



Odorant and Gustatory Receptors in the Tsetse Fly *Glossina morsitans morsitans*

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Abstract

Tsetse flies use olfactory and gustatory responses, through odorant and gustatory receptors (ORs and GRs), to interact with their environment. *Glossina morsitans morsitans* genome ORs and GRs were annotated using homologs of these genes in *Drosophila melanogaster* and an *ab initio* approach based on OR and GR specific motifs in *G. m. morsitans* gene models coupled to gene ontology (GO). Phylogenetic relationships among the ORs or GRs and the homologs were determined using Maximum Likelihood estimates. Relative expression levels among the *G. m. morsitans* ORs or GRs were established using RNA-seq data derived from adult female fly. Overall, 46 and 14 putative *G. m. morsitans* ORs and GRs respectively were recovered. These were reduced by 12 and 59 ORs and GRs respectively compared to *D. melanogaster*. Six of the ORs were homologous to a single *D. melanogaster* OR (DmOr67d) associated with mating deterrence in females. Sweet taste GRs, present in all the other Diptera, were not recovered in *G. m. morsitans*. The GRs associated with detection of CO₂ were conserved in *G. m. morsitans* relative to *D. melanogaster*. RNA-sequence data analysis revealed expression of GmmOR15 locus represented over 90% of expression profiles for the ORs. The *G. m. morsitans* ORs or GRs were phylogenetically closer to those in *D. melanogaster* than to other insects assessed. We found the chemoreceptor repertoire in *G. m. morsitans* smaller than other Diptera, and we postulate that this may be related to the restricted diet of blood-meal for both sexes of tsetse flies. However, the clade of some specific receptors has been expanded, indicative of their potential importance in chemoreception in the tsetse.

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Introduction

Trypanosomiasis management has been a longstanding development preoccupation in sub-Saharan Africa, with tsetse fly control constituting the cornerstone in this effort [1]. Since all tsetse species are able to transmit trypanosomes, the critical determinant of transmission is their obligate blood feeding. Tsetse flies select their hosts through visual and olfactory signals, a process that is mediated by olfactory and gustatory receptors. Tsetse flies navigate their environment by detecting and responding to volatiles and non-volatile cues (odors and tastants). Artificial bait technologies, based on tsetse olfactory responses to natural cues and blends of synthetic versions that mimic those of their natural hosts in the field, have successfully been applied in tsetse control because of their relatively high specificity, low cost, community acceptability, and ability to slow down tsetse re-invasion from adjacent areas [2,3]. These technologies are environment friendly [4], and applicable for riverine and savanna species of tsetse flies [5,6]. The attractants include various phenolic derivatives [7–9], carbon dioxide, acetone, 1-octen-3-ol, and

vertebrate host breath, skin and urine extracts [10–12]. Interestingly, 1-octen-3-ol is a constituent of the chemical profile from *Lantana camara*, an invasive plant to which tsetse flies are attracted [13]. The response to olfactory cues has also been exploited in design of tsetse repellents [14,15]. The repellents include guaiacol (methylphenols), δ -octalactone and methylketones [16–18] and 2-methoxy-4-methylphenol [14]. Natural differential responses among tsetse species and even between sexes and allopatric populations of the same species have been observed [18–22], which have stimulated research and design to enhance the efficiencies of the existing attractant-based bait technologies, to develop new ones based on repellent blends ('push' tactics) from refractory animals, and to integrate these into 'push-pull' strategies. Different *Glossina* species exhibit different olfactory uniqueness' and this may partly account for the observed gradation of preferences for particular hosts. For instance, riverine tsetse species (such as *G. fuscipes fuscipes*, *G. palpalis* and *G. tachinoides*) prefer feeding on reptilian hosts compared to their savanna relatives (*G. morsitans morsitans*, *G. pallidipes*) that feed largely on ungulates and other large mammals [6]. Larviposition

Author Summary

Tsetse flies navigate their environments using chemosensory receptors, which permit them to locate hosts, mating partners, and resting and larviposition sites. The genome of *G. m. morsitans* was interrogated for coding genes of odorant receptors (ORs) and gustatory receptors (GRs) that express in antennae and maxillary palp, and detect the volatile and soluble chemical signals. The signals are then transmitted to the central nervous system and translated to phenotypes. Majority of these genes in *G. m. morsitans* were spread across different scaffolds, but a few were found to occur in clusters, which suggested possible co-regulation of their expression. The number of ORs and GRs were much reduced in the *G. m. morsitans* genome, including the apparent loss of receptors for sugar when compared to selected Diptera. There was also an apparent numerical expansion of some receptors, presumably to maximize on their restricted blood-meal diet. The annotation of the chemoreceptor package of *G. m. morsitans* provides a resource for investigating key activities of tsetse flies that could be exploited for their control.

pheromones (n-pentadecane and n-dodecane) from exudates of mature larvae are also known to attract and induce gravid *G. m. morsitans* and *G. m. centralis* females to aggregate and deposit larvae [23]. Research on response to tastants in tsetse flies are limited, but point to their potential application in tsetse control [10,24]. In all, responses to odors and tastants in tsetse have established utility in tsetse control that can be augmented with better understanding of the molecular factors that underpin these responses.

Molecular factors mediating the olfactory and gustatory responses in the tsetse flies are poorly understood. However, research on other insects indicates that the odors and tastants in the environment are generally detected in peripheral sensory neurons by distinct odorant and gustatory receptors (ORs and GRs) [25–28]. These receptors are divergent members of a superfamily characterized by seven transmembrane domains, and share low sequence conservation among them except at the C-terminus region that coincides with the seventh trans-membrane domain [29]. The ORs and GRs are thought to have evolved as parallel chemoreceptors across diverse organisms [26]. Each OR is expressed in olfactory receptor neurons (ORNs) within maxillary palpi and antennae [25,30–32]. The ORs generally have multiple introns and are very divergent with poor structural conservation within and across insect orders and species [33,34], which potentially reflect diverse olfaction related preferences within the orders and species. However, a canonical co-receptor commonly referred to as Orco remains highly conserved across insect orders [35–38], a phenomenon that may be associated with its role in proper tuning of odor specificity and activation necessary for appropriate signal transduction in the neurons [39]. The GRs are generally expressed in gustatory receptor neurons (GRNs) within gustatory organs [40] in response to soluble taste and contact pheromones [41,42]. However, some GRs are expressed in antennal dendrites and respond to carbon dioxide, potentially implicating them in olfaction [40,43]. The GRs are more conserved in sequence and structure than the ORs [44,45] probably due to comparatively smaller search space among cues associated with GRs than ORs. The diversity among the ORs and GRs in tsetse can potentially shed light on the natural differential responses observed among them [12,17,18,20–29], with potential application in tsetse control. To improve or develop new approaches of vector management, an understanding of the

molecular attributes of GRs and ORs and their potential roles in tsetse ecology is essential.

This study was initiated to (1) comparatively annotate and catalogue ORs and GRs in *G. m. morsitans* (GMOY1.1), (2) establish evolutionary distance between *G. m. morsitans* ORs or GRs and those in especially *D. melanogaster*, and (3) examine relative expression of the ORs and GRs in the *G. m. morsitans*. The assembly has been estimated to be over 99% complete based on the software Core Eukaryotic Genes Mapping Approach (CEGMA) [46] and manually sequenced BACs data. The assembly is currently undergoing genome-wide manual curation and annotation by the International Glossina Genome Initiative (IGGI) consortium.

Methods

Retrieval and annotation of *G. m. morsitans* OR and GR gene models

Coding sequences (CDS) of ORs and GRs in *Drosophila melanogaster* were obtained from FlyBase5.13 [47] and used to isolate their respective homologs in the *G. m. morsitans* genome (GMOY1.1) at VectorBase [48] using tBLASTx algorithm [49]. Scaffolds encoding the homologs were searched for and retrieved at a cut-off e-value $<1.0e-05$. Whole transcriptome illumina 84 million RNA sequence reads generated from female *G. m. morsitans* [50] were mapped onto the scaffolds using default settings in CLC Genomics workbench suite Version 4.8 (CLC Bio, Aarhus, Denmark). Gene loci of putative *Glossina* homologs were curated in the scaffold sequences flanking the tBLASTx hits, and intron/exons modeled using the RNA-seq mappings. The predicted gene models were viewed and edited using Artemis v13.2.12 [51] where, intron/exon boundaries were edited using motifs GT for 5' donor site, and AG for 3' acceptor site. The start codon (ATG) for each gene model was fixed at the 5' end and the reading frame terminated at the first of any of the stop codons (TAA, TGA, or TAG). Sequences shorter than average size of known insect ORs (370 aa) were marked as incomplete if they lacked start or stop codons. Sequences with poorly conserved functional domains were considered as pseudogenes.

The homologs were validated through sequence-based searches for presence of ORs or GRs specific 7tm-6-olf-recept or 7tm-7-olf-recept [29,52] domains respectively. The homologs were probed for the domains using DELTA BLAST algorithm [53] against the conserved domains databases (CDD) [54], and presence of alpha helix trans-membrane domains validated using TMHMM server v2.0 [55]. Additionally, all the putative ORs or GRs were validated, using BLAST2GO analyses [56] against the non-redundant Swiss-Prot database [57]. The curated gene models were assigned annotation identifiers by comparing them with automated transcript feature models obtained from the Glossina community annotation portal at VectorBase [48] and edited using Artemis genome viewer tool [51]. The models without automated prediction matches and identifiers were manually built using the Artemis gene build tool window [51] and given unique temporary annotation identifiers. In this respect, features for gene, exons, mRNA, and CDS were created for such gene models. The Glossina gene models were assigned putative gene names where GmmOR* and GmmGR* were adopted for *G. m. morsitans* odorant receptors and gustatory receptors respectively (the asterisk (*) being an identifier number). The annotated gene model features were submitted to the VectorBase community annotation portal for *G. m. morsitans* [48] for integration into genome database; nevertheless, a list of annotated amino acid coding sequences is presented in supplementary Dataset S1, and a list of associated

Table 1. Annotated ORs and GRs in *G. m. morsitans* and other selected insect species.

| Insect | ORs | GRs | Reference |
|------------------------------|---------------|-----------|--------------------|
| <i>D. melanogaster</i> | 60 (2)* | 60 (13)* | [25–27,29,42] |
| <i>G. m. morsitans</i> | 46 (3) | 14 | This study. |
| <i>An. gambiae</i> | 79 | 76 | [52,76] |
| <i>Ae. aegypti</i> | 100(31) | 79 | [74,77] |
| <i>Apis mellifera</i> | 163 (11) | 10 (3) | [41] |
| <i>Nasonia vitripennis</i> | 225 (76) | 47 (11) | [80] |
| <i>Camponotus floridanus</i> | 352 (55) | 46 (17) | [75] |
| <i>Harpegnathos saltator</i> | 347 (30) | 17 (4) | [75] |
| <i>Tribolium castaneum</i> | 265 (76) | 220 (25) | [78,79] |

Figures in parentheses are numbers incomplete genes and or pseudogenes of the receptors.

*- in parentheses are alternatively spliced forms.

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gene identities in Table S2. The *G. m. morsitans* receptor repertoires were evaluated against those documented for *D. melanogaster*, *Anopheles gambiae*, *Aedes aegypti*, *Apis mellifera*, *Nasonia vitripennis*, *Camponotus floridanus*, *Harpegnathos saltator* and *Tribolium castaneum* (references in Table 1).

Phylogenetic analyses of ORs and GRs in *G. m. morsitans* and selected Diptera

Multiple Sequence Comparison by Log-Expectation (MUSCLE) tool [58] was used to align GmmORs and GmmGRs with homologs in *D. melanogaster*, and the alignments edited using Jalview web-server [59]. The secondary structures in the alignments were predicted using JPred program [60]. Phylogenetic cluster inference was done using Maximum Likelihood approach with best fitting Wheelan and Goldman+Freq (WAG+F) model [61], which was chosen as the best ranked from a panel of all amino acid model tests run in MEGA5 [62]. The initial tree was automatically generated and bootstrapped with 500 iterations. The evolutionary rate difference among sites was modeled using a discrete Gamma distribution (5 categories (+G, parameter = 4.2651)). The rate variation model allowed for some sites to be evolutionarily invariable ([+I], 0.8705% sites). All positions with less than 95% site coverage were eliminated and branch nodes determination set at very strong. Evolutionary analyses were conducted using the MEGA5 suite [62].

Comparative analyses of expression profiles of *G. m. morsitans* ORs and GRs

The expression profiles of *G. m. morsitans* ORs and GRs gene loci were determined using whole transcriptome 84 million illumina RNA-sequence reads [50]. The RNA-seq reads were mapped onto the *G. m. morsitans* ORs or GRs nucleotide coding sequences (CDS) in CLC Genomics Workbench (CLC Bio, Aarhus, Denmark) via RNA-seq analysis pipeline with default settings. The expression profiles were presented as reads per kilobase of exon model per million mapped reads (RPKM) for each receptor sequence [63].

Results

Most of the gene loci of *G. m. morsitans* ORs and GRs were scattered amongst the scaffolds. Fifty percent of *G. m. morsitans*

OR genes were encoded as single-copies on their respective scaffolds. The remainder were encoded in pairs or triplets per scaffold. Five *G. m. morsitans* OR loci (GmmOR6/7/8, GmmOR18/19, GmmOR22/25, GmmOR27/28 and GmmOR41/42) were located in tandem on their respective scaffolds. Similarly, five *G. m. morsitans* GR genes clustered on a single scaffold. The rest were encoded as single-copies on their respective scaffolds. All *G. m. morsitans* GR loci were annotated as complete genes.

Gene models for *G. m. morsitans* OR and GR and their annotation

Numbers of OR and GR gene loci recovered in *G. m. morsitans*, relative to those published in other insects are summarized in Table 1. Similar to most insects, the *G. m. morsitans* has more ORs loci than GRs loci, with the exception of *D. melanogaster* where the numbers are equal. However, the *G. m. morsitans* ORs are fewer than those documented in all the insects evaluated, including *D. melanogaster*. A similar trend was exhibited in *G. m. morsitans* GRs, except in relation to *A. mellifera*. Annotation of *G. m. morsitans* ORs and GRs are summarized in Table 2. The lengths of *G. m. morsitans* ORs varied between 260 and 541 amino acids, while those of *G. m. morsitans* GRs ranged from 309 to 514 amino acids. The number of exons ranged between two and eight or 12 in GRs and ORs respectively. The predicted genome structures are given in Figure S1. The frequency of detectable trans-membrane domains was also variable, with proteins having six trans-membrane domains representing about one half of all genes. The *G. m. morsitans* ORs (57%, 26 out of 46) were homologous to nine *D. melanogaster* ORs. Similarly, most of the *G. m. morsitans* GRs (57%, 8 out of 14) were homologous to three *D. melanogaster* GRs genes. The remainder of the *G. m. morsitans* GRs had one-to-one homology with a single *D. melanogaster* specific homolog. Reciprocal blasts onto non-redundant protein databases for both *G. m. morsitans* ORs and GRs are summarized in Supplementary material – Table S1). GmmGR3 and GmmGR4 were also homologous to *An. gambiae* orthologs, while GmmGR5, GmmGR8 and GmmGR13 had homologs to genes in other *Drosophila* species. The *G. m. morsitans* ORs pseudogenes were scanty, representing 7% of the ORs genes recovered. Only GmmOR5 had alternative splice variants. The 7tm-6-olfct-rcpt domain was detected in all *G. m. morsitans* ORs, and the 7tm-7-chem-rcpt domain was detected in five ORs (GmmOR17, GmmOR21, GmmOR24, GmmOR38 and GmmOR39). The 7tm-7-chem-rcpt domain was also detected in all the *G. m. morsitans* GRs.

Phylogenetic analysis of *G. m. morsitans* ORs and GRs with other insects

Phylogenetic relationships between *G. m. morsitans* ORs and GRs and their counterparts in *D. melanogaster* are summarized in Figure 1. Most of the *G. m. morsitans* ORs and GRs clustered with their respective ORs and GRs orthologs with a bootstrap support of over 80%. The *G. m. morsitans* OR14, OR15 and OR16 were homologous to a drosophila larvae receptor, Or45a. The *G. m. morsitans* co-receptor (Orco) (GmmOR1) had 100% bootstrap support homology to *D. melanogaster* homolog, Or63b, and was a single copy in the genome, similar to other insects investigated (data not shown). There was an expanded cluster of ORs in *G. m. morsitans* (GmmOR41-46), relative to a single *D. melanogaster* homolog, Or67d (Figure 1A), which also had multiple copies in *An. gambiae*, *Cu. quinquefasciatus*, *Ae. aegypti*, *Tribolium castaneum* (Data not shown). The *G. m. morsitans* and *D. melanogaster* GRs clustered into four groups (Figure 1B). Four *G. m. morsitans* GRs (GmmGR1-

Table 2. Annotations of odorant and gustatory receptor genes in *G. m. morsitans* and their homologs in *D. melanogaster*.

| <i>G. m. morsitans</i> Genes | Length (AA) | Exons | TMMs | Gene ID | <i>Dmel</i> orthologs/Accession Number |
|------------------------------|-------------|-------|------|------------|--|
| GmmOR1 | 521 | 8 | 6 | GMOY005610 | DmOr83b/CG10609 |
| GmmOR2 | 394 | 3 | 7 | GMOY005796 | DmOr2a/CG3206 |
| GmmOR3 | 387 | 3 | 3 | GMOY004772 | DmOr19a/CG18859 |
| GmmOR4 | 384 | 2 | 7 | TMP_Or4 | DmOr59a/CG9820 |
| GmmOR5* | 442 | 4 | 5 | GMOY012018 | DmOr33b/CG16961 |
| GmmOR6 | 387 | 4 | 5 | GMOY009475 | DmOr42b/CG12754 |
| GmmOR7 | 406 | 3 | 6 | TMP_Or7 | DmOr42b/CG12754 |
| GmmOR8 | 389 | 4 | 6 | TMP_Or8 | DmOr42b/CG12754 |
| GmmOR9 | 409 | 3 | 6 | TMP_Or9 | DmOr42b/CG12754 |
| GmmOR10 | 444 | 3 | 6 | TMP_Or10 | DmOr46a/CG33478 |
| GmmOR11 | 341 | 3 | 6 | GMOY010761 | DmOr46a/CG33478 |
| GmmOR12 | 340 | 3 | 3 | GMOY009271 | DmOr94b/CG17241 |
| GmmOR13 | 391 | 6 | 6 | GMOY003312 | DmOr82a/CG31519 |
| GmmOR14 | 341 | 3 | 6 | GMOY001365 | DmOr45a/CG1978 |
| GmmOR15 | 446 | 4 | 7 | TMP_Or15 | DmOr45a/CG1978 |
| GmmOR16 | 387 | 4 | 6 | TMP_Or16 | DmOr45a/CG1978 |
| GmmOR17 | 541 | 12 | 8 | GMOY005386 | DmOr69a/CG33264 |
| GmmOR18 | 420 | 8 | 6 | TMP_Or18 | DmOr63a/CG9969 |
| GmmOR19 | 385 | 8 | 7 | GMOY012322 | DmOr63a/CG9969 |
| GmmOR20# | 269 | 7 | 6 | TMP_Or20 | DmOr85b/CG11735 |
| GmmOR21 | 465 | 5 | 2 | GMOY011399 | DmOr83a/CG10612 |
| GmmOR22# | 296 | 4 | 5 | TMP_Or22 | DmOr49a/CG13158 |
| GmmOR23 | 331 | 4 | 5 | TMP_Or23 | DmOr85b/CG11735 |
| GmmOR24 | 388 | 3 | 6 | GMOY010839 | DmOr85c/CG17911 |
| GmmOR25 | 385 | 3 | 6 | GMOY012357 | DmOr56a/CG12501 |
| GmmOR26 | 418 | 4 | 5 | TMP_Or26 | DmOr85b/CG11735 |
| GmmOR27 | 415 | 3 | 6 | GMOY008038 | DmOr67c/CG14156 |
| GmmOR28# | 260 | 2 | 7 | TMP_Or28 | DmOr92a/CG17916 |
| GmmOR29 | 438 | 3 | 4 | TMP_Or29 | DmOr67a/CG12526 |
| GmmOR30 | 361 | 3 | 6 | TMP_Or30 | DmOr67a/CG12526 |
| GmmOR31 | 435 | 7 | 5 | TMP_Or31 | DmOr24a/CG11767 |
| GmmOR32 | 450 | 5 | 7 | GMOY005084 | DmOr13a/CG12697 |
| GmmOR33 | 353 | 6 | 5 | GMOY005479 | DmOr49b/CG17584 |
| GmmOR34 | 360 | 7 | 4 | GMOY011902 | DmOr30a/CG13106 |
| GmmOR35 | 392 | 5 | 6 | TMP_Or35 | DmOr43a/CG1854 |
| GmmOR36 | 343 | 7 | 6 | TMP_Or36 | DmOr43a/CG1854 |
| GmmOR37 | 430 | 4 | 4 | TMP_Or37 | DmOr74a/CG13726 |
| GmmOR38 | 371 | 5 | 6 | TMP_Or38 | DmOr47b/CG13206 |
| GmmOR39 | 403 | 3 | 6 | GMOY004392 | DmOr88a/CG14360 |
| GmmOR40 | 284 | 5 | 6 | GMOY012356 | DmOr56a/CG12501 |
| GmmOR41 | 386 | 4 | 6 | GMOY006480 | DmOr67d/CG14157 |
| GmmOR42 | 386 | 4 | 5 | GMOY006479 | DmOr67d/CG14157 |
| GmmOR43 | 389 | 4 | 5 | TMP_Or43 | DmOr67d/CG14157 |
| GmmOR44 | 390 | 4 | 6 | GMOY006265 | DmOr67d/CG14157 |
| GmmOR45 | 385 | 4 | 7 | GMOY007896 | DmOr67d/CG14157 |
| GmmOR46 | 348 | 4 | 3 | GMOY003305 | DmOr67d/CG14157 |
| GmmGR1 | 425 | 3 | 6 | GMOY007472 | DmGr21a/CG13948 |
| GmmGR2 | 514 | 7 | 6 | GMOY011510 | DmGr22b/CG31931 |
| GmmGR3 | 425 | 6 | 6 | TMP_Gr5 | DmGr21a/CG13948 |

Table 2. Cont.

| <i>G. m. morsitans</i> Genes | Length (AA) | Exons | TMMs | Gene ID | <i>Dmel</i> orthologs/Accession Number |
|------------------------------|-------------|-------|------|------------|--|
| GmmGR4 | 496 | 8 | 6 | GMOY008001 | DmGr63a/CG14979 |
| GmmGR5 | 467 | 5 | 7 | GMOY004207 | DmGr66a/CG7189 |
| GmmGR6 | 443 | 4 | 8 | GMOY011615 | DmGr28b/CG13788 |
| GmmGR7 | 402 | 3 | 7 | GMOY006209 | DmGr28b/CG13788 |
| GmmGR8 | 407 | 2 | 6 | TMP_Gr4 | DmGr22e/CG31936 |
| GmmGR9 | 348 | 5 | 4 | GMOY011903 | DmGr2a/CG18531 |
| GmmGR10 | 458 | 4 | 7 | GMOY003231 | DmGr33a/CG17213 |
| GmmGR11 | 450 | 3 | 6 | TMP_Gr3 | DmGr22b/CG31931 |
| GmmGR12 | 375 | 2 | 8 | TMP_Gr2 | DmGr32a/CG14916 |
| GmmGR13 | 457 | 2 | 6 | TMP_Gr1 | DmGr22b/CG31931 |
| GmmGR14 | 309 | 3 | 6 | TMP_Gr6 | DmGr22b/CG31931 |

GmmOR – *Glossina morsitans morsitans* odorant receptor; GmmGR– *G. m. morsitans* gustatory receptor; TMM- Trans-membrane helices; GMOY – *Glossina morsitans* Yale strain; TMP_Or – Provisional odorant receptor ID; TMP_Gr – Provisional gustatory receptor ID; DmOr- *Drosophila melanogaster* odorant receptor; DmGr- *D. melanogaster* gustatory receptor;

*- longest alternative splice variant in locus OR5;
 #- pseudogene.

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4) clustered with homologs of CO₂ receptors, Gr21a and Gr63a in *D. melanogaster*; GmmGR6-7 and GmmGR14, though distantly, clustered with an unusual splice variant DmelGr28a/28b;

GmmGR5, 8–12 were homologous to bitter taste-related sensors in *D. melanogaster*; and GmmGR13 clustered distantly to Dmel-Gr58a/58b homologs, whose functions are unknown.

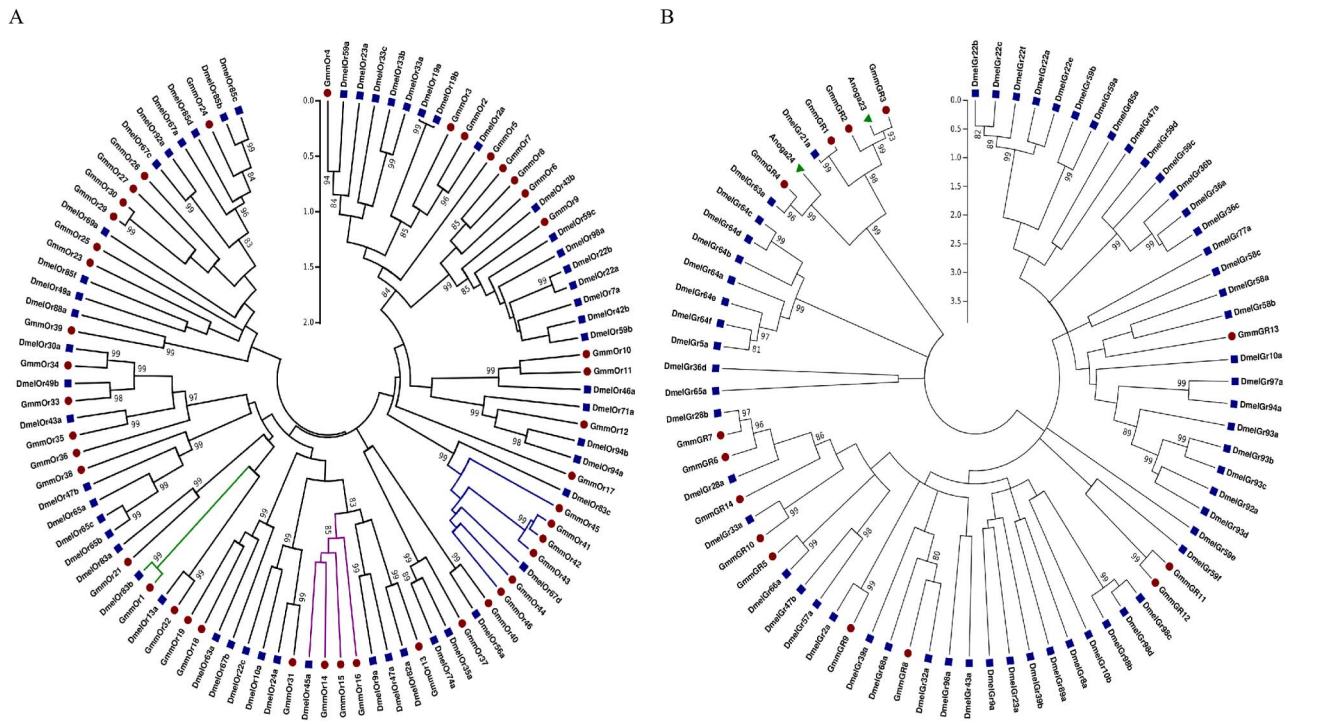
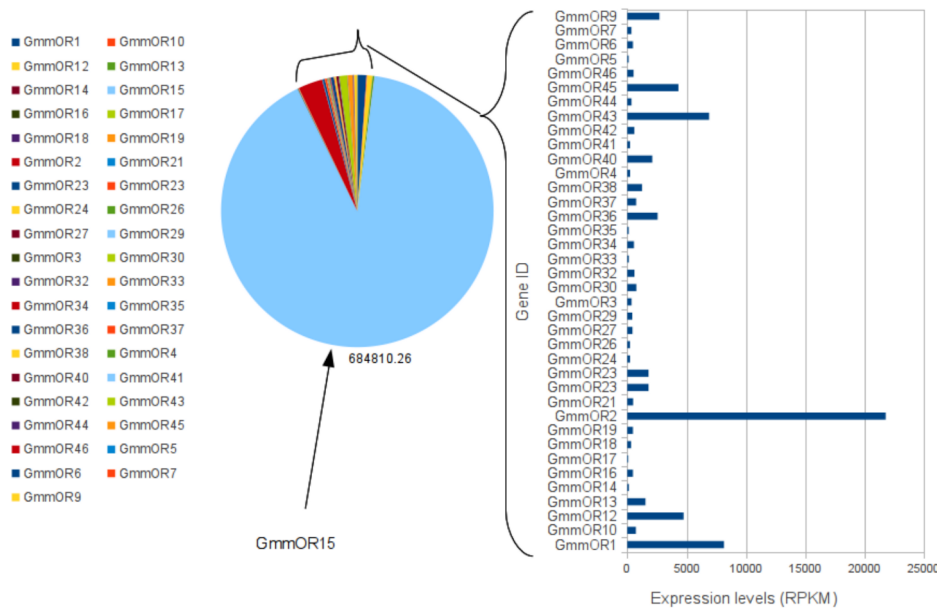


Figure 1. Phylogenetic analyses of ORs or GRs in *G. m. morsitans* and selected Diptera. (A) Maximum likelihood (ML) tree for GmmORs and DmelORs; branches annotated blue is an expanded clade orthologous to DmelOr67d; purple branches is the clade orthologous to DmelOr45a; and green branches indicate the orco cluster. (B) Maximum likelihood tree for GmmGRs and DmelGRs. In both trees, blue labels are *D. melanogaster* receptors and red labels *G. m. morsitans* receptors (green labels are *An. gambiae* CO₂ receptors). Phylogenetic cluster inferences were deduced using Maximum Likelihood approach with best fitting Wheelan And Goldman+Freq (WAG+F) model [59]. Evolutionary analyses were conducted using MEGA5 suite [60].

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A.



B.

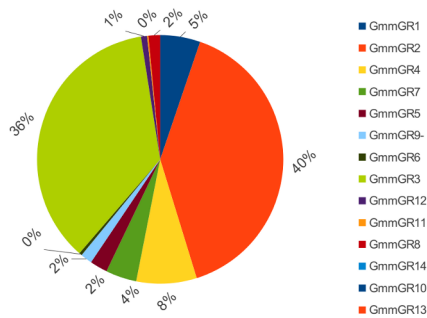


Figure 2. Glossina chemoreceptor expression abundances by RNA-seq data in RPKM. (A) Expression abundances of *GmmORs*. There was no sufficient data to support profiles for *GmmOR8*, 11, 25, 31, and 39. *GmmOR15* had abundant transcriptome data of 90.746% relative to sequence reads that mapped onto *GmmORs*. **(B)** Expression abundances of *GmmGRs*. Expression profiles for *GmmGR6*, *GmmGR11* and *GmmGR13* were not detected; *GmmGR2* and *GmmGR3* accounted for 40% and 36% respectively of the total transcripts considered for *GRs*. doi:10.1371/journal.pntd.0002663.g002

Relative expression profiles of *G. m. morsitans* ORs and GRs

Relative expression profiles of the *G. m. morsitans* ORs and GRs gene loci are summarized in Figure 2. Among the *G. m. morsitans* ORs, expression of *GmmOR15* was surprisingly most predominant, accounting for more than 90% of the total RNA-sequence data supporting expression of the ORs. *GmmOR15* is homologous to Or45a gene in *D. melanogaster*. About 5% of RNA-sequence data provided supporting evidence for expression of *GmmOR2*, *GmmOR1* (Orco homolog), *GmmOR43* and *GmmOR9*. Expressions of *GmmOR8*, *GmmOR11*, *GmmOR25*, *GmmOR31*, and *GmmOR39* were not detected in the available RNA-sequence dataset (Figure 2A). Amongst the GRs, *GmmGR1-4* had the best RNA-sequence data expression support (Figure 2B).

Discussion

Specific groups of the *G. m. morsitans* ORs and GRs were clustered within selected scaffolds. Similar clusters of genes

performing common and related functions have been observed among chemosensory genes in *D. melanogaster* [41,42,44], and more recently among twelve *G. m. morsitans* major milk proteins associated with lactation [50]. Since genes within clusters are generally co-regulated and can lead to joint gene expression [29,34,64], the individual clusters of ORs and GRs might be under common regulatory mechanisms and in response to common or related stimuli. The ORs and GRs in *G. m. morsitans* were fewer than those documented in most insects evaluated (Table 1) [65,66]. Additionally, specific ORs and GRs in *D. melanogaster* (nine and three ORs and GRs respectively) appear to have been expanded in *G. m. morsitans*, representing more than half of the chemoreceptors.

The factors underlying the apparent reductions and expansions of these receptors in the tsetse are unknown. However, it can be postulated that the obligate blood feeding of the tsetse fly (restricted to vertebrate hosts) relative to *D. melanogaster* (with expansive fruit species hosts) might have necessitated evolutionary selection for specific chemoreceptor loci relevant to discriminate

among limited host choices. We know also that environmental factors can determine host choice, as tsetse have been shown to have an acquired preference to specific hosts encountered early in life [67]. Notably, other blood-feeders, such as mosquitoes also seek a variety of plant sources for sugar as energy source, while tsetse flies derive their energy from the amino acids proline and alanine [68]. The *G. m. morsitans* OR15 (GmmOR15) accounted for more than 90% of the OR expression data. This OR is homologous to DmelOr45a, whose product has been, associated with an escape response in *D. melanogaster* larvae [69]. The function of this OR in tsetse was not determined; nonetheless it is notable that the source of RNA sequence data was a reproductively active adult female. Hence, it is possible that the GmmOR15 is in some way associated with larval activity.

Similarly, the GmmGR1-4 cluster was most prominent among the GRs homologous to CO₂ receptors in *D. melanogaster*. These GRs may be associated with host seeking and may have a duplicate role in olfaction. These receptors may putatively be associated with attractive responses elicited by the savanna tsetse species, including *G. m. morsitans* [10]. From the foregoing, it is evident that tsetse seems to prioritize and invest on a select few chemoreceptor genes towards their adaptive behaviors. Indeed, a heavy investment in specific genes is not uncommon in insects [70–73]. The *G. m. morsitans* OR1 (homologous to *Orco*) was the most conserved amongst the *G. m. morsitans* ORs, not surprising since such conservation has been observed in other insects [74] probably due to its critical role in modulating responses of the other receptors.

In conclusion, when examined against other blood feeders, which also take sugar sources from plants (e.g. *An. gambiae* and *Ae. aegypti*), the *G. m. morsitans* has a reduced repertoire of ORs and GRs genes. There is a complete loss of receptors for sugar, and a heavy investment in some chemoreceptors, such as those associated with detection of CO₂. These observations offer opportunities to develop control tools exploiting these unique adaptations.

Supporting Information

Dataset S1 Annotated amino acid sequences of *Glossina morsitans morsitans* ORs and GRs. Each sequence name is

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followed by annotation identity (previous temporary identity), scaffold number and the coordinates within the scaffold. (PDF)

Figure S1 *Glossina m. morsitans* ORs and GRs genome structure. Most of the gene loci were encoded as singlets on their respective scaffolds. Some loci were encoded in tandem in their respective scaffolds suggestive of possible joint expression regulation. All genes had multiple exons ranging from two to eight or 12 in GRs or ORs respectively. (PDF)

Table S1 Reciprocal blast of *G. m. morsitans* ORs and GRs to non-redundant protein database at NCBI. The names of annotated gene loci, scaffold identity, gene location within the scaffold, in bracket (*) refers to the coding strand where (–), reverse and (+), forward strands. The reciprocal blast gives the distribution of homology to other insects including *D. melanogaster*. (PDF)

Table S2 List of *Glossina m. morsitans* gene names and their associated identities. The vectorbase identities (GMOY....) has since replaced the *Glossina* community annotation identities (TMP0....) that were computationally derived. Both annotation identities and phylomedb identities can be used as seed queries to retrieve their related *Glossinia* phylomedb 182 trees (www.phylomedb.org/?q = user/28). (PDF)

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Author Contributions

Conceived and designed the experiments: GFOO POM AC DKM. Performed the experiments: GFOO. Analyzed the data: GFOO HMR SRGN. Contributed reagents/materials/analysis tools: POM AC DKM. Wrote the paper: GFOO POM AC SRGN HMR DKM. Designed *Glossina morsitans morsitans* gene name/symbol nomenclature: HMR.

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