Population Genomics of Infectious and Integrated *Wolbachia pipientis* Genomes in *Drosophila ananassae*

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**Abstract**

Coevolution between *Drosophila* and its endosymbiont *Wolbachia pipientis* has many intriguing aspects. For example, *Drosophila ananassae* hosts two forms of *W. pipientis* genomes: One being the infectious bacterial genome and the other integrated into the host nuclear genome. Here, we characterize the infectious and integrated genomes of *W. pipientis* infecting *D. ananassae* (wAna), by genome sequencing 15 strains of *D. ananassae* that have either the infectious or integrated wAna genomes. Results indicate evolutionarily stable maternal transmission for the infectious wAna genome suggesting a relatively long-term coevolution with its host. In contrast, the integrated wAna genome showed pseudogene-like characteristics accumulating many variants that are predicted to have deleterious effects if present in an infectious bacterial genome. Phylogenomic analysis of sequence variation together with genotyping by polymerase chain reaction of large structural variations indicated several wAna variants among the eight infectious wAna genomes. In contrast, only a single wAna variant was found among the seven integrated wAna genomes examined in lines from Africa, south Asia, and south Pacific islands suggesting that the integration occurred once from a single infectious wAna genome and then spread geographically. Further analysis revealed that for all *D. ananassae* we examined with the integrated wAna genomes, the majority of the integrated wAna genomic regions is represented in at least two copies suggesting a double integration or single integration followed by an integrated genome duplication. The possible evolutionary mechanism underlying the widespread geographical presence of the duplicate integration of the wAna genome is an intriguing question remaining to be answered.

**Key words:** lateral gene transfer, horizontal gene transfer, whole-genome sequencing, endosymbiont.

**Introduction**

The alpha-proteobacteria *Wolbachia pipientis* (Lo et al. 2007) is an intracellular bacterium that infects both arthropods and filarial nematodes (Werren et al. 2008). *Wolbachia pipientis* is an obligate symbiont in filarial nematodes; its elimination results in sterility or lethality of the nematodes (Taylor et al. 2005; Slatko et al. 2010). On the other hand, the symbiosis between *W. pipientis* and its arthropod hosts is mainly facultative (but see Hosokawa et al. 2010; Nikoh et al. 2014 for exceptions) and in most cases *W. pipientis* is parasitic toward its arthropod hosts. The parasitism causes harmful fitness consequences to its host mainly by controlling the host reproductive processes for its own benefit (Stouthamer et al. 1999).

Within a single species transmission of *W. pipientis* is usually maternal (mother to offspring) which requires the endosymbiont to localize within the host germline to be successfully transmitted to the next generation (Serbus et al. 2008). Due to the extensive association *W. pipientis* has with its host germline cells, opportunities of lateral gene transfer (LGT) are predicted to occur between the host eukaryotic genome and the endosymbiotic bacterial genome. LGT (sometimes referred as horizontal gene transfer) is the process of genetic exchange between two evolutionary divergent organisms. LGT between *W. pipientis* and its host has been identified both from the host to the endosymbiont (Woolfit et al. 2009) and from the endosymbiont to the host (Dunning Hotopp 2011).

After LGT, the recipient organism acquires novel genetic material and if the newly acquired genetic material improves the fitness of the recipient species, it will spread through the population ultimately fixing within the species (Long et al. 2013). For example, genes transferred from prokaryotes to eukaryotes have been postulated to be the source of novel genes that gave selective advantage to eukaryotes during
adaptation to novel ecological niches (Keeling and Palmer 2008). However, the efficiency of the eukaryotic transcriptional machinery on a prokaryotic gene is questionable. Most LGT products from prokaryotes into eukaryotes are expected to have minimal functionality in the eukaryotic genome eventually accumulating mutations and becoming pseudogenes. Thus, the evolutionary consequence of these LGT elements is an intriguing question to examine.

Evidence of *W. pipientis* LGT (W\(^L\)\(G\)T) into eukaryotic nuclear genomes, or sometimes referred as nuclear Wolbachia transfers, has been detected in various natural hosts of *W. pipientis* (Dunning Hotopp 2011). W\(^L\)\(G\)T was discovered in the genome sequence of the filarial nematode *Brugia malayi* (Dunning Hotopp et al. 2007) where many of the W\(^L\)\(G\)T genes from *W. pipientis* infecting *B. malayi* (wBm) were found to be degenerating suggesting their nonfunctionality. However, a recent study has found as many as 227 W\(^L\)\(G\)T genes and genetic fragments with high sequence similarity to the wBm genome scattered across the host *B. malayi* genome (Ioannidis et al. 2013), and some of these W\(^L\)\(G\)T were hypothesized to retain functionality in the new eukaryotic genome based on the detection of transcripts in different developmental stages of *B. malayi*. There are other examples of W\(^L\)\(G\)T in nematodes, such as in the parasitic nematode *Onchocerca volvulus*, where the W\(^L\)\(G\)T predates the speciation with its sister species *Onchocerca ochengi* (Fenn et al. 2006).

In arthropods, W\(^L\)\(G\)T has been discovered in beetles (Kondo et al. 2002; Aikawa et al. 2009), fruit flies (Dunning Hotopp et al. 2007), mosquitoes (Klasson, Kambris, et al. 2009), parasitoid wasps (Werren et al. 2010), and the tsetse fly (Doudoumis et al. 2012; International Glossina Genome Initiative 2014). The W\(^L\)\(G\)T in nematodes is characterized by small genetic fragments scattered across the host genome (Ioannidis et al. 2013). In contrast, arthropod W\(^L\)\(G\)T is characterized by both large and small segments of genomic DNA integrated into the host genome. For arthropods, the functional consequences of these W\(^L\)\(G\)T products are debatable. Transcription of some of the W\(^L\)\(G\)T genes was detected by quantitative and reverse transcription polymerase chain reaction (PCR), albeit at very low expression levels compared with control genes (Dunning Hotopp et al. 2007; Nikoh et al. 2008). In addition, a recent RNA sequencing study detected only a few W\(^L\)\(G\)T originating reads out of the total *Drosophila ananassae* RNA pool (Kumar et al. 2012), suggesting that the observed gene expression could be background transcriptional noise. Thus, with potential nonfunctionality, these W\(^L\)\(G\)T elements were suggested as transient evolutionary phenomena that are ultimately decaying to become noncoding junk DNA in the host genome (Blaxter 2007). Indeed, in the case of the beetle *Callosobruchus chinesis*, many of the W\(^L\)\(G\)T elements were found degenerated and pseudogenized (Nikoh et al. 2008). However, evidence of LGT from other bacterial endosymbionts has shown that the integrations of single genes or operons have become functional in host genomes, such as aphids (Nikoh et al. 2010), mealy bug (Husnik et al. 2013), and psyllids (Sloan et al. 2014).

The vinegar fly *D. ananassae* shows the most extreme case of W\(^L\)\(G\)T; the whole genome of *W. pipientis* infecting *D. ananassae* (wAna) was estimated to have integrated into its host genome (Dunning Hotopp et al. 2007). Further, by sequencing several wAna loci, Choi and Aquadro (2014) have shown that the integrated wAna (wAna\(^T\)\(G\)T) genome exists in *D. ananassae* lines sampled from much of the host’s geographic range. The study also noted that both infectious wAna (wAna\(^R\)\(G\)T) and wAna\(^T\)\(G\)T genes were highly similar to each other for the regions examined, suggesting that the wAna invasion and whole-genome integration might be relatively recent. However examination of the host *D. ananassae* mitochondriod DNA (mtDNA) variation, which is in complete linkage disequilibrium with *W. pipientis* due to their shared maternal inheritance (Hurst and Jiggins 2005), revealed the original wAna invasion more likely originated in the ancestral population of *D. ananassae* (Choi and Aquadro 2014). These results illustrated that sequencing of only a few loci does not reveal sufficient variation for resolving the joint evolution of wAna\(^R\)\(G\)T and wAna\(^T\)\(G\)T genomes within *D. ananassae*. However, studies utilizing whole-genome sequencing have been fruitful in elucidating the coevolutionary and population genomics of *W. pipientis* infecting *Drosophila melanogaster* (wMel) with its host (Richardson et al. 2012; Chrostek et al. 2013; Early and Clark 2013), suggesting that whole-genome sequencing is a better method for studying of *W. pipientis* genomics.

Here, we have described the evolution of the wAna\(^R\)\(G\)T and wAna\(^T\)\(G\)T genomes by whole-genome sequencing host *D. ananassae* flies that harbor either the wAna\(^R\)\(G\)T or the wAna\(^T\)\(G\)T genome. Using well-established computational pipelines, reads originating from the wAna genomes were separated out for further analysis. The wAna\(^R\)\(G\)T genome comparisons, with additional data from mitochondrial genomic variation, reveal that wAna\(^R\)\(G\)T is stably maternally transmitted by its *D. ananassae* host corroborating the findings of Choi and Aquadro (2014). In addition, we have discovered several nucleotide substitutions and large structural variations that distinguish several wAna\(^R\)\(G\)T genome types. In contrast, the wAna\(^T\)\(G\)T genome is pseudogene-like, accumulating large numbers of highly deleterious mutations. Interestingly, analysis of single nucleotide and copy number variation indicates that the majority of the wAna\(^T\)\(G\)T regions has a minimum of two copies integrated into the *D. ananassae* genome consistent with independent results reported recently for two lines of *D. ananassae* by Klasson et al. (2014). The whole-genome integration and duplication of the wAna genome are surprising due to its widespread geographic occurrence despite its nonfunctional pseudogenome like characteristics.
Materials and Methods

**Genome-Sequenced D. ananassae Strains**

All D. ananassae strains examined in this study originated from the study of Choi and Aquadro (2014). Eight strains with evidence of the wAnaINF genome (Cebu, GB1, HNL0501, KMU1, OGS-98K1, RC102, TBU136, and VAV150) and seven strains with evidence of only the wAnaITG genome (BKK13, D38, EZ104, PNP1, TB43, TBU3, and T18) were examined (fig. 1). Each D. ananassae strain that was positive for wAna genes by PCR was treated with the antibiotic tetracycline with concentrations of 200 μg/ml to cure them of wAnaINF. Then, the cured D. ananassae strains were screened by PCR again for the wAna gene and from here we defined a D. ananassae strain to have the wAnaITG genome if no wAna genes were detected after curing, and as a strain having the wAnaITG genome if wAna genes were detected after curing. As it is possible for the D. ananassae strains with the wAnaITG genome to also have the wAnaINF genome, we avoided sequencing both genomes by only genome sequencing cured wAnaITG carrying D. ananassae strains.

For each strain, DNA was extracted from whole bodies of 12–15 female D. ananassae flies using the Qiagen DNEasy blood and tissue kit. To prepare for genome sequencing, the purified DNA were processed by the Biotechnology Resource Center Genomics Facility at Cornell University (http://www.biotech.cornell.edu/biotechnology-resource-center-brc, last accessed August 2015) using Illumina TruSeq DNA sample library prep kit as paired-end 2 × 150 bp samples. DNA libraries were barcoded and pooled for whole-genome sequencing using the Illumina HiSeq 2500 rapid run mode at the Biotechnology Resource Center Genomics Facility at Cornell University.

Reference Genome-Based Realignment and Alignment Statistics Calculation

The program Trimmomatic version 0.32 (Bolger et al. 2014) was used to preprocess the raw reads to trim any remaining adapter sequences and for quality control purposes (Parameters used: ILLUMINACLIP:2:30:10:5, LEADING:3, TRAILING:3, MINLEN:70, and SLIDINGWINDOW:4:15). The Program FastQC version 0.11.1 (http://www.bioinformatics.bbsrc.ac.uk/projects/fastqc, last accessed August 2015) was then used to heuristically check and visualize the quality of the raw reads.

The quality-controlled raw reads were then realigned to a reference genome using the program BWA-MEM version 0.7.10 (Li 2013) using default parameters. Currently, a high-quality wAna genome is unavailable, so we used the reference genome sequence of W. pipientis infecting Drosophila simulans strain Riverside (wRi) (Klasson, Westberg, et al. 2009). A previous study has shown high nucleotide identity between the genomes of wAna and wRi (Salzberg et al. 2005). The D. ananassae reference mitochondria genome (GenBank accession number BK006336.1; Montooth et al. 2009) was used to extract D. ananassae mitochondrial reads.

Genome-wide coverage was calculated by counting the number of reads aligned on a given site using the program genomeCoverageBed from the bedtools version 2.17.0 suite (Quinlan and Hall 2010). Total number of reads aligning to each reference genome was calculated using the program saamtools version 1.0 (Li et al. 2009). For all wAna samples, read coverage per site was normalized against the mean read coverage of the longest scaffold assembled from the draft D. ananassae nuclear genome (scaffold 13340; GenBank accession number CH902617.1).

wAnaINF Copy Number Variation Analysis

The program CNVnator version 0.3 (Abyzov et al. 2011) was used to detect regions with significant changes in the copy number of the wAnaINF-aligned BAM files. CNVnator divides the whole genome into bins to calculate the mapping density and find regions with significantly different read depth. For the wAnaINF sample, genome-wide coverage was equal for most of the regions except for the few candidate regions with variation in the copy number (see Results section) so that CNVnator was appropriate to detect those copy number variations. In contrast, the wAnaITG samples had varying genome coverage throughout the genome making CNVnator an unsuitable method; thus, we used experimental methods to directly quantify the differences in the genome coverage (see below).

Quantitative PCR to Analyze wAnaITG Genome Copy Number

To verify the variable read coverage observed in the wAnaITG genomes (see Results section), quantitative PCR (qPCR) was conducted on three D. ananassae strains with the wAnaITG genome (BKK13, D38, and T18). Based on our wAnaITG genome coverage results, a low coverage region (LOW) corresponding to gene YP_002726693.1 (coordinate: 40,653–41,915) and high coverage region (HIGH) corresponding to gene YP_002726693.1 (coordinate: 998,610–1,000,511) were selected as candidate regions and compared with the D. ananassae rp49 gene. qPCR primers were designed using the online version of the program primer3plus (Untergasser et al. 2012; http://primer3plus.com/cgi-bin/dev/primer3plus.cgi, last accessed August 2015), and primer sequences are provided in supplementary table S1, Supplementary Material online. qPCR reactions were conducted using iTaq Universal SYBR Green Supermix (Biorad), and fluorescence was detected using the ViIA7 Real-Time PCR system (Life Technology). PCR mixes were denatured at 95 °C for 10 min followed by a 40 cycle amplification stage, which consisted of 95 °C for 15 s followed by a data collection step of 53 °C for 1 min. A melt curve stage was conducted at the end with 95 °C
**Fig. 1.**—Geographic origin of samples analyzed in this study. *Drosophila ananassae* strains with wAna\textsuperscript{INF} genomes are indicated with black colored stars and the localities are Cebu (Cebu, Philippines), GB1 (Mauritius), HNL0501 (Oahu, USA), KMJ1 (Kumejima, Japan), OGS-98K1 (Ogasawara, Japan), RC102 (Rwanda), TBU136 (Tonga), and VAV150 (Vava’u, Tonga). *Drosophila ananassae* strains with wAna\textsuperscript{ITG} genomes are indicated with red colored stars and the localities are BKK13 (Bangkok, Thailand), D38 (Coimbatore, India), EZ104 (Ethiopia), PNP1 (Phnom Pen, Cambodia), TB43 (Trinity Beach, Australia), TBU3 (Tonga), and TI8 (Thursday Island, Australia).
Single females from each of the three examined *D. ananassae* strains (BKK13, D38, and T18) were combined and extracted for DNA using our previous methods of DNA extraction. This master mix was then serially diluted to fit a standard curve for the three genes (LOW, HIGH, and rp49) and estimate the efficiency of the primers. This standard curve was then used to convert the observed C_t values for each sample into its relative concentration. Afterwards, the HIGH and LOW relative concentrations were compared with the relative concentration of rp49. Three biological replicates and three technical replicates were conducted for each *D. ananassae* strains. All qPCR reactions were conducted in the same plate to keep the reaction condition and detection of fluorophores consistent.

Detection of Nucleotide and Structural Variations

BAM files generated from the previous reference genome-based raw read realignment step were then processed according to the guidelines of GATK’s BestPractice version 3.3 for downstream analysis (https://www.broadinstitute.org/gatk/guide/best-practices, last accessed August 2015). Briefly, this process involves removing duplicate reads using the program Picard version 1.115 (http://broadinstitute.github.io/picard/, last accessed August 2015), and realigning regions around insertion and deletions (INDELS) using the GATK suite version 3.2.2 (https://www.broadinstitute.org/gatk/, last accessed August 2015). The processed BAM file was then used to call INDEL and single nucleotide polymorphism (SNP) variants using the joint genotyping method of GATK. The program VariantFiltration from the GATK suite was then used to apply hard filters on the raw variant calls (parameters for INDEL filtering: QD < 2.0, FS > 200.0, ReadPosRankSum < −20.0; parameters for SNP filtering: QD < 2.0, FS > 60.0, MQ < 40.0, HaplotypeScore > 13.0, MappingQualityRankSum < −12.5, ReadPosRankSum < −8.0; please see https://www.broadinstitute.org/gatk/gatkdocs/index [last accessed August 2015] for details on parameters). Additionally for the SNP filtering stage, the previous INDEL calls were used to mask SNP calls that coincided with the INDEL position as these may cause false polymorphism calls because of alignment errors. SNP variants detected among our genomes and used for downstream analysis are reported in supplementary data, Supplementary Material online, in a tab-delimited format. Raw VCF files for the SNP and INDELS are available upon request.

To detect small- and large-sized structural variations, specifically INDELS and inversions, we used the program pindel version 0.2.5 (Ye et al. 2009), which uses a pattern growth algorithm to detect structural variations at base pair resolution. After the structural variations were predicted by pindel the pindel2vcf from the pindel suite was used to filter out variants that were predicted with less than four reads as well as any variants due to homopolymer and microsatellite repeats as these could represent sequencing errors (pindel2vcf parameters: -e 4 -ir 2 -il 10 -pr 1 -pl 10).

The functional effects of each SNPs and INDELS were predicted using the program snpEff version 4.0 e (Cingolani et al. 2012).

**Phylogenetic Analysis**

Nucleotide variants that were called using above methods were then used for phylogenetic analysis. Phylogenetic reconstruction was carried out using the maximum-likelihood method implemented in PhyML version 3.0 (Guindon et al. 2010), the HKY (Hasegawa–Kishino–Yano) model of DNA substitution (Hasegawa et al. 1985), and substitution rate variation across sites modeled with a gamma distribution with four rate categories (commonly known as the HKY + G model). Confidence of the inferred phylogenetic tree was estimated using 1,000 bootstrap replicates. Trees were plotted and manipulated using FigTree ver 1.4.0 (http://tree.bio.ed.ac.uk/software/figtree/, last accessed August 2015) and the R package phytools (Revell 2012).

**Preparation of wAna\(^{TG}\) Genome for Phasing Heterozygous Genotypes**

We initially assumed that the wAna\(^{TG}\) genome existed in a diploid state in *D. ananassae* strains with evidence of the integration, and thus attempted phasing sites called as hetereozygotes by first making the cytotype of the wAna\(^{TG}\) genome
haploid. Male *D. ananassae* strains with the wAna\textsuperscript{ITG} genome were mated to virgin females of a *D. ananassae* strain that had neither the wAna\textsuperscript{INF} genome nor wAna\textsuperscript{ITG} genome. Crossing males with the wAna\textsuperscript{ITG} genome to females lacking both wAna\textsuperscript{INF} and wAna\textsuperscript{ITG} genomes prevents any maternal factors (i.e., *W. pipientis*) from being inherited, and only the paternal nuclear DNA will be transmitted to the next generation. Both males and females of the cross were cured of *W. pipientis* with 200 µM tetracycline for four generations to clear wAna\textsuperscript{INF} genomes. We collected only the F1 progeny females in case there were integrations in both the autosome and X chromosome, thus retaining equal coverage between the two different chromosomal backgrounds. Additionally to keep one copy of the haploid wAna\textsuperscript{ITG} genome for downstream phasing analysis, we chose a single virgin female for DNA extraction and subsequent genome sequencing.

Single female flies were individually homogenized with a clean pestle in a lysis buffer (50 mM Tris–HCL ph8.2, 100 mM ethylenediaminetetraacetic acid, 100 mM NaCl, 1% SDS) then treated with Proteinase K and RNase I. DNA extraction was conducted using a standard phenol–chloroform–isoamyl protocol followed by an overnight ammonium acetate DNA precipitation. A detailed protocol is available at request. The purified DNA was then prepared for genome sequencing following the same protocol previously described.

### In Silico Prediction of Breakpoints between wAna\textsuperscript{ITG} and Host Nuclear Genome

In an effort to determine breakpoints between the wAna genomic fragments and the host genome, we treated those breakpoints as translocation events (i.e., exchange of chromosomal DNA between nonhomologous chromosomes) and used the program Lumpy ver 0.2.6 (Layer et al. 2014) to detect the breakpoints between the wAna genome and the host genome. Lumpy combines multiple signals from read-pair, split-reads, and read-depth of a sample to identify structural variations. We realigned our original raw reads to a reference genome that included a subset of the reference *D. ananassae* CAF1 assembly (Clark et al. 2007) and the wRi genome. Only a subset of scaffolds was selected because the *D. ananassae* reference genome is largely a draft genome that includes thousands of unassembled scaffolds. Further, it is possible that some scaffolds contain *W. pipientis* sequences that were accidently incorporated because the sequenced line of *D. ananassae* contained *W. pipientis*. In fact, Salzberg et al. (2005) were able to reassemble a draft wAna genome from the raw sequence repository of the *D. ananassae* reference genome. We selected large scaffolds that were identified as syntenic to the *D. melanogaster* Muller elements (Schaefter et al. 2008) (see supplementary table S3, Supplementary Material online, for full list of scaffolds; GenBank assembly accession number: GCA_000005115.1). Using this custom reference genome, paired-end reads that contain both the bacterial and host sequences were aligned to their appropriate genome. Next, discordant paired-end and split-end reads were extracted from the BAM files using samtools. These reads were then processed using the pairend_distro.py script from the Lumpy suite to estimate the library size, mean, and standard deviation. The sample statistics of the library were then prepared as a configuration file with the following Lumpy parameters to detect translocation events using two different libraries: 1) Discordant paired-end library analysis: Read length = 150; min nonoverlap = 150; discordant z = 4; back distance = 20; weight = 1; id = 1; min mapping threshold = 40; and 2) split reads library analysis: Back distance = 20; weight = 1; id = 2; min mapping threshold = 40. Inferred breakpoints between *D. ananassae* and wAna were only used if they were supported by reads with a minimum mapping quality of 40, had evidence from both paired-end and split-end read libraries, and had a minimum of six supporting reads.

Our analysis revealed that Lumpy identified many breakpoints in which the host *D. ananassae* sequence corresponded to potential transposable elements (TEs) (described further in Results). To further characterize these breakpoints, we used the program RepeatMasker version open 4.0 (http://www.repeatmasker.org/, last accessed August 2015; developed by A.F.A. Smit, R. Hubley, and P. Green) to identify repetitive DNA and TEs across the reference *D. ananassae* genome. The *Drosophila* RepBase repeat library Update 20140131 (http://www.girinst.org/repbase/, last accessed August 2015) was used to identify specific TE families.

### Results

#### Raw Read Alignment Statistics

Whole adult flies representing each of the 15 strains of *D. ananassae* that were previously screened for the presence of wAna\textsuperscript{INF} and wAna\textsuperscript{ITG} genomes (Choi and Aquadro 2014) were genome sequenced, and all wAna originating reads were computationally extracted using the reference wRi genome (alignment statistics are shown in table 1). Raw reads from both wAna\textsuperscript{INF} and wAna\textsuperscript{ITG} genomes comprised on average 2% of the total reads. However, the proportion of wAna reads was variable between the two genomes; the wAna\textsuperscript{INF} genomes had more variability among lines, ranging from 1.3% of the total reads in strain HNL0501 to 3.8% in strain TBU136. Mirroring this result, the average read depth (RD\textsubscript{avg}) of the wAna\textsuperscript{INF} genomes, normalized relative to the RD\textsubscript{avg} of the *D. ananassae* nuclear genome, also varied from 1.7-fold higher in HNL0501 wAna\textsuperscript{INF} to 5.2-fold higher in TBU136 wAna\textsuperscript{INF}. For wAna\textsuperscript{ITG}, a single genome integration of wAna would be expected to have an RD\textsubscript{avg} equal to that of the *D. ananassae* nuclear genome. However, the RD\textsubscript{avg} for wAna\textsuperscript{ITG} was roughly 2.6-fold higher.
Alignment statistics for the mtDNA genome indicated 0.3–0.7% of the total raw reads originated from the mtDNA for both D. ananassae strains with the wAnaINF and wAnaITG genomes. Deep genome coverage was observed for the mtDNA, and compared with the D. ananassae nuclear genome, the D. ananassae strains with the wAnaITG genomes had more mitochondrial genome copies than D. ananassae strains with the wAnaINF genomes (table 1). However, this difference is probably due to the tetracycline treatment of wAnaITG carrying D. ananassae to eliminate any residual wAnaINF genomes. A previous study has shown that tetracycline increases mtDNA copy in Drosophila that was not infected with W. pipientis (Ballard and Melvin 2007).

Analysis of wAnaINF and wAnaITG Genome Coverage

The program CNVnator was used to detect regions with significant changes in the copy number among the wAnaINF samples. As expected from the wAnaINF genome coverage plots, no regions with significant changes in the copy number were detected for D. ananassae strains Cebu, GB1, KMJ1, OGS-98K1, and RC102. In the wAnaINF genomes of strains TBU136 and VAV150, regions between coordinates 0.61–0.63 and 1.12–1.14 Mb were estimated to have a 0.5-fold decrease (P < 0.0001) in copy number compared with their genome-wide background coverage. In the HNL0501 strain, the wAnaINF genome was predicted to have a significant 2-fold increase in copy number (P < 0.0001) between coordinates 0.57–0.63 and 1.07–1.14 Mb. The two regions between 0.61–0.63 and 1.12–1.14 Mb that showed copy number variation overlapped among wAnaINF genomes from strains HNL0501, TBU136, and VAV150.

In contrast to the wAnaINF genomes, the wAnaITG genomes showed strikingly heterogeneous coverage across the genome (fig. 3), with low coverage in the beginning of the genome increasing to a maximum coverage around coordinates 1 Mb and subsequently decreasing. Regions with low and high coverage were also correlated across all seven wAnaITG genomes. The normalized read depth across the wAnaITG genomes indicated that the majority of the regions had at least double the read depth compared with the D. ananassae nuclear genome. We verified that the high and low coverage regions of the wAnaITG segments represented actual differences in the genome sequence copy number using qPCR. A candidate

Table 1
Raw Read Realignment Statistics for mtDNA and wAna Genomes

| Strain   | Total Reads | wAna Originating Reads | Proportion of wAna Reads | wAna RDavg | wAna Ratio | mtDNA Originating Reads | mtDNA RDavg | mtDNA Ratio |
|----------|-------------|------------------------|--------------------------|------------|------------|-------------------------|-------------|-------------|
| Raw reads from wAnaINF genome | | | | | | | | |
| BKK13    | 35,914,931  | 721,253                | 0.020                    | 74.4       | 2.5        | 238,401                 | 0.003       | 2,376.8     |
| D38      | 40,631,930  | 987,102                | 0.024                    | 101.8      | 3.1        | 299,310                 | 0.007       | 2,977.6     |
| EZ104    | 29,830,655  | 613,044                | 0.021                    | 63.2       | 2.6        | 216,173                 | 0.007       | 2,150.7     |
| PNP1     | 90,416,482  | 1,766,984              | 0.020                    | 180.3      | 2.4        | 291,938                 | 0.003       | 2,681.3     |
| TB43     | 28,871,004  | 621,913                | 0.022                    | 64.0       | 2.7        | 208,458                 | 0.007       | 2,071.6     |
| TBU3     | 39,325,872  | 1,001,506              | 0.025                    | 103.4      | 3.2        | 292,405                 | 0.007       | 2,908.0     |
| TIB      | 37,948,414  | 853,466                | 0.022                    | 88.0       | 2.8        | 265,647                 | 0.007       | 2,645.9     |
| Median   | 37,948,414  | 853,466                | 0.022                    | 88.0       | 2.7        | 265,647                 | 0.007       | 2,645.9     |
| Raw reads from wAnaITG genome | | | | | | | | |
| Cebu     | 141,458,302 | 2,971,843              | 0.021                    | 304.7      | 2.6        | 528,382                 | 0.004       | 5,262.5     |
| GB1      | 33,180,346  | 640,906                | 0.019                    | 64.3       | 2.6        | 228,571                 | 0.007       | 2,121.2     |
| HNL0501  | 19,121,957  | 243,774                | 0.013                    | 24.9       | 1.7        | 101,483                 | 0.005       | 1,008.5     |
| KMJ1     | 76,690,462  | 1,457,894              | 0.019                    | 149.2      | 2.3        | 208,845                 | 0.003       | 1,894.7     |
| OGS99-K1 | 30,352,531  | 709,801                | 0.023                    | 72.1       | 3.3        | 78,842                  | 0.003       | 784.6       |
| RC102    | 82,433,612  | 1,387,896              | 0.017                    | 141.5      | 2.1        | 246,883                 | 0.003       | 2,258.8     |
| TBU136   | 20,721,322  | 794,974                | 0.038                    | 81.3       | 5.2        | 81,676                  | 0.004       | 811.3       |
| VAV150   | 20,791,631  | 535,975                | 0.026                    | 54.4       | 3.5        | 122,822                 | 0.006       | 1,153.5     |
| Median   | 31,766,439  | 752,388                | 0.020                    | 76.7       | 2.6        | 165,834                 | 0.004       | 1,524.1     |

NOTE.—RDavg, average read depth of a sample; Ratio, value obtained by dividing the mean depth of the wAna or mtDNA genome to the mean depth of D. ananassae nuclear genome (scaffold 13370).
FIG. 2.—Genome-wide read coverage for the eight wAnaINF genomes. Per-site read depth normalized against the D. ananassae nuclear genome (scaffold 13340) coverage is shown against the reference wRi genome coordinate. Note that regions of tRNA and rRNA, comprising 0.48% of the genome, were expected to have spurious reads from other endosymbiotic bacteria, leading to spikes of read coverage and were ignored from subsequent analysis (see text).
Fig. 3.—Genome-wide read coverage for the seven wAna\textsuperscript{fg} genomes. Per-site read depth normalized against the D. ananassae nuclear genome (scaffold 13340) coverage is shown against the reference wRI genome coordinate. Note that regions of tRNA and rRNA, comprising 0.48% of the genome, were expected to have spurious reads from other endosymbiotic bacteria, leading to spikes of read coverage and were ignored from subsequent analysis (see text).
high copy region (HIGH) at coordinate 998,610–1,000,511 and candidate low copy region (LOW) at coordinate 40,653–41,915 were each compared with the *D. ananassae* nuclear gene *rp49* in three representative strains (BKK13, D38, and TI8). Based on genome coverage results compared with the *D. ananassae* *rp49* gene, LOW was expected to have a 1.5-fold increase whereas HIGH was expected to have a 3.5-fold increase in copy number (fig. 4). Qualitatively, the qPCR results were concordant with the genome coverage results: For all three strains, copy number for LOW was 0.8-fold that of the *rp49* gene, whereas HIGH was estimated to be 2.6-fold higher than *rp49*. Thus, the difference in read coverage across the wAnaITG genome was due to differences in the copy number of wAnaITG integrated into the *D. ananassae* nuclear genome.

### Analysis of wAnaINF Genome Variants

Due to the relatively high read depth for both mtDNA and wAna genomes, variant detection programs were used to call SNPs, INDELs, and inversion variations for wAnaINF both within and among strains of *D. ananassae*. The program HaplotypeCaller from the GATK suite was used to call single nucleotide variants. This program assumes diploidy of the organism when calling each variants genotype. However, wAnaINF genomes are expected to be haploid, and most of the variants are predicted to be called as homozygous genotypes within a strain. Sites called as heterozygous within a single strain could be due to infection by multiple wAna genomes, a duplicated region differing in one duplicate by a mutation, or a false variant call from sequencing error. Levels of variation among the eight wAnaINF genomes were assessed as follows: A site was considered monomorphic when all eight individuals have the same base call and polymorphic when the base calls for at least one line differ from the others. Polymorphic sites that had at least one line with a heterozygote call were also tabulated and were abbreviated as PSHet.

From the total 1,445,873-bp reference wRi genome, base calls were made for 1,194,063 bp (82.6%) for the sample of wAnaINF genomes (table 2). Initially base calls from the wAnaINF genomes were compared with the approximately 7,000-bp regions that Choi and Aquadro (2014) reported as monomorphic using Sanger sequencing, and we verified those regions to be monomorphic in our new data as well. Compared with the reference wRi genome from *D. simulans*, we detected ten fixed differences for the wAnaINF genomes suggesting that these are wAnaINF-specific mutations.

![FIG. 4](https://example.com/fig4.png)

**FIG. 4.**—Computational and qPCR estimates of copy number for the LOW and HIGH regions of the wAnaITG genome. Relative copy number of LOW and HIGH region is compared with *D. ananassae* *rp49*. The x axis represents read depth of LOW and HIGH region divided by read depth of *rp49*. The y axis represents relative concentration of LOW and HIGH region divided by relative concentration of *rp49*.
homologous regions of the reference genome and cause false heterozygote calls (as was observed in the rRNA and tRNA regions). We conservatively chose to discard PS\textsubscript{HET} sites that were clustered within 10-bp windows among the wAna\textsuperscript{INF} genomes. Among the eight wAna\textsuperscript{INF} genomes, there were 125 polymorphic sites where eight polymorphic sites had at least one line of wAna\textsuperscript{INF} called as a heterozygote. Among these eight PS\textsubscript{HET}, seven of the eight polymorphic sites had a single line called as a heterozygote whereas the remaining lines were called as the reference allele. This suggested that those lines with the heterozygote calls were likely to be false variant calls. A single PS\textsubscript{HET} had five of the eight lines called as a heterozygote but this was located within a pseudogenized gene (WRi_p10660). Thus, we infer that heteroplasmy was minimal and that only a single strain of wAna infected each D. ananassae strain. The average genome-wide number of pairwise nucleotide differences per site ($\pi$) was $3.65 \times 10^{-5}$ among the eight wAna\textsuperscript{INF} genomes.

The program pindel was used to identify small-sized INDELs (<100 bp) across each wAna\textsuperscript{INF} genome. Among the eight wAna\textsuperscript{INF} genomes, we detected a total of ten polymorphic sites with small-sized INDELs (table 2). Four of these polymorphic sites were a PS\textsubscript{HET}; however, for each PS\textsubscript{HET} the line with the heterozygote call had an INDEL with a single base pair deletion. This suggested that sequencing errors could have caused the false variant calls in those lines with a site called as a heterozygous INDEL.

We also used pindel to detect large structural variations, such as large deletions (>100 bp) and inversions, that were both polymorphic and fixed (compared with the reference WRi genome) among the wAna\textsuperscript{INF} genomes (supplementary tables S4 and S5, Supplementary Material online). The sizes of these predicted large deletions ranged from 619 to 2,544 bp. Examining the gene annotations of these large deletions, most involved the loss of a TE that existed in the reference wRi genome. Interestingly some of the inversions had predicted breakpoints that coincided with the same genome coordinates as for the large deletions. When the program lumpy was used to detect structural variations, no significant inversions or large deletions were found.

We designed PCR primers that flanked the breakpoints found by pindel and used PCR amplification to verify the computationally predicted structural variations (see supplementary table S2, Supplementary Material online, for genome coordinates of the PCR primers). Nine of the predicted large structural variations were PCR amplified and results are shown in figure 5A. There were seven large structural variations (SV1–SV7) that were polymorphic among the eight wAna\textsuperscript{INF} samples. The remaining two large structural variations (SV8 and SV9) were deletions of a TE in the reference wRi genome, and this deletion was observed in each of our wAna\textsuperscript{INF} samples as the PCR product was smaller than predicted from the wRi reference genome. When compared with the computational prediction, six of the large deletions detected by pindel (SV2, SV3, SV4, SV7, SV8, and SV9) were verified by PCR to be true large deletions. The three inversions computationally inferred by pindel (SV1, SV5, and SV6; supplementary table S2, Supplementary Material online) were examined by trying to PCR amplify one side of the predicted breakpoints. Results showed that these pindel-predicted inversions were not inversions but in fact were insertions of a large genetic sequence, likely from a TE.

### Analysis of wAna\textsuperscript{TG} Genome Variants

In preparation of this manuscript, Klasson et al. (2014) reported evidence of extensive duplication of the wAna\textsuperscript{TG} genome from two samples of D. ananassae originating from Hawaii, the United States, and Mumbai, India. A third sample from Java, Indonesia, on the other hand, had no evidence of duplications for the wAna\textsuperscript{TG} genome. Further, they have conducted in situ hybridization experiments to determine the integration site of wAna\textsuperscript{TG} genome and suggested its localization to the fourth chromosome of D. ananassae (see “Investigating the integration site of the wAna\textsuperscript{TG} genome” for our computational analysis of verifying the fourth chromosome as the site of integration). Compared with Klasson et al. (2014) we have sampled from a wider geographic distribution and discovered that all of our samples had evidence of increased copy number throughout the wAna\textsuperscript{TG} genome (figs. 3 and 4). As Klasson et al. (2014) did not examine the nucleotide or structural variations of the wAna\textsuperscript{TG} genome, we next analyzed the genome variants existing in our sample of wAna\textsuperscript{TG} genomes.

Using HaplotypeCaller to call variant base calls for each wAna\textsuperscript{TG} genome of D. ananassae could result in a site being called as a heterozygote either because the two allelic copies of a specific wAna\textsuperscript{TG} sequence differ, or because the two duplicated regions on the same chromosome differ from each other (much like paralogs of a single gene might differ). Thus, among the wAna\textsuperscript{TG} genomes we also examined PS\textsubscript{HET} carefully.

Analysis of levels of nucleotide variation among the seven wAna\textsuperscript{TG} genomes compared with the eight wAna\textsuperscript{INF} genomes (table 2) revealed both greater differentiation from the wRi reference genome and more variation among wAna\textsuperscript{TG} genomes. Analyzing a total of 1,228,381 bp (85.0%), the wAna\textsuperscript{TG} genomes had 42 fixed differences from the reference wRi genome (compared with only 10 for wAna\textsuperscript{INF}).

Like the variant detection from the wAna\textsuperscript{INF} genomes, we took a conservative approach to filter out potentially false variants called as heterozygotes by examining each wAna\textsuperscript{TG} genome using a 10-bp sliding window, and masking windows with more than three variant base calls. These windows were considered as variant clusters likely due to misalignments and were thus ignored for downstream analysis. After our hard filtering we found 2,259 polymorphic sites among the 7 wAna\textsuperscript{TG} genomes, with 2,234 (98.9%) of these being PS\textsubscript{HET}
The remaining 25 polymorphic sites had a singleton variant. In contrast to results for 7 kb of the wAnaITG genome found to be invariant by Sanger sequencing (Choi and Aquadro 2014), our whole-genome analysis revealed a much higher number of polymorphic sites in other regions of the genome; this contrast will be addressed in the Discussion section.

The seven wAnaITG genomes had a higher number of polymorphic sites with small-sized INDELs (<100 bp) compared with the eight wAnaINF genomes. Among the seven wAnaITG genomes the majority of the INDEL polymorphic sites was a PSHET (114 of 122 polymorphic sites). However, for each PSHET the line with the heterozygous INDEL call had a variant of size 13 bp on average. This suggested that in contrast to the heterozygous INDEL calls from wAnaINF genomes, those in wAnaITG were not false INDEL calls from sequencing errors.

Among the wAnaITG genomes, a total of ten large deletions (>100 bp) were detected using Pindel and all were fixed between the wAnaITG samples and the wRi reference genome (supplementary table S4, Supplementary Material online). All of these large deletions were the same variants that were predicted in our wAnaINF genome analysis (fig. 5A). We verified these large deletions in the wAnaITG samples using the same primers from the PCR analysis of the wAnaINF large structural variation. Results showed that unlike the wAnaINF genomes, there were no variations in large deletions among the seven wAnaITG genomes (fig. 5B). Further, PCR results showed no large INDEL that was able to differentiate the wAnaINF or wAnaITG genomes and all large INDEL variants...
among the wAna\textsuperscript{ITG} genomes existed within the wAna\textsuperscript{INF} genomes (fig. 5A and B).

Analysis of Mitochondrial Genome Variants

For the \textit{D. ananassae} mitochondrial genomes, we analyzed 14,905 bp (99.9%) of the total 14,920 bp of the \textit{D. ananassae} mtDNA genome (table 2). We discovered a total of 165 polymorphic sites, each with a single nucleotide variant among the mitochondrial genomes from both \textit{D. ananassae} with the wAna\textsuperscript{INF} and wAna\textsuperscript{ITG} genomes. We compared our computationally called polymorphic sites with the Sanger-sequenced mtDNA polymorphic sites of Choi and Aquadro (2014), and were able to verify 34 of the 35 segregating sites suggesting a high true positive rate of variant calling for our new results. Out of the total polymorphic sites, there were 11 PSHET where the lines called as a heterozygote were all from individuals with the wAna\textsuperscript{INF} genomes. However, all 11 PSHET were from only a single line called as a heterozygote suggesting that those sites were called as heterozygotes due to sequencing errors. Compared with the reference \textit{D. ananassae} mitochondrial genome, individuals with the wAna\textsuperscript{ITG} genomes had 25 fixed mtDNA differences whereas individuals with the wAna\textsuperscript{INF} genomes had 5 fixed differences; thus, the reference mitochondrial genome was more similar to the mtDNA haplotypes associated with the wAna\textsuperscript{INF} genomes. \textit{Drosophila ananassae} strains with only the wAna\textsuperscript{ITG} genomes had twice as many singletons as individuals with only the wAna\textsuperscript{INF} genomes, whereas the average mitochondrial genome-wide $\pi$ across all \textit{D. ananassae} strains was 2.84 $\times$ 10\textsuperscript{-3}. No significant small- and large-sized INDELs or inversions were detected in the mtDNA.

Functional Classification of Polymorphic Site Variants among wAna\textsuperscript{INF} and wAna\textsuperscript{ITG} Genomes

We used the program snpEff to infer functional consequences of the single nucleotide and small INDEL variants observed among the wAna\textsuperscript{INF} and wAna\textsuperscript{ITG} genomes relative to the reference wRi genome (table 3). We found 52.8% of the polymorphic single nucleotide variant sites among the wAna\textsuperscript{INF} genomes segregated for nonsynonymous variants, all of which resulted in missense mutations. The remaining 47.2% of the single nucleotide polymorphic sites segregated variants of synonymous or intergenic sites. In contrast, among the wAna\textsuperscript{ITG} genomes, 61.1% of polymorphic sites segregated for single nucleotide variants resulting in nonsynonymous changes, with 56.3% of those variants being missense mutations and 4.8% of those variants being nonsense mutations.

For small-sized INDELS, we observed ten polymorphic sites among the wAna\textsuperscript{INF} genomes (table 2) with 9 out of 10 of them occurring in intergenic regions. A single polymorphic site occurred in the coding region with a variant that caused an in-frame deletion (table 3). Among the wAna\textsuperscript{ITG} genomes there were 122 polymorphic sites with small-sized INDELS (table 2), 60.5% of which occurred within a coding region. Of those, 54.9% had a variant resulting in a frameshift mutation and 15.6% had a variant causing an in-frame INDEL change (table 3). Thus, the wAna\textsuperscript{ITG} genomes had an increased number of polymorphic sites with single nucleotide variants and small-sized INDELS that were predicted to cause a strongly deleterious effect (i.e., nonsense mutation and frameshift mutations) in a functioning wAna genome.

Phylogenomic Analysis of mtDNA and wAna Genomes

The single nucleotide variants observed for mtDNA and wAna were used to estimate the phylogeny of each genome we sampled, using a maximum-likelihood method from the program PhyML. The phylogenies of both the host \textit{D. ananassae} mtDNA genome as well as wAna were examined because, like the \textit{W. pipientis} genome, mtDNA is also maternally inherited yet has a higher mutation rate which provides more phylogenetically informative mutations for analysis (Richardson et al. 2012; Early and Clark 2013; this study). Many methods of phylogenetic reconstruction, however, are not suited to deal with heterozygous sites (Sota and Vogler 2003), which is problematic for the wAna\textsuperscript{ITG} genomes where we found many sites to be called as heterozygotes within each line. Phasing the heterozygote calls using statistical methods was almost impossible as 98.9% of the total polymorphic sites among the wAna\textsuperscript{ITG} genomes were a PSHET. Thus, only sites...
that were called as homozygous were examined from each wAna$^{TG}$ genome, assuming that those were representative of the variants that existed in the wAna genome before integrating into the host genome as wAna$^{TG}$.

Qualitatively the mtDNA genome phylogeny (fig. 6A) was concordant with the mtDNA phylogeny from the study of Choi and Aquadro (2014), which was estimated using the same D. ananassae samples from this study but with Sanger sequences of only three mtDNA genes. No evidence of phylogenetic clustering was observed among the mtDNA haplotypes originating from D. ananassae strains with the wAna$^{INF}$ or wAna$^{TG}$ genomes.

As expected for maternally inherited organelles and endosymbionts, the branching patterns observed among the wAna$^{INF}$ genomes were congruent with the phylogenetic relationship estimated for the mtDNA (fig. 6A and B). Interestingly, the copy number analysis of specific wAna$^{INF}$ genomic regions (fig. 2), large structural variation analysis (fig. 5A), and the wAna$^{INF}$ nucleotide phylogeny (fig. 6B), which were inferred independently from one another, gave congruent phylogenetic results. For example, wAna$^{INF}$ genomes from strain TBU136 and VAV150 had a significant decrease in copy number in two regions of the genome compared to those expected from the distribution of the PCR-verified large structural variations (fig. 5A). SV1 and SV2 suggested that the wAna$^{INF}$ genomes were different from wAna$^{INF}$ of strains Cebu, HNL0501, and KMJ1; whereas all other large structural variants suggested that the wAna$^{INF}$ genomes were most related to wAna$^{INF}$ genomes from Cebu, HNL0501, and KMJ1.

Analysis of Sites Called as Heterozygotes for Each wAna$^{TG}$ Genome

Among the wAna$^{TG}$ genomes the majority of the polymorphic sites with a single nucleotide variant or small-sized INDEL were PSHET (table 2) and filtered out for most of our downstream analysis. To analyze these heterogeneous sites, for each wAna$^{TG}$ genome we attempted to phase the variation by sequencing a single F1 offspring of each wAna$^{TG}$ containing strain (while absent of wAna$^{INF}$) of D. ananassae that had been crossed to a strain lacking both integrated and infectious forms of wAna (see Materials and Methods for details). We expected these F1 progeny to have only a haploid complement of the wAna$^{TG}$ genome that would have allowed phasing of sites with a heterozygote base call. Each site that was called as a heterozygote in the diploid wAna$^{TG}$ genome was then examined in the haploid wAna$^{TG}$ genome. Surprisingly, the majority of the sites initially called as heterozygotes in the haploid wAna$^{TG}$ genome remained called as heterozygotes in the haploid wAna$^{TG}$ genome F1 offspring (table 4).

We next investigated whether for each wAna$^{TG}$ genome the increased number of sites called as heterozygotes could be explained by an increase in copy number of specific regions of the wAna$^{TG}$ genome (figs. 3 and 4). Duplication and triplication of regions would result in paralogs among which mutations could accumulate resulting in what would simply look like a heterozygote among Illumina reads. A sliding window analysis was conducted by dividing each wAna$^{TG}$ genome into nonoverlapping 50,000 bp windows, and for each window the average genome coverage and total number of single nucleotide variants were counted. Across all seven
wAnaITG genomes results showed a significant ($P < 0.0001$) positive correlation between the average genome coverage and the total number of single nucleotide variants in a given window (fig. 7A) as would be expected by varying numbers of paralogs of wAnaITG.

Next, each site called as a heterozygote was individually examined by counting the total read coverage and the total number of reads with the alternative allele supporting that heterozygote in strains of D. ananassae. These reads should remain 50:50 in proportion regardless of whether the site was called as a heterozygote from next-generation sequencing is expected by varying numbers of paralogs of wAnaITG.

Next, each site called as a heterozygote was individually examined by counting the total read coverage and the total number of reads with the alternative allele supporting that heterozygote in strains of D. ananassae. These reads should remain 50:50 in proportion regardless of whether the site was called as a heterozygote from next-generation sequencing. In contrast to this expectation, analysis of our wAnaITG genome results showed a significant ($P < 0.0001$) negative correlation between total read coverage and their proportion of reads originating from the alternative allele (fig. 7B). These results also support the hypothesis that variation in the number of paralogs of wAnaITG accounts for the large number of sites called as heterozygotes in strains of D. ananassae with the wAnaITG genome.

### Investigating the Integration Site of the wAnaITG Genome

In an effort to localize the integration site of wAnaITG into the D. ananassae nuclear genome, we used the program lumpy to detect chimeric raw paired-end reads where one end mapped to the wAnaITG genome whereas the other end mapped to the D. ananassae nuclear genomes. Reads from wAnaINF were also examined as a potential negative control for our in silico experiment, expecting no evidence of integration in the D. ananassae nuclear genome, as previous PCR screens have not detected any evidence of wAnaITG genes for our wAnaINF samples (Choi and Aquadro 2014).

Four of the seven wAnaITG samples had significant evidence of a breakpoint existing between the wAnaITG and D. ananassae nuclear genome (supplementary table S6, Supplementary Material online). There were multiple wAna breakpoints with overlapping genome coordinates among the wAnaITG samples, whereas only a single breakpoint was detected at a single location around 23,595,400–23,595,600 bp at scaffold_13340 from chromosome 2 L (Muller element E) of the D. ananassae nuclear genome (supplementary table S6, Supplementary Material online). Due to its unique localization it was designated as the candidate site for the W. pipientis–D. ananassae integration (DanaITG). PCR primers were designed to flank the computationally predicted putative DanaITG region and amplified in all 15 D. ananassae samples examined in this study plus the reference D. ananassae strain. Surprisingly, PCR results showed the same size bands in the wAnaINF and wAnaITG samples; however, the reference strain did not show any PCR bands (supplementary fig. S1A, Supplementary Material online). Sanger sequencing the putative DanaITG region from strains TB43 (D. ananassae line with wAnaITG) and KMJ1 (D. ananassae line with wAnaINF) indicated that the reference genome contained a unique DNA sequence at this location (supplementary fig. S1B, Supplementary Material online). A BLAST (Basic Local Alignment Search Tool) search of this sequence matched the long terminal repeats (LTRs) of a retrotransposon (supplementary fig. S1C, Supplementary Material online) and the breakpoints predicted by lumpy spanned this LTR region (supplementary fig. S1B, Supplementary Material online). In fact, the program RepeatMasker indicated that this region between 23,595,477 and 23,595,672 at scaffold_13340 corresponds to the Pao family of LTR retrotransposons (Xiong et al. 1993). Thus, multiple regions of the wAnaITG genome had breakpoints matching with the LTR of the host D. ananassae retrotransposon.

The Cebu strain was the only wAnaINF genome-carrying D. ananassae line with a lumpy-predicted DanaITG region that was different from the wAnaITG genomes. However, PCR amplification using primers designed to flank the Cebu wAnaINF–DanaITG region could not verify the computational predictions (results not shown).

Klasson et al. (2014) have recently shown evidence that wAnaITG sequence could be detected on chromosome 4 (Muller element F) of D. ananassae by fluorescent in situ hybridization (FISH) to mitotic chromosomes. We thus reanalyzed our wAnaITG genome sequence data with lumpy to see whether we could identify potential breakpoints between the wAnaITG genome and D. ananassae scaffolds identified as Muller F from a previous study (Bhutkar et al. 2008; see supplementary table S7, Supplementary Material online, for names of scaffold). These Muller F scaffolds were not part of the reference D. ananassae genome sequence we initially
used for our lumpy analysis. No significant breakpoints were detected in any of the eight wAna\textsuperscript{INF} genomes. However, all seven wAna\textsuperscript{TG} genomes had significant breakpoints (supplementary table S8, Supplementary Material online) with a total of 31 breakpoints found between the genomes of wAna\textsuperscript{TG} and D. ananassae Muller F scaffolds. Results from RepeatMasker indicated that for 28 of the 31 predicted breakpoints, the D. ananassae region of Muller F scaffolds corresponded to a TE or a retrotransposon (supplementary table S8, Supplementary Material online) and not a unique sequence.

Discussion

We have used next-generation sequencing to analyze the molecular evolution and phylogenomics of two different forms of W. pipientis associated with D. ananassae: 1) The bacterial originating and maternally transmitted wAna\textsuperscript{INF} genomes and 2) the host nuclear genome integrated wAna\textsuperscript{TG} genome. The wAna\textsuperscript{INF} and wAna\textsuperscript{TG} genomes showed distinct biological differences and characteristics.

Genomic Diversity within the wAna\textsuperscript{INF} Genomes

Our previous study of wAna\textsuperscript{INF} DNA sequence diversity based on Sanger sequencing failed to find any variability for 7 kb sampled from the wAna\textsuperscript{INF} genomes infecting eight geographically diverse strains of D. ananassae (Choi and Aquadro 2014). However, full genome sequencing reported here did find single nucleotide, INDEL, and copy number variation among these eight wAna\textsuperscript{INF} genomes, though for nucleotide variability at levels a 100-fold lower than those of the host mtDNA genomes. Congruent phylogenies for both the wAna\textsuperscript{INF} genomes and the maternally inherited host mtDNA suggest that the main mode of transmission for wAna\textsuperscript{INF} is through maternal transmission, an observation that has also been reported for W. pipientis in other species of Drosophila (e.g., Richardson et al. 2012; Early and Clark 2013).

Our analyses of copy number variation (fig. 2), large structural variation (fig. 5A), and nucleotide diversity (fig. 6) revealed several genomic strains of wAna\textsuperscript{INF} that could potentially differ functionally from each other. For example, wAna\textsuperscript{INF} from D. ananassae strain HNL0501 was the only strain to have a significant increase in copy number around a prophage region at genome coordinates 0.57–0.61 and 1.07–1.12 Mb (fig. 2). This prophage, named wRi-WOB (Klasson, Westberg, et al. 2009), exists as two identical duplicates in the wRi genome around genome coordinates 0.57–0.61 and 1.07–1.12 Mb, suggesting that the wAna\textsuperscript{INF} from D. ananassae strain HNL0501 has four copies of wRi-WOB. The functions of prophages in W. pipientis have not been fully described, but they can transpose and frequently horizontally transfer among divergent W. pipientis strains (Masui et al. 2000; Gavotte et al. 2007). Due to their lytic ability, the density of prophages has been suggested to control the strength of W. pipientis-mediated cytoplasmic incompatibility by decreasing the titer of W. pipientis infection (Bordenstein et al. 2006).
Interestingly, wAna\textsuperscript{INF} from \textit{D. ananassae} strain HNL0501 had the lowest wAna originating reads and normalized average read depth (\textit{table 1}). Assuming this low read depth reflects a low wAna\textsuperscript{INF} titer in \textit{D. ananassae} strain HNL0501, then the increased copy number of prophages may have caused the lowered bacterial load in this strain.

Another region with differences in copy number variation (0.61–0.63 and 1.12–1.14 Mb; \textit{fig. 2}) had only half of the typical copy number in wAna\textsuperscript{INF} of \textit{D. ananassae} strains TBU136 and VAV150 but double the typical copy number in \textit{D. ananassae} strain HNL0501. The two regions are mostly identical in sequence and contain coding sequences that are repetitive (Klasson, Westberg, \textit{et al.} 2009). The functional consequence of this variation in copy number is unknown; however, in wMel copy number variation of a 21-kb region has been suggested to cause pathogenicity in the virulent Popcorn strain wMelPop (Chrostek \textit{et al.} 2013; Chrostek and Teixeira 2015). Because copy number variation in the \textit{W. pipientis} coding region could lead to physiological responses in the host, future studies will be necessary to fully understand the potential host effects caused by the copy number variations observed across the wAna\textsuperscript{INF} genome.

As \textit{W. pipientis} genomes have an unusually high abundance of mobile genetic elements compared with other endosymbionts (Moran and Plague 2004), it is not surprising that all of the large structural variations we detected in the wAna\textsuperscript{INF} genome involved genes with transposase ability (\textit{fig. 5}; \textit{supplementary tables S2 and S3, Supplementary Material online}). This variation can be used for genotyping variability within wAna\textsuperscript{INF} as has been reported in studies of wMel, where the absence or presence of these mobile elements has been used as genotyping tools to differentiate within wMel strains (Riegler \textit{et al.} 2005; Woolfit \textit{et al.} 2013). Here, we have identified several TE absences or presences that were both polymorphic and fixed among the wAna\textsuperscript{INF} genomes. The presence/absence of polymorphic TE insertions was generally consistent with the nucleotide substitution-inferred phylogenetic tree, suggesting that these TE polymorphisms also are phylogenetically informative and can be a useful fast way to genotype different wAna\textsuperscript{INF} strains.

Further, these variants can be used to describe \textit{W. pipientis} strain variation in other hosts infected with strains highly similar to wAna. For example, wRi is not only very similar to wAna but also very similar to the \textit{W. pipientis} infecting \textit{Drosophila suzukii} (Siozios \textit{et al.} 2013), suggesting a very recent horizontal transfer of \textit{W. pipientis} among the three host species (\textit{D. ananassae}, \textit{D. simulans}, and \textit{D. suzukii}). The phylogenetically informative variants found in our study could be used to characterize the \textit{W. pipientis} diversity among these three host species, and potentially determine the coevolutionary history of the \textit{W. pipientis} infection among these three species.

Finally, variation in the proportion of raw reads originating from the wAna\textsuperscript{INF} genomes and average read depth of wAna\textsuperscript{INF} genomes (\textit{table 1}) suggest differences in bacterial titer among the \textit{D. ananassae} hosts. The biological significance of this variation in wAna\textsuperscript{INF} load is unknown but previous studies have shown various effects that are \textit{W. pipientis} density-dependent including level of cytoplasmic incompatibility (Clark and Karr 2002), male killing effect (Unckless \textit{et al.} 2009), and differences in antiviral protection for its host (Osborne \textit{et al.} 2012; Chrostek and Teixeira 2015). These phenotypes are likely to be under selection. Thus, the observed differences in bacterial titer among \textit{D. ananassae} strains could have larger biological and evolutionary consequences. In addition, as the host genotype can also lead to differences in \textit{W. pipientis} titer (Kondo \textit{et al.} 2005; Serbus \textit{et al.} 2011), it is important to understand the underlying coevolutionary history between \textit{W. pipientis} and its host to understand the biology of \textit{W. pipientis} (i.e., Chrostek \textit{et al.} 2013). The genomic resources and results from this study can help advance future studies of the genetic interaction between wAna\textsuperscript{INF} and its host \textit{D. ananassae}.

**Evolution of the wAna\textsuperscript{ITG} Genomes**

In all seven of our \textit{D. ananassae} samples with evidence of wAna\textsuperscript{ITG}, we found not only the whole wAna genome integrated into the host genome but also that specific regions were present in as many as seven copies (\textit{fig. 3}). During the preparation of this manuscript, Klasson \textit{et al.} (2014) independently reported evidence of extensive duplication in the wAna\textsuperscript{ITG} genome originating from their Hawaii and India strains of \textit{D. ananassae}. In our study, however, we have genome sequenced \textit{D. ananassae} samples from an even broader geographic range (Africa, southeast Asia, west Asia, and south Pacific islands) that encompasses both the ancestral and peripheral range of \textit{D. ananassae} (Das \textit{et al.} 2004; Schug \textit{et al.} 2007). Additional phylogenetic analyses together with genotyping using large structural variation allowed us to demonstrate that all wAna\textsuperscript{ITG} genomes were closely related to each other (\textit{figs. 5B and 6}). These results suggest that a single wAna genotype initially integrated into the host genome and subsequently dispersed throughout much of the worldwide range of \textit{D. ananassae}. Based on our phylogenetic and large structural variation results, the wAna\textsuperscript{ITG} genomes are most closely related to the wAna\textsuperscript{INF} infecting \textit{D. ananassae} strains Cebu, HNL0501, and KMJ1, suggesting that the common ancestor of the wAna\textsuperscript{INF} infecting these three strains was closely related to that which integrated into the host genome.

Interestingly the wAna\textsuperscript{ITG} genomes examined in this study had on average twice the \textit{D. ananassae} nuclear genome coverage (\textit{table 1}), whereas Klasson \textit{et al.} (2014) reported the wAna\textsuperscript{INF} samples from India and Indonesia to have far less coverage than the \textit{D. ananassae} nuclear genome. In addition, the wAna\textsuperscript{ITG} genome from Indonesia lacked the heterogeneous increased copy number, which was observed in all wAna\textsuperscript{ITG} genomes in this study (\textit{fig. 3}). This
underrepresentation of wAnaITG DNA in the India and Indonesia samples (Klasson et al. 2014) could be due to temporal differences in the integration of the wAna genome. Through an unknown mechanism the initial wAnaITG genome could have accumulated increased copy number and overrepresentation of itself over time. Then, the wAnaITG genomes examined in this study and the Hawaii strain from Klasson et al. (2014) could represent an ancient integrated wAnaITG genome. The increased number of mutations observed in the wAnaITG genome compared with the wAnaNE genome further suggests it to be an ancient integration (table 2). On the other hand, the wAnaITG genomes from India and Indonesia (Klasson et al. 2014) could be from a more recent integration event. This further suggests that the original wAna genome type integrated into the host genome is polymorphic, and potentially is an ongoing process in D. ananassae occurring multiple times. Here, the structural variations identified from this study could test whether multiple wAna genome types have integrated into the host nuclear genome.

The wAnaITG genomes have many characteristics of a pseudogene (Li et al. 1981; Miyata and Hayashida 1981) with an elevated proportion of nonsynonymous relative to synonymous (and intergenic) variants, and many segregating variants were predicted to have strongly deleterious effects if present in a functional wAna genome (table 3). These observations lead us to conclude that the wAnaITG genome is becoming (if not already has become) a large pseudogene after integrating into the host D. ananassae nuclear genome.

Regions with higher copy number in the wAnaITG genome also have more mutations (fig. 7A) due to the increased number of copies and sites accumulating mutations. In addition, our analysis of genome coverage (figs. 3 and 4) and unphased sites computationally called as heterozygotes in the wAnaITG genome (table 4), further supports the interpretation that the majority of the wAnaITG genome exists in at least two copies in the host genome (Klasson et al. 2014). Although copy number is variable across wAna genomic regions, the increased and decreased copy number regions were correlated across all seven wAnaITG genomes (fig. 3). This suggested that after the integration, specific regions of the wAnaITG first increased in copy number then subsequently dispersed worldwide. A recent study using FISH to mitotic chromosomes using the high copy region of the wAnaITG genome as a probe has found evidence of the integration only at a single location on the D. ananassae fourth chromosome (Klasson et al. 2014). The presence of only a single site of hybridization suggests that the increased copy number is likely to be due to increased tandem duplications of the wAnaITG regions.

wAnaITG regions with multiple copies could have a mutation occurring on one of the multiple paralogs, and this would be computationally called as a heterozygote as the multiple copies are collapsed into a single copy in the reference wRi genome. Compared with regions with low copy number, regions with higher copy number across the wAnaITG genome had more sites called as heterozygotes with lower proportions of reads from the alternative allele than from the reference allele (fig. 7B). This pattern would result if the rate of increase in copy number of specific wAnaITG regions is higher than the rate of nucleotide mutation.

The rapid increase in copy number also explains why our current full genome sequencing allowed us to detect numerous polymorphisms, whereas our previous analysis which had used Sanger sequencing of 7 kb from several wAnaITG gene regions found only three segregating sites (Choi and Aquadro 2014). The lower proportion of alternative to reference reads for many of the sites called as heterozygotes (fig. 7B) would be reflected in the Sanger sequencing chromatograms as different sized double peaks. We have reexamined the chromatograms of Choi and Aquadro (2014) and verified many PSHet identified in this study actually had double peaks in the chromatogram, albeit one peak was always much smaller than the other (results not shown), which normally would be discarded as noise. Thus, we believe that many of our computationally detected heterozygous base calls in the wAnaITG genome are not due to sequencing errors.

Using in silico analysis coupled with PCR verification, we found no convincing evidence of breakpoints between the wAnaITG and D. ananassae nuclear unique sequence from the euchromatic chromosomes X, 2, and 3. The only breakpoints discovered were between multiple wAnaITG regions and an LTR of a D. ananassae retrotransposon, suggesting that the host retrotransposons were actively integrating into various regions of the integrated wAnaITG genome. This has been noted previously (Dunning Hotopp et al. 2007) and our further analysis with the D. ananassae fourth chromosome scaffolds also corroborated this result. The fourth chromosomal scaffolds have a higher frequency of retrotransposons and TEs than D. ananassae scaffolds from chromosome X, 2, and 3 (Leung et al. 2015). Thus, many of the putative-predicted breakpoints between the D. ananassae fourth chromosome and wAnaITG genome (supplementary table S8, Supplementary Material online) are likely artifacts caused by D. ananassae TEs that have inserted into the wAnaITG genome. Consistent with the results previously reported by Dunning Hotopp et al. (2007), evidence of frequent integrations of D. ananassae TEs into the wAnaITG genome complicates the placement of wAnaITG genomic fragments in the D. ananassae genome. Long-read sequencing technology (e.g., Pacific Biosciences) and additional in situ hybridization experiments may in the future provide insight into the precise integration site for the wAnaITG genome.

Our analyses of the wAnaITG genomes thus lead to several intriguing conclusions: 1) Although pseudogenes are rare across Drosophila species (Clark et al. 2007), the wAnaITG of D. ananassae seems to have become a large pseudogene after the integration; 2) Drosophila shows a strong bias toward deletion of nonfunctional genes (Petrov et al. 1996) but
the wAnaITG genome shows evidence of extensive duplications despite its likely lack of functionality; and 3) more than 2% of the genomic reads from D. ananassae originated from the pseudogene-like wAnaITG genome (table 1) which we infer to have had minimal negative fitness consequences as evidenced by its wide geographic distribution (Choi and Aquadro 2014). Thus despite the apparent loss of functionality across the wAnaITG genome, some paralogs among the multiple wAnaITG copies could be under positive selection leading to the observed pattern of spreading across multiple D. ananassae populations.

Conclusion

The wAnaINF and wAnaITG genomes show drastically different evolutionary and population genomics. We discovered diverse nucleotide and structural variants among wAnaINF genomes that showed phylogenetic relationships consistent with a strict maternal inheritance after an initial single interspecific infection within D. ananassae. Similarly, the wAnaITG genome appears to have arisen from a single integration of one wAnaINF variant not long after the initial infection of the species. Subsequent to the initial integration, the majority of the wAnaITG genome had at least doubled its copy number within the D. ananassae host genome. Additionally after the integration of the wAnaITG genome, it appears to have become a large pseudogene accumulating substantially more mutations than the cytoplasmic wAnaINF genome, including many of which would be predicted to be strongly deleterious in a functioning wAna genome. Although it is possible that the wAnaITG has spread across a wide geographic distribution through genetic drift and/or migration alone, further study of the wAnaITG genomes from additional geographically diverse strains of D. ananassae is needed to distinguish hypotheses as to what has apparently caused the geographically wide-spread distribution of wAnaITG despite its apparent loss of functionality.

Supplementary Material

Supplementary data, figures S1, and tables S1–S8 are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org/).

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