Genetic changes of *Plasmodium vivax* tempers host tissue-specific responses in *Anopheles stephensi*

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A B S T R A C T

Recently, we showed how an early restriction of gut flora proliferation by *Plasmodium vivax* favors immune-suppression and *Plasmodium* survival in the gut lumen (Sharma et al., 2020). Here, we asked post gut invasion how *P. vivax* interacts with individual tissues such as the midgut, hemocyte, and salivary glands, and manages its survival in the mosquito host. Our data from tissue-specific comparative RNA-Seq analysis and extensive temporal/spatial expression profiling of selected mosquito transcripts in the uninfected and *P. vivax* infected mosquito’s tissues indicated that (i) a transient suppression of gut metabolic machinery by early oocysts; (ii) enriched expression of nutritional responsive proteins and immune proteins against late oocysts, together may ensure optimal parasite development and gut homeostasis restoration; (iii) pre-immune activation of hemocyte by early gut-oocysts infection via REL induction (p < 0.003); and altered expression of hemoocyte-encoded immune proteins may cause rapid removal of free circulating sporozoites from hemolymph; (iv) while a strong suppression of salivary metabolic activities, and elevated expression of salivary specific secretory, as well as immune proteins together, may favor the long-term storage and survival of invaded sporozoites. Finally, our RNA-Seq-based discovery of 4449 transcripts of *Plasmodium vivax* origin, and their developmental stage-specific expression modulation in the corresponding infected mosquito tissues, predicts a possible mechanism of mosquito responses evasion by *P. vivax*. Conclusively, our system-wide RNA-Seq analysis provides the first genetic evidence of direct mosquito-*Plasmodium* interaction and establishes a functional correlation.

1. Introduction

The evolution and adaptation of adult female mosquitoes to blood-feeding have a significant influence on their reproductive outcome and disease transmission dynamics. The sexual cycle of malaria-causing *Plasmodium* immediately begins after the ingestion of gametocytes containing blood meal by anopheline mosquitoes (Bennink et al., 2016; Kuehn and Pradel, 2010; Talman et al., 2004). Within the lumen of the midgut, male and female gametes fuse to form a zygote, which then transforms into motile ookinete (Aly et al., 2009; Bennink et al., 2016). Eventually, within 24h ookinetes traverse through the gut epithelium to reach the basal lamina either through intracellular and/or intercellular routes and transform into tiny oocysts (Baton and Ranford-Cartwright, 2004; Han et al., 2000). During this phase, mosquitoes impart early defense responses through nitration of the midgut and activation of signaling pathways (Ramphul et al., 2015). Midgut nitration results in modification of ookinete surfaces, which makes them perceptible to mosquito complement-like system (Garver et al., 2013; Shiao et al., 2006). Whereas signaling pathways show different responses towards different *Plasmodium* species, such as the IMD pathway, work more effectively against *P. falciparum* than *P. berghei*, and the Toll pathway is more responsive to *P. berghei*, and *P. gallinaceum* (Kumar et al., 2004; Cirimotich et al., 2010). Active pathways boost the rapid production of immune peptides i.e. AMPs (Cecropin, gambicin, etc.), which are responsible for parasite killing.

The majority of ookinetes killing is targeted at the basal side of midgut epithelium, through an indirect hemocyte-mediated cellular response such as lysis or melanization (Smith et al., 2016; Belachew, 2018). A recent study showed that during the immune active complement system, the hemocyte responds through microvesicles (Castillo et al., 2017). Once they reached the basal lamina beneath the epithelium and enveloped by basal lamina, the surviving ookinetes rapidly transform

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into oocytes. These hidden young oocytes gradually mature to round shaped large oocytes and remain protected from any external response until they burst up into millions of sigmoid-shaped sporozoites. The underlying mechanism of how maturing oocytes (i) are least affected by any external responses such as immune responses caused by hemocytes; (ii) regulates their time-dependent developmental transformation to mature oocytes; (iii) trigger sporozoites release outside the basal lamina by rupturing the oocytes, is not well known.

Once released, millions of free circulatory sporozoites (fcSpz) are rapidly killed by the immune blood cells i.e. hemocytes. Despite a substantial reduction in the population of sporozoites (~85 fold) (Gouagna et al., 1998), a significant number of parasites succeed to specifically invade the salivary glands and survive within them. Several hemocyte-mediated mechanisms such as melanization, lysis, engulfment, and toxic anti-Plasmodium immune effector molecules have been proposed to kill the free circulating sporozoites, but the molecular nature of these interactions is unknown (De et al., 2018; Juli Hillyer et al., 2003; Smith et al., 2016). Even for salivary invasion, Plasmodium needs salivary gland proteins such as saglin, MAEBL, and PGRP (Kariu et al., 2002). A recent study also showed that Plasmodium invasion induces several salivary immune proteins such as SP14D1, Ficolin, SP24D, PGRP, and LRIM (Roth et al., 2018). However, the genetic basis of how sporozoites (i) manage to avoid hemocyte-mediated killing responses, (ii) guide its movements for salivary invasion, and (iii) enhances their virulence in the salivary environment is not well understood (Mueller et al., 2010; Roth et al., 2018; Vanderberg et al., 1975; Mikolajczak et al., 2008; Sato et al., 2011).

In our preceding study (see Sharma et al., 2020) we demonstrated that in the gut lumen, Plasmodium vivax follows a unique strategy of immunosuppression by disabling gut flora proliferation. Here, we further decode and establish a functional correlation of how the post-gut invasion of P. vivax tempers the tissue-specific molecular responses. Using a comprehensive RNA-Seq analysis of P. vivax-infected midgut, hemocyte, and salivary glands, we not only identified mosquito transcripts but also captured a large pool of P. vivax transcripts expressed in respective tissues. Detailed annotation and transcriptional profiling revealed that P. vivax can avoid immuno-physiological responses by altering its genetic architecture. Manipulating tissue-specific immuno-physiology of mosquitoes may limit the Plasmodium development and hence the transmission.

2. Materials and methods

Technical overview and workflow of the project are presented in Supplementary Material Fig. S1 document (Data Sheet-1).

2.1. Mosquito rearing

The cyclic colony of the mosquito An. stephensi was maintained at 28 °C ± 2, RH = 80% in the insectary fitted with a simulated dawn and dusk machine with a 12 h Light/Dark cyclic period at NIMR. For egg maturation and regular gonotrophic cycle maintenance, mosquitoes were fed on rabbit blood. All protocols for rearing, maintenance of the mosquito culture were approved by the ethical committee of NIMR (ECR/C14/C6). For egg maturation, all protocols for rearing, maintenance of the mosquito culture were approved by the Ethics committee of NIMR, Delhi (ECR/C14/C6/2012/41). An equal number of age-matched overnight starved 4–5 days old female A. stephensi mosquitoes were fed on uninfected or P. vivax infected blood samples, using optimized artificial membrane feeding assay (AMFA). Fully engorged mosquitoes were maintained at optimal insectarium conditions and positive infection was confirmed by standard mercurichrome staining of gut oocysts readily observed under a compound microscope (Musiime et al., 2019). Post confirmation of P. vivax infection, ~20–25 mosquitoes were dissected at 3–6days post-infection (DPI) and 8–10 DPI for midgut, 9–12 DPI for hemocytes, and 12–14 DPI for salivary glands in PBS and collected in Trizol reagent.

### Tissue-specific RNA-Seq library preparation & sequencing:

Total RNA was isolated from each dissected tissue sample using the standard trizol method. Due to technical limitation, we followed a well-established and in-house optimized sample pool protocol, with the add-in of a SMART- PCR based mRNA pre-amplification protocol available with Clontech SMARTTM kit (Dixit et al., 2009, 2011, 2015a; Palomares et al., 2019). Approximately, ~ 1 µg of purified total RNA, was subjected to construct a full-length double-stranded cdNA library. Chances of over-amplification and optimal normalization were achieved by PCR cycle numbers optimization for individual samples, followed by validation through Real-time PCR strategy (see Data Sheet1). Quality of the ds-cDNA was checked on Bioanalyzer 2100 using Agilent DNA HS chip Sequencing of whole transcriptomes was performed on Illumina NextSeq, where a constant amount of 200 ng of each ds cDNA samples was subjected to Covaris shearing followed by end-repair of overhangs resulting from shearing. The paired-end cDNA sequencing libraries were prepared using Illumina TruSeq Nano DNA HT Library Preparation Kit as per the described protocol. The end-repair fragments were A-tailed, adapter-ligated, and then enriched by a limited number of PCR cycles. Library quantification and qualification was performed using DNA High Sensitivity Assay Kit and sequenced using 2 × 150 PE chemistry on NextSeq for generating ~ 1 GB of data per sample.

### RNASeq database generation and assembly:

Following sequencing raw data were filtered using Trimmomatic v0.30. All the reads with adapter contamination and reads with low-quality value i.e. an average QV less than 20 (QV < 20) were filtered. Proper filtration high-quality clean reads of all tissue-specific samples were assembled with Trinity software (release r2013-02-25) on optimized parameters. Due to poor annotation of mosquito An. stephensi reference genome, we followed the De-Novo strategy to assemble individual tissue-specific transcriptome databases. CD-HIT-EST (Version 4.6) was used to remove the shorter redundant transcripts when they were entirely covered by other transcripts with 90% identity. From assembled contigs/transcripts, CDS were predicted from the longest reading frame using the Trans-decoder. Subsequently, the predicted CDS were annotated as per the technical plan presented in Fig. S1 (see Data sheet1). To validate the quality of assembly, we randomly selected at least five transcripts of sizes ranging from >1000bp (large), 500-1000bp (medium), and 300-500bp (short) from each RNA-seq database, and manually performed homology using BLASTx search analysis against both NCBI/NR database and BLASTx as well as mosquito specific Genome/transcripts databases, as described earlier (Sharma et al., 2015a). A maximum BLASTx homology (>80% identity BLASTx homology (>99% identity with full coverage), and global BLASTx based species distribution analysis showing the dominant match with anopheline mosquito species, together provide enough confidence to accept and process the denovo assembly database for functional annotation (see Data sheet1).

### GO annotation and molecular cataloging:

Post database assembly, the predicted CDS FASTA file of individual tissues of P. vivax infected or uninfected control samples, were processed for functional prediction analyses and molecular cataloging through using Blast2GO PRO 3.0 software (Conesa et al., 2005). A comprehensive functional annotation integrated with Blast2GO analysis was used for the construction of Immuno-Physiological and Cellular Ontology databases. Basic BLASTx based homology search analysis was valuable to assign accession IDS and predict the closest species homology to the transcript encoded proteins. Next, GO mapping was carried out to retrieve GO terms for all the BLASTx functionally annotated CDS, where assignments were used to classify the functions of the predicted CDS and defined ontology terms representing gene product properties, which could be grouped into three main domains: Biological process, Molecular Function, and Cellular Component. To retrieve GO terms for annotated
CDS, the BLASTX resulted in accession IDs were used to retrieve gene names or symbols, followed homology search in the species-specific entries of the gene-product tables or via direct accession IDs search against dbxref table of the database.

For immune cataloging, all predicted CDS were subjected to blast analysis against An. gambiae, D. melanogaster, Ae. aegypti and Culex pipiens Immune database (http://cegg.unige.ch/Insecta/immunodi), as described earlier (Thomas et al., 2016). Then, transcripts with E-value < e⁻¹⁰ were shortlisted, cataloged, and compared to select transcripts for expression profiling. To catalog tissue-specific P. vivax gene from a pre-analyzed whole transcripts database (NR/BLASTX), the best match Plasmodium transcripts were retrieved as the FASTA file. A standard GO annotation was performed, and P. vivax transcripts were analyzed for functional prediction.

Digital Gene Expression analysis: The high-quality reads for each sample was mapped on their respective set of CDS using CLC Genomics Workbench for read count calculation. The common hit accesses based on BLAST against the NR database were identified for differential gene expression analysis. The differential gene expression was carried out by using the DESeq R package. CDS were further classified as up and down-regulated based on their log fold change (FC) value calculated by formula, FC = Log 2 (Treated-Control), FC values greater than zero were considered up-regulated whereas less than zero were down-regulated.

The level of expression is represented as a log2 ratio of transcript abundance between control (uninfected) and experimental samples (P. vivax infected). Differentially expressed genes identified in control and experimental conditions were analyzed by hierarchical clustering. A heat map was constructed using the log-transformed and normalized value in FPKM (fragments per kilobase of exon per million fragments mapped) of genes based on Pearson uncentered correlation distance as well as based on complete linkage method, and the significance for differentially expressed genes was calculated at a P-value < 0.05. We compared and estimated the tag dispersion, assuming each tissue-specific two samples from different conditions with comparable gene expression for common transcripts. As both the samples contained RNA extracted from tissues of approximately ~25 mosquitoes each and pooled to form one single sample, a quantifiable estimation of gene expression was expected with a minimum chance of aberrations. Our DESeq data analysis with a single replicate was valuable to identify differentially expressed genes (Alaux et al., 2011; Wang et al., 2010; Sharma et al., 2015; Assefa et al., 2020), which was subsequently validated by large scale gene expression analysis by real-time PCR. Full technical details and reports of sequence analysis with representative sample data as well as DGE analysis validation are presented in supplemental data (see Data Sheet1).

Transcriptional Profiling analysis: For spatial and temporal expression analysis of DGE selected transcripts of mosquito and/or P. vivax origin were monitored through Real-Time PCR analysis, as described previously (De et al., 2018; Thomas et al., 2016; Sharma et al., 2020). Briefly, the target tissues from ~15 to 20 uninfected or P. vivax infected (average infection level i.e. 50-130 oocysts/midgut) mosquitoes were collected in trizol and isolated ~1 μg total RNA was subjected to cDNA through Reverse-Transcription protocol (Verso cDNA synthesis kit@Thermo Scientific, USA). Prior transcriptional profiling, the quality of each cDNA reaction was evaluated by amplification of internal control gene Actin or rp57 through standard RT-PCR and agarose gel analysis. Only the qualified cDNA templates were further used for relative gene expression profiling by BIORAD CFX96 Real-time systems, using SYBR® FAST universal SYBR® qPCR master mix. PCR cycles included initial denaturation at 95 °C for 5 min, then further 40 cycles of 95 °C for 10 s, 52 °C for 15 s, and 72 °C for 22 s. After each cycle, fluorescence was detected at the 72 °C step. Then further final steps include 65 °C for 0 sec and 95 °C for 5 sec. Each experiment was conducted in triplicate to minimize variation, except for hemocytes which were tested in duplicate, especially due to technical limitation of sample collection. The actin gene was used as an internal control for normalizing the relative expression data, and analysis was performed by 2–ΔΔCt method (Sharma et al., 2020).

Statistical analysis: For statistical analysis “test” sample data was compared with the ‘control’ data set. Since, we compared each ‘test’ data with a single ‘control’, and therefore statistically analyzed using Student’s t-test only. Significant differences were considered at the P < 0.05 level.

3. Results

Working hypothesis development and RNA-Seq data generation: Previous studies targeting individual tissues, either midgut or salivary glands, have been valuable to understand the mosquito-parasite interactions. But still, there are several unresolved questions regarding how (i) each tissue viz. midgut, hemocytes, and salivary glands together coordinate and manage the challenge of Plasmodium infection; and (ii) how Plasmodium avoids the tissue-specific host immunity and quickly respond and adapt to environmental changes in different tissues for its survival and transmission. To partly answer and resolve tissue-specific molecular complexity, we developed a working hypothesis. We opined that when blood meal itself significantly alters mosquitoes ‘metabolic physiology’, Plasmodium infection may cause an additional burden of immune activation. Thus, mosquitoes may need to follow a dual management strategy for overcoming the challenge of a metabolic shift in response to fast gut engorgement (De et al., 2019), as well as limiting the parasite population. Alternatively, for its survival, parasites may either suppress or temper/modulate the host tissue responses.

To test and evaluate the hypothesis, we designed a strategy to capture a molecular snapshot of the three different tissues directly interacting with Plasmodium vivax. After the establishment of an artificial membrane feeding protocol, we fed mosquitoes with clinically diagnosed P. vivax-infected patient’s blood samples (0.5–2% gametocytaemia). In our regular experience, we noticed an average infection intensity of 50–130 oocysts/midgut 5–6 days post-infection. But in a few experimental studies, we noticed a super-infection of P. vivax, raising the average infection intensity to 300–380 oocysts/midgut (Fig. 1). Since, for our RNAseq project, facing a challenge of technical limitation of getting ‘superinfection’ mosquito samples, we cautiously followed a sample pool, and add-in SMART-cDNA technology, which not only advantageous to discover tissue-specific P. vivax transcripts but also found reliable to validate the DEG data using ‘average infection’ samples for transcriptional profiling in triplicates (Palomares et al., 2019; Assefa et al., 2020; see Supplemental data sheet1/Table S9).

Accordingly, from infected 20–25 individual mosquitoes, we dissected and pooled targeted midgut, hemocytes, and salivary gland tissues, and performed a tissue-specific comparative RNA-Seq analysis. As per the technical design, we sequenced two midgut samples covering early (3–6 days) pre-mature to maturing oocysts, and late (8–10 days) fully matured/bursting oocysts; one hemocyte/hemolymph sample pooled from 9 to 12 DPI, a time window matching free circulatory sporozoites infection stage; and one sample of salivary glands 12–14 DPI (Fig. 1; see PanelA-B). For comparative study, we sequenced the naïve blood-fed mosquitoes’ tissues collected from the same aged mosquitoes. In the case of the midgut, we sequenced only one sample 3–4 days’ post blood meal (also see Sharma et al., 2020). From the seven RNA-Seq libraries we generated and analyzed a total of 28942203 reads (ST1). Table 1 represents the complete stat of the sequencing and analysis of each tissue-specific RNA-Seq data. Due to the unique nature of RNA-Seq sample i.e. ‘superinfection’, we mainly aimed to understand the genetic basis of parasite development in the mosquito, along with a very basic knowledge that how P. vivax infection may influence tissue-specific immune-physiological responses.

Plasmodium vivax alters the molecular architecture of mosquito tissues: To understand that, how P. vivax infection influences tissue-specific molecular responses, we compared the global change in the transcript abundance of ‘uninfected’ and ‘infected’ blood-fed mosquito tissues. Our initial attempt of mapping of cleaned reads (filtered low quality, microbial, and P. vivax origin) to the available draft reference
genome was unsuccessful. Alternatively, we mapped all the high-quality reads against the \textit{denovo} assembled reference map, as described earlier (De et al., 2018; Sharma et al., 2015; Thomas et al., 2016). Though, a read density map analysis revealed that \textit{P. vivax} infection causes a significant suppression in the salivary glands and midgut transcripts, but caused a greater shift in the read density of the infected hemocyte

![Image](99x286 to 497x738)

**Fig. 1.** Technical presentation of mosquito infection validation and tissue collection strategy for tissue-specific RNA-Seq analysis: (a/i-ii) a comparative microscopic analysis of average infected and superinfected mosquito midgut, six days’ post \textit{Plasmodium vivax} infection, (see “ruler needle pointer” of microscope eyepiece, highlighting the presence of gut oocysts as dark red circle); (b/i-ii) Microscopic view of early and late oocyst development in the mosquito midgut; (c): (i) Microscopic view of \textit{Plasmodium} sporozoites in the mosquito salivary glands; (ii) Molecular confirmation of parasite infection in the mosquito by RT-PCR and agarose gel electrophoresis Uninfected (C) & Infected (I) midgut cDNA samples were subjected to 28 PCR cycle amplification using mosquito specific Actin & parasite-specific 18S primers. M = Molecular marker.

| Sl. No. | Sample Name  | No. of Reads | No. of Contigs | Total transcripts | Transcripts with BLAST Hits (NR) | Annotated CDS analyzed |
|--------|--------------|--------------|----------------|-------------------|---------------------------------|------------------------|
| 1.     | MG-C         | 4673408      | 74,548         | 11159             | 10219 (91%)                     | 6205                   |
| 2.     | MG-INF (3-6D) | 3625905     | 52,187         | 12065             | 11011 (91%)                     | 6813                   |
| 3.     | MG-INF (8-10D) | 5399936     | 42,212         | 7585              | 7222 (95%)                      | 7496                   |
| 4.     | HC-C         | 3827903      | 30,959         | 6236              | 6044 (96%)                      | 2970                   |
| 5.     | HC-INF       | 3253689      | 18,462         | 3492              | 3333 (95%)                      | 3418                   |
| 6.     | SG-C         | 3782999      | 64,966         | 7674              | 6640 (89%)                      | 6776                   |
| 7.     | SG-INF       | 4396363      | 32,110         | 4492              | 4100 (91%)                      | 2743                   |

Table 1
Complete sequencing stat of tissue-specific RNAseq libraries.
transcripts (Supplemental data sheet-2/Fig. S1). To further uncover the molecular and functional nature of the encoded proteins, we carried out a comprehensive GO annotation, cataloged, and extensively profiled tissues specific shortlisted genes altered in response to P. vivax infection.

Midgut response to Plasmodium oocysts development: A comparative catalog of biological process (L4) revealed that early maturing oocyst development coincides with the suppression of the majority of midgut proteins (ST1). Although oocyst development exceptionally induces the expression of proteins encoding cellular catabolic, organic substance catabolic, and organic acid metabolic processes, it also restricts the expression of proteins with regulatory functions (Fig. 2a). Interestingly, when compared to naive midgut, late-stage oocysts showed a re-enrichment of the gut proteins with common functions (Fig. 2a). Venn diagram analysis of annotated transcripts showed that 877, 1036, and 1251 transcripts uniquely restricted their expression to naïve, early, and late oocysts in the infected mosquito guts, respectively, (Fig. 2c). A heat map analysis of differentially expressed common transcripts showed a significant alteration in response to early oocysts infection (Fig. 2e). Taken together, these observations suggest that the fast-developing oocysts of P. vivax may cause a biphasic modulation of gut metabolic machinery, allowing early suppression and late recovery for the maintenance of homeostasis.

To further clarify how Plasmodium vivax infection alters gut immunophysiological responses, we identified, cataloged, and compared the expression of selected transcripts in response to a blood meal and P. vivax infection. A comparative gut-immunome analysis showed an increased percentage of immune transcripts, except for a few classes such as Autophagy, TAK, IAP, PGRP, and IKKG (Fig. 2b). Additionally, we also observed an exclusive enrichment of several classes of immune proteins such as CTL, Gambicin, GNBP, IMD, CTMLA, and IML, in response to early infection (ST2). As expected, blood meal causes a transient change in immune gene expression (S1), but in the P. vivax-infected midgut samples, we observed a multifold enrichment of Gambicin expression after 48hr and late induction of other AMPs such as C1, C2, and D1 (see Sharma et al., 2020). Together, these data suggest that a time-dependent action of distinct AMPs against P. vivax is necessary to delimit the gut-specific oocysts’ development. Furthermore, the rapid induction of transcripts encoding Folliculin, Trehalase, and Sterol carrier indicated their possible role in managing nutritional imbalance during P. vivax development (Fig. 2d).

Hemocyte response to free circulatory sporozoites: Infected mosquito hemocytes showed a slight enrichment of transcripts (1408) encoding the diverse nature of proteins (ST1). The expression of 1128 transcripts linked to organelle organization and ribonucleoprotein complex biogenesis functions remains uniquely associated with infected hemocytes (Fig. 3a). Interestingly, we observed at least 966 transcripts encoding proteins of cellular catabolic and organic acid metabolic processes, which remained restricted to naïve blood-fed mosquito hemocytes. Surprisingly, contrary to this in the gut, similar categories of proteins were exclusively induced in response to early oocysts infection (see Figs. 2 and 3). FPKM (Fragments Per Kilobase of transcript per Million mapped reads) based heat map analysis further showed a unique modulation of common transcripts, with restricted expression either in the naïve or infected mosquito hemocytes (Fig. 3e).

Since mosquito hemocytes mount a highly specific cellular immune response, we next targeted to decode the molecular nature of the immune interaction between mosquito hemocytes and free circulating sporozoites (fSPZ) of P. vivax. A comparative immunome (ST2) analysis indicated that fSPZ causes a greater suppression of many immune family proteins, except for the marked enrichment of GPX, TPX, Diperinet, LYSC, PGRP, and ML family proteins (Fig. 3b). Interestingly, like the midgut, infected mosquito hemocytes also showed an exclusive induction of similar classes of immune proteins such as CTL, Gambicin, Defencin, IAP, and IMD. (Fig. 3d). Significant enrichment of REL than NOS coincided with early oocysts development in the gut (Fig. 3e). We also observed an exclusive induction of the physiologically active class of immune proteins such as ApolIII, Hexamerin, and FREP13 during the 8th day of P. vivax infection (Fig. 3d). While, in our recent study we further observed that the expression of FREP12 significantly upregulated after 10 days of P. vivax infection, highlighting its possible role in immunity against free circulating sporozoites (Chauhan et al., 2020).
Salivary glands response to *P. vivax* infection: In contrast to midgut and hemocytes, *P. vivax* infection impaired the molecular architecture of the salivary glands by delimiting the expression of common salivary transcripts (ST1). Additionally, we also observed restricted expression of primary metabolic process proteins in uninfected mosquito salivary glands (Fig. 4a). Venn diagram analysis further indicated that a large pool of 3069 (60%) transcripts were restricted to uninfected mosquito salivary glands and the remaining 699 (12%) transcripts showed a unique appearance to sporozoites invaded salivary glands (Fig. 4c). A heat map data analysis further suggested that salivary sporozoites may keep a stronghold on the salivary metabolic machinery, possibly favoring its survival (Fig. 4e).

Next, to test how the salivary immune system influences sporozoite development and survival, we cataloged and compared the relative percentage of immune transcripts (ST2). We noticed an exclusive appearance of four classes of immune family proteins, such as FADD, Gambicin, GNBP, and SCRC in the invaded salivary glands (Fig. 4b). The expression of TAK1, CASP, HOP, and STAT immune family proteins remained restricted to naïve mosquito salivary glands (Fig. 4d). Transcriptional profiling of selected immune transcripts such as Gambicin, Cecropin, and SP-24 D showed a high induction in the salivary glands (Fig. 4e). Surprisingly, several other salivary secretory proteins such as anopheleine, *D7 Family*, ion transporter family proteins, and 53.7 kDa also
showed significant up-regulation, suggesting their anti-Plasmodium role against *P. vivax* infection.

**Deep sequencing identifies *P. vivax* transcripts:** The above analysis allowed us to hypothesize that when *Plasmodium* sp. switches from one stage to another, it may follow a unique strategy to regulate tissue-specific metabolic and immuno-physiological responses. The surprising finding of a large pool of 4,449 transcripts of *P. vivax* origin, distinctly expressed in their respective infected mosquito tissues (Table 2), encouraged us to establish a genetic relationship of mosquito-parasite interaction. We noticed that 73.8% of *P. vivax* transcripts were associated with oocysts (early and late), while 74 transcripts are originated from free circulatory sporozoites, and 1,133 transcripts from salivary sporozoites (ST3).

| Target infection stage | Total *P. vivax* transcripts | Hypothetical/Unknown | Unique/stage-specific | The qualified best match to other *Plasmodium* |
|------------------------|------------------------------|-----------------------|-----------------------|-----------------------------------------------|
| MG(OOC-E)              | 3208                         | 1265                  | 1943                  | 2200                                          | 539                                      |
| MG(OOC-L)              | 75                           | 65                    | 15                    | 9                                             | 10                                       |
| HC(fcsSPZ)             | 74                           | 57                    | 17                    | 6                                             | 7                                        |
| SG(sgp-SPZ)            | 1133                         | 797                   | 336                   | 162                                           | 113                                      |
| TOTAL                  | 4449                         | 2184                  | 2311                  | 2377                                          | 669                                      |

Table 2: Detailed information of *P. vivax* transcripts retrieved from mosquito transcriptomes.

Molecular changes of *P. vivax* may facilitate its survival and transmission: To establish a functional correlation, we annotated, cataloged, and compared the relative abundances of the *P. vivax*-specific transcripts. Initially, putative transcripts encoding *Plasmodium* homolog proteins were filtered out from the pre-analyzed RNA-Seq dataset. Each developmental stage of *P. vivax* revealed a large pool of hypothetical proteins, especially in early oocysts and salivary sporozoites (Table-2), suggesting complex biology is yet to unravel. Further, GO annotation revealed that each developmental stage carries an enriched transcript abundance of the genes linked to cellular as well as metabolic processes (Fig. 5a). Surprisingly, free circulatory sporozoites not only showed a selective enrichment for the family proteins encoding ‘Response to Stimulus’ and ‘Localization’ but also unraveled an exclusive appearance of signaling and detoxification linked genes in the hemocytes.

Since the rupturing of late oocysts releases millions of fcsSPZ, which are highly vulnerable to the hemocyte cellular immune response, we hypothesize that *Plasmodium* must have a unique strategy to defend its survival. An in-depth functional annotation analysis (BP-Level 4), clearly demonstrated that despite having a low number of transcripts, late oocysts and fcsSPZ encoded a more diverse nature of proteins than early oocysts and salivary sporozoites (Fig. 5b). A Venn diagram analysis further reveals that the molecular nature of *Plasmodium* encoded proteins is significantly altered when it switches from one stage to another, particularly late oocysts (LO) to fcsSPZ.

Remarkably, a cross tissue comparison showed a restricted expression of a large number of proteins in each developmental stage of *P. vivax* (Fig. 6a–c). Interestingly, a total of 2,200 transcripts showed restricted expression to early oocysts (EO), while 9 transcripts were exclusively expressed to the late oocysts (LO) stage. Out of the total 74 transcripts, at
least 6 showed restricted expression to the fcSPZ stage and 162 transcripts expression exclusively remained in the salivary invaded sporozoites (Fig. 6d). Though the exact mechanism of this developmental transformation is yet unknown. But an FPKM-based relative expression analysis and transcriptional profiling of selected common transcripts further suggested that the ability to alter genetic makeup may enable P. vivax to misguide tissue-specific immuno-physiological responses and survival in the mosquito host (Fig. 6e–f).

Finally, we targeted the catalog of well-annotated key genes that may have a potential role in parasite development and transmission. To do this, we performed a comprehensive literature search analysis and shortlisted transcripts with high FPKM, but a restricted expression to a particular tissue (Fig. 7). Though an ongoing independent detailed comparison of P. vivax transcript database with other parasites such as P. falciparum and P. berghei, yet to unravel genetic differences (unpublished), current findings provide a valuable resource for P. vivax transcripts, especially free circulatory sporozoites directly interacting with mosquito hemocytes, for future functional characterization.

4. Discussion

Evolution and adaptation to blood-feeding is an essential requirement for mosquitoes’ reproductive success (Attardo et al., 2005; Hansen et al., 2014; Kokoza et al., 2001). However, Plasmodium parasites take the benefit of mosquitoes’ blood-feeding behavior for the completion of the sexual cycle and disease transmission (Thomas, 2012; Lacroix et al., 2005; Schwartz and Koella, 2001). Thus, to win the developmental race, we opined that Plasmodium must have the ability to manipulate immuno-physiological responses of directly interacting tissues, such as the midgut, hemocytes, and salivary glands of mosquitoes (Cator et al., 2012). It has been well established that during its development inside susceptible mosquito hosts, the population dynamics of Plasmodium significantly altered (Vezilier et al., 2012). But holistically the molecular nature of tissue-specific host-parasite interactions, covering gut-hemocyte-salivary gland-Plasmodium together in a single study, experimentally remains poorly understood (Molina-Cruz et al., 2012). Using a system-wide RNAseq analysis, here we now uncover the tissue-specific genetic basis of mosquito-Plasmodium interaction.

Initial profiling of AMPs/non-AMPs in response to uninfected blood-feeding showed a transient influence in the tested tissue, suggesting a general role in the maintenance of physiological homeostasis (Fig. 54). However, surprisingly, when compared to an uninfected counterpart, each tissue shows a unique relationship of metabolic alteration in response to P. vivax infection. Our functional annotation of gut-RNAseq data demonstrated that post-gut invasion, the mosquito can regulate and recover from the acute harms caused by fast-developing early oocysts. However, many naturally selected refractory anopheline species can cease the development of parasites through melanization (Molina-Cruz et al., 2012; Simoes et al., 2017). It is plausible to predict that even in the susceptible strain, the controlled regulation of gut-parasite interplay is key for the survival of both mosquito and parasite (Dong et al., 2009). Studies also suggest that nutritional deprivation may have a direct impact...
on the gut oocysts development and mosquitoes’ reproductive outcome, although the mechanism is yet to unravel (Liu et al., 2013; Yang et al., 2019). Our observation of early induction of transcripts regulating gut-specific nutritional homeostasis, such as Folliculin, Trehalase, and Sterol carrier suggested that maturing oocysts may cause host nutritional resources imbalance (Baba et al., 2006; Schekman, 2013; Shukla et al., 2015). However, the delayed elevation of gut immune transcripts may counterbalance the negative impact of rapidly exiting sporozoites into the hemolymph. Thus, disrupting this relationship may favor the development of new tools to numb parasites either by delimiting the nutritional demand and/or enhancing gut immunity (Shea-Donohue et al., 2017).

Once left the gut epithelium, millions of fcSPZ directly encounter and are cleared by hemocytes, but the molecular nature of this interaction is not fully understood (Jull Hillyer et al., 2003; Jaramillo-Gutierrez et al., 2009). Our data suggest a major shift in transcript abundance, especially encoding structural and catabolic activity associated proteins, indicated the hyperactivity of hemocytes to control Plasmodium sporozoites population. The enriched number of hemocyte-specific immune transcripts and their time-dependent elevated expression together indicated that a hyper immunity of hemocytes is essential for the majority of sporozoite clearance. Although the reason for how this hyper-response is achieved by hemocyte is unclear, however, our observation of increased expression of REL compared to NOS against early P. vivax oocysts supports the idea that pre-immune activation of hemocyte may exist (De et al., 2018; Kwon and Smith, 2018). An indirect anti-Plasmodium response via hemocyte and REL immune signaling have also been suggested in the An. gambiae, P. berghoi, and P. falciparum model (Hillyer and Estévez-Lao, 2010). Surprisingly, however, it remains unclear how a fraction of fcSPZ succeeds in avoiding the hyperimmune response of hemocytes and invade salivary glands.

Our RNA-Seq data analysis and annotation of salivary-parasite interactions indicated that Plasmodium sporozoites not only impair the metabolic machinery but also enrich nucleic acid binding transcriptional activities related proteins. For successful blood meal acquisition, salivary glands release a cocktail containing nucleic acid binding factors such as Apyrase (King et al., 2011), D7 family proteins (Calvo et al., 2006), and nucleotide transferase (Dhar and Kumar, 2003; Dixit et al., 2009; Vogt et al., 2018). Thus, it is plausible to hypothesize that Plasmodium infection may enhance host-seeking behavioral activities by stimulating the expression of nucleic acid binding factors (Chen et al., 2008). Additionally, increased expression of salivary immune transcripts indicated that an active local immune response is essential to restrict salivary invaded sporozoite (IS) populations.

As a proof of concept, our data suggested that even after mounting an effective tissue-specific immuno-physiological response, mosquitoes fail to disrupt the Plasmodium sporogonic cycle. Hence, we propose that Plasmodium parasites are clever enough to dodge the mosquito responses by wise manipulation of their own molecular architecture. Retrieval of a large pool of P. vivax transcripts originating from distinct developmental stages provided an opportunity to anticipate the molecular dynamics facilitating its survival. An initial observation of more than 50% transcripts encoding hypothetical proteins indicated that a deep understanding of the P. vivax sporogonic cycle is still obscure. However, an observation of a significant difference in the molecular repertoire of

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**Fig. 7.** Representative catalog of RNA-Seq identified P. vivax-specific transcripts dominantly expressing in distinct mosquito tissues.
stage-specific *P. vivax* genes further strengthens our hypothesis that the *Plasmodium* parasite has the unique ability to misguide the track and trap system of mosquitoes.

5. Conclusion

For its successful survival and transmission, at every stage of development inside the mosquito host, *Plasmodium vivax* negotiates multiple tissues. Independent studies targeting individual tissues have been valuable (Roth et al., 2018; Boonkaw et al., 2020), but conceptually, we have unresolved major questions about how a mosquito’s tissue-specific actions manage the challenge of *Plasmodium* infection and/or how *Plasmodium* manages to avoid tissue-specific responses. For the first time, we demonstrate and establish that *P. vivax* follows a smart strategy of genetic makeup change to misguide and evade, even a highly sophisticated immune barrier of each tissue. We hypothesize and propose that an unharmed tissue-specific molecular wave of negotiations and actions by genetic changes benefits *P. vivax* successful development and transmission in the mosquito host. Further establishing a functional correlation may lead to the identification of mosquitoes as well as *P. vivax* specific crucial genetic factors for target selection and designing a new molecular strategy for malaria control.

Data deposition

The sequences of the individual tissue samples were submitted to the NCBI with accession numbers SRR8580009 (MG-Inf 8-10D); SRR8580010 (MG-C); SRR8580011 (MG-Inf 3-6D); SRR8501353 (HC-C); SRR8501352 (HC-PI); SRR8476334 (SG-PI); SRR8476333 (SG-C).

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Authors’ contribution

ST, SK, CC, KCP, VP, RKD conceived scientific hypothesis and designed the experiments, ST, SK, CC performed tissue i.e. midgut, salivary gland, and hemocyte specific experiments, DS, PS, TT, JR, TDD, DSR contributed to design and performing the experiments, data acquisition, writing, and editing; ST, SK, CC, KCP, VP, RKD data analysis and interpretation, data presentation, contributed reagents/materials/analysis tools, wrote, reviewed, edited, and finalized MS. All authors read and approved the final manuscript.

Authors summary

Malaria transmission dynamics are heavily influenced by mosquito-parasite interactions. When passing through tissue-specific barriers, *Plasmodium* has to compromise by losing its population, but the genetic relationship is unknown. To win the developmental race, *Plasmodium* needs to overcome two important immuno-physiological barriers. The first one accounts for an indirect 24–30 h long pre-invasive gut-microbe-parasite interaction in the gut lumen. The second one follows a direct post-gut invasive 14–18 days long interaction with the midgut, hemocyte, and salivary glands. During the pre-invasive phase of interaction, we showed that *Plasmodium vivax* follows an immunosuppression strategy by restricting microbial growth in the gut lumen (Sharma et al., 2020). Here, we demonstrate that successful developmental progression by *P. vivax* is accompanied by the manipulation of tissue-specific metabolic responses and altering its genetic makeup. This strategy not only clears off the multifaceted mosquito’s tissue-specific immune responses but also favors *Plasmodium*’s survival and transmission. Comprehending this tissue-specific interaction between host and parasite at the molecular level could provide a new tool to intervene in the *Plasmodium* life cycle within a vector.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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