Transcriptome profiles of *Anopheles gambiae* harboring natural low-level *Plasmodium* infection reveal adaptive advantages for the mosquito

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Anopline mosquitoes are the sole vectors for the *Plasmodium* pathogens responsible for malaria, which is among the oldest and most devastating of human diseases. The continuing global impact of malaria reflects the evolutionary success of a complex vector-pathogen relationship that accordingly has been the long-term focus of both debate and study. An open question in the biology of malaria transmission is the impact of naturally occurring low-level *Plasmodium* infections of the vector on the mosquito's health and longevity as well as critical behaviors such as host-preference-seeking.

To begin to answer this, we have completed a comparative RNAseq-based transcriptome profile study examining the effect of biologically salient, salivary gland transmission-stage *Plasmodium* infection on the molecular physiology of *Anopheles gambiae* s.s. head, sensory appendages, and salivary glands. When compared with their uninfected counterparts, *Plasmodium* infected mosquitoes exhibit increased transcript abundance of genes associated with olfactory acuity as well as a range of synergistic processes that align with increased fitness based on both anti-aging and reproductive advantages. Taken together, these data argue against the long-held paradigm that malaria infection is pathogenic for anophelines and, instead suggests there are biological and evolutionary advantages for the mosquito that drive the preservation of its high vectorial capacity.

Globally, malaria remains the most endemic infectious mosquito-borne disease, with over 500 million cases per annum, and putative vectors present in almost 100 countries worldwide placing up to 40% of the world's population at risk. Human malaria is the result of pathogenic infection by five species of unicellular *Plasmodium* (Haemosporidia: Plasmodiidae). Of utmost importance is *Plasmodium falciparum*, which causes severe health complications and the highest human mortality rate. *Plasmodium* parasites are solely vectored between humans by mosquitoes within the genus *Anopheles*. *Anopheles* mosquitoes are anautogenous; females require a blood meal for reproduction and replenishment of energy stores. Importantly, *An. gambiae* adult females are only infectious to humans when the *Plasmodia* they harbor complete a multi-stage sporogonic cycle and successfully invade the mosquito's salivary glands. Despite the complexity of the interlaced *Anopheles-Plasmodium* life cycles and the inherent ecological and environmental challenges, it is often surprising that in malaria-endemic regions of Sub-Saharan Africa, where there is a high disease prevalence, the percentage of infectious mosquitoes within surveys of *Anopheles* populations is actually very low, typically ranging from 1 to 10%. Blood feeding, which is carried out solely by adult females, is a central factor that positively affects the complex *Anopheles-Plasmodium* paradigm. As such, *Anopheles* blood-meal host-seeking and preference are critical behaviors for regulating the synergistic processes that ultimately impact the mosquito's vectorial capacity. Indeed, several studies support a hypothesis that pathogenic malaria in humans and *Plasmodium* infection of vectors may modulate mosquito behavior and physiology to increase the likelihood of pathogen transmission. However, many of these and other efforts to examine the relationship between *Anopheles* and *Plasmodium* have for experimental simplicity...
often resorted to employing unnaturally high mosquito infection intensities that might have artifactually altered
the natural relationships between *Anopheles* and *Plasmodium*\textsuperscript{11–14}.

The vast majority of anopheline mosquitoes that ingest *Plasmodium* gametocyte-infected blood meals never
become infectious\textsuperscript{15}. In controlled laboratory studies, only 5–10\% of ingested *Plasmodium* gametocytes suc-
cessfully differentiate into ookinetes; of those, less than 0.3\% form oocysts, giving rise to approximately 1–2
oocysts per mosquito midgut\textsuperscript{15,16}. This closely mirrors West African field studies indicating that the majority of
malaria-infected mosquitoes are within that range and only rarely are trapped mosquitoes observed with more
than 5 oocysts per midgut\textsuperscript{17}. In that light, it is reasonable to suggest that when investigating salient biological
relationships between *Anopheles* and *Plasmodium* in the laboratory, the relative intensity of mosquito infections
should be low to better align with these natural levels. Paradoxically, the extremely high malaria infection inten-
sities that are often employed in laboratory studies to provide robust levels of infection prevalence are likely to
significantly alter the *Anopheles-Plasmodium* paradigm potentially giving rise to biologically irrelevant shifts in
mosquito gene expression, physiology, and behavior.

In order to assess the impact of malaria infection on chemosensory processes of *An. gambiae* we have deliber-
ately employed low gametocytemia and, thereby, biologically relevant *P. falciparum* blood meals to examine how
salivary-gland-stage sporozoite infection influences the transcriptome of female *An. gambiae* heads (including
sensory appendages) and salivary glands, henceforth referred to as ‘head’. These data suggest that natural levels
of malaria infection align with enhanced olfactory sensitivity that likely impacts chemosensory driven host
preference. Moreover, and surprisingly, natural levels of *Plasmodium* infection also appear to provide a broad
and synergistically beneficial modulation of mosquito fitness that underlies and may indeed promote the evo-
lutionary stability of the *Anopheles-Plasmodium* paradigm that continues to drive global malaria transmission.

**Results and discussion**

**Experimental design and establishment of optimal gametocytemia.** Our study was designed to
rigorously compare the impact of biologically salient *P. falciparum* infections on the head (encompassing brain,
sensory appendages, and salivary glands) transcriptome of a laboratory strain of the Afro-tropical malaria vec-
tor mosquito *An. gambiae* s.s. While every effort has been made to rigorously examine the impact of naturally
occurring *P. falciparum* infections on vector mosquitoes, it must be noted there are likely to be inherent differ-
ences between established laboratory colonies and diverse field populations of both pathogen and vectors. We
nevertheless believe the broad implications of this study remain salient to understanding the natural biology of
malaria infected anophelines.

An experimental paradigm incorporating triplicate biological replicates (21 *An. gambiae* heads per replicate)
was designed wherein 5-day-old females were blood fed on either *P. falciparum*-infected (treatment) or naïve
(control) human blood meals and thereafter reared for 18 days prior to collection and tissue dissection for down-
stream molecular analyses. This allows for completion of the mosquito sporogonic cycle as well as sporozoite
invasion of the salivary glands in treated (infected) mosquitoes (Fig. 1A), thus ensuring that mosquitoes harboring
sporozoites in their salivary glands are examined. Despite of the experimental challenges resulting from sig-
nificant reductions in prevalence we prioritized utilizing low-level intensities representative of natural biological
infections\textsuperscript{17}. To achieve this goal, approximately 14\% of the treatment (infected) mosquitoes were sampled 8 days
after blood meals, and their midguts examined for the presence of *P. falciparum* oocysts following established
protocols\textsuperscript{19}. These studies indicated that mosquitoes that had blood fed on human blood supplemented with *P*

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**Figure 1.** Schematic representation of experimental methodology. (A) Timeline for establishment of
*Plasmodium falciparum* sporozoite infected treatment and naive control *Anopheles gambiae* s.s. mosquito lines.
(B) Total RNA and gDNA extraction treatment and control protocols for PCR *P. falciparum* sporozoite infection
validation and RNA sequencing.
falciparum NF54W at a final gametocytemia of 0.15%, 0.05%, and 0.01% resulted in median/upper threshold oocyst counts that were significantly higher than the optimal levels of 1-oocyst/midgut median and 3–6 oocysts/midgut upper levels that we required. Iterative efforts established an optimal gametocyte dilution of 0.008% to meet those criteria, which was thereafter used for all treatments in this study (Table 1). The heads (including sensory appendages and salivary glands) were used for preparation of total RNA (for sequencing) and genomic DNA (gDNA) for PCR validation of the presence of \textit{P. falciparum} sporozoite-treated individual mosquitoes. These PCR-based analyses utilized primers designed to detect the \textit{P. falciparum} circumsporozoite (CS) gene as well as the \textit{An. gambiae} ribosomal protein s7 (AgRsp7) gene as an internal standard. This analysis revealed that of the 176 treatment mosquitoes that took infected blood meals and survived to 23 days of age, 65 were confirmed as having salivary-gland-stage sporozoites, representing an overall prevalence of ~37% (Fig. 2). Consistent survivor

**Table 1.** Results from midgut oocyst counts of 8-d.o. \textit{Plasmodium falciparum} infected treatment. \textit{Anopheles gambiae} s.s. mosquitoes.

| \textit{Plasmodium falciparum} Gametocytemia (%) & An. gambiae s.s Rearing Cage No. | Oocyst counts | Prevalence (%) |
|-----------------------------------------------|---------------------------------|---------------|---------------|
| \textit{P. falciparum} 0.15%, Cage 1 | 29 | 25.7 | 45 | 7 | 100 |
| \textit{P. falciparum} 0.15%, Cage 2 | 12.5 | 14.3 | 45 | 10 | 90 |
| \textit{P. falciparum} 0.15%, Cage 3 | 22 | 32.6 | 53 | 7 | 100 |
| \textit{P. falciparum} 0.15%, Cage 4 | 11.5 | 13 | 25 | 8 | 100 |
| \textit{P. falciparum} 0.15%, Cage 5 | 13 | 12.9 | 19 | 7 | 85.7 |
| \textit{P. falciparum} 0.15%, Cage 6 | 13 | 11.4 | 22 | 7 | 85.7 |
| \textit{P. falciparum} 0.01%, Cage 1 | 4.5 | 4.8 | 10 | 8 | 87.5 |
| \textit{P. falciparum} 0.01%, Cage 2 | 3 | 4.4 | 11 | 7 | 85.7 |
| \textit{P. falciparum} 0.01%, Cage 3 | 2.5 | 2.8 | 1 | 8 | 87.5 |
| \textit{P. falciparum} 0.01%, Cage 4 | 4 | 4 | 1 | 9 | 88.9 |
| \textit{P. falciparum} 0.008%, Cage 1 | 1 | 1 | 3 | 7 | 57.1 |
| \textit{P. falciparum} 0.008%, Cage 2 | 1 | 1.7 | 6 | 9 | 66.7 |
| \textit{P. falciparum} 0.008%, Cage 3 | 1 | 1.2 | 5 | 9 | 55.6 |
| \textit{P. falciparum} 0.008%, Cage 4 | 1 | 1.4 | 3 | 11 | 63.6 |
| \textit{P. falciparum} 0.008%, Cage 5 | 1 | 1.1 | 3 | 10 | 60.0 |
| \textit{P. falciparum} 0.008%, Cage 6 | 1 | 1.4 | 4 | 12 | 66.7 |

**Figure 2.** (A) \textit{Plasmodium falciparum} circumsporozoite (CS) primer set schematics indicating expected PCR amplification products. (B) Agarose gel (1.5%) electrophoresis of amplified products using CS primer set. Lanes 1–6 and 10–16 are a representative subset of the 176 examined \textit{P. falciparum} sporozoite infected treatment \textit{Anopheles gambiae} s.s. mosquitoes with a \textit{P. falciparum} (+) positive and (−) negative control (blank). Lane Ld, 100 bp DNA size marker ladder.
rates of ~50% were observed between treatment and control An. gambiae lines. The observed survival rate parallels previous studies documenting 20-day-old An. gambiae laboratory survival rates20.

**RNAseq/Bioinformatics.** Transcriptome profiling was conducted using total RNA extracted from triplicate An. gambiae treatment (infected; T1–3) and control (uninfected; C1–3) heads. Following sequence trimming and quality control bioinformatic pipelines, individual data sets were aligned to the An. gambiae PEST strain genome. Sequencing of the T1, T2, and T3 P. falciparum-infected (treatment) An. gambiae head libraries generated a total of over 254 million reads mapping to 83%, 83%, and 85% respectively of the An. gambiae genome (VectorBase-53_AgambiaePEST; Table 2). Likewise, Illumina sequencing of the C1, C2, and C3 control (uninfected) An. gambiae head libraries generated a total of 426 million reads mapping to 81%, 80%, and 81% respectively of the An. gambiae genome (VectorBase-53_AgambiaePEST; Table 2). Uniquely mapped reads were counted using the program HTSeq. DESeq2 statistical analysis identified 5765 annotated transcripts present in significantly different abundances between treatment and control replicates (p < 0.05; Fig. 3A). These annotated transcripts represent a transcriptional shift of 36% of the An. gambiae head transcriptome. Of these, 2776 transcripts were present in significantly higher abundances in treatment replicates and the remaining 2989 transcripts present in significantly higher abundance in control replicates (p < 0.05). With such a large transcriptional shift observed between treatment and control An. gambiae heads, both principal component analysis (PCA) and Pearson's correlation coefficient (PCC) calculations were performed to verify validity. PCC calculations for the three treatment and three control An. gambiae head libraries returned r values of 0.9429 and 0.9770, respectively, indicating a strong association among biological replicates (Table 2). Additionally, PCA determined that 83% of observed variance between the six library datasets was attributed to the presence/absence of P. falciparum infection, with negligible variance attributed to biological replication (Fig. 3B).

**Table 2.** Statistics of RNA sequencing for Plasmodium falciparum sporozoite infected treatment and naïve control Anopheles gambiae s.s. libraries. PCC denotes Pearson's correlation coefficient.

| Replicate | Total no. reads | Mapped reads | No. transcripts | PCC (r) | Coverage |
|-----------|-----------------|--------------|-----------------|--------|----------|
| Treatment | T1 65,613,838   | 54,459,486 (83%) | 12,194 (77%) | 0.9429 | 70x      |
|           | T2 123,249,032  | 102,296,697 (83%) | 12,799 (81%) | 131x   |          |
|           | T3 656,683,884  | 558,181,30 (85%) | 12,533 (79%) | 70x    |          |
| Control   | C1 166,457,374  | 123,178,457 (74%) | 12,848 (81%) | 0.9770 | 177x     |
|           | C2 110,612,110  | 818,529,61 (74%) | 12,625 (80%) | 118x   |          |
|           | C3 149,580,984  | 107,698,308 (72%) | 12,719 (81%) | 160x   |          |

**Figure 3.** (A) Scatterplot of log fold changes vs. mean normalized counts generated using HTSeq and DESeq2 for Plasmodium falciparum sporozoite infected treatment and naïve control library replicates (p < 0.05). (B) Principal component analysis (PCA) of the 3 sporozoite infected treatment and 3 naïve control library replicates. Gray circles represent control replicates; blue circles represent treatment replicates.

GO annotation using OmicsBox assigned 4976 GO-terms (Level II) to 2832 transcripts of the 5765 differentially abundant transcripts (control and treatment libraries). Of the total annotated transcripts, 1291 were elevated in treatment libraries and 1541 transcripts elevated in control libraries, with approximately 10% of the
transcripts having no known function or characterization (Fig. 4). Transcripts with no clear molecular function via GO were further BLASTed using Blast2GO and the GeneBank non-redundant protein database to assist in further functional analysis. The resultant assigned GO annotations are representative of a diverse array of biological processes and molecular functions that make interpretation difficult. Taken together, these analyses reveal a broad transcriptome profile shift that spans a wide range of cellular processes, some of which are likely to impact the behavior, physiology and ultimately the vectorial capacity of *An. gambiae* carrying biologically relevant *P. falciparum* sporozoite infections.

*Plasmodium* manipulation of mosquito chemosensory mechanisms. To examine the potential for behavioral modification of infected mosquitoes, our initial focus was to examine transcriptome modulations directly relevant to the chemosensory systems of *An. gambiae*. These systems collectively play an essential role in establishing and maintaining critically important behaviors, including the blood-meal host preference of female mosquitoes.

Odorant receptors. The transcript abundance of several *An. gambiae* odorant receptors (AgORs) were highly differentiated between *P. falciparum* sporozoite-infected (sporozoite; treatment) and uninfected (control) heads. Indeed, 60 of the 71 AgORs and the OR co-receptor (Orco) that were detected in sequencing libraries were consistently more abundant in heads of infected mosquitoes, with 19 displaying statistically significant induction (Fig. 5A). Transcripts encoding the remaining 11 AgORs (AgOR1, AgOR3, AgOR4, AgOR26, AgOR36, AgOR45, AgOR46, AgOR52, AgOR60, AgOR62 and AgOR72) showed an insignificant trend of less abundance in samples from infected mosquitoes (Fig. 5A). With the near complete representation of AgORs (95% of annotated *An. gambiae* ORs), it is evident that sporozoite infection significantly impacts the levels of these ORs within mosquito chemosensory structures. The majority of anopheline ORs are functionally characterized, and we have previously integrated OR functional RNAseq data to model the receptivity profile for the antennae of *An. gambiae* following their initial bloodmeal. Transcriptome profiling studies together with behavioral analyses support the hypothesis that the sensitivity of the anopheline olfactory system is directly associated with the relative abundance of olfactory transcripts. Applying this approach here suggests that the large number of significantly abundant AgOR transcripts associated with infected *An. gambiae* indicates substantial increases in odorant receptivity (Fig. 6). In fact, AgOR receptivity modelling identified 70 odorants, many of which are components of human sweat, with significantly increased receptivity in *Plasmodium*-infected mosquitoes than in uninfected controls. Of these, several [including L-lactic-acid, 1-octen-3-ol and 4-methylphenol] are human sweat components that have been shown to be attractive kairomones for host-seeking female anophelines. Notably, lactic acid showed a 217% increase in AgOR receptivity in treatments. Furthermore, the significant increases in receptivity to 1-butanol (247%), 3-methyl-1-butanol (260%), and 2,3-butanedione (252%) align with laboratory olfactometry and semi-field studies, indicating that these compounds significantly increase the number of anopheline collected (vs. catches) when added to MMX traps baited with CO2 and a basic blend of amine/carboxylic acid attractants.

These putative increases of olfactory acuity in *Plasmodium*-infected anopheline mosquitoes are consistent with behavioral bioassay and surveillance studies that determined *Plasmodium* infections to increase mosquito bite frequency while maintaining host-preference plasticity. That said, our receptivity modelling does not align with recent studies that examined peripheral olfactory responses in *P. falciparum*-infected *An. gambiae*. It is likely that the significantly higher infection intensities used in that study over-challenged *An. gambiae* infection-responsive systems, such as the innate immune system, resulting in unintended fitness consequences.
altered host-seeking and blood-feeding behaviors\textsuperscript{31}. Because our studies avoid impacts arising from non-natural infection intensities, they can more confidently reveal a clear effect of biologically relevant malaria infections on the anopheline olfactory system. The effect of increasing host seeking and host feeding synergistically support both parasite and mosquito survivorship as well as maximizing vectorial capacity. Interestingly, the presence of polycistronic OR genes in \textit{Anopheles}\textsuperscript{32} raises the hypothesis that \textit{Plasmodium} parasites may have evolved to specifically activate the transcription factors regulating polycistronic ORs to efficiently induce large-scale olfactory modulation. Indeed, all of the AgORs identified in polycistronic cluster 1\textsuperscript{B}\textsuperscript{32} are present in our sequencing libraries, and one of these AgORs\textsuperscript{17} was significantly over-represented in malarial mosquitoes; the increased abundance of the remaining five AgORs (13, 15, 16, 47, 55) are just below statistical significance (Fig. 5A).

**Odorant binding proteins/chemosensory proteins.** Odorant binding proteins (OBPs), chemosensory proteins (CSPs) and sensory appendage proteins (SAPs) have been postulated to mediate the solubility and transport of odorant molecules within the sensilla lymph\textsuperscript{33}. A total of 57 AgOBPs/AgCSPs/AgSAPs were identified in sequencing datasets, with 47 AgOBPs/AgCSPs more abundant in heads of infected mosquitoes (17 significantly) while two AgOBPs (9 and 25) and a single AgCSP (5) were significantly under-represented relative to uninfected controls (Fig. 5B). These findings are consistent with microarray-based transcriptome studies in \textit{Ae. aegypti} that determined dengue viral infection to increase abundance of a subset of OBPs that was correlated to increased host feeding and vectorial capacity\textsuperscript{34}. It is reasonable to postulate that \textit{Plasmodium} infection similarly impacts anopheine mosquitoes to potentially increase host feeding, synergistically supporting parasite and mosquito survivorship as well as maximizing vectorial capacity\textsuperscript{34}. It is reasonable to postulate that \textit{Plasmodium} infection similarly impacts anopheine mosquitoes to potentially increase host feeding, synergistically supporting parasite and mosquito survivorship as well as maximizing vectorial capacity. AgCSP5 (−0.72-fold; \(p\)-adj = 0.014; AGAP029127), AgOBP9 (−0.54-fold; \(p\)-adj = 0.030; AGAP000278), and AgOBP25 (−0.40-fold; \(p\)-adj = 0.018; AGAP012320) are the only significantly under-represented OBP/CSP transcripts (Fig. 5B). In any case, the precise role of AgCSP5, AgOBP9, and AgOBP25 along with those of other OBP/CSP genes in \textit{An. gambiae} olfaction remains enigmatic.

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**Figure 5.** Chemosensory differential transcript abundances following relevant sporozoite infections. Chemosensory transcripts that were present at significantly higher (blue) or lower (yellow) levels in \textit{Plasmodium falciparum} sporozoite infected treatments; non-differentially expressed chemosensory transcripts are denoted as zeros (black). Chemosensory genes within each family organized by adjusted \(p\)-value and subsequently arrayed left to right from most abundant to least abundant based on FPKM values (quartile bars above each image). (A) Odorant receptor family (OR). (B) Odorant binding protein family (OBP). Chemosensory co-receptors are identified by red boxes. Log\(_2\) scale indicates transcript abundances that were significantly higher (blue) or lower (yellow) in sporozoite infected treatments (Trt) vs. naïve controls (Cnt).
Figure 6. Calculated changes in *Anopheles gambiae* odorant receptor (AgOR) mediated odorant receptivity following relevant sporozoite infections. Graphic representation of conceptualized differences in odorant receptivity for 70 odors in *Plasmodium falciparum* sporozoite infected treatments (Trt) vs. naïve controls (Cnt). Response characteristics were determined using known *A. gambiae* odorant receptor responses in heterologous expression systems and weighted by relative AgOR expression levels for each corresponding receptor. Results are sorted high to low. Scale bar show calculated increases (blue).

Gustatory receptors. Phylogenetic-based annotation along with appendage-specific transcriptome profiling of *A. coluzzii*, *A. gambiae*, and *Ae. aegypti* gustatory receptors (GRs) correlate several orthologous GR genes with the gustatory responses of putative gustatory receptor neurons (GRNs) on the proboscis and labelum. That said, there is a paucity of direct functional data for the majority of *A. gambiae* gustatory receptors (*AgGRs*)\(^{23,37,38}\). In contrast, *AgGR22* (AGAP00999), *AgGR23* (AGAP003098), and *AgGR24* (AGAP001915), which make up a complex that is required for volatile carbon dioxide (CO\(_2\)) detection critical for blood-feeding behaviors, have been well characterized\(^{39}\). The differential transcriptome profiles for *AgGR22* (0.28-fold), *AgGR23* (0.31-fold), and *AgGR24* (0.34-fold) are only slightly more abundant in sporozoite-infected head transcriptomes than in controls (Fig. 7A, red bar) and, not surprisingly, these changes fail to meet the rigorous significance thresholds employed here. This suggests that *Plasmodium* infection does not significantly impact CO\(_2\) sensitivity and is consistent with previous electrophysiology-based studies that showed *P. berghei* infections had no effect on *A. stephensi* maxillary palp neuron sensitivity to CO\(_2\)^{40}. Taken together, it appears likely that *P. falciparum* parasites do not manipulate early, long-range anopheline host-seeking behaviors, such as CO\(_2\) detection\(^{41}\).

Beyond the CO\(_2\) receptors, we observed a considerable shift in the AgGR transcriptome profile between heads of uninfected and those of sporozoite-infected mosquitoes. A total of 44 AgGRs were differentially detected; transcripts for 40 AgGRs showed a higher abundance and the remaining 4 were less abundant in heads of sporozoite-infected mosquitoes than in controls (Fig. 7A). Of these differentially abundant AgGRs, 13 were present at significantly higher levels and none at significantly lower levels in treatments. While the most significantly
abundant, AgGR11 (3.43-fold), and seven other AgGRs that are significantly more abundant in treatments remain unannotated, several AgGRs with homology to functionally characterized sugar-responding GRs from *Drosophila* display significantly higher transcript abundances than controls, suggesting a higher sensitivity to sugary substrates in treatments. These include the DmGR5a orthologs AgGR14 (1.94-fold; p-adj = < 0.001; AGAP006399) and AgGR21 (0.88-fold; p-adj = 0.05; AGAP003260); the DmGR43a ortholog AgGR25 (0.83-fold; p-adj = 0.05; AGAP004727); and the DmGR68a ortholog AgGR9 (2.90-fold; p-adj = < 0.0001; AGAP009805; Fig. 7A). It is reasonable to hypothesize that because glucose is the primary energy source for both mosquitoes and the *Plasmodium* sporozoites, it would be advantageous to manipulate its acquisition, uptake and transport, thereby synergistically supporting sporozoite and mosquito health, survival and by extension, malaria transmission. This differential increase in putative sugar-responsive AgGRs as a consequence of salivary gland *P. falciparum* infections also correlated with a significant increase in the abundance of several sugar transporters (Supplementary Information). While this is the first identification of a *P. falciparum*-mediated increase in abundance

**Figure 7.** Chemosensory differential transcript abundances following relevant sporozoite infections. Chemosensory transcripts that were present at significantly higher (blue) or lower (yellow) levels in *Plasmodium falciparum* sporozoite infected treatments; non-differentially expressed chemosensory transcripts are denoted as zeros (black). Chemosensory genes within each family organized by adjusted p-value and subsequently arrayed left to right from most abundant to least abundant based on FPKM values (quartile bars above each image). (A) Gustatory receptor family (GR). (B) Ionotropic receptor family (IR). (C) Transient receptor potential family (TRP). (D) Pickpocket protein family (PPK). (E) Sensory neuron membrane protein family (SNMP). Chemosensory co-receptors are identified by red boxes. The three CO₂ gustatory receptors are identified by a red line. Log, scale indicates transcript abundances that were significantly higher (blue) or lower (yellow) in sporozoite infected treatments (Trt) vs. naïve controls (Cnt).
of non-CO₂ responsive AgGRs, our data are consistent with previous studies highlighting the overabundance of sugar transporters in the salivary glands of Plasmodium-infected An. gambiae and their importance for sporozoite survival⁵⁵. The remaining transcript, AgGR47 (2.60-fold; p-adj = 0.05; AGAP005514), found significantly abundant in treatments is a DmGR66a ortholog and putative functions in bitter/caffeine detection⁴⁸. It is reasonable to speculate that Plasmodium sporozoites in Anopheles mosquitoes upregulate the deterrent effects of such AgGR neuronal responses to prevent ingestion of detrimental bitter nutrient sources that have been shown to temporally inhibit mosquito sugar and water neurons and reduce nectar feeding of An. quadrimaculatus⁴⁶.

Shifting mosquito gustatory profiles to increase sensitivity to bitter and behaviorally aversive semiochemicals therefore increases the likelihood of blood-meal (and or nectar) feeding and consequently pathogen transmission. It is likely that this effect contributes to increased sensitivity and the marked decrease in mortality rate of An. gambiae harboring P. falciparum sporozoites when exposed to DEET (N, N-diethyl-3-méthylbenzamide), a widely used repellent, in comparison with uninfected An. gambiae and other pyrethroid test substrates⁴⁴,⁴⁵. Lastly, the increased sensitivity of AgGR47 observed in sporozoite-infected An. gambiae relative to naïve controls may reflect age-related nutrient deficiencies in controls. In aging Drosophila, nutrient deprivation dramatically alters feeding behaviors, including the depotentiation of responses to bitter⁴⁶.

Iontropic receptors. Iontropic receptors (IRs) are an ancient family of chemosensory receptors evolutionarily derived from ionotropic glutamate receptors⁴⁹ that are associated with olfactory responses to amines and carboxylic acids that have been identified as important semiochemicals closely associated with host-preference and seeking⁵⁰. As is true in other insects, AgIR8a, AgIR25a and AgIR76b likely function as co-receptors along with odorant-recognition AgIRs that ‘tune’ responses to host cues⁵¹,⁵². In An. gambiae, AgIR25a together with AgIR76b and AgIR8a function in the detection of amines and carboxylic acids, respectively⁵³ both of these chemical classes have been directly implicated in anopheline host seeking⁵⁴,⁵⁵. Interestingly, the abundance of AgIR76b transcripts was significantly enhanced (0.53-fold; p-adj = 0.05; AGAP011968) in infected mosquito heads while the levels of AgIR25a and AgIR8a were not significantly different (Fig. 7B). Transcripts for 38 of the 49 odorant-tuning AgIRs detected here were found to be more abundant in sporozoite-infected head transcriptomes; only 11 were downregulated compared with uninfected controls (Fig. 7B). Of those upregulated, 18 were significantly more abundant but none were significantly less abundant in treatment groups than controls. Inasmuch as many AgIRs are uncharacterized, including those with the most pronounced and significant differential abundance (AgIR149, AgIR168, and AgIR203), the implications of these differentially up-/downregulated IRs are difficult to assess. Nevertheless, functional information for a small subset of AgIRs is available and others can be derived from analyses of clear orthologs in Ae. aegypti, An. gambiae, An. sinensis and Drosophila melanogaster. This allows us to identify several AgIRs significantly more abundant in treatment groups (Fig. 7B) to be associated with both gustation (AgIR7x (3.11-fold; p-adj = 0.03; AGAP013520), AgIR93a (0.52-fold; p-adj = 0.05; AGAP000256), AgIR60a (2.48-fold; p-adj = 0.05; AGAP011943) and host-seeking behaviors (AgIR75g (1.96-fold; p-adj = 0.03; AGAP013085))⁵⁶. In addition, we observed three receptors localized to the antenna (AgIR31a (1.43-fold; p-adj = 0.03; AGAP009014), AgIR68a (2.00-fold; p-adj = 0.003; AGAP007951), and AgIR411 (1.44-fold; p-adj = 0.03; AGAP004432)⁵⁷. While persistence of host seeking and blood feeding might well be advantageous for Plasmodium, it would only be beneficial for their anopheline hosts if these behavioral shifts promoted the acquisition of a successful blood meal.

Transient receptor potential channels. As is true for all insects, An. gambiae transient receptor potential (AgTRP) family members are associated with a variety of sensory modalities such as chemosensation, gustation, thermosensation, hygrosensation, mechanosensation, vision, and intracellular signaling⁵⁸. A total of nine AgTRPs were identified in sequencing libraries, five of which are more abundant (three significantly) in heads of sporozoite-infected mosquitoes while four AgTRPs (three significantly) are under-represented relative to uninfected controls (Fig. 7C). Of these, AgTRP (canonical) is 0.70-fold; p-adj = 0.0002; AGAP000349), AgTRP (canonical; 0.67-fold; p-adj = 0.0004; AGAP008435), and AgTRPM (melastatin; 0.75-fold; p-adj = 0.01; AGAP006825) are all active in thermosensation and feeding. TRPc has been identified as having several physiological roles in insects, including signal transduction, phototransduction, and gustatory sensitivity to CO₂. TRPM mediates Drosophila nociception behaviors and putatively critical behaviors for An. gambiae viability and survival⁵⁹. Given the critical roles that AgTRPC and AgTRPM are likely to play in mosquito host seeking and blood feeding, manipulation of these transcripts would likely be advantageous for Plasmodium to ensure maximum host contact rate and transmission. TRP-Painless and TRP-Pyrexia function in thermosensation and have been documented to mediate avoidance of noxious high temperatures in D. melanogaster⁶⁰. It is possible the significantly reduced abundance of AgTRP-Painless (− 0.78-fold; p-adj = 0.01; AGAP001243) and AgTRP-Pyrexia (− 0.75-fold; p-adj = 0.0003; AGAP010269) in treatment groups compared with naïve controls is related to a loss of nociception.

The remaining An. gambiae TRP identified as significantly differentially abundant between treatments and controls functions as a mechanoreceptor with roles in Drosophila hygrosensation. AgTRP-Waterwitch which is significantly less abundant in treatment tissues (− 0.70-fold; p-adj = 0.001; AGAP000361) is homologous to the DmTRP-Waterwitch that functions in dry-air detection⁶¹. Moisture detection, or importantly the lack thereof, mediates flight and host-seeking behavior in anopheline mosquitoes such that augmenting mosquito traps with moist air significantly increases flight activity and attraction of wild Aedes, Anopheles and Culex mosquitoes⁶². It is likely that upregulation of chemosensory TRPs as well as the inverse regulation of Ag-Waterwitch optimizes persistent host seeking and blood feeding and is consistent with studies showing that Anopheles mosquitoes harboring Plasmodium sporozoites exhibit more persistent host-seeking behavior than uninfected controls⁶³.
Pickpocket channel proteins/sensory neuron membrane proteins. We identified 18 *An. gambiae* pickpocket channel proteins (AgPPKs) of which transcripts were predominately more abundant in heads of sporozoite-infected mosquitoes (Fig. 7D). Two of these AgPPKs, AgPPK13-like (0.97-fold; p-adj = 0.004; AGAP007945) and AgPPK31-like (1.47-fold; p-adj = 0.05; AGAP000657), showed significant enrichment. In *D. melanogaster*, PPKs have been implicated to function in salt and water taste, although the behavioral implications of these characterized PPKs is unclear. Similarly, while aedine PPKs are hypothesized to function in a variety of sensory modalities, such as chemosensation, hygrosensation and mechanosensation, anopheline PPKs remain largely uncharacterized and the implications of these differentially abundant PPKs remain unclear. Sensory neuron membrane proteins (SNMPs) are both colocalized with ORs and broadly expressed throughout *D. melanogaster* and are postulated to have both chemosensory and non-chemosensory functions with particular roles in dipteran and lepidopteran pheromone detection. Both AgSNMP1 and AgSNMP were identified in sequencing libraries, although without significance (Fig. 7E). These and other anopheline SNMPs remain uncharacterized, and the implications of differentially abundant AgSNMPs are therefore unclear.

General neuronal function. Insect neurotransmitters comprise acetylcholine (ACh), gamma-aminobutyric acid (GABA), glutamate (Glu), and select biogenic amines such as dopamine, histamine, octopamine, serotonin and tyramine as well as their respective membrane receptors. In total, 11 subunits for nicotinic ACh receptors (nAChr) and muscarinic ACh receptor subunits (mAChr) were identified with 5 Ag-nAChr and 1 Ag-mAChr significantly overabundant in the heads of sporozoite-infected *An. gambiae*: Ag-mAChr2 (0.75-fold; p-adj = 0.001; AGAP004675), Ag-nAChr2 (0.96-fold; p-adj = < 0.0001; AGAP002972), Ag-nAChr3 (1.18-fold; p-adj = < 0.0001; AGAP003229), Ag-nAChr5 (1.25-fold; p-adj = < 0.0001; AGAP000588), Ag-nAChr6 (1.731-fold; p-adj = < 0.0001; AGAP002152), and Ag-nAChrβ1 (0.90-fold; p-adj = < 0.0001; AGAP000966; Fig. 8A). Only Ag-nAChr9 (−1.23-fold; p-adj = < 0.0001; AGAP009493) was significantly diminished in heads of sporozoite-infected mosquitoes compared with naïve mosquitoes (Fig. 8A). While insect and especially mosquito AChRs (nicotinic and muscarinic) are not well characterized, it has been reported that with age, AChr responses become significantly diminished as a function of oxidative damage and neurodegeneration. In *Drosophila* this is associated with aged-related immune system dysfunction. As such, the increased abundance of AgAChrs in sporozoite-infected *An. gambiae* suggests that malaria infected anopheline mosquitoes are resistant.

Figure 8. Neurotransmitter differential transcript abundances following relevant sporozoite infections. Neurotransmitter transcripts that were present at significantly higher (blue) or lower (yellow) levels in *Plasmodium falciparum* sporozoite infected treatments; non-differentially expressed neurotransmitter transcripts are denoted as zeros (black). Neurotransmitter genes within each family organized by adjusted p-value and subsequently arrayed left to right from most abundant to least abundant based on FPKM values (quartile bars above each image). (A) Acetylcholine receptor family. (B) Gamma aminobutyric acid receptor family. (C) Glutamate receptor family. (D) Histamine receptor family. (E) Serotonin receptor family. (F) Octopamine receptor family. (G) Dopamine receptor family. Log2 scale indicates transcript abundances that were significantly higher (blue) or lower (yellow) in sporozoite infected treatments (Trt) vs. naïve controls (Cnt).
to both aged-related immune system dysfunction and neurodegeneration. A similar logic could apply to the significant overabundance of acetylcholinesterase (Ace; AgAce, 1.18-fold; p-adj = < 0.0001; AGAP000466) identified in treated An. gambiae relative to naïve controls (Fig. 9A). As such, the increased levels of AChrs in treatment groups could also represent altered baseline levels within mosquitoes that as a result are not yet susceptible to the effects of aging and senescence.

Transcripts for both ionotropic (GABA-A) receptors and metabotropic (GABA-B) receptors were significantly more abundant in treatment groups when compared with naïve controls: AgGABA-A (0.80-fold; p-adj = 0.0007; AGAP006028), AgGABA-B (0.38-fold; p-adj = 0.02; AGAP000038), and AgGABA-B2 (1.14-fold; p-adj = < 0.0001; AGAP004595; Fig. 8B). Transcripts for the GABA processing enzyme glutamate decarboxylase (Gad) displayed a dissimilar differentiation as the abundance of transcripts encoding AgGad1 (− 2.92-fold; p-adj = < 0.0001; AGAP005866) and AgGad2 (− 1.10-fold; p-adj = 0.0001; AGAP008904) were significantly diminished in treatment groups compared with controls (Fig. 9A). In Drosophila, GABA mechanisms help stabilize neuronal function and maintain tissue homeostasis. As such, the decreased abundance of AgGABAs and increased abundance of AgGads in naïve controls may actually reduce aging-related neurodegeneration to help maintain neuronal homeostasis.

Similarly, transcripts for NMDA and non-NMDA types of anopheline glutamate receptors were identified in sequencing libraries and predominately under-represented in heads of infected An. gambiae. Of the eight

**Figure 9.** Neuropeptide differential transcript abundances following relevant sporozoite infections.

Neuropeptide transcripts that were present at significantly higher (blue) or lower (yellow) levels in Plasmodium falciparum sporozoite infected treatments; non-differentially expressed neuropeptide transcripts are denoted as zeros (black). Neuropeptide genes within each family organized by adjusted p-value and subsequently arrayed left to right from most abundant to least abundant based on FPKM values (quartile bars above each image). (A) Neurotransmitter processing enzyme family. (B) Neuropeptide family. (C) Neuropeptide receptor family. (D) Neuron transporter family. Log2 scale indicates transcript abundances that were significantly higher (blue) or lower (yellow) in sporozoite infected treatments (Trt) vs. naïve controls (Cnt).
glutamate receptor transcripts identified, none was significantly abundant in heads of sporozoite-infected mosquitoes; in contrast, six NMDA and non-NMDA glutamate receptors—AgGluRIa (−1.27-fold; p-adj = < 0.0001; AGAP000803), AgGluRIId (−0.54-fold; p-adj = 0.003; AGAP002797) AgGluRIle (−0.81-fold; p-adj = < 0.0001; AGAP012417), AgNMDAR1 (−1.36-fold; p-adj = < 0.0001; AGAP001478), AgNMDAR2 (−0.52-fold; p-adj = 0.005; AGAP012419) and AgNMDAR3 (−0.98-fold; p-adj = 0.002; AGAP005527)—were significantly diminished in sporozoite-infected mosquitoes compared with naïve controls (Fig. 8C). It is possible that in light of the ability of these receptors to function as immunomodulators of inflammatory responses the over-abundance of NMDA and non-NMDA glutamate receptors in naïve mosquitoes reflects either an aging-related immune system dysfunction or neuronal pathology. Transcripts encoding the receptors for several peptide biogenic amines, including dopamine, histamine, octopamine, serotonin and tyramine, which function as anopeline neuromodulators and neurotransmitters were identified in our libraries. With the exception of two pairs of histamine and octopamine receptor transcripts that were similarly up-/downregulated, most biogenic amine receptors showed no significant differential abundance in heads of sporozoite-infected An. gambiae relative to naïve controls (Fig. 8D–G).

Transcripts encoding several biogenic amine processors and transporters were identified in similarly up-/downregulated pairs with no clear distinction between treatment and naïve controls (Fig. 9A, D). Insect neuropeptides are essential neuromodulators that regulate a variety of physiological and behavioral functions. Several neuropeptide transcripts identified in sequencing libraries were shown to be significantly overabundant in the heads of sporozoite-infected An. gambiae (Fig. 9B). Interestingly, there were also four neuropeptide transcripts and their correlating receptors that were significantly diminished in treatment heads compared with naïve controls, allatostatin 1 (AgAs1; −0.50-fold; p-adj = 0.001; AGAP003712), allatostatin 2 (AgAs2; −0.41-fold; p-adj = 0.01; AGAP010157), FMRFamide (AgFMRF, −1.34-fold; p-adj = < 0.0001; AGAP005518) and sickie (AgSickie; −1.66-fold; p-adj = <0.0001; AGAP009424), that are directly or indirectly correlated with insect stress and immune response (Fig. 9B, C). In Ae. aegypti, allatostatin interacts with hemocytes to increase humoral immune responses to combat cellular stress and inflammation. FMRFamide signaling in Drosophila increases in response to cellular stress to promote stress-induced sleep and cellular recovery to help bolster immune function. Sickie acts directly on Drosophila innate immunity by activating the transcription factor Relish and subsequent transcription of immune defense antimicrobial peptides (AMPs). Neuropeptide F (NPF) also acts indirectly to modulate stress responses through activation of downstream signaling pathways. In dipterans, NPF independently activates insulin signaling pathways and coordinate the target or rapamycin (TOR) pathway. TOR pathway is an insect nutrient-sensing pathway that regulates cellular metabolism and homeostasis in response to nutrient flux and stress. TOR signaling naturally increases with age and increasing cellular dysfunction and stress. With a substantial number of neuropeptides/hormones and their correlating receptors significantly abundant in heads of sporozoite-infected An. gambiae and functioning to maintain healthy mosquito cells and tissues, the decreased abundance of these select four neuropeptides in infected mosquitoes is puzzling. If the ‘normal’ transcriptome profile of naïve mosquitoes reflects changes associated with age-related immune system dysfunction and the loss of cellular homeostasis, or some other age-related pathology, then one could hypothesize that infected mosquitoes might be refractory to those effects.

Immune system. Activation of the immune system in response to infection is energetically costly and likely results in decreased fitness reflected by other life history traits with the potential to impact vectorial capacity. Mosquito anti-Plasmodium immunity is stage specific and involves diverse defense processes, although the extent of immune defense in salivary-gland-stage infected mosquitoes, and especially the head and salivary glands (post sporozoite salivary gland invasion), remains unclear. Since Plasmodium, unlike arboviruses, does not establish infection in the mosquito head, the immunity-related transcriptome changes observed in our study are likely to mostly derive from the combined head (including sensory appendages) and salivary gland transcriptomes. In heads of sporozoite-infected and control sequencing libraries, several components of An. gambiae humoral and cellular, innate immune defenses were identified. An. gambiae Pattern Recognition Receptors (PRRs) include 6 fibrinogen proteins (FBGs), 7 3-glucan binding proteins (3GNBPs), 7 peptidoglycan recognition proteins (PGRPs), 24 leucine-rich repeat immune proteins (LRIMs) including Plasmodium infection responsive leucine-rich proteins, 13 thioester-containing proteins (TEPs), and 2 members of C-type lectin family proteins (CTLs)—AgCTL4 and AgCTLM2 (Fig. 10A, B). A large portion of the PRR transcripts were not present at significant differential abundances between heads of sporozoite-infected and control mosquitoes; instead, neither of the transcripts encoding AgCTLM2 or AgCTL4 displayed significant differential abundance.

Anopeline TEPs are antimicrobial and antiparasitic PRRs complement-like factors that drive pathogen clearance through lysis and melanization. Of those, only AgTEP13 (0.34-fold; p-adj = 0.03; AGAP008407) was significantly more abundant in heads of sporozoite-infected mosquitoes, and AgTEP1 (−1.95; p-adj = < 0.0001; AGAP010815) and AgTEP12 (−0.94; p-adj = 0.011; AGAP008654) significantly diminished in treatment groups compared with controls (Fig. 10A). While the roles of TEP12 and TEP13 in anopheline immune responses are not currently known, the well-studied TEP1 functions as an enhancer of mosquito phagocytosis. Since TEP1 increases, the increased abundance of AgTEP1 in naïve controls likely reflects aging-related immune system dysfunction as aging decreases phagocytic efficiency and contributes to immunosenescence. LRIMs and APLs are PRRs that mediate innate immune surveillance and response. Transcripts for 20 LRIMs and APLs were downregulated in heads of infected mosquitoes relative to naïve controls, while 4 of the 24 An. gambiae LRIMs were more abundant in treatments. Of the 4 AgLRIMs upregulated, 2 were significantly more abundant, whereas 9 of the 20 LRIMs and APLs were significantly less abundant in treatments than controls (Fig. 10B). These data are largely consistent with previous studies on the transcriptional regulation of LRIMs.
Figure 10. Immune response differential transcript abundances following relevant sporozoite infections. Immune response transcripts that were present at significantly higher (blue) or lower (yellow) levels in *Plasmodium falciparum* sporozoite infected treatments; non-differentially expressed immune response transcripts are denoted as zeros (black). Immune response genes within each group organized by adjusted p-value and subsequently arrayed left to right from most abundant to least abundant based on FPKM values (quartile bars above each image). (A, B) Pattern recognition proteins. (C) Modulator proteins. (D, E) Effector proteins. Log2 scale indicates transcript abundances that were significantly higher (blue) or lower (yellow) in sporozoite infected treatments (Trt) vs. naïve controls (Cnt).
and APLs in response to *Plasmodium*, gram-negative/gram-positive bacteria, and fungal challenge. While AgLRIM12 (0.72-fold) and AgLRIM5 (0.82-fold; ACOM043147) were both significantly more abundant in heads of infected mosquitoes they are not yet conclusively associated with *Plasmodium* infection; it is therefore possible the observed differential abundance of AgLRIM reflect non-immunity responses or potential aging effects that are only partially impacted by *P. falciparum* infection.

Similarly identified AgFBGs, which function in phagocytosis and complement activation, were predominately diminished in treatments relative to naive controls, with AgFBG (− 1.47-fold; p-adj = 0.015; AGAP011231), AgFBG (− 1.98-fold; p-adj = 0.004; AGAP010869), AgFBG (− 1.29-fold; p-adj = 0.02; AGAP004917) and AgFBG (− 1.29-fold; p-adj = 0.007; AGAP006914) significantly reduced (Fig. 10B). In *Drosophila*, aging is associated with increased microbial loads and dysbiosis that trigger increased expression of immune genes and stronger immune activation. Excessive immune activation ultimately results in immune dysfunction and decreased lifespan. In that context, the large number of AgLRIMs, AgAPs and AgFBGs that are more abundant in naive control transcriptomes may reflect aging-related dysfunction of the *An. gambiae* immune system. Accelerated immune system dysfunction in naïve controls also rationalizes the observed AgPGRP transcript abundances identified in replicate sequencing libraries. Our data indicate that transcripts for two AgPGRPs that have been characterized as negative regulators of immune IMD signaling defense, AgPGRP-LB (1.46-fold; p-adj = 0.02; AGAP001212) and AgPGRP-LD (0.69-fold; p-adj = 0.05; AGAP005552: Fig. 10A), are significantly more abundant in heads of sporozoite-infected mosquitoes, suggesting a suppression of the IMD immune pathway. The IMD signaling initiation receptor AgPGRP-LC (− 0.25-fold; p-adj = 0.02; AGAP005203), identified as significantly reduced in treatments compared with controls, further supports the presence of diminished IMD signaling in sporozoite-infected *An. gambiae*. PGRP immune regulator deficiency results in unregulated IMD signaling causing dysplasia, inflammation, apoptosis, and decreased lifespan. Preservation of immune homeostasis would be advantageous for both *Anopheles* hosts and *Plasmodium* parasites to maintain vectorial capacity. It is possible that *An. gambiae* challenged with biologically relevant, low-level *P. falciparum* results in a mild and prolonged engagement of the mosquito immune system that ultimately primes the system for protection from secondary infection and importantly, the negative impact of aging-related immune system degeneration. Indeed, *Plasmodium*-induced anophele immunological priming may represent an advantageous selective adaptive pressure that maximizes mosquito host fitness and, consequently, vectorial capacity.

Serine proteases constitute the majority of mosquito immune system moderators that are activated in response to pathogen recognition. While there are hundreds of serine proteases functioning in a variety of mosquito molecular processes, eight CLIP-domain serine proteases (CLIPs)—SPCLIP1, CLIPA2, CLIPA7, CLIPA8, CLIPA14, CLIPA28, CLIPB14, and CLIPB15—mediate *P. berghei* melanization. No AgCLIP transcripts were significantly more abundant in heads of sporozoite-infected mosquitoes, although eight were less abundant in treatment groups for which mRNAs of AgCLIPA7 (− 1.34-fold; p-adj < 0.0001; AGAP011792), AgCLIPA14 (− 1.2-fold; p-adj < 0.0001; AGAP011788) and AgCLIPA28 (− 0.92-fold; p-adj = 0.02; AGAP010968) were significantly under-represented relative to uninfected controls (Fig. 10C). These data were expected because anophele CLIPs predominately moderate melanization of rodent *Plasmodium* ookinetes and are not associated with immune defenses targeting sporozoites or human *Plasmodium*. Indeed, the significant abundance of AgCLIPs in uninfected middle-aged *An. gambiae* may result from aging-related immune dysfunction. The absence of elevated levels of AgCLIPs highlights another putative benefit of *Plasmodium*-mediated immune priming. Another way to moderate immune effector mechanisms such as melanization is performed by Serpins (serine protease inhibitors). Levels of two Serpins, AgSerpin6 (− 0.64-fold; p-adj = < 0.0001; AGAP00212) and AgSerpin10 (− 0.42; p-adj = 0.03; AGAP005246), that are upregulated in response to *Plasmodium* ookinetes, were significantly under-represented compared with uninfected controls (Fig. 10C). As melanization is not expected to be active in *An. gambiae* carrying salivary-gland-stage *P. falciparum* infections, there would be no requirement for Serpin modulation, and such genes as Serpin6 and Serpin10 would no longer be required at this timepoint. Likewise, putative immune dysfunction in uninfected controls may impair proper regulation of melanization and, as such, Serpin transcription.

A wide range of immune effectors are downstream of PRR-controlled pathways and other recognition pathways that include anti-microbial peptides (AMPs), reactive oxygen species (ROS), reactive nitrogen species (RNS), including prophenoloxidase (PPOs), and apoptosis- and phagocytosis-related genes. In *Anopheles*, five classes of AMPs including attacins, defensins, diptericins, cecropins, and gambicins have been described. In our data set, three defensins (Def), three cecropins (Cec), one gambicin (Gam) and one attacin (Att) are present (Fig. 10E). Of these, only two defensins and Cec1 were significantly differentially abundant between heads of sporozoite-infected and uninfected control mosquitoes, with AgDef2 (0.96-fold; p-adj = < 0.0001; AGAP004632), AgDef4 (1.96-fold; p-adj = 0.005; AGAP005416) and AgCec1 (1.78-fold; p-adj = 0.001; AGAP006913) all significantly abundant in treatment groups compared with controls. Interestingly, these shifts plus the significantly increased abundance of AgGNBP2 (1.46-fold; p-adj = 0.02; AGAP001212; Fig. 10A), a PRR that co-localizes with *P. falciparum*, are the only other indication of an *An. gambiae* immune response to *P. falciparum* infection in this study. Of these, only AgCec1 is associated with anti-sporozoite activity. AgDef1 and AgDef4 demonstrate antibacterial activity but have not shown any anti-*Plasmodium* activity. Interestingly, studies in *Ae. aegypti* have determined that differential Def abundances can be observed without immune challenge, and often after pathogen clearance, suggesting alternative functions for the AMPs. As such, the role of AgDef in response to sporozoite infections remains unclear. ROS and RNS are vital effectors of anophele immune defenses active against malaria parasites. ROS formation in dipterans is postulated to involve NADPH oxidase (NOX)-based production of superoxide anions that are transformed into H₂O₂ by superoxide dismutase (SOD) and subsequent production of ROS. NOX is also likely required for heme peroxidase (HPX) and nitric oxide synthase (NOS) activation, both of which potentiate nitration and generations of RNS. Interestingly, the transcripts for all the enzymes associated with ROS/RNS system and their potentiators were largely under-represented in sporozoite-infected
head transcriptomes compared with uninfected controls: AgMnSOD1 (−0.88-fold; p-adj = 0.001; AGAP010517), AgCuSOD2 (−0.38-fold; p-adj = 0.006; AGAP005234), AgHPX2 (−1.20-fold; p-adj = 0.0001; AGAP009033) and AgNOX-DUOX (−0.63-fold; p-adj = 0.03; AGAP009978). In that light, it is logical to expect the significant under-representation of transcripts encoding catalase, glutathione peroxidase and thioredoxin peroxidase detoxifying enzymes observed in heads of sporozoite-infected An. gambiae when compared with uninfected controls (Fig. 10E).

In Anopheles and other mosquitoes, CLIPs activate PPOs, the primary effectors mediating melanization. In that light, the lack of differentially abundant AgCLIP transcripts aligns with the observation that AgPPGs were predominately under-represented in heads of sporozoite-infected mosquitoes compared with uninfected controls. AgPPO2 (−1.70-fold; p-adj = 0.001; AGAP006258), AgPPO5 (−1.30-fold; p-adj = < 0.0001; AGAM012616), AgPPO6 (−1.38-fold; p-adj = 0.002; AGAP004977), and AgPPO9 (−1.50-fold; p-adj = 0.006; AGAP004978) all significantly diminished in An. gambiae carrying salivary-gland-stage sporozoites (Fig. 10). The higher PPO levels in heads of uninfected mosquitoes is surprising and may reflect a higher background of metabolic stress. However, this pattern is consistent with the fact that melanized sporozoite-stage Plasmodium has never been observed. In Ae. aegypti, allatostatin neuropeptide interacts with hemocytes and may have subsequently increased AgPPO expression to address dysbiosis and inflammation resulting from IMD destabilization. This finding is particularly interesting because it correlates mosquito immune system fitness with neuropeptide production and has significantly broader implications for aging-related immune system dysfunction. It also provides a plethora of anaphelne molecular mechanisms that putatively synergize with Plasmodium immune priming, thereby providing a significant advantage that ultimately facilitates malaria transmission.

Apoptosis, autophagy and phagocytosis genes are important effectors in response to viral and pathogen invasion through programmed cell death of infected cells and clearance of cellular debris and pathogen particles. In dipterans, apoptosis is induced through the activation of the long caspases. Dronc and Dredd, and inhibition of inhibitors of apoptosis (IAPs) that in turn regulate autophagy genes, with expression of both increasing concurrently. Transcripts for AgIAP, AgDronc, AgDredd, 18 autophagy-related proteins (AgApgs), and 2 phagocytosis signaling proteins were identified (Fig. 10D). AgDronc, AgDredd, and AgIAP were all under-represented in heads of sporozoite-infected mosquitoes when compared with controls, although none significantly (Fig. 10D). AgApgs were predominately under-represented in treatment groups compared with controls, with only AgApg8 (0.20-fold; p-adj = 0.03; AGAP002685) and AgApg18B (0.59-fold; p-adj = 0.003; AGAP005910) significantly abundant in treatment groups, and 8 AgApgs significantly diminished in treatment groups (Fig. 10D): AgApg2 (−0.57-fold; p-adj = 0.004; AGAP004092), AgApg3 (−0.69-fold; p-adj = < 0.0001; AGAP011582), AgApg6 (−0.37-fold; p-adj = 0.02; AGAP003858), AgApg7 (−0.49-fold; p-adj = 0.001; AGAP010303), AgApg7A (−1.14-fold; p-adj = < 0.0001; AGAP008637), AgApg13 (−1.48-fold; p-adj = < 0.0001; AGAP005715), AgApg16L (−0.67-fold; p-adj = 0.007; AGAP002315), and AgApg18A (−2.01-fold; p-adj = 0.03; AGAP002685) and AgApg18B (0.59-fold; p-adj = 0.003; AGAP005910) significantly abundant in treatment groups, and 8 AgApgs significantly diminished in treatment groups (Fig. 10D). As is the case for Toll-mediated signaling, the anopheline IMD immune signaling pathway involves binding of the PRR, PGRP-LC, which triggers activation of NK-kb transcription factor (Rel2) and production of AMPs. Transcripts for all components of IMD signaling, including PGRP-LC, were identified in sequencing libraries, although AgPGRP-LC (−0.26-fold; p-adj = 0.02; AGAP005203) was the sole transcript significantly diminished in heads of sporozoite-infected mosquitoes (Figs. 10A, 11B). Even the modest differential abundance of AgPGRP-LC, which is a positive regulator of IMD signaling and PGRP expression, provides a rationale for the lack of significantly differential IMD signaling pathway transcripts in sporozoite-infected mosquitoes (Fig. 11B). In any case, the destabilization of anopheline IMD signaling pathway components would putatively render the immune signaling system a poor target for Plasmodium immunological priming, as the long-lasting immune response would be unpredictable and difficult to adapt.

The anopheline JAK-STAT signaling pathway has been implicated to function in a variety of dipteran developmental processes, as well as in antimicrobial, antiparasitic and antiviral immune defenses. It is negatively regulated by PIAS and suppressor of cytokine signaling (SOCS) and induced by the binding of Unpaired (Upd) to Domeless (Dome) to form a receptor dimer that, in turn, activates Janus Kinase (JAK) or its homolog Hopscotch (Hop) and ultimately STAT which triggers the production of genes regulated by the JAK-STAT pathway and...
NOS72. Transcripts for AgSOC5 (0.58-fold; p-adj = 0.008; AGAP029624) and AgUPD3 (2.39-fold; p-adj = 0.007; AGAP013506) were significantly more abundant in treatment groups, and AgDome (−0.33-fold; p-adj = 0.006; AGAP029053), AgHop (−1.26-fold; p-adj = 0.003; AGAP008354; JAK homolog) and AgSTAT2 (−0.31-fold; p-adj = 0.02; AGAP000099) were significantly under-represented compared with uninfected controls (Fig. 11B).

As is the case for IMD pathways, anopheline JAK-STAT signaling primarily targets pre-sporozoite stages of *Plasmodium*, making the lack of JAK-STAT activation in the salivary-gland-stage infected *An. gambiae* examined here expected80. This is particularly evident by the significant abundance of JAK-STAT's primary negative regulator AgSOC in heads of sporozoite-infected mosquitoes. Dipteran SOCs deactivate the Hop-Dome dimer, block STAT activation, and target components of the Hop-Dome complex for degradation98. As such, the increased abundance of AgUPD3 in treatment groups may reflect replenishment activity to restore UPD3 levels following *P. falciparum* midgut invasion or simply maintenance of normal UPD3 equilibrium. In *Drosophila*, immune system deregulation and mitochondrial degeneration significantly increase JAK-STAT expression in response to growing systemic inflammation and oxidative stress associated with aging. Excessive JAK-STAT activation ultimately leads to a loss of tissue homeostasis and cell death77. The active inhibition of JAK-STAT signaling in infected mosquitoes would explain the significantly diminished abundance of AgSTAT2 and AgHop and may also reflect aging-related dysfunction in uninfected controls. As such, the anopheline JAK-STAT signaling pathway may represent a target for *Plasmodium* priming, reducing the detrimental effects of aging-related immune deregulation thereby synergistically benefiting both parasite and host.

**Figure 11.** Immune response differential transcript abundances following relevant sporozoite infections. Immune response transcripts that were present at significantly higher (blue) or lower (yellow) levels in *Plasmodium falciparum* sporozoite infected treatments; non-differentially expressed immune response transcripts are denoted as zeros (black). Immune response genes within each group organized by adjusted p-value and subsequently arrayed left to right from most abundant to least abundant based on FPKM values (quartile bars above each image). (A) Toll and spatzle proteins (SPZ). (B) IMD and JAK-STAT proteins. Log2 scale indicates transcript abundances that were significantly higher (blue) or lower (yellow) in sporozoite infected treatments (Trt) vs. naive controls (Cnt).
p-adj = < 0.0001; AGAP007873) and AgTORC1 (−0.51-fold; p-adj = 0.003; AGAP010035)—are significantly diminished in heads of sporozoite-infected An. gambiae compared with naïve controls (Fig. 12A). Additionally, a transcript for the TOR kinase inhibitor tuberous sclerosis complex 2 (AgTSC2; −0.46-fold; p-adj = < 0.001; AGAP003445), which in Drosophila inhibits TOR signaling and acts on fat bodies as a mechanism of lifespan expansion, was found to be significantly diminished in treatment groups compared with controls (Fig. 12A)99.

The AMP-activated kinase (AMPK) signaling pathway maintains insect energy homeostasis by regulating ATP consumption and production; it is activated in response to low intracellular AMP levels, prompting ATP re-accumulation by reducing costly anaerobic processes as well as activating phosphofructokinase (Pfk), thereby increasing glycogenolysis100. Interestingly, the transcripts for both AgAMPK (−0.75-fold; p-adj = < 0.0001; AGAP002686) and AgPfk (−0.33-fold; p-adj = 0.05; AGAP007642) were significantly diminished in heads of sporozoite-infected An. gambiae compared with non-infected mosquitoes (Fig. 12B). In contrast, AMPK over-abundance in An. stephensi is associated with increased innate immune responses and significant alterations in mitochondrial biogenesis101. In aging D. melanogaster, increased levels of AMPK improve tissue homeostasis via an anti-inflammatory mechanism to promote longevity102. It is possible that, compared with treatment transcriptomes, the age-related increased immune response observed in heads of naïve An. gambiae results in reduced nutrient and energy homeostasis. This hypothesis is also supported by the significantly diminished abundance of hepatocyte nuclear factor 3 (AgHNF3; −2.17-fold; p-adj = < 0.0001; AGAP001671) and hepatocyte nuclear factor 4 (AgHNF4; −0.98-fold; p-adj = 0.005; AGAP002155) in heads of sporozoite-infected An. gambiae mosquitoes.
compared with naïve controls. In Ac. Aegypti, RNAi depletion of HNF4 was shown to result in significant downregulation of transcripts encoding triacylglycerol catabolism, which is essential for fat body metabolism.

The head transcriptomes of naïve An. gambiae controls provide substantial evidence of a robust endemically sterile inflammatory response as well as decreased mitochondrial function, increased mitochondrial damage and mitophagy. In D. melanogaster, mitochondrial dysfunction is closely associated with decreased mitochondrial transcript levels of genes such as cytochrome oxidase C (COX) and cytochrome B1 (CB). Indeed, in heads of sporozoite-infected An. gambiae, transcripts for COX and CB were present at very high levels, significantly more abundant than in uninfected controls (Fig. 12C). In general, naïve An. gambiae displayed a significant overabundance of transcripts associated with ubiquitination and autophagy, indicating a higher degree of cell dysfunction than in the treatment group (Fig. 12F). For example, PTEN-induced kinase 1 (PINK1), a serine/threonine kinase that specifically targets damaged mitochondria for autophagy, is significantly diminished in Plasmodium-infected An. gambiae compared with non-infectious mosquitoes (− 0.66-fold; p-adj = 0.0004; AGAP004315) (Fig. 12B), which may be an indication of a higher degree of mitochondrial dysfunction in naïve aged mosquitoes.

In insects, the forkhead box O (FOXO) transcription factor signaling pathway is an important moderator of protein quality control, stress resistance, and lifespan extension pathways. FOXO functions with its transcriptional target translation initiation factor, 4E-binding protein (4E-BP), to moderate proteostasis in Drosophila by removing damaged/abnormal proteins. While Ag4E-BP was not significantly different, AgFOXO (− 2.36-fold; p-adj = < 0.0001; AGAP000662) was significantly diminished in heads of sporozoite-infected An. gambiae compared with naïve controls (Fig. 12D). Similarly, the heat-shock transcription factor (AgHsTF; − 1.00-fold; p-adj = < 0.0001; AGAP029908) and numerous subsequently induced heat shock protein transcripts (Hsps) were all significantly diminished in treatment groups compared with naïve control An. gambiae (Fig. 12E). In diterans, protein abnormalities resulting from oxidative stress gradually increase with age and are associated with an increase in HsTF and Hsp production to combat injury from abnormal and malformed proteins. Lastly, consistent with these transcriptome shifts, our analyses have also uncovered several transcripts involved in ubiquitin-mediated proteosome protein degradation pathways, the abundance of which are significantly diminished in sporozoite infected mosquitoes (Fig. 12F). When viewed collectively, it is likely these differentially abundant transcripts are indicative of an accumulation of abnormal proteins with detrimental effects in naïve, uninfected mosquitoes that importantly does not occur in age-matched counterparts that harbor salivary-gland-stage Plasmodium sporozoites after low intensity, biologically relevant blood-meal infections.

Conclusions
This study is one of the first examinations of the effects of biologically relevant Plasmodium sporozoite-stage infections on anopheline mosquitoes where the impact of Plasmodium infection on vector behavior and physiology significantly drives vectorial capacity. To gain insight into this process in as biologically relevant a manner as possible, we have carried out a set of comprehensive comparative analyses of the transcriptome profile of the head (including sensory appendages) and salivary gland tissues of female anophelines that have undergone low-intensity P. falciparum infections representative of those occurring naturally in disease-endemic regions. To our eyes, the significant reduction in prevalence and other experimental challenges associated with these studies are outweighed by the necessity to examine laboratory-based Plasmodium infections under the most biologically relevant conditions possible. As might be expected, these infections have broad effects on transcript abundance (as a partial proxy for differential gene expression) across a large number of gene families. Here, we have focused on those that impact neuronal function and chemosensation as well as age-related immunity and metabolic homeostasis.

The transcriptome profile shifts we have uncovered reveal a variety of systematic impacts that broadly suggest there are synergistic advantages for mosquitoes harboring salivary gland sporozoites (and therefore likely to be malarious) that align with significant increases in their vectorial capacity (Fig. 13). These notably include an expected increase in chemosensory sensitivity that could reasonably lead to more successful host seeking of infectious mosquitoes, along with a surprisingly distinctive transcriptome alteration that collectively aligns with an anti-aging paradigm that would likely provide a distinct advantage to these mosquitoes. When compared with similarly aged, uninfected mosquitoes, the synergistic advantage of naturally relevant malaria infections provides at least a partial rationale for the enduring persistence of Plasmodium pathogens and, indeed, global malaria.

Materials and methods
Ethics statement. All animal work was conducted in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health (NIH), USA. The protocols and procedures used in this study were approved by the Animal Care and Use Committee of the Johns Hopkins University (Permit Number: MO18H82) and the Johns Hopkins School of Public Health Ethics Committee. Commercial anonymous human blood was used for parasite cultures and mosquito feeding; thus, informed consent was not required, and the use of human blood for these types of experiments is exempt from human subject research.

Mosquito rearing and plasmodium infections. An. gambiae s.s. (An. gambiae Keele strain) mosquitoes were reared and maintained at 27 °C and 80% humidity with a 12-h light:12-h dark cycle at Johns Hopkins School of Public Health insectary (Baltimore, MD) according to standard procedures. Newly emerged, 0-day-old female An. gambiae were split into nine independent groups (rearing cages; six infected or treatment, and three non-infected or control). All control and treatment groups were given their first and only bloodmeal at 5 days old. Treatment groups were fed live P. falciparum NF54 wild-type (NF54W) gametocyte cultures at a final gametocytemia of 0.008% in human blood mix (provided by the Johns Hopkins Institute Core Facility,
Baltimore, MD) through glass membrane feeders at 37 °C (in the insectary chamber at 27 °C) for 1 h as previously described18,110. In brief, because gametocyte cultures take at least 2 weeks to produce mature gametocytes that are infectious to *Anopheles* mosquitoes the quantification of gametocytemia was carried out manually using blood smears at day 15 after setting up the gametocyte cultures. Exflagellation was also manually assessed to validate the infectious nature of gametocyte cultures. Day 16 gametocyte cultures with a gametocytemia (only mature gametocytes were counted) of 2.1% was centrifuged at 1800 rpm for 4 min to remove the culture medium and diluted with reconstituted whole blood prepared by mixing packed O+ red blood cells (Interstate Blood Bank, Inc.) 1:1 by volume with O+ human serum (Interstate Blood Bank, Inc.) to give a final gametocytemia for each mosquito blood feeding.

After feeding, treatment and control groups were independently sorted at 4 °C and unfed and partially fed mosquitoes were removed and disposed. The remaining fully engorged mosquitoes in treatment and control groups were incubated for 18 days in an insectary maintained at 27 °C and 80% relative humidity, with a photoperiod of 12 h light:12 h dark, with dusk and dawn periods of 1 h each at the beginning of each scotophase. All groups had continuous access to sterile distilled water and a 10% sucrose/distilled water solution throughout the duration of the study. The infection of *P. falciparum* was confirmed at the oocyst stage by sampling at least 8–10 mosquitoes from each of the 6 treatment groups (rearing cages) at 8 days post infection (Table 1). In total, 58 randomly selected mosquitoes were dissected in PBS, stained with 0.1% PBS-buffered mercurochrome (MilliporeSigma, Burlington, MA) and examined under a light-contrast microscope (Olympus Life Science, Waltham, MA). At the oocyst stage, *P. falciparum* infection prevalence of the 58 randomly selected infected mosquitoes was 62%, with a median count of two oocysts per mosquito midgut (SEM = 0.33).

**Mosquito dissections and nucleic acid extractions.** After collection, each mosquito in this study was individually stored prior to manual trans-section of heads (including sensory appendages) and thorax including salivary glands, referred to as 'head' or 'head tissues', from body/carcass which were stored for future use (Fig. 1B). Total RNAs extracted from the heads of established treatment- and control-group *An. gambiae* mosquitoes were used for Illumina NovaSeq sequencing. Total DNAs were also extracted from the heads of treatment mosquitoes for PCR to validate the *P. falciparum* infection at sporozoite stage in individual mosquitoes. All head tissues were collected from control- and treatment-group *An. gambiae* mosquitoes immediately following the 18-day incubation period (Fig. 1). Dissections of control- and treatment-group mosquito heads were performed over 3 days; each group was handled on a different day to prevent cross contamination. All mosquitoes were handled wearing gloves and all dissection equipment washed with glassware detergent, autoclaved, and decontaminated with RNase Away (MilliporeSigma, Burlington, MA). To eliminate potential hemolymph contamination during mosquito dissections, all equipment was sterilized using 90% ethanol and decontaminated with RNase Away (MilliporeSigma, Burlington, MA) in between individual mosquito handling. To maintain tissue integrity, dissections were performed on cold-anesthetized mosquitoes, and work limited to one group—treatment or control. For treatment groups, the head with intact salivary glands from one mosquito was dissected into a distinct numbered and cataloged 1.5-mL test tube containing 200µL Tri Reagent (MilliporeSigma, Burlington, MA). The smaller starting volume of Tri Reagent (MilliporeSigma, Burlington, MA) aided subsequent tissue homogenization. PCR infection validation studies on treatment-group mosquitoes required that all dissected heads remain separated. Prior to dissections, each treatment-group *An. gambiae* mosquito was randomly assigned an identification number used for labeling of all associated dissected tissues. In total, the heads of 176 treatment mosquitoes were collected into 176 distinct labeled tubes. For control lines, the heads from 30 mosquitoes within a single group were dissected and pooled into one labeled 1.5-mL test tube containing 400 µL Tri Reagent (MilliporeSigma, Burlington, MA). This process was repeated for the two remaining control groups. All sample...
Total DNA concentrations and purities were measured using NanoDrop (Thermo Fischer, Waltham, MA) and according to the manufacturer's protocol using the DNEasy Blood and Tissue Kit (Qiagen, Germantown, MD). The samples were processed in small batches of ten to preserve DNA integrity. A batch of treatment sample tubes was randomly selected samples per replicate (Fig. 1). The resulting aqueous phase (containing DNAs) from each sample was carefully transferred into a coordinately labeled, new sterile 1.5-mL test tube containing 200µL 70% ethanol and thoroughly mixed by pipetting. Total DNA concentrations and purities were measured using NanoDrop (Thermo Fischer, Waltham, MA) and any detected phenol contaminants removed by 1-butanol extraction. Control-sample total RNAs were kept separate to allow for three distinct biological replicates pertinent to RNAseq analyses. For RNA extractions, treatment sample tubes were processed in small batches of ten to preserve RNA integrity. A batch of treatment sample tubes was thawed on ice, centrifuged at 13,000 RPM for 5 min at 4 °C, and any remaining aqueous phase was removed and disposed of to prevent RNA contamination of DNA extracts. Then, 200µL back extraction buffer was added to each sample tube and hand mixed by inversion for 3 min. For complete phase separation, sample tubes were centrifuged at 13,000 RPM for 30 min at 4 °C. The resulting aqueous phase (containing RNAs) from each sample was carefully transferred into a coordinately labeled, new sterile 1.5-mL test tube containing 200µL 70% ethanol and thoroughly mixed by pipetting. Total RNAs were extracted from the aqueous phase/ethanol solutions according to the manufacturer's protocol using the PureLink RNA Mini Kit (Thermo Fischer, Waltham, MA). This process was repeated for all 179 treatment and 3 control sample tubes. Total RNA concentrations and purities were measured using NanoDrop (Thermo Fischer, Waltham, MA) and any detected phenol contaminants removed by 1-butanol extraction. Control-sample total RNAs were kept separate to allow for three distinct biological replicates pertinent to RNAseq analyses. For the treatment replicates, after each individual sample was confirmed to contain *P. falciparum* sporozoites via PCR validation, three distinct pools were prepared for RNAseq analyses. Ultimately, 65 of the 176 treatment heads were positively confirmed for *P. falciparum* infection and correlating total RNAs pooled into three replicates (21 randomly selected samples per replicate).

DNA extractions were performed for all 179 treatment sample tubes containing remaining interphase and organic phase solutions to perform PCR *P. falciparum* validation studies. Like RNA extractions, treatment sample tubes were processed in small batches of ten to preserve DNA integrity. A batch of treatment sample tubes was thawed on ice, centrifuged at 13,000 RPM for 5 min at 4 °C, and any remaining aqueous phase was removed and disposed of to prevent RNA contamination of DNA extracts. Then, 200µL back extraction buffer was added to each sample tube and hand mixed by inversion for 3 min. For complete phase separation, sample tubes were centrifuged at 13,000 RPM for 30 min at 4 °C. The resulting aqueous phase (containing DNAs) from each sample was carefully transferred into a coordinately labeled, new sterile 1.5-mL test tube containing 200µL 70% ethanol and thoroughly mixed by pipetting. Total DNAs were extracted from the aqueous phase/ethanol solutions according to the manufacturer's protocol using the DNEasy Blood and Tissue Kit (Qiagen, Germantown, MD). Total DNA concentrations and purities were measured using NanoDrop (Thermo Fischer, Waltham, MA) and any detected phenol contaminants removed by 1-butanol extraction. The total gDNA was finally resuspended at a final concentration of 10 ng/µL and kept at − 20 °C for further PCR experiments.

**PCR-based plasmodium infection validation.** PCR experiments were performed to validate the presence of *P. falciparum* sporozoites within each treatment sample. PCR experiments used total DNAs extracted from dissected heads and salivary glands of individual treatment *An. gambiae* female mosquitoes (Ref. Mosquito Dissections and Nucleic Acid Extractions). PCR was performed using an MJ Thermal Cycler (Bio-Rad Inc, Hercules, CA) in combination with the DreamTaq Mastermix technology (Thermo Scientific, Waltham, MA) according to the manufacturer's protocol. PCR products were visualized on a 1% agarose gel using the QuickGene DNA isolation kit. PCR was performed using an MJ Thermal Cycler (Bio-Rad Inc, Hercules, CA) in combination with the DreamTaq Mastermix technology (Thermo Scientific, Waltham, MA) according to the manufacturer's protocol. PCR products were visualized on a 1% agarose gel using the QuickGene DNA isolation kit and second, mechanically by sonication (Q700; QSonica, Newtown, CT) to ensure thorough head tissue and cell disruption. Sonication was performed in short 10- to 20-s bursts with samples on ice to minimize nucleic acid shearing. To prevent cross contamination, one pellet pestle was used per sample tube, and the sonicator probe was thoroughly sterilized using 90% ethanol and decontaminated with RNase Away (MilliporeSigma, Burlington, MA) in between individual tube processing. Following complete tissue homogenization, RNA extractions were performed. An additional 200µL Tri Reagent (MilliporeSigma, Burlington, MA) was added to each treatment sample tube bringing the total volume to 400 µL, equivalent to control sample volumes. Sample tubes were then hand shaken and incubated at room temperature for 5 min. A 100-µL aliquot of UltraPure phenol:chloroform:isoamyl alcohol (25:24:1, v/v; Thermo Fischer, Waltham, MA) was added to each of the tubes, which were then vigorously hand shaken for 1 min and incubated at room temperature for 5 min. For complete phase separation, sample tubes were centrifuged at 13,000 RPM for 15 min at 4 °C. The resulting aqueous phase was carefully transferred into a coordinately labeled new RNase-free 1.5-mL test tube containing 100µL 70% ethanol and thoroughly mixed by pipetting. Sample tubes containing the remaining interphase and organic phase were frozen at − 70 °C in preparation for DNA extractions. Total RNAs were extracted from the aqueous phase/ethanol solutions according to the manufacturer's protocol. Total RNA concentrations and purities were measured using NanoDrop (Thermo Fischer, Waltham, MA) and any detected phenol contaminants removed by 1-butanol extraction. Control-sample total RNAs were kept separate to allow for three distinct biological replicates pertinent to RNAseq analyses. For the treatment replicates, after each individual sample was confirmed to contain *P. falciparum* sporozoites via PCR validation, three distinct pools were prepared for RNAseq analyses. Ultimately, 65 of the 176 treatment heads were positively confirmed for *P. falciparum* infection and correlating total RNAs pooled into three replicates (21 randomly selected samples per replicate).

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**CS forward (Fw):** 5′-AGCAGAAGATATGTTACGAACAAAACA-3′.
**CS reverse (Rv):** 5′-ACCTTTGGCCATTACCTTGATTG-3′.
**AgRsp7 Fw:** 5′-AGATTCGAACTCTTGGTGGTGC-3.
**AgRsp7 Rv:** 5′-ACAACAACATCGAAGGATACGA-3′.

All PCR primers were validated and optimized in preliminary studies. PCR experiments were conducted for all treatment samples, with a correlating negative control (blank) and positive *P. falciparum* infected positive control (Fig. 2). Treatment samples producing CS PCR products were reconfirmed to be positively infected through a second follow-up PCR experiment.

**Illumina NovaSeq paired-end sequencing.** Deep sequencing utilizing Illumina RNAseq technology (Illumina, San Diego, CA) was performed at Hudson Alpha Discovery Genomic Services Laboratory (Huntsville, AL). Poly(A) mRNAs were isolated separately for library preparation and barcoding from approximately 500 ng of total RNA collected from each of the three control replicates and approximately 300 ng of total RNA.
collected from each of the three treatment replicates. Final mRNAs were fragmented, and cDNA was synthesized, amplified, digested, and purified using Illumina TruSeq chemistry protocols (Illumina, San Diego, CA). In sequencing, for each of the six libraries—three treatment and three controls—was labelled with a 16-bp barcode to identify sequencing data generated from each library [treatment replicate 1 (T1) = TTGAGACTCAGGG, treatment T2 = CGTCTAAGTGTCGC, treatment T3 = AACTGATGAGGCC, control C1 = AGACTTTAACATCT, control C2 = GCCTAAGTACGGC, and control C3 = TTGAGACTCCTC TAA]. Barcoded treatment and control replicate libraries were hybridized across four lanes of two S4 flow cells for cBot (Illumina, San Diego, CA) cluster generation and sequencing (paired-end, 2 × 150 bp; total 300 cycles). A total of 36 raw data sets of sequencing outputs, 6 per library, were generated using the Illumina software assembler (Illumina, San Diego, CA).

**Illumina bioinformatics.** The 36 control and treatment replicate Illumina NovaSeq data sets generated were cleaned and quality trimmed (Q30) using Trimmomatic v0.33113, then verified for quality control using FastQC v.0.11.9 (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/). Control and treatment replicate data sets were aligned to the An. gambiae PEST strain genome (Vectorbase-53_AgamiaBePest_Genome.fa, https://vectorbase.org) using STAR v.2.7.5 (https://github.com/exobin/STAR), with An. gambiae PEST strain gene annotations (Vectorbase-53_AgamiaBePestGtf, https://vectorbase.org) to map splice junctions, 2-pass mapping, –sjdbOverhang set to 149 bp, MAPQ set to 10, and a preliminary UNIX concatenate script result in six output BAM alignment files: control R1, control R2, control R3, treatment R1, treatment R2, and treatment R3. HTSeq v.0.11.2114, an alignment-based count algorithm, was used to quantify transcript counts for all six BAM alignment files with default parameters and mapping validation. DESeq2 (Bioconductor v.3.8) and R v.3.5.1 were used to calculate mean variance and obtain differentially abundant transcripts between control and treatment replicates using default parameters with added shrinkage and Wald’s hypothesis tests. Gene ontologies (GO) of differentially abundant transcripts were obtained from OmicsBox as GenBank non-redundant protein (nr) database containing all available mosquito genomes with an expected value (e-value) cutoff of less than 0.01. All GO annotations were thereafter manually confirmed via hypothesis tests. Gene ontologies (GO) of differentially abundant transcripts were obtained from OmicsBox using the GenBank non-redundant protein (nr) database containing all available mosquito genomes with an expected value (e-value) cutoff of less than 0.01. All GO annotations were thereafter manually confirmed via hypothesis tests.

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Author contributions

A.L.C. and L.J.Z. designed the study. A.L.C. and Y. D. performed the experiments. A.L.C and D.R. analyzed data and prepared figures; A.L.C, D.R., Y.D., G.D. and L.J.Z. interpreted the results and wrote the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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