Gene-expression profiling discriminates between cerebral malaria (CM)-susceptible mice and CM-resistant mice.
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Gene expression profiling discriminates cerebral malaria-susceptible mice and cerebral malaria-resistant mice

Short title: Gene expression profiles in mouse malaria

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Abstract

The development of mouse cerebral malaria (CM) upon *Plasmodium berghei* ANKA infection is under genetic control. Brain gene expression patterns were investigated in well-defined genetically CM-resistant (BALB/c and DBA/2) and CM-susceptible (C57BL/6 and CBA/J) mouse strains by using cDNA microarrays. By combining transcriptional profiling with rigorous statistical methods and cluster analysis, we identified a set of 69 genes that perfectly discriminated between mouse strains and between CM-resistant and CM-susceptible mice. The analysis of Gene Ontology terms revealed that the genes clustered and related to susceptibility to CM preferentially belonged to some biological process classes, such as those pertaining to immune responses. Using a false discovery rate of 5% and Welch t-test, we identified 31 genes with consistent differential expression between CM-resistant and CM-susceptible mice. These data indicate that microarray analysis may be useful to identify candidate genes potentially responsible for resistance or susceptibility to CM.

Key words: cerebral malaria, microarrays, mouse, brain, resistance, statistics, immunopathology
Introduction

Malarial disease is a major public health problem in the developing countries. Cerebral malaria (CM) is one of the most severe complications and a major cause of death [1,2]. A large body of evidence indicates that malaria infection and disease have heritable components, both in the human disease and in murine models. In the search for human genetic factors involved in malaria pathology or protection, several genes have been associated with resistance to severe malaria, and genes controlling parasitemia and mild malaria have been mapped [3-6]. The role of some genes has been demonstrated in mouse models and new malaria resistance genes that have been located on mouse genome remain to be identified [1,7].

To investigate the pathogenesis of CM, several animal models have been established, including mice infected by *Plasmodium berghei* ANKA (PbA) [7,8]. Although experimental CM models cannot reproduce all the features of human CM, several observations in mouse CM have been extended and confirmed in human disease. These include behavioural changes, histopathological features in brain and retina, biochemical changes in brain, blood-brain barrier function, immunological responses, and expression of molecules in brain and retina [1].

Inbred mouse strains markedly differ in their susceptibility to *Plasmodium* infection and disease. In the murine model of PbA infection, BALB/c and DBA/2 mice are resistant to CM (CM-R) while C57BL/6 and CBA/J mice are susceptible to the neurological syndrome (CM-S). Our working hypothesis was that CM resistance/susceptibility differences may be due to genetic differences influencing gene expression. In particular, since CM is associated with an abnormal increase in the proinflammatory immune responses, it is conceivable that genes involved in inflammation are up-regulated.

Global expression analysis of 5053 genes was employed to identify differences in the brain expression between CM-R mice (BALB/c and DBA/2 mice) and CM-S mice (C57BL/6 and CBA/J).
We report here that CM-R mice and CM-S mice infected by PbA differ in their gene expression, the gene expression profile being discriminant of resistance or susceptibility to CM.

**Materials and Methods**

**Mouse Strains and Phenotyping**

Six to 8 weeks old BALB/c, DBA/2 (CM-R strains), C57BL/6 and CBA/J (CM-S strains) female, were obtained from IFFA CREDO (Ch. River Lab, France) and kept in our facilities. Five from each strain were infected by i.p. injection of *P. berghei* ANKA. Parasitemia was monitored daily on blood smear. The CM-S mice developed a neurological syndrome which occurred 6 to 7 days after parasite inoculation with a cumulative mortality of 100 %. The CM-R mice die during the 3rd or the 4th week of infection, with severe anaemia and hyperparasitemia [7].

**Organ Sampling and Histology**

Brains were taken from CM-R and CM-S mice when the CM-S mice develop CM. Brains were completely removed and cut in two parts: one part was frozen in RNALater (Qiagen, TM) until RNA analysis, and the other one was embedded in Tissue Tek (Leica), snap frozen in liquid nitrogen and kept at -80°C until histopathological examination of cryosections. Histology consisted in specific antibody immunohistochemistry for VCAM-1 (Clone 429, BD Pharmingen) and ISCBP1 (C19, TEBU Bio). Afterwards, slides were photographed and protein expression was quantified using LUCIA software (Nikon) [8].

**RNA Isolation and cDNA Microarray Hybridizations**

Total RNA from brains was extracted using TRIzol reagent (Gibco-BRL, Life Technologies). The quality of RNA was confirmed on a formaldehyde agarose gel, and the concentration of RNA was determined by reading absorbance at 260/280 nm. Each mRNA sample extracted from an individual sample was run on a single microarray. All microarray procedures were done at our microarray core facility (http://tagc.univ-mrs.fr/). cDNA microarrays were designed and
prepared as described [9]. The microarrays used in this study were composed of 5053 sequences. PCR amplification was performed as previously described [9], and PCR products were spotted onto nylon membranes (Hybond-N+, Amersham) with a GMS-417 arrayer (Affymetrix, Santa Clara, CA). About 20% of the genes included in this clone set are represented by two or more different cDNA clones, providing internal controls to assess the reproducibility of gene expression measurements. Microarrays were hybridized with $^{33}$P-labelled probes, first with an oligonucleotide sequence common to all spotted PCR products (5'TCACACAGGAAACAGCTATGAC-3'), then after stripping, with complex probes made from 5 µg of retrotranscribed total RNA. Probe preparations, hybridizations and washes were carried out as described previously [9]. After 48h hybridization, arrays were scanned with a FUJI BAS5000 machine at 25 µm resolution.

**Microarray Data Analysis**

Hybridization signals were quantified using ArrayGauge software (Fuji Ltd, Tokyo, Japan). All images were carefully inspected, and spots with overestimated intensities due to neighborhood effects were manually excluded. The data from 1 sample (CBA/J mouse) indicated a lower hybridization quality, so the sample was omitted from subsequent analyses. The data were filtered such that only spots with intensities that were two times greater than the median background in either microarray were used in the analysis, and the signal intensities were then corrected to take into account the amount of spotted DNA and the variability of experimental conditions, as described [10]. Of the 5053 spotted clones, we selected the clones that had detectable expression levels in at least 80% of the experiments ($n=2145$). Unsupervised hierarchical clustering investigated relationships between samples and relationships between genes. It was applied to data log-transformed and median-centred using the Cluster and TreeView programs (average linkage clustering using Pearson’s uncentered correlation as similarity metric) [11]. Using supervised SVM (Support Vector Machine) method, we
investigated the discrimination between CM-R and CM-S mice based on gene expression profiles [12].

Statistical analyses were performed using the TIGR MeV (MultiExperiment Viewer) v3.1 software (http://www.tm4.org/mev.html). Figure 1 shows an outline of data analysis. A one-way ANOVA and SAM (Significant Analysis of Microarrays) procedures [13] were applied to look for strain- and CM-R/CM-S-specific variation in gene expression in the full data set. Welch t-statistics was used to compare CM-R and CM-S mice [14]. Multiple test corrections were performed [15].

Gene annotation of all 2145 analysable genes/ESTs was obtained using EASE (Expression Analysis Systematic Explorer) software [16]. This program which allows a biological interpretation of gene clusters on the basis of GO terms was used to assess whether specific biological pathways were over-represented among the differentially expressed genes and within specific gene clusters. A score based on Fisher Exact test reports the probability that the prevalence of a particular theme within a cluster is due to chance alone given the prevalence of that theme in the population of all genes under study.

All data are MIAME compliant and have been submitted to ArrayExpress database (www.ebi.ac.uk/arrayexpress).

**Results**

To identify genes differentially expressed between CM-S and CM-R mice, we combined unsupervised and supervised clustering methods with several statistical approaches (figure 1). All the CBA/J mice \((n=4)\) and C57BL/6 mice \((n=5)\) developed CM while the BALB/c mice \((n=5)\) and DBA/2 mice \((n=5)\) did not, in spite of similar levels of parasitemia \((10.4\% \pm 6.4)\). The mRNA from the 19 different brain mouse samples was hybridized with cDNA arrays. The mRNA samples from each strain represented biological replicates. The degree of correlation between
gene expression levels of two samples from each strain was high (Pearson’s correlation coefficient R mean=0.89).

To analyze the overall expression patterns, we used an unsupervised hierarchical clustering method that groups genes on the vertical axis and samples on the horizontal axis on the basis of similarity in their expression profiles (figure 2A). The similarities are summarized in a dendrogram, in which the pattern and length of the branches reflects the relatedness of the samples (figure 2B). On this basis, 8 of 10 CM-R mice and 9 of 9 CM-S mice were clustered together, respectively. The discrimination between CM-R and CM-S mice was highly significant (Fisher exact test : $P<.0001$). We assumed that 2 CM-R mice were incorrectly classified because of confounding genes, which were not differentially expressed between mouse groups.

We, therefore, performed one-way ANOVA to look for strain- and CM-R/CM-S-specific variation in gene expression in the full data set (figure 1). We generated empirical $P$ values for each gene. Then, we selected all genes having $P <.05$, and we defined a set of 292 informative genes. We performed further unsupervised hierarchical clustering of samples based on the expression of the selected genes (figure 2C). As shown in figure 2D, this procedure successfully classified 10/10 CM-R mice and 9/9 CM-S mice. All mice of each strain were also correctly classified.

To ascertain the accuracy of the prediction of CM molecular signature, we used a “one-out-iterative cross-validation” testing procedure developed by the supervised SVM method. Iteratively, one of the 19 mouse samples was removed from the group, and was classified on the basis of the correlation between its expression profile and the median profile of samples from CM-R mice and CM-S mice. The prediction was considered correct if it corresponded to the actual group. Using this approach, we correctly classified 10 of 10 CM-R mice and 9 of 9 CM-S mice, and we confirmed the results of unsupervised hierarchical clustering.
To search for significant genes based on differential expression between mouse groups, we carried out more stringent statistical methods, which control the percentage of genes identified by chance. We applied two SAM procedures on the full data set to pick out mouse strain-specific and CM-R/CM-S-specific genes (figure 1). Differentially expressed genes were selected by using a false discovery rate (FDR) of 5%. Multi-class SAM yielded a list of 50 genes that showed a differential expression between mouse strains. Two-class SAM yielded a list of 40 genes that appeared to be differentially expressed between CM-R and CM-S mice. Twenty-one genes belonged to both lists, indicating that the effect of both strain and CM phenotype was detected. We, therefore, found 69 significant genes, which belong to the preliminary list of 292 informative genes.

Unsupervised hierarchical clustering of samples based on the expression of the 69 significant genes successfully classified all the BALB/c, DBA/2, C57BL/6, and CBA/J mice on the one hand, and all the CM-R and CM-S mice on the other hand (figure 3). In the same way, 10 of 10 CM-R mice and 9 of 9 CM-S mice were correctly classified by using supervised SVM method. We observed 7 clusters that were either strain-specific or CM-R/CM-S-specific. Clusters A and E grouped DBA/2 and CBA/J strains vs BALB/c and C57BL/6 strains. Clusters C and G discriminated C57BL/6 and BALB/c mice, respectively. Clusters B, D, and F separated CM-R and CM-S mice.

We further analyzed the genes that we selected as CM-R/CM-S-specific by using the two-class SAM (figure 1). The 40 genes were assessed for bias due to strain-specific variation in gene expression. First, using Welch t-test, we searched for genes that were differentially expressed in brain samples from BALB/c and DBA/2 CM-R strains on the one hand, and C57BL/6 and CBA/J CM-S strains on the other hand. Then, we took into account these differences when we analyzed differential expression between CM-R and CM-S strains.

Among the 40 genes selected as CM-R/CM-S-specific, we found 28 genes, the expression of which was not different between BALB/c and DBA/2 CM-R strains nor between C57BL/6 and
CBA/J CM-S strains (table 1). We showed by using Welch t-test and a FDR of 5% that the expression of 11 genes was lower in CM-S mice than in CM-R mice (figure 3, cluster F), and that the expression of 17 genes was higher in CM-S mice than in CM-R mice (figure 3, clusters B and D).

Among the 40 genes selected as CM-R/CM-S-specific, we found 12 genes, the expression of which was different either between BALB/c and DBA/2 CM-R mice or between C57BL/6 and CBA/J CM-S mice (table 2). Using Welch t-test and a FDR of 5%, we found that the expression of $Icsbp1$ and $Vcam1$ was lower in CM-R strains than in C57BL/6 and CBA/J CM-S strains, and that the expression of $Ccl27$ was higher in CM-S strains than in BALB/c and DBA/2 CM-R strains. The expression of the other genes did not significantly differ between CM-R and CM-S (table 2). In all, we identified 31 genes associated with CM.

The analysis of the GO terms of genes composing the different identified clusters by the EASE software showed an overrepresentation of genes involved in defence response (cluster D, $P<.0004$; cluster F, $P<.008$) and immune response (cluster D, $P<.0002$), compared to the population of all genes under study. For instance, the cluster D contains $Samhd1$, $Serping1$, $Icsbp1$, $C1qb$, $Ifit3$ and $Ccl27$, which are annotated with the “immune response” GO term.

By using immunochemistry, we further analyzed the expression of two genes associated with susceptibility to CM. We chose $Icsbp1$, which is a crucial component of the IFN$\gamma$ response, and $Vcam1$, which is a TNF-induced gene. The anti-ICSBP1 specific immunostaining, however, was not efficient enough to perform quantification. In contrast, the results were satisfactory for anti-VCAM-1 antibody (figure 4), and we found that $Vcam1$ was significantly overexpressed in CM-S mice as compared to CM-R ($P<.0001$, Student t-test).

**Discussion**
By combining transcriptional profiling with rigorous statistical methods and cluster analysis, we searched for differences in brain gene expression between CM-S (C57BL/6 and CBA/J) and CM-R (BALB/c and DBA/2) mice. We used 5 biological replicates per group to take into account variability between mice. This allowed us to search for genes, which show small but significant changes in expression, and which may be biologically important. The approach used by others was rather to perform one or two microarrays for each condition or to pool the samples in order to minimize the experimental variability. Moreover, the selection of genes is generally based on non-statistically motivated criteria, such as « 2-fold-change » [17-19]. On the basis of a FDR of 5%, we identified a set of 69 genes, the expression of which was significantly different between mouse strains and/or between CM-R and CM-S mice. The cluster analysis showed that this set of genes fully discriminated between mouse strains and between CM-R and CM-S mice.

As several mouse malaria resistance loci have been mapped, we looked at the chromosomal localization of the selected genes. From the 69 selected genes, 67 had a reliable chromosomal location, the 2 missing data corresponding to 2 ESTs. Among the 67 genes, several were located at chromosomal regions previously shown to contain susceptibility loci for cerebral malaria caused by PbA. *Usf1* and *Pafah1b1* were located at the chromosome 1 region and the chromosome 11 region, which contain Berr1 and Berr2, respectively [20]. In addition, *Nfatc1* and *Egr1* were located at chromosome 18 that harbours the locus Esmr also controlling resistance to mouse CM [21]. We also found that *Igf2r, Fkbp5, Lst1,* and *H2-D1* were located at the chromosome 17 region harbouring the Cmcs locus [22]. Interestingly, the same chromosome 17 region was reported to contain genes controlling resistance to *P. chabaudi* (Char3) [23]. Moreover, three other chromosomal regions containing a locus controlling resistance to *P. chabaudi*, namely Char1 (chromosome 9), Char2 (chromosome 8) and Char4 (chromosome 3) contained *Tcf12, Sntb2, Fnta, Icsbp1* and *Vcam1*, which we found differentially expressed between CM-R and CM-S mice [23]. It should be stressed that the chromosomal regions linked to malaria resistance are large. Therefore, microarray analyses should be helpful in identifying
candidate genes lying within those genetic regions and showing a differential expression between CM-R and CM-S mice. Cis-acting polymorphisms may affect both gene expression levels and CM resistance.

We identified a cluster of genes up-regulated in C57BL/6 CM-S strain, compared to the CM-R strains and also the CBA CM-S strain. This is consistent with the data indicating that some CM mediators partly differ in CBA and C57BL/6 mice. In particular, it is thought that TNF may be a critical mediator of CM in CBA mice [24], while LTA may be a critical mediator of CM in C57BL/6 mice [25]. In the same way, we identified a cluster of genes specifically up-regulated in BALB/c CM-R mice, suggesting that some CM-R genetic factors may differ between BALB/c and DBA/2 CM-R mice.

We identified three gene clusters associated with resistance or susceptibility to CM. Using Welch t-test and multiple test correction, we further confirmed that the expression of 31 genes significantly differ between CM-R and CM-S mice. This suggests that these genes and related physiological pathways may be critical in malaria pathogenesis in mice. To explore the possible functional role of these genes, we analyzed the occurrence of the GO terms within the clusters. The EASE analysis revealed that the genes clustered preferentially belonged to the two following biological process classes: defence response and immune response. Interestingly, the GO terms are organized in a hierarchical way [26], and the over-represented terms within the clusters of interest were related. Indeed, a number of known genes involved in the immune system and in the pro-inflammatory response had an expression associated with resistance or susceptibility to CM.

In particular, Gzmb, Igf2r, Ctla2a, and C1qb were found to be up-regulated in CM-S mice. Gzmb (Ctla1) encodes granzyme B in cytotoxic T cells and Natural Killer cells [27,28], and induces apoptosis by caspase activation. Igf2r has been reported to bind Gzmb and to be essential for cytotoxic T cell-mediated apoptosis [29]. This is consistent with the data showing that perforin-deficient C57BL/6 mice are resistant to CM. This indicates that the perforin/granzyme cytolytic pathway may be a major mechanism employed by CD8+ T cells during CM [30,31]. Ctla2a is
expressed in mouse activated T cells [32]. $Clqb$ that is the complement component 1 beta gene has also been shown to be up-regulated after permanent focal ischemia in the mouse [33]. In addition, overexpression of cyclooxygenase-2 (COX-2), which is expressed in mouse CM brain [34], leads to selective induction of $Clqb$ expression in brain [35]. One might expect that the presence of locally produced complement components within brain is likely to induce a potent inflammatory response, and to contribute to brain damage.

$Pafhab1$, which was down-regulated in CM-S mice, encodes a subunit Platelet Activating Factor acetylhydrolase, which regulates the level of platelet activating factor (PAF) in brain [36]. It has been suggested that PAF could play a pathophysiological role in severe malaria through activating platelets [37]. Platelets that are sequestered in cerebral microvessels are thought to be involved in endothelial cell damage and in malaria pathogenesis [1].

We identified several genes known to be related to IFN or TNF responses. $Samhd1$, $Ifit3$ and $Lst1$ are interferon-regulated genes [38-40], and $Icsbp1$, $Stat1$ are important components of the cellular response to IFNγ. $Nfatc1$ binds to $Icsbp1$ to promote the transcription of $Il12B$ [41], and to $Egr-1$ to promote the transcription of $Tnf$ and $Il2$ [42]. $Vcam-1$ and $Ccl27$ are TNF-induced genes [43,44], while $Fkbp5$ encodes a heat shock protein (Hsp) 90-binding immunophilin implicated in the TNF signal transduction pathway [45]. We found that $Samhd1$, $Ifit3$, $Icsbp1$, $Stat1$, $Nfatc1$, $Fkbp5$, and $Vcam-1$ expression level was nominally higher in the brain of CM-S mice than in the brain of CM-R mice. We analyzed the expression of $Vcam-1$ and $Icsbp1$ to validate the DNA microarray data by using immunohistochemistry method. We were unable to detect $Icsbp1$, indicating that the monoclonal antibody (mAb) did not perform well on cryosections. In contrast, $Vcam-1$ results confirmed the correlation between $Vcam-1$ expression level and susceptibility to CM. In the same way, Lou et al [46] showed that TNF induced a higher level of $ICAM-1$ and $VCAM-1$ expression in CBA/J (CM-S) than in BALB/c (CM-R) brain microvascular endothelial cells (MVECs). In addition, Sexton et al [47] recently showed that interferon-regulated gene transcripts or IFN regulatory factors, such as $Stat1$, strongly increased
in brain from C57BL/6 (CM-S) mice infected by PbA. These results are in line with the current knowledge of the physiological pathways involved in malaria pathogenesis. The role of IFNγ, TNF and their receptors in mouse CM has been demonstrated by using neutralizing mAb [48] or gene targeting [49-51]. Moreover, human TNF and IFNγR1 polymorphisms are associated with susceptibility to CM [52,53]. High levels of TNF and IFNγ have been measured in patients with severe malaria [48,54], and TNFR2 complex has been observed on the surface of MVECs in human CM [1]. NK cells, CD4+ T cells and γδ T cells were shown to produce IFNγ in in vitro models of P. falciparum infection [55-57]. In mice, NK T cells or NK cells could be important sources of IFNγ, since the introduction of C57BL/6 (CM-S) strain NKC genes into a BALB/c (CM-R) strain results in an increased production of IFNγ, an increased Th1 response, and susceptibility to CM [58].

Here, we show that there are gene expression profiles associated with susceptibility to mouse CM. We identified several genes, the expression of which significantly differs between CM-S and CM-R mice. The concordance between our data and those from literature with respect to the function of genes strengthens the validity of our results. This suggests that these genes -or genes involved in the same physiological pathway- are implicated in malaria pathogenesis in mice and can be considered as candidate genes in humans. Microarray analyses should provide new insights into the knowledge of key genes governing the response to malaria.

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**Figure legends**

**Figure 1.**

**A schematic outline of step-wise data analysis.** SAM, Significant Analysis of Microarrays. FDR, False Discovery Rate. HCL, Hierarchical Clustering. SVM, Support Vector Machine. CM, cerebral malaria. CM-R, CM-resistant mice. CM-S, CM-susceptible mice. A one-way ANOVA was applied to look for strain- and CM-R/CM-S-specific variation in gene expression in the full data set. Empirical $P$-values were computed for each gene by bootstrapping from 10,000 permutations. All genes having $P<.05$ were selected for unsupervised and supervised clusterings.

To estimate the accuracy of prediction of the “CM molecular signature”, we applied the “one-out-iterative cross-validation” testing procedure developed by the supervised SVM method. Briefly, the algorithm withholds one of the 19 mice samples, builds a training set based only on the remaining samples, and predicts the class of the withheld sample. The process was repeated for each sample, and a Fisher exact test was used to assess the significance of the classification.

From the full data set, significant differential gene expression was determined using SAM method. The SAM algorithm gives an estimate of the FDR, which is the proportion of false positives among all of the genes initially identified as being differentially expressed. The multi-class and two-class unpaired SAM procedures were used to identify mouse strain-specific and CM-R/CM-S-specific genes, respectively, on the basis of 10,000 bootstrapped permutations. Missing values were imputed using a K-nearest-neighbor algorithm. Multiple testing problem was addressed by choosing a FDR of 5%. The significant genes with differential expression between CM-R and CM-S mice were extracted from the SAM results with Welch t-test for small samples with unequal variances in the two groups. Empirical $P$-values were computed for each gene from 10,000 bootstrapped permutations. All genes having $P<.05$ were selected and we applied a FDR of 5%.
Figure 2.

**Gene expression profiles in CM-R mice and CM-S mice.** *A*, Hierarchical clustering of 19 brain tissue samples representing the four mouse strains (BALB/c, DBA/2, C57BL/6, CBA/J) and 2145 cDNA clones (full data set) based on mRNA expression levels. Each row represents a gene and each column represents a sample. The normalized expression value for each gene was depicted according to the colour scale at the bottom. Red and green indicate expression levels above and below the median, respectively. Grey indicates missing data. Dendograms of samples (above matrix) and genes (to the left of matrix) represent overall similarities in gene expression profiles. 

*B*, Dendogram of samples representing the results of the same global hierarchical clustering applied to the 19 brain tissue samples. Two groups of samples were defined: CM-R mice (BALB/c, DBA/2) in blue and CM-S mice (C57BL/6, CBA/J) in red. 

*C*, Hierarchical clustering of the 19 brain tissue samples using expression levels of 292 informative genes differentially expressed between the four mouse strains. This group of informative genes was extracted from the full data set (*n*=2145) by an one-way ANOVA test on the basis of 10,000 permutations (*P*<.05). 

*D*, Dendogram of the samples representing the results of the same hierarchical clustering applied to the 292 informative genes.

Figure 3.

**Hierarchical classification of strain-specific and CM-R and CM-S significant genes.** Hierarchical clustering of the 19 brain tissue samples using expression levels of 69 significant genes differentially expressed between the four mouse strains (BALB/c, DBA/2, C57BL/6, CBA/J). This group of significant genes was extracted from the full data set (*n*=2145) by a SAM procedure on the basis of 10,000 permutations, and a FDR of 5%. The Clone ID and the UniGene Symbol are indicated for each gene. Coloured bars to the right indicate the location of clusters (A-G).
Figure 4.

**VCAM-1 expression in brain vessels of CM-S and CM-R mice.** A, Histological pictures of CM-S and B, CM-R mice. VCAM-1 expression was expressed for each vessel as a ratio of the red chromogen area out of the total vessel lumen. C, VCAM-1 was significantly overexpressed in CM-S mice (Student t-test).
Figure 1.
Figure 2.
Figure 4.
Table 1. Cerebral malaria (CM) specific genes obtained from Welch t-test and FDR 5%.

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<td>Budding uninhibited by benzimidazoles 3 homolog (S. cerevisiae)</td>
<td>1226577</td>
<td>1343463</td>
<td>3.31</td>
<td>.0016</td>
</tr>
<tr>
<td>Dnajb4 (D)</td>
<td>DnaJ (Hsp40) homolog. subfamily B. member 4</td>
<td>1225671</td>
<td>1914285</td>
<td>3.68</td>
<td>.0016</td>
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<tr>
<td>Ctla2 (D)</td>
<td>Complement component 1. q subcomponent. beta polypeptide</td>
<td>1225655</td>
<td>88224</td>
<td>3.74</td>
<td>.0006</td>
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<tr>
<td>Fkbp5 (D)</td>
<td>FK506 binding protein 5</td>
<td>1243723</td>
<td>104670</td>
<td>4.69</td>
<td>.0006</td>
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<tr>
<td>Ifit3 (D)</td>
<td>Interferon-induced protein with tetratricopeptide repeats 3</td>
<td>598493</td>
<td>1101055</td>
<td>4.46</td>
<td>.0002</td>
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<tr>
<td>Gzmb (D)</td>
<td>Granzyme B</td>
<td>1378751</td>
<td>109267</td>
<td>4.44</td>
<td>.0009</td>
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<tr>
<td>Nfatac1 (D)</td>
<td>Nuclear factor of activated T-cells. cytoplasmic. calcineurin-dependent 1</td>
<td>575679</td>
<td>102469</td>
<td>4.37</td>
<td>.0013</td>
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<tr>
<td>Dusp11 (F)</td>
<td>Dual specificity phosphatase 11 (RNA/RNP complex 1-interacting)</td>
<td>1226939</td>
<td>1919352</td>
<td>-4.87</td>
<td>.0001</td>
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<tr>
<td>Lst1 (F)</td>
<td>Leukocyte specific transcript 1</td>
<td>621304</td>
<td>1096324</td>
<td>-4.37</td>
<td>.0008</td>
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<tr>
<td>Igh-6 (F)</td>
<td>Immunoglobulin heavy chain 6 (heavy chain of IgM)</td>
<td>1263881</td>
<td>96448</td>
<td>-4.03</td>
<td>.0011</td>
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<tr>
<td>H2-D1 (F)</td>
<td>Histocompatibility 2. D region locus 1</td>
<td>576114</td>
<td>95896</td>
<td>-3.83</td>
<td>.0013</td>
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<tr>
<td>Ccr4l (F)</td>
<td>CCR4 carbon catabolite repression 4-like (S. cerevisiae)</td>
<td>1263702</td>
<td>109382</td>
<td>-3.54</td>
<td>.0004</td>
</tr>
<tr>
<td>Palp1b (F)</td>
<td>Platelet-activating factor acetylhydrolase. isoform 1b. beta1 subunit</td>
<td>1263702</td>
<td>109520</td>
<td>-4.79</td>
<td>&lt;.00001</td>
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<tr>
<td>Slc25a19 (F)</td>
<td>Solute carrier family 25 (mitochondrial deoxynucleotide carrier). member 19</td>
<td>1263222</td>
<td>1914533</td>
<td>-4.66</td>
<td>.0004</td>
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<tr>
<td>9630015D15Rik (F)</td>
<td>RIKEN cDNA 9630015D15 gene</td>
<td>1244014</td>
<td>2140359</td>
<td>-3.67</td>
<td>.0036</td>
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<tr>
<td>Pcnxl3 (F)</td>
<td>Pecanex-like 3 (Drosophila)</td>
<td>1263238</td>
<td>1861733</td>
<td>-4.18</td>
<td>.0013</td>
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<tr>
<td>Gcn11 (F)</td>
<td>GCN1 general control of amino-acid synthesis 1-like 1 (yeast)</td>
<td>1226847</td>
<td>2444248</td>
<td>-4.60</td>
<td>.0004</td>
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<tr>
<td>Fnta (F)</td>
<td>Farnesyltransferase. CAAX box. alpha</td>
<td>1263688</td>
<td>104683</td>
<td>-3.92</td>
<td>.0007</td>
</tr>
</tbody>
</table>

**NOTE.** The genes are listed following the clustering order in figure 2. CM-R, CM-resistant mice. CM-S, CM-susceptible mice.

* Positive score: CM-S mice gene expression was higher than CM-R mice gene expression.

Negative Score: CM-R mice gene expression was higher than CM-S mice gene expression.
Table 2. Cerebral malaria (CM) specific genes with a strain effect obtained from Welch t-test and FDR 5%.

<table>
<thead>
<tr>
<th>Group*, Gene symbol</th>
<th>Name</th>
<th>CloneID</th>
<th>MGI ID</th>
<th>Welch t-test Score</th>
<th>Empirical P value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Group A</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Cc21a (C)</em></td>
<td>Chemokine (C-C motif) ligand 21a (leucine)</td>
<td>576610</td>
<td>1349182</td>
<td>8.18</td>
<td>&lt;.00001*</td>
</tr>
<tr>
<td><em>Tle4 (C)</em></td>
<td>Transducin-like enhancer of split 4, homolog of Drosophila E(spl)</td>
<td>576586</td>
<td>104633</td>
<td>6.73</td>
<td>.00033*</td>
</tr>
<tr>
<td><em>Icsbp1 (D)</em></td>
<td>Interferon consensus binding protein 1</td>
<td>577563</td>
<td>96395</td>
<td>2.47</td>
<td>.033*</td>
</tr>
<tr>
<td>1110008H02Rik (D)</td>
<td>RIKEN cDNA 1110008H02 gene</td>
<td>1226403</td>
<td>1921074</td>
<td>6.30</td>
<td>&lt;.00001*</td>
</tr>
<tr>
<td><em>Vcam1 (D)</em></td>
<td>Vascular cell adhesion molecule 1</td>
<td>576563</td>
<td>98926</td>
<td>6.29</td>
<td>&lt;.00001*</td>
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<tr>
<td><strong>Group B</strong></td>
<td></td>
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<td></td>
</tr>
<tr>
<td><em>Ccl27 (D)</em></td>
<td>Chemokine (C-C motif) ligand 27</td>
<td>1446118</td>
<td>1343459</td>
<td>6.31</td>
<td>&lt;.00001*</td>
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<tr>
<td><em>Arc (G)</em></td>
<td>Activity regulated cytoskeletal-associated protein</td>
<td>3327010</td>
<td>88067</td>
<td>-5.73</td>
<td>.0005*</td>
</tr>
<tr>
<td><strong>Group C</strong></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>C630013N10Rik (C)</em></td>
<td>RIKEN cDNA C630013N10 gene</td>
<td>1225973</td>
<td>2443079</td>
<td>10.30</td>
<td>&lt;.00001*</td>
</tr>
<tr>
<td><em>Cd21b (C)</em></td>
<td>Chemokine (C-C motif) ligand 21b (serine)</td>
<td>1379475</td>
<td>1349183</td>
<td>11.74</td>
<td>&lt;.00001*</td>
</tr>
<tr>
<td><em>Ptgg1 (C)</em></td>
<td>Pituitary tumor-transforming 1</td>
<td>1243983&lt;sup&gt;d&lt;/sup&gt;</td>
<td>1353578</td>
<td>15.05</td>
<td>&lt;.00001*</td>
</tr>
<tr>
<td>2610005L07Rik (F)</td>
<td>RIKEN cDNA 2610005L07 gene</td>
<td>1262936</td>
<td>1914283</td>
<td>-4.53</td>
<td>&lt;.00001*</td>
</tr>
</tbody>
</table>
NOTE. The genes, the expression of which significantly differed between CM-R and CM-S mice, are shown in **bold**. A false discovery rate (FDR) of 5% was used.

- **Group A**: the expression of the gene differed between C57BL/6 and CBA/J mice.
- **Group B**: the expression of the gene differed between BALB/c and DBA/2 mice.
- **Group C**: the expression of the gene differed between C57BL/6 and CBA/J mice, and between BALB/c and DBA/2 mice.

- Positive score: CM-S mice gene expression was higher than CM-R mice gene expression.
- Negative score: CM-R mice gene expression was higher than CM-S mice gene expression.

- Significant results with a FDR of 5% are indicated by an asterisk.
- Results are shown for this clone. Similar results were obtained for the other clone.