RESEARCH

Dual host-parasite transcriptomes of apicomplexan Eimeria falciformis and its natural mouse host

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Abstract

First part title: Text for this section. **Second part title:** Text for this section.

Keywords: Parasite, apicomplexa, RNA-seq, transcriptome, life-cycle, interaction; article; author

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Introduction

Text and results for this section, as per the individual journal's instructions for authors.

Results

Text for this section ...

1 Experimental overview

Replicate average of transcripts as order of magnitude for *E. falciformis*/mouse are indicated in Table.... Columns in table represent mouse strains used in infection experiments. Rows represent timepoints post infection plus oocyst and sporozoite samples. The upper part of the table shows data for first infection, and oocyst and sporozoite data. The lower part shows data for challenge infection. Averages were calculated after sample exclusions (see Methods). For exact values, see Table 1.

Table 1 Experimental overview with number of read for parasite/mouse indicated.

Day, 1st infection	NMRI	C57BL/6	Rag1-/-
0 (control)	$10^4 / 10^8$	$10^2 / 10^8$	$10^2 / 10^7$
3	$10^5 / 10^8$	NA	NA
5	$10^7 / 10^8$	$10^5 / 10^7$	$10^5 / 10^7$
7	$10^{8} / 10^{7}$	NA	NA
Oocysts	10^8 / NA	NA	NA
Sporozoites	10^7 / NA	NA	NA
Day, 2nd infection			
0 (control)	$10^3 / 10^8$	NA	NA
3 `	$10^4 / 10^8$	NA	NA
5	$10^4 / 10^8$	$10^5 / 10^7$	$10^5 / 10^7$
7	$10^6 / 10^8$	NA	NA

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Kasemo et al. Page 2 of 11

2 mRNA abundance differences between different experimental groups

Table 2 mRNA abundance differences between different experimental groups.

Day post infection comparisons	Ef genes different (FDR≤0.01)	Mouse genes different (FDR≤0.01/0.05)
NMRI 0 vs NMRI 3 NMRI 0 vs NMRI 5 NMRI 0 vs NMRI 7 NMRI 3 vs NMRI 5 NMRI 3 vs NMRI 7 NMRI 5 vs NMRI 7 C57BL/6 0 vs C57BL/6 5 Rag1-/- 0 vs Rag1-/- 5	NA NA NA 111 1385 1895 NA NA	274 1736 2802 1 1407 873 914
First and second infection comparisons		
NMRI 3 1st vs NMRI 3 2nd NMRI 5 1st vs NMRI 5 2nd NMRI 7 1st vs NMRI 7 2nd C57BL/6 1st vs C57BL/6 2nd (day 5) Rag1-/- 1st vs Rag1-/- 2nd (day 5)	0 0 0 0 0	5 1 902 mouse mouse
Mouse strain comparisons		
NMRI vs C57BL/6 NMRI vs Rag1-/- C57BL/6 vs Rag1-/- Day post infection, parasite relevant comparisons	22 0 0	NA* NA* 356
Oocysts vs NMRI 3 Oocysts vs NMRI 5 Oocysts vs NMRI 7 Oocysts vs sporozoites Sporozoites vs NMRI 3 Sporozoites vs NMRI 5 Sporozoites vs NMRI 7	3310 3605 3085 3421 1663 1605 2473	NA NA NA NA NA NA

 $E.\ falciformis$ data overview of pairwise comparisons and number of genes with differently abundant mRNAs per comparison. NMRI followed by number indicates day post infection (e.g. NMRI3 = $E.\ falciformis$ genes from NMRI mouse day 3 post infection). Genes with Benjamini-Hochberg corrected p-values ≤ 0.01 as implemented in edgeR are indluced. NAs are missing samples or not applicable for the species. NA* is due to missing NMRI day 0 sample from first infection.

Kasemo et al. Page 3 of 11

2.1 Life cycle analysis... E. falciformis

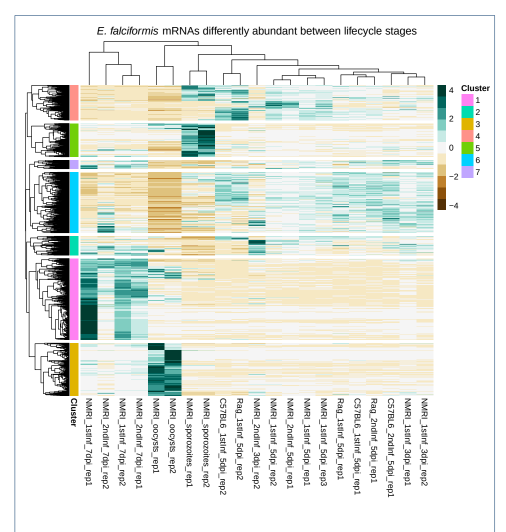


Figure 1 *E. falciformis* mRNAs with significantly different abundance at different times post infection in NMRI mice (see Table ...). *E. falciformis* samples from seven days p.i. cluster together (NMRI mice only). For *E. falciformis* day 7, NMRI, the pattern is most pronounced in cluster 1 (up). Within this cluster, the second replicate from the challenge infection has a deviant profile (compare mouse data in Figure 6). Distinct groups of genes also define sporozoites (cluster 6: up) and oocysts (cluster 3: up, cluster 7: down). mRNA profiles on days three and five p.i. from all three mouse strains cluster together. These samples are distinct from oocysts, NMRI day 7 p.i., and sporozoites. The latter cluster closer with days 3 and 5 than with oocysts or day 7.

Kasemo *et al.* Page 4 of 11

2.2 Parasite development in mouse data

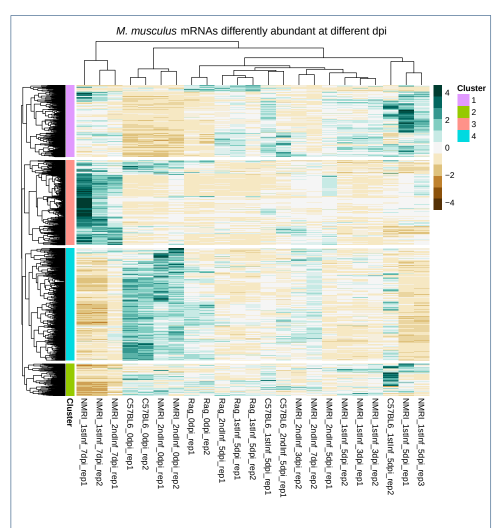
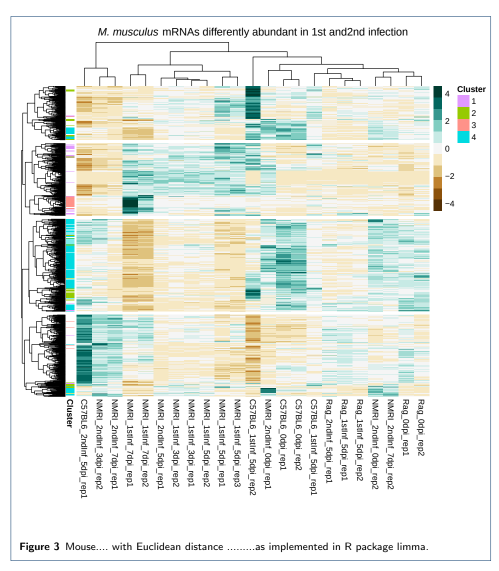


Figure 2 Mouse mRNAs with significantly different abundance at different times post infection in NMRI mice (see Table 3). Three out of four samples from day 7 p.i. (NMRI only) cluster together. These samples are characterized by genes in cluster 2 (up) as well as 3 and 4 (down). The fourth day 7 sample (challenge infection, second replicate) clusters with day 3 and 5 samples but upon visual inspection of cluster 4 diplays a profile similar to non-infected mice. The same sample is abnormal in the parasite profile (see Figure 5). NMRI and C57BL/6 uninfected samples cluster together, defined by clusters 3, 4 (up), and 1 (down). Day 3 and 5 samples cluster together, with Rag1-/- forming a separate group and Rag1-/- non-infected most distant in this sample cluster. All non-infected samples share a high mRNA abundance in cluster 3.

Kasemo et al. Page 5 of 11

2.3 First versus second infection, mouse



GO term enrichments in heatmap clusters

The annotations referred to here are inferred from orthologs in other Eimeria spp. or in $T.\ gondii$

$Data\ overview$

In total, xx reads were sequenced. For each sample, mouse read numbers were 10^7 - 10^8 Quality filters applied resulted in xx reads being removed from further analysis

Preparation for invasion in oocysts

The mRNA profile in the oocyst stage is mainly determined by highly abundant genes in cluster 4. Overrepresented GO-terms in this cluster are enriched by ortholog genes to peptidases, microneme localized proteins reported to be involved in invasion, genes associated with adhesion in protozoans (and with clotting in higher

Kasemo et al. Page 6 of 11

eukaryotes), and genes that are annotated to be involved in amino acid biosynthesis. Aminopeptidase N ('related' annotation) is the reported ortholog for three genes with abundant mRNAs in oocysts. In humans, this enzyme has been reported to cleave peptides bound to major histocompatibility complex, MHC, II (UniProt reference if we want to keep this... but does any secretion happen from oocysts...? Or is this too far-fetched to be interesting?).

A Thrombospondin type 1 domain-containing protein ortholog is highly abundant in cluster 4 (high abundance in oocysts). Thrombospondin type 1 domains have been reported in E. tenella microneme localizing proteins, MIC, e.g. MIC4 (Tomley01) In E. tenella MIC4 mRNA was reported in sporozoites where it localizes to the apical end, and in late schizonts and late oocyst stages, when sporozoites are forming. (Tomley01). For the same gene, the T. gondii annotation is Sushi domain-containing protein, which is also the ortholog annotation of another gene in this cluster. In the related malaria parasite P. falciparum the apical sushi protein, ASP, (which has a sushi domain) localizes to micronemes in merozoites but not other stages (OKeeffe05). Limulus clotting factor C, Coch-5b2 (Cochlin) and Lgl1, LCCL, (syn. F5/8 domain) domains are associated discoidin lectin domains and thereby with adhesion. (Pfam entries for 'LCCL domain' and 'Discoidin domain', May 2016). Taken together, this indicates that the thrombospondin and sushi-domain genes (EfaB_MINUS_4114.g412 and EfaB_PLUS_1425.g183) are involved in sporozoite invasion in E. falciformis and that the mRNAs are transcribed and available before excystation. A role in merozoite re-invasion in E. falciformis is not indicated by our data. The LCCL domain annotation and thrombospondins role in higher eukaryotes also indicates that adhesion or preparation for adhesion is important in oocysts. We suggest (speculate...?) that the thrombospondin annotated ortholog and the LCCL domain-containing protein (EfaB_MINUS_11233.g986) are involved in cell adhesion in E. falciformis.

Amino acid biosynthesis in oocysts

High abundance of aminotransferase mRNAs indicate amino acid biosynthesis or preparation for the same in oocysts (cluster 4). We identify D-3-phosphoglycerate dehydrogenase and alanine dehydrogenase orthologs, which are enzymes contributing to L-serine and L-alanine production, respectively. A putative *Eimeria* spp. cystathionine beta-synthase, CBS, in this cluster also indicates de novo cysteine production. Alkyl sulfatase mRNA is also abundant in oocysts. Generally, this enzyme enables an organism to exploit organic sulfur to produce and incorporate inorganic sulfur into the amino acids cysteine and methionine, when no inorganoc sulfur is available.

'Embryonic development' Nicalin 1, patched family protein (hedgehog)

Oocysts contain mRNA for fatty acid catabolism

MmgE/PrpD is overrepresented in oocysts. The enzyme is important for propionate catabolism in the 2-methylcitric acid cycle and has been shown to be used by the intestinal intracellular bacterium Salmonella typhimurium to generate pyruvate (Horswill99). Propionate is one of two most abundant small-chain fatty acids in the gut along with butyrate. Both fatty acids are largely produced as degradation

Kasemo et al. Page 7 of 11

products from food by commensal bacteria (Sun13). Sharing the intestines as a niche with S. typhimurium it is possible that also E. falciformis uses Mmg/PrpD to exploit available propionate for pyruvate production.

Oocyst highly abundant mRNAs are downregulated in sporozoites

Interestingly, the genes described above which are thought to be involved in amino acid biosynthesis and invasion are highly abundant in oocysts but are underrepresented in schizont stages (day 3 and day 5 samples) and even in sporozoites. An average abundance was detected on day 7 for these genes, indicating a role in either gametes or early oocyst formation. This pattern supports the suggestion that these specific mRNAs (cluster 4) for invasion and biosynthetic processes are prepared (and possibly expressed) in the oocyst stage but are no longer detectable in the cell at the timepont when the protein is assumed to be in use (sporozoites and merozoite stages). Therefore, correlating mRNA prevalence with biological function at the timepoint when mRNAs are detected must be done with care.

Down in sporozoites and oocysts -¿ cluster 3.......

Oocysts: profile for 3, 5, 6:

Sporozoites: 3 and 5 $\,$

Day 7: 5 and 7

Specific genes.... Enolase 2, encoded by Eno2, is among the downregulated genes in oocysts and sporozoites. In *T. gondii* the paralog Eno1 is strongly associated with the cyst (bradyzoite) stage and Eno2 is associated with tachyzoite stages (kibe05). It is therefore expected that this mRNA is underrepresented in oocysts and our data also show that the same is true in sporozoites for *E. falciformis*. (TK: If important we could look specifically for Eno1 in cluster 4).

Down in sporozoites and oocysts -ż cluster 3......

Motility-related mRNAs indicate gamete development on day 7

Two clusters contain genes with mRNAs highly abundant on day 7 p.i; cluster 1 and 2. Dynein, kinesin and tubulin are annotations highly represented among orthologs of genes in both these clusters. The annotations indicate an important role for motility at this timepoint, probably reflecting development of microgametes. In addition, in cluster 2, there are two 'EF-hand domain containing proteins' annotations as well as caltractin, centrin-1, and troponin annotations. Caltractin and centrin-1 are associated with the centrosome and structure and function of microtubuli in mammals, and troponin is linked to muscle function (UniProt). Also potentially linked to motility is the occurrence of growth arrest specific protein 8, Gas8, which in the mouse has been reported to be highly expressed in the testes and important for mouse sperm function (Yeh02).

Other genes among the 38 indicate carbon fixation (glycolysis/gluconeogenesis) or conversions of nucleoside phosphates. In addition, a Ras family protein, RNA polymerase II transcription initiation factor and Sec23 and Sec24 were among orthologs identified in *E. falciformis* cluster 2.

In cluster 1, carbon metabolism genes are represented by 6-phosphogluconate dehydrogenase and glycogen phosphorylase family protein 1. UDP-glucose 4-epimerase Kasemo et al. Page 8 of 11

and a miloride-sensitive amine oxidase are reported as upregulated in game tocytes in E. tenella by RNA-seq (Walker15) and suggested by those authors to play a role in cyst wall synthesis.

Microneme proteins highly expressed on day 7 p.i.

Unintuitively for a protozoan organism, seven out of eight GO biological process terms in cluster 1 are associated with wound healing and blood coagulation. An explanation is offered by some of the orthologs to the three E. falciformis genes responsible for these terms. In protozoa, e.g., other Eimeria spp. and Toxoplasma gondii orthologs are annotated as 'Micronemal protein MIC4, related' (E. tenella) and more generally for several other protozoa, 'PAN domain containing proteins'. The PAN domain is found in the plasminogen/hepatocyte growth factor family and in coagulation factor XI family (REF), explaining why terms related to blood coagulation are enriched by these genes. Later publications on T. qondii (Marchant 12) also associate PAN domains and proteins in apicomplexan parasites with micronemes and therefore invasion. In our case, this is peculiar, since the enrichment appears on day 7 p.i.. A possible role at this timepoint is suggested by work on the fungi Sclerotinia sclerotiorum where Yu et al. reported an important role for PAN domain proteins in cell wall integrity (Yu12). This role for MIC proteins has to our knowledge not been investigated in apicomplexan parasites. The PAN domain domain has also been reported to be common in nematodes such as Caenorhabditis elegans, however the function is not understood. (Thordai99) The other two GO terms in the cluster of day seven upregulated genes are DNA replication and DNA replication initiation, which most likely reflects late stage schizogony or gamete formation. Six genes contribute to this enrichment and orthologs are either annotated as DNA replication licencing factors, DNA polymerases or minichromosome maintenance proteins 2/3/5/7, Mcm2/3/5/7.

Gene and sample patterns by hierarchical clustering

Samples (columns) cluster into two major clusters where day 7 p.i. samples form one group distinct from other samples. In the second group, oocsts and sporozoites are distinct and sporozoites cluster most closely with day 3 and 5 p.i. samples. Day 3 and 5 p.i. samples also cluster into two groups, of which one contains all NMRI day 5 p.i. samples. Apart from this, the two day 3 and 5 p.i. sample clusters have no obvious patters.

For gene clusters (rows), the two groups with high mRNA abundance on day 7 p.i. (cluster 1 and 2) do not cluster most closely with each other, but with the cluster for high mRNA abundance in oocysts (cluster 1 association) and with the cluster for high mRNA abundance in sporozoites (cluster 2 association).

Discussion

In our analysis we demonstrate which biological processes are dominant in different life cycle stages of *E. falciformis* in the mouse. The RNAseq transcriptome provided here allows for detailed analysis of genes involved in those processes, providing candidates for life stage specific marker in *Eimeria* spp. research.

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Kasemo et al. Page 9 of 11

3 Methods

3.1 Mice and infection procedure

Three strains of mice were used in our experiments: NMRI (Charles River Laboratories, Sulzfeld, Germany), C57BL/6 (), and Rag1-/- on C57BL/6 background (gift from Susanne Hartmann, FU?). Animal procedures were performed according to the German Animal Protection Laws as directed and approved by the overseeing authority Landesamt fuer Gesundheit und Soziales (Berlin, Germany). Animals where infected as described by Schmid et al., (Schmid12), but tapwater was used instead of PBS for administration of oocysts. Briefly, NMRI mice were infected two times, which will be referred to as first and second infection. For the first infection, 150 sporulated oocysts were administered in 100 L by oral gavage. During the first infection of 60 mice, all animals were weighed every day. On day zero, before infection, as well as on day three, five and seven days post infection, dpi, caeca from 3-4 sacrificed mice per time point were collected. Epithelial cells were isolated as described in Schmid et al.(schmid12). For challenge infection, mice recovered for four weeks before second infection. Recovery was monitored by weighing and visual inspection of fur. For the second infection, 1500 sporulated oocysts were applied by oral gavage. Three mice were used as non-second infection control, referred to as day 0, second infection.

3.2 Oocyst purification for infection and sequencing

Sporulated oocysts were purified by flotation from feces stored in potassium dichromate and administered orally in 100 uL tapwater. One *E. falciformis* isolate, *E. falciformis* Bayer Haberkorn 1970, was used for all infections and parasite samples. The strain is maintained through passage in NMRI mice in our facilities as described elsewhere (schmid12).

3.3 Sporozoite isolation

Sporozoites were isolated from sporocysts by excystment. For this, sporocysts were incubated at 37C in DMEM containing 0.04% tauroglycocholate (MP Biomedicals) and 0.25% trypsin (Applichem) for 30 min. Sporozoites were purified by the method of Schmatz et al (schmatz—).

3.4 RNA extraction

Total RNA was isolated from infected epithelial cells, sporozoites and sporulated oocysts using Trizol according to the manufacturers protocol (Invitrogen). High quality what is the meaning of 'high quality' here? RNA was used to produce an mRNA library using the Illuminas TruSeq RNA Sample Preparation guide. stolen from genome paper Sporozoites were stored in 1 mL Trizol until RNA-isolation. Total RNA was isolated using the PureLink RNA Mini Kit (Invitrogen) and reverse transcribed into cDNA.

3.5 Sequencing, sequence quality assessment and alignment

cDNA samples were sequenced by either GAIIX or Illumina Hiseq 2000 as specified in SI xx (both unstranded). A fastq_quality_filter (FASTQ-toolkit, version 0.0.14, available at https://github.com/agordon/fastx_toolkit.git) was applied to Illumina

Kasemo et al. Page 10 of 11

Hiseq 2000 samples after replacing "N" bases by "." annotation. A phred score of 10 was applied. We further set q = 60. These settings require that nine out of ten bases or more are correct in at least 60% of the bases for each read.

3.5.1 Alignment and reference genomes

We used the published *Mus musculus* mm10 assembly (Genome Reference Consortium Mouse Build 38, GCA_00001635.2) as reference genome including annotations for mouse data. The *E. falciformis* genome (Heitlinger14) was downloaded from ToxoDB (Gajria07). For the alignment, the mouse and parasite genome files were merged into a dual reference genome, and files including mRNA sequences from both species were aligned against the dual reference genome using TopHat2 (version 2.0.14, Trapnell09) with -G specified, and a Bowtie2 (version 1.1.2, Langmead12) index of the dual genome. Single-end and pair-end sequence samples were aligned separately with library type 'fr-unstranded' specified for pair-end samples. Import into R was enabled by the R package Ballgown, which requires bam files to be processed by Tablemaker (Frazee15), in our case used with -qW -G specified. Tablemaker in turn makes use of Cufflinks (version 2.1.1, Trapnell10).

3.5.2 Data normalisation and sample exclusions

Count data was normalised in R, using the upperquartile method as implemented in edgeR (version 3.14.0). Briefly, normalisation assumes a negative binomial distribution of data and dispersed variance for genes with higher mean abundance. The normalisation shrinks the dispersion of variance towards a Poisson (linear) relationship between mean and variance. We evaluated the density distribution of our data and two samples contradicted the assumed negative binomial distribution and were excluded from further analysis: NMRI_1st_3dpi_rep1 and NMRI_2nd_5dpi_rep1 (SI ...).

3.6 Differentially abundant mRNAs used for hierarchical clustering

A selection of differently abundant mRNAs are used for hierarchical clustering of *E. falciformis* life cycle relevant genes. In each comparison (see Table 3), the 500 differentially abundant mRNAs with lowest FDR are selected. In the next step, the 500 mRNAs from each comparison (or less) are joined (4935 genes) and only unique genes are selected (1618 genes). The 22 genes in the NMRI vs C57BL/6 comparison are not included in the *E. falciformis* life cycle analysis.

All parasite samples were included in heatmap of genes detected as differentially abundant in timepoint comparisons.

On scale bar in heatmaps, 0 is mean mRNA abundance for each gene (row). Up (green) and down-regulation (brown) numbers denote number of standard deviations from row mean. Hierarchical clustering was performed using with Euclidean distances, method 'complete' (R package base).

Table 3 Differentially abundant E. falciformis mRNAs used for hierarchical clustering.

Data description	Number of genes
Sum of 1st infection NMRI sample differences	4935
(including oocysts and sporozoites) Used in hierarchical clustering (heatmap)	1618

Kasemo et al. Page 11 of 11

Competing interests

The authors declare that they have no competing interests.

Author's contributions

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Figures

Figure 4 Parasite read sequences per mouse read sequences for each experimental condition. The more advanced the infection is, the higher the ratio of parasite reads is, reflecting parasite replication between day zero and day seven.compare 1st and 2nd patterns between Rag and NMRI/C57BL/6? Values are mean of replicates (n=2, if * n=1) on log10 scale. Each sample (replicate) consists of mRNA from three different mice.

Tables

Table 4 Sample table title. This is where the description of the table should go.

	B1	B2	B3
A1	0.1	0.2	0.3
A2			
А3			

Additional Files

Additional file 1 — Sample additional file title

Additional file descriptions text (including details of how to view the file, if it is in a non-standard format or the file extension). This might refer to a multi-page table or a figure.

 $\label{eq:Additional} \mbox{Additional file 2} \mbox{$-$ Sample additional file title} \\ \mbox{Additional file descriptions text.}$