

# Divergent and Conserved Elements Comprise the Chemoreceptive Repertoire of the Nonblood-Feeding Mosquito *Toxorhynchites amboinensis*

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Accepted: October 7, 2014

## Abstract

Many mosquito species serve as vectors of diseases such as malaria and yellow fever, wherein pathogen transmission is tightly associated with the reproductive requirement of taking vertebrate blood meals. *Toxorhynchites* is one of only three known mosquito genera that does not host-seek and initiates egg development in the absence of a blood-derived protein bolus. These remarkable differences make *Toxorhynchites* an attractive comparative reference for understanding mosquito chemosensation as it pertains to host-seeking. We performed deep transcriptome profiling of adult female *Toxorhynchites amboinensis* bodies, antennae and maxillary palps, and identified 25,084 protein-coding “genes” in the de novo assembly. Phylogenomic analysis of 4,266 single-copy “genes” from *T. amboinensis*, *Aedes aegypti*, *Anopheles gambiae*, and *Culex quinquefasciatus* robustly supported *Ae. aegypti* as the closest relative of *T. amboinensis*, with the two species diverged approximately 40 Ma. We identified a large number of *T. amboinensis* chemosensory “genes,” the majority of which have orthologs in other mosquitoes. Finally, cross-species expression analyses indicated that patterns of chemoreceptor transcript abundance were very similar for chemoreceptors that are conserved between *T. amboinensis* and *Ae. aegypti*, whereas *T. amboinensis* appeared deficient in the variety of expressed, lineage-specific chemoreceptors. Our transcriptome assembly of *T. amboinensis* represents the first comprehensive genomic resource for a nonblood-feeding mosquito and establishes a foundation for future comparative studies of blood-feeding and nonblood-feeding mosquitoes. We hypothesize that chemosensory genes that display discrete patterns of evolution and abundance between *T. amboinensis* and blood-feeding mosquitoes are likely to play critical roles in host-seeking and hence the vectorial capacity.

**Key words:** *Toxorhynchites*, mosquito, chemosensation, chemosensory receptor, transcriptome, RNA-seq, host-seeking, blood feeding, disease vector, chemical ecology.

## Introduction

The notorious reputation garnered by many mosquito species as human disease vectors is exclusively attributable to the concomitant behaviors of host-seeking and blood feeding (hematophagy) exhibited by adult female mosquitoes. These blood meals provide the female with a concentrated nutritive source that initiates egg development and allows the completion of the reproductive cycle. Notably, there are three nonhematophagous mosquito genera (*Malaya*, *Topomyia*, and

*Toxorhynchites*) that lack a reproductive requirement for blood meals and do not exhibit host-seeking behavior(s). Of these, *Toxorhynchites* is the most widely distributed and comprised over 90 species found throughout the tropical areas of Africa, Asia, the South Pacific, and the Americas. Adult *Toxorhynchites* are physically much larger than other blood-feeding mosquitoes and feed exclusively on sugar sources. Moreover, the mouthparts (labium) of adult *Toxorhynchites* are noticeably elongated and curved downward, a presumed

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morphological refinement toward nectar feeding that also might render them less amenable to piercing skin (Clements 1999).

Although *Toxorhynchites* is a morphologically distinctive genus within the family Culicidae, its precise placement among other genera is uncertain. Previous studies that included *Toxorhynchites* have produced several incongruent phylogenies. For example, early phylogenetic studies based on morphologic characters positioned *Toxorhynchites* as its own subfamily within Culicidae, whereas subsequent molecular studies placed *Toxorhynchites* at various locations within the subfamily Culicinae (Miller et al. 1997; Harbach and Kitching 1998; Mitchell et al. 2002; Shepard et al. 2006; Reidenbach et al. 2009). The precise phylogenetic position of *Toxorhynchites* within Culicidae is important as it would allow inference as to whether the blood-feeding phenotype represented a trait that was lost in *Toxorhynchites* or a novel adaptation that occurred only in certain Culicidae lineages, after their divergence from *Toxorhynchites*. Given that hematophagy is thought to have arisen and been lost independently multiple times in dipterans, the order of events within mosquito taxa is not immediately clear (Wiegmann et al. 2011).

*Toxorhynchites* mosquitoes are also interesting from a chemoreception perspective because, unlike the vast majority of hematophagous mosquitoes, adults have no reproductive requirement to engage in host-seeking behaviors. Host-seeking in mosquitoes is facilitated through a range of sensory cues and is largely mediated through the chemoreception of host-associated kairomones. Consequently, a characterization of the chemosensory gene repertoire of *Toxorhynchites* and its comparison to transcriptome profiles from blood-feeding mosquitoes is expected to yield insights into the molecular aspects of the chemoreceptive requirements that encompass host-seeking or blood-feeding behaviors.

Thus far, studies of chemoreceptor expression and function in mosquitoes have been limited to vector species where the identification of large gene families and the fundamental properties of chemosensory signaling continue to be elucidated. Odorant receptors (ORs), variant ionotropic receptors (IRs), and gustatory receptors (GRs) comprise the major families of chemoreceptors defined to date and no studies of chemoreceptors in nonblood-feeding mosquitoes have been published. To address these open questions regarding the ecology, evolution, and comparative genomics of *Toxorhynchites*, we have sequenced total RNA from the bodies of female *Toxorhynchites amboinensis* along with each of the two major chemosensory organs of mosquitoes, the antennae and the maxillary palps. Our data facilitated the construction of a transcriptome of the most abundant transcripts present within the adult female *T. amboinensis* as a whole, along with the characterization of chemosensory-associated transcripts that are enriched within the head appendages. To do so, we leveraged extant genomic resources of disease vector mosquitoes *Anopheles gambiae* (vector of malaria [Holt et al.

2002]), *Aedes aegypti* (vector of yellow fever and dengue [Nene et al. 2007]), and *Culex quinquefasciatus* (vector of West Nile virus [Arensburger et al. 2010]) to annotate the assembled *T. amboinensis* transcriptome as well as place *T. amboinensis* into a well-defined phylogenetic context. Finally, we have compared the transcript abundance patterns of the major chemoreceptor gene families in the olfactory tissues of three mosquito species to further refine our understanding of the molecular bases for host-seeking behaviors. This study brings together a de novo assembly of the *T. amboinensis* transcriptome and a novel comparative analysis of chemoreceptor evolution and expression between hematophagous and nonhematophagous mosquitoes.

## Materials and Methods

### Mosquitos and Mosquito Rearing

*Toxorhynchites amboinensis* eggs were obtained from the US Centers for Disease Control and Prevention Dengue Laboratory in San Juan, Puerto Rico. Eggs were hatched in distilled water in an upright incubator at 26°C and 75% relative humidity under a 12:12 light dark cycle. *Toxorhynchites amboinensis* larvae were reared on a diet of *Ae. aegypti* or *An. gambiae* larvae, with prey developmental stage (L1–L4) approximating that of predator. Pupae were transferred to small cups of clean distilled water and allowed to eclose inside a 100 × 50 × 50 cm cage. Adults were provided apple slices plus cotton balls soaked with a solution of 10% sucrose, 10% honey ad libitum. Females were given access to a small, black melamine bowl, approximately 50 mm in diameter, partially filled with 100 ml distilled water plus 1% oak leaf infusion for oviposition.

### RNA Isolation and RNA Sequencing

Approximately 200 antennae or maxillary palps were removed using surgical forceps from randomly selected 7- to 14-day-old adult females and placed into Trizol reagent for total RNA extraction. Additionally, 20 bodies from the same females, minus antennae and palps, were treated similarly. Messenger RNA (mRNA) isolation and cDNA library preparation were carried out using the Illumina mRNA sequencing kit (Illumina Inc., San Diego, CA). Libraries were sequenced by the HudsonAlpha Institute for Biotechnology (Huntsville, AL) using an Illumina HiSeq2000.

### Data Preprocessing and De Novo Transcriptome Assembly

Initial quality assessment of raw RNA sequencing (RNA-seq) data was performed using FastQC (version 0.10.1; <http://www.bioinformatics.babraham.ac.uk/projects/fastqc/>, last accessed October 22, 2014) and several issues were identified, including adapter contamination and low quality reads. To ensure the quality of transcriptome analyses, the following

preprocessing procedures were applied to RNA-seq data from each tissue: 1) Reads that failed the Illumina “chastity” filter (a measurement of the overall signal purity of reads) were first removed, 2) reads with adapter contamination were trimmed using Trimmomatic (version 0.22; Lohse et al. 2012), 3) reads were mapped against known *T. amboinensis* ribosomal RNA sequences and sequenced mitochondrial genomes of Culicidae species (downloaded from National Center for Biotechnology Information [NCBI] at <http://www.ncbi.nlm.nih.gov/genomes/OrganelleResource.cgi?taxid=7157>, accessed July, 2012) using Bowtie 2 (version 2.0.0; Langmead and Salzberg 2012) to remove reads that are derived from rRNA/mitochondrial genes, and 4) a final quality-based trimming was performed using Trimmomatic (version 0.22) with the following parameters: LEADING:3 TRAILING:3 SLIDINGWINDOW:4:15 MINLEN:25. All abovementioned data filtrations were conducted in paired-end mode. Cleaned data from all three tissues were combined together and assembled using Trinity (version 2012-06-08; Grabherr et al. 2011) to generate a de novo transcriptome assembly. Trinity was run with default settings and only reported transcripts of 200 bp or longer. No filtering of low-abundance transcripts was performed in order to maximize the detection of chemosensory genes as some of them may have low transcript abundances.

### Gene Identification

To annotate protein-coding genes in the transcriptome assembly, open reading frames (ORFs) of at least 300 bp long were first extracted from all assembled transcripts in all six frames, allowing for overlap. For each ORF, the translated protein sequence was compared against the NCBI nonredundant (NR) protein database using BLASTP (version 2.2.26+; Altschul et al. 1997) with an e value cutoff of 1e-5 and the bit score of the best match was recorded. Results of this similarity search were later used to generate the taxonomic distribution of best matches for annotated genes. ORFs with significant coding potential were also identified from the complete set of ORFs using the TransDecoder utility of Trinity.

This initial set of annotation contains a high level of redundancy as more than half of the annotated transcripts are alternative splicing isoforms (i.e., transcripts reconstructed from the same Trinity component which ideally represent one gene). To avoid this redundancy in subsequent sequence and expression analyses, a reference transcriptome assembly was created by selecting one representative transcript for each Trinity subcomponent. The representative transcripts were determined as following: 1) For each transcript, the highest bit score sum was calculated by comparing all possible combinations of nonoverlapping ORFs identified on that transcript, including both the ORFs with homologs in the NR database and the ORFs with coding potential; 2) for each Trinity component, the transcript with the highest bit score sum was

selected as the representative; and 3) the longest one was chosen if multiple isoforms have the same bit score sum. For each representative transcript, the ORFs in the combination that gave rise to the highest bit score sum were considered annotated genes; the one with the longest total ORF length was selected if multiple combinations have the same bit score sum.

### Functional Annotation

Two approaches were employed to obtain functional information about annotated *T. amboinensis* genes. First, translated protein sequences of all genes were compared against the UniProtKB/Swiss-Prot database (release 2013\_11) using BLASTP with an e value cutoff of 1e-5; each *T. amboinensis* genes was assigned the gene ontology (GO) terms associated with its best match in the database. Second, as a more conservative strategy, InterProScan 5 (data version 38.0; Quevillon et al. 2005; Hunter et al. 2012) was used to identify conserved protein domains encoded by *T. amboinensis* genes and the GO terms associated with protein domains were transferred to corresponding genes. Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway annotations of *T. amboinensis* genes were generated using the KEGG Automatic Annotation Server. Genes encoding (retro-)transposons were identified using TransposonPSI (version 08222010; <http://transposonpsi.sourceforge.net/>, last accessed October 22, 2014).

### Phylogenetic Analysis and Divergence Time Estimation

OrthoMCL (version 2.0.3; Li et al. 2003) was used to identify ortholog groups between *T. amboinensis* and four other dipteran species, including *An. gambiae* (version AgamP3.6), *Ae. aegypti* (version AeagL1.3), *C. quinquefasciatus* (version CpipJ1.3), and *Drosophila melanogaster* (version 5.48). Gene annotations of the three mosquitoes and the fruit fly were downloaded from the websites of VectorBase (<https://www.vectorbase.org>, last accessed October 22, 2014) and FlyBase (<http://flybase.org/>, last accessed October 22, 2014), respectively, and only the longest isoform of each gene was used in this study. The ortholog groups that contain a single gene from each of the five species were used to infer the phylogenetic relationship between species. For each of these groups, a codon-based alignment of the coding sequences was created using MAFFT (version 7.037b; Katoh and Standley 2013) with the “auto” option and poorly aligned regions in the alignment were removed using trimAl (version 1.4; Capella-Gutierrez et al. 2009) with the “automated1” option. Maximum-likelihood (ML) trees were constructed from both individual alignments and a concatenated alignment of all single-copy ortholog groups using RAxML (version 7.4.2; Stamatakis 2006) with the “GTRGAMMA” model and 100 bootstrap replicates were carried out to evaluate the reliability of tree topologies.

The 144 orthologous groups that recovered the same ML tree topology as the concatenated analysis with full support were used for the divergence time estimation using BEAST2 (version 2.0.2; Bouckaert et al. 2014). The analysis was performed with the following settings: Partitions, data set was partitioned by codon positions; site model, GTR+GAMMA (four categories); clock model, relaxed clock log normal; and tree priors, birth–death model. The separation of mosquitoes and *Drosophila* at 227 Ma (95% confidence interval 210–244 Ma), as suggested by a recent study (Wheat and Wahlberg 2013), was used as a calibration point. Four independent runs were carried out, each with 100 million generations and one tree being sampled every 10,000 generations. Convergence check and tree summarization were performed using Tracer (version 1.5) and TreeAnnotator (version 2.0.2), respectively.

### Chemosensory Gene Family Analysis

The annotation of chemosensory genes was performed following a previous protocol (Zhou et al. 2012). In brief, previously reported chemosensory genes from *An. gambiae*, *Ae. aegypti*, *C. quinquefasciatus*, and *D. melanogaster* were used as queries in TBLASTN searches against the complete *T. amboinensis* transcriptome assembly. Putative chemosensory gene coding regions were identified after filtering out low-scoring blast hits. For each region, the query sequence that yields the highest bit score was selected as reference to perform homology-based gene prediction using GeneWise (version 2.2.0; Birney et al. 2004) to avoid fragmented gene models due to unspliced introns. All gene models were manually inspected and modified if needed, and redundant models derived from isoforms were removed.

For each of the *OR/GR/IR* families, protein sequences of genes in the four mosquitoes were aligned using MAFFT with a high accuracy option (–maxiterate 1000 –genafpair). The multiple sequence alignments were manually curated and poorly aligned regions were removed using trimAl with the “automated1” option. ML trees were constructed using RAxML with the PROTGAMMALG model and 100 bootstrap replicates. Trees were annotated and displayed using the website of iTOL (Letunic and Bork 2011). *Toxorhynchites amboinensis* *OR/GR/IR* genes were named (supplementary table S7, Supplementary Material online) according to their (co-)orthologs in *Ae. aegypti* (or *An. gambiae* if the former is absent; e.g., *TaOr7* is the ortholog of *AaOr7* and *AgOr7*). Lineage-specific duplicates were further distinguished by appending a unique number identifier following the orthologs number (e.g., *TaOr3.1* and *TaOr3.2* are co-orthologs to *AaOr3*)

### Transcript Abundance Calculation

Gene expression values for *An. gambiae* samples were taken directly from our previous study on nonblood-fed female chemosensory appendages (Pitts et al. 2011). Gene expression

values for *Ae. aegypti* were generated by mapping read sets from nonblood-fed female antenna (Sequence Read Archive [SRA] depositions: SRR1103960, SRR1103962, and SRR1103964) and palp (courtesy of Dr Joseph Dickens [USDA, ARS]) to the current version of the genome using Bowtie2/TopHat2 (Langmead et al. 2009) and then counting the reads mapped to each gene from the resulting alignment file to the *Ae. aegypti* genome features (Vectorbase; version 2.2 with misassigned transcript IDs removed). Gene expression values for *T. amboinensis* were generated by mapping the reads used in the assembly to the Trinity transcriptome assembly (representative isoforms) using Bowtie2. In all cases, normalized gene expression levels were calculated for each tissue in terms of reads per kilobase per million reads mapped (RPKM) (Mortazavi et al. 2008).

### Data Availability

The raw sequence reads data generated in this study are available through the NCBI SRA database under the BioProject accession number: Pending. The complete transcriptome assembly of *T. amboinensis* will be available through the NCBI Transcriptome Shotgun Assembly database with the accession number: Pending. The protein and coding sequences as well as functional annotations of all annotated *T. amboinensis* genes are available at <http://dx.doi.org/10.6084/m9.figshare.1092617> (last accessed October 22, 2014).

## Results and Discussion

### A Comprehensive Transcriptome Assembly of *T. amboinensis*

In-depth RNA-seq profiling of antennae, maxillary palps, and whole bodies of female *T. amboinensis* generated more than 90 million pairs of raw reads from each sample, most of which passed rigorous quality control procedures (supplementary table S1, Supplementary Material online). Collectively, we obtained approximately 300 million pairs of high-quality sequence reads and created a de novo transcriptome assembly that was 267.6 Mb in size and consisted of 303,383 transcripts belonging to 199,064 Trinity components (table 1), which ideally each represents one gene. These numbers are an order of magnitude higher than that of annotated transcripts and genes in other available mosquito genomes. However, alternative assembly strategies and programs consistently generated comparable numbers of transcripts and components (supplementary table S2, Supplementary Material online), suggesting that these surprisingly large numbers were unlikely due to artifacts/redundancies in the assembly. Our presumption is that the vast majority of these assembled transcripts is noncoding.

We then annotated protein-coding regions (hereafter referred to as “genes” unless otherwise indicated) from the transcriptome assembly. All genes represent continuous

**Table 1**Summary of *Toxorhynchites amboinensis* Transcriptome Assembly

|                       | Complete Assembly | Protein-Coding Components |                         |
|-----------------------|-------------------|---------------------------|-------------------------|
|                       |                   | All Isoforms              | Representative Isoforms |
| Total size            | 267.6 Mb          | 171.21 Mb                 | 41.0 Mb                 |
| Number of components  | 199,064           | 21,605                    |                         |
| Number of transcripts | 303,383           | 80,552                    | 21,605                  |
| Longest transcript    | 28,920 bp         | 28,920 bp                 | 28,908 bp               |
| N50 transcript length | 2,082 bp          | 3,575 bp                  | 3,041 bp                |
| GC content            | 41.51%            | 43.42%                    | 43.86%                  |
| CEGMA completeness    | 248/248           | 241/248                   | 240/248                 |

NOTE.—The completeness of *T. amboinensis* transcriptome assembly was evaluated using CEGMA based on the presence/absence of 248 conserved eukaryotic genes. All the seven genes missed in annotated transcripts (all isoforms) and seven of the eight genes missed in annotated transcripts (representative isoforms) are less than 300 bp long, which is the lower limit of gene length to be considered in our annotation.

ORFs supported by either sequence homology to known protein-coding genes or significant protein-coding potential as determined by a likelihood-based approach. All subsequent analyses in this study were based on genes rather than transcripts because a nonnegligible portion (approximately 13.5%) of the transcripts contained more than one gene (supplementary fig. S1A, Supplementary Material online), which may confound the investigation of gene expression and evolution. In total, we identified 21,605 components each of which contains at least one annotated gene (table 1) about half of which (10,195) contain multiple transcripts (supplementary fig. S1, Supplementary Material online). We used homology criteria (see Materials and Methods) to select the most representative transcript for each component, resulting in a final annotation set composed of 25,084 genes (table 2).

Approximately 80% of the annotated *T. amboinensis* genes had at least one homolog in the NCBI NR database (conserved genes). Not surprisingly, the best match for more than 90% of *T. amboinensis* genes was with sequences from insects, with *Ae. aegypti*, *C. quinquefasciatus*, and *An. gambiae* being the top three source species in descending order (fig. 1A). Previous studies have revealed significant contributions of transposable elements (TEs) to the three sequenced mosquito genomes (Holt et al. 2002; Nene et al. 2007; Arensburger et al. 2010); strikingly, almost half of the *Ae. aegypti* genome assembly was composed of various types of TEs. Similarly, we found 4,888 *T. amboinensis* genes (approximately 20%) likely encode (retro-) transposon-related proteins, which cover many major types of TEs (table 2 and supplementary table S3, Supplementary Material online).

The remaining 20% of genes lacked detectable homologs in the database, but are likely protein coding according to their sequence signatures. These types of genes are also termed “orphan” and have been recently documented in a wide range of organisms (reviewed in Tautz and Domazet-

**Table 2**Summary of *Toxorhynchites amboinensis* Transcriptome Annotation

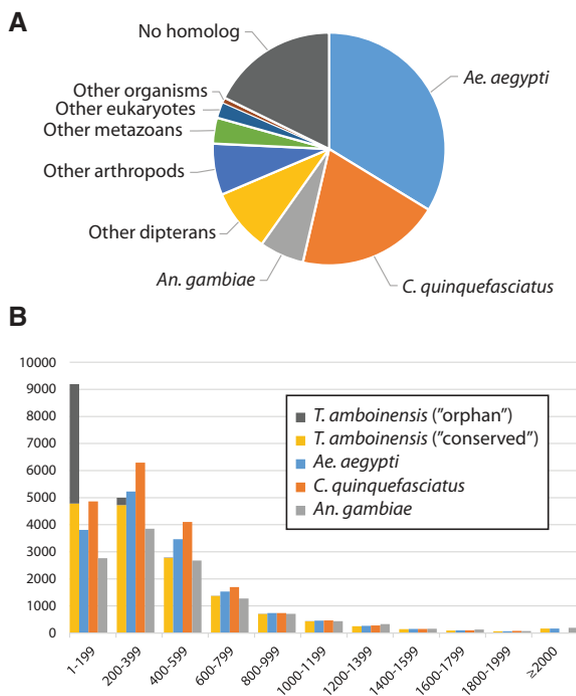
|                                 |                               |
|---------------------------------|-------------------------------|
| Total Gene Count                | 25,084                        |
| Genes with homolog in NR        | 20,278                        |
| Genes with coding potential     | 4,770                         |
| Genes coding (retro-)transposon | 4,888                         |
| Total gene length               | 26.7 Mb                       |
| Longest gene length             | 24,453 bp (8,151 amino acids) |
| Median gene length              | 663 bp (221 amino acids)      |
| GC content                      | 46.79%                        |

Loso 2011), including insects (Wissler et al. 2013). For example, orphans comprise approximately 5% of the annotated genes in each of the three mosquito genomes (Wissler et al. 2013). Such genes may be fast evolving and thus too divergent from their homologs in other species to be recognized solely based on sequence similarity. Alternatively, orphans may represent novel genes that recently evolved in the *T. amboinensis* lineage, after its divergence from other mosquitoes. In general, transcripts of orphan genes were less abundant than those of conserved genes in *T. amboinensis* (supplementary fig. S2, Supplementary Material online). That said, more than 1,500 of them had transcript abundances above the transcriptome median level in at least one of the three samples, and over 2,000 showed significant differential enrichment among samples. In addition, approximately 300 orphan genes encode hypothetical protein products that are longer than 200 amino acids, with the largest one comprising 1,002 residues (fig. 1B). Altogether, the data suggest that some of these orphan genes may indeed be functional and have contributed to the morphological and physiological differences between *T. amboinensis* and its relatives.

### Completeness of Transcriptome Assembly of *T. amboinensis*

The transcriptome assembly and gene annotations generated here constitute valuable resources for the study of *T. amboinensis* biology, with the caveat that, to some extent, the assembly is likely incomplete. Indeed, although we have performed in-depth transcriptome profiling on whole bodies and two chemosensory appendages of female *T. amboinensis*, genes that have low abundance levels or show restricted expression in specific tissues and/or developmental stages beyond our sample set will be missed or at the very least, underrepresented. However, these limitations aside, it is reasonable to posit that the transcriptome assembly generated here is relatively comprehensive based on the following lines of evidence (see supplementary note, Supplementary Material online, for details).

An evaluation of the assembly completeness using Core Eukaryotic Gene Mapping Approach (CEGMA) (Parra et al. 2007) showed that all of the 248 conserved eukaryotic



**Fig. 1.**—Summary characteristics of *T. amboinensis* proteins identified in the transcriptome assembly. (A) Taxonomic distribution of best hits of annotated *T. amboinensis* proteins in the NCBI NR protein database. (B) Length distribution of proteins identified in the transcriptome assembly of *T. amboinensis* and sequenced genomes of three blood-feeding mosquitoes, including *Ae. aegypti*, *C. quinquefasciatus*, and *An. gambiae*. For *T. amboinensis*, the “conserved” and “orphan” proteins refer to proteins with and without homolog in the NR database, respectively.

genes can be found in the complete transcriptome assembly (table 1). In addition, the number and length distribution of genes annotated in the *T. amboinensis* transcriptome assembly closely resemble those in the three sequenced mosquito genomes analyzed here (fig. 1B). More importantly, our parallel de novo transcriptome assembly analysis in *An. gambiae* (using much smaller transcriptome data sets but following the same assembly and annotation protocol) was able to recover more than 70% of *An. gambiae* genes, more than half of which was recovered at full length. Taken together, these considerations indicate that the *T. amboinensis* transcriptome assembly generated here likely represents a comprehensive genomic resource. Sequences and transcript abundances of all annotated *T. amboinensis* genes are available along with detailed functional annotations, including GO, protein domains, and metabolic pathways (see Data Availability).

Compared with other mosquitoes, *T. amboinensis* larvae are predatory whereas adults are very large and do not take blood meals, which suggests that salient aspects of its development, immunity, metabolism, and neuronal systems would likely be unique. In this regard, a comprehensive transcriptome assembly also enabled a genome-wide survey of gene families

of interest. For example, we have identified 354 immunity-related genes in *T. amboinensis*, which covered all major pathways currently known in mosquitoes and this number is similar to the total number of immunity genes reported in the three mosquito genomes (Waterhouse et al. 2007; Bartholomay et al. 2010) (supplementary table S4, Supplementary Material online; also see later section on chemosensory gene families). In conclusion, the sequence and functional information generated here is expected to significantly facilitate future studies of this nonblood-feeding mosquito as well as comparative analyses with its blood-feeding relatives.

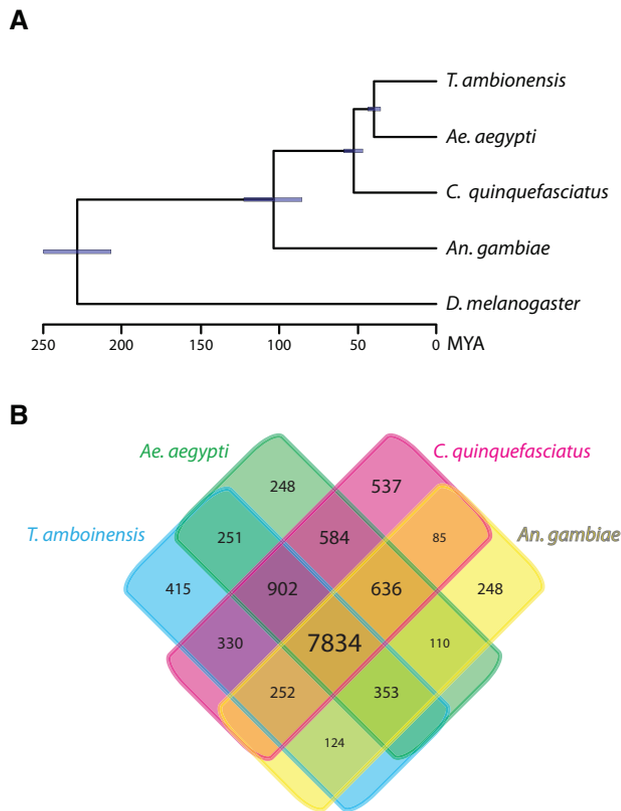
### Resolving the Phylogenetic Placement of *T. amboinensis* within Culicidae

The transcriptome assembly of *T. amboinensis* provides a unique opportunity to examine its precise placement within Culicidae using genome-scale data. We constructed orthologous gene groups from the four mosquitoes plus *D. melanogaster*, which was included as outgroup, and identified 4,266 genes that are single-copy in all five species. The concatenated maximum-likelihood analysis of these single-copy genes (6,807,855 nucleotide positions in total) resulted in a fully resolved tree placing *T. amboinensis* sister to *Ae. aegypti* (fig. 2A) relative to the other two mosquitoes. This grouping of *T. amboinensis* and *Ae. aegypti* was also strongly supported by most of the single gene phylogenies (supplementary fig. S3, Supplementary Material online). Furthermore, using the 144 genes that fully supported the concatenated ML tree, we recovered the same topology in a Bayesian analysis and estimated that the ancestors of *T. amboinensis* and *Ae. aegypti* separated about 39.6 Ma, soon after the estimated divergence of the ancestors of *Ae. aegypti* and *C. quinquefasciatus* at about 52.5 Ma (fig. 2A).

The genome-scale data presented here display overwhelming support for a sister relationship between *T. amboinensis* and *Ae. aegypti*. Additional phylogenomic studies with broader taxon sampling will be helpful to further evaluate this relationship, as only four mosquito species were included here. Nevertheless, the phylogeny proposed here provides a solid evolutionary context for comparative studies between *T. amboinensis* and other mosquitoes. When one considers that most mosquito genera are populated with species that are capable of taking blood meals for reproduction, the more derived position of *T. amboinensis* suggested here indicates that blood feeding is an ancestral trait that has been lost in the lineage leading to *T. amboinensis*.

### Characterization of the Chemosensory Gene Repertoire of *T. amboinensis*

The transition from the ancestral hematophagous state to the derived nonblood-feeding state in *T. amboinensis* is expected to be accompanied by changes in lifestyle requirements, most notably the loss of a need to engage in host-seeking



**FIG. 2.**—Phylogenetic analyses of orthologous genes supporting a sister relationship between *T. amboinensis* and *Ae. aegypti*. (A) Chronogram showing the phylogenetic relationships and estimated divergence times between *T. amboinensis* and other mosquitoes. *Drosophila melanogaster* was included as an outgroup, and its separation with Culicidae at 227 Ma (95% confidence interval: 210–244 Ma) was used as a calibration point in the divergence time estimation. The purple bar on each node indicates the 95% confidence interval of estimated divergence time. The same tree topology was recovered in a concatenated ML analysis of 4,266 single-copy orthologous genes with maximum supports for all nodes, and also received overwhelming supports from ML analyses of individual genes. (B) A Venn diagram showing the number of orthologous gene groups shared between four mosquito species.

behavior(s). Inasmuch as these behaviors are strongly mediated by chemosensory cues in hematophagous mosquitoes, it is reasonable to ask whether or not the repertoire of peripherally expressed, chemosensory genes in *T. amboinensis* may have changed concomitantly with the shift in feeding behavior. With a high quality transcriptome assembly of *T. amboinensis* available and its phylogenetic affiliation resolved, we now have the opportunity to carry out comparative analyses of chemosensory genes between blood-feeding and non-blood-feeding mosquitoes.

A large number of chemosensory genes, including 87 ORs, 21 GRs, and 38 IRs were identified in the transcriptome assembly of *T. amboinensis* (also see [supplementary table S5, Supplementary Material online](#), for information on additional

**Table 3**

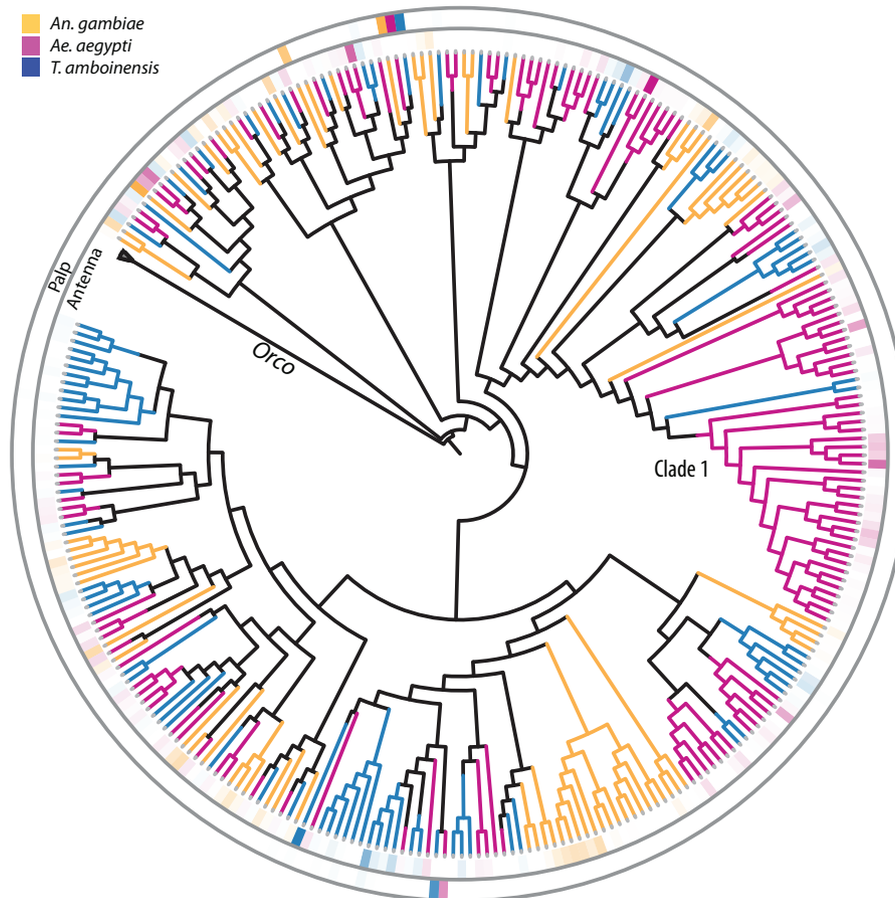
Number of Chemosensory Genes in Mosquitoes

|                                   | OR  | GR | IR      |
|-----------------------------------|-----|----|---------|
| <i>Anopheles gambiae</i>          | 75  | 61 | 46 (22) |
| <i>Culex quinquefasciatus</i>     | 178 | 65 | 69 (45) |
| <i>Aedes aegypti</i>              | 127 | 70 | 95 (37) |
| <i>Toxorhynchites amboinensis</i> | 87  | 21 | 38 (31) |

NOTE.—Numbers in parenthesis indicate antennal IRs.

chemosensory genes). In comparison to its closest relative, *Ae. aegypti*, we identified fewer chemosensory genes in *T. amboinensis*, which may reflect excessive loss in *T. amboinensis* (table 3) or may be attributed to genes that are present in the genome but whose transcripts were absent in the samples included in this study. In fact, a previous transcriptome profile in *An. gambiae* indicated that many chemosensory genes have very low expression levels in adult chemosensory appendages and whole bodies of both sexes (Pitts et al. 2011). Moreover, several *An. gambiae* OR (AgOr) genes are specifically expressed during larval stages or other tissues (Xia et al. 2008; Pitts et al. 2011). Indeed, 31 of the 38 IR genes identified in the *T. amboinensis* transcriptome assembly belong to a subfamily (“antennal” IRs) whose members in other insects have been shown to be expressed in antennae (Croset et al. 2010); this number is close to that in *Ae. aegypti* which has many more IR genes ([supplementary table S5, Supplementary Material online](#)).

To better understand the changes in the chemosensory gene repertoire of *T. amboinensis*, we reconstructed the evolutionary histories of mosquito genes in the three major chemosensory receptor gene families, namely ORs, GRs, and IRs. We focused on these gene families, which together comprise the chemosensory receptome, because their functional roles in chemosensation are much better characterized compared with other gene families such as odorant-binding proteins. This analysis reveals that most chemosensory receptor genes in *T. amboinensis* have clear (co-)orthologs in *Ae. aegypti*. In the OR gene family phylogeny (fig. 3), we identified 31 well-supported clades (bootstrap support  $\geq 70\%$ ) that include both *T. amboinensis* and *Ae. aegypti* ORs (*AaOrs*), each of which likely represents one single gene in the common ancestor of the two mosquitoes; 27 of them also contain ORs in *C. quinquefasciatus* and/or *An. gambiae*. In total, the 31 clades encompass 58 of the 87 *T. amboinensis* OR genes (*TaOrs*). Among these clades, *T. amboinensis* and *Ae. aegypti* have the same number of genes in 16 clades, including 13 one-to-one orthologous pairs, and they differ by only one gene in nine clades (fig. 3 and [supplementary fig. S5, Supplementary Material online](#)). Similar results were obtained from the phylogenetic analyses of GR and IR families ([supplementary figs. S6 and S7, Supplementary Material online](#)); 19 of 21 *T. amboinensis* GR genes (*TaGrs*) and 35 of 38 *T.*



**Fig. 3.**—OR gene tree and their relative expression in the antennae and palp of *T. amboinensis*, *Ae. aegypti*, and *An. gambiae*. Gene phylogeny of the OR-family of chemoreceptors. Branches are color coded by the respective species and the heat maps at the branch tips represent the RPKM-percent composition of each family member within the entirety of that family's expression within the tissue of each species. The outer heat map ring displays the tissue expression profile in maxillary palps and the inner heat map ring displays the tissue expression profile in antennae. *Orco* was used to root the tree and the *Orco* clade was collapsed to emphasize differences between the tuning ORs.

*amboinensis* IR genes (*Talrs*) have well-supported (co-) orthologs in *Ae. aegypti*, many of which are one-to-one orthologous pairs (13 *TaGr*s and 15 *Talr*s).

These data suggest that the chemosensory receptor gene repertoire of *T. amboinensis* (as recovered from its transcriptome assembly) is relatively stable compared with the other three mosquito genomes. In particular, six OR clades, eight GR clades, and nine IR clades are well supported and each is comprised by a single gene in each of the four mosquitoes. Among these single-copy clades are several genes with well-characterized or potentially important chemosensory functions, such as *AaOrco* (for convenience, here we use the name of *Ae. aegypti* gene in each clade) which encodes the highly conserved OR coreceptor, *AaGr1-3* which encodes receptors for carbon dioxide (Kent et al. 2008), *AaGr7* and *AaGr11* which encode candidate sugar receptors (Kent et al. 2008), *AaGr14* which is orthologous to a *D. melanogaster Gr* (*DmGr66a*) required for caffeine response (Moon et al. 2006; Kent et al.

2008), as well as a number of aforementioned conserved "antennal" IR genes, including *Aalr8a*, *Aalr21a*, *Aalr25a*, *Aalr31a*, *Aalr40a*, *Aalr64a*, *Aalr75i*, *Aalr76b*, and *Aalr93a*.

There are also several apparent differences between the chemosensory gene repertoires of *T. amboinensis* and *Ae. aegypti*. There are eight *TaOr*s, two *TaGr*s, and three *Talr*s whose (co-)orthologs are missing in the *Ae. aegypti* genome, but can be found in the other two, more distant mosquito genomes, indicating lineage-specific loss in *Ae. aegypti*. In addition, 21 *TaOr*s do not have clear orthologs in any of the three blood-feeding mosquitoes, which may suggest gene loss in these species; alternatively, it is possible that these genes have experienced relatively rapid sequence divergence and thus their relationship with other genes has become more difficult to resolve. Furthermore, we found a relatively modest expansion of genes in *T. amboinensis* relative to *Ae. aegypti* in two OR clades (six *TaOr*s vs. two *AaOr*s in one clade, and five *TaOr*s vs. one *AaOr*s in the other) and one IR

clade (six *Talrs* vs. two *Aalrs*). Conversely, more clades in the three families have higher numbers of *Ae. aegypti* genes than observed in *T. amboinensis*. Most strikingly, one such clade in the *OR* family included 40 *AaOrs* and only 2 *TaOrs*. Importantly, a large number of *Ae. aegypti* chemosensory genes lack orthologs in the transcriptome assembly of *T. amboinensis*, including 30 *AaOrs* and 43 *AaGr*s.

#### Chemoreceptive Transcript Abundance in *T. amboinensis*

To contextualize these newly annotated *ORs*, *IRs*, and *GRs* within the olfactory tissues of *T. amboinensis*, we next calculated the relative transcript abundance levels in terms of RPKM. We then compared the chemoreceptor transcriptome profile of *T. amboinensis* with both that of its close relative *Ae. aegypti* as well as *An. gambiae*. To accommodate the fact that *de novo* assembly is unable to account for *T. amboinensis* chemoreceptors that yield little to no transcriptional signal (i.e., those that may be present in the genome but are otherwise not discernable), we have limited our quantitative analysis to transcripts that showed a minimum abundance of two RPKM. This level is designated as being “detectable” and is applied to all data sets analyzed here.

#### Odorant Receptors

*ORs* appear to have evolved exclusively within insects and form heteromeric ion channel complexes consisting of at least two subunits: One a highly conserved coreceptor (*Orco*) whereas the other (*ORx*) belongs to a class of highly divergent ligand-specifying partners (Clyne et al. 1999; Vosshall et al. 1999; Larsson et al. 2004; Sato et al. 2008; Missbach et al. 2014). *Orco* is required for membrane localization of *OR* complexes, whereas *ORx* confers the odor tuning properties (Benton et al. 2006). Within this paradigm, ligands for numerous *AgOrs* and a more limited number of *ORs* from *AaOrs* and *C. quinquefasciatus* *ORs* (*CqOrs*) have been identified (Carey et al. 2010; Pelletier et al. 2010; Wang et al. 2010).

#### Antennae

The tuning *OR* composition of the *T. amboinensis* antenna is complex with 67 of the 86 annotated *TaOrs* showing abundance at reliably detectable levels, with the majority of those occurring above the transcriptome wide median level of abundance. Forty-seven of the detectable tuning *TaOrs* (70%) showed relative abundances above the median level of all antennal transcripts (5.39 RPKM), a very similar number to the 52 *AaOrs* (52% of the 91 detectable *AaOrs*) above the transcriptome median in *Ae. aegypti* antennae (6.02 RPKM; supplementary table S6, Supplementary Material online). Therefore, although the vast majority of the antennal *OR* population in both *T. amboinensis* and *Ae. aegypti* appears comprised nearly identical numbers of *ORs*, there are twice as many *AaOrs* detectable below the transcriptome median.

This is potentially important as the presence of this additional, low abundance *AaOr* complexity may represent a salient difference in the effective odor receptivity between these two species.

Next, we compared the antennal expression profile of *ORs* of the three mosquito species within the context of their evolutionary history to investigate the degree to which species-specific *ORs* contribute to the transcriptional profile of their respective species (fig. 3). Analyzed in this way, the *OR* abundance profiles within each species' antennae show a nearly equal compositional division between those *ORs* that are conserved among all three species and *ORs* that are species-specific. In *An. gambiae*, *Ae. aegypti*, and *T. amboinensis*, approximately 50% of the summed expression levels of tuning *OR* abundances are attributable to conserved *ORs* whose orthologs show expression in the antennae of all three species. Clades containing orthologs to such pan-Culicidae conserved *ORs* as *AgOr11*, *AgOr35*, *AgOr39*, and *AgOr80* account for over one-third of the total *OR* abundance in their respective species alone. Perhaps, the best-characterized example of functionally conserved *ORs* is the orthologous group containing *AgOr2* (*AaOr2*, *CqOr2*, *TaOr2*) and *AgOr10* (*AaOr10*, *CqOr10*, *TaOr10*) and that have retained similar odorant sensitivities in their respective species (Hughes et al. 2010; Pelletier et al. 2010; Bohbot et al. 2011). In the antennae, *Or2/Or10* comprises 2%, 3 and 4% of the total abundance of tuning *ORs* in *Ae. aegypti*, *An. gambiae* and *T. amboinensis*, respectively. This high degree of interspecific sequential, functional and transcriptional conservation suggests a common role for *ORs* in these mosquito species, and their persistence in *T. amboinensis* further suggests that this role ranges beyond host-seeking.

The remaining half of antennal tuning *OR* abundance is attributable to subfamily or lineage-specific *ORs* within each species. For example, a large, species-specific expansion (fig. 3, Clade1) accounts for approximately 20% of the total level of tuning *ORs* in *Ae. aegypti*. Similarly, the Culicinae-specific clade containing an expansion of 17 *TaOrs* and just 7 *AaOrs* accounts for over 30% of the total level of tuning *ORs* in *T. amboinensis*. This expansion includes such *TaOrs* as *TaOr13.3*, *TaOr3.2*, and *TaOr71.1*, the latter being the single, most highly abundant *TaOr* in the antennae. Although *TaOr71.1* alone accounts for 17.8% of all tuning *ORs* (57 times that of the transcriptome median), the sum of all seven *AaOr* orthologs in this expansion account for just over 3% of tuning *ORs* in *Ae. aegypti*.

The *Orco* coreceptor represented the most highly abundant *OR* family member in each of the three species. As reported previously, levels of *Orco* in the chemosensory tissues of two Anophelinae mosquitoes were consistently close to one-to-one as compared with the sum total of all tuning *ORs* in anopheline antennae, suggesting a possible mechanism of *Orco-OR* complex regulation within olfactory receptor neurons (*ORNs*) (Pitts et al. 2011; Rinker, Pitts, et al. 2013; Rinker,

Zhou, et al. 2013). In the two Culicinae analyzed here this ratio remained close to one-to-one, although interestingly, *Orco* abundance in the antennae appears proportionally higher than the summed levels of tuning *OR* in both *Ae. aegypti* (1.40) and in *T. amboinensis* (1.30). This is in contrast to the relationship observed in the Anophelinae and suggests that *Orco* may be regulated differently between the two mosquito subfamilies.

### Palp

*OR* expression in the maxillary palps of mosquitoes is largely restricted to basiconic sensilla present on the most distal segment. Each of these sensilla is innervated by three ORNs, two of which stereotypically express one of a pair of tuning *ORs* (Lu et al. 2007); one ORN expresses a conserved ortholog of *Or8* (e.g., *AgOr8*, *AaOr8*, *CqOr8*) whereas the second contains a tuning *OR* that appears specific to either the Anophelinae subfamily (e.g., *AgOr28*) or Culicinae subfamily (e.g., *AaOr49*, *CqOr49*).

Similarly, the palps of *T. amboinensis* displayed only two detectable tuning *TaOrs*: *TaOr8* and *TaOr49* (fig. 3). In terms of levels of *OR* abundance, it is notable that the ratio of *TaOrco* to the summed RPKM of the two tuning *TaOrs* (2.02) is nearly identical to that seen in *Ae. aegypti* palp (1.90) but is reciprocal to the relationship in *An. gambiae* palp (0.89). This again recapitulates the quantitative relationships we observed between *Orco* and the tuning *ORs* in the antennae of these species, and its presence in a tissue that is well characterized as having only two tuning *ORs* further suggests a difference in regulatory mechanisms, rather than the existence of unannotated *ORs* in Culicinae.

### Ionotropic Receptors

The *IRs* are the most ancient of the three chemoreceptive gene families, yet are the most recently discovered (Benton et al. 2009; Croset et al. 2010; Liu et al. 2010) and the least well characterized in terms of their localization and function. *IRs* as a class appear particularly tuned to detection of ammonia, amines, acids and alcohols, and are expressed in ORNs, which may or may not also express *ORs* (Benton et al. 2009). In *Drosophila*, *IRs* appear to be expressed in specific ORNs of the antenna and proboscis, with the distribution of *IR* transcripts in the mosquito suggesting similar enrichments in the antenna (Pitts et al. 2011; Leal et al. 2013; Rinker, Pitts, et al. 2013) and labellum (Sparks et al. 2014).

Functionally, the *IR* gene family resembles *ORs* in that it is split into expansive groups of *IRs* that seem to confer odorant specificity, and several putative *IR* coreceptors (orthologs to *Dmlr25a*, *Dmlr8a*, and *Dmlr76b*) that are highly conserved and are likely to be necessary for proper *IR* function (Croset et al. 2010).

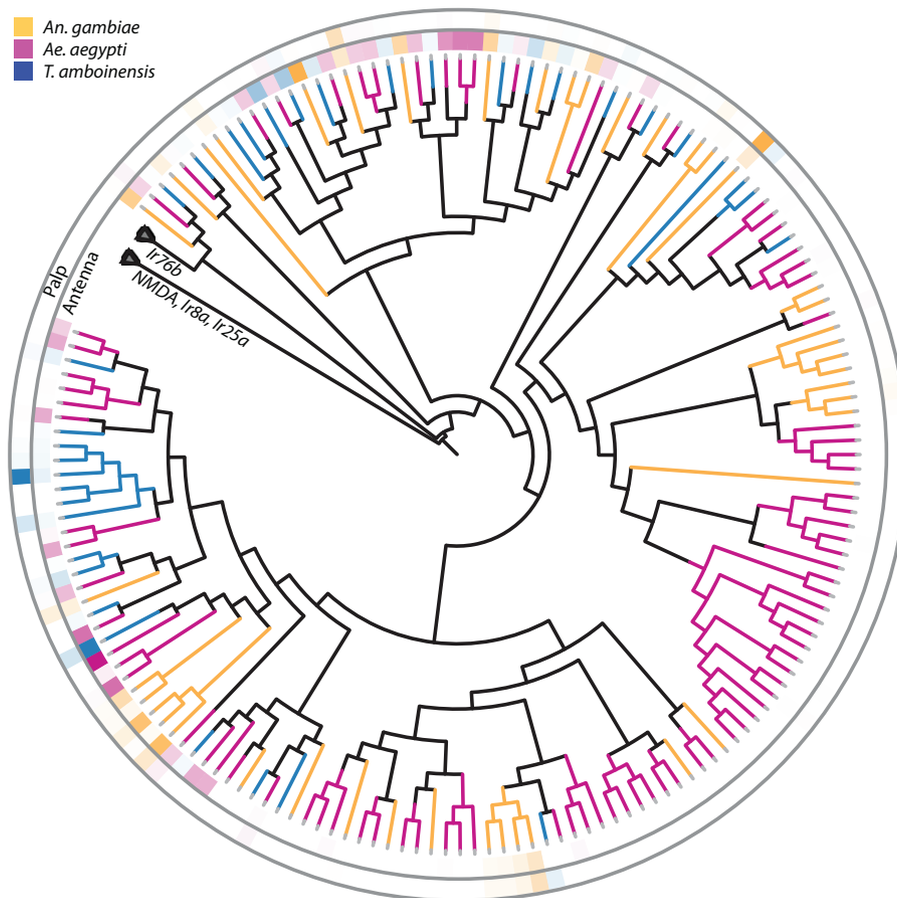
### Antennae

The *IR* composition of the antennae of all three species was very similar. Twenty-eight ligand specifying *Talr* genes were reliably detected in *T. amboinensis* antennae, nearly equal to the numbers seen in *An. gambiae* (27) and *Ae. aegypti* (26). Orthologs to all three putative, *IR* coreceptors were found to have very high transcript abundances, consistent with transcriptomic studies in the antenna of other insects (Pitts et al. 2011; Zhou et al. 2014). Indeed the most highly abundant *IR* transcripts in the antennae of each of the three species were the homologs to *Ir25a* and *Ir76b*, with *Ir8a* consistently showing an abundance level of approximately one-third that of *Ir25a*. Moreover in both of the Culicinae, the ratio of total RPKM for the three *IR* coreceptors to the RPKM total of the remaining *IRs* was remarkably consistent (0.998 in *Ae. aegypti* and 0.997 in *T. amboinensis*), a correlation reminiscent of the conserved *Orco*/tuning *OR* relationship in *Anopheles* and implying an interdependency between these two *IR* subcategories.

The *IRs* appear to differ markedly from the *ORs* with regard to their evolution and transcriptome profiles. When the abundance levels of antennal *IRs* are viewed within the context of the evolutionary history of the *IR* gene family, it appears that greater than 90% of the antennal abundance of ligand specifying *IRs* is derived from those residing in single-copy, three species clades (fig. 4). For example, the most highly abundant *IRs* in *An. gambiae* (*AgIr75l*), *Ae. aegypti* (AAEL008587), and *T. amboinensis* (*Talr41r*) each have orthologs in the other two species that are also moderately or highly expressed within those species. Furthermore, there are few notable instances of detectable *IRs* that appear to be species-specific. Only *An. gambiae* shows a modest level of some members of an *IR* clade (*AgIr41n*, *AgIr41c*, *AgIr41t.2*) that has no discernable orthologs in Culicinae. Overall, the underrepresentation in terms of abundance of lineage-specific *IRs* within mosquito species appears to confirm the observation that even though members of the *IR* gene family rapidly duplicate and diversify within species, most of those *IRs* that are detectable are highly conserved (Benton et al. 2009).

### Palp

Although primarily concentrated in the antennae, *IR* transcripts have also been reported in the maxillary palps of both *An. gambiae* (Pitts et al. 2011) and *Ae. aegypti* (Bohbot et al. 2014). *Toxorhynchites amboinensis* palps show a similar profile with a very limited number of *IRs* being detectable along with what may be the stereotypical absence (Pitts et al. 2011; Guo et al. 2013) of the *IR8a* coreceptor in insect palps. More strikingly, the *Ir76b* ortholog—the second putative *IR* coreceptor expressed in the palps of both *An. gambiae* and *Ae. aegypti*—is also not detectable within the palps of *T. amboinensis* and this absence may



**Fig. 4.**—*IR* gene tree and their relative expression in the antennae and palp of *T. amboinensis*, *Ae. aegypti*, and *An. gambiae*. Gene phylogeny of the *IR*-family of chemoreceptors. Branches are color coded by the respective species and the heat maps at the branch tips represent the RPKM-percent composition of each family member within the entirety of that family's expression within the tissue of each species. The outer heat map ring displays the tissue expression profile in maxillary palps and the inner heat map ring displays the tissue expression profile in antennae. *NMDA* receptors were used to root the tree and the *NMDA* clades were collapsed to emphasize differences between the ligand specifying *IRs*.

indicate a significant shift in the chemoreceptive capacity in the palp of this particular lineage.

### Gustatory Receptors

GRs, along with ORs, are the other class of seven transmembrane chemoreceptors that have been identified in insects (Hill et al. 2002; Bohbot et al. 2007; Kent et al. 2008; Kent and Robertson 2009). As their name implies, they are most associated with contact chemosensation although several of the family members play a role in olfaction and host-seeking (Kent and Robertson 2009; McMeniman et al. 2014).

### Antennae

In mosquitoes, *GR* transcripts are detectable in the antennae but always at very low levels (Pitts et al. 2011; Rinker, Pitts, et al. 2013). In *T. amboinensis* we observed nine detectable *TaGr*s in the antennae, seven of which were present at

abundance levels greater than the transcriptome wide median. In contrast, only two *AaGr*s and five *AgGr*s were detectable, and in both cases none reached an RPKM level above their respective transcriptome median values.

All of the detectable *TaGr*s had orthologs in both *Ae. aegypti* and *Culex*. Of the most abundant antennal *TaGr*s, the top three, *TaGr2*, *TaGr23* and *TaGr21* (RPKMS of 15.0, 13.6, and 10.3, respectively), had detectable orthologs in *Ae. aegypti* antennae (*AaGr14*, *AaGr2*, and *AaGr7*, respectively). Notably, none of the detectable *TaGr*s had any clear orthologs within *An. gambiae*.

### Palp

*GR* expression in the maxillary palps of mosquitoes has been localized to the non-OR expressing ORNs innervating the basiconic sensilla in *An. gambiae* and *Ae. aegypti* (Lu et al. 2007; Erdelyan et al. 2012). Orthologs of two of the three *GR*s

expressed here (*AaGr1/AaGr3*, *AgGr22/AgGr24*) are necessary for CO<sub>2</sub> reception in *D. melanogaster* (*Dm21a/Dm63a*) (Dahanukar et al. 2007; Jones et al. 2007; Kwon et al. 2007), and the presence of a third *GR* in the basiconic sensilla of mosquitoes has been postulated to confer enhanced sensitivity of CO<sub>2</sub> in *An. gambiae* (Lu et al. 2007). Each of the three *GRs* (*AaGr1/AaGr2/AaGr3*, *AgGr22/AgGr23/AgGr24*, and *TaGr1/TaGr2/TaGr3*) shows high sequence conservation between species and appears to function together to confer sensitivity to CO<sub>2</sub> as well as other semiochemicals in basiconic sensilla of the palp (Kellogg 1970; Grant et al. 1995; Tauxe et al. 2013). This is significant inasmuch as CO<sub>2</sub> appears to be an important long-range activator of mosquito host-seeking (Gillies 1980; Takken and Knols 1999).

All three *GRs* are expressed at very high levels in the palp in each of the three species (supplementary fig. S4, Supplementary Material online). Moreover, the pattern of expression between orthologous *GRs* appears conserved for *TaGr1* and *TaGr2* with each detected at nearly equal ratios to one another in all species. In contrast, the abundance of *TaGr3* relative to the level of the most abundant palpal *Gr* was markedly lower in *T. amboinensis* (40%) than in either *An. gambiae* (56%) or *Ae. aegypti* (68%). Because *Gr3* is essential for CO<sub>2</sub> detection in flies and confounds host-seeking in *Ae. aegypti* (McMeniman et al. 2014), the lower relative abundance of *TaGr3* could hint at an alteration in the sensitivity of this particular mode of *GR*-mediated chemoreception.

The expression analysis of chemoreceptive genes present on the antennae and palps shows that the peripheral chemoreceptive repertoire of female *T. amboinensis* mosquitoes displays two broad tendencies that are also observed in the other two Culicidae analyzed here. First, we see that the overwhelming majority of detectable *GRs* and *IRs*, along with at least half of the detectable *ORs* consist of genes that have clear orthologs in the transcriptomes of all three taxa. Second, we see that the remaining half of the *OR* composition of the antennae in each of the three taxa consist of lineage-specific *OR* genes. Together, these observations suggest a generalizable model of peripheral chemoreceptor variation that exists between mosquito species. One rationale for this is that the expression of conserved *GRs*, *IRs*, and *ORs* provides a Culicidae-wide chemoreceptive foundation upon which lineage-specific expansions of *ORs* can then dynamically and independently explore sequence and odor space within each lineage.

Beyond the generalities described above, the transcriptome profile of *T. amboinensis* also displays subtle characteristics that appear more specific to the Culicinae as well as some which distinguish it from *Ae. aegypti*. First, in all chemosensory tissues, *T. amboinensis* and *Ae. aegypti* show a modest yet inverted relationship, between the amounts of *Orco* transcript and total tuning *OR* transcripts relative to *An. gambiae*, suggesting alternative modes of *OR* regulation within the two subfamilies. Second, *T. amboinensis* displays the conserved

presence of an ortholog for *AaOr49/CqOr49* in the palp along with the absence of any *AgOr28* ortholog. These trends may represent two, broad divergences in stereotypic *OR*-expression that distinguish Anophelinae from Culicinae.

Such examples of chemoreceptive similarities between *T. amboinensis* and other mosquito species hint at conserved ranges of olfactory tuning that reflect the constraints of chemical ecology common to all Culicidae. In this light, behaviorally salient chemosensory signals such as plant, host, and oviposition site cues likely share many similarities in their odor profiles and may underlie the prevalence of the observed set of ancient and abundant chemoreceptors. Indeed, such a shared chemoreceptive repertoire of interspecific oviposition cues would be of particular importance to *T. amboinensis* whose life cycle depends heavily on depositing eggs in water sources that contain the larvae of other mosquitoes.

Where *T. amboinensis* appears to show some distinction from *Ae. aegypti* is in the number and relative abundance of certain chemoreceptive genes. First, the palp of *T. amboinensis* is lacking the detectable presence of an ortholog to *Ag/Aalr76b* which, as a putative *IR* coreceptor, could indicate an overall shift in functional *Talr* diversity in the palp. Second, *T. amboinensis* females appear to have a markedly reduced antennal *OR* diversity than *Ae. aegypti*, and this diversity is lacking specifically in the numbers of lineage-derived *OR* genes. Such examples of chemoreceptive “simplification” in *T. amboinensis* could be the result of the loss of blood feeding in this lineage, with the reduction in the variety of tuning *ORs* supporting the role *ORs* play in conferring host specificity in mosquitoes (DeGennaro et al. 2013).

## Conclusions

In this study we have performed a comparative analysis of the transcriptome profiles between *T. amboinensis* and three disease vector mosquitoes, with a focus on chemosensory gene families. Through in-depth RNA-seq of chemosensory appendages and whole bodies of adult female *T. amboinensis*, we obtained a comprehensive de novo transcriptome assembly, and confidently resolved for the first time the phylogenetic position of *T. amboinensis* thereby providing an important context for subsequent comparative studies. The transcriptome assembly, together with the detailed functional annotations we have generated (e.g., protein domains and GO), represents a valuable genomic resource that will facilitate future studies of *T. amboinensis*. In the transcriptome assembly of *T. amboinensis*, we identified a large chemosensory gene repertoire that is comparable to other mosquitoes. Phylogenetic analyses of the three major chemosensory receptor gene families revealed both clades that have maintained stable copy numbers in the four mosquitoes, as well as those with differential gain-and-loss among these blood-feeding and nonblood-feeding mosquitoes.

Subsequent transcript abundance analysis revealed that *T. amboinensis* is generally similar to other mosquito species while lacking certain chemoreceptive components present in the other taxa analyzed. Specifically, we observed a reduction in the numeric variety of antennal *ORs* as compared with *Ae. aegypti*. This marked depletion in *OR* complexity co-occurs with the retention of a large number of conserved mosquito *ORs*, indicating that *T. amboinensis* retains a core repertoire of expressed *ORs* that complement sets of conserved *GRs* and *IRs*. In the palp, we also observed the absence of one of the putative *IR* coreceptors along with indications of a potential decrease in  $\text{CO}_2$  receptivity.

Taken together, our analyses show that in its divergence from *Ae. aegypti*, *T. amboinensis* not only lost its reproductive requirement for the taking of blood meals but also experienced a measurable, diminution in the qualitative complexity of peripheral chemoreceptive coding. Chemoreceptors that appear to have been conserved in both copy number and abundance may be centrally involved in such common phenotypic behaviors as nectar feeding and ovipositing that reflect the constraints of chemical ecology.

## Supplementary Material

Supplementary note, tables S1–S7, and figures S1–S7 are available at *Genome Biology and Evolution* online (<http://www.gbe.oxfordjournals.org/>).

## Acknowledgments

The authors thank Dr Joseph Dickens (USDA, ARS) for palpal RNA-seq reads from *Ae. aegypti*, and Manuel Amador (CDC, Puerto Rico) for providing *T. amboinensis* eggs. They are also indebted to the members of the Rokas/Zwiebel labs for helpful discussions and critical readings. This work was conducted in part using the resources of the Advanced Computing Center for Research and Education at Vanderbilt University and was supported by funds provided by the National Science Foundation (DEB-0844968) and the National Institutes of Health (NIAID, AI105619) to A.R., as well as grants from the Innovation and Discovery in Engineering and Science (IDEAS) program of Vanderbilt University and from the National Institutes of Health (NIAID, AI056402) to L.J.Z. This work was also supported by the NIDCD through an NRSA (F31 DC012991) to D.C.R.

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Associate editor: Jay Storz