RESEARCH ARTICLE

Insight into the Transmission Biology and Species-Specific Functional Capabilities of Tsetse (Diptera: Glossinidae) Obligate Symbiont Wigglesworthia

Rita V. M. Rio, Rebecca E. Symula, Jingwen Wang, Claudia Lohs, Yi-neng Wu, Anna K. Snyder, Robert D. Bjornson, Kenshiro Oshima, Bryan S. Biehl, Nicole T. Perna, Masahira Hattori, and Serap Aksoy

Department of Biology, West Virginia University, Morgantown, West Virginia, USA; Department of Ecology and Evolutionary Biology, Yale University, New Haven, Connecticut, USA; Division of Epidemiology of Microbial Diseases, School of Public Health, Yale University, New Haven, Connecticut, USA; Department of Computer Science and W. M. Keck Biotechnology Resource Laboratory, Yale University, New Haven, Connecticut, USA; Graduate School of Frontier Sciences, The University of Tokyo, Kashiwa, Chiba, Japan; and Department of Genetics, University of Wisconsin—Madison, Madison, Wisconsin, USA

ABSTRACT Ancient endosymbionts have been associated with extreme genome structural stability with little differentiation in gene inventory between sister species. Tsetse flies (Diptera: Glossinidae) harbor an obligate endosymbiont, Wigglesworthia, which has coevolved with the Glossina radiation. We report on the ~720-kb Wigglesworthia genome and its associated plasmid from Glossina morsitans morsitans and compare them to those of the symbiont from Glossina brevipalpis. While there was overall high synteny between the two genomes, a large inversion was noted. Furthermore, symbiont transcriptional analyses demonstrated host tissue and development-specific gene expression supporting robust transcriptional regulation in Wigglesworthia, an unprecedented observation in other obligate mutualist endosymbionts. Expression and immunohistochemistry confirmed the role of flagella during the vertical transmission process from mother to intrauterine progeny. The expression of nutrient provisioning genes (thiC and hemH) suggests that Wigglesworthia may function in dietary supplementation tailored toward host development. Furthermore, despite extensive conservation, unique genes were identified within both symbiont genomes that may result in distinct metabolomes impacting host physiology. One of these differences involves the chorismate, phenylalanine, and folate biosynthetic pathways, which are uniquely present in Wigglesworthia morsitans. Interestingly, African trypanosomes are auxotrophs for phenylalanine and folate and salvage both exogenously. It is possible that W. morsitans contributes to the higher parasite susceptibility of its host species.

IMPORTANCE Genomic stasis has historically been associated with obligate endosymbionts and their sister species. Here we characterize the Wigglesworthia genome of the tsetse fly species Glossina morsitans and compare it to its sister genome within G. brevipalpis. The similarity and variation between the genomes enabled specific hypotheses regarding functional biology. Expression analyses indicate significant levels of transcriptional regulation and support development- and tissue-specific functional roles for the symbiosis previously not observed in obligate mutualist symbionts. Retention of the genetically expensive flagella within these small genomes was demonstrated to be significant in symbiont transmission and tailored to the unique tsetse fly reproductive biology. Distinctions in metabolomes were also observed. We speculate an additional role for Wigglesworthia symbiosis where infections with pathogenic trypanosomes may depend upon symbiont species-specific metabolic products and thus influence the vector competence traits of different tsetse fly host species.

Microbial symbioses with eukaryotic hosts are ubiquitous. Despite the wide prevalence of symbiosis, obligate associations, where partners are inextricable from and entirely dependent on one another, are relatively rare. Endosymbionts residing within host cells are typically vertically transmitted between generations with high fidelity, coupling partners, and this results in an intimate, specialized association over evolutionary time. Such symbiotic associations can enable hosts to acquire new metabolic capabilities and thus thrive in novel niches. This idea was first hypothesized for insects that subsist on nutrient-poor phloem sap, where symbionts supplement dietary deficiencies (1). Similar nutritional symbioses have since been identified in many insects with nutritionally restricted diets, including tsetse flies, carpenter ants, and many plant-feeding insects (2). Based on whole-genome sequences, the obligate symbiont genomes have all been drastically reduced in size in comparison to those of their free-living
relatives and display high A+T bias (3, 4). These traits are thought
to have arisen through relaxed natural selection and resulting
genome deterioration (5). The obligate symbionts of aphids,
carpenter ants, and tsetse flies, *Buchnera*, *Blochmannia*, and
*Wigglesworthia*, respectively, form a close lineage in the gamma-
proteobacteria and are believed to have independently established
host relations. Despite their close relatedness, the extant genomes
of the symbionts have undergone drastic, yet distinct, adaptive
reductions. It is thought that genes retained in each small genome
are necessary for functional capabilities that complement host
physiology and ecology, with gene inventory having some speci-
city to the host lineage (5).

Every tsetse fly species is associated with a distinct *Wiggleswor-
thia glossinidius* lineage (6). Phylogenetic analysis of *Wiggleswor-
thia* shows concordant history between symbiont lineages and
their host species, indicating partner coevolution. Further, molec-
ular clock methods suggest that this symbiosis is about 80 million
years old (6). In addition to *Wigglesworthia*, tsetse fly laboratory
colonies and some natural populations can harbor a commensal
bacterial symbiont, *Sodalis glossinidius*, of relatively recent estab-
lishment (7, 8) and parasitic *Wolbachia* infections (9, 10).

Tsetse flies feed exclusively on nutrient-poor vertebrate blood.
Unlike many other insects, tsetse flies display viviparous repro-
duction (deposition of live late-stage larvae rather than eggs),
where the mother develops a single oocyte at a time and then
carries and nourishes the resulting embryo and larva in an intra-
uterine environment. The mother undergoes parturition to a fully
developed larva that quickly pupates and remains dormant for
about a month prior to adult metamorphosis. Thus, throughout
its developmental cycle, the tsetse fly is solely dependent on its
vertebrate host blood diet. The obligate mutualist *Wigglesworthia*
is thought to complement the exclusive blood diet of its host.

*Wigglesworthia* resides intracellularly in bacteriocytes, which
form the bacteriome organ in the anterior midgut (see Fig. 1). In
the bacteriocyte cytoplasm, the symbionts live free and are not
surrounded by host membranes. In addition to the bacteriome,
extracellular *Wigglesworthia* is also detected in the milk gland lu-
men (11, 12). Extracellular *Wigglesworthia*, along with *Sodalis*, is
maternally transmitted to the tsetse fly’s intrauterine progeny
through milk secretions synthesized via the modified accessory
gland (milk gland) that connects to the uterus (13). Without
*Wigglesworthia*, tsetse fly females are reproductively sterile. Given
that the exclusive tsetse fly blood diet is low in vitamins, coupled
with data from dietary supplementation experiments of
antibiotic-fed (symbiont-free) tsetse flies, a putative role in vita-
mim metabolism has been suggested for the symbionts (14).

**FIG 1** Localization of *Wigglesworthia* within the tsetse fly. *Wigglesworthia* resides within the bacteriome organ (top left) and is found free in the cytoplasm of specialized cells known as bacteriocytes (top right). (Bottom) *Wigglesworthia* is present extracellularly in the milk gland tissue. (Bottom left) Fluorescence in situ hybridization (FISH) staining for *Wigglesworthia*. (Bottom right) Schematic drawing based on FISH results shown on the left. DAPI staining indicates nuclei in bacteriocytes and milk gland tubule cells. Pink fluorescent rhodamine staining shows *Wigglesworthia* within bacteriocytes and in milk gland lumen.
addition to host dietary supplementation through vitamin provisioning, the presence of *Wigglesworthia* is essential for the tsetse fly’s immune system maturation. It has been possible to develop flies that lack *Wigglesworthia* by maintaining fertile tsetse fly females on ampicillin-supplemented blood meals (11). This is because the antibiotic ampicillin does not affect the intracellular forms of *Wigglesworthia* within the bacteriome but can clear the extracellular *Wigglesworthia* from tsetse fly milk. The resulting progeny of ampicillin-treated females lack *Wigglesworthia* (*Gmm*<sup>mWig</sup>) but retain the commensal *Sodalis*. In comparison to their normal counterparts, *Gmm*<sup>mWig</sup>-adults are highly susceptible to trypanosome midgut infections (11) and microbial challenge (15). Thus, it appears that when larvae develop without *Wigglesworthia*, cellular immunity is particularly compromised in the emerging adult progeny (15).

The genome of *Wigglesworthia gossinidia* characterized from *Glossina brevipalpis* (referred to here as WGB) is about 697 kb in size and has a small plasmid (pWgb). The genome encodes 621 predicted coding sequences (CDSs) and displays a high (82%) adenine-thymine (A+T) bias (16, 17). It is possible that the high A+T content of *Wigglesworthia* resulted from the loss of repair and recombination functions such as the SOS, base excision, and nucleotide excision repair system (*uvrABC*). Surprisingly, the important gene coding for the DNA replication initiation protein DnaA was missing from the WGB genome—an observation previously unprecedented in eu- bacteria. More than 10% of the retained CDSs are involved in the biosynthesis of cofactors, prothetic groups, and carriers, supporting *Wigglesworthia*’s genetic contributions in the *de novo* metabolism of biotin, thia-zole, lipoic acid, flavin adenine dinucleotide (riboflavin, B<sub>2</sub>), folate, pantothenate, thiamine (B<sub>1</sub>), pyridoxine (B<sub>6</sub>), and protoheme and further substantiating the role of *Wigglesworthia* in host dietary supplementation (14). In addition to providing its tsetse fly host with vitamins, comparative genome studies with *Sodalis* indicate that *Wigglesworthia* may also provide thiamine to *Sodalis*, which lacks the thiamine biosynthetic pathway but has retained the transporter for acquiring thiamine (18). The functional complementation of symbiont genomes has been postulated to reduce competition between microbes, as well as prevent the possibility of symbiont genome replacement especially during the early establishment of a dual symbiosis (2).

Here we describe the genome of a second *Wigglesworthia* species isolated from *Glossina morstans morstans* (referred to here as WGM). Phylogenetic molecular clock analyses suggest that tsetse fly host species of WGB and WGM have been distinct for 50 to 80 million years (6). We compare the genome structures and gene inventories of WGM and WGB and explore evolutionary patterns in the genes which may contribute to functional variation within their respective tsetse fly host species. We describe the expression of *Wigglesworthia* genes that may be significant for tsetse fly nutrition through development. Lastly, we provide support for the role of flagella during the crucial symbiont maternal transmission process. We discuss similarities and differences between the two genomes that may ultimately affect important host physiological processes, including varying vector competence of the tsetse fly host species.

**RESULTS**

**Features of the WGM genome.** The genome of WGM consists of a circular chromosome of 719,535 bp (with a guanine-plus-cytosine [G+C] content of 25%) and a single plasmid of 5,198 bp. The putative origin of replication, without a clear G+C skewing and diagnostic DnaA boxes, was assigned to the same A+T-rich region upstream of the *gidA* locus as WGB (Fig. 2A and B). Table 1 summarizes the general features of the WGM genome, relative to those of other insect endosymbionts, including that of WGB, the relatively recently established genus *Sodalis*, the related ancient obligate symbiont, *Blochmannia*, of carpenter ants, and two *Buchnera* symbionts from different aphid hosts, respectively (7, 16, 19–21). Both the genome size and the exceptionally low G+C content of WGM were comparable to those reported for other ancient endosymbionts, including WGB. Annotation revealed that, similar to the other small obligate genomes, the coding content of WGM is high (83.9%) with 620 predicted CDSs at an average length of 979 bp (Table 1). The high A+T bias of the WGM chromosome was reflected in the higher average predicted isoelectric points of putative proteins, as was noted in WGB (9.84 in obligates versus 7.2 in *Sodalis*). Like WGB, WGM has two identical copies of each of the rRNA genes (*rrsH* and *rrlB*) (Fig. 2B). Similar to those of other symbionts, these rRNA genes have higher G+C contents, 49.3% and 45.8%, respectively, than protein coding genes. A total of 11 recognizable pseudogenes were identified in WGM (see Table S1 in the supplemental material), and these were distributed throughout the genome.

**Comparison of WGM and WGB genome structures.** Alignment of the two *Wigglesworthia* genomes indicates high chromosomal synteny, as was previously described for the *Buchnera* (22) and *Blochmannia* (23) genomes. However, since the divergence of WGB and WGM, a chromosomal inversion has occurred in one of the lineages (Fig. 2A). The inversion, which can be interpreted as an either 550-kb or 170-kb inversion due to the circular chromosome of *Wigglesworthia*, occurs approximately 150 kb from the *gidA* locus, in proximity to the origin of replication (Fig. 2B). An inversion in proximity to the origin of replication could create imbalanced replicches between the *Wigglesworthia* genomes. The inversion is flanked on either end by the rRNA genes *rrsH* and *rrlB*. In both *Wigglesworthia* genomes, within this G+C-rich region, and specifically within the *rrsH* gene, is a sequence that is nearly identical to the *Escherichia coli* Chi recombination hot spot (24). The sequence differs by only a single base (in bold): *E. coli*, GCTGGTGG; *Wigglesworthia*, TCTGGTGG. Since the RecA protein, which has been retained in both *Wigglesworthia* genomes, has the highest affinity to the TGG repeats in *E. coli* (25), we propose that this site (~480 bp into *rrsH*) likely demarcates the inversion site.

**Comparative analysis of WGB and WGM contents.** Comparative analyses of the WGB and WGM genomes reveal a shared set of 599 CDSs (Fig. 3A). Both genomes have retained pathways involved in B vitamin biosynthesis, including biotin (B<sub>7</sub>), thiazole (B<sub>1</sub>), riboflavin (B<sub>2</sub>), pantothenate (B<sub>5</sub>), and pyridoxine (B<sub>6</sub>). However, genetic components involved in the synthesis of co-balamine (B<sub>12</sub>) and nicotinate (B<sub>3</sub>) appear to be absent from WGM and WGB. Genes necessary for the synthesis of a complete flagel-lum apparatus have also been preserved in both genomes. Since the release of the WGB genome annotation, genes exhibiting high sequence identity to the Wg001 to Wg003 orphan genes have been reported in other host-associated bacteria. Wg001 to Wg003 are homologous to a putative transmembrane protein, an endonu-
FIG 2  Linearized comparison of the genomes of WGB and WGM. (A) Genome alignment as represented from Mauve. Each colored block represents a locally collinear block (LCB) of DNA that has not undergone rearrangement within its boundaries. Bar height indicates average nucleotide similarity within a region. The green LCB is inverted, as indicated by the relative reverse orientation of the block in each genome. (B) Annotation of the WGM genome. Each gene is shown as a trapezoid with the straight side representing the start codon. Genes above the line are on the positive strand, while genes below the line are on the negative strand. The shaded area indicates the inverted region relative to the WGB genome. Genes are labeled and color coded according to the different functional categories assigned.
clease, and an integral membrane protein, respectively. Notably, these genes have also been retained in WGM. Unlike most bacteria, WGM and WGB both lack dnaA, suggesting gene loss in the ancestral lineage prior to host diversification.

Unique genes (i.e., those lacking in the sister genome) were identified in WGM and WGB (Fig. 3B compares the unique gene inventories). Notably, significant differences in the distribution of functional categories of unique genes were observed between WGB and WGM (Kolmogorov-Smirnov test, \(p < 0.05\)). These 19 and 21 genes, respectively, and their putative biological roles are listed in Table S2 in the supplemental material. In addition, the positions of these genes within the WGM genome are highlighted in Fig. 2. The retention of these unique genes suggests metabolic distinctions within the proteomes of the two Wigglesworthia sister species. Notably, analysis of the WGM-specific gene set reveals the presence of a complete shikimate biosynthetic pathway in which 3-deoxy-d-arabino-heptulosonate-7-phosphate can be converted into chorismate (see Fig. S2 in the supplemental material). This pathway is degraded in the WGB genome, where only aroG and aroK homologs are still detectable. Acting downstream of the chorismate pathway, WGM also contains pabA, pabB (encoding aminodeoxychorismate synthases II and I, respectively), and pabC (encoding 4-amino-4-deoxychorismate lyase), which catalyzes the reaction from chorismate to \(p\)-aminobenzoate, an essential component in folate biosynthesis (see Fig. S2 in the supplemental material). The WGM genome also contains an aspC homolog that can also be used following chorismate biosynthesis toward phenylalanine production.

WGM plasmid and putative functions. The WGM plasmid (pWgm) is 5,198 bp in length with a G+C content of 24%. pWgm has six CDSs (Fig. 3A in the supplemental material), which are homologous to the six CDSs previously identified in pWgb (16). The four CDSs, with only a minor frequency of indels, encode a spermidine acetyltransferase (pWgm open reading frame 1 [ORF1], 177 amino acids [aa]; WGPWb0002, 174 aa), a putative mechanosensitive channel protein (encoded by yggB; pWgm ORF3, 282 aa; WGPWb0005, 280 aa), a putative heat shock protein (pWgm ORF5, 137 aa; WGPWb0006, 133 aa), and a conjugative transfer surface exclusion lipoprotein (pWgm ORF6, 243 aa; WGPWb0003, 239 aa). The remaining two CDSs, which exhibit higher sequence variation, encode a replication protein A,

### Table 1: Comparison of genome features of various insect bacterial endosymbionts

| Species described | Chromosome size (bp) | No. of plasmids [size (bp)] | Protein coding (%) | G+C content (%) | No. of predicted CDSs | No. of rRNA operons | tRNA | No. of pseudogenes | Average ORF length (bp) |
|-------------------|----------------------|-----------------------------|-------------------|----------------|----------------------|---------------------|------|------------------|------------------------|
| Wigglesworthia spp. |                      |                             |                   |                |                      |                     |      |                  |                        |
| W. morsitans      | 719,535              | 1 (5,198)                   | 83.9              | 25             | 620                  | 2                   | 34   | 11               | 979                    |
| W. brevipalpis    | 697,742              | 1 (5,280)                   | 89                | 22.5           | 618                  | 2                   | 34   | 8                | 988                    |
| Sodalis glossinidiusb | 4,171,146          | 3 (121,356)                 | 50.9              | 54.7           | 2,432                | 7                   | 69   | 972              | 873                    |
| Blochmannia floridanusc | 705,557          | 0                           | 83.2              | 27.4           | 583                  | 1                   | 37   | 6                | 1,007                  |
| Bacteroides strains |                    |                             |                   |                |                      |                     |      |                  |                        |
| BAp                | 640,681              | 1 (15,044)                  | 88                | 26.3           | 583                  | 1                   | 32   | 12               | 988                    |
| BSGf               | 641,454              | 2 (~22,267)                 | 84.5              | 26.2           | 545                  | 1                   | 38   | 33               | 978                    |

a As reported by Akman et al. (16) and with our own revision.

b As reported by Toh et al. (7).

c As reported by Gil et al. (21).

d Bacteroides strains BAp and BSG were selected because of similar 16S rRNA genetic distances between isolates relative to W. morsitans and W. brevipalpis.

e B. aphidica strain isolated from Acyrthosiphon pisum as reported by Shigenobu et al. (19).

f B. aphidica strain isolated from Schizaphis graminum as reported by Tamas et al. (20).

g Tryptophan plasmid size as reported by Lai et al. (62).
RepA (pWgm ORF2, 239 aa), that has a 36-aa deletion at the 5’ end in comparison to its pWgb homolog, and a hypothetical protein (pWgm ORF4, 311 aa) that contains 12 nonsynonymous changes occurring within the first 20 aa of the sequence relative to its pWgb homolog.

**Maternal transmission of Wigglesworthia to intrauterine progeny.** Similar to the WGB genome, that of WGM has retained the capacity to synthesize functional flagella. To better understand the biological role of flagella, we quantified transcripts for flagellin (fliC), which encodes the filament subunit of bacterial flagella, and motility protein A (motA), which confers motility functions on flagella, using quantitative reverse transcription-PCR (qRT-PCR) and immunohistochemistry approaches (Fig. 4). We quantified gene expression in the maternal gut bacteriome organ, in different stages of intrauterine larvae (L1 to L3) and in the corresponding mothers’ carcasses representing milk glands, and in young (newly deposited) and old (prior to eclosion) pupae. Within the tsetse fly mother, motA and fliC were expressed only in the carcass, apparently by WGM bacteria that are extracellular in the milk gland organ (Fig. 4A). We also detected the expression of flagellar components in the intrauterine larvae carried by the mothers and in the young pupae immediately postdeposition (Fig. 4A). The expression levels of both flagellar genes were highest in the L1 stage of the intrauterine larvae and in the carcasses (milk glands) of the corresponding mothers. Flagellum-specific expression in larvae decreased during development and was lowest during pupal development. Neither fliC nor motA expression was detected in adult bacteriomes. Immunohistochemistry analysis with antibodies specific for WGM flagellin also confirmed the expression profile. No flagellin was detected in the intracellular WGM in the maternal gut bacteriome, whereas flagellin was observed in milk gland cells and in the newly formed bacteriome organ in the intrauterine larva (Fig. 4B).

**Functional biology of Wigglesworthia.** To understand the regulation and functions of Wigglesworthia genes during different host developmental stages, we examined the expression profiles of two genes (hemH and thiC) associated with heme and thiamine biosynthesis; respectively, which may be involved in host nutrient supplementation. We also evaluated the expression of groEL, which encodes a chaperonin that may compensate for the higher-frequency protein misfoldings typically associated with an accelerated mutation rate (Fig. 5). Interestingly, all of the genes except groEL exhibited tissue- and host development-specific transcriptional regulation. The thiC and hemH genes showed similarities in their transcriptional regulation, and their expression was highest during the pupal stage of host development. However, thiC expression was higher than hemH expression in the adult bacteriome organ (intracellular stage). In contrast, during intrauterine larval development (L1 and L2), the hemH level was significantly higher than that in the adult bacteriome. Interestingly, the hemH levels in the different larval stages (L1 to L3) were similar to those observed in the corresponding milk gland samples obtained from the mother (MomL1 to MomL3; Fig. 5). The chaperonin encoded by groEL was expressed more consistently throughout all of the host stages examined, presumably due to its required assistance in protein folding throughout host development. Thus, the symbiont genes analyzed were subject to spatial and temporal transcriptional regulation during host development.

**Molecular evolution.** Rates of synonymous (dS) and nonsynonymous (dN) nucleotide changes in genes common to WGM and WGB were estimated to identify potential targets of selection. Both the dN and dS methods (dnds and dndsml) estimated com-

![FIG 4](image_url)
parable values (see Table S3 in the supplemental material), so the data were discussed without the application of a maximum-likelihood-based correction. Rather than a direct inference of positive selection, we identified genes that have a higher rate of non-synonymous change than the rest of the genome. Twenty-one gene comparisons were excluded from the dN and dS analysis due to length differences of >100 bp (see Table S3 in the supplemental material). Only one of these genes was found in the six plasmid orthologs. Of these 21 genes, 7 had dN and dS values that were greater than 2 standard deviations above the mean, and most of these genes had large deletions, suggesting that these loci are not under positive selection but are undergoing degradation. Potential targets of selection are summarized in Table 2. Only two genes (cspE and acpP) were identified as likely targets of purifying selection, although a few additional genes had relatively lower dN and dS values (Table 2; Fig. 6; see Table S3 in the supplemental material). Cold shock proteins (cspE) are associated with the maintenance of cellular function in cold temperatures and have been observed to function under osmotic stress (26). Acyl carrier proteins (acpP) are involved in cellular metabolism, particularly in fatty acid synthesis (27). Twenty-one genes were found to have significantly high values of dN and dS relative to the remainder of the genome (mean dN and dS = 0.2617), although only a single gene (fliK) was found to have a dN and dS value of >1, suggesting the influence of positive selection on this gene (Table 2). Importantly, genes that are potentially targets of selection were not restricted to a single area of the genome (Fig. 6) and one of these genes was found in the plasmid (WgpWb003). The functions of the genes varied, but notably, three genes with relatively higher dN and dS values that were involved in flagellar biosynthesis (fga, fliM, fliK) and six cell surface-associated genes (ppiD, yraP, ompF, yfO, tolA, and imp) were included. The plasmid gene WgpWb003 has homology to the traT gene that encodes a highly cell surface-exposed lipoprotein specified by F-like plasmids known to impede the conjugative transfer of similar or identical plasmids (28). These loci encode proteins that are exposed to the host environment, and thus, the higher dN and dS values may indicate selection for varying host immunological backgrounds.

DISCUSSION

Despite almost 80 million years of evolutionary distance, comparative analyses of the WGM and WGB genomes reveal similarly reduced size, almost complete synteny with the exception of one large inversion event, and a large set of genes shared by the two symbiont species. This is largely analogous to what has been described for other obligate insect endosymbionts (20, 23). This genome evolutionary process noted in obligate symbionts is distinct from what has been observed for free-living microbes, where significant diversity is driven by horizontal gene transfer on a background of gradual genome sequence drift. Despite high conservation, the two Wigglesworthia genomes display several unique capabilities, which are indicative of an adaptive evolutionary process. In particular, the putative proteomes indicate various metabolic capabilities in chorismate, phenylalanine, and folate biosynthesis, which in turn may affect their host physiology, including host vector competence (ability to transmit pathogenic trypanosomes). Furthermore, distinct from other obligate mutu-

FIG 5  qRT-PCR of hemH, groEL, and thiC expression during tsetse fly development. Gene expression results are shown for hemH (A), groEL (B), and thiC (C) in the adult bacteriome (BAC), different stages of intrauterine larvae (L1, L2, and L3), newly deposited pupae (Pu-early), late pupae (Pu-late), and carcasses of mothers that carry the corresponding intrauterine larvae (Mom-L1, Mom-L2, and Mom-L3). All data were normalized to the ribosomal gene rpsC Asterisks indicate statistically significant differences between the bacteriome and different developmental stages. *** P < 0.0001; * P < 0.05.
alist symbiont systems, Wigglesworthia displays significant transcriptional regulation, including the expression of functional flagella in the extracellular forms present in mother’s milk, which are apparently transmitted to host progeny. Significant levels of gene regulation at the transcriptional level have not been previously described in other ancient endosymbionts.

Eleven WGM genes demonstrated extensive (≥50%) truncation compared to their E. coli orthologs and were annotated as pseudogenes (see Table S1 in the supplemental material). Only three of these pseudogenes (ftsK, nusB, and nlpB) are similarly truncated in the WGB genome, suggesting ongoing, but relatively minor, gene degradation. Previously, Degnan et al. (23) questioned the pseudogene designation within the Blochmannia genome since the sequence conservation of affected genes suggests that they may still potentially encode functional proteins. Similarly, a majority of WGM pseudogenes (i.e., 6 out of 11) have frameshifts consisting of only 1 or 2 indels, including WGM thiI, which has previously been shown to be transcribed within adult bacteriomes (18). Various molecular processes occurring during transcription or translation may restore protein function in asso-

### Table 2: Summary of genes found to have notable dN/dS ratios

| Gene  | p distance | Genome position | G+C content | dN/dS ratio | Function                                      |
|-------|------------|-----------------|-------------|-------------|-----------------------------------------------|
| cspE  | 0.0931     | 632650          | 0.3286      | 0.0000      | Cold shock protein                            |
| acpPb | 0.0109     | 101980          | 0.3376      | 0.1325      | Acyl carrier protein                          |
| ppID  | 0.3635     | 662377          | 0.1675      | 0.5300      | Outer membrane protein folding                |
| tif   | 0.3235     | 384266          | 0.1907      | 0.5335      | Protein chain elongation                      |
| flagA | 0.4000     | 45952           | 0.2153      | 0.5377      | Flagellar biosynthesis                        |
| yrB   | 0.3957     | 454967          | 0.1657      | 0.5387      | Unknown                                       |
| cyoD  | 0.3519     | 665149          | 0.1545      | 0.5519      | Electron carrier                              |
| rnmF  | 0.2965     | 568641          | 0.2259      | 0.5526      | Methyltransferase                             |
| recC  | 0.3888     | 530384          | 0.1648      | 0.5547      | DNA helicase                                  |
| flIM  | 0.3302     | 58850           | 0.1729      | 0.5598      | Flagellar energizing component                |
| yraP  | 0.3909     | 311493          | 0.1968      | 0.5657      | Lipoprotein                                   |
| ygFZ  | 0.3791     | 327763          | 0.1849      | 0.5674      | Folate-dependent regulation                   |
| ygB   | 0.3389     | 304193          | 0.1685      | 0.5927      | Transmembrane protein                         |
| folB  | 0.4453     | 297053          | 0.1861      | 0.5936      | Dihydronicotin aldolase component             |
| ampF  | 0.4047     | 522497          | 0.1680      | 0.6185      | Outer membrane porin                          |
| yfO   | 0.3556     | 214973          | 0.1380      | 0.6389      | Outer membrane protein component              |
| holAc | 0.4176     | 639753          | 0.1293      | 0.6892      | DNA polymerase subunit                        |
| tolAc | 0.3966     | 360872          | 0.1627      | 0.7200      | Membrane-anchored protein                     |
| ycaZc | 0.4678     | 406494          | 0.1746      | 0.7509      | Putative protease                             |
| tig   | 0.4068     | 656418          | 0.1369      | 0.7526      | Molecular chaperone                           |
| impC  | 0.4183     | 20329           | 0.1418      | 0.8048      | Envelope biosynthesis                         |
| flKc  | 0.4699     | 60369           | 0.1383      | 1.0489      | Flagellar hook protein                        |
| WgpWb003 | 0.3717 | Plasmid         | 0.2069      | 0.5659      | Transfer surface lipoprotein                  |

* a: Genes without superscripts have dN/dS ratios that are ≥2 standard deviations above the mean.
  b: Gene with a notably low dN/dS ratio.
  c: Gene ≥3 standard deviations above the mean.
  d: p distance is the proportion of differences between the two species.
  e: G+C content corresponds to that of the WGM gene copy.

**FIG 6** Summary of dN/dS ratio calculations relative to genome position. Genes putatively influenced by purifying selection are represented by blue diamonds. Genes within 2 standard deviations of the mean dN/dS ratios are represented by red squares. Genes with >2 and >3 standard deviations from the mean dN/dS ratio are represented by yellow triangles and green circles, respectively.
evolution with the ident genome suggest early acquisition followed by a lengthy co-background. Wigglesworthia both the WGM and WGB genomes suggests their importance in ecology. The retention of these genes and genome elements by reported for the gene content and order suggests exposure to similar evolutionary formity between pWgm and pWgb in size, G/C content, and gene content and order suggests exposure to similar evolutionary processes over time. The stasis of the Wigglesworthia plasmids, relative to gene content and order, is in contrast to the versatility reported for the Buchnera extrachromosomal elements (32, 33) and may be attributable to particularities of insect host species ecology. The retention of these genes and genome elements by both the WGM and WGB genomes suggests their importance in Wigglesworthia biology and the symbiosis within the tsetse fly host background.

Only a small set of genes occurs in only one of the Wigglesworthia genomes (i.e., unique genes). These genes are either absent or still identifiable as a pseudogene in the sister genome (Fig. 3B; see Table S2 in the supplemental material). The retention of these unique genes may provide insight into the functional adaptation and evolution of the endosymbionts following tsetse fly divergence. However, the genomes of the obligate symbionts investigated to date have undergone drastic size reduction and appear to continuously lose genes due to random reductive processes (2, 4). Thus, the suite of unique genes in each of the Wigglesworthia genomes may reflect remnants of these random processes rather than species-specific interactions. Despite this, the presence of constituents of genetic pathways, which are widely dispersed throughout the host chromosome, does argue for selection favoring the retention of these loci. Whether these genes encode functional pathways and how they factor into host biology and ecology remain to be examined.

Interestingly, although the WGB and WGM genomes retain similar numbers of unique genes (Fig. 3B), these genes span a variety of functional classes. When the unique genes were classified by functional relevance, categories such as information transfer, regulation, transport, and hypothetical were quantitatively comparable (see Table S2 in the supplemental material). In relation to DNA processing (information transfer), the two Wigglesworthia genomes demonstrated distinctions, particularly in light of recombination and repair. Exclusively encoded within WGB are uvrD, involved in nucleotide excision repair and methyl-directed mismatch repair, recF, the single-stranded-DNA-specific exonuclease necessary for many recombination events (34), yggF, a putative Holliday junction resolvase (35), and the nucleotide exchange factor, greE (36). Meanwhile, mutY, which is involved in the correction of error-prone DNA synthesis due to oxidative stress (37), is present in WGM. Whether these differences in DNA repair and recombination reflect particular advantages in different host environments is unknown. It is possible that the retention of a suite of recombination-related genes by the WGB genome, including recA found in Wigglesworthia and Blattabacterium spp., may have contributed to the chromosomal inversions noted in both species. The absence of the recA gene, in particular, in many ancient endosymbiont genomes has been suggested to contribute to the chromosomal stability noted by the absolute conservation of genome colinearity (20). Genetic loci associated with the stabilization and maturation of ribosomal subunits, b2511 and rimM, were also differentially retained within the WGB and WGM genomes, respectively.

In addition, some of the unique genes retained in each genome (surA, ygcS, fisL, bacA, brrNQ, and b2817 in WGB and zmaA and yfgd in WGM) encode cell surface-associated proteins. Symbiotic surface proteins have been shown to be pivotal in the homeostasis of host-microbe relations, suggesting a possible role for these proteins in host species adaptation processes (38, 39). Unlike Buchnera, which is enclosed in host-derived vacuoles in bacteriocytes, Wigglesworthia lies free within the host cell cytosol in the bacteriome organ and has an extracellular stage in the accessory glands. Thus, cell surface proteins may be particularly relevant in host-symbiont interactions for Wigglesworthia symbiosis. In support of their divergence, signatures of Darwinian positive selection mean of 0.65 ± 0.04 (standard error) were noted in several Wigglesworthia membrane-associated proteins (encoded by ppdE, yraP, ompF, yfIO, tolA, and imp).

Some aspects of the WGM genome suggest that WGM can perform novel functions compared to WGB (see Table S2 in the supplemental material). The WGM unique gene set includes poxA and yjek, which are functionally coordinated in the posttranslational modification of elongation factor P (EF-P), which is involved in protein synthesis (40). In a recent survey where 725 bacterial genomes were analyzed, all possessed an efp gene but only 28% possessed both the poxA and yjek genes. In other organisms, including WGB, EF-P may be modified by another pathway or the translation machinery may have been adapted to cope with the lack of EF-P modification (40). Analysis of the WGM specific gene set also reveals distinct metabolic capabilities such as the presence of a complete shikimate biosynthetic pathway. Choriomate is required for the synthesis of all aromatic amino acids, as well as other vitamins and cofactors (41). Unlike E. coli and Salmonella enterica Typhi, which utilize a type I 3-dehydroquinate dehydratase (i.e., aruD) as the third enzymatic reaction in the shikimate pathway, WGB encodes a type II 3-dehydroquinate dehydratase (aroQ) that is shorter but orthologous to genes found in Helicobacter pylori, Yersinia pestis, and Mycobacterium tuberculosis and in other symbiont genomes such as those of Buchnera and Blochmannia. Interestingly, these type II 3-dehydroquinase dehydratases are homologous to fungal catabolic 3-dehydroquinases (42). Given that WGB is associated with the most ancestral tsetse fly species (G. brevipalpis), it is likely that unique genes in WGM were lost in WGB following the divergence of the WGM host lineage, likely due to random gene loss. Alternatively, unique genes in WGM may have been acquired by lateral transfer following host speciation, but such events are thought to be negligible in the evolution of obligate endosymbionts due to their intracellular localization and reduced recombination rates (2). Thus, the origin of the shikimate biosynthetic pathway requires further investiga-
tion into its absence or presence within different Wigglesworthia species.

Downstream of the chorismate pathway, WGM also contains genes involved in folate and phenylalanine biosynthesis. Intriguingly, African trypanosomes, i.e., *Trypanosoma brucei brucei* (43), are unable to synthesize phenylalanine and folate yet encode transporters to salvage both from the host environment. Whether these genomic differences between WGM and WGB contribute to variation in chorismate, phenylalanine, and folate biosynthetic capabilities and are involved in the higher vector competency (44, 45) of *G. morsitans* warrants further investigation.

Both WGB and WGM are genetically capable of flagellar synthesis, with associated genes demonstrating dN and dS values approximately equivalent to those of the remainder of the genome (dN and dS average of 0.28 ± 0.04 compared to a genome-wide average of 0.2617 ± 0.005; see Table S3 in the supplemental material), suggesting the potential for selection to be acting to preserve genes of biological importance. Here we demonstrate that fliC and motA, which are associated with structural and motility functions, respectively, are specifically expressed at particular host life stages, notably, during the maternal transmission process and functions, respectively, are specifically expressed at particular host life stages. In contrast, the chaperonin expressed in the midgut there is excess heme acquired through the blood diet, while Wigglesworthia-synthesized heme may help provision iron during intrauterine and pupal developmental stages. In contrast, the chaperonin encoded by groEL was expressed more consistently throughout all of the host stages examined, presumably due to its required assistance in protein folding. Our ongoing experiments where global gene expression is being investigated from different developmental stages will shed further light on symbiont functional biology and transcriptional regulation, as well as host–symbiont dialogue.

Comparison of the WGM and WGB genomes indicates high levels of synteny and functional conservation. Despite this, similarity and variation in genome composition between WGB and WGM have allowed us to make and test specific hypotheses regarding the functional biology of *Wigglesworthia* (e., flagellar expression, nutritional supplementation) and form the basis of future experimental analyses. For example, a high dN in the genes of obligate symbionts may be indicative of genome degradation or diversifying selection. Examination of additional *Wigglesworthia* genomes will now shed light on these processes. Our expression studies with the thiC, hemH, groEL, fliC, and motA genes indicate significant levels of transcriptional regulation and development- and tissue-specific functional roles for the symbiosis previously not observed in other obligate symbionts. Genome-wide analyses of gene expression in different host developmental stages and tissues are needed to better understand host-symbiont cross talk. In addition to tsetse fly host nutrient provisioning, the presence of *Wigglesworthia* during larval development has been associated with host immune maturation. Based on comparative genome analysis, we speculate another possible role for *Wigglesworthia* symbiosis where infections with pathogenic trypanosomes may depend upon symbiont species-specific metabolic products and thus influence the vector competence traits of different tsetse fly host species.

**MATERIALS AND METHODS**

**Insects and DNA preparation.** Genomic DNA from *Wigglesworthia sp.* (WGM) harbored by the tsetse fly *G. m. morsitans* was prepared. The *G. m. morsitans* colony maintained in the insectary at Yale University was originally established from puparia originating from fly populations in Zimbabwe. Flies are maintained at 24°C and 50 to 55% relative humidity. Flies are maintained at 24°C and 50 to 55% relative humidity and received defibrinated bovine blood every 48 h by an artificial membrane system (51). The bacteriome organs were isolated from about 1,000 adult females by dissection, bacteriocytes were released by gentle homogenization of the tissue, and DNA was isolated as previously described (16).

**Sequencing methodology.** The genome sequence of WGM was determined by the whole-genome shotgun strategy using Sanger sequencing. Genomic DNA was amplified by multiple-displacement amplification using a REPLI-g Midi kit (Qiagen) to obtain a sufficient amount of DNA for sequencing. The amplified genomic DNA was sheared using a HydroShear (Gene Machine). DNA fragments were fractionated by agarose gel electrophoresis and subcloned into vector plasmid pTS1 (Nippon Gene) to construct a shotgun library with an average insert size of 3 kb for sequencing using a 3730xl sequencer (Applied Biosystems). Template DNA was prepared by PCR with Ex-Taq (Takara Bio) on an aliquot of the bacterial culture to amplify the insert DNA of each clone. We produced 9,984 reads by sequencing both ends of the clones, giving 9.4-fold coverage. The assembly generated 14 contigs. Gap closing and resequencing of low-quality regions in the assembled data were performed by PCR, primer walking, and direct sequencing of appropriate plasmid clones. The overall accuracy of the finished sequence was estimated to have an error rate of less than 1/10,000 bases (Phrap score of ≥40).

**Genome annotation and alignment.** The rapid annotation using subsystems technology (RAST) server (52) was used for automated gene prediction and annotation of the WGM genome sequence. Predictions of ortholog between WGM and *E. coli* K-12 strain MG1655 were performed using a BLASTP reciprocal best-hit analysis with a threshold cutoff of 30% amino acid identity and requiring at least 60% of both proteins in the alignment. Because the *E. coli* MG1655 genome has been manually cu-
rated, resulting in high-quality and more-comprehensive gene annotations than automated processes can generate, the product names and gene names were transferred to orthologous genes from WGM. These annotations, alongside the ones from RAST, are available through the ASAP database at http://asap.ahabs.wisc.edu/asap/home.php (53). Genome sequences of WGM and WGB were aligned using Mauve with the match seed weight parameter increased to 21, allowing for a more accurate alignment of AC-rich genomes (54, 55), and orthologous genes were extracted using the export ortholog function. The WGM genome sequence was circularly permuted based on the Mauve alignment to the corresponding start site from WGB.

Manual annotation. For this analysis, no ORF smaller than 50 codons was considered a gene. Each WGM and WGB CDS predicted to be unique based on Mauve alignment was manually reanalyzed based on the results of BLAST (56) and FASTA (57) sequence comparisons using the nonredundant database at NCBI. To determine if lineage-specific orthologs were present, nucleotide sequences of unique CDSs, flanking orthologs, and the intervening sequences from both WGB and WGM were manually examined by MacClade 4.08. Unique CDS nucleotides were then aligned with the intervening sequence and translated to inspect the amino acid alignment. All unique CDSs, relative to either the WGM or the WGB genome, have been classified into one of the class identifiers based on hierarchical cellular functions of MultiFun (58) available in the ECOCYC database (http://www.ecocyc.org). Metabolic pathways were reconstructed using the reference pathways available for E. coli at EcoCyc and Kegg (59). CDSs proposed to be absent from WGM were similarly manually verified using WGB gene sequences. Manual nucleotide and amino acid sequence alignments were performed in MacClade 4.08. Sequences were determined to be orthologous (have shared ancestry) if nucleotide and amino acid sequences were similar and if start and stop codons were present in approximately the same position within the alignment.

Pseudogene annotation. Final manual inspection identified adjacent ORFs representing fragments of the same gene and truncated ORFs; therefore, CDSs less than half the length of their functional homologs in related species were categorized as pseudogenes. All of the pseudogenes identified in WGM and their functional classes are shown in Table S1 in the supplemental material.

Plasmid annotation. Identification of loci orthologous to the WGB plasmid was performed with BLAST at NCBI. A graphic display of the WGM plasmid map was generated using PlasMapper (60).

Molecular evolution. Orthologous CDSs were retrieved from WGM and WGB using alignment coordinates from Mauve. Manually annotated orthologous sequences were extracted by hand. Start and stop codons were removed from each pair of orthologs, and nucleotide sequences were then translated into amino acid sequences in MatLab. Using the Needleman-Wunsch algorithm (61), amino acid sequences were aligned with a gap penalty opening of 12 and a gap extension penalty of 4. The following calculations were made for each ortholog pair using the Bioinformatics Toolbox in MatLab: p distance, GC content (for each gene), dN/dS ratio, and maximum-likelihood-corrected dN/dS ratio. Once aligned by codons, sequences were converted back to nucleotides and the proportion of nucleotide differences (p distance) were calculated using the seqpdist function in MatLab. Alignments of sequences with large p distances (many nucleotide comparisons differed) were manually verified in MacClade.

To examine evidence of selection on genes or regions of the genome, the ratio of the number of nonsynonymous substitutions to the number of synonymous substitutions (dN/dS ratio) was calculated using the seqpdist, dnds, and dndsm1 functions in MatLab. The dndsm1 function incorporates a model of sequence evolution to minimally account for multiple hits in sequences. Typically, in comparisons of dN/dS ratios, low values suggest that genes are under purifying selection while high values are indicative of positive selection. This study is limited by the comparisons of genes between only two genomes, so evidence of selection does not explicitly incorporate the organism’s evolutionary history. Therefore, we instead examined the dN/dS ratio relative to the entire genome of each of the two organisms. Genes that were found to have a dN/dS ratio significantly higher or lower than the mean of the genome comparisons were identified as those that differed from the mean by 2 to 3 standard deviations. Symbiotic genomes commonly are subjected to deletions and accumulation of nonsynonymous mutations that eventually lead to gene loss (5). As a result, examination of the dN/dS ratios of genes that have substantial deletions and are potentially no longer functional may also accumulate large numbers of nonsynonymous mutations and therefore generate false positives in our survey of dN/dS ratios. Therefore, genes that differed between the two genomes by >100 bp were excluded from final dN/dS ratio presentation (see Table S3 in the supplemental material).

FlC immunostaining and microscopy. Antibodies were generated against E. coli His-tagged recombinant Flc protein. Primers were designed to amplify the coding region from bp 591 to 1001 of the WGB flc gene. Primer design included restriction sites to facilitate directional, in-frame cloning into the pET-28a 6×His tag expression vector (Novagen, Madison, WI) (primer sequences: WgbFlc forward, 5′-AGC ATGAGCTCGAATTGATAAAGGACCA; WgbFlc reverse, 5′-AG CATCTGAGGATCTTGGTAAATATTTGAA). The pET-28a—WgbFlc constructs were transformed into E. coli BL21, and recombinant protein expression was induced by treatment of cultures with 100 μM isopropyl-1-β-D-thiogalactopyranoside. Bacteria were lysed by sonication, and products were analyzed by SDS-PAGE. RecFlc was found predominantly in the insoluble fraction as inclusion bodies. Inclusion bodies were solubilized in binding buffer in the presence of 6 M urea and purified by using nickel resin under a denaturing conditions protocol (Novagen His- Bind kit). RecFlc proteins were subsequently purified by SDS-PAGE, and gel slices were provided for commercial antisera production (Cocalico Biologicals).

For immunohistochemistry, tissues were dissected and fixed for about 1 week in 4% paraformaldehyde. Samples were then dehydrated, embedded in paraffin, cut into 5-μm thick sections, and mounted on poly-L-lysine-coated microscopy slides. After being dewaxed for 2 × 15 min in methylocyclohexane and 2 × 10 min in ethanol, samples were air dried and rehydrated in 1× phosphate buffer saline containing 0.01% Tween 20 (PBST). After 1 h of blocking in 3% bovine serum albumin in 1× PBST at room temperature, sections were incubated in WGB Flc antibody solution (1:500 in 1× PBST) overnight at 4°C. Following three 10-min washes in PBST, slides were incubated in anti-rabbit Alexa 488 antibody (Molecular Probes; diluted 1:500 in PBST) for 1 h at room temperature in the dark. Sections were then washed again three times for 10 min each time in PBST, rinsed in water, and air dried in the dark. They were then mounted in GelMound mounting medium, which contained 4′,6-diamidino-2-phenylindole (DAPI) and covered with coverslips. Microscopic analyses were conducted using a Zeiss Axioskop 2 microscope equipped with an Infinity 1 USB 2.0 camera and software (Lumenera Corporation). Fluorescent images were taken using a fluorescence filter set with fluorescein- and DAPI-specific channels.

Real-time qRT-PCR gene expression analyses. For Wigglesworthia gene expression, gene-specific primers were used to quantify hemH, motA, groEL, flc, and thiC transcripts. The rpsC gene was used for normalization. qRT-PCR was performed with an iCycler iQ real-time PCR detection system (Bio-Rad, Hercules, CA) using the primer sets and conditions described in Table S1 in the supplemental material. The normality of sample means from each treatment was determined by Shapiro-Wilk test prior to t test analysis. Values are represented as the mean ± the standard error of the mean, and statistical significance was determined using a Student’s t test and Microsoft Excel software.

Nucleotide sequence accession number. The sequence data obtained in this study have been deposited in GenBank under project accession no. CP003315.
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SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at http://mbio.asm.org

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