothorax*, coincides with geographic structure, raising the possibility that molecular methods may also provide insight into the distribution of subspecies as well (Figure 1). Taken together, these data show that comparative transcriptome analysis of single mosquito mNGS data can provide critical information regarding the identity and diversity of circulating mosquitoes.

Comprehensive and quantitative analysis of non-host sequences detected in single mosquitoes

To understand the distribution of microbial species within the microbial cargo of mosquitoes, we first examined the overall proportion of non-host reads assembled into contigs that could be assigned to viral, bacterial, and eukaryotic taxa. A detailed overview of the analysis we applied to identify, assemble, classify, and quantify all the non-host contigs and their associated read counts is provided in Materials and Methods (summarized in Supplemental Figure 3). Details of the per mosquito breakdown of non-host read assignment across high level taxonomic categories are provided in Supplemental Figure 4 (underlying data for Supplemental Figure 4 can be found in Supplemental Data file 3). Figure 2 provides a quantitative treemap overview of how the assembled non-host reads mapped across the viral, prokaryotic, and eukaryotic taxa (see Supplemental Data File 2 for data underlying Figure 2 and Supplemental Figure 5, a higher resolution treemap overview). In sum, we were able to classify, to at least kingdom level, 77% of the 13 million non-host reads that assembled into contigs with more than two reads.

Diverse known and novel RNA virus taxa dominate the mosquito microbiota.

We found that the vast majority of the non-host reads of mosquitoes assembled into contigs corresponding to complete viral genomes (11 million reads of 13 million total non-reads; Figure 2, all blocks in the treemap annotated with suffix “-viridae”). Positive sense single stranded RNA viruses made up the most abundant class of detected viruses (7.4 million reads of the 11 million viral reads; Figure 2 blocks labeled *Solemoviridae*, *Luteoviridae*, *Tombusviridae*, *Narnaviridae*, *Flaviviridae*, *Virgaviridae*, and *Filoviridae*), negative sense single stranded RNA viruses made up the next most abundant virus category (2.25 million reads of the 11 million viral reads; Figure 2 blocks labeled *Peribunayviridae*, *Phasmaviridae*, *Phenuiviridae*, *Orthomyxoviridae*, *Chuviridae*, *Rhabdoviridae*, and *Ximnoviridae*), and double-stranded RNA viruses formed the third most abundant virus category (0.94 million reads of the 11 million reads; Figure 2 blocks labeled *Chrysoviridae*, *Totiviridae*, *Partitiviridae*, and *Reoviridae*). In many cases, multiple independent isolates of complete viral genomes were recovered across the individual mosquito specimens. In all, a total of 70 distinct viral taxa were recovered, 46 of which correspond to distinctly divergent novel viruses (Table 1). Intriguingly, only 10 of the 24 previously described viral taxa have been previously recovered from mosquitoes in California (Table 1, rows highlighted in gray (Chandler et al., 2015; Sadeghi et al., 2018)). Although a number of the known and novel viral species we detected corresponded to additional viral species from families previously thought to only infect plants and fungi, (e.g., the *Chrysoviridae*, *Totiviridae*, *Luteoviridae* and *Solemoviridae*, Table 1), emerging evidence from mosquito and broader insect metatranscriptomics has indicated that these viral taxa are tightly associated with, if not actually infecting, mosquitoes and other insects (Shi et al., 2017, 2016; Xiao et al., 2018b).

The balance of additional reads that could be assigned to viral taxa corresponded to reads assembled into contigs that were clearly viral in origin but incomplete by either not associated with an RNA-dependent polymerase (0.37 million reads, Figure 2, light gray block labeled “uncurated viruses”) or associated with contigs aligning to viral taxa that were detected at levels too low to visualize on the treemap. This latter set of low abundance viral taxa corresponded to several types of DNA viruses, such as nucleocytoplasmic large DNA viruses, members of the *Polydnoviridae*, *Alphaculovirus*, *Nudiviridae*, and *Circovirus*-like sequences, and phages (data not shown). Some of these low abundance viral taxa reflect *bona fide* infections, while others are likely the result of indirect infections. For example, 6

distinct types of *Botourmiaviridae*, a family of viruses primarily known to infect fungi, were all detected at very low levels in a single mosquito that was also found to be co-infected with a more likely Botourmiaviridae host species, the ergot fungus, *Claviceps*.

***Trypanosomatidae* and vertebrate species are major constituents of the eukaryotic taxa.**

An additional 2.2 million of the 13 million non-host reads assembled into contigs that mapped to non-viral taxa. Just under 1 million reads could be assigned to eukaryotic taxa (0.9 million reads total, Figure 2, bottom left row of boxes). Members of *Trypanosomatidae* comprised 50% of these reads. Slightly more than half (0.25 million reads) of the *Trypanosomatidae* reads correspond to the subfamily *Leishmaniinae*, that encompasses multiple species known to infect insects and vertebrates. The second most abundant group of eukaryotes detected in the dataset were *Bilateria* (animals), with 0.20 million reads made up of mammals (*Boreoeutheria*, 73,000 reads) and birds (*Aves*, 51,000 reads), followed by invertebrates (*Ecdysozoa*, 36,000 reads (not shown); see Supplemental Figure 5 for higher resolution details of these and other notable lower abundance eukaryotic taxa detected). The reads derived from vertebrate taxa almost certainly belong to blood meal hosts, which we investigate in detail below. Fungal and plant contigs, made up the remainder of the eukaryotic reads we captured from individual mosquito sequencing, with 79,000 and 62,000 total reads, respectively.

***Wolbachia* species make up the majority of prokaryotic taxa.**

Prokaryotic contigs encompassed 0.7 million non-host reads. Among the prokaryotic taxa detected, *Wolbachia*, a known endosymbiont of *Culex quinquefasciatus* (Werren et al., 2008), comprised most of the reads (0.22 million reads, Figure 2, bottom central row of brown-hued blocks). Various other bacterial taxa were detected at lower abundance; i.e. members of Alphaproteobacteria, Gammaproteobacteria, Terrabacteria group, and Spirochaetes (*Spiroplasma culicis* (73,000 reads), a bacterial species previously detected in *Culex* mosquitoes (Cechová et al., 2004; Duguma et al., 2019), makes up 68% of Spirochaete reads). A higher resolution overview of the lowest common ancestor (LCA) species we could assign within each of these 4 broad categories is provided in Supplemental Figure 5 and Supplemental Data File 2. Interestingly, these results largely agreed with data obtained for the *Culex* and *Aedes* species in prior sequencing studies involving more directed capture of prokaryotic and eukaryotic taxa via 16S rRNA metabarcoding of mosquitoes collected in Thailand (Thongsripong et al., 2018).

Ambiguous and metagenomic “dark matter” sequences are present.

A significant portion of the non-host reads assembled into contigs with sequences that were taxonomically ambiguous. Approximately 0.5 million reads assembled into contigs with a lowest common ancestor (LCA) assigned to the taxonomy nodes of “root” or “cellular organisms” (Figure. 2, unlabeled light gray box). A much larger fraction of non-host reads -- approximately 4.4 million reads -- corresponded to metagenomic “dark matter”, i.e. contigs without any recognizable sequence homology to previously published sequences. Contig co-occurrence analysis across the individual mosquito sequence results (see main text below) allowed us to identify additional viral contigs from this set of

contigs, contributing 0.4 million reads to the total tally of detectable viral reads in the mosquito microbiota.

Together these data establish the utility of our comprehensive single mosquito mNGS analyses to define the composition and diversity of the mosquito metatranscriptome. The sensitivity of our analysis reveals endogenous constituents of the mosquito microbiome, the source of their blood meals, and the potential human and animal pathogens they carry, and even viruses that selectively infect mosquito-associated fungi.

Identifying constituents of the mosquito microbiota

To define components of the mosquito microbiota and investigate the potential variation between individuals, we next analyzed the distribution of the viral, prokaryotic, and microbial eukaryotic taxa detectable across individual mosquitoes. All confidently called contigs for microbial taxa detected within each mosquito were compiled, and the fraction of non-host reads aligning to each contig was computed to estimate the composition and proportions of microbial agents detectable within each mosquito. Figure 3 displays those agents that were detectable at a level above 1% of non-host reads plotted as bars for each individual mosquito analyzed. The data for read counts per taxon per sample and detected viral taxa can be found in Supplemental Data Files 4 and 5, respectively. Notably, given the sensitivity of our data and analytical pipeline (see Materials and Methods), we could confidently call contigs even when supported by <1% of the non-host reads (i.e., <<0.02% of total reads). These are plotted as dark circle symbols above the x-axis coordinate for the central *Wolbachia* panel of Figure 3, and grey bars in the *Trypanosomatidae*, *Apicomplexa*, and *Nemotoda* panels at the base of Figure 3 (underlying data for these selected bacterial and eukaryotic microbial taxa are provided in Supplemental Data Files 6 and 7, respectively). The data reveal unprecedented insight into the heterogeneity in the microbiota associated with individuals, between mosquito species and collection sites.

Viral diversity and prevalence measured in single mosquitoes

Given their predominance and potential relevance to human disease, we first examined the 70 unique viral taxa on a single mosquito basis (Table 1, Supplemental Data File 4). Of these, only 24 were closely related or identical to previously identified mosquito viruses. The remaining 46 viral genomes shared less than 85% amino acid sequence identity to any publicly available viral sequences. Despite this divergence, family-level sequence conservation of genomic features allowed us to confirm complete genome sequence recovery. For example, conserved sequences were identified at the 5' and 3' ends of bunyavirus segments in a novel peribunya-like virus (Udune virus, Table 1) with 28% to 76% amino acid identity to its closest relative (Supplemental Figure 6 and Supplemental Data File 8). Only a single mosquito harbored this virus, and at a relatively low abundance of reads (0.02% of total reads). Combining this approach with the co-occurrence investigation detailed below (Figure 6 and main text), we were able to identify and assign the 45 additional novel genomes to specific viral taxa. Thus, single mosquito analysis provides a highly sensitive approach to detect new and divergent viral species circulating in mosquito populations, even when present at low prevalence or abundance.

Single mosquito analysis also shed light on the variability in the composition and number of viral reads both within and across mosquito species (Figure 3, top panel), and their corresponding collection sites (indicated by colored bar plot at the bottom of the plots). Importantly, among single mosquitoes,

co-infections predominated, with 88% of mosquitoes harboring 2 or more (median 3) distinct viral taxa (Figure 4A, see Supplemental Data File 9 for underlying data). Focused analysis of viral species within single mosquitoes provides the opportunity to examine the proportion of viruses within each of these co-infections, which in turn can inform and extend our understanding of the distribution patterns of known and emerging novel viruses within the mosquito population, and the associated co-infecting viruses. Figure 3, shows the wide range in the number and type of viruses that are detected across individual mosquitoes. For instance, several *Culex* mosquito species stand out as outliers harboring only a single viral species encompassing >50% of the non-host reads assembled into contigs in that mosquito (Figure 3, top panel: *Iflaviridae* species [dark green bars] in *Culex tarsalis*; *Tombusviridae* and *Virgaviridae* species [light blue bars] in *Culex erythrothorax*). At the other end of the spectrum are multiple examples of individual mosquitoes that do not stand out with regard to viral read abundance, yet still harbor a mixture of 4 or more viruses (Figure 3, top panel - see especially *Culex tarsalis* and *Culiseta* species plots). Other viruses that are detected broadly across diverse mosquito species (Figure 3, top panel, see the *Solemoviridae/Luteoviridae* [yellow bars], *Narnaviridae* [blue bars], *Virgaviridae* [light blue bars], and *Dicistroviridae/Iflaviridae* [dark green bars]). Interestingly in some mosquito species, these viruses are the predominant proportion of the non-host reads assembled into viral contigs, while in other mosquito species where these viruses are detected, they make up only a minor fraction. The opposite pattern of virus distribution where viral species are restricted to a single mosquito species, for example the *Partitiviridae* and *Reoviridae* among the *Culiseta* species (Figure 3, top panel, right edge of plot, dark purple bars). These distinct patterns of viral distribution point to potentially testable hypotheses as to their causes, such as mosquito species susceptibility or competence to vector a virus, the potential pathogenicity of a given virus (or mixture of viruses), or factors in the environment, such as food sources or weather. Regardless of the ultimate source of this variability, such insights are only possible by analyzing mosquitoes individually rather than in bulk.

This variation is particularly relevant when we consider that viral abundance is often calculated based on bulk mosquito sequencing, which does not provide information about the prevalence or heterogeneity in abundance of a virus across the mosquito population. Importantly, we find that the average abundance of a virus (*i.e.* the average number of reads across a set of mosquitoes) is not necessarily predictive of the prevalence of that virus (*i.e.* the number of mosquitoes in which it occurs). For example, *Culex narnavirus* 1 and *Culex pipiens*-associated Tunisia viruses were found at similar abundance in *Culex erythrothorax* mosquitoes obtained from the same collection site in West Valley; however, the latter was 3 times more prevalent (30% vs 90%, Figure 4B; see Supplemental Data File 10 for underlying data). A more global view of viral diversity and prevalence across mosquito species is shown in (Figure 4C and summarized in Supplemental Data File 11) which plots the fraction of individuals infected for each mosquito species and each virus detected in our study. This quantitative and comprehensive analysis of the prevalence of mosquito-borne viruses would not be possible without single mosquito sequencing, yet provides critical epidemiological information needed to manage the transmission of mosquito-borne viruses. For example, the sampled mosquito genera (*Culex*, *Aedes*, and *Culiseta*) have distinct viromes, with only four viruses shared across genus boundaries and even then, only Merida virus (-ssRNA) and *Culex* iflavi-like virus 4 (+ssRNA) are shared by *Aedes* and *Culex* mosquitoes. Within each genus, viruses appear to be largely unique to species, though some overlap is detectable (Figure 4C), potentially reflecting greater similarities in ecology and physiology (Longdon et al., 2014) that enable an easier flow of viruses between populations.

Exploring the impact of *Wolbachia* endosymbionts

We restricted our single mosquito analysis of detectable prokaryotes to *Wolbachia* given its abundance and evidence suggesting that as an endosymbiont it could impact the microbiota of its mosquito hosts. *Wolbachia* was detected in 32 mosquitoes belonging to *Culex quinquefasciatus*, *Culex pipiens*, and *Aedes albopictus* species (Figure 3, middle panel, black bars and circle symbols; see Supplemental Data File 6 for underlying data). These observations are consistent with previous reports of wild-caught mosquito species that are naturally infected with *Wolbachia* (Kittayapong et al., 2000; Rasgon and Scott, 2004). Among these 3 species, *Wolbachia* was detected in all or nearly all of the mosquitoes. Thus, it was not possible to draw definitive conclusions regarding whether the presence or absence of *Wolbachia* influenced the composition of detectable co-occurring viral taxa among these mosquitoes. However, the fraction of non-host reads assigned to *Wolbachia* per mosquito varied dramatically, from <1% to 74% (Figure 3, middle panel, black circles and black bar plots, respectively), and revealed interesting trends that would require further validation. For example, for *Ae. albopictus*, individuals with higher levels of detectable *Wolbachia* (Figure 3, central panel: samples with black bars) exhibited lower numbers of reads from single stranded positive sense RNA viruses (*Solemonviridae* and *Luteoviridae*) than individuals with a lower percentage of *Wolbachia* reads (Figure 3, central panel: samples with black circles). Similarly, higher levels of *Wolbachia* in *Culex pipiens* mosquitoes also generally correlated with lower viral abundance. Although not statistically significant, given the low sample numbers and lack of *Wolbachia* positive and negative individuals, these data again demonstrate the potential utility of sequencing individuals.

Prevalence of eukaryotic microbes and pathogens in single mosquitoes

Although we detected fungi, plants and other eukaryotes in our analyses (Figure 2), we focus here on three potentially human pathogenic species: *Trypanosomatidae*, which was the most abundant eukaryotic taxon detected and contains established pathogens of both humans and birds; *Apicomplexa*, which encompasses the causative agents of human and avian malaria; and *Nematoda*, which contain filarial species that cause heartworm in canines and filarial diseases in humans (Supplemental Data File 7).

Twelve mosquitoes (8%) were found to harbor *Trypanosomatidae* taxa (Figure 3, bottom panel). We detected sequences corresponding to monoxenous (e.g. *Crithidia* and species), dixenous (*Trypanosoma*, *Leishmania* species), as well as the more recently described *Paratrypanosoma confusum* species. Of the *Trypanosomatidae*-positive mosquitoes, eight were *Culex erythrothorax* mosquitoes, while the remaining four were *Culex pipiens* and 2 *Culex tarsalis* (Figure 3, bottom panel). Notably, all were collected from the same trap site in Alameda County, albeit at different times, providing insight into, in this case, a limited distribution and potential prevalence of *Trypanosomatidae* within the mosquito population.

We investigated the distribution of the *Apicomplexa* contigs and reads, as this phylum encompasses the *Plasmodium* genus, which includes several pathogenic species that cause avian and human malaria. Within our single mosquito dataset, we identified 8 mosquitoes with *Apicomplexa* contigs (Figure 3, bottom panel). These corresponded to 3 *Aedes aegypti* mosquitoes and 1 *Culex quinquefasciatus* mosquito, both collected in San Diego, and 2 *Culex erythrothorax* mosquitoes, 1 *Culex pipiens* mosquito, and 1 *Culex tarsalis* mosquito collected in Alameda County. Only the *Culex quinquefasciatus* mosquito

harbored *Apicomplexa* reads at a level above 1% of total non-host reads. Interestingly, this mosquito harbored *Wolbachia*, but no viruses could be detected.

Finally, we examined taxa falling under *Nematoda*, a phylum that encompasses a diverse set of more than 50 filarial parasites of humans and animals. Here, we saw evidence of *Nematoda* carriage in 3 *Culex* mosquitoes: 2 *Culex tarsalis* and 1 *Culex pipiens* (Figure 3, bottom panel). Two of these mosquitoes were collected in Alameda County and showed very low levels *Nematoda* (< 1% of non-host reads, Figure 3, bottom panel, dark gray square symbols). In the third mosquito, a *Culex tarsalis* collected in Coachella Valley, the *Nematoda* made up 2% of the non-host reads (Figure 3, bottom panel gray bar with red outline).

Together these data reveal the diversity and prevalence of microbial species harbored within single mosquitoes and establish the comprehensive nature and sensitivity of single mosquito metagenomic analysis.

Blood meals and associated microbes

As vectors, mosquitoes transfer the pathogenic microbes they carry from one animal to another as they feed. Identifying the sources of these blood meals can provide critical information regarding the animal reservoir of these vector-pathogens and the paths of transmission. Therefore, we next investigated the possibility of identifying the blood meal host directly from mNGS. We restricted this analysis to the 60 mosquitoes from Alameda County, as they were selected for visible blood-engorgement. For 45 of the 60 mosquitoes, there was at least one contig with an LCA assignment to the phylum *Vertebrata* (range = 1-11 contigs, with 4-12,171 supporting reads). To assign a blood meal host for each of these mosquitoes, we compiled their corresponding *Vertebrata* contigs and selected the lowest taxonomic group consistent with those contigs. For all samples, the blood meal call fell into one of five broad categories (Figure 5, underlying data and detailed summary provided in Supplemental Files 12 and 14, respectively): even-toed ungulates (*Pecora*), birds (*Aves*), carnivores (*Carnivora*), rodents (*Rodentia*), and rabbits (*Leporidae*). For 10 samples, we were able to identify the genomic source at the species level, including rabbit (*Oryctolagus cuniculus*), mallard duck (*Anas platyrhynchos*), and raccoon (*Procyon lotor*).

The potential blood meal sources identified were broadly consistent with the habitats where the mosquitoes were collected. For the 25 samples collected in or near the marshlands of Coyote Hills Regional Park, we compare our calls to the wildlife observations in iNaturalist, a citizen science project for mapping and sharing observations of biodiversity. iNaturalist reports observations consistent with all five categories, including various species of squirrel, rabbit, raccoon, muskrat, and mule deer. The mosquitoes with blood meals in *Pecora* are likely feeding on mule deer, as no other ungulate commonly resides in that marsh (iNaturalist, 2020).

We also investigated whether bloodborne pathogens of the blood meal source were detectable. We performed a hypergeometric test for association between each blood meal category and each microbial taxon (see Materials and Methods and Supplemental Data File 13). The only statistically significant association ($p = 0.0005$, Bonferroni corrected) was between *Pecora* and *Anaplasma*, an intracellular erythroparasite transmitted by ticks. *Anaplasma* was detected in 11 of the 20 samples with *Pecora*. This striking co-occurrence suggests a possible burden of anaplasmosis in the local deer population. Additionally, we detected evidence for three other bloodborne pathogens which, because of the small number of observations, could not pass the threshold of statistical significance. These included an orbivirus closely related to those known to infect deer, a *Trypanosoma* species previously found in birds,

and the apicomplexans *Plasmodium* and *Eimeria* from species known to infect birds (See Materials and Methods). The likely hosts of these pathogens were also concordant with the blood meal calls. Thus, sensitive and comprehensive metagenomic analysis of single mosquitoes not only provides information as to paths of transmission, it also provides a tool to detect emerging pathogens within animal communities in their environments.

Recovery and assignment of previously unrecognizable viral genome segments and species within the orthomyxovirus family

Although many new viruses can be identified in bulk samples, the majority of these are identified only via their conserved RNA-dependent RNA polymerase (RdRp). Recovering complete genomes for segmented viruses from bulk samples is challenging, as genes that are not highly conserved may be unrecognizable by sequence homology. Moreover, the assignment of putative segments to a single genome can be confounded if the pooled libraries are derived from mosquitoes with multiple infections of related segmented viruses.

By sequencing many individual mosquitoes, we can exploit the fact that all segments of a segmented virus will co-occur in the samples where that virus is present and be absent in samples where the virus is absent. Applying these criteria to our data analysis should enable the identification of previously unidentified viral genome segments. To do this, we first grouped all contigs that were longer than 500 nucleotides into clusters of highly homologous contigs, then grouped these clusters by co-occurrence across all of the 148 individual mosquitoes sampled (Figure 6A). Importantly, this required only the sequence information from the study, without using any external reference. We then scanned each cluster for sequences containing a viral RdRp domain (see Materials and Methods). For each RdRp cluster, we consider any other contig cluster whose sample group overlaps the set of samples in the viral RdRp cluster above a threshold of 80% as a putative segment of the corresponding virus. A cluster-by-sample heatmap for all segments co-occurring with RdRps resulted in 27 candidate complete genomes for segmented viruses (Supplemental Figure 7, underlying data provided in Supplemental Data Files 15A and 15B). For the 79 of the 96 putative segments recognizable by homology to published sequences (colored in black), these groupings into genomes were accurate. This supports the notion that the remaining 17 putative segments (colored in red), which lack homology to any known sequences at either nucleotide or amino acid level, may indeed be part of viral genomes. Combined, these putative segments represented 8% of the metagenomic “dark matter” portion of the reads in the study.

Our co-occurrence analysis enabled the discovery of new viral segments and new viral species within the segmented *Orthomyxoviridae* family (Figure 6B; see Supplemental File S16 for underlying data). Orthomyxoviruses are segmented viruses (ranging from 6 to 8 segments) including influenza viruses, isaviruses, thogotoviruses, and quaranjaviruses that infect a range of vertebrate and arthropod species. Quaranjaviruses are largely found in arthropods, and in this study, we identified four quaranjaviruses, two of which were previously observed in mosquitoes collected outside California (Wuhan Mosquito Virus 6 (WMV6) (Li et al., 2015; Shi et al., 2017) and Guadeloupe mosquito quaranja-like virus 1 (GMQV1) (Shi et al., 2019)) and two, which we have named Ūsinis virus and Astopletus virus, were previously unknown.

Thus, for WMV6 and GMQV1 detected here, we observed all of the previously identified segments (Li et al., 2015; Shi et al., 2017), as well as two additional segments (which we named hypothetical 2 and hypothetical 3) for WMV6 and five for GMQV1 (PA, gp64, hypothetical, hypothetical 2, hypothetical 3)

(Supplemental Figure 8A and 8B). We confirmed the existence of the 2 putative segments for WMV6 by assembling homologous segments from reads in two previously published datasets describing this virus. For GMQV1, we were able to find reads in NCBI's short read archive entries that are similar at the amino acid level to putative protein products of the two new segments; however, there was not sufficient coverage to reconstruct whole segments. Furthermore, phylogenetic trees constructed separately for each of the eight segments of WMV6 have similar topologies (See tanglegram, Figure 7; see Supplemental Data File 17 for underlying data), suggesting that the two new putative segments have evolved in conjunction with the previous six, bringing the total number of segments for each genome to eight.

For the two quaranjaviruses discovered in this study, *Usinis virus* and *Astopletus virus*, the co-occurrence analysis also produced 8 segments, 5 and 4 of which, respectively, were recognizable by alignment to NCBI reference sequences. The hypothetical 2 and hypothetical 3 segments we identified from this set of four quaranjavirus genomes are too diverged from one another to align via BLASTx, but they do share cardinal features such as sequence length, ORF length, and predicted transmembrane domains (Supplemental Figure 8A-D). Intriguingly, this set of four viruses are part of a larger clade of quaranjaviruses (Supplemental Figure 9, Supplemental Data File 18). It is nearly certain that the remaining seven viruses in this clade also have eight segments and quite likely that all quaranjaviruses share this genome organization hinted at in earlier studies (Zeller et al., 1989).

The high rate of viral co-infections detected among the single mosquitoes we analyzed (Figure 4A) indicated a concomitant high likelihood that multiple mosquitoes could harbor more than one segmented virus, and potentially confound our co-occurrence analysis. However, the co-occurrence threshold of 0.8 that we applied was sufficient to deconvolve those segments into distinct genomes in all cases but one. There were 15 mosquito samples containing both *Usinis virus*, an orthomyxovirus with eight segments (three of which were unrecognizable by BLASTx) and *Barstukas virus*, a Phasma-like bunyavirus, with one additional sample where only *Barstukas virus* was found (Figure 6B, top two blocks). In this case, we were able to disentangle the genomes of these two viruses using additional genetic information: *Barstukas virus* contains all three segments expected for a bunyavirus (L, GP, and NP), all of which had BLASTx hits to other Phasma-like viruses, while the unrecognizable segments of *Usinis virus* shared features with the other quaranjaviruses in the study (as described above).

Co-occurrence reveals unknown genome segments of *Culex narnavirus 1*

Beyond detection of missing genome segments for known segmented viruses, the co-occurrence analysis also revealed additional genome segments in "dark matter" contig clusters for viruses with genomes previously considered to be non-segmented. A striking example is an 850 nucleotide contig cluster that co-occurred with the *Culex narnavirus 1* RdRp segment in more than 40 mosquitoes collected from diverse locations across California (Supplemental Figure 7, Supplemental Data Files 15A and 15B). Like the RdRp segment, the putative new second segment shares the exceptional feature of ambigrammatic open reading frames (ORFs), i.e. a distinct ORF encoded by the reverse complementary RNA strand (Supplemental Figure 8E). The phylogenetic tree topology for the set of 42 putative second segments is similar to the tree for the RdRp segments, suggesting co-inheritance (Supplemental Figure 10A, Supplemental Data File 19). Moreover, we were able to recover nearly identical contigs from previously published mosquito datasets, all of which also contained the *Culex narnavirus 1* RdRp segment. This provides strong evidence that this otherwise unrecognizable sequence is a genuine *Culex*

narnavirus 1 segment, which we refer to here as the “Robin” segment, given it’s consistent, but underappreciated presence.

Since the Narnaviruses were first described in fungi (Hillman and Cai, 2013) and recent studies have shown other eukaryotes can serve as Narnavirus hosts (Charon et al., 2019; Dinan et al., 2019; Göertz et al., 2019; Richaud et al., 2019), we investigated whether this virus co-occurred with a potential non-mosquito host. However, there was no significant co-occurrence with a non-mosquito eukaryotic taxon, or between the abundance of *Culex* narnavirus 1 and abundance of fungi (Supplemental Figure 10B, Supplemental Data File 20). Thus, it is likely that mosquitoes serve as direct hosts of the *Culex* narnavirus 1, whose genomes we show here consist of two, still enigmatic, ambigrammatic RNA segments.

Discussion

We demonstrate how mNGS of single mosquitoes, together with reference-free analyses and public databases, provides—in a single assay—critical and actionable epidemiological information. This includes quantitative information regarding circulating mosquito species, pathogen prevalence, and co-occurrence of diverse known and novel viruses, as well as prokaryotes, eukaryotes, blood meal sources and their potential pathogens. We are able to identify and confirm, using public data, the existence of previously unknown segments of both known and novel viruses, focusing on four quaranjaviruses and *Culex* narnavirus 1 as examples. In the context of an emerging disease, where knowledge about vectors, pathogens, and reservoirs is lacking, the techniques described here can be applied to rapidly provide actionable information for public health surveillance and intervention decisions. While unbiased sequencing of individual mosquitoes is not currently practical or appropriate in all contexts, advances in lab automation and rapidly decreasing costs of mNGS technologies are expected to increase the affordability and practicality of single mosquito sequencing in the near future.

Inferring biology from sequence in the context of an incomplete reference

The power of metatranscriptomic NGS depends on the ability to extract biological information from nucleic acid sequences. For both bulk and single mosquito sequencing studies, the primary link between sequence and biology is provided by public reference databases, and thus the sensitivity of these approaches will depend crucially on the quality and comprehensiveness of those references. In practice, even the largest reference databases, such as nt/nr from NCBI, represent a small portion of the tree of life. Consequently, sequences derived from a sample of environmental or ecological origin, often exhibit only a low percent identity to even the best match in a database. Here, we manage that uncertainty by assigning a sequence to the lowest common ancestor of its best matches in the reference database. However, there is a fundamental limit to the precision of taxonomic identification from an incomplete reference.

An advantage of single mosquito sequencing is that it offers an orthogonal source of information: the ability to recognize nucleic acid sequences detected in many samples even when they have no homology to a reference sequence. This allowed us to associate unrecognizable sequences with viral polymerases, generating hypothetical complete genomes. The strategy of linking contigs that co-occur across samples is utilized in analysis of human and environmental microbiomes, where it is referred to as “metagenomic binning” (Breitwieser et al., 2019; Roumpeka et al., 2017). Using this approach, we identified previously

unknown genome segments establishing that the genome of a large clade of quaranjaviruses (those descended from the common ancestor of WMV6 and GMQV1), like distantly related influenza A and B viruses, consists of 8 segments. We also discovered a second ambigrammatic RNA encoded by the *Culex* narnavirus that in retrospect was identifiable in multiple previously published mosquito datasets. In sum, we pulled 8% of the reads in the metagenomic “dark matter” fraction of our dataset into the light. The putative complete genomes we identified were supported retrospectively by public datasets and can be further validated by biological experiments or approaches such as short RNA sequencing that indicate a host antiviral response (Aguiar et al., 2015; Waldron et al., 2018).

Another advantage of single mosquito sequencing is the ability to supplement, or potentially circumvent, visual species identifications using molecular data. Accurate mosquito species identification is essential for the control of mosquito-borne diseases, as pathogen competence is often limited to a range of species, such as various *Aedes* species for Zika, dengue, and chikungunya viruses, and *Anopheles* species for malaria. Also, the primary mosquito species responsible for vectoring a disease can vary geographically—West Nile virus has been detected in 65 mosquito species, but a narrow range of *Culex* species drives transmission of the virus. Field validation of which mosquito species carry which pathogens in a specific geographic area informs targeted analysis and control of that species (Petersen et al., 2013). Here, though only 3 of the 10 collected mosquito species had complete genome references, it was possible to estimate pairwise SNP distances between samples in a reference-free way and perform an unsupervised clustering. The clusters were 95% concordant with the visual mosquito species calls, and discordant outliers were easy to detect and correct. This approach generalizes to any collection of metatranscriptomes containing multiple representatives of each species. Accurate mosquito identification is essential for selecting the appropriate strategy and materials to control viremic mosquitoes. In a bulk pool of mosquitoes, the microbiota from any miscalled specimens would be blended in with the correctly labeled ones, making it difficult or impossible to deconvolute host species after the fact. By correctly identifying the host range of a known or novel pathogen in a given area, vector control can be appropriately targeted for the prevention of disease.

Distribution of microbes within mosquito populations

Once sequences have been mapped to taxa, it is relatively straightforward to characterize the composition of the microbes within a circulating population of mosquitoes. This information can inform basic research and epidemiologic questions relevant for modeling the dynamics of infectious agents and the efficacy of interventions. A key parameter is the prevalence of a microbe, which cannot be inferred from bulk data. For the 70 viruses in this study, the prevalence ranged from detection in one mosquito (peribunya-like Udune virus) to detection in all 36 *Culex tarsalis* samples in the study (Marma virus).

For some questions, the prevalence data supplied by single mosquito sequencing is helpful for experimental design. For example, in our dataset, *Wolbachia* was either absent or endemic in each mosquito species sampled. Thus, although a trend between the amount of *Wolbachia* relative to viral diversity and load was detectable across samples that harbored *Wolbachia*, it was not possible to detect a statistically significant effect of *Wolbachia* on virome composition or abundance within any species. Nonetheless, our data establish that single mosquito sequencing could address such questions via more extensive, targeted sampling of mosquito populations where *Wolbachia* (or any other agent of interest) is expected to have an intermediate prevalence. This information would be invaluable, as the introduction of *Wolbachia* might be a useful biological agent to suppress viral transmission by mosquitoes (Moreira et al., 2009).

Blood meal sources and xenosurveillance

The identification of blood meal hosts is important for understanding mosquito ecology and controlling mosquito-borne diseases. Early field observations were supplemented by serology, and, more recently, molecular methods based on host DNA. Currently, the most common method of blood meal identification is targeted PCR enrichment of a highly-conserved "barcode" gene, such as mitochondrial cytochrome oxidase I, followed by sequencing (RATNASINGHAM and HEBERT, 2007; Reeves et al., 2018). To monitor specific relationships between mosquito, blood meal, and pathogen, studies have combined visual identification of mosquitoes, DNA barcode identification of blood meal, and targeted PCR or serology for pathogen identification (Batovska et al., 2018; Boothe et al., 2015; Tedrow et al., 2019; Tomazatos et al., 2019). Here, we extend the spectrum of molecular methods, and show that unbiased mNGS of single mosquitoes can identify blood meal hosts, while simultaneously validating the mosquito species and providing an unbiased look at the pathogens. This allows for both reservoir identification, which seeks to identify the unknown host of a known pathogen, and xenosurveillance, which seeks to identify the unknown pathogens of specific vertebrate populations (Grubaugh et al., 2015). For example, in this study we found a high prevalence of the tick-borne pathogen *Anaplasma* in mosquitoes that had likely ingested a blood meal from deer. In one deer-fed mosquito, we found Lobuck virus, a novel orbivirus isolate that belongs to a clade of viruses implicated in a disease of commercially farmed deer reported in Missouri, Florida, and Pennsylvania (Ahasan et al., 2019b, 2019a; Cooper et al., 2014). Our data suggest that mosquito species are a potential vector for such orbiviruses. For these analyses, it was crucial that single mosquitoes were sequenced—if the mosquitoes had been pooled, it would not have been possible to associate potential vertebrate pathogens with a specific blood meal host.

A critical role for public data in public health

This study would have been impossible without rich public datasets containing sequences, species, locations, and sampling dates. These provided the backbone of information allowing us to identify the majority of our sequences. Citizen scientist resources, such as the iNaturalist catalog of biodiversity observations, was a valuable complement, providing empirical knowledge of species distributions in the mosquito collection area that resolved the ambiguity we detected in sequence space.

In sum, complementing conventional analyses of mosquito pools and field observations of mosquitoes and the animals they bite with single mosquito mNGS can provide valuable complementary information to enhance the evidence base for distinct interventions to control mosquito-borne infectious diseases. As shown here, single mosquito mNGS can map an uncharted landscape related to the movement of pathogens between mosquitoes and their reservoirs. This can inform the deployment of targeted detection or surveillance assays for both established and emerging mosquito-borne pathogens across large geographical areas or animal reservoir populations. As mosquitoes and their microbiota continue to evolve and migrate, posing new risks for human and animal populations, these complementary approaches will empower scientists and public health professionals.

Data and Code Availability

Raw and assembled sequencing data are deposited in NCBI Bioproject PRJNA605178. Code is available on Github at <https://github.com/czbiohub/california-mosquito-study>. Derived data (including all contigs) and supplementary data are available on Figshare at <https://doi.org/10.6084/m9.figshare.11832999.v2>

Materials and Methods

Mosquito collection

Adult mosquitoes were collected at sites indicated in using encephalitis virus survey (EVS) or gravid traps that were baited with CO₂ or hay-infused water, respectively. The collected mosquitoes were frozen using dry ice or paralyzed using triethyl amine and placed on a -15 °C chill table or in a glass dish, respectively, for identification to species using a dissection microscope. Identified female mosquitoes were immediately frozen using dry ice in deep well 96-well plates and stored at -80 °C or on dry ice until the nucleic acids were extracted for sequencing

RNA Preparation

Individual mosquitoes were homogenized in bashing tubes with 200uL DNA/RNA Shield (Zymo Research Corp., Irvine, CA, USA) using a 5mm stainless steel bead and a TissueLyserII (Qiagen, Valencia, CA, USA) (2x1min, rest on ice in between). Homogenates were centrifuged at 10,000xg for 5min at 4 °C, supernatants were removed and further centrifuged at 16,000xg for 2min at 4 °C after which the supernatants were completely exhausted in the nucleic acid extraction process. RNA and DNA were extracted from the mosquito supernatants using the ZR-Duet™ DNA/RNA MiniPrep kit (Zymo Research Corp., Irvine, CA, USA) with a scaled down version of the manufacturer's protocol with Dnase treatment of RNA using either the kit's DNase or the Qiagen RNase-Free DNase Set (Qiagen, Valencia, CA, USA). Water controls were performed with each extraction batch. Quantitation and quality assessment of RNA was done by the Invitrogen Qubit 3.0 Fluorometer using the Qubit RNA HS Assay Kit (ThermoFisher Scientific, Carlsbad, CA, USA) and the Agilent 2100 BioAnalyzer with the RNA 6000 Pico Kit (Agilent Technologies, Santa Clara, CA, USA).

Library Prep and Sequencing

Up to 200ng of RNA per mosquito, or 4uL aliquots of water controls extracted in parallel with mosquitoes, were used as input into the library preparation. A 25pg aliquot of External RNA Controls Consortium (ERCC) RNA Spike-In Mix (Ambion, ThermoFisher Scientific, Carlsbad, CA, USA) was added to each sample. The NEBNext Directional RNA Library Prep Kit (Purified mRNA or rRNA Depleted RNA protocol; New England BioLabs, Beverly, MA, USA) and TruSeq Index PCR Primer barcodes (Illumina, San Diego, CA, USA) were used to prepare and index each individual library. The quality and quantity of resulting individual and pooled mNGS libraries were assessed via electrophoresis with the High Sensitivity NGS Fragment Analysis Kit on a Fragment Analyzer (Advanced Analytical Technologies, Inc.), the High-Sensitivity DNA Kit on the Agilent Bioanalyzer (Agilent Technologies, Santa Clara, CA, USA), and via real-time quantitative polymerase chain reaction (qPCR) with the KAPA Library Quantification Kit

(Kapa Biosystems, Wilmington, MA, USA). Final library pools were spiked with a non-indexed PhiX control library (Illumina, San Diego, CA, USA). Pair-end sequencing (2 x 150bp) was performed using an Illumina NovaSeq or Illumina NextSeq sequencing systems (Illumina, San Diego, CA, USA). The pipeline used to separate the sequencing output into 150-base-pair pair-end read FASTQ files by library and to load files onto an Amazon Web Service (AWS) S3 bucket is available on GitHub at <https://github.com/czbiohub/utilities>.

Mosquito Species Validation

To validate and correct the visual assignment of mosquito species, we estimated SNP distances between each pair of mosquito transcriptomes by applying SKA (Split Kmer Analysis) (Harris, 2018) to the raw fastq files for each sample. The hierarchical clustering of samples based on the resulting distances was largely consistent with the visual assignments, with each cluster containing a majority of a single species. To correct likely errors in the visual assignment, samples were reassigned to the majority species in their cluster, resulting in 7 changes out of 148 samples and one species assignment for a sample lacking a visual assignment.

Host and Quality Filtering

Raw sequencing reads were host- and quality-filtered and assembled using the IDseq (v3.2) (Kalantar et al., 2020) platform <https://idseq.net>, a cloud-based, open-source bioinformatics platform designed for detection of microbes from metagenomic data.

Host Reference

We compiled a custom mosquito host reference database made up of:

1. All available mosquito genome assemblies under NCBI taxid 7157 (*Culicidae*; n=41 records corresponding to 28 unique mosquito species, including 1 *Culex*, 2 *Aedes*, and 25 *Anopheles* records) from NCBI Genome Assemblies (accession date: 12/7/2018).
2. All mosquito mitochondrial genome records under NCBI taxid 7157 available in NCBI Genomes (accession date: 12/7/2018; n=65 records).
3. *Drosophila melanogaster* genome (GenBank GCF_000001215.4; accession date: 12/7/2018).

Mosquito Genome Assembly and mitochondrial genome accession numbers and descriptions are detailed in Supplemental Data file `mosquito_genome_refs.txt`.

Read Filtering

To select reads for assembly, we performed a series of filtering steps using the IDSeq platform:

Filter Host 1

Remove reads that align to the host reference using the Spliced Transcripts Alignment to a Reference (STAR) algorithm.

Trim Adapters

Trim Illumina adapters using trimmomatic.

Quality Filter

Remove low-quality reads using PriceSeqFilter.

Remove Duplicate Reads

Remove duplicate reads using CD-HIT-DUP.

Low-Complexity Filter

Remove low-complexity reads using the LZW-compression filter.

Filter Host 2

Remove further reads that align to the host reference using Bowtie2, with flag very-sensitive-local.

The remaining reads are referred to as "putative non-host reads." A detailed description of all parameters is available in the IDseq documentation.

<https://github.com/chanzuckerberg/idseq-dag/wiki/IDseq-Pipeline-Stage-%231:-Host-Filtering-and-QC>

Assembly

The putative non-host reads for each sample were assembled into contigs using SPADES (Bankevich et al., 2012) with default settings. The reads used for assembly were mapped back to the contigs using Bowtie2 (Langmead and Salzberg, 2012) (flag very-sensitive), and contigs with more than 2 reads were retained for further analysis.

Taxonomic Assignment

We aligned each contig to the nt and nr databases using BLASTn (Altschul et al., 1990) (discontinuous megablast) and PLAST (a faster implementation of the BLASTx algorithm), respectively. (The databases were downloaded from NCBI on Mar 27, 2019.) Default parameters were used, except the E-value cutoff was set to 1e-2. For each contig, the results from the database with a better top hit (as judged by bitscore) are used for further analysis.

For contigs with BLAST hits to more than one species, we report the lowest common ancestor (LCA) of all hits whose number of matching aligned bases alignment length*percent identity is no less than the number of aligned bases for the best BLAST hit minus the number of mismatches in the best hit. (In the

case that the same segment of the query is aligned for all hits, this condition guarantees that the excluded hits are further from the best hit than the query is.)

For 172,244 contigs there were strong BLAST hits to *Hexapoda*, the subphylum of arthropods containing mosquitoes. This is likely a consequence of the limited number and quality of genomes used in host filtering, and all contigs with an alignment to *Hexapoda* of at least 80% of the query length or whose top hit (by e-value) was to *Hexapoda* were discarded from further analysis.

Contigs with no BLAST hits are referred to as "dark contigs".

For RNA viruses, where complete or near-complete genomes were recovered, a more sensitive analysis was performed.

Viral Polymerase Detection and Segment Assignment

Alignments of viral RNA-dependent RNA polymerases used to detect domains were downloaded from Pfam. These were RdRP_1 (PF00680, *Picornavirales*-like and *Nidovirales*-like), RdRP_2 (PF00978, *Tymovirales*-like and Hepe-Virga-like), RdRP_3 (PF00998, *Tombusviridae*-like and *Nodaviridae*-like), RdRP_4 (PF02123, *Toti*-, *Luteo*-, and *Sobemoviridae*-like), RdRP_5 (PF07925, *Reoviridae*-like), Birna_RdRp (PF04197, *Birnaviridae*-like), Flavi_NS5 (PF00972, *Flaviviridae*-like), Mitovir_RNA_pol (PF05919, *Narnaviridae*-like), Bunya_RdRp (PF04196, *Bunyavirales*-like), Arena_RNA_pol (PF06317, *Arenaviridae*-like), Mononeg_RNA_pol (PF00946, *Mononega*- and *Chuviridae*-like), Flu_PB1 (PF00602, *Orthomyxoviridae*-like). Hidden Markov model (HMM) profiles were generated from these with HMMER (v3.1b2; <http://hmmer.org/>) and tested against a set of diverged viruses, including ones thought to represent new families. Based on these results only the RdRP_5 HMM was unable to detect diverged Reovirus RdRp, such as Chiqui virus. An additional alternative Reovirus HMM (*HMMbuild* command) was generated by using BLASTp hits to Chiqui virus, largely to genera *Cypovirus* and *Oryzavirus*, aligned with MAFFT (Katoh et al., 2005) (E-INS-i, BLOSUM30).

All contigs of length >500 base pairs were grouped into clusters using a threshold of $\geq 99\%$ identity (CD-HIT-EST (Li and Godzik, 2006)). Representative contigs from each cluster were scanned for open reading frames (standard genetic code) coding for proteins at least 200 amino acids long, in all six frames with a Python script using Biopython (Cock et al., 2009). These proteins were scanned using HMM profiles built earlier and potential RdRp-bearing contigs were marked for follow up. We chose to classify our contigs by focusing on RdRp under the assumption that *bona fide* exogenous viruses should at the very least carry an RdRp and be mostly coding-complete. Contigs that were not associated with an RdRp or coding-complete included Cell fusing agent virus (*Flaviviridae*, heavily fragmented) and Phasma-like nucleoprotein sequences (potential piRNAs) in a few samples.

Co-Occurrence

For each cluster whose representative contig contained a potential RdRp, we identified its putative viral segment from CD-HIT clusters whose set of samples overlapped the set of samples in the RdRp cluster at a threshold of 80%. (That is, a putative segment should be present in at least 80% of the samples that RdRp is present in, and RdRp should be present in at least 80% of the samples that the putative segment is present in).

In cases where a singleton segmented (bunya-, orthomyxo-, reo-, chryso-like, *etc*) virus was detected in a sample we relied on the presence of BLASTx hits of other segments to related viruses (*e.g.* diverged orthobunyavirus). We thus linked large numbers of viral or likely viral contigs to RdRps representing putative genomes for these lineages.

Final Classification

There were 1269 contigs identified as viral either by RdRp detection or co-occurrence, and the resulting species-level calls are used for further analysis in lieu of the LCA computed via BLAST alignments. This included 338 “dark contigs” which had no BLAST hits, 748 with LCA in Viruses; the LCAs for the remainder were Bacteria (9), and Eukaryota (4), and Ambiguous (170), a category including (including root, cellular organisms, and synthetic constructs). Reads are assigned the taxonomic group of the contig they map to.

Water Controls and Contamination

There are many potential sources of contaminating nucleic acid, including lab surfaces, human experimenters, and reagent kits. We attempt to quantify and correct for this contamination using 8 water controls. We model contamination as a random process, where the mass of a contaminant taxon t in any sample (water or Mosquito) is a random variable X_t . We convert from units of reads to units of mass using the number of ERCC reads for each sample (as a fixed volume of ERCC spike-in solution was added to each sample well). We estimate the mean of X_t using the water controls. We say that a taxon observed in a sample is a possible contaminant if the estimated mass of that taxon in that sample is less than 100 times the average estimated mass of that taxon in the water samples. Since the probability that a non-negative random variable is greater than 100 times its mean is at most 1% (Markov's inequality), this gives a false discovery rate of 1%. For each possible contaminant taxon in a sample, all contigs (and reads) assigned to that taxon in that sample were excluded from further analysis. A total of 46,603 reads were removed as possible contamination using this scheme. (Human and mouse were identified as the most abundant contaminant species.)

For every sample, "classified non-host reads" refer to those reads mapping to contigs that pass the above filtering, *Hexapoda* exclusion, and decontamination steps. "Non-host reads" refers to the classified non-host reads plus the reads passing host filtering which failed to assemble into contigs or assembled into a contig with only two reads.

Treemap

Treemaps (e.g. Figure 2) are a way of visualizing hierarchical information as nested rectangles whose area represents numerical values. To visualize the distribution of reads amongst taxonomic ranks, we first split the data into two categories: viral and cellular. For cellular taxonomic ranks (Bacteria, Eukaryotes, Archaea and their descendants) we assigned all reads of a contig to the taxonomic compartment the contig was assigned (see above, "Taxonomic Assignment"). For viral taxa we relied on the curated set of viral contigs coding for RdRp and their putative segments, where a putative taxonomic rank (usually family level) had been assigned. All the reads belonging to contigs that comprised putative

genomes were assigned to their own compartment in the treemap, under the curated rank. Additional compartments were introduced to either reflect aspects of the outdated and potentially non-monophyletic taxonomy which is nevertheless informative (*e.g.* positive- or double-strandedness of RNA viruses) or represent previously reported groups without an official taxonomic ID on public databases (*e.g.* Narna-Levi, Toti-Chryso, Hepe-Virga, *etc.*).

To prototype the cellular part of the treemap, all taxonomic IDs encountered along the path from the assigned taxonomic ID up to root (*i.e.* the taxonomic ID's lineage) were added to the treemap. Based on concentrations of reads in particular parts of the resulting taxonomic treemap, prior beliefs about the specificity of BLAST hits, and information utility, this was narrowed down to the following taxonomic ranks: *Bacteria*, *Wolbachia*, *Gammaproteobacteria*, *Spirochaetes*, *Terrabacteria* group, Fungi, *Boroetheria*, *Aves*, *Trypanosomatidae*, *Leishmaniinae*, *Viridiplantae*.

Microbiota distribution in single mosquitoes

In Figure 3, the denominators are non-host reads. The numerators are numbers of reads from contigs with confident assignments. For viruses, these contigs came from viral curation or co-occurrence. For *Wolbachia* and eukaryotes, these contigs had LCA assignment within the *Wolbachieae* tribe (taxid: 952) and *Eukaryota* superkingdom (taxid: 2759), respectively, and had a BLAST alignment where the percentage of aligned bases was at least 90%. Groups within viruses, *Wolbachia*, and eukaryotes were excluded for a given sample if the cumulative proportion of non-host reads was less than 1%. Samples were excluded if the total proportions of non-host reads belonging to viruses, *Wolbachia*, or eukaryotes were all below 1%.

Blood meal calling

For each of the 60 blood fed mosquito samples from Alameda County, we selected each contig with LCA in the subphylum *Vertebrata*, excluding those contained in the order *Primates* (because of the possibility of contamination with human DNA). For each sample, we identified the lowest rank taxonomic group compatible with the LCAs of the selected contigs. (A taxonomic group is compatible with a set of taxonomic groups if it is an ancestor or descendent of each group in the set.) For 44 of the 45 samples containing vertebrate contigs, this rank is at class or below; for 12 samples, it is at the species level. Each taxonomic assignment falls into one of the following categories: *Pecora*, *Aves*, *Carnivora*, *Rodentia*, *Leporidae*. In Figure 5, each sample with a blood meal detected is displayed according to which of those categories it belongs (Underlying data for figure 5 are provided in Supplemental Data Files 12 - 14). The remaining sample, CMS001_022_Ra_S6, contained three contigs mapping to members of *Pecora* and a single contig with LCA *Euarchothoglires*, a superorder of mammals including primates and rodents; we annotate this sample as containing *Pecora*.

Notably, 19 samples contain at least one contig with LCA in genus *Odocoileus* and another contig with LCA genus *Bos*. While the lowest rank compatible taxonomic group is the infraorder *Pecora*, it is likely that a single species endemic in the sampled area is responsible for all of these sequences. Given the observational data in the region (described in the main text), that species is likely a member of *Odocoileus* whose genome diverges somewhat from the reference.

Phylogenetic analyses

We chose a single Wuhan mosquito virus 6 genome from our study (CMS001_038_Ra_S22) as a reference to assemble by alignment the rest of the genome of strain QN3-6 (from SRA entry SRX833542 as only PB1 was available for this strain) and the two small segments discovered here for Australian segments (from SRA entries SRX2901194, SRX2901185, SRX2901192, SRX2901195, SRX2901187, SRX2901189, and SRX2901190) using Magic-BLAST (Boratyn et al., 2018). Due to much higher coverage in Australian samples, Magic-BLAST detected potential RNA splice sites for the smallest segment (hypothetical 3) which would extend the relatively short open reading frame to encompass most of the segment. Sequences of each segment were aligned with MAFFT (Auto setting) and trimmed to coding regions. For hypothetical 3 segment we inserted Ns into the sequence near the RNA splice site to bring the rest of the segment sequence into frame.

PhyML (Guindon et al., 2010) was used to generate maximum likelihood phylogenies under an HKY+ Γ_4 model (Guindon et al., 2010; Hasegawa et al., 1985; Yang, 1994). Each tree was rooted via a least-squares regression of tip dates against divergence from root in TreeTime (Sagulenko et al., 2018). Branches with length 0.0 in each tree (arbitrarily resolved polytomies) were collapsed, and trees untangled and visualized using baltic21 (<https://github.com/evogytis/baltic>).

To generate the Culex narnavirus 1 tanglegrams, 42 sequences of RdRp and 42 co-occurring Robin segment sequences from our samples and three previously published RdRp sequences (MK628543, KP642119, KP642120) as well as their three corresponding Robin segments assembled from SRA entries (SRR8668667, SRR1706006, SRR1705824, respectively) were aligned with MAFFT and trimmed to just the most conserved open reading frame (as opposed to its complement on the reverse strand). Maximum likelihood phylogenies for both RdRp and Robin segments were generated with PhyML with 100 bootstrap replicates under an HKY+ Γ_4 substitution model. The resulting phylogenies were mid-point rooted, untangled and visualized using baltic21 (<https://github.com/evogytis/baltic>).

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Figures

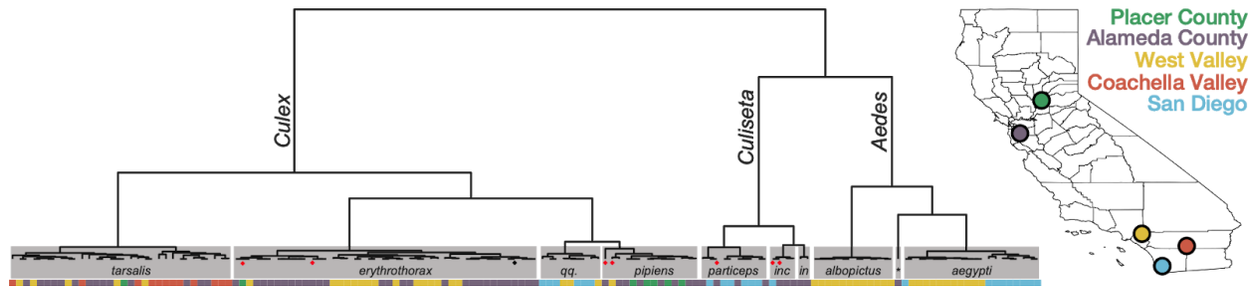


Figure 1. Whole transcriptome analysis for mosquito species identification. Hierarchical clustering of pairwise single-nucleotide polymorphism (SNP) distances between whole transcriptome sequences from the 148 mosquitoes included in this study estimated using SKA (Harris, 2018). The inferred mosquito species for each cluster (text in gray boxes) is the consensus of the species calls made by visual inspection during sample collection for samples in that cluster (*qq.* = *Culex quinquefasciatus*, *part.* = *Culiseta particeps*, *inc* = *Culiseta incidens*, *in* = *Culiseta inornata*, *albo* = *Aedes albopictus*, * = *Aedes dorsalis*). Red dots below the nodes on the tree highlight mismatches (n=7) between consensus transcriptome species call and initial visual species call; black dot indicates a sample missing a visual species call. Color bar below the tree shows the collection location for each sample, coded according to the California map legend at right.

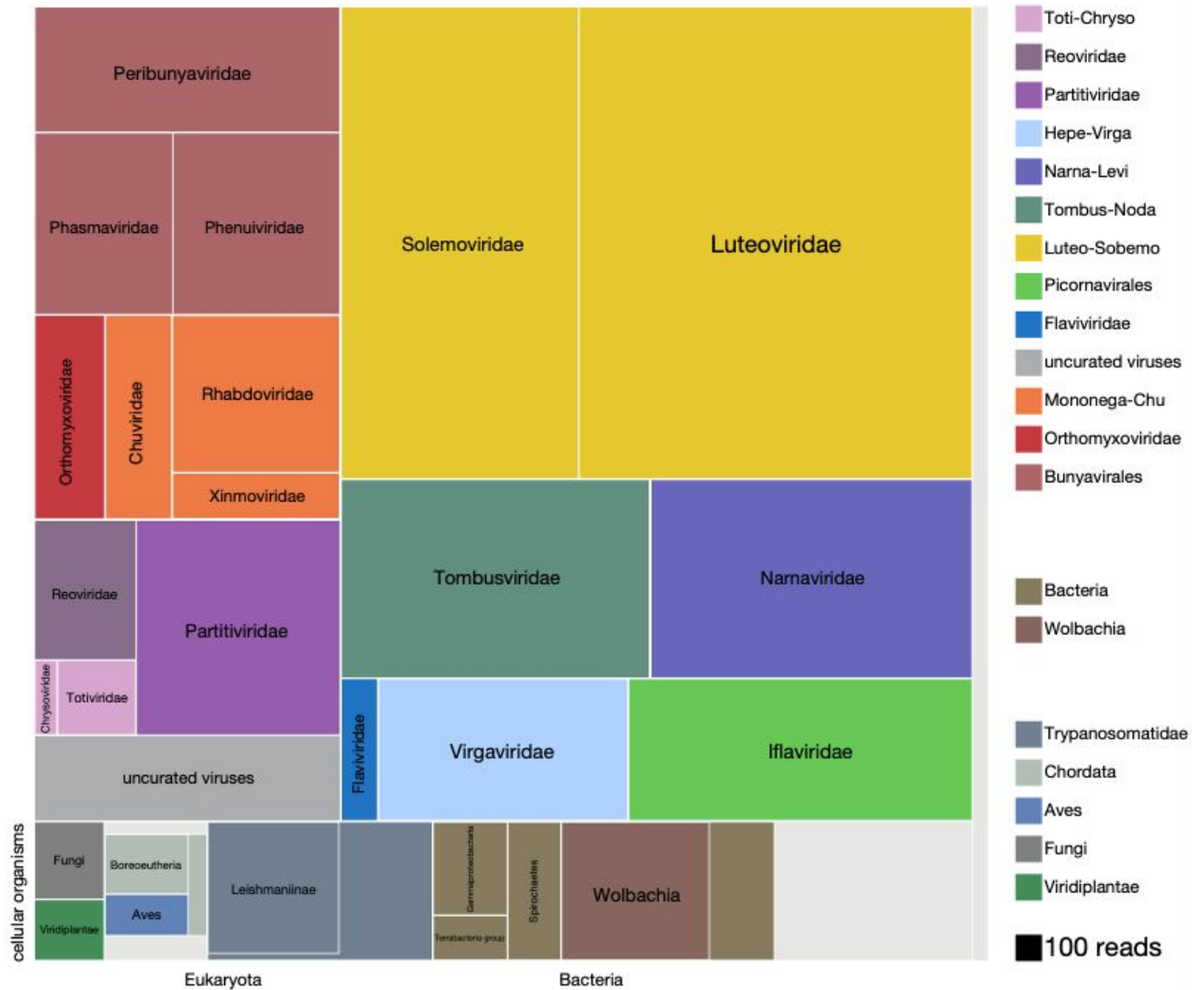


Figure 2. Viruses dominate the microbial signature of mosquitoes. Treemap plot of the proportion of reads recovered among non-host contigs assembled from the 148 single mosquitoes, with viral taxa making up the top portion of the plot above “cellular organisms” label at lower left edge of image, which designates reads assembled into contigs encompassing prokaryotic taxa, the eukaryotic taxa and the taxonomically ambiguous “cellular organisms” that were not possible to assign to a higher resolution (light gray box). The block areas are plotted in proportion to the number of reads assembled into contigs that could be assigned to a given taxa (see area scale, legend). The gray block labeled “uncurated viruses” corresponds to the number of reads assembled into contigs that were clearly viral in origin, but difficult to further resolve due fragmented homology and/or the lack of an associated RNA-dependent RNA polymerase.

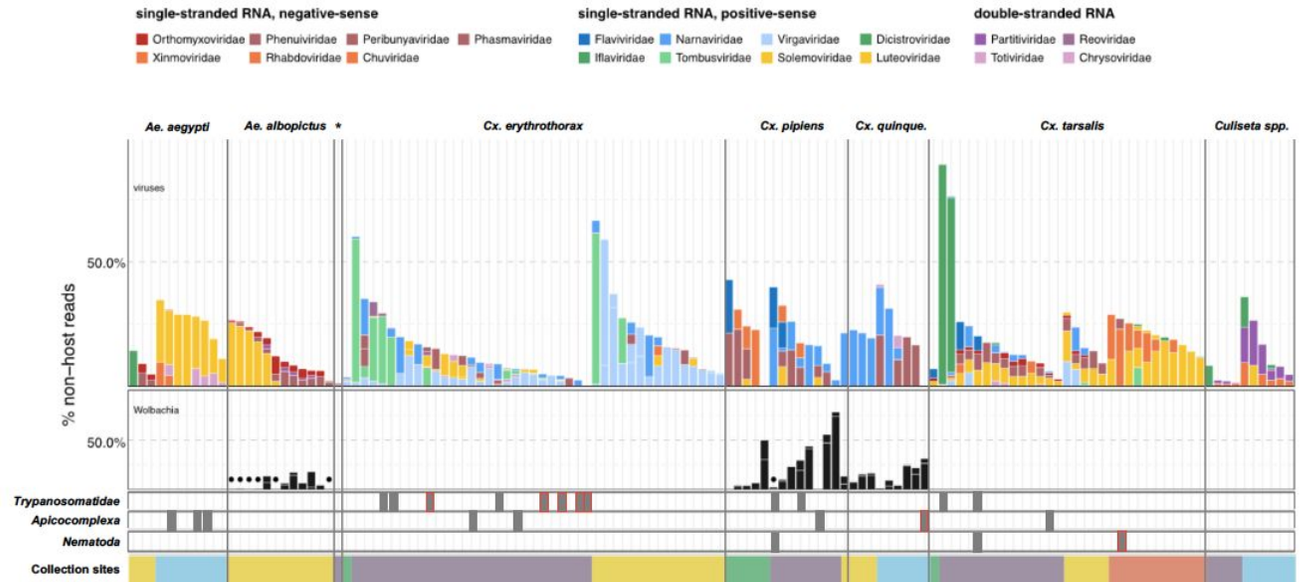


Figure 3. Single mosquito sequencing reveals the heterogeneity of microbiota between individuals, species and locations. Plotted bars report the proportions of non-host reads from individual mosquitoes that are supported by $\geq 1\%$ of non-host reads corresponding to assembled contigs identified as viruses (top panel), *Wolbachia* (middle panel), and selected eukaryotic microbes (bottom panel). Species are color coded as indicated in the legend (top of graph). Plotted symbols on the *Wolbachia* and eukaryotic panels indicate microbes confidently identified, but present at $< 1\%$ of non-host reads (middle panel, black circles = *Wolbachia* taxa; bottom panels, gray bars are plotted in samples with *Trypanosomatidae*, *Apicomplexa* or *Nematoda* taxa; gray bars outlined in red indicate detection of these taxa at $\geq 1\%$ of non-host reads). Samples were clustered by mosquito species (top labels) and ordered by: i) collection site location from north to south (indicated at the bottom, colored as in Figure 1) and ii) viral abundance (descending order, left to right).

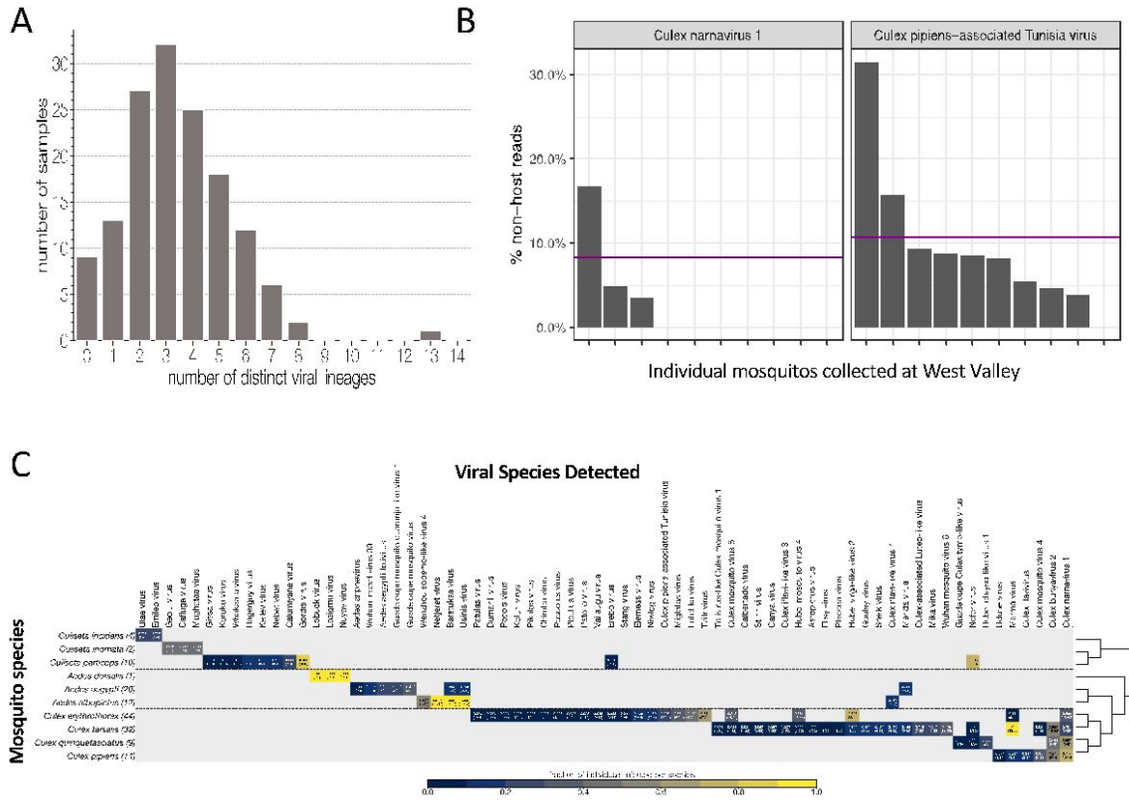


Figure 4. Quantifying viral diversity in and prevalence among single mosquitoes (A) Distribution of mosquitoes within the study in which no, one, or multiple viral lineages were detectable. The sample with 13 distinct viral lineages is sample CMS002_053a that contains six *Botourmia*-like viruses thought to primarily infect fungi and in which evidence of an ergot fungus was detected. (B) An example of viruses with similar bulk abundance, but different prevalence. Both *Culex narnavirus 1* and *Culex pipiens*-associated *Tunisia virus* were found among the 10 *Culex erythrothorax* mosquitoes collected at the same collection site in West Valley. The bulk abundances were calculated by the mean % non-host reads averaged across the 10 mosquitoes for *Culex narnavirus 1* and *Culex pipiens*-associated *Tunisia virus* were 8.3% and 10.6%, respectively (as indicated by the red line). However, the prevalence (i.e. the percent of single mosquitoes carrying the virus) was markedly different (C). Global analysis of viral prevalence measured in this study. For each virus, the fraction of individuals infected within each species was calculated, shown on a color scale. Mosquito species arranged according to a phylogeny based on the cytochrome c oxidase subunit I (COI) gene.

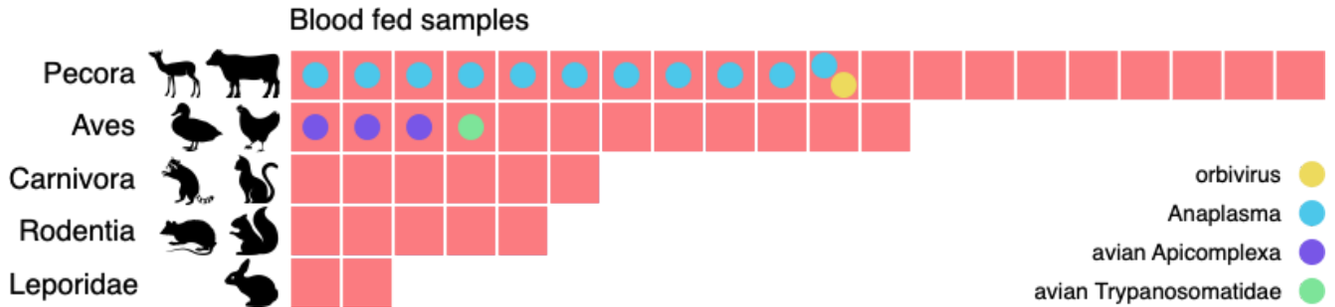


Figure 5. Metagenomic identification of sources of blood meals in individual mosquitoes Consensus taxonomic calls of vertebrate contigs for 45 of 60 blood fed mosquitoes collected in Alameda County. The remaining 15 samples had no vertebrate contigs. Red blocks represent individual mosquito samples; colored circles represent co-occurring contigs matching Orbivirus, Anaplasma, Avian Apicomplexa and Avian Trypanosomatidae representing possible bloodborne pathogens of the blood meal host.

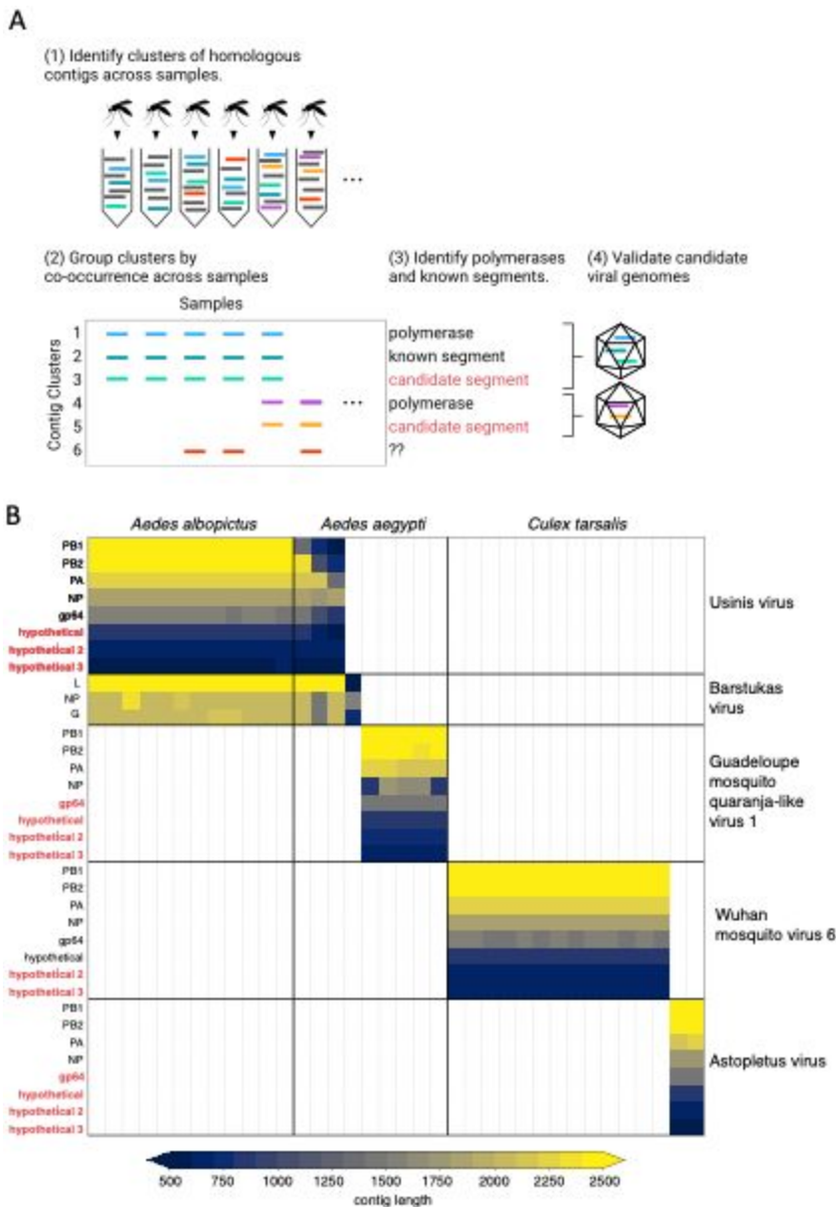


Figure 6. Bringing viral genomic dark matter into the light through single mosquito next generation metagenomic sequencing. (A) Previously unrecognized viral genome segments were identified among unaligned “dark matter” contigs using co-occurrence analysis, which assumes that all segments of a segmented virus will co-occur in the samples where that virus is present and be absent in samples where the virus is absent. (B) Matrix of contigs derived from four distinct Orthomyxoviruses and one Phasma-like virus that were detected via their distinct co-occurrence pattern across mosquitoes. Rows are clusters of highly similar (>99% identity) contigs and columns are individual mosquito samples. Light grey vertical lines delineate mosquito samples, dark black vertical lines indicate boundaries between mosquito species of each sample. Dark horizontal lines delineate segments comprising viral genomes. Labels on the right indicate viruses, with genomes delineated by horizontal lines. Guadeloupe mosquito quaranja-like virus 1 and Wuhan mosquito virus 6 were previously described and *Ūsinis*, *Barstukas* and *Astopletus* were named here. At left, plain text indicates putative labels for homologous clusters; black text indicates segments identifiable via homology (BLASTx) and red text indicates contig clusters that co-occur with identifiable segments but themselves have no identifiable homology to anything in GenBank. The Phasma-like *Barstukas* virus exhibits a nearly perfect overlap with *Ūsinis* virus (except for one sample in which *Ūsinis* was not found) but is identifiable as a Bunya-like virus due to having a three-segmented genome with recognizable homology across all segments to other Phasma-like viruses. Cells are colored by contig lengths (see color scale legend), highlighting their consistency which is expected of genuine segments. Deviations in detected contig lengths (e.g., *Aedes aegypti* samples that harbor shorter *Ūsinis* virus genome segments) reflect the presence of partial or fragmented contig assemblies in some of the samples.

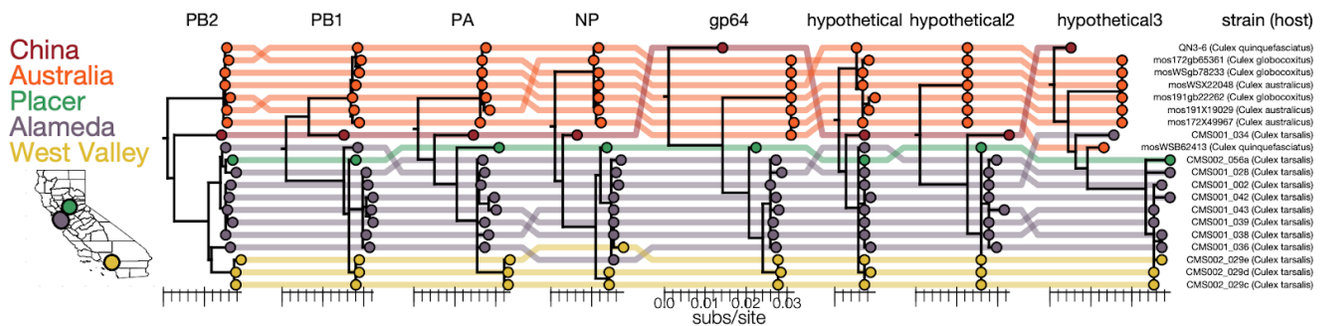
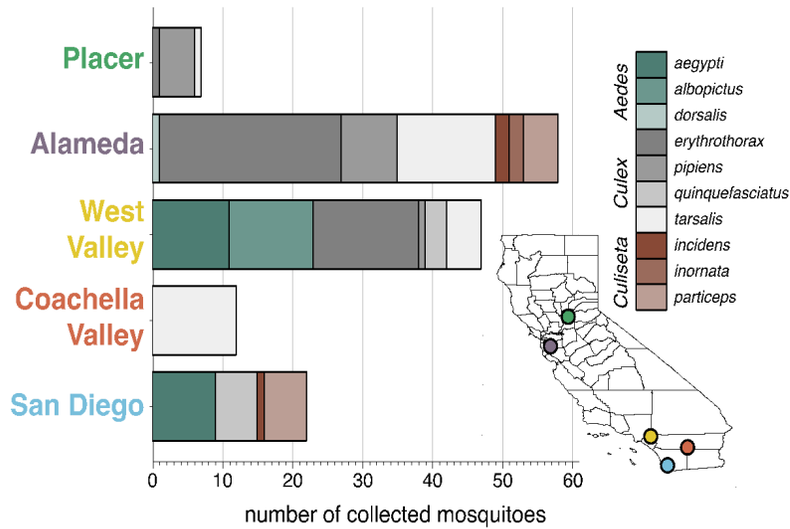
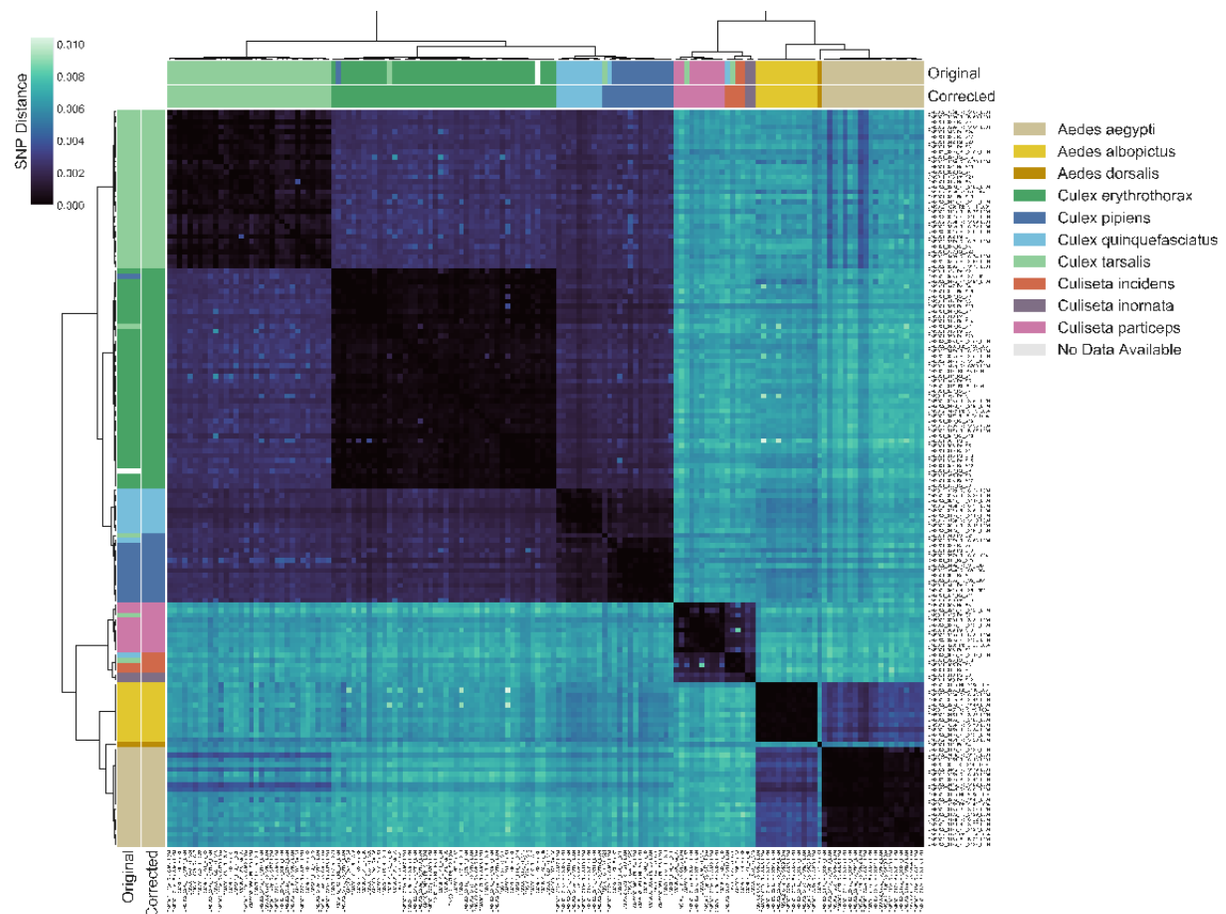


Figure 7. Genomic evidence for segment reassortment and intercontinental spread of mosquito-borne quaranjaviruses (A) The Chinese strain (QN3-6) was originally described from a single PB1 sequence, while Australian (orange) viruses were described as having 6 segments. In this study we report the existence of two additional smaller segments (named hypothetical 2 and hypothetical 3) which we have assembled from our samples and the SRA entries of Chinese and Australian samples. Strains recovered in California as part of this study are colored by sampling location (Placer County in green, Alameda County in purple, West Valley in yellow). Strain names and hosts are indicated on the far right with colored lines tracing the position of each tip in other segment trees with crossings visually indicating potential reassortments.

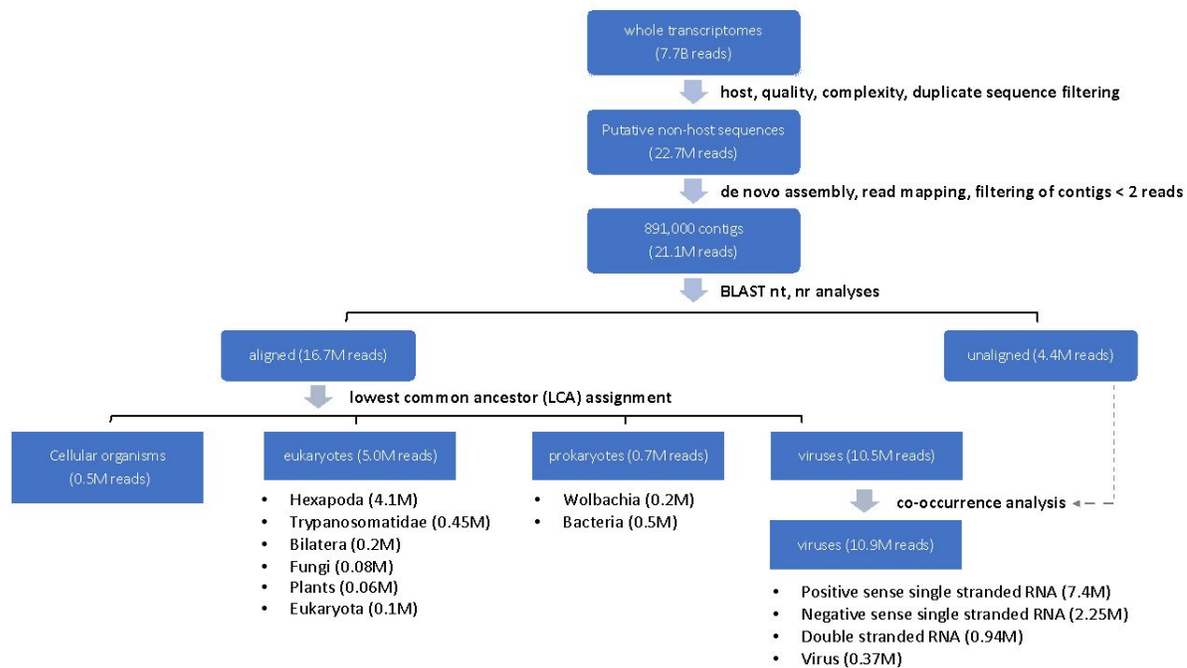
Supplemental Figures



Supplemental Figure 1. Diversity of individual mosquitoes collected across California. Plot shows the number of each genus and species of mosquitoes collected across 5 regions in California: Placer County, Alameda County, West Valley, Coachella Valley, and San Diego, as indicated by the correspondingly colored dots on the map, lower right). Colors in the bar graphs correspond to mosquito species as indicated in the legend.

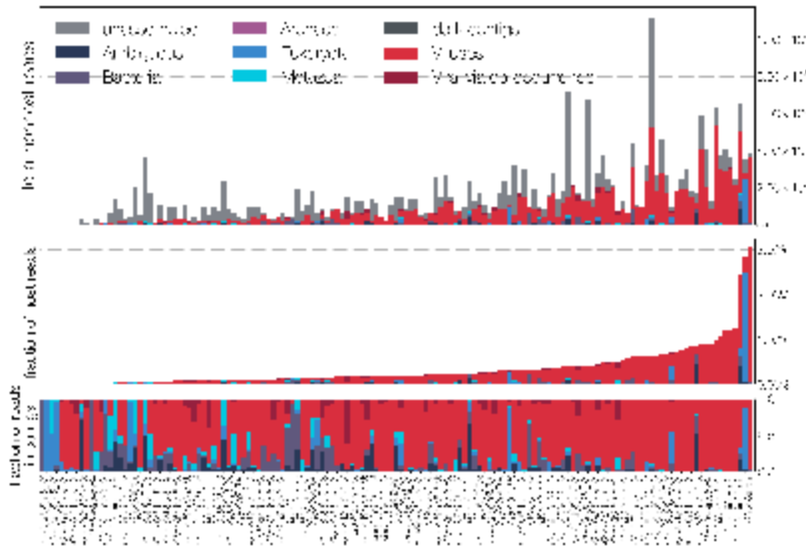


Supplemental Figure 2: Transcriptome-based identification of mosquito species. Hierarchical clustering of pairwise single-nucleotide polymorphism (SNP) distances between samples estimated using SKA (Harris, 2018), ranging from 0 (no SNPs detected, black) to 0.01 (1% of comparable sites had SNPs, blue). Outside color bar indicates visual species label for each sample (Visual), inside color bar indicates consensus transcriptome species label for each sample's cluster (Computed). Computed mosquito species calls were used when Visual and Computed calls were discordant.

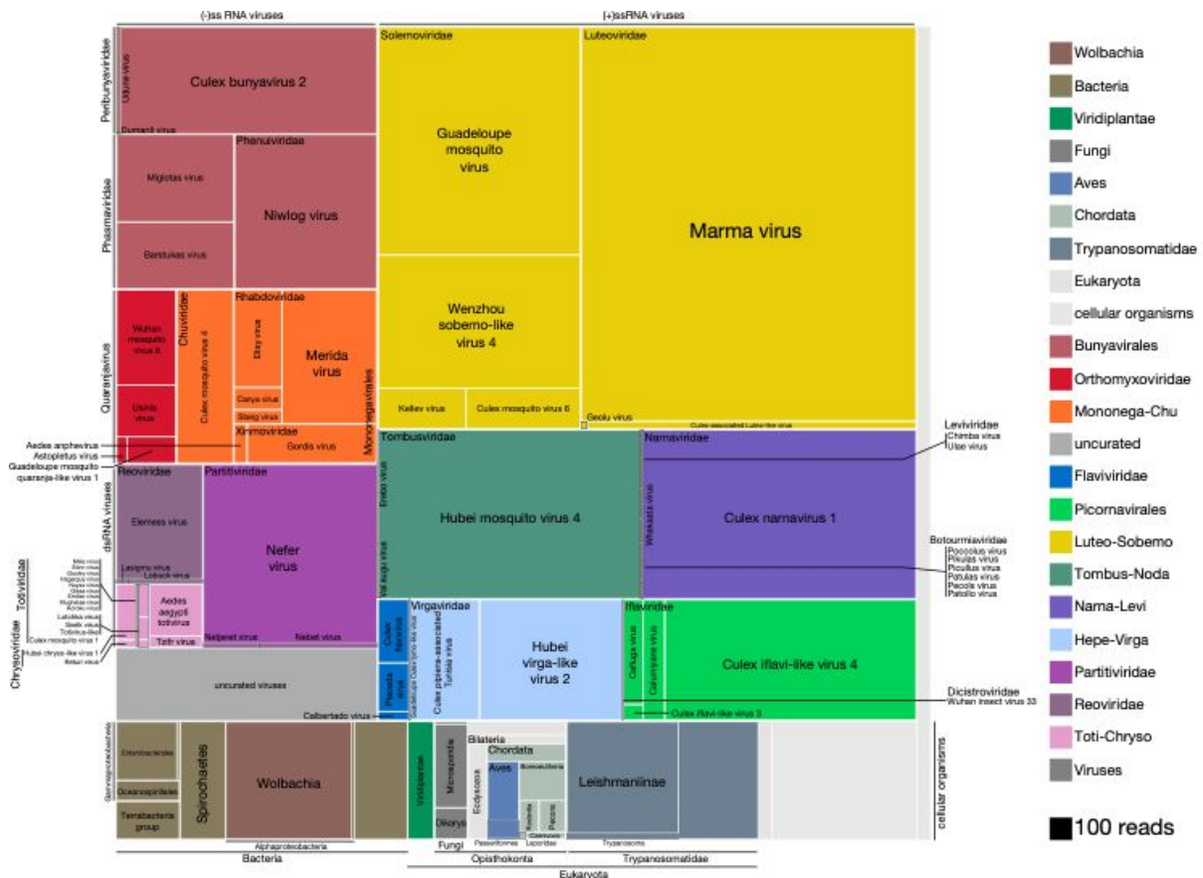


Supplemental Figure 3: Summary of single mosquito mNGS analysis pipeline with sequence recovery yields. A set of 7.78 billion reads served as input transcriptome sequence data for analysis of the taxonomic distribution of 148 single mosquitoes sequenced in this study. We first filtered host, low quality, and low complexity sequences from each transcriptome dataset. A total set of 22.7 million reads passed all of these filters. A set of 891,000 contigs were *de novo* assembled from 21.1 million of these reads (93%). To classify and quantify the reads assembled into the contigs into viral, eukaryotic or prokaryotic taxa, we aligned each contig to the NCBI nucleotide and non-redundant protein databases. Recovery of complete or near complete genomes of RNA viruses facilitated viral taxonomic assignment directly from the BLAST results. We further curated contigs likely to code for viral RNA-dependent RNA polymerases (RdRps) and contigs that were physically unlinked but showed evidence of co-occurrence with viral RdRps (labeled “co-occurrence analysis”, above; details in Materials and Methods). In contrast, only a fraction of a genome was generally detected among contigs aligning to prokaryotic and eukaryotic taxa. Here, we employed a lowest common ancestor (LCA) analysis of the BLAST results for these contigs (Materials and Methods) to infer their corresponding taxon assignments.

We found that a significant proportion of reads assembled into contigs did not readily align to microbial species or plausible bloodmeal hosts. One set of 4.1 million reads contributed to contigs classified as *Hexapoda*. These likely correspond to mosquito host sequences too diverged from available mosquito genome reference sequences to be recognized at the host filtering step. As such, these reads were excluded from further analysis. An additional set of 4.4 million reads were assembled into contigs corresponding to metagenomic “dark matter”, *i.e.* contigs without recognizable homology to previously published sequences. Co-occurrence analysis (see main text), allowed us to identify additional viral contigs from this set of contigs, contributing 0.4 million reads to the viral reads tally. This provided a final set of 13.4 million reads assembled into contigs with BLAST alignments to enable analysis of the distribution of viral, bacterial, or eukaryotic taxa detectable among all the mosquitoes analyzed for this study.

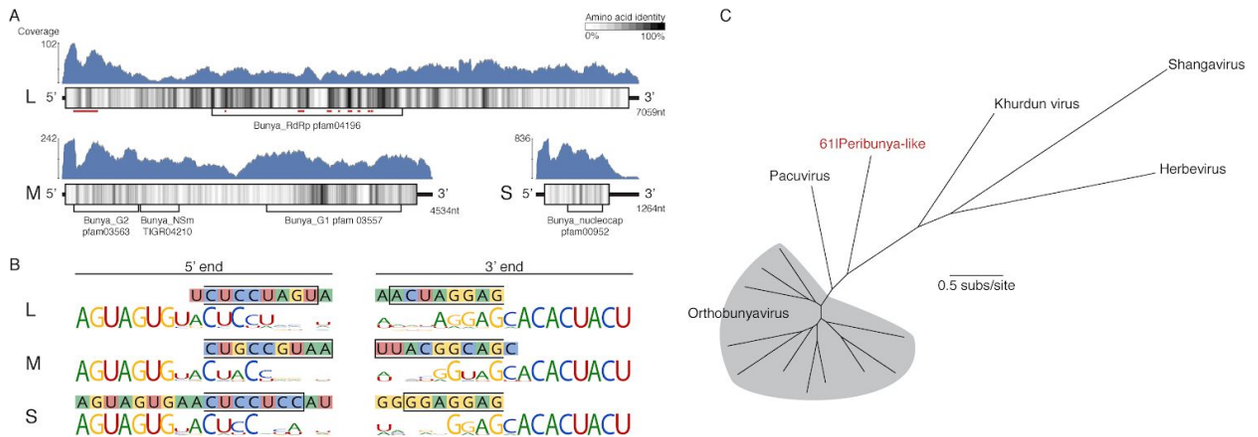


Supplemental Figure 4: Summary of all reads in all sampled mosquitoes A breakdown of reads for each sample, into unassembled (those which did not assemble into contigs), dark (those which assembled but were not identifiable taxonomically), Ambiguous (those with LCA at root or cellular organisms), Bacteria, Archaea, Eukaryota, Metazoa, Viruses (via alignment to reference), and Viral via co-occurrence analysis (See Figure 6). Top panel is a stacked bar decomposing the total reads per sample. Middle panel shows the break-down as a fraction of all reads in a sample. Bottom panel shows the fractional breakdown among assembled reads.

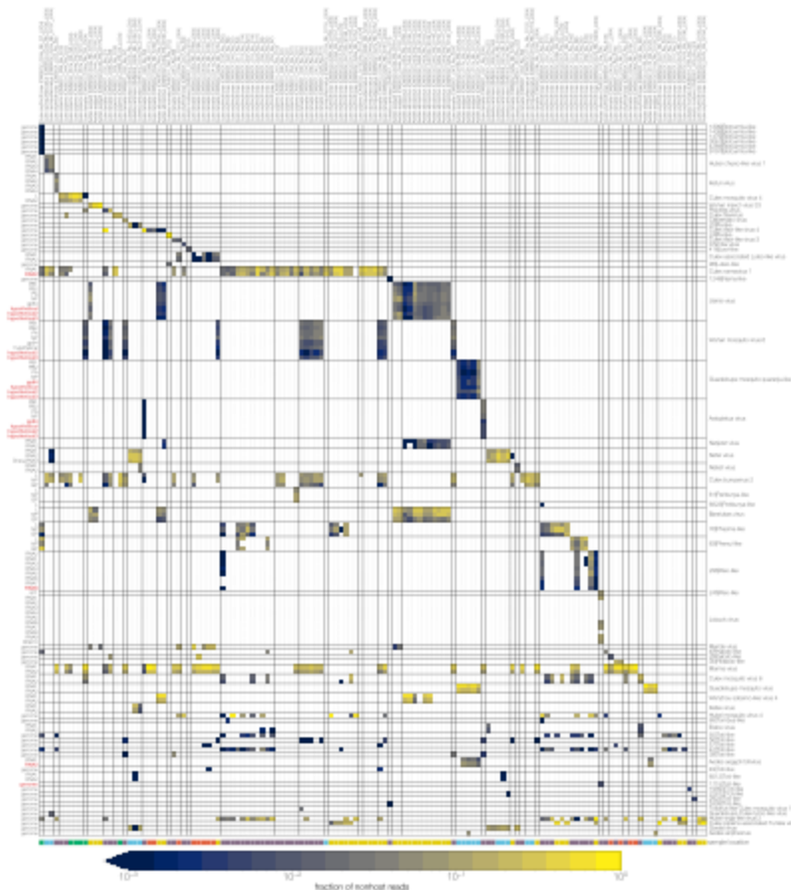


Supplemental Figure 5: High resolution breakdown the microbial signature of mosquitoes. Shown is the read count distribution across non-host taxonomic groups detected among the set of 148 mosquitoes. Treemap overview of taxa detected among the assembled contigs are plotted in squares shaded according to taxon identity (legend) and sized according to number of reads supporting each contig (scaled as indicated, bottom right). Due to the uncertainty in prokaryotic and eukaryotic taxonomic assignments, cellular organisms are assigned to the lowest plausible taxonomic rank. Each distinct viral lineage is displayed as its own compartment. Reddish hues indicate negative-sense RNA viruses; purple hues indicate double-stranded RNA viruses; blue, green and

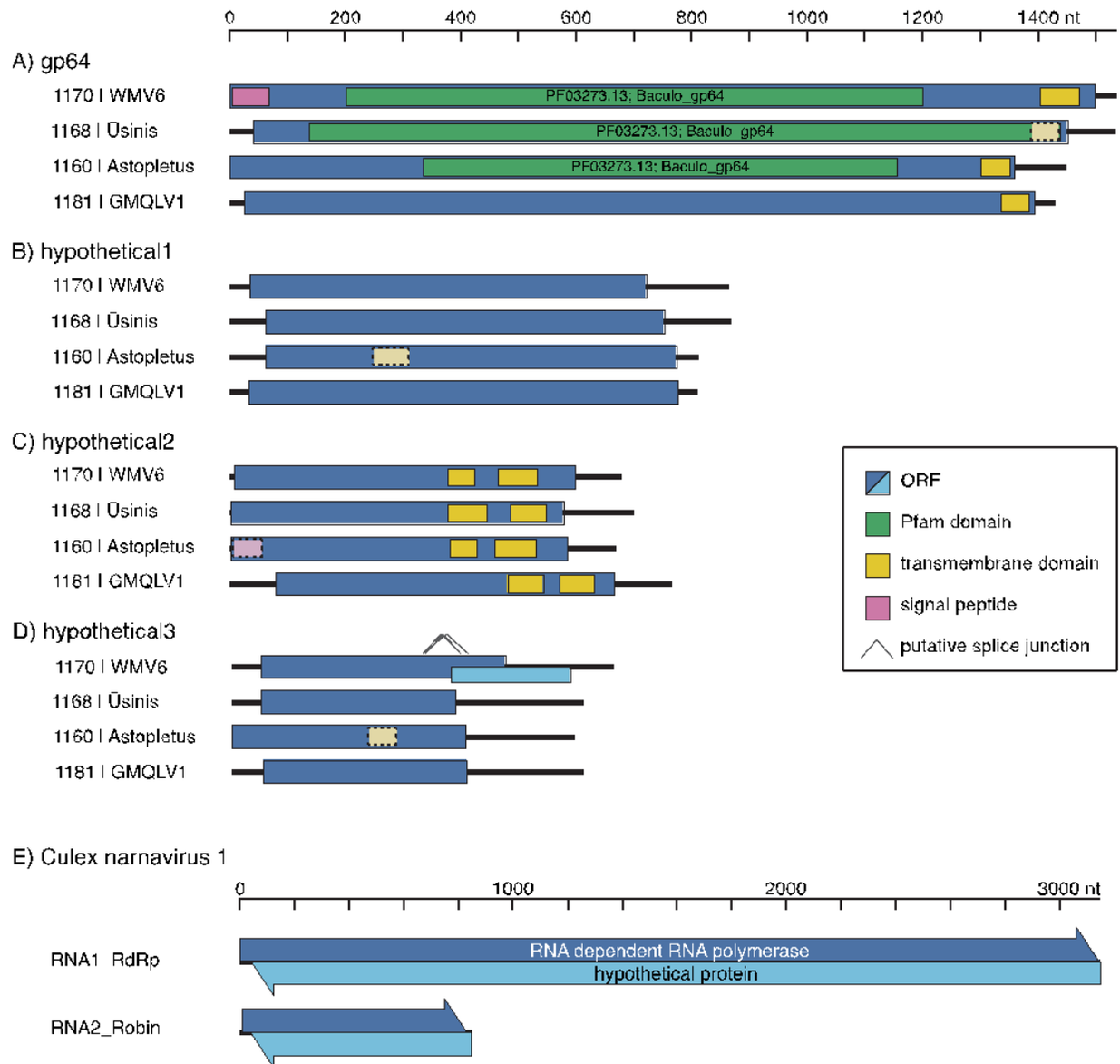
yellow hues indicate positive-sense RNA viruses. The "uncurated viruses" box (light grey, lower left) indicates contigs that were clearly viral in origin but too fragmented and/or not associated with an RNA-dependent RNA polymerase.



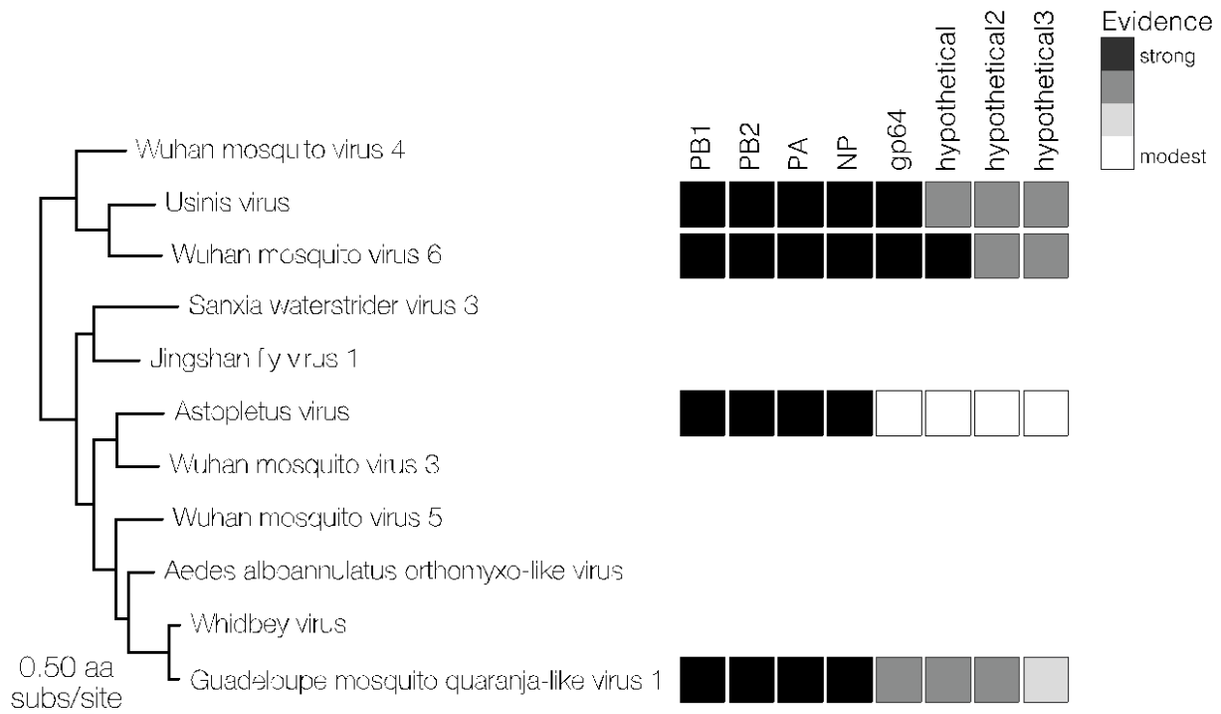
Supplemental Figure 6. Analysis of novel peribunya-like virus showing completeness. (A) Read coverage in blue across the length of each genome segment. Diagrams of genome segments display the longest ORF and indicate amino acid identity with a sliding window of 15 amino acids. Regions of homology to known domains are indicated by Pfam labels, and on the L segment, the locations of canonical RdRp motifs are indicated by red underlining. (B) Ends of the assembled contigs are shown (sequence in 11led boxes). Regions complementary between 5' and 3' end for each segment are indicated by a black outline. These contig ends were aligned to the conserved end sequences of orthobunya and nearby peribunya viruses (logo diagram shows conservation after alignment of reference sequences). (C) Maximum likelihood phylogenetic tree with representative orthobunyaviruses and nearby peribunyavirus



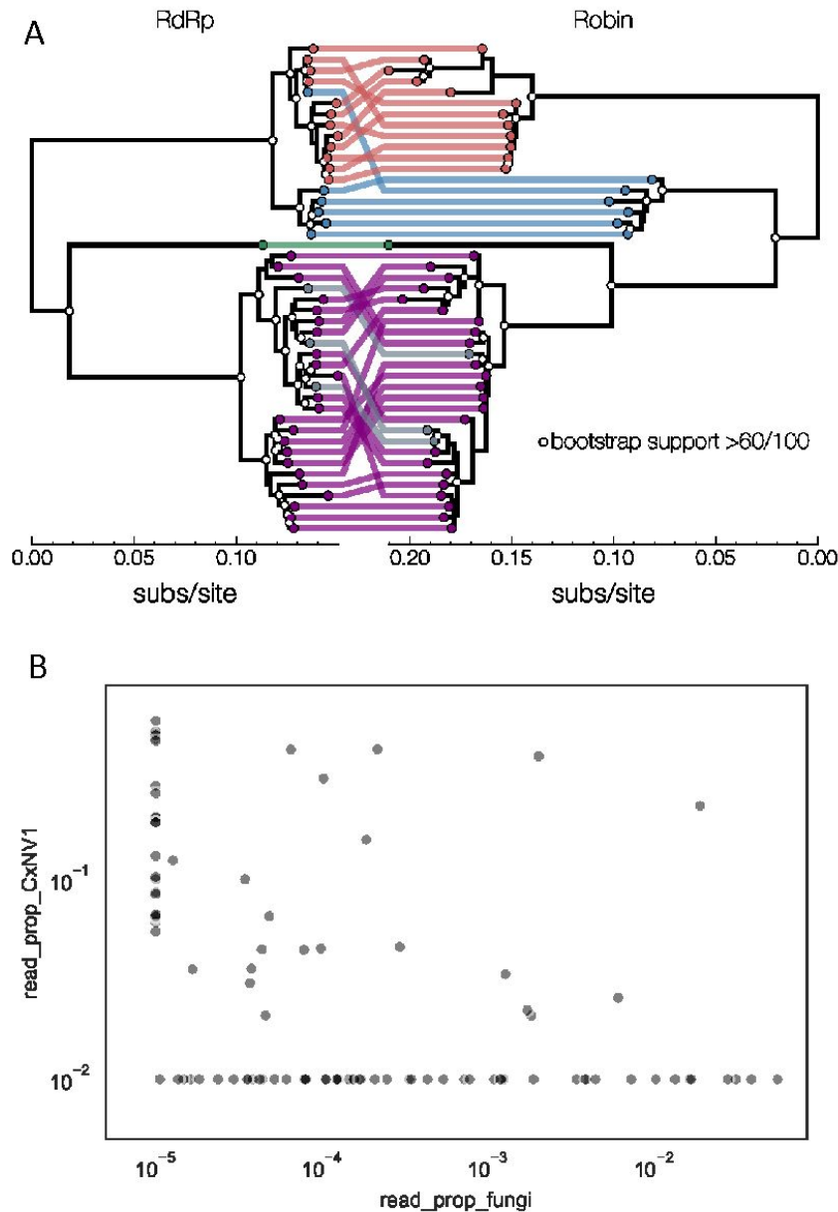
Supplemental Figure 7. Identifying novel RNA segments in the 'dark matter' by co-occurrence. Rows indicate groups of contigs clustered based on nucleotide identity (>99%). Columns are samples with computed mosquito species indicated. Cells are colored by the fraction of non-host reads in the sample with color bar legend at the bottom. Clusters of rows likely comprising a single genome are separated from other such putative genomes by black horizontal lines with identified or proposed virus names. Black vertical lines delineate blocks of neighboring samples with the same computed mosquito species. Sampling locations are indicated by colored rectangles at the bottom, above the color bar legend.



Supplemental Figure 8. Co-occurrence in individual mosquitoes allows identification of novel RNA genome segments in mosquito-borne viruses Diagrams of newly discovered genome segments for quaranjaviruses identified in this study, including Wuhan Mosquito Virus 6 (WMV6), Usinis, Astopletus, and Guadeloupe Mosquito, Quaranja-Like Virus 1 (GMQLV1) (panels A-D, respectively), and Culex narnavirus 1 (panel E), are shown (displayed 5' to 3', left to right, not aligned to each other). (A-D) The longest ORFs of the new quaranjavirus segments were analyzed for the presence of transmembrane domains and similarity to known domains (see Methods for details). Weaker support for these predictions is indicated by lighter color and dashed border. In the hypothetical 3 segment of WMV6, putative splicing events could result in a jump to a frameshifted ORF, which would substantially increase the coding region of the segment. (E) In both segments of the Culex narnavirus 1, long uninterrupted ORFs are found on both RNA strands (arrowheads indicate 5' to 3' direction), and in each case the codon boundaries are aligned.



Supplemental Figure 9. RdRp-based maximum likelihood tree spanning the quarantaviruses in this study for which 8 segments were recovered. The 8 boxes in the graphic to the right of the tree correspond to each of these 8 genome segments, which are labeled at top right. Levels of evidence for the existence of each segment is highlighted by a grayscale shading of the segment boxes, ranging from strong (black) to modest (white), based on: homology to BLAST hits from NCBI nt/nr database (black); assembly of contigs >500nt from SRA samples containing the RdRp, or co-occurrence in >10 samples in our data (dark grey); reads mapping from SRA and co-occurrence in 5 samples in our data (light grey); or co-occurrence in <5 samples and shared protein domains (white).



Supplemental Figure 10. Co-occurrence analysis enables identification of novel second narnavirus ambigrammatic RNA segment. (A) Narnavirus Robin segment. tanglegram shows two midpoint rooted nucleotide phylogenies of *Culex narnavirus 1* - RdRp segment on left and Robin segment on right. Tips are colored based on visual presence of clusters on longer branches - clusters in red, blue, green, and purple are sequences reported in this study. Grey tips in the RdRp phylogeny were published by other studies, and their Robin segment counterpart was assembled here from corresponding SRA entries. Reassortment between the two segments is largely restricted to lineages within clusters, but reassortment also took place in either the red or the blue lineage. One possibility is a replacement of a red/blue lineage RdRp with a blue/red lineage or a replacement of red Robin with a diverged and unsampled lineage. White dots on nodes indicate bootstrap support >60 out of 100 replicates. Scale at the bottom of each tree is nucleotide substitutions per site. (B) Proportion of non-host reads mapped to contigs of *Culex narnavirus 1* versus proportion of non-host reads mapped to contigs of any fungus. The lack of correlation argues against an indirect infection of fungi by the narnavirus. Each point represents a single mosquito. For visualization purposes, values of zero are displayed at 1e-6 for fungi and 1e-3 for *Culex narnavirus*.

Table 1. Complete genomes of known and novel viral taxa recovered in this study

genome type	scientific name*	viral family	novel?	number detected overall	number in <i>Aedes</i> samples	number in <i>Culex</i> samples	number in <i>Culiseta</i> samples
single stranded positive sense RNA	Patello virus	<i>Botourmiaviridae</i>	TRUE	1	0	1	0
	Piculus virus	<i>Botourmiaviridae</i>	TRUE	1	0	1	0
	Poccolus virus	<i>Botourmiaviridae</i>	TRUE	1	0	1	0
	Pikulas virus	<i>Botourmiaviridae</i>	TRUE	1	0	1	0
	Pecols virus	<i>Botourmiaviridae</i>	TRUE	1	0	1	0
	Patulas virus	<i>Botourmiaviridae</i>	TRUE	1	0	1	0
	Wuhan insect virus 33	<i>Dicistroviridae</i>	FALSE	3	3	0	0
	Placeda virus	<i>Flaviviridae</i>	TRUE	3	0	3	0
	Culex flavivirus	<i>Flaviviridae</i>	FALSE	3	0	3	0
	Calbertado virus	<i>Flaviviridae</i>	FALSE	1	0	1	0
	Culex iflavi-like virus 4	<i>Iflaviridae</i>	FALSE	6	2	4	0
	Calumiyane	<i>Iflaviridae</i>	TRUE	3	0	0	3
	Culex iflavi-like virus 3	<i>Iflaviridae</i>	FALSE	2	0	2	0
	Cafluga virus	<i>Iflaviridae</i>	TRUE	1	0	0	1
	Chimba virus	<i>Leviviridae</i>	TRUE	1	0	1	0
	Ulae virus	<i>Leviviridae</i>	TRUE	1	0	0	1
	Culex-associated Luteo-like virus	<i>Luteoviridae</i>	FALSE	7	0	7	0
	Geolu virus	<i>Luteoviridae</i>	TRUE	1	0	0	1
	Culex namavirus 1	<i>Narnaviridae</i>	FALSE	43	0	43	0
	Whakaata virus	<i>Narnaviridae</i>	TRUE	1	0	0	1
	Marma virus	<i>Solemoviridae</i>	FALSE	39	0	39	0
	Culex mosquito virus 6	<i>Solemoviridae</i>	FALSE	15	0	15	0
	Guadeloupe mosquito virus	<i>Solemoviridae</i>	FALSE	8	8	0	0
	Wenzhou sobemo-like virus 4	<i>Solemoviridae</i>	FALSE	7	0	0	0
	Kellev virus	<i>Solemoviridae</i>	TRUE	2	0	0	2
	Hubei mosquito virus 4	<i>Tombusviridae</i>	FALSE	18	0	18	0
	Erebo virus	<i>Tombusviridae</i>	TRUE	4	0	3	1
	Vai augu virus	<i>Tombusviridae</i>	TRUE	2	0	2	0
	Guadeloupe Culex tymo-like virus	<i>Tymoviridae</i>	FALSE	1	0	1	0
	Hubei virga-like virus 2	<i>Virgaviridae</i>	FALSE	36	0	36	0
	Culex pipiens-associated Tunisia virus	<i>Virgaviridae</i>	FALSE	13	0	13	0
	Culex mosquito virus 4	<i>Chuviridae</i>	FALSE	6	0	6	0
	Usinis virus	<i>Orthomyxoviridae</i>	TRUE	15	15	0	0
	Wuhan mosquito virus 6	<i>Orthomyxoviridae</i>	FALSE	13	0	13	0
	Guadeloupe mosquito quaranja-like virus 1	<i>Orthomyxoviridae</i>	FALSE	5	5	0	0
	Astopletus virus	<i>Orthomyxoviridae</i>	TRUE	2	0	2	0
	Culex bunyavirus 2	<i>Peribunyaviridae</i>	FALSE	30	0	30	0
	Udune virus	<i>Peribunyaviridae</i>	TRUE	1	0	1	0
	Dumani virus	<i>Peribunyaviridae</i>	TRUE	1	0	1	0
	Barstukas virus	<i>Phasmoviridae</i>	TRUE	16	0	0	0
Miglotas virus	<i>Phasmoviridae</i>	TRUE	16	0	16	0	
Nwlog virus	<i>Phenuiviridae</i>	TRUE	8	0	8	0	
Merida virus	<i>Rhabdoviridae</i>	FALSE	9	4	5	0	
Stang virus	<i>Rhabdoviridae</i>	TRUE	4	0	4	0	
Elisy virus	<i>Rhabdoviridae</i>	TRUE	3	0	3	0	
Canya virus	<i>Rhabdoviridae</i>	TRUE	2	0	2	0	
Gordis virus	<i>Xinmoviridae</i>	TRUE	9	0	0	9	
Aedes anphevirus	<i>Xinmoviridae</i>	FALSE	2	0	0	0	
Hubei chryo-like virus 1	<i>Chrysoviridae</i>	FALSE	2	0	2	0	
Keturi virus	<i>Chrysoviridae</i>	TRUE	1	0	1	0	
Netjeret virus	<i>Partitiviridae</i>	TRUE	11	11	0	0	
Nefer virus	<i>Partitiviridae</i>	TRUE	10	0	2	8	
Nebet virus	<i>Partitiviridae</i>	TRUE	2	0	0	2	
Elemess virus	<i>Reoviridae</i>	TRUE	6	0	6	0	
Lasigmu virus	<i>Reoviridae</i>	TRUE	1	0	0	0	
Lobuck virus	<i>Reoviridae</i>	TRUE	1	1	0	0	
Tzifr virus	<i>Totiviridae</i>	TRUE	27	0	27	0	
Lotchka virus	<i>Totiviridae</i>	TRUE	19	0	19	0	
Mika virus	<i>Totiviridae</i>	TRUE	10	0	10	0	
Aedes aegypti totivirus	<i>Totiviridae</i>	FALSE	5	5	0	0	
Snelk virus	<i>Totiviridae</i>	TRUE	4	0	4	0	
Gouley virus	<i>Totiviridae</i>	TRUE	4	0	4	0	
Stinn virus	<i>Totiviridae</i>	TRUE	2	0	2	0	
Wap virus	<i>Totiviridae</i>	TRUE	2	0	0	2	
Nuyav virus	<i>Totiviridae</i>	TRUE	1	1	0	0	
Mughataa virus	<i>Totiviridae</i>	TRUE	1	0	0	1	
Koroku virus	<i>Totiviridae</i>	TRUE	1	0	0	1	
Emileo virus	<i>Totiviridae</i>	TRUE	1	0	0	1	
Gissa virus	<i>Totiviridae</i>	TRUE	1	0	0	1	
Totivirus-like Culex mosquito virus 1	<i>Totiviridae</i>	FALSE	1	0	1	0	

*gray rows highlight viruses previously detected in California (Chandler, JA et al., 2015 and Sadeghi, M et al., 2018)