Wolbachia reduces virus infection in a natural population of Drosophila

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Abstract

Wolbachia is a maternally transmitted bacterial symbiont that is estimated to infect approximately half of arthropod species. In the laboratory it can increase the resistance of insects to viral infection, but its effect on viruses in nature is unknown. Here we report that in a natural population of Drosophila melanogaster, individuals that are infected with Wolbachia are less likely to be infected by viruses. By characterising the virome by metagenomic sequencing and then testing individual flies for infection, we found the protective effect of Wolbachia was virus-specific, with the prevalence of infection being up to 15% greater in Wolbachia-free flies. The antiviral effects of Wolbachia may contribute to its extraordinary ecological success, and in nature the symbiont may be an important component of the antiviral defences of insects.

Introduction

Wolbachia is an alphaproteobacterium that lives within the cytoplasm of arthropod cells and is maternally transmitted. It infects approximately half of arthropod species, and many strains manipulate host reproduction, most commonly by inducing cytoplasmic incompatibility (CI). CI allows Wolbachia to invade populations by causing embryonic mortality when uninfected females mate with infected males, hence conferring a selective advantage to infected females. Wolbachia can also protect Drosophila species against RNA viruses. Combined with Wolbachia’s ability to invade populations due to CI, this provides a way to modify natural insect populations to make them resistant to viral infections. Wolbachia has been transferred from Drosophila to the mosquito Aedes aegypti, where it limits the replication of dengue virus as well as chikungunya, yellow fever, Zika and West Nile viruses. When Wolbachia-infected mosquitoes were released into the wild, the bacterium spread through the mosquito populations by CI, and large field trials have shown substantial reductions in dengue incidence in the human population.

While the antiviral effects of Wolbachia have great value in the field of public health, their ecological importance is far from clear. As Wolbachia is estimated to infect 52% of arthropod species, it may be a major component of antiviral defences in nature. However, studies on the antiviral effects of Wolbachia have largely been performed under laboratory conditions and frequently with artificial routes of infection. Wolbachia protects wild mosquitoes against dengue virus, but here Wolbachia has been artificially transferred into the mosquito, resulting in an activation of immune defences that is not typical of natural host-Wolbachia associations. Furthermore, two studies of natural populations of Drosophila melanogaster have failed to find evidence of Wolbachia protecting infected insects against viral infection, so there is currently no evidence that Wolbachia is a natural antiviral defence of insects.

If antiviral protection is present in nature, Wolbachia may frequently be a mutualist that defends its host against infection. This may explain why Wolbachia strains that do not cause CI and have no obvious phenotypic effect can invade and be maintained in populations. For example, the Wolbachia strain wAu spread through Australian populations of Drosophila simulans despite not causing CI. In the same host species, the wRi Wolbachia strain has evolved to become a mutualist, but the cause of the fitness benefit is unknown. The benefits provided by antiviral protection could also allow CI-inducing strains of Wolbachia to invade new populations and species. Theory predicts that CI can only invade when local infection frequencies becomes sufficiently high to offset imperfect maternal transmission and infection costs. However, recent data suggested that Wolbachia can spread from arbitrarily low frequencies.
This can be explained if there is a fitness advantage for the host caused by Wolbachia, which may be its antiviral effects.

Results

*Wild Drosophila melanogaster* harbour a diverse community of viruses

We collected 1014 male *D. melanogaster* from an orchard in Connecticut, USA and extracted RNA from single flies. To characterize the diversity of viruses in this population in an unbiased way, we pooled RNA from groups of 23 flies to generate 40 RNAseq libraries. These were mapped to the published genome sequences of *D. melanogaster*, the Wolbachia strain wMel and known *Drosophila* viruses. The unmapped reads were then assembled to identify novel *Drosophila*-associated viruses (see methods for inclusion criteria).

We identified 30 viruses associated with *D. melanogaster* in this population (Figure 1). There was a wide range of abundance, with approximately 260,000 times more reads from the most abundant virus relative to the least abundant virus (Figure 1). Seventeen of the viruses we identified, including the twelve most abundant ones, have previously been described as infecting *Drosophila melanogaster*.

We identified thirteen viruses that have not been associated with *D. melanogaster* before. We reconstructed the phylogeny of these viruses based on predicted protein sequences, and refer to them by the name of the virus family (Supplementary Figure 1). One of these viruses belongs to the *Flaviviridae* and is closely related to Hermitage virus from *Drosophila immigrans*. One virus from the order *Picornavirales* is closely related to Basavirus sp. A novel virus belonging to the *Tymoviridae* is closest to Bee Macula−Like virus 2, which has been detected in several wild bee species. Four novel viruses identified within the *Totiviridae* clustered with Ahus virus from *Culex* mosquitoes, Keenan toti-like virus from the fly *Sarcophaga impatiens* and *Leishmania* RNA virus-like. One virus was a negative sense RNA virus related to *Drosophila unispina* virus 1. Five viruses belong to the *Narnaviridae*, and these were related to a virus from a fungus (*Plasmopara viticola* lesion associated narnavirus 2), an arthropod (*Serbia narna−like* virus 4-like) and a trypanosome (*Leptomonas seymouri* RNA virus-like). As viruses will be present in the food, environment and pathogens of flies, we would caution that the presence of these viruses in our samples does not mean they infected *D. melanogaster*, although the close relationship of many of them to other arthropod viruses suggests that some do (Supplementary Figure 1).

We used our RNAseq data to design PCR primers that matched the eleven viruses present in all our libraries, and tested the panel of 1014 individual flies for infection by quantitative PCR. Viral infection is common, with 93% of flies infected with at least one virus (N=938, including data only from samples tested for all 11 viruses). This infection rate was driven by the high prevalence of Galbut virus and Vera virus, which infect 68% and 75% of flies respectively (Figure 1). These belong to the Partitiviridae, a family of viruses with segmented double-stranded RNA genomes. Galbut virus, which has previously been reported to infect most wild *D. melanogaster*, is efficiently vertically transmitted through both males and females, likely explaining its high prevalence. Seven other viruses infected over 10% of flies (Figure 1). The viruses that we assayed by PCR cover a diversity of taxonomic groups, including a double-stranded DNA virus (*Kallithea* virus), a negative-sense RNA virus (*Drosophila melanogaster* sigmavirus),
two dsRNA viruses (Vera and Galbut viruses) and six positive-sense RNA viruses (La Jolla, Craigies Hill, Motts Mill, Nora, Dansoman, Thika, Kilifi and Drosophila A viruses).

**Wolbachia protects wild flies against viral infection**

Seventy-one percent of the flies carried Wolbachia (N=1014), and these flies were infected with fewer viruses. Wolbachia-free flies were infected with a mean of 2.85 viruses, which is 15% more than the number of viruses detected in Wolbachia-infected flies (2.48 viruses; Wilcoxon rank sum test: $W = 101030, p = 0.0005$), suggesting the Wolbachia is protecting flies against infection in nature.

We estimated the prevalence of each virus in Wolbachia-free and Wolbachia-infected flies, and found there are no cases where the symbiont completely blocks viral infection (Figure 2a). To quantify the level of protection we calculated the risk that a Wolbachia-free fly was infected with a virus relative to the risk of a fly carrying Wolbachia (Figure 2b). In 9 out of 10 cases the risk of infection was greatest in Wolbachia-free flies (Figure 2a and 2b), and for two viruses this effect was statistically supported (Figure 2A and 2B; $p_{mcmc}<0.001$). These were a positive-sense RNA virus—Motts Mill virus—where the Wolbachia-free flies were 2.73 times more likely to be infected, and the dsRNA partitivirus Vera virus, where Wolbachia-free flies were 1.19 times more likely to be infected (Figure 2b). For both of these viruses we repeated the PCR tests of all the samples using an independent set of primers to verify these results (Supplementary Figure 2 a and b).

As well as reducing the likelihood that flies are infected, Wolbachia could reduce viral loads in infected flies. To investigate this, we examined viral loads among the virus-infected flies. For nine of the ten viruses there is no significant difference between the Wolbachia-infected and Wolbachia-free flies (Supplementary Figure 3; $p>0.01$). However, Galbut virus loads were significantly lower in the presence of Wolbachia (Figure 2c; $p=0.0007$). Comparing the distribution of viral loads, it is clear that this is caused by a minority of flies with strongly reduced viral loads in the Wolbachia-infected flies, while most individuals have similar loads (Figure 2c). Furthermore, this result still holds if the viral load was not normalised to rpl32 mRNA levels, indicating that it is not an artefact of Wolbachia affecting expression of the reference gene we used ($F=14.47, \text{ d.f.}=1,632, p=0.0002$).

**Discussion**

We have found that Wolbachia protects wild Drosophila against viral infection, with Wolbachia-infected flies carrying on average 0.37 fewer viruses. As viruses are common in natural insect populations, this phenotype may benefit many Wolbachia-infected insects and partly explain the extraordinary ecological success of Wolbachia. If the magnitude of this benefit is sufficient to outweigh the fitness cost of carrying Wolbachia, the symbiont will become a mutualist that can invade populations in the absence of other phenotypes. Establishing whether this is the case is particularly important as the Wolbachia strains that provide the greatest anti-viral protection tend to be associated with the highest fitness costs, as both traits depend on the density of Wolbachia in insect cells. However, even if the benefits of antiviral protection are insufficient to make Wolbachia a mutualist and there remains a net fitness cost, then the antiviral phenotype can still reduce this cost, making it more likely that Wolbachia can invade populations as a reproductive parasite.
The effect of Wolbachia on host fitness will depend not only on the reduction in viral prevalence and titre, but also on how harmful virus infection is to the fly. Of the three viruses affected by Wolbachia, only the phenotypic effects of Galbut virus infection have been reported. Under laboratory conditions this virus had only very modest effects on lifespan and fecundity 32. If we speculate that these results hold for other viruses affected, and given that Wolbachia-infected flies carrying ~0.37 fewer viruses, the magnitude of any fitness benefit might be so small as to have minimal impact on Wolbachia dynamics. However, harsh competitive conditions can increase the cost of infection, and these may be common in the field. For example, flies infected with the Drosophila melanogaster sigmavirus appear healthy in the laboratory. However, in the field or under competitive laboratory conditions it is estimated to reduce fitness by 20-30% 33,34. If this was the case for the viruses affected by Wolbachia, then the benefits of antiviral protection could be as high as 10%. This is comparable to the fitness benefit of wAu that allowed it to invade populations of Drosophila simulans in the absence of CI 18.

An important caveat to this study is that we only investigated males, as we could not reliably morphologically identify female D. melanogaster to the species level. However, because Wolbachia is maternally transmitted, it is antiviral protection in females that will have the greatest effect on the symbiont’s fitness and population dynamics. Therefore, an important question for the future is whether similar levels of antiviral protection are seen in female hosts.

Our results contrast with three previous that failed to find any effect of Wolbachia on the natural viral community of Drosophila. The first of these was a study designed to characterize the diversity of viruses infecting D. melanogaster and D. simulans, and the authors suggest their sampling design means they have low power to detect the effects of Wolbachia 16. The second study investigated D. melanogaster, but used considerably smaller sample sizes than us and reared the flies for one or more generations in the laboratory at 19°C before testing them 15. It was later discovered that the antiviral effect of wMel is greatly reduced at this temperature 35. Finally, another study investigated D. simulans but used comparatively small sample sizes that are unlikely to detect effects of the size we observed 36.

The microbiome plays a key role in protecting animals against infection, and in insects this role is frequently played by specialized heritable endosymbionts that function alongside the immune system as an integral component of the animal’s defences against infection 37. For the first time, our results demonstrate Wolbachia naturally protects wild insects against infection and should therefore be regarded as a defensive symbiont. Because Wolbachia is so common in terrestrial arthropods 1 it may be an important component of antiviral defence in many species. This has the potential to affect the population biology of beneficial and pest insects, disease transmission by vector species, and the evolution of insect immune defences 38.
Methods

Field collection

Flies were collected at Lyman Orchards in Middlefield, CT, USA, a common field site to collect natural *Drosophila melanogaster* populations \(^{39,40}\). From the 4th to the 6th of September 2018, we collected a total of 1014 *D. melanogaster* males by aspirating and netting over fermenting dropped peaches. We collected males as they can be identified to species level morphologically and individually preserved them in RNAlater™ reagent a few hours after field collecting.

RNA preparation and *Wolbachia* screening

RNA was isolated from single flies using Trizol™ (ThermoFisher, 15596018) extraction as previously described \(^{41}\). RNA pellets were re-suspended in 10µl nuclease free water (ThermoFisher, AM9930) and stored at -80°C. Half of the RNA from each fly was saved for library preparation and half was reverse transcribed with Promega GoScript reverse transcriptase and random hexamer primers. cDNA was diluted 1:10 with nuclease free water. RT-qPCR was performed on an Applied Biosystems StepOnePlus system using Sensifast Hi-Rox Sybr kit (Bioline) with the following PCR cycle: 95°C for 2 minutes followed by 40 cycles of: 95°C for 5 seconds followed by 60°C for 30 seconds. Each sample was tested for *Wolbachia* infection by amplification of a segment of the gene *atpD* by RT-qPCR using primers CCCTATCTTAAAGGAGGAAA and AATCCTTTATGAGCTTTTGC \(^{31}\). To normalise estimates of *Wolbachia* and virus loads we also amplified the fly gene *RpL32* using primers TGCTAAGCTGTCGCACAAATGG and TGCGCTTGTTCGATCCGTAAC \(^{42}\).

Library preparation and RNA sequencing

Single fly RNA samples were combined into 40 different pools, each pool contained samples from 23 individual flies to give a total volume of 69µl per pool. The RNA from each pool was quantified using Qubit RNA HS assay kit (ThermoFisher, Q32852). RNAseq libraries were prepared from each RNA pool as follows: Ribosomal RNA was depleted using a Ribo-Zero Gold rRNA Removal Kit (Human/Mouse/Rat) (Illumina, MRZG12324). Between 620ng of RNA for the lowest and 1800ng of RNA for the highest sample in a total volume of 28µl was used. To this was added 8µl of Ribo-zero removal solution and 4µl of Ribo-Zero reaction buffer. The protocol was followed according to the manufacturer's recommendation. The rRNA-depleted RNA was cleaned up using ethanol precipitation and the resulting pellet was re-suspended in 5µl of nuclease free water.

RNAseq libraries were prepared using the NEB Next Ultra II Directional RNA Library Prep kit for Illumina (New England Biolabs, E7760L) according to the manufacturer's recommendations. All 5µl of the rRNA-depleted sample was used for each library. Adapters used were from KAPA Single-Indexed Adapter kits KK8701 and KK8702, the 30µM stock was diluted to 1.87µM before use. 8 cycles of PCR were used to amplify the libraries. Libraries were quantified using Qubit HS DNA quantification kit (ThermoFisher, Q32854). The final concentration of libraries was 11-29ng/µl in a volume of 20µl. The quality of the libraries was assessed using Bioanalyzer HS DNA kit (Agilent, 5067-4626) according to the manufacturer's instructions. The relative quantity of the libraries was ascertained using qPCR: 3 x 1:1000.
dilutions were made from each library by adding 1µl of library to 1ml of 10mM Tris-HCl pH8.0 with
0.05% Tween 20. 2µl of each dilution was used in a qPCR reaction using primers, IS5.reamp.P5: AATGATACGGCCACCCGA and IS6.reamp.P7: CAAGCAGAAGACGGCATACGA. Libraries were normalized to the concentration of the lowest in the pool by diluting in 0.1x TE buffer and combined into 3 separate pools of 13 or 14 libraries. The multiplexed library pools were quantified by Qubit HS DNA as above and assessed for quality and average fragment length using a Bioanalyzer HS DNA kit as above. The concentration of each pool was calculated and then diluted to 20nM by adding the appropriate quantity of 0.1 x TE buffer before sequencing. Paired-end RNA sequencing reads from 40 libraries were obtained. Libraries were sequenced on three lanes of the Illumina HiSeq4000 with paired end 150bp reads. Quality control of the raw RNA sequencing reads was implemented with TrimGalore-0.6.0 (http://www.bioinformatics.babraham.ac.uk/projects/trim_galore/).

Mapping to published genomes

The bioinformatic analyses are summarized in Supplementary Figure 4. Trimmed reads were mapped to combined genomes of Drosophila melanogaster, Wolbachia strain wMel and viruses isolated from or associated with flies in the genus Drosophila (Supplementary Figure 4; Round 1 Mapping). To account for genetic variation in the viral population, the viral sequences included all the sequences deposited in GenBank. Mapping was carried out with STAR-2.6.0 with default settings. Uniquely and multiple mapped reads were collected and counted for D. melanogaster, Wolbachia and each virus. Multiple mapped reads were counted only once as a randomly selected location where they had mapped.

Virus discovery

To reduce the size of the dataset, unmapped reads from all libraries were pooled and mapped to Ribosomal RNAs (rRNA) database downloaded from SILVA, including both SSU and LSU datasets, using bowtie2- v2.3.5.1. The rRNA reads were removed from the pooled reads. Trinity-v2.8.4 was then used to assemble transcript sequences from the pooled RNAseq reads with minimum contig length set at 200 nucleotides. Assembled contigs with open reading frames no shorter than 30 amino acids identified by TransDecoder (https://github.com/TransDecoder/TransDecoder) were collected, and subsequently blasted to NCBI non-redundant protein database and viral non-redundant protein database using DIAMOND blastx. Contigs with blast top results corresponding to viral origins in both databases were identified as candidate viral contigs, and were selected to be assembled into longer contigs using Sequencher 4.5 (http://www.genecodes.com), followed by manual curation (Sequencher contigs).

These candidate viral contigs were once more queried against the NCBI non-redundant protein database using DIAMOND blastx to identify closely related viruses for inclusion in phylogenetic analyses. Novel viruses where the top blast hit in GenBank did not infect eukaryotes were excluded from downstream analyses. Where available, RNA-dependent RNA-polymerase protein sequences of these related viruses were then used to construct phylogenetic trees. Multiple sequence alignment was done using the M-Coffee mode in T-Coffee. Phylogenies were estimated using PhyML with LG substitution model and
nearest neighbour interchanging during the tree search. We identified numerous novel viruses that clustered within the Mitoviridae in the phylogenetic tree, and these were excluded as they may have been infecting other organisms such as yeasts and were mostly uncommon.

**Viral abundance in RNAseq data**

Trimmed RNA reads from the 40 libraries were mapped to the same sequences as before (Drosophila melanogaster, Wolbachia and Drosophila related viruses, with all published sequences included) combined with the new viral contigs assembled from this population. Again, mapping was performed using STAR-2.6.0 with default settings (Supplementary Figure 4; round.2 mapping). We counted the reads mapping to D. melanogaster, Wolbachia and each virus. Multiple mapped reads were counted once to a randomly selected mapped location.

The count of reads mapping to Grom virus (D. obscura) and Machany virus (D. obscura) read counts were positively correlated with that of their close relatives, respectively Motts Mill virus (D. melanogaster) and Kilifi virus (D. melanogaster) 22, suggesting miss-mapping (Supplementary Figure 2). Therefore, Grom virus and Machany virus read counts were reclassified into their respective relatives. Twyford virus was excluded from analyses as it is likely a virus of the fungal pathogen Entomophthora muscae 51. Drosophila immigrans sigmavirus (DImmSV), which infects approximately 38% of D. immigrans flies 52, was excluded as there was evidence to suggest low levels of D. immigrans contamination in the RNAseq libraries, and the count of D. immigrans mitochondrial COI reads was positively correlated with the count of DimmSV reads (Supplementary Figure 2). Contamination could have arisen in the field, during collection or in the laboratory. In the most heavily contaminated library, the number of reads mapping to D. immigrans COI was less than 0.2% of the number of COI reads mapping to D. melanogaster.

We used our PCR data (see below) to identify pairs of contigs that were likely segments of the same viral genome. First, there was a strong correlation between the abundance of a new viral contig we identified and Vera virus ($r=0.99$, $p<10^{-10}$) (Supplementary Figure 2A), suggesting these are two segments of the same Partitiviridae genome. The abundance of Galbut virus and Chaq virus were also strongly correlated ($r=0.41$, $p<10^{-10}$), but in this case many flies were infected with Galbut but not Chaq. This agrees with previous data suggesting Chaq virus is either a satellite virus of Galbut virus or an ‘optional’ segment of the Galbut virus genome 21. We therefore refer to this sequence as Galbut (Chaq) virus.

**Virus prevalence**

Quantitative PCR (qPCR) was used to determine the presence and load of each virus in each sampled fly. Primers were designed in Primer-BLAST, which uses the Primer3 and BLAST, setting Drosophila melanogaster as the organism to check specificity 53,54. For virus primer design, we used out RNAseq data to ensure there was no polymorphism in the first 5 bp in the 3’ end of each primer 55. A degenerate base was used when a polymorphism was present elsewhere in the primer region with a minor allele frequency over 10%. No more than one degenerate site per primer was allowed. The efficiency with which each primer amplified viral RNA was estimated using a serial dilution of template cDNA. The complete list the primers, their efficiency, and the amplified product size can be found in Supplementary
Table 1. To verify results for Vera and Motts Mill viruses we repeated the PCR tests of all the samples using an independent set of primers (Supplementary Figure 2). Amplifications by qPCR were carried out with primer at a final concentration of 0.25 μM, using SensiFAST SYBR Hi-ROX master mix (Bioline) and 2 μL of a single-fly cDNA in a total volume of 10 μL. Reactions were performed in 96 well plates, including in each run six positive controls using cDNA library used in RNAseq as template and two template-free negative controls. The reactions were done using a StepOnePlus Real-Time PCR System in the following conditions: 95 °C for 2 min, 40 cycles at 95 °C for 5 s, 60 °C for 30 s. The product of reaction was submitted to melting curve analysis to check the target-specific amplification, and samples where the melting curve was anomalous were discarded. To calculate relative viral load, we used the amplification of the host transcript \( RpL32 \) (see above). Because primers for the viruses and the endogenous genes have approximate similar efficiencies, we calculated viral titer from the cycle thresholds \( (Ct) \) as \( 2^{ΔCt} \), where \( ΔCt = Ct_{RPL32} - Ct_{virus} \).

Statistics and reproducibility

The effect of Wolbachia on the probability that flies were infected by viruses was estimated using a generalised linear mixed model implemented using the \( R \) package \( MCMCglmm \) \(^{56} \), which uses Bayesian Markov chain Monte Carlo (MCMC) techniques. The binary response variable was whether or not a single fly tested positive for a given virus, which was treated as a binomial response with a logit link function. The model included a single fixed effect—whether or not a fly was infected with Wolbachia. The first random effect in the model was the identity of the individual fly being tested. The second random effect was the identity of the virus being tested for. For this random effect, separate variances were estimated for Wolbachia-infected and Wolbachia-free flies, and the covariance was set to zero (specified as ‘idh(wolbachia):virus’ in \( MCMCglmm \)). We used inverse Wishart priors \( (V=1, nu=0.002) \). We estimated the prevalence of viruses in Wolbachia-infected and Wolbachia-free flies from the random effects of the model, and these estimates were transformed from the logit scale back into proportions. Credible intervals were obtained as the 95% highest posterior density of these random effects. To investigate if there was an effect of Wolbachia on flies being infected with a given virus, we calculated the proportion of samples from the MCMC chain where the viral prevalence in Wolbachia-infected samples is less than the prevalence in Wolbachia-free samples. The risk ratio was estimated by dividing the random-effects estimate of the prevalence in Wolbachia-infected flies by the estimate in Wolbachia-free flies for each sample from the MCMC chain, and then calculating the mean (posterior mean) and 95% highest posterior density (95% credible interval) of these numbers.

Data availability statement

The RNAseq data has been submitted to the NCBI Sequence Read Archive under the BioProject number PRJNA728554. The assembled contigs of novel \( D. \ melanogaster \) associated viruses are available in GenBank (MZ852356 to MZ852369). The data underlying Figure 1 and 2 is available in Supplementary Data 1 (figure 1), Supplementary Data 2 (virus prevalence), Supplementary Data 3 (risk ratios) and Supplementary Data 4 (viral load).
Code availability statement

The code used for the bioinformatic analysis is available on the Github Repository at https://doi.org/10.5281/zenodo.5525968.

Figure captions

Figure 1. Viruses associated with wild D. melanogaster. The total number of RNAseq reads that map to each virus (left). The prevalence of selected viruses estimated using quantitative PCR to test single flies for infection (right). Error bars are 95% confidence intervals.

Figure 2. Viral prevalence and load in Wolbachia-free and Wolbachia-infected flies. (a) The prevalence of viruses in male D. melanogaster. The bars are the posterior means of the random effect estimates of a glm. The p values are posterior probabilities that the prevalence differs in Wolbachia-free and Wolbachia-infected flies, estimated from the glm. (b) The risk of viral infection in Wolbachia-free flies relative to Wolbachia-infected flies. Values above 1 indicate that Wolbachia-free flies are more likely to be infected. The points are posterior means and the error bars are 95% credible intervals estimated from a glm. (c) Viral load of Galbut virus in flies with and without Wolbachia. Viral load is measured by quantitative PCR relative to the Rpl32 mRNA. The P-value is the result of a one-way ANOVA.

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Author Contributions

RC and FMJ designed the study. RC, ACP and JPD collected the data, SDD, FMJ and RC analysed the data. FMJ and RC wrote the manuscript with inputs from all other authors.
Competing interests

The authors declare no competing interests.
Known D. melanogaster virus

- Vera virus
- La Jolla virus
- Drosophila A virus
- Craiges Hill virus
- Galbut virus
- Motts Mill virus
- Nora virus
- Dansoman virus
- Thika virus
- Galbut (Chaq) virus
- Kilifi virus

Drosophila melanogaster sigmavirus

- Bloomfield virus
- Vesanto virus

Drosophila associated picorna–like virus

- Brandeis virus

Drosophila associated tymovirus

- Kallithea virus
- Linvill Road virus

Drosophila associated narnavirus 1

Drosophila associated narnavirus 2

Drosophila associated anphevirus

Drosophila associated totivirus 2

Drosophila associated totivirus 3

Drosophila associated narnavirus 4

Drosophila associated narnavirus 5

Drosophila associated totivirus 4

Drosophila associated totivirus 5

Drosophila associated narnavirus 3

Novel D. melanogaster virus

- Motts Mill virus
- Galbut virus
- Craigie’s Hill virus
- Drosophila associated flavivirus 7
- Drosophila associated flavivirus 8
Cogni et al. report a protective effect of *Wolbachia* on viral infection of a natural *Drosophila* population through metagenomics sequencing of the virome. This study provides new evidence showing that *Wolbachia* confers viral resistance in natural population of flies.