Long read single cell RNA sequencing reveals the isoform diversity of *Plasmodium vivax* transcripts

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Abstract

*Plasmodium vivax* infections often consist of heterogenous populations of parasites at different developmental stages and with distinct transcriptional profiles, which complicates gene expression analyses. The advent of single cell RNA sequencing (scRNA-seq) enabled disentangling this complexity and has provided robust and stage-specific characterization of *Plasmodium* gene expression. However, scRNA-seq information is typically derived from the end of each mRNA molecule (usually the 3’-end) and therefore fails to capture the diversity in transcript isoforms documented in bulk RNA-seq data. Here, we describe the sequencing of scRNA-seq libraries using Pacific Biosciences (PacBio) chemistry to characterize full-length *Plasmodium vivax* transcripts from single cell parasites. Our results show that many *P. vivax* genes are transcribed into multiple isoforms, primarily through variations in untranslated region (UTR) length or splicing, and that the expression of many isoforms is developmentally regulated. Our findings demonstrate that long read sequencing can be used to characterize mRNA molecules at the single cell level and provides an additional resource to better understand the regulation of gene expression throughout the *Plasmodium* life cycle.

Author summary

Single-cell RNA-sequencing is a valuable tool for identifying gene expression differences among cells present in one sample. However, scRNA-seq data is usually generated by sequencing the 3’ end of mRNA molecules after poly-A capture, which complicates assigning reads to specific genes for organisms with poorly annotated UTRs, and prevents identifying differences in isoform expression. Here, we utilize a modified version of 10X scRNA-seq technology to characterize full-length transcripts using PacBio sequencing from both sporozoite and blood stages of *Plasmodium vivax*. These data allow us to predict full-length, stage-specific, transcripts for *P. vivax*, as well as to identify variations in UTR usage throughout the *P. vivax* life cycle.
**Introduction**

*Plasmodium vivax* is the second most common cause of human malaria worldwide and was responsible for 4.5 million clinical cases of malaria in 2020 [1]. Despite these numbers, *P. vivax* research lags behind that of *P. falciparum*, in part due to the difficulty to propagate the parasites in vitro [2,3]. Genomic techniques, such as genome [4–9] and transcriptome [10–12] sequencing, have improved our knowledge of *P. vivax* biology, but are hindered by the complexity of most blood-stage *P. vivax* infections: multiple genetically-distinct parasites are often simultaneously present in one infection and, due to the lack of, or incomplete, sequestration of *P. vivax* stages, all intraerythrocytic developmental stages concurrently circulate in the blood. Experimental infections of non-human primates using monkey-adapted strains of *P. vivax* [13,14] provide a robust system to study the regulation of all blood stages in vivo using monoclonal and well-characterized parasites. However, the simultaneous presence of multiple developmental stages in the blood, each with their own regulatory profiles, remains a major challenge. Short-term *ex vivo* cultures [10,11] and statistical inferences of the stage composition [10–12,15,16] have been used to circumvent this issue but suffer from limited resolution and possible artefacts.

Characterization of the gene expression of single cells (scRNA-seq) provides an elegant alternative and has been successfully applied to different *Plasmodium* species [17–23], including *P. vivax* [24,25]. However, many scRNA-seq assays rely on the capture and sequencing of the 3’ ends of polyadenylated transcripts [26,27] and, consequently, only a short portion of each transcript is sequenced and the data generated provide little information about transcript isoforms and alternative splicing [11,12,15,28]. Additionally, it can be difficult to assign one signal to a specific gene since scRNA-seq reads typically derive from the 3’ untranslated regions (3’-UTRs) which are incompletely annotated in *P. vivax* [11,12,15,24,28].

Full-length isoform sequencing (iso-Seq) using long-read technologies, such as PacBio, enables reliable characterization of gene isoforms and their UTRs [29–35]. This method has also been applied to single cells for identifying cell-type specific isoforms in a high throughput manner [30,34,35] and improving analysis of subsequent scRNA-seq data [34].

Here we combine long- and short read sequencing of scRNA-seq libraries to characterize *P. vivax* isoforms throughout the intraerythrocytic life cycle as well as in sporozoites. The standard short read Illumina sequencing of scRNA-seq libraries provides a robust description of the developmental stage of each *P. vivax* parasite captured, while sequencing the same mRNA molecules using PacBio long reads enables characterizing full-length transcripts present in these cells. The data generated allow to better annotate the 5’- and 3’-UTRs of *P. vivax* genes, to comprehensively characterize the transcripts expressed, and to identify stage-specific isoforms. Overall, our results demonstrate that single cell long read sequencing, in conjunction with short read sequencing, provides a robust method for comprehensively characterizing mRNA sequences at the single cell level and identifying isoforms involved in the regulation of specific *Plasmodium* stages.

**Results and discussion**

Characterization of single cell *P. vivax* blood-stage and sporozoite transcriptomes

We obtained blood-stage parasites from two *Saimiri boliviensis* monkeys infected with the Chesson strain of *P. vivax* and prepared 10X Genomics 3’-end single cell RNA sequencing (scRNA-seq) libraries after enrichment of infected red blood cells (see Material and Methods). After generating 57,550,235–63,399,045 short reads per sample, we successfully mapped 72–
74% of all reads to the *P. vivax* P01 genome sequence [36]. While full-length mRNAs are captured and converted into cDNA in the 10X droplets, only the 3’-end of each transcript is sequenced due to the cDNA fragmentation occurring during library preparation (Fig 1). All 3’-end scRNA-seq reads should therefore derive from the last ~300 bp of the transcripts (see e.g., S1 Fig). However, only 27–33% of the reads mapped within annotated *P. vivax* genes and 47–52% mapped more than 500 bp away from any annotated gene (S1 Table). These results are consistent with previous analyses [24] and highlight that many genes and/or their 3’-UTRs remain incompletely annotated in the *P. vivax* genome. After removing PCR duplicates and stringent quality control filters, we obtained information about 949 and 1,807 single cell blood-stage transcriptomes, each characterized by more than 5,000 unique reads (Table 1). To characterize which blood stages were present in each infection, we analyzed these transcriptomes using principal component analysis and showed that several asexual and sexual developmental stages were present in both samples, although with some differences in their relative proportions (S2 Fig).

We also prepared two scRNA-seq libraries from salivary gland sporozoites dissected from *Anopheles stephensi* and *Anopheles freeborni*, respectively, fed on *Saimiri* monkeys infected...
with *P. vivax* parasites. Out of the 63,039,836 and 79,688,059 reads generated from each library, 3–7% mapped to the *P. vivax* genome sequence, similar to the numbers obtained previously for *P. berghei* sporozoites [21]. 25–26% of those reads mapped within annotated *P. vivax* genes and 58–59% mapped more than 500 bp away from any gene (S1 Table). After removing PCR duplicates and stringent quality control filters, we obtained information about 2,609 and 2,363 single cell transcriptomes, each characterized by more than 250 unique reads (Tables 1 and S1). We used a lower threshold for the sporozoite analysis due to the lower number of *P. vivax* reads recovered as a result of the overwhelming presence of mosquito RNA (but the cutoff is comparable to those previously used for analyzing sporozoite scRNA-seq data [21]). In contrast to the heterogeneity of the blood stage parasites, these salivary gland sporozoites formed a relatively homogeneous population (S2 Fig).

**Long read sequencing of scRNA-seq libraries provides full-length transcript information**

Since only the 3’-end of each transcript is usually sequenced in scRNA-seq experiments, it is sometimes difficult to assign the scRNA-seq reads to a specific gene, or to identify signals derived from different gene isoforms [24]. We therefore used cDNA from the same four 10X scRNA-seq libraries before fragmentation to generate full-length isoform sequences using the PacBio chemistry (see Fig 1 and Material and Methods for details).

From each blood sample, we generated 3,390,270 and 3,570,838 circular consensus sequences (CCS) that each derived from at least 10 passes of sequencing of an individual cDNA molecule. 50–64% of these sequences carried a 10X barcode matching one of the cells characterized by Illumina sequencing, and >99% of those sequences mapped unambiguously to the *P. vivax* genome (Tables 1 and S1). After removal of PCR duplicates, we ended up with 1,688,745 and 1,523,456 unique reads (each derived from a unique mRNA molecule) for further analysis (Table 1). While the Illumina and PacBio library preparations diverged after cDNA amplification, the same barcoded cDNA molecules were used for both experiments (Fig 1) and the numbers of reads obtained with each technology for each individual cell were highly correlated (S3 Fig).

For the sporozoite samples, we generated 2,578,778 and 1,935,640 circular consensus sequences. 5–8% of these sequences carried a 10X barcode matching one of the cells characterized by Illumina sequencing, and 50–65% of those sequences mapped unambiguously to the *P. vivax* genome. After removal of PCR duplicates and additional QCs, we ended up with 106,531 and 40,977 unique reads respectively for further analysis (Tables 1 and S1). Similar to the pattern observed for the blood stage samples, the numbers of reads obtained by Illumina and PacBio sequencing for each sporozoite were highly correlated (S3 Fig) although, due to the lower sequencing depth, the overall number of cells with PacBio information was limited to 2,449 and 1,670.

We then summarized these mapped reads into *P. vivax* transcripts and, to avoid including sequences that may represent partially degraded molecules or technical artefacts, we only considered predicted transcripts represented by more than 10 PacBio unique reads in both samples of the same type (e.g., in both blood-stage samples). Overall, we identified a total of 9,982 transcripts from blood stage parasites and 784 from sporozoites (Tables 1 and S2), with an average transcript length of 2,432 bp from blood stages and 2,054 bp from sporozoites and ranging from 100 to 20,506 bp and from 197 to 6,809 bp, respectively. (Note that the transcript prediction algorithm occasionally leads to artefacts when transcripts in the same orientation overlap, which seems to be the case for the 20 kb transcript).
Most transcripts encode proteins corresponding to the genome annotation

Out of the 9,982 blood stage transcripts, 8,550 transcripts were predicted to encode more than 100 amino acids and 5,368 of those (63%) were predicted to encode a full-length protein (including a start and stop codon) (S1 Table). The median length of these full-length protein sequences was 291 amino acids, significantly smaller than the length of the protein sequences annotated in the latest version of the P01 P. vivax genome [36] (median of 401 amino acids) (S4 Fig). This discrepancy is possibly due to the lower 10X reverse transcriptase processivity that may hamper synthesis of extended cDNA molecules and prevent recovery of transcripts from long genes (note that most full-length transcripts obtained from PacBio sequencing match the annotated reference, see below). Of the 784 sporozoite transcripts, 563 transcripts were predicted to encode more than 100 amino acids and 344 of those (60%) were predicted to encode a full-length protein (including a start and stop codon), with a median length of 267 amino acids (S4 Fig and S1 Table).

We then compared the protein sequences predicted from these full-length transcripts to all P. vivax protein sequences annotated in the P01 genome (see S5 Fig, Materials and Methods). Most transcripts were predicted to encode protein-coding sequences highly similar to those annotated in the genome: 4,553 of the blood-stage transcripts (84.8%) and 254 of the sporozoite transcripts (73.8%) were identical to an annotated P. vivax protein-coding sequence for >90% of their length (S6 Fig, see e.g., Fig 2A). Additionally, 378 blood-stage (7.1%) and 23 sporozoite transcripts (6.7%) partially matched known protein-coding sequences and were identical over 50–90% of their sequence. While some of these transcripts could represent instances where the current gene annotation is possibly incorrect (see e.g., Fig 2B), in 253 of these cases (63%) another isoform in our dataset matched the entire annotated protein-coding sequence, suggesting that these differences represent incomplete annotations of transcript isoforms rather than incorrect annotations (see also below). One example is the cytochrome b5-like heme/steroid binding protein (PVP01_0716500) that is transcribed as annotated in the genome by blood-stage parasites but, due to an alternative start site, is transcribed into a shorter mRNA by sporozoites, resulting in a shorter predicted protein (Figs 2C and 57).

Finally, 436 blood-stage (8.1%) and 67 sporozoite transcripts (19.5%) aligned over less than 50% of their length to annotated protein sequences but those transcripts were very short (less than 200 amino acids) and/or had low support from the PacBio reads and likely represented artifacts or fragmented transcripts. The complete list of predicted transcripts, their sequence and their identity to reference transcripts is presented in S2 Table.

Most P. vivax transcripts have extended UTRs that often contain introns

Even for the transcripts highly similar to the annotated protein-coding genes, the PacBio sequences add novel information by providing a detailed description of the 5'- and 3'-UTRs which, despite recent efforts [28], remain incompletely annotated in P. vivax. Consistent with previous reports [11,15], our data showed extensive UTRs in many genes (Fig 3). We observed that 5' - and 3'-UTRs were roughly similar in length in transcripts expressed by blood-stage parasites, while 5'-UTRs were, on average, slightly longer than 3'-UTRs in sporozoite transcripts (745 vs 731 bp in blood stages [p-value = 0.3149] and 762 vs 650 bp in sporozoites [p-value = 0.0076]). To evaluate how this improved characterization of UTRs would affect future analyses of scRNA-seq data, we compared the percentage of scRNA-seq reads mapped to the currently annotated P. vivax genes with the percentage mapped to annotations supplemented by our predicted isoforms (available in S1 Data). While only 27–33% of the scRNA-seq reads mapped within previously annotated genes for the blood stage parasites and 25–26% for the sporozoites, including the PacBio predictions raises these figures to 69–77% and 65–70%,
Fig 2. Examples of full-length isoform sequencing using PacBio and resulting transcript predictions. (A) Example of protein-coding transcripts matching the current gene annotations. The figure shows 5.7 kb of chromosome 13 containing two annotated *P. vivax* genes, the ubiquitin fusion degradation protein 1 (PVP01_1330900) and an ATP synthase-associated protein (PVP01_1331000). The blue horizontal bars at the bottom shows the annotations for these genes in plasmoDB v54, while the top panel shows the PacBio reads mapping to this locus (each red horizontal line is a unique read mapped to the positive strand with the grey lines indicating spliced introns). Note that, while the PacBio reads support a shorter 3'-UTR than annotated for PVP01_133100, the predicted protein coding sequences are identical to the ones annotated. (B) Example of protein coding transcript differing from the current gene annotation. The figure shows 5 kb of chromosome 13 surrounding the rhoptry-associated protein 1 (RAP1, PVP01_1338500). The PacBio reads (mapped to the negative strand and displayed in blue) support the presence of an unannotated intron (red box), leading to additional predicted coding sequences upstream of this intron, and a different protein than annotated in the genome (thick blue bars at the bottom). (C) Example of two isoforms with different predicted protein coding sequences. The top panel shows that blood-stage parasites express a transcript for cytochrome b5-like heme/steroid binding protein (PVP01_0716500) identical to the annotated protein-coding sequence (although with a shorter 5'-UTR). The middle panel shows that *P. vivax* sporozoites express this gene from a different start site (red box) resulting in a shorter transcript and a different predicted protein. (Note also the presence of an unannotated, and alternatively spliced, intron in the 3'UTR).

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respectively (S1 Table). Reanalysis on an independent scRNA-seq dataset generated from another *P. vivax* blood stage infection (AMRU-I from [24]) confirmed these results, improving the mapping of reads to annotated transcripts from 30% to 88%.

While UTRs have not been extensively studied in *Plasmodium*, in many eukaryotes they contain important regulatory elements which often affect mRNA stability [37–41]. While the length of the 5’-UTR does not appear to be associated with the level of the transcript expression determined using the short reads ($p = 0.315$), the 3’-UTR length was negatively correlated with expression level ($p = 2.8 \times 10^{-6}$), although with a very low coefficient of correlation (Spearman’s Rho = -0.0798) (S8 Fig). In an attempt to identify regulatory elements in the UTRs, we compared the abundance of all possible 5-mers in the 5’-UTRs, 3’-UTRs and promoter regions but failed to detect significant enrichment (S9 Fig). More in-depth analyses will be required to understand the function of these UTRs but the data presented here will provide a solid foundation to implement these studies.

Interestingly, out of the 5,368 full-length blood stage protein-coding transcripts, 1,072 (20%) had at least one intron in the UTRs: 419 transcripts had at least one intron in the 5’-UTR, 568 in the 3’-UTR and 89 had introns in both UTRs (see e.g., S10 Fig). Of these, 322 transcripts had more than one intron in their UTRs. Of the 344 full-length sporozoite transcripts, 26 had introns in the UTRs, 12 had at least one intron in the 5’-UTR, 11 in the 3’-UTR and 3 had introns in both UTRs. Five of these transcripts had more than one intron in the 5’- or 3’-UTR. Remarkably, in blood-stage parasites the levels of expression of the transcripts with an intron in the UTR were, on average, 3-fold higher than those of genes with no UTR intron (Wilcoxon rank test, $p$-value $< 2.2 \times 10^{-16}$), suggesting that presence of introns in the UTR may be associated with increased mRNA stability in *P. vivax*. Furthermore, genes with introns in
their coding sequences were also three times more likely to have introns in the UTR ($\chi^2 = 206.78$, p-value $< 2.2e-16$), which may suggest that UTR splicing is mechanistically associated with splicing of coding sequences, possibly due to a more effective recruitment of the splicing machinery at those transcripts. To evaluate whether the presence of UTR introns was more frequent, at a specific developmental stage, we assigned each gene to the blood stage where it was most abundantly expressed. The proportion of genes with introns in their UTR was significantly different among stages ($\chi^2 = 26.141$, p-value $= 8.9 \times 10^{-6}$), with 11% and 18% of the genes most expressed in early and late trophozoites having an UTR intron, respectively, compared to only 3% and 8% of the genes expressed in female gametocytes and schizonts (note that genes with multiple isoforms were excluded from this analysis as it is difficult to robustly quantify the relative expression of isoforms). Finally, we tested whether genes containing introns in their UTRs disproportionally belonged to a specific pathway but failed to detect any gene ontology enrichment (FDR $< = 0.1$, S3 Table).

**Transcript isoforms are common in P. vivax and can be expressed in a stage-specific manner**

The 5,368 full-length protein coding transcripts derived from blood-stage parasites were transcribed from 2,869 genes: 1,687 genes (59%) were transcribed into a single isoform, while 1,182 (41%) showed evidence of multiple isoforms (and out of those, 719 genes were expressed in two isoforms, 247 in three and 216 were transcribed in four or more isoforms). Most isoforms ($n = 819, 70\%$) encoded the same protein sequence and differed only in their UTR length: 1,024 (87%) genes with isoforms differed in their 5'UTR length and 1,008 (85%) differed in the 3'UTR length, with 450 genes with isoforms differing in their UTR introns (Table 2, see also S10 Fig for an example). 363 genes showed evidence of isoforms that were predicted to encode for different proteins, due to an alternate protein coding start (299), alternative end (311), and/or exon skipping (67) (Table 2).

In sporozoites, the 344 full-length protein coding transcripts were transcribed from 238 genes: 211 genes (89%) were transcribed into a single isoform, while 27 (11%) showed evidence of multiple isoforms (20 genes were expressed in two isoforms and three in three isoforms and four in four or more). All but eight isoforms were predicted to encode the same protein. Five of these genes had an alternate coding start and six had an alternate coding stop resulting in the change in coding sequence, and all had an exon skip or truncation (Table 2).

Since the full-length transcript data derived from molecules characterized by scRNA-seq, these data provide a unique opportunity to preliminarily examine whether different isoforms were expressed at different stages of the parasite development. Using the 10X cell barcodes, we determined the developmental age of each cell using pseudotime (determined from the Illumina scRNA-seq data). We only considered in this analysis 636 genes that had two (or more) isoforms expressed in at least 50 individual cells each. 123 (19%) of these genes showed evidence of expressing isoforms according to the parasite development (S4 Table). These stage-specific isoforms included 62 and 35 genes with differences in 5'- or 3'-UTR length (see e.g., Figs 4 or S11A) and/or changes in coding sequences ($n = 40$) (S11B Fig). These findings suggest that *P. vivax* utilizes alternative start or termination of transcription as a means of transcriptional or translational regulation between stages.

| Table 2. Summary of isoform types from PacBio predictions. |
| Sample | No. of genes | Genes with >1 isoforms | Alt 5’-UTR | Alt 3’-UTR | Alt Coding Seq | Alt Coding Start | Alt Coding End |
|--------|--------------|------------------------|------------|------------|----------------|-----------------|---------------|
| Blood  | 2,870        | 1,182                  | 1,024      | 1,008      | 363            | 299             | 311           |
| Sporozoites | 238        | 27                     | 8          | 9          | 8              | 5               | 6             |

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Conclusion

Long read PacBio sequencing coupled with short read Illumina sequencing of single cell RNA-seq libraries provide a robust method to detect and characterize full-length transcripts and to identify mRNA isoforms expressed at different times. These simple modifications of existing laboratory protocols for generating 10XGenomics scRNA-seq can be easily applied to any eukaryotic cells and could be invaluable to examine variations in gene expression in organisms that are difficult to study in the laboratory and display a complex life cycle. This study also yielded novel insights on the variations and regulation of gene expression in \textit{P. vivax}, which provides a solid framework to improve our understanding of \textit{P. vivax} biology and regulation. This study notably highlighted the presence of extensive and incompletely annotated UTRs, and of ubiquitous UTR introns, that were sometimes expressed in a stage-specific manner. In particular, the observation that different stages used different start sites for transcribing the...
same gene provides new insights on how gene expression may be regulated throughout the Plasmodium life cycle. In addition, the extensive variations observed in 3'-UTR lengths and the presence of introns in these regions might indicate important roles in mRNA stability or translation efficiency and, consequently, in the regulation of Plasmodium parasite biology.

Materials and methods

Ethics statement

All animal procedures were conducted in accordance with the National Institutes of Health (NIH) guidelines and regulations [42], under approved protocols by the National Institute of Allergy and Infectious Diseases (NIAID) Animal Care and Use Committee (ACUC) (Animal study NIAID LMVR15). Animals were purchased from NIH-approved sources and transported and housed according to Guide for the Care and Use of Laboratory Animals [42].

Animal studies and sample collection

We infected two splenectomized Saimiri boliviensis monkeys with the Chesson strain of P. vivax using parasitized erythrocytes from cryopreserved stocks. Once they developed a parasitemia >0.1%, we collected 1 mL of blood from the femoral vein of each monkey after anesthesia with 10 mg/kg of ketamine and processed the blood samples on MACS LS columns as previously described [24]. Note that this enrichment procedure, that relies on the paramagnetic properties of hemozoin generated by maturing blood-stage parasites, fails to capture ring stage parasites.

Two blood samples from two additional Saimiri boliviensis monkeys coinfected with a NIH-1993 clone [24] and the Chesson P. vivax strain were used for membrane feeding of Anopheles stephensi and Anopheles freeborni. Salivary glands sporozoites were collected from each feeding at 21 days post-feed: 50 female mosquitoes were anesthetized on ice and their salivary glands dissected in PBS under a stereomicroscope. The salivary glands were transferred to a low-retention tube (Protein LoBind Tube; Eppendorf) containing PBS, homogenized with a disposable pestle, spun down, washed, resuspended, and quantified.

10X single cell RNA-sequencing library preparation and sequencing

An estimated 3,000 infected red blood cells or sporozoites from each sample were loaded onto a 10X Chromium controller to prepare scRNA-seq libraries according to the manufacturer’s instructions. We then generated, from each library, 57–75 million paired-end reads using an Illumina NovaSeq.

In addition, an aliquot of the cDNA prior to fragmentation was amplified by eight additional cycles of PCR before preparation of a PacBio library using the SMRTBell Express kit 2.0. We then generated 196–328 million reads from each library using a PacBio Sequel II.

Short read analysis and single cell characterization

Following Illumina sequencing, the short reads were processed as described previously [24]. Briefly, we mapped all reads to the P01 P. vivax genome [36] using Hisat2 [43] with the default parameters except for a maximum intron length of 5,000 bp. We then removed PCR duplicates by identifying reads with identical barcode, unique transcript identifier (UMI) and mapping coordinates. We assigned each unique read to i) a cell based on its barcode and ii) a 500 bp window based on its genomic position. Only cells defined by more than 5,000 unique reads in blood stages and 250 unique reads in sporozoites were further analyzed. Count matrices and principal component analysis was performed using in-house scripts.
Long read analysis

The entire analytical pipeline for processing and analyzing the long-read data is described in S5 Fig. Briefly, the raw PacBio reads were first collapsed into circular consensus sequencing (CCS) using smrtanalysis [44] and only CCS supported by more than 10 passes were considered for further analysis. We then compared the 10X barcodes of each CCS reads with those obtained after Illumina sequencing and kept all reads matching the barcodes of one of the cells characterized by more than 5,000 unique Illumina reads. To account for the higher error rate of PacBio sequencing we allowed up to one nucleotide mismatch in the barcode sequence. After trimming the 10X adapters, 10X barcodes and the polyA tails using custom scripts, we mapped all CCS reads to the \textit{P. vivax} P01 genome using minimap2 [45] using the cdna parameters and a k of 14. PCR duplicates were removed as described above. Only reads mapped to the P01 genome over 50% of their length were kept for further analysis.

Transcript and protein identification

Transcript prediction was performed separately for each sample utilizing Stringtie2 [46,47]. Mapping files were divided by forward and reverse reads, and into single and multiple exon reads. Each of the four files was run separately using Stringtie2 long read default parameters. Only transcripts supported by more than 10 reads in each sample were considered for further processing (separately for the blood-stage and sporozoite samples). Transcript predictions were then compared and collapsed across samples into a single gtf for the blood-stage and sporozoite samples using gffcompare and custom scripts. We then used Transdecoder [48] to identify putative protein coding sequences from the predicted transcript sequences. Finally, we compared the predicted protein sequences to those annotated in the most current \textit{P. vivax} P01 genome (v54) using BlastP and custom scripts.

Stage-specific transcript analysis

To identify isoforms expressed in a stage-specific manner, we first assigned each PacBio read to a specific isoform with BLAT, using all predicted stringtie transcripts (including non-coding and incomplete transcripts) and considering the match with the greatest overall identity (and only considering reads aligned with >90% identity as aligned). We then identified the pseudotime of the cell expressing this transcript by matching the GEMS barcode to the Illumina data. For each gene that had two (or more) isoforms, each expressed in more than 50 cells, we then tested for qualitative differences in isoform expression associated with the parasite development by comparing the pseudotime ranks of the cells expressing the first isoform with the pseudotime ranks of the cells expressing the second isoform using a Kolmogorov-Smirnov test.

Supporting information

S1 Fig. Example of Illumina and Pacbio data generated from the same scRNA-seq library. The top panel shows the data generated by Illumina sequencing and displays typical peaks corresponding to the 3'-end of each expressed transcript. The middle panel shows data generated, from the same mRNAs, using PacBio sequencing and illustrates how generating full-length transcripts improves interpretation of scRNA-seq data for organisms with incomplete gene annotations. (TIF)

S2 Fig. Principal component analysis showing the relationships among individual parasite cells characterized by scRNA-seq (using the Illumina data). Top row: each dot is a single cell
blood stage transcriptome, and is displayed based on its gene expression profile and colored according to the expression of stage markers (with cells being assigned to marker with the highest expression): red—early trophozoites, green—late trophozoites, purple—schizonts, turquoise—female gametocytes. Note that few ring stage parasites are included (if any) due to the enrichment method used (see Material and Methods). Bottom row: single cell sporozoite data.

S3 Fig. Correlation between the number of reads obtained by Illumina and PacBio. The scatterplot shows the correlation between the number of Illumina reads (x-axis) and PacBio reads (y-axis) obtained from each cell (individual black dots). Each panel represents the data for a different sample.

S4 Fig. Predicted protein length distributions. Distribution of the length (in amino acids) of the protein-coding sequences predicted from the PacBio transcripts (in red) and of the protein-coding sequences annotated in the P01 P. vivax genome (in blue).

S5 Fig. Summary of the bioinformatic pipeline used for processing the PacBio reads and for predicting transcripts.

S6 Fig. Comparison of the predicted proteins with annotated proteins. Distribution of the percentage alignment (x-axis) of the predicted protein coding sequences with the most similar protein sequence annotated in the P01 genome. Left: blood-stage transcripts. Right: sporozoite transcripts. (Note that the y-axis is cut and the right-most bars (perfect matches) go to 400 and 250 for the left and right panels, respectively).

S7 Fig. Stage-specific expression of the isoforms of the cytochrome b5-like heme/steroid binding protein (PVP01_0716500). The left PCA shows blood-stage and sporozoite parasites jointly displayed according to their gene expression profiles. The right figure shows the same PCA, with each parasite colored based on the cytochrome b5-like heme/steroid binding protein isoform expression: red—“sporozoite” isoform, blue—“blood-stage” isoform.

S8 Fig. UTR length and expression. Correlation between the length of a transcript’s UTR (x-axis, in bp) and its level of expression determined by Illumina data (y-axis). Note that only genes expressing a single isoform are included in this analysis.

S9 Fig. UTR kmer analysis. Comparison of the abundance of all 5-mers in gene promoters, 5'-UTRs and 3'-UTRs. Note that most motifs with different abundance (i.e., deviating from the diagonales) are either repeated sequences or encoding for a start codon (ATG).

S10 Fig. Example of a transcript with unannotated UTR introns. The figure shows PacBio reads (in blue) corresponding to the annotated mRNA for coenzyme Q-binding protein COQ10 homolog (PVP01_0113000) but with an unannotated intron in the 3'-UTR (red box) as well as, for a subset of the mRNAs, a second intron in the 5'-UTR (blue box).
S11 Fig. Examples of isoforms expressed in a stage-specific manner. Each panel shows the PacBio reads mapped to a selected locus and split in four groups according to the stage of the parasites they derived from: early trophozoites, late trophozoites, schizonts and female gametocytes (from top to bottom). (A) Early trophozoites express the Ham 1-like protein (PVP01_0316500) from a more upstream TSS than the other stages and the resulting transcripts have a longer 5’-UTR containing five introns (red box). (B) The isoforms expressed from suppressor of kinetochore protein 1 (PVP01_1105000) result into different predicted protein coding sequences: some transcripts expressed exclusively in early trophozoites retain the third intron (red box) leading to a different open reading frame (blue bars at the bottom).

S1 Table. Sequencing and transcript predictions results. Table listing numbers from each step of sequencing analysis and transcript prediction.

S2 Table. List of all predicted transcripts. Table listing all predicted transcripts, their nucleotide sequence, amino acid sequence (if applicable), most similar PVP01 gene name and description (if applicable), their UTR lengths, and the number of introns.

S3 Table. Results of Gene Ontology analysis. Top hits from GO analysis.

S4 Table. List of predicted genes with differentially expressed isoforms.

S1 Data. Gene predictions. gtf file of all predicted protein coding transcripts combined with the current reference.

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References

1. World Malaria Report 2021. Geneva: World Health Organization; 2021. Licence: CC BY-NC-SA 3.0 IGO.
2. Anstey NM, Guerra CA, Yeung S, Price RN, Tjitra E, White NJ. Vivax Malaria: Neglected and Not Benign. Am J Trop Med Hyg. 2007 Dec 1; 77(6 Suppl):79–87. PMID: 18165478
3. Bermúdez M, Moreno-Pérez DA, Arévalo-Pinzón G, Curiñor H, Patarroyo MA. Plasmodium vivax in vitro continuous culture: the spoke in the wheel. Malar J. 2018 Dec 17;17(1):301. https://doi.org/10.1186/s12936-018-2456-5 PMID: 30126427
4. Carlton JM, Adams JH, Silva JC, Bidwell SL, Lorenzo H, Caler E, et al. Comparative genomics of the neglected human malaria parasite Plasmodium vivax. Nature. 2008 Oct 4; 455(7214):757–63. https://doi.org/10.1038/nature07327 PMID: 18643361
5. Neatsey DE, Galinsky K, Jiang RHY, Young L, Sykes SM, Siff S, et al. The malaria parasite Plasmodium vivax exhibits greater genetic diversity than Plasmodium falciparum. Nat Genet. 2012 Sep;44(9):1046–50. https://doi.org/10.1038/ng.2375 PMID: 22863733
6. Chan ER, Barnwell JW, Zimmerman PA, Serre D. Comparative Analysis of Field-Isolate and Monkey-Adapted Plasmodium vivax Genomes. Escalante AA, editor. PLoS Negl Trop Dis. 2015 Mar 13;9(3):e0003566. https://doi.org/10.1371/journal.pntd.0003566 PMID: 25768941
7. Pearson RD, Amato R, Auborn S, Miotto O, Almagro-Garcia J, Amaratunga C, et al. Genomic analysis of local variation and recent evolution in Plasmodium vivax. Nat Genet. 2016 Aug;48(8):959–64. https://doi.org/10.1038/ng.3599 PMID: 27348299
8. Auborn S, Benavente ED, Miotto O, Pearson RD, Amato R, Grigg MJ, et al. Genomic analysis of a pre-elimination Malaysian Plasmodium vivax population reveals selective pressures and changing transmission dynamics. Nat Commun. 2018 Dec;9(1):2585. https://doi.org/10.1038/s41467-018-04965-4 PMID: 29968722
9. Popovici J, Friedrich LR, Kim S, Bin S, Run V, Lek D, et al. Genomic Analyses Reveal the Common Occurrence and Complexity of Plasmodium vivax Relapses in Cambodia. Miller LH, editor. mBio. 2018 Mar 7;9(1):e01888–17.
10. Bozdach Z, Mok S, Hu G, Imwong M, Jaidee A, Russell B, et al. The transcriptome of Plasmodium vivax reveals divergence and diversity of transcriptional regulation in malaria parasites. Proc Natl Acad Sci. 2008 Oct 21;105(42):16290–5.
11. Zhu L, Mok S, Imwong M, Jaidee A, Russell B, Nosten F, et al. New insights into the Plasmodium vivax transcriptome using RNA-Seq. Sci Rep. 2016 Apr;6(1):20498. https://doi.org/10.1038/srep20498 PMID: 26980307
12. Kim A, Popovici J, Menard D, Serre D. Plasmodium vivax transcriptomes reveal stage-specific chloroquine response and differential regulation of male and female gametocytes. Nat Commun. 2019 Dec;10(1):371. https://doi.org/10.1038/s41467-019-08312-z PMID: 30670687
13. Collins WE. Nonhuman primate models. II. Infection of Saimiri and Aotus monkeys with Plasmodium vivax. Methods Mol Med. 2002;72:85–92. https://doi.org/10.1385/1-59259-271-6:85 PMID: 12125161
14. Young Martin D., Porter James A., Johnson Carl M. Plasmodium vivax Transmitted from Man to Monkey to Man. Science. 1966 Aug 26; 153(3739):1006–7. https://doi.org/10.1126/science.153.3739.1006 PMID: 4958954
15. Kim A, Popovici J, Vantaux A, Samreth R, Bin S, Kim S, et al. Characterization of P. vivax blood stage transcriptomes from field isolates reveals similarities among infections and complex gene isoforms. Sci Rep. 2017 Dec 7;7(1):7761. https://doi.org/10.1038/s41598-017-07275-9 PMID: 28798400
16. Tebben K, Dia A, Serre D. Determination of the Stage Composition of Plasmodium Infections from Bulk Gene Expression Data. Langelier CR, editor. mSystems. 2022 Aug 30;7(4):e00258–22.
17. Reid AJ, Talman AM, Bennett HM, Gomes AR, Sanders MJ, Illingworth CJR, et al. Single-cell RNA-seq reveals hidden transcriptional variation in malaria parasites. eLife. 2018 Mar 27; 7:e33105. https://doi.org/10.7554/eLife.33105 PMID: 29580379

18. Poran A, Nötzel C, Aly O, Mencia-Trinchant N, Harris CT, Guzman ML, et al. Single-cell RNA sequencing reveals a signature of sexual commitment in malaria parasites. Nature. 2017 Nov; 551(7678):95–9. https://doi.org/10.1038/nature24280 PMID: 29094698

19. Howick VM, Russell AJC, Andrews T, Heaton H, Reid AJ, Natarajan K, et al. The Malaria Cell Atlas: Single parasite transcriptomes across the complete Plasmodium life cycle. Science. 2019 Aug 23; 365(6455):eaaw2619.

20. Brancucci NMB, De Niz M, Straub TJ, Ravel D, Sollelis L, Birren BW, et al. Probing Plasmodium falciparum sexual commitment at the single-cell level. Wellcome Open Res. 2018 Oct 17: 3:70–70. https://doi.org/10.12688/wellcomeopenres.14645.4 PMID: 30320226

21. Bogale HN, Pascini TV, Kanatani S, Sá JM, Wellems TE, Sinnis P, et al. Transcriptional heterogeneity and tightly regulated changes in gene expression during Plasmodium berghei sporozoite development. Proc Natl Acad Sci. 2021 Mar 9; 118(10):e2023438118.

22. Real E, Howick VM, Dahalan FA, Wittmer K, Cudini J, Andraid-Brown C, et al. A single-cell atlas of Plasmodium falciparum transmission through the mosquito. Nat Commun. 2021 Dec; 12(1):3196. https://doi.org/10.1038/s41467-021-23434-z PMID: 34045457

23. Ruberto AA, Bourke C, Merienne N, Obadia T, Amino R, Mueller I. Single-cell RNA sequencing reveals developmental heterogeneity among Plasmodium berghei sporozoites. Sci Rep. 2021 Dec; 11(1):4127. https://doi.org/10.1038/s41598-021-82914-w PMID: 33619283

24. Sá JM, Cannon MV, Caleon RL, Wellems TE, Serre D. Single-cell transcription analysis of Plasmodium vivax blood-stage parasites identifies stage- and species-specific profiles of expression. Duffy M, editor. PLoS Biol. 2020 May 4; 18(5):e3000711. https://doi.org/10.1371/journal.pbio.3000711 PMID: 32365102

25. Ruberto AA, Bourke C, Vantaux A, Maher SP, Jex A, Witkowski B, et al. Single-cell RNA sequencing of Plasmodium vivax sporozoites reveals stage- and species-specific transcriptomic signatures. Mireji PO, editor. PLoS Negl Trop Dis. 2022 Aug 4; 16(8):e0010633. https://doi.org/10.1371/journal.pntd.0010633 PMID: 35926062

26. Baran-Gale J, Chandra T, Kirschner K. Experimental design for single-cell RNA sequencing. Brief Funct Genomics. 2018; 17(4):8. https://doi.org/10.1093/bfgp/elx035 PMID: 29126257

27. Zheng GXY, Terry JM, Belgrader P, Ryvkin P, Bent ZW, Wilson R, et al. Massively parallel digital transcriptional profiling of single cells. Nat Commun. 2017 Apr; 8(1):14049. https://doi.org/10.1038/s41467-016-014049 PMID: 28091601

28. Siegel SV, Chappell L, Hostetler JB, Amarantunga C, Suon S, Böhme U, et al. Analysis of Plasmodium vivax schizont transcriptomes from field isolates reveals heterogeneity of expression of genes involved in host-parasite interactions. Sci Rep. 2020 Dec; 10(1):16667. https://doi.org/10.1038/s41598-020-73662-7 PMID: 33028892

29. Sharon D, Tilgner H, Grubert F, Snyder M. A single-molecule long-read survey of the human transcriptome. Nat Biotechnol. 2013 Nov; 31(11):1099–14. https://doi.org/10.1038/nbt.2705 PMID: 24108091

30. Tian L. Comprehensive characterization of single-cell full-length isoforms in human and mouse with long-read sequencing. 2021; 24. https://doi.org/10.1108/s13059-021-02525-6 PMID: 34763716

31. Shields EJ, Sorida M, Sheng L, Sieriebriennikov B, Ding L, Bonasio R. Genome annotation with long RNA reads reveals new patterns of gene expression and improves single-cell analyses in an ant brain. BMC Biol. 2021 Dec; 21(1):74. https://doi.org/10.1186/s12915-021-01188-w PMID: 34838024

32. Westoby J, Artemov P, Hemberg M, Ferguson-Smith A. Obstacles to detecting isoforms using full-length scRNA-seq data. Genome Biol. 2020 Dec; 21(1):74. https://doi.org/10.1186/s13059-020-01981-w PMID: 32293520

33. Rhoads A, Au KF. PacBio Sequencing and Its Applications. Genomics Proteomics Bioinformatics. 2015 Oct; 13(5):278–89. https://doi.org/10.1016/j.gpb.2015.08.002 PMID: 26542840

34. Healey HM, Bassham S, Crespo WA. Single-cell Iso-sequencing enables rapid genome annotation for scRNAseq analysis. Sanchez Alvarado A, editor. Genomics. 2022 Mar 3; 220(3):iyac017. https://doi.org/10.1038/s41467-021-01403-w PMID: 35143654

35. Gupta I, Collier PG, Haase B, Mahfouz A, Joglekar A, Floyd T, et al. Single-cell isoform RNA sequencing characterizes isoforms in thousands of cerebellar cells. Nat Biotechnol. 2018 Dec; 36(12):1197–202. https://doi.org/10.1038/nbt.4259 PMID: 30320766

36. Auburn S, Böhme U, Steinbiss S, Trimmeranto H, Hostetler J, Sanders M, et al. A new Plasmodium vivax reference sequence with improved assembly of the subtelomeres reveals an abundance of pir
37. Adjalley SH, Chabbert Christophe D, Klaus Bernd Pelechano Vicent Steinmetz Lars M. Landscape and Dynamics of Transcription Initiation in the Malaria Parasite Plasmodium falciparum. Cell Rep. 2016 Mar 15;(14):14. https://doi.org/10.1016/j.celrep.2016.02.025 PMID: 26947071

38. Breitbart RE, Andreadis A, Nadal-Ginard B. ALTERNATIVE SPlicing: A Ubiquitous Mechanism for the Generation of Multiple Protein Isoforms from Single Genes. Annu Rev Biochem. 1987 Jun 1; 56(1):467–95. https://doi.org/10.1146/annurev.bi.56.070187.002343 PMID: 3304142

39. Pavlovic Djuranovic S, Erath J, Andrews RJ, Bayguinov PO, Chung JJ, Chalker DL, et al. Plasmodium falciparum translational machinery condones polyadenosine repeats. eLife. 2020 May 29; 9:e57799. https://doi.org/10.7554/eLife.57799 PMID: 32469313

40. Li J, Cai B, Qi Y, Zhao W, Liu J, Xu R, et al. UTR introns, antisense RNA and differentially spliced transcripts between Plasmodium yoelii subspecies. Malar J. 2016 Dec; 15(1):30. https://doi.org/10.1186/s12936-015-1081-9 PMID: 26791272

41. Yeoh LM, Lee VV, McFadden GI, Ralph SA. Alternative Splicing in Apicomplexan Parasites. Sibley D, Garsin DA, editors. mBio. 2019 Feb 26; 10(1):e02866–18. https://doi.org/10.1128/mBio.02866-18 PMID: 30782661

42. Hawkins P, Morton DB, Burman O, Dennison N, Honess P, Jennings M, et al. A guide to defining and implementing protocols for the welfare assessment of laboratory animals: eleventh report of the BVAAWF/FRAME/RSPCA/UFAW Joint Working Group on Refinement. Lab Anim. 2011 Jan 1; 45 (1):1–13. https://doi.org/10.1258/la.2010.010031 PMID: 21123303

43. Kim D, Langmead B, Salzberg SL. HISAT: a fast spliced aligner with low memory requirements. Nat Methods. 2015 Apr; 12(4):6. https://doi.org/10.1038/nmeth.3317 PMID: 25751142

44. Bayega A, Fahiminiya S, Olkonomopoulos S, Ragoussis J. Current and Future Methods for mRNA Analysis: A Drive Toward Single Molecule Sequencing. Methods Mol Biol Clifton NJ. 2018; 1783:209–41. https://doi.org/10.1007/978-1-4939-7834-2_11 PMID: 29767365

45. Li H. Minimap2: pairwise alignment for nucleotide sequences. Bioinformatics. 2018 May 10; 34(18):7. https://doi.org/10.1093/bioinformatics/bty191 PMID: 29750242

46. Kovaka S, Zimin AV, Pertea GM, Razaghi R, Salzberg SL, Pertea M. Transcriptome assembly from long-read RNA-seq alignments with StringTie2. Genome Biol. 2019 Dec; 20(1):278. https://doi.org/10.1186/s13059-019-01910-1 PMID: 31842956

47. Pertea M, Kim D, Pertea GM, Leek JT, Salzberg SL. Transcript-level expression analysis of RNA-seq experiments with HISAT, StringTie and Ballgown. Nat Protoc. 2016 Sep; 11(9):1650–67. https://doi.org/10.1038/nprot.2016.095 PMID: 27560171

48. Haas BJ, Papanicolaou A, Yassour M, Grabherr M, Blood PD, Bowden J, et al. De novo transcript sequence reconstruction from RNA-seq using the Trinity platform for reference generation and analysis. Nat Protoc. 2013 Aug; 8(8):1494–512. https://doi.org/10.1038/nprot.2013.084 PMID: 23845962