

Into the Wild: Parallel Transcriptomics of the Tsetse-*Wigglesworthia* Mutualism within Kenyan Populations

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Abstract

Tsetse flies (Diptera: Glossinidae) have medical significance as the obligate vectors of African trypanosomes. In addition, tsetse harbor a simple gut microbiota. A predominant gut microbiota member, the Gammaproteobacterium *Wigglesworthia* spp., has coevolved with tsetse for a significant portion of *Glossina* radiation proving critical to tsetse fitness. Although multiple roles have been described for *Wigglesworthia* within colony flies, little research has been dedicated towards functional characterization within wild tsetse. Here, dual RNA-Seq was performed to characterize the tsetse-*Wigglesworthia* symbiosis within flies captured in Nguruman, Kenya. A significant correlation in Gene Ontology (GO) distribution between tsetse and *Wigglesworthia* was observed, with homogeneous enrichment in metabolic and transport categories, likely supporting a hallmark of the symbiosis-bidirectional metabolic exchange. Within field flies, highly transcribed *Wigglesworthia* loci included those involved in B vitamin synthesis and in substrate translocation, including amino acid transporters and multidrug efflux pumps, providing a molecular means for interaction. The universal expression of several *Wigglesworthia* and *G. pallidipes* orthologs, putatively involved in nutrient provisioning and resource allocation, was confirmed in sister tsetse species. These transcriptional profiles varied through host age and mating status likely addressing varying symbiont demands and also confirming their global importance within *Glossina*. This study, not only supports symbiont nutrient provisioning roles, but also serves as a foundation for insight into novel roles and molecular mechanisms associated with vector-microbiota interactions. The role of symbiont B vitamin provisioning towards impacting host epigenetics is discussed. Knowledge of vector-microbiota interactions may lead to the discovery of novel targets in pest control.

Key words: tsetse, *Wigglesworthia*, symbiosis, microbiota.

Introduction

Bacteria are the most prevalent animal symbionts (Moya et al. 2008). Among the best examples of reciprocal selection occurring between animals and bacteria are the intimate alliances of insects and their ancient obligate endosymbionts (Douglas 1998; Moran and Sloan 2015). These associations drive evolutionary modifications between partners that serve to enhance the functional roles that form the pinnacle of the species-species integration. Only a small fraction of insects harbor these long-established and essential associations (Douglas 1998; Moran et al. 2008), primarily those that require nutritional supplementation due to limited and

unbalanced diets deficient in vitamins or amino acids, such as exclusive feeding on blood, phloem, or xylem. These symbionts are often localized to specific host tissues or organs (often called the bacteriome) that maximize functional efficiency by centralizing activity. Despite the functional characterization of the critical roles provided by these symbioses, little is known concerning the molecular dialog between host and symbiont that may facilitate integration in the wild.

Tsetse flies (Diptera: Glossinidae) are medically significant as the obligate vector of African trypanosomes (*Trypanosoma* spp.), the etiological agents of Human African trypanosomiasis (HAT) and Animal African trypanosomiasis (AAT).

In addition to potential infections with trypanosomes, tsetse harbor a naturally simple digestive tract microbiota consisting of only a few bacterial species (Weiss et al. 2011; Aksoy et al. 2014). These bacteria have distinct coevolutionary histories with tsetse and, likewise, contributory roles towards their insect host biology (reviewed in Rio et al. [2016]). Two Gammaproteobacteria are associated with the digestive track of the tsetse fly; *Wigglesworthia* spp. and *Sodalis glossinidius* (Aksoy 1995a; Cheng and Aksoy 1999). Infections with *Wolbachia* are also common in tsetse reproductive tissue (Cheng et al. 2000). These parasitic *Wolbachia* infections may cause the arrest of embryonic development following incongruous mating in a condition known as cytoplasmic incompatibility (Alam et al. 2011). The simplicity in the microbiota composition is likely generated through two unique facets of tsetse fly biology; the obligate vertebrate blood feeding lifestyle of adults and the live birth of progeny following intrauterine larval development (known as adenotrophic viviparity [Benoit et al. 2015]), both of which severely curtail environmental microbial exposure.

The most predominant member of the tsetse alimentary tract microbiota (Rio et al. 2006; Aksoy et al. 2014) is the obligate mutualist *Wigglesworthia* spp. that has codiversified with tsetse for a significant portion of *Glossina* radiation, dating back 50–80 million years (Chen et al. 1999). The small *Wigglesworthia* genome (~0.7 Mb) retains only those genes necessary for habitation within the tsetse (Akman et al. 2002; Rio et al. 2012). Both the tsetse and *Wigglesworthia* partners rely on each other for evolutionary persistence, exemplified by their interwoven reproductive fitness. The *Wigglesworthia* symbiont is tightly harnessed to the tsetse host due to extreme reductive genome evolution (Akman et al. 2002; Rio et al. 2012) that makes a free-living lifestyle highly unlikely. Likewise, removal of the *Wigglesworthia* symbionts from the tsetse fly through selective antibiotic cocktails severely impedes fly reproduction (Hill et al. 1973; Nogge 1978; Nogge 1981) that can be partially restored through B vitamin cocktails or yeast supplementation (Nogge 1981; Pais et al. 2008) indicating essential metabolic interdependency between these partners.

Within tsetse, intracellular *Wigglesworthia* are localized to a specialized organ (bacteriome) at the anterior gut (Aksoy 1995a). A second *Wigglesworthia* population is harbored exclusively within milk glands that empty into the uterus (Attardo et al. 2008). These modified accessory glands found only within females, provide nourishment to the larva while also serving as a vessel for vertically transmitting gut microbiota (Ma and Denlinger 1974; Attardo et al. 2008; Balmand et al. 2013). To date, most of our understanding of the tsetse-*Wigglesworthia* association has been generated primarily from experimental assays conducted with colony flies reared under controlled rearing conditions. In this setting, key *Wigglesworthia* roles towards tsetse reproduction, digestion, development and immunological priming have been identified (reviewed in Rio et al. [2016]).

Studies that investigate the relationship between animals and their microbiota are largely performed using model systems that have been maintained in laboratory settings for extensive periods. Little research has been dedicated towards validating whether these model systems accurately reflect wild counterparts within their native habitats. Field-based studies may be used to validate or disprove conclusions obtained through the microbiota studies of captive or artificially bred animals. Equally as important, these field-based studies can also identify novel functional significance of the microbiota towards host biology that may be masked, or simply unattainable, whereas rearing in captivity.

In the efforts to deepen our understanding of wild animals and their microbiota interactions within their natural ecology, a parallel Illumina-based RNA-Seq approach was used to examine the molecular integration of the tsetse fly-*Wigglesworthia* mutualism in field collected flies. Here, we characterize the genome activity of the tsetse-*Wigglesworthia* association within the bacteriomes of wild caught adult tsetse (*Glossina pallidipes*) captured in Nguruman, Kenya. To further assess the universality of RNA-Seq results across tsetse species, we characterize the expression profiles of genes associated with novel putative functional roles for *Wigglesworthia* and *Glossina* in sister tsetse species. These high-throughput transcriptome analyses provide a global picture of host and symbiont interconnectedness by providing insights into important functional roles including concerted transcriptional profiles involved in nutrient provisioning and resource allocation within their natural habitat. These studies aim to identify molecular candidate targets that disrupt the crucial symbiosis within the medically important tsetse fly providing novel vector control strategies.

Materials and Methods

Insect Collection

Adult tsetse flies, *Glossina pallidipes* (Diptera: Glossinidae) were isolated from Nguruman escarpment, Kenya in March 2015 using Ngu traps. Bacteriomes were microscopically removed and placed in RNA_{later} (Invitrogen, CA). Flies included in this study were all trypanosome free as verified through microscopic analyses of their midguts.

Molecular Phylogenetic Analyses

The bacteriomes of *G. pallidipes* field collected flies were dissected and DNA was extracted following the Holmes–Bonner protocol (Holmes and Bonner 1973). DNA samples were subject to PCR amplification with general eubacterial 16S rRNA primers, 27F' and 1492R' (Lane 1990; Weisburg et al. 1991) ($T_a=50^\circ\text{C}$; 28 cycles). Amplicons were ligated into pGEM-T vector (Promega, WI) and *Escherichia coli* JM109 cells (Promega, WI) were transformed. Colonies were verified for a 16S rRNA insertion and sequenced at the West Virginia

University's Department of Biology Genomics Center on an ABI 3130xl analyzer Applied BiosystemsCA) using a 3.1 BigDye protocol (Applied Biosystems). All sequences were quality trimmed and assembled into contigs using CLUSTALW (available at www.ebi.ac.uk/Tools/msa/muscle/; last accessed January 3, 2017). A consensus 16S rRNA nucleotide sequence for the *G. pallidipes Wigglesworthia* isolate was deposited in the NCBI Genbank database under accession number MF148851.

The evolutionary model used for Bayesian analyses (General time-reversible plus invariant sites plus gamma; GTR + I + G) of 16S rRNA sequences was determined using the Akaike Information Criterion in MRMODELTEST 2.3 (Nylander 2004). Bayesian analyses were performed in MRBAYES 3.2.6 (Ronquist and Huelsenbeck 2003) with the number of categories used to approximate the gamma distribution set at four. Additionally, six Markov chains (Larget and Simon 1999) were run for 3,000,000 generations. Posterior probability (PP) values were calculated, with stabilization of model parameters (i.e., burn-in) occurring at 2,800,000 generations (standard deviation of split frequencies < 0.01). Every 100th tree following stabilization (burn-in) was sampled to calculate a 50% majority-rule consensus tree. All trees were constructed using the program FIGTREE v1.4.3 (<http://tree.bio.ed.ac.uk/software/figtree/>; last accessed January 3, 2017).

RNA Extraction, Illumina Library Preparation, Sequencing, and Genome Aligning

The bacteriomes of 12 mature tsetse flies were pooled for one biological sample, resulting in the construction of six biological samples analyzed from male and female adult samples. Bacteriome samples were homogenized and total RNA was extracted using a MasterPure RNA purification kit (Epicentre, Madison, WI) according to the manufacturer's protocol for tissue samples. DNA was removed from the RNA samples using a Turbo DNA-free kit (Ambion, Austin, TX) following the rigorous DNase treatment option. RNA of sufficient quality for cDNA synthesis was confirmed using the Agilent 2000 Bioanalyzer RNA Nano chip. RNA samples were subsequently processed with a Ribo-Zero magnetic kit for Gram-negative bacteria (Epicentre, Madison, WI) according to the manufacturer's protocol. The resulting mRNA-enriched RNA was then purified using an RNeasy MinEluteCleanup kit (Qiagen, Valencia, CA) and eluted in RNase-free H₂O. The eluted RNA (~1 µg) was then processed using a Kapa stranded mRNA-seq kit (Kapa Biosystems, Wilmington, MA), with the omission of the poly(A) pulldown, by the WVU Genomics Core Facility. The resulting cDNA libraries were sequenced using the Illumina HiSeq 1000 platform (2 by 51 bp) at Marshall University. Following sequencing, raw reads were postprocessed in order to remove Illumina adapters/primer sequences.

FASTQC analysis was performed on the RNA-Seq data sets to validate read quality. In order to capture both

Wigglesworthia and tsetse fly reads, the Kallisto–Sleuth pipeline (Bray et al. 2016) was first used for the identification and quantification of *G. pallidipes*-specific read counts based on the genome available at Vectorbase (<https://www.vectorbase.org/organisms/glossina-pallidipes>, last accessed February 16, 2015, Gpal1.2 version). Following the parsing out of tsetse-specific reads from the total pool of bacteriome reads, the remaining sequences were mapped to the *Wigglesworthia morsitans* genome (NC_016893.1; Rio et al. 2012) using the STAR alignment tool (Dobin and Gingeras 2015). TPM (Transcripts per Million) was used as a measure of gene expression (Li et al. 2010). Relative fold differences in gene expression between samples were determined as a ratio of each TPM.

Differential Expression and Gene Ontology Analyses

Differential expression profiles of specific loci between male and female bacteriomes were identified using DESeq (Anders and Huber 2010), Kallisto–Sleuth, and ANOVA with an internal multiple tests correction in R using custom scripts (only performed for the *Wigglesworthia* data set with scripts available upon request). Transcripts were considered differentially expressed if showing an *adjusted P value* ≤ 0.05. A web-scraping Python script (available at https://github.com/nathantspencer/auto_vectorbase; last accessed October 26, 2016.) merged biologically relevant information obtained from VectorBase to all tsetse genes with significant differences in expression levels between the bacteriomes of different sexes. For those loci that lacked any associated VectorBase annotation, NCBI blastx analyses to the nonredundant protein sequences (nr) database were performed with results filtered to retain only hits with an *E-value* of < 1e⁻¹⁰ and a BitScore of >50. TRANSPORTDB 2.0 (Elbourne et al. 2017) (<http://www.membranetransport.org/transportDB2/index.html>; last accessed June 15 2017) was used to describe the predicted cytoplasmic transport protein complement within the *Wigglesworthia* (WGM) genome with those identified then used to assess transporter expression within field flies. Further, sequences were assigned to Gene Ontology (GO) terms falling within biological process, molecular function and cellular component according to GO hierarchy using BLAST2GO (Conesa et al. 2005; <https://www.blast2go.com/home>; last accessed March 14 2017). Differentially expressed GO terms were identified using Fisher's exact test followed by a False Discovery Rate (FDR) corrections. Raw RNA-Seq data were uploaded to the National Center for Biotechnology Information (NCBI) Sequence Read Archive (SRA) database (Bio-project PRJNA387614).

Real-Time qRT-PCR Gene Expression Analyses of Select *Wigglesworthia* Genes

Tsetse flies (*G. morsitans*) were reared in standard conditions and mated as described previously (Snyder et al. 2015). Tsetse flies, of known ages and female fecundity status, were sacrificed and bacteriomes dissected. Total RNA was isolated

following the TRIzol protocol (Invitrogen, CA) and treated with DNaseI (Ambion, TX). First strand cDNA synthesis was performed with 140 ng total RNA, a 2 μ M primer cocktail of gene specific 3' end primers (supplementary table S6, Supplementary Material online) and Superscript Reverse Transcriptase II (Invitrogen) following manufacturer instructions. qRT-PCR was performed with SsoFast EvaGreen supermix (Bio-Rad), 0.4 mM gene-specific primers designed by the Primer3 server (supplementary table S6, Supplementary Material online), and 2 μ l of cDNA template in a Bio-Rad CFX96 real-time PCR detection system (Bio-Rad, CA) with 35 amplification cycles. Primers were checked with the BLAST tool at NCBI to exclude potential unspecific amplification. Additionally, primer specificity was confirmed by a melting curve analysis where the dwell temperatures increased from 65 to 95 $^{\circ}$ C in 0.5 $^{\circ}$ C increments every 5 s. Primer efficiency was evaluated using the standard curve method and ranged from 90 to 110%. The threshold cycle ($2^{-\Delta\Delta CT}$) method was used to calculate relative expression. The *Wigglesworthia rpsC* (30S ribosomal subunit) was used as the reference gene. At least five individual bacteriomes were processed per group, with each sample being analyzed in triplicate, and the average quantification cycle (CT) obtained. The fold change in gene expression, as compared with the same sex teneral stage (i.e., newly emerged, nonfed adult), was determined for each sample. Negative controls were included in all amplification reactions. Values are represented as the mean \pm the standard error of the mean (SEM) with statistical significance determined with ANOVA followed by Tukey's multiple comparisons post hoc analyses.

Investigation of Tsetse Transporter Expression within Tsetse Tissues

To examine the expression of the *Glossina* species orthologs of reduced folate carrier and thiamine transporter (i.e., GPAI043750 and GPAI022398, respectively), semiquantitative reverse transcription PCR (RT-PCR) analyses were performed for GMOY005445, GMOY009200, GFUI022020, and GFUI042523. Tsetse flies were sacrificed and tissues (bacteriomes and heads) dissected. Total RNA was isolated following the TRIzol protocol (Invitrogen, Carlsbad, CA), treated with DNase I (Ambion), and verified to be free of DNA contamination through PCR using RNA only as the template. First-strand cDNA synthesis was performed with 140 ng RNA, a 2 μ M primer cocktail of gene-specific 3' end primers and SuperScript II reverse transcriptase (Invitrogen). Second-strand synthesis was performed with primers specific for the complementary 5' end of the gene and 2 μ l of cDNA template for 35 amplification cycles. Primers for GMOY005445 included: forward: 5'-CTCAAAGCCACCACCTTGTT-3', reverse: 5'-CAACGATGACAAGACGGCTA-3'; for GMOY009200: forward: 5'-GAAGGTGATTGGTGGACTGG-3', reverse: 5'-TGAGGATACAGAGGCAGGAA-3'; for GFUI022020: forward:

5'-TAGCGGCTACACAAATGCCA-3', reverse: 5'-AGCCACA CGTCTTCAACGAT-3'; and for GFUI042523: forward: 5'-TCGGGTAACATCGCACACAA-3', reverse: 5'-CTTCCT CCAAATGACGGGCT-3'. The constitutively expressed β -Tubulin gene (forward: 5'-GATAACGAGGCCCTGTACGA-3', reverse: 5'-GATAACGAGGCCCTGTACGA-3') was used as a loading control and to verify RNA integrity. At least six bacteriomes were processed per group. Negative controls were included in all amplification reactions.

Statistical Analyses

All statistical analyses were performed with GraphPad Prism software (version 6.0) with the identification of specific analyses performed indicated throughout text.

Results

General Transcriptome Features

Six cDNA libraries were generated from the total homogenates of sex-specific bacteriomes isolated from Nguruman, Kenya field-captured adult tsetse flies. These libraries consisted of the homogenized bacteriomes of same sex adults of different ages, mating, and feeding states all obtained within a 24 h sampling period in March 2015. The libraries were sequenced using Illumina HiSeq technology, resulting in \sim 23 million high-quality paired-end reads (2 x 51 bp in length) for each biological sample, with a combined total for all libraries of 140,008,786 cleaned reads. Barcodes and adapters were trimmed during the demultiplexing process. A total of 139,635,056 reads (\sim 99.7% of cleaned reads) were identified as having tsetse origin whereas 373,730 reads (\sim 0.3% of cleaned reads) mapped to *Wigglesworthia* genes (fig. 1A). There were no significant differences in the mean abundance of either *Wigglesworthia* or tsetse reads between male and female specific libraries (Student's *t*-test; $P = 0.7952$ and $P = 0.9302$, respectively).

Wigglesworthia-Based Analyses

To investigate the phylogenetic relationship of the *G. pallidipes* *Wigglesworthia* isolate to other Gammaproteobacteria, we first performed Bayesian analyses based on 16S rRNA sequences generated through this study and additional sequences obtained from the GenBank database. Tsetse bacteriomes, obtained from *G. pallidipes* collected from Nguruman, Kenya, were dissected and DNA extracted for the PCR amplification and sequencing of eubacteria 16S rRNA. Single amplicons with highest nucleotide identities to the *Wigglesworthia* 16S rRNA isolated from other tsetse species were consistently obtained when amplifying from the bacteriome-generated DNA. Phylogenetic analyses of 16S rRNA sequences, generated through Bayesian analyses, indicates a close relation with a high posterior probability value (0.9483) between the

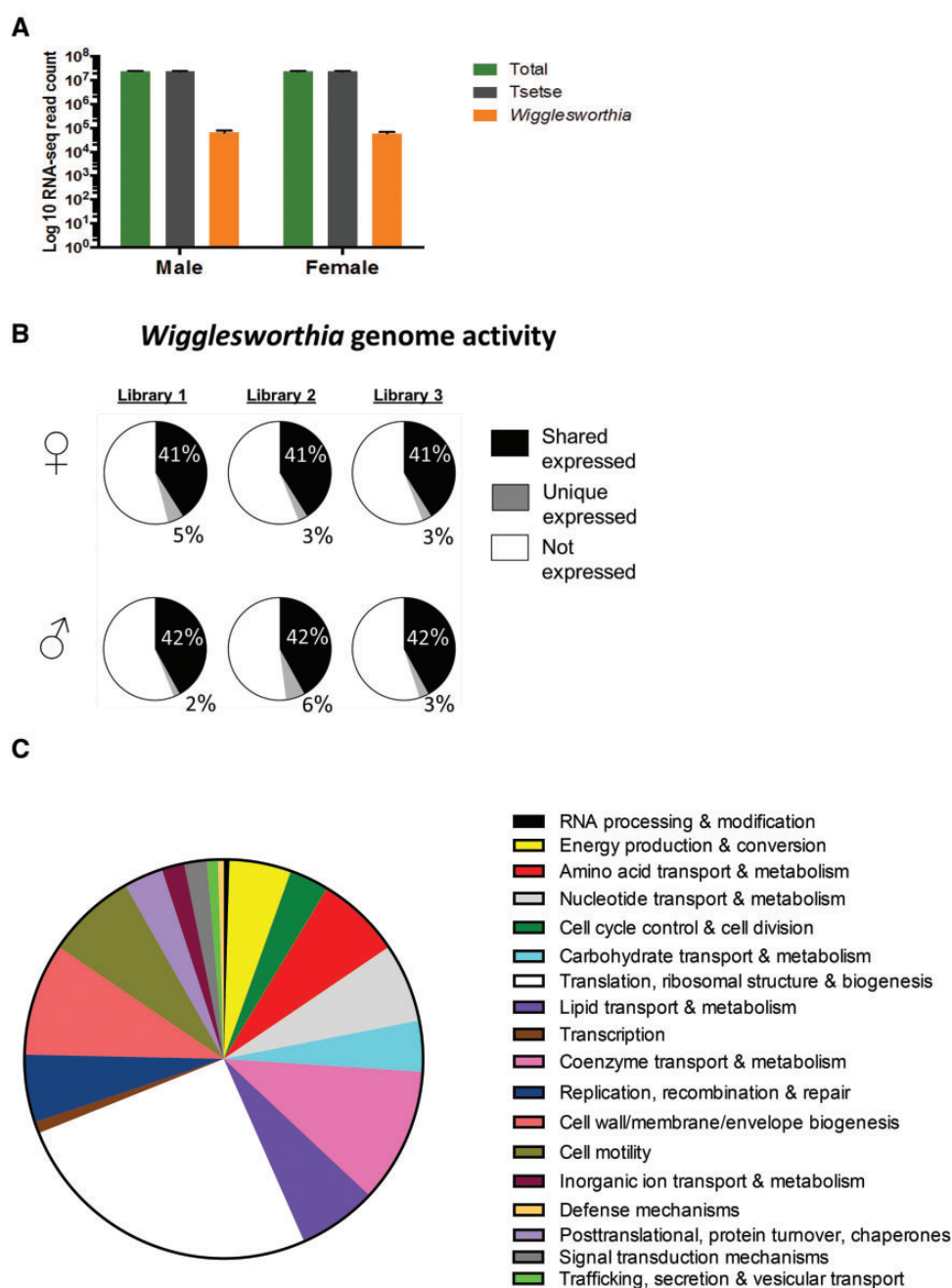


FIG. 1.—A summary of RNA-Seq libraries. (A) Number (Log10 base) of total RNA-Seq reads mapping to either tsetse (*G. pallidipes*) or *Wigglesworthia* reference genomes. 1 Standard Error of the Mean (S.E.M.) bars are depicted. (B) Percentage of the *Wigglesworthia* genome found to be expressed within each of the bacteriome libraries, shared and uniquely (relative to other same sex libraries) expressed proportions are indicated. (C) A total of 273 genes lacked reads in any library, the distribution of their Clusters of Orthologous Genes (COGs) designations are conveyed in the pie chart.

Wigglesworthia G. pallidipes and *G. morsitans* (WGM) isolates (supplementary fig. S1, Supplementary Material online). Depicting a substantially more distant relation to the *Wigglesworthia* isolate from *G. pallidipes* is the *G. brevipalpis* *Wigglesworthia* isolate (WGB, the other available reference genome). WGB is located within an adjacent sister clade reflective of the ancestral state of *G. brevipalpis* within the tsetse lineage

(Petersen et al. 2007; Dyer et al. 2008) and codiversification of this obligate symbiont through tsetse radiation (Chen et al. 1999). The higher sequence identity between the *Wigglesworthia G. pallidipes* and WGM 16S rRNA sequences (98% sequence identity vs. 95% sequence identity between *G. pallidipes* and WGB) validated the use of the WGM isolate as a reference genome for the mapping of Illumina reads.

All libraries had equivalent levels (44–48%) of the *Wigglesworthia* genome expressed (fig. 1B) with no significant differences between male and female bacteriomes in the percentage of the genome being expressed (Chi-square; $P = 0.6390$). Small portions of each library (2–6%) were exclusive in expression relative to other libraries within the same sex (fig. 1B). A total of 418 genes were expressed within at least 1 bacteriome library, corresponding to ~60% of the *Wigglesworthia* genome being transcribed during tsetse adulthood. Moreover, a total of 273 *Wigglesworthia* loci, distributed among 18 Cluster of Orthologous Genes (COGs), lacked Illumina reads within all the bacteriome libraries (fig. 1C and supplementary table S1, Supplementary Material online). The most widely represented COG lacking expression was translation, ribosomal structure and biogenesis, representing ~26% of COGs.

Gene Ontology (GO) Annotation

BLAST2GO (Conesa et al. 2005) was used to assign 370 protein-coding *Wigglesworthia* genes to associated Gene Ontology (GO) classifications for cellular components, molecular functions and biological processes. The resulting 1,106 GO terms ranged from levels 2 to 14 (supplementary fig. S2, Supplementary Material online), including parent and child terms, with 403 accessions within biological process, 538 in molecular function, and 165 in cellular components. A total of 23 GO terms were obtained when examining root level 4, with 3 to cellular components, 14 to biological processes and 6 to molecular functions (fig. 2A). Among cellular components, cytoplasm was the most enriched category (~58% of all genes within cellular components). Within biological processes, cellular nitrogen compound metabolism (~15% of all genes within biological processes) was the top process. Lastly within molecular functions, nucleotide binding and anion binding (~21% of all genes within molecular functions for both categories) was the top process identified. An enrichment analysis using Fisher's exact test with a false-discovery rate (FDR) correction found no significant differences between male and female bacteriomes (Fisher's exact test, $P > 0.05$) in the number of *Wigglesworthia* genes expressed and their respective GO distribution. Within all adult bacteriomes, there was a universal enrichment of expressed *Wigglesworthia* genes with GO terms associated with metabolism and transport (i.e., 57% of total GO categories at root level 4). Cellular nitrogen compound metabolism was the most highly represented GO category (~10% of all genes with GO terms at root level 4), a group that includes loci involved in the synthesis of amino acids, vitamins and cofactors.

To examine whether GO classifications differed in their mean expression levels, categories were averaged in their respective TPM values and comparatively analyzed (fig. 2B). Individual Student's *t*-test analyses found no significant differences ($P > 0.05$) in the mean TPM values of individual GO

categories between male and female bacteriomes (fig. 2B). Additionally, no significant statistical differences in the mean expression between GO categories within a sex (for males; $P = 0.9968$, for females; $P = 0.9908$) were observed.

Tsetse Sex-Specific Differences in *Wigglesworthia* Expression

Principal Component Analysis (PCA) revealed that only 24% (PC2) of the *Wigglesworthia* count variation between male and female bacteriome libraries (fig. 3A) could be accounted for by sex. The variation in symbiont expressed genes between tsetse male and female bacteriomes is likely driven by genes that either lacked expression or were overexpressed in the bacteriome of one sex relative to the other. A total of nine genes was expressed by *Wigglesworthia* exclusively within female or male bacteriomes (table 1), albeit with relatively low TPM values (i.e., 8.1–34.6, whereas the average TPM of libraries was 2,392). Within only female bacteriomes, *Wigglesworthia* reads pertaining to; *lolD*, an ATP binding component/lipoprotein transporter, *pyrG*, a CTP synthetase and WIGMOR_RS03465, a putative membrane protein transporter, were identified. Within only male bacteriomes, reads associated with; *trkA*, a potassium uptake protein, *pyrI*, a component of pyrimidine biosynthesis, *dnaB*, a DNA helicase, *rseP*, a zinc metalloprotease, *ahpC*, an alkyl hydroperoxide reductase, and *mrdb*, involved in cell elongation/peptidoglycan synthesis were found. Additionally, a small number of *Wigglesworthia* genes significantly differed in expression between female and male bacteriomes (fig. 3B and supplementary table S2, Supplementary Material online). Upon comparison of the results generated by the different analyses (DESeq, Kallisto-Sleuth and R analyses), a consensus of four *Wigglesworthia* loci were found to differ in expression between tsetse sexes; *flhD*, flagellar transcriptional activator, *motA*, the proton conductor of flagella motor, WIGMOR_RS02980, a cysteine ABC transporter binding protein and *infB*, a translation initiation factor IF2, which were all more highly expressed within female bacteriomes.

Due to the little variation in expression between tsetse sexes, we decided to then pool all the libraries to characterize global *Wigglesworthia* expression patterns of adult bacteriomes within field tsetse. The genes, *groEL* and *cspE*, had the highest abundance of mapped reads with TPM values of 70- and 75-fold higher than the average TPM, respectively. The *groEL* gene encodes a member of the chaperonin-60 family and has previously been associated with constitutive overproduction by *Wigglesworthia* throughout tsetse development (Aksoy 1995b) and also by other insect-associated obligate endosymbionts (Ishikawa and Yamaji 1985; Charles et al. 1997). Chaperonins assist in the correct assembly of proteins, including the refolding of denatured polypeptides, and their translocation (Goloubinoff et al. 1989; Goloubinoff et al. 1989; Nakamoto et al. 2014), which may be of particular

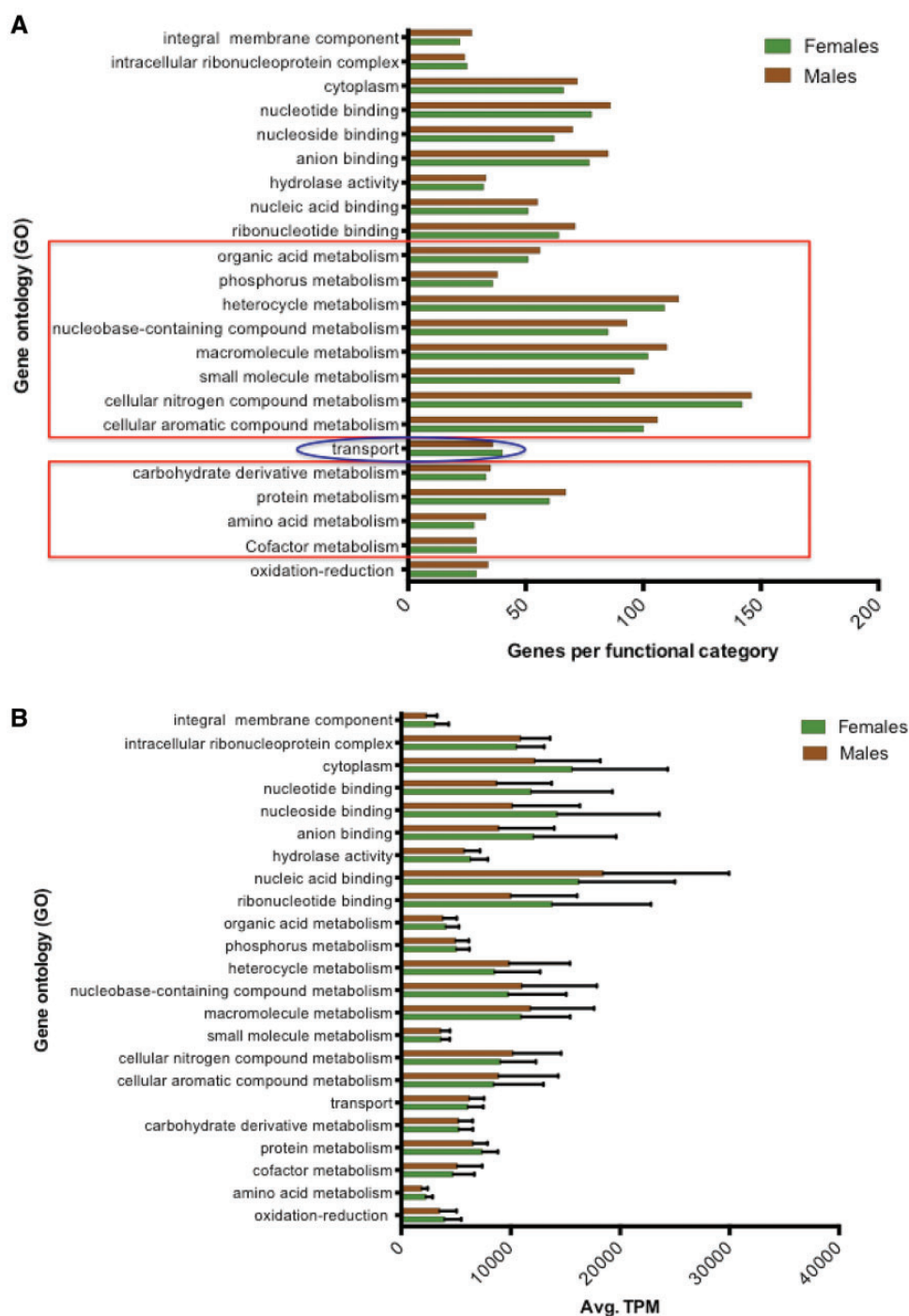


Fig. 2.—Gene ontology (GO) distribution of *Wigglesworthia* transcriptome. (A) GO classifications at root level 4 for each *Wigglesworthia* protein-coding gene ($n = 370$) expressed. GO terms were determined using BLAST2GO followed by enrichment analyses using Fisher's exact test with a false rate discovery correction. Red boxes indicate GO classifications associated with metabolism and the blue circle highlights transport. (B) GO classifications normalized to average TPM values. There were no statistically significant differences in mean TPM values either within a given category between male and female bacteriomes or between different categories within the same sex.

importance given the high mutational biases within the genomes of many of these ancient obligate symbionts (Fares et al. 2002, 2005; Tokuriki and Tawfik 2009; McCutcheon and Moran 2011). Previous reports of *groEL* as

one of the more highly transcribed *Wigglesworthia* genes serve to validate our RNA-Seq results. Interestingly, *cspE*, annotated as a cold shock protein and DNA-binding transcriptional repressor, is the gene under the highest purifying

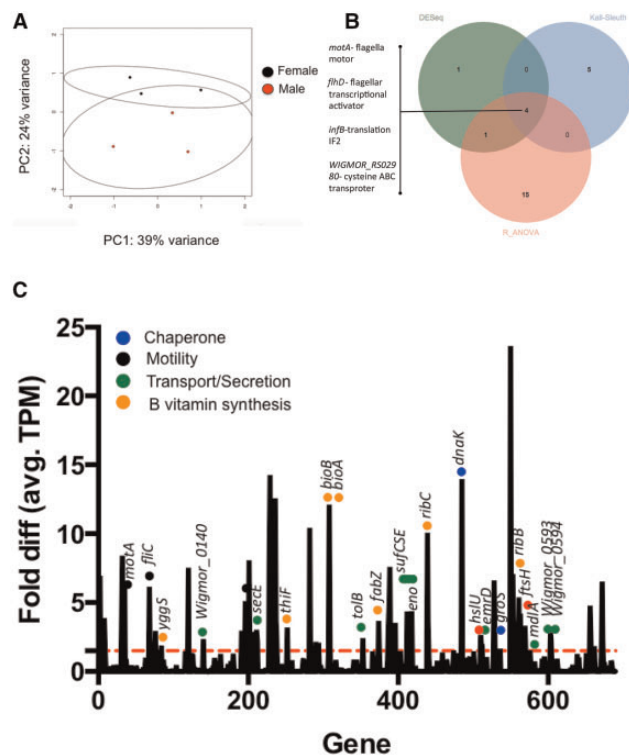


FIG. 3.—Tsetse sex-specific differences in *Wigglesworthia* expression. (A) Principal components analysis (PCA) of *Wigglesworthia* expression within male and female bacteriomes. Each point represents the *Wigglesworthia* transcriptomic signature of one RNA-Seq library. The directionality of separation was observed in the Principal Component (PC) 2, with 24% variance. 95% confidence circles are indicated. (B) A small number of differentially expressed *Wigglesworthia* genes were found between female and male bacteriomes. A consensus of 4 *Wigglesworthia* loci was found to be significantly higher expressed in female bacteriome libraries utilizing different analytical methods; DESeq (Differential expression of RNA-Seq data), Kall-Sleuth (Kallisto-Sleuth), and R-ANOVA (R statistical package, Analysis of variance). (C) Histogram showing the distribution of fold diff (avg. TPM) across the *Wigglesworthia* genome for compiled RNA-Seq libraries. The *Wigglesworthia* genome is depicted linearized, colored circles are associated with specific functional groups, the red dashed line indicates 1.5 fold diff. (avg. TPM).

selection ($dN/dS = 0$) within the *Wigglesworthia* genomes (Akman et al 2002; Rio et al. 2012). Upon excluding the saturating signals of *groEL* and *cspE* which introduce bias in the determination of the average TPM values, 65 loci were found to have ≥ 1.5 fold higher TPM values in comparison to the average TPM (fig. 3C and supplementary table S3, Supplementary Material online). Many of these genes are associated with roles as chaperones (*dnaK*, *groES*), proteases involved in the synthesis of the cell envelope (*hslU* and *ftsH*), supporting active division by *Wigglesworthia* within tsetse bacteriomes, and in cellular motility (*motA* and *fliC*). Additional genes, exhibiting ≥ 1.5 fold higher TPM values in comparison to the average TPM, included those putatively involved in protein translocation (*secE*), amino acid transport

(*Wigmor_0593*, *Wigmor_0594*), and multidrug transport pumps (*emrD*, a member of the major facilitator superfamily [MFS] and *mdlA*, a member of the ATP-binding cassette [ABC] superfamily), which have likely been repurposed for the transfer of molecules other than antibiotic drugs. In other Gram-negative bacteria, including *Pseudomonas aeruginosa*, *Neisseria gonorrhoeae*, and *Salmonella typhimurium*, these efflux pumps have been reused to excrete host-produced toxins, enabling the adaptation and survival within these niches (reviewed in Sun et al. [2014]). Loci involved in the recognition of environmental stimuli (the sensor histidine kinase *cpxA*) and towards B vitamin synthesis (i.e., thiamine [B1], riboflavin [B2], pyridoxine [B6], biotin [B7], and folate [B9]) also exhibited >1.5 fold TPM values. Further, loci involved in the assembly of [Fe-S] clusters (*sufCSE*) had >1.5 fold TPM values. In *Wigglesworthia*, the sulfur utilization factor (*suf*) unit is the only intact Fe-S cluster biosynthetic machinery available within the genome, with the exception of the cysteine desulfurase, *iscS*, traditionally associated with the iron sulfur cluster (Isc) formation system (Johnson et al. 2005). Fe-S clusters are present in proteins with a great variety of functions including electron transfer, posttranscriptional gene regulation, providers of free iron, DNA damage repair, and substrate binding during catalysis (reviewed in Brzoska et al. [2006]). Moreover, Fe-S containing proteins are direct players in many key aspects of metabolism, participating in the biosynthesis of essential cofactors and enzymes. For example, B vitamin metabolism involves Fe-S cluster containing enzymes. Specifically, the pyrimidine moiety for the thiamine cofactor is generated through the conversion of AIR into HMP-P, which is catalyzed by ThiC, an enzyme that holds a 4Fe-4S cluster (Raschke et al. 2007; Chatterjee et al. 2008). Thiamine biosynthesis also requires *iscS* as a S donor via ThiI (Palenchar et al. 2000), as well as the participation of another Fe-S protein called ThiH (Leonardi et al. 2003; Park et al. 2003; Leonardi and Roach 2004). Similarly, the last step in biotin synthesis is catalyzed by biotin synthase (BioB), another iron-sulfur cluster containing enzyme. At this step, the sulfur atom comes from one of two distinct iron-sulfur clusters present in the enzyme, a 2Fe-2S and a 4Fe-4S (Jarrett 2015), for the conversion of dethiobiotin to biotin (Ugulava et al. 2001; Jameson et al. 2004).

Lastly, 38 of the 55 putative transporters ($\sim 69\%$) identified within the *Wigglesworthia* genome using TRANSPORTERDB (Elbourne et al. 2017) had Illumina reads mapping back to these loci (supplementary fig. S3, Supplementary Material online). A total of 10 of these 38 ($\sim 26\%$) transporters were identified as having ≥ 1.5 TPM in comparison to the average TPM of bacteriome libraries, including the aforementioned efflux pumps (*mdlA* and *emrD*), a GlnQ family glutamine transport ATP-binding protein (*Wigmor_0594*), a cysteine ABC transporter substrate binding protein (*Wigmor_0593*) and several components of energy generating ATP synthase.

Table 1*Wigglesworthia* Genes Exclusively Expressed within the Bacteriomes of Either Female or Male *G. pallidipes*^a

COG Category	Gene	Sequence Identifier	Mean TPM	Description
Female				
Cell wall/membrane/ envelope biogenesis	<i>loID</i>	WIGMOR_RS00540	21.5	ATP binding component lipoprotein transporter
Nucleotide transport and metabolism	<i>pyrG</i>	WIGMOR_RS02070	13.9	CTP synthetase
Hypothetical		WIGMOR_RS03465	25.9	Putative membrane protein transporter
Male				
Nucleotide transport and metabolism	<i>trkA</i>	WIGMOR_RS00660	8.1	Potassium uptake protein
	<i>pyrI</i>	WIGMOR_RS00770	34.6	Pyrimidine biosynthesis
Replication, recomb. and repair	<i>dnaB</i>	WIGMOR_RS01295	11.6	DNA helicase
	<i>rseP</i>	WIGMOR_RS01905	9.7	Zinc metalloprotease
Defense mechanisms	<i>ahpC</i>	WIGMOR_RS01355	26.4	Alkyl hydroperoxide reductase
Cell cycle, control and division	<i>mrdB</i>	WIGMOR_RS3065	14.3	Cell elongation/ peptidoglycan synthesis

^aExpression observed in ≥ 2 libraries.

qPCR Validation of *Wigglesworthia* Gene Expression within Heterospecific Tsetse

To further verify the transcriptional activity of genes, we describe the expression of select genes within the bacteriomes of a heterospecific tsetse fly, *G. morsitans*. A total of 8 *Wigglesworthia* genes (*ribB* (3,4-dihydroxy-2-butanone-4-phosphate synthase), *ribC* (riboflavin synthase, alpha subunit), *bioB* (biotin synthase), *bioA* (adenosylmethionine-8-amino-7-oxononanoate aminotransferase), *mdIA* (putative ATP-binding component of multidrug ABC transporter), *cspE* (DNA binding transcriptional repressor), *fliC* (flagellin), and *Wigmor_0594* (GlnQ family glutamine transport ATP binding protein) identified by RNA-Seq analyses to have ≥ 1.5 fold greater than average TPM were chosen for real time quantitative gene expression assays. Because we were interested in identifying whether *Wigglesworthia* symbionts within different tsetse species also actively transcribed these genes and, further, whether specific adult ages and/or fecundity status were likely driving an increase in library TPM values (information that could not be captured with our field collections), these analyses were performed with RNA isolated from tsetse colony flies of known ages and mating status.

Similar to *Wigglesworthia* transcript abundance within *G. pallidipes* adult bacteriomes, the *G. morsitans* colony flies exhibited expression of all eight loci throughout adulthood (fig. 4). Only 1 gene, *bioB*, a biotin synthase, demonstrated no significant differences in gene expression upon the comparison of flies of different adult ages and female fecundity status. Although in both sexes, there was an increase in *bioB* expression as tsetse aged, a trend even more significantly apparent with *bioA* (involved in the conversion of 8-amino-7-oxononanoate to 7,8-diamino nonanoate within the biotin [B7] metabolic pathway), indicating a higher demand for symbiont-mediated production of biotin through the progression of tsetse adulthood. Further, there was a significantly higher expression of *Wigglesworthia bioA* within the bacteriomes of 2-week-old mated relative to same age virgin

females supporting an additional role for biotin during pregnancy. The *cspE* locus, encoding a cold shock protein/DNA binding transcriptional repressor, demonstrated significant differences in expression only between different aged males with 4-week-old males exhibiting the highest fold change in gene expression relative to younger males. Four other *Wigglesworthia* loci examined; *Wigmor_0594*, *ribB*, *ribC*, and *mdIA*, exhibited significant expression differences within the bacteriomes of female adults of different ages and mating status, but no significant differences upon the comparison of distinctly aged males, suggesting more prominent roles in female biology. The two *Wigglesworthia* loci involved in riboflavin (B2) synthesis, *ribB*, involved in the conversion of ribulose 5-phosphate to 3,4-dihydroxy-2 butanone-4-phosphate, and *ribA*, mediating the final step in riboflavin synthesis from 6,7-dimethyl-8-ribityl lumazine, had the highest level of expression within the bacteriomes of 2 week pregnant females. The higher expression of these genes suggests a greater need for symbiont-produced riboflavin during pregnancy potentially increasing maternal resources and aiding in embryogenesis. The two loci involved in transport functions, *mdIA* and *Wigmor_0594*, had significantly higher expression levels within the bacteriomes of 4-week-old virgin females and teneral and 2-week-old mated females, respectively. Interestingly, *fliC* had significantly higher expression within the bacteriomes of younger relative to older flies within both sexes ($P \leq 0.05$). Cumulatively, these results characterize the gene expression dynamics within a heterospecific tsetse and show that *Wigglesworthia* orthologs remain transcribed and may additionally vary in transcript abundance to adapt to the needs of different aged adults and fecundity status. These biological differences may have also driven the increase in read abundance within our RNA-Seq libraries from the field. Importantly, these analyses provide insight into specific symbiont biochemical processes that vary in activity depending on specific host traits that could not be obtained through the homogenization of bacteriomes of adult flies, which were

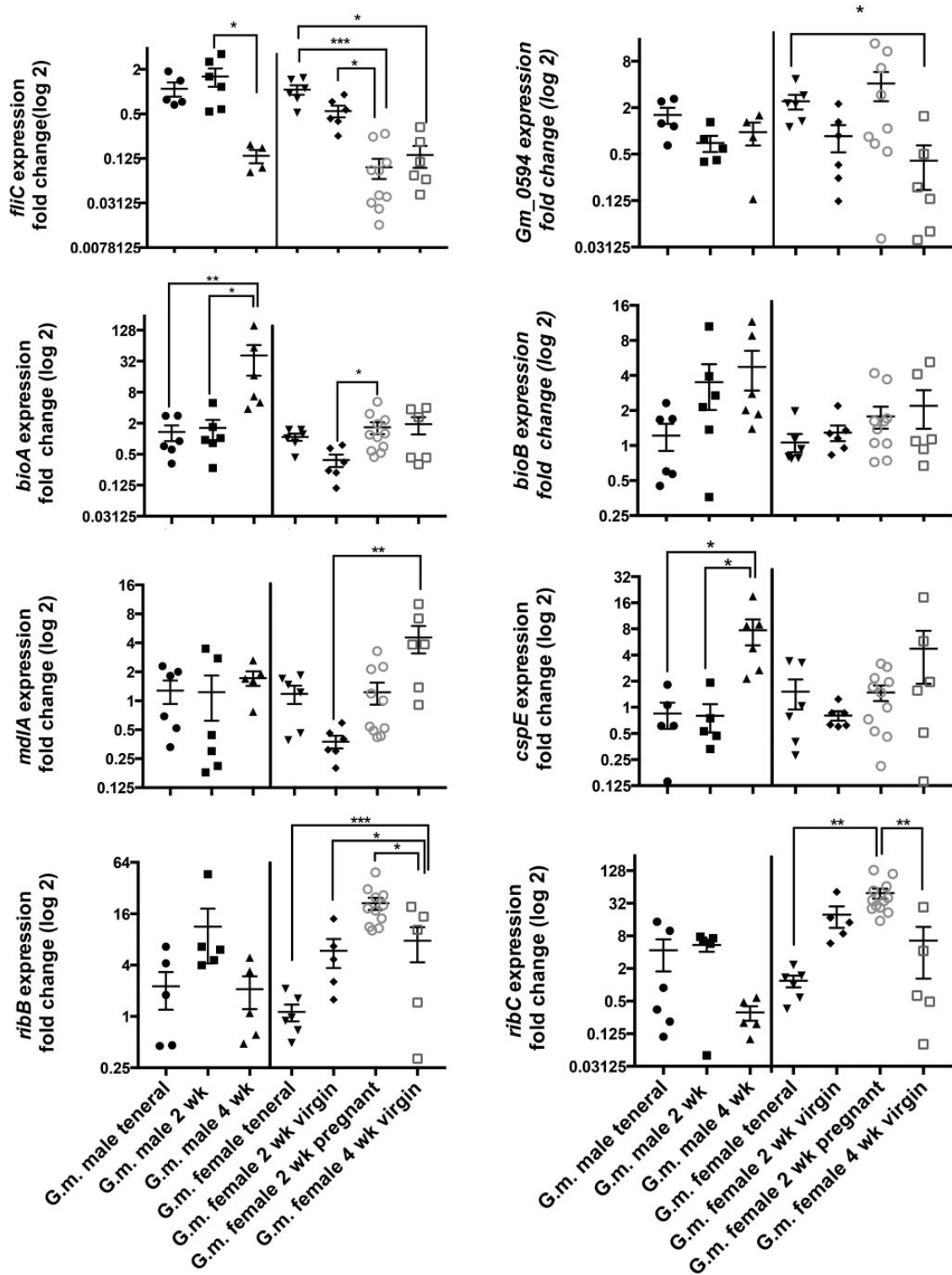


Fig. 4.—Expression kinetics of *Wigglesworthia* genes within a sister tsetse species, *G. morsitans morsitans*. qRT-PCR based fold change gene expression was analyzed through the threshold cycle ($2^{-\Delta\Delta CT}$) method. The results for *fliC*, *Wigmor_0594*, *ribB*, *ribC*, *mdIA*, *cspE*, *bioB*, and *bioA* were normalized to *rpsC* within the bacteriomes of various aged adult males, females and mating status and compared with same sex tenerals. Each point represents the expression (Average of triplicate Cts) of an individual fly with mean (represented by the horizontal bar) and ± 1 Standard Error of the Mean (SEM) indicated. $N > 5$ bacteriomes per group, asterisks indicate statistically significant differences between the bacteriomes. *** $P < 0.001$, ** $P < 0.01$ and * $P < 0.05$; Gm: *Glossina morsitans*, wk: week.

pooled regardless of age or mating status, within our field studies. Like many insect-symbionts, the small and compact genome of *Wigglesworthia* is a result of a long and complex evolutionary history associated with tsetse (reviewed in McCutcheon and Moran 2011). It is hypothesized that the majority of genes retained in *Wigglesworthia*'s small genome simply preserve the relationship with the host. Yet, the demands on the symbiosis, and consequently effects towards impacting symbiont gene expression, may vary in response to host age, mating and nutritional status.

G. pallidipes-Based Analyses

A total of 14,119 *G. pallidipes* gene transcripts had Illumina reads mapping back, representing ~71% of the total potential tsetse transcripts (16 Feb 2015, VectorBase, <https://www.vectorbase.org/organisms/glossina-pallidipes/iaea/gpali12>) being expressed within the bacteriome organ. Female flies had reads to 594 more gene transcripts than males (13,098 vs. 12,504), with 1,600 and 1,000 transcripts unique to females and male bacteriomes, respectively.

GO Annotation

BLAST2GO (Conesa et al. 2005) was used to assign 13,786 protein-coding tsetse genes to associated GO classifications for cellular components, molecular functions and biological processes. The resulting 23,771 GO terms ranged from levels 2 to 15 (supplementary fig. S4, Supplementary Material online), including parent and child terms, with 8,405 accessions within biological process, 10,840 in molecular function, and 4,509 in cellular components. A total of 18 GO terms for tsetse transcripts were obtained when examining root level 4, with four to cellular components, eight to biological processes, and six to molecular functions (fig. 5A). Among cellular components, integral component of membrane was the most enriched category (~32% of all genes within cellular components). Within biological processes, macromolecule metabolic process (~22% of all genes within biological processes) was the most enriched process. Lastly within molecular functions, nucleic acid binding (~19% of all genes within molecular functions) was the top process identified. Within all adult bacteriomes, and highly mirroring the *Wigglesworthia* symbiont profile (fig. 5A), there was a universal enrichment of expressed tsetse genes with GO terms associated with metabolism and transport (i.e., 44% of total GO categories at root level 4) likely facilitating the coordination and shuffling of metabolite production between these two organisms during the symbiosis. More specifically, GO category proportions between tsetse and *Wigglesworthia* have a positive correlation to each other when analyzed using simple linear regression (SLR) ($r = 0.4149$, $P = 0.0438$ for a test of 0 SLR slope). Lastly, multiple expressed tsetse loci were associated with roles in conveying substrates into and out of tsetse

bacteriocytes with a number relevant to vitamin transport including thiamine (thiamine pyrophosphate carrier proteins [GPAI038258 and GPAI008049], and thiamine transporter 1 [GPAI022398], riboflavin [solute carrier family 52, riboflavin transporter member 3-A {GPAI045057}], proton-coupled folate transporter [GPAI036223], reduced folate carrier [GPAI043750], solute carrier family 52, riboflavin transporter, member 3-A [GPAI045057] and multivitamin transporters [GPAI012112, GPAI014644, and GPAI028931]. Using Web Apollo [Lee et al. 2013] to identify orthologs within other tsetse species, we also confirmed that reduced folate carrier [GPAI043750] and thiamine transporter-1 (GPAI022398) were both expressed within the bacteriomes of adults of two other tsetse species, *G. morsitans* and *G. fuscipes* (fig. 6) supporting the significance of these genes throughout *Glossina*. Further, the expression of both these loci within the head samples of *G. fuscipes* indicates a systemic expression that may address the ubiquitous nature of vitamin requirements throughout the body.

To examine whether categories differed in their mean expression levels between males and females, GO classifications were averaged in their respective TPM values and comparatively analyzed (fig. 5B). Individual Mann–Whitney analyses found no significant differences ($P > 0.05$) in the mean TPM values of distinct GO categories expressed between tsetse male and female bacteriomes (fig. 5B). Interestingly, female flies had genes binning within the nucleus GO category ($n = 713$) where as males lacked any associated loci within this category. Within a sex, multiple GO categories significantly differed in their mean TPM values ($P < 0.0001$ for both males and female flies, supplementary table S4, Supplementary Material online contains detailed P values).

Tsetse Sex-Specific Differences in Gene Expression within Bacteriomes

To identify tsetse genes that were found to differ in expression within the bacteriomes of the different sexes, DESeq and Kallisto–Sleuth analyses were used for the comparison of transcriptome libraries. Kallisto–Sleuth identified 82 loci, whereas DESeq identified 32 genes that were differentially expressed with statistical significance, with 17 in consensus between the two approaches (fig. 5C and supplementary table S5, Supplementary Material online). Of the loci found to be in consensus between the two analyses, the majority (13/17, 76%) exhibited significantly higher expression within male bacteriomes, including orthologs to suppressor of zeste 2, Su(z)2 (GPAI038413), multiple trypsins (GPAI012727 and GPAI012728), loci involved in lipid metabolism; low density lipoprotein receptor adaptor protein 1 (GPAI030993) and lipin (GPAI004503), and orthologs to the previously described male significant (Fontaine et al. 2003; Blagden et al. 2009); outer dense fiber protein 3-like protein 2 (GPAI039186) and a La related protein (GPAI010220). Within female bacteriomes,

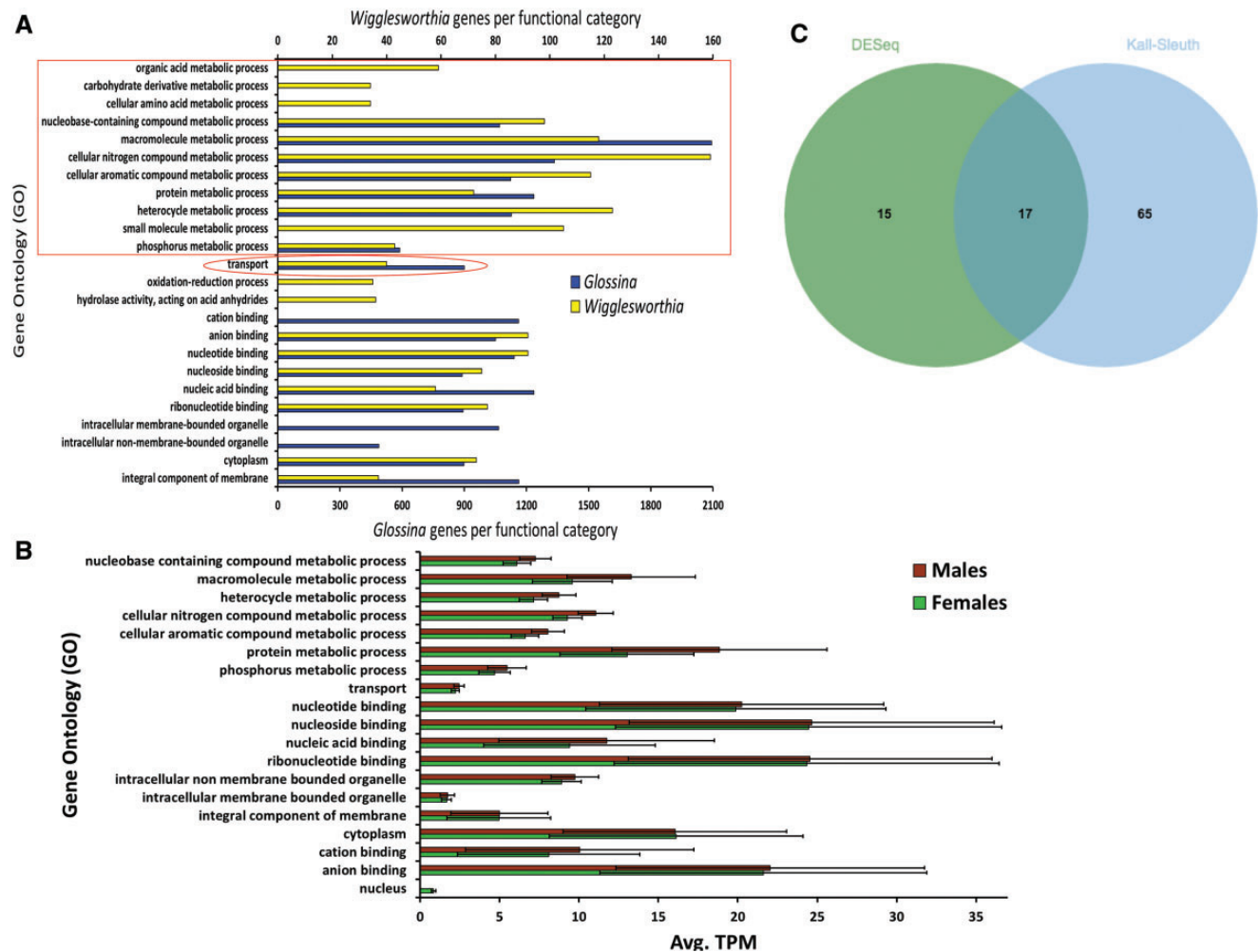


Fig. 5.—Gene ontology (GO) distribution of *G. pallidipes* transcriptome. (A) GO classifications at root level 4 for each *G. pallidipes* and *Wigglesworthia* gene expressed. The red box indicates GO classifications associated with metabolism and the circle highlights transport. (B) GO classifications of expressed tsetse genes normalized to average TPM values. There were no statistically significant differences in mean TPM values within a given category between male and female bacteriomes (Mann–Whitney; $P > 0.05$). A Kruskal–Wallis one-way analysis of variance (ANOVA) and Dunn’s multiple comparisons test were performed to determine whether mean TPM values differed between categories within a sex. F -tests were applied to assess the homogeneity of variances. The normality of TPM distributions was determined with a goodness-of-fit test. Mean TPM values were logtransformed to satisfy normality. SEMs are depicted. (C) A small number of differentially expressed *G. pallidipes* genes were found between female and male bacteriomes. A consensus of 17 *G. pallidipes* loci was found to significantly differ between male and female bacteriomes upon comparisons utilizing different analytical methods.

tsetse loci putatively involved in immunity related salivary c-type lectin (GPAI033252) and the transcriptional regulator protein FAM76A (GPAI025219) demonstrated significantly higher expression upon comparison to males.

Discussion

Animal alliances with bacteria are significant sources of biological diversification, having shaped evolutionary history (McFall-Ngai et al. 2013) as well as facilitating or constraining future adaptation to environmental change (Kiers et al. 2010; Wernegreen 2012; Kikuchi et al. 2016). Much of what we know about host–microbiota interactions is generated from the lab environment with the use of artificially bred or captive

animals. Here, we characterize the transcriptomes obtained through parallel Illumina deep sequencing of two obligate symbiotic partners, the tsetse fly, *G. pallidipes*, and its ancient bacterial symbiont, *Wigglesworthia*, from Kenyan field populations. Although much has been discovered on the functional contributions of this important member of the tsetse microbiota through empirical investigations of established tsetse fly lines, little research has been dedicated towards understanding the coordinated activities between partners in the wild.

Through empirical studies involving colony flies, *Wigglesworthia* symbionts have been identified as indispensable players in tsetse energy processing by nutritionally supplementing the strict diet of its host with B vitamins including thiamine (B1), pyridoxine (B6) and folate (B9) (Snyder et al.

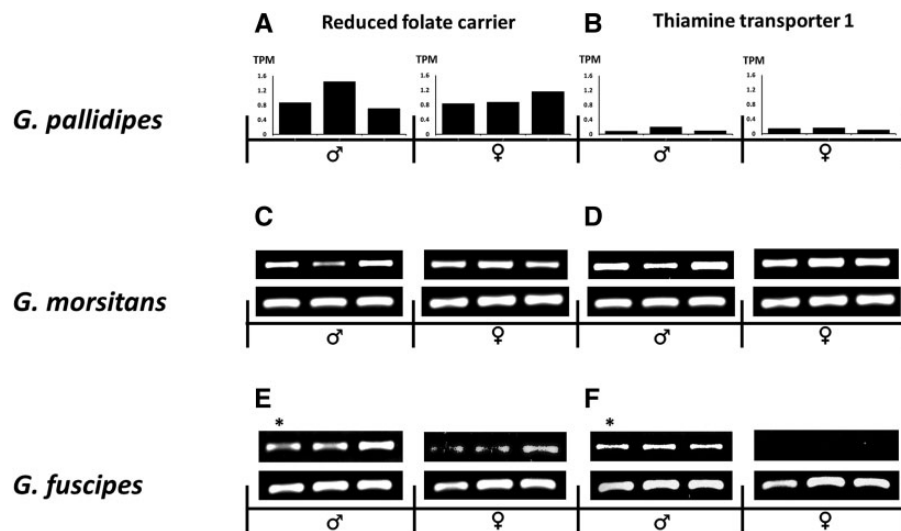


FIG. 6.—Semiquantitative RT-PCR analysis of tsetse transporters within tissues of different *Glossina* species. Columns show orthologs of reduced folate carrier (GPAI043750) and thiamine transporter 1 (GPAI022398). Rows correspond to the *Glossina* species indicated with orthologs identified using VectorBase and Web Apollo. The results shown are representative samples ($n > 6$) of each group. (A and B) TPM of the transporter loci in the six bacteriome libraries from Nguruman, Kenya field flies. (C–F) panels show the corresponding expression in two species of 2 week old tsetse flies from colony lines of *G. morsitans* (WVU) and *G. fuscipes* (Yale). Top row corresponds to the indicated transporter while the bottom row is β -tubulin expression (serving as a RNA integrity control). All samples are bacteriomes, except for those labeled with an asterisk (*), which represents RNA isolated from heads.

2010, 2012, 2015; Michalkova et al. 2014) which are lacking in vertebrate blood. As *Wigglesworthia* is essential for the development and fecundity of its tsetse host, we hypothesize that genes expressed within the bacteriome of field tsetse are integral components of the host–symbiont interface mediating nutrient synthesis and transfer, communication and regulatory control.

The interchange of metabolites within a microbial symbiosis is often the motivating factor for its formation and subsequent persistence. In the tsetse-*Wigglesworthia* association, the bacteria lie free within the cytoplasm of bacteriocytes (Aksoy 1995a), very different from many other endosymbionts that are surrounded by a secondary membrane which is host derived (Hinde 1971; von Dohlen et al. 2001; Baumann 2005). An additional distinction from other endosymbionts is that *Wigglesworthia*, despite drastic genome reductions, still encodes products that are integral to Gram-negative cell wall structure including lipopolysaccharide and peptidoglycan (Akman et al. 2002; Rio et al. 2012). Thus for *Wigglesworthia*-generated metabolites to gain access into the host cytoplasm of bacteriocytes, these substances would only have to pass through the bacterial dual membrane. Our field results indicate transcriptional activity for the majority of *Wigglesworthia* transporters and even more robust expression for transporters involved in multidrug efflux, glutamine and cystine ABC-binding and ATP synthases. These transporters are likely key mediators in the biochemical communication and metabolite translocation between partners. Following export from the *Wigglesworthia* cells, systemic distribution of provisioned nutrients is still necessary to address the

nutritional demands of other body regions. We demonstrate, that in both field and colony flies of different species, tsetse transcribe various transporters within the bacteriome whose orthologs have been characterized to be involved in the specific transfer of vitamins/cofactors and likely mediating their dispersal to other tissues. Further, these transporters are also expressed in other fly tissue, indicating a role in the widespread distribution of symbiont-produced nutrients. Future studies should examine how tsetse and *Wigglesworthia* coordinate these different metabolic and physiological steps, particularly in regards to time, space and stoichiometry. In an analogous insect system, involving aphids and their ancient obligate *Buchnera* bacterial symbionts, the insect hosts have been shown to control the biosynthesis of essential amino acids through substrate feedback inhibition, specifically by regulating the supply of the precursor glutamine into bacteriocytes (Price et al. 2014). Here, we also show, using flies of known age and mating status, *Wigglesworthia* transcriptional activity does fluctuate likely addressing differences in nutrient requirements through host age and reproduction.

The majority of genes found to be expressed by both *Wigglesworthia* and tsetse within bacteriomes are involved in metabolism and transport, indicating that, similar to what has been demonstrated with tsetse colony lines, the production and transfer of nutrients is also pinnacle to this association within the field. An obvious symbiotic exchange between tsetse and *Wigglesworthia* is in amino acids (*Wigglesworthia* is an auxotroph for the majority of amino acids) and B vitamins (blood meals are deficient) as indicated through genome analyses, several lab based empirical studies and with the field

studies described here. For instance, thiamine synthesis may either involve the integration of pyridoxal phosphate (B6), or purine metabolism, with cysteine that is likely provided by either the tsetse host or from the blood meal and imported through the *Wigglesworthia* cystine ABC transporter. These pathways converge for condensing the phosphorylated derivatives of 4-amino-5-hydroxymethylpyrimidine (HMP(PP)) and 4-methyl-5-(β -hydroxyethyl) (THZ(P)) moieties for the formation of thiamine monophosphate (TMP). TMP can then be supplied to tsetse and the neighboring *Sodalis* symbionts (Snyder et al. 2010, 2012), which are both thiamine auxotrophs. Similarly, the majority of enzymatic activity necessary for biotin and riboflavin biosynthesis was characterized within the bacteriome tissues of field flies. It is important to note that there were steps that were lacking within *Wigglesworthia* transcriptome libraries in B vitamin biosynthesis pathways that could be due to a multitude of factors including; contributions by the tsetse host or *Sodalis*, inadequacy in the depth of coverage of transcriptome libraries, the spatial/temporal stability of the mRNA and/or protein (i.e., shorter half-lives), or the possibility of a not yet described enzyme performing the necessary interconversion.

Symbionts may have major implications towards host epigenetics (Kim et al. 2016). For example, the parasitic *Wolbachia* manipulates a host microRNA to interfere with a DNA methyltransferase thereby affecting Dengue virus manipulation in *Aedes aegypti* (Zhang et al. 2013). It remains to be seen whether *Wigglesworthia* metabolite production may also affect tsetse epigenetics, especially when this symbiont is the major or sole provider of B vitamins. For example, biotin may be added to histones to induce posttranslational modification that affects gene expression by changing the accessibility of genes towards transcription factors (Smith et al. 2007; Filenko et al. 2011). Additionally, folate provides methyl groups used for DNA methylation (Crider et al. 2012). Within insects, epigenetic modifications have been associated with significant biological processes including aging (Lin et al. 2005; Lockett et al. 2016) and caste determination (Lyko et al. 2010). Our field studies not only increase our basic knowledge of tsetse biology, but additionally, strongly support results generated with colony lines in regards to the nutrient exchange involved in the symbiosis. The tsetse microbiota may ultimately be used for innovative symbiont-based control mechanisms beyond paratransgenesis (Aksoy et al. 2008). Research efforts focused on eliminating essential microbiota members or inhibiting their important functional roles for vector fitness, particularly by altering life span, fertility or vector competency, may provide new paradigms towards tsetse-specific suppression methods.

Supplementary Material

Supplementary data are available at *Genome Biology and Evolution* online.

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