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Pathogenesis of congenital cytomegalovirus infection : finding prognostic markers and correlates of protection

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Citation

Rovito, R. (2018, October 16). *Pathogenesis of congenital cytomegalovirus infection : finding prognostic markers and correlates of protection*. Retrieved from <https://hdl.handle.net/1887/66319>

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Author: Rovito, R.

Title: esis of congenital cytomegalovirus infection : finding prognostic markers and correlates of protection

Issue Date: 2018-10-16

CHAPTER

IMPACT OF CONGENITAL CYTOMEGALOVIRUS INFECTION ON TRANSCRIPTOMES FROM ARCHIVED DRIED BLOOD SPOTS IN RELATION TO LONG-TERM CLINICAL OUTCOME

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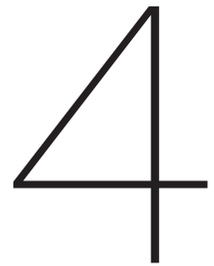
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ABSTRACT

Congenital Cytomegalovirus infection (cCMV) is the leading infection in determining permanent long-term impairments (LTI), and its pathogenesis is largely unknown due to the complex interplay between viral, maternal, placental, and child factors. The cellular activity, considered to be the result of the response to exogenous and endogenous factors, is captured by the determination of gene expression profiles. In this study, we determined whole blood transcriptomes in relation to cCMV, CMV viral load and LTI development at 6 years of age by using RNA isolated from neonatal dried blood spots (DBS) stored at room temperature for 8 years. As DBS were assumed to mainly reflect the neonatal immune system, particular attention was given to the immune pathways using the global test. Additionally, differential expression of individual genes was performed using the voom/limma function packages. We demonstrated feasibility of RNA sequencing from archived neonatal DBS of children with cCMV, and non-infected controls, in relation to LTI and CMV viral load. Despite the lack of statistical power to detect individual genes differences, pathway analysis suggested the involvement of innate immune response with higher CMV viral loads, and of anti-inflammatory markers in infected children that did not develop LTI. Finally, the T cell exhaustion observed in infected neonates, in particular with higher viral load, did not correlate with LTI, therefore other mechanisms are likely to be involved in the long-term immune dysfunction. Despite these data demonstrate limitation in determining prognostic markers for LTI by means of transcriptome analysis, this exploratory study represents a first step in unraveling the pathogenesis of cCMV, and the aforementioned pathways certainly merit further evaluation.

4.1. INTRODUCTION

Human Cytomegalovirus (CMV) is one of the most common causes of congenital viral infection, leading to a significant number of children with permanent disabilities. The overall birth prevalence of congenital CMV infection (cCMV) in industrialized countries is between 0.6% and 0.7% (1, 2). Among the congenitally infected infants, 12.7% are estimated to have symptoms at birth, ranging from mild, such as petechiae, to severe, such as microcephaly (1, 2). An estimated 40-58% of these symptomatic children develop permanent long-term disabilities, such as hearing loss, cognitive and motor developmental delay (1). Although symptomatic neonates have a considerable risk to develop permanent long-term impairments (LTI), approximately 13% of the asymptomatic children will also develop permanent LTI (1). Despite the current insights into the clinical outcome of cCMV, the multifactorial process that determines whether a child is symptomatic at birth or will develop LTI is largely unknown.

The control of cCMV, and cCMV-related disease, may be the result of a complex interaction between viral, maternal, placental, fetal and child factors (3). The clinical impact of cCMV has mainly been evaluated in relation to maternal factors, such as the CMV immune status before pregnancy or the time of vertical transmission. The vertical transmission rate is higher among women without prior CMV infection than among previously exposed women (2), indicating that pre-existing immunity can be protective. Vertical transmission occurring in the first 20 weeks of pregnancy leads to a worse clinical outcome than transmission occurring later in pregnancy (4, 5). The latter is probably related to an increased susceptibility to infection due to fetal organogenesis, and a still developing fetal immune system. Although the pathogenesis of LTI is poorly understood, the fetal and neonatal immune system likely play an important role in controlling the infection, thereby influencing LTI development (3). Several studies have demonstrated a CMV-specific adaptive immune response in congenitally infected children, such as γ and δ T cells or B cells (6-10), as well as an innate immune response (11, 12). However, only few studies have evaluated these responses in relation to clinical outcome at birth, whereas the majority has not done so in relation to LTI development. An increase of NK cells was observed in congenitally infected children, and their frequency was higher in those who were symptomatic at birth (11). In proteomic studies, an increase of macrophage-derived cytokines was observed in congenitally infected children, whereas an increase of α -defensin was observed in those who were asymptomatic at birth (12). Moreover, the cytokine profile of congenitally infected children, both asymptomatic and symptomatic, was different from that of their mothers with primary infection (13).

The gene expression profile captures a snapshot of the cellular activity which is the result of the response to genetic, environmental and epigenetic factors (14). After having established, through forensic studies, that reliable RNAs can be extracted from dried stains, a considerable amount of studies focused on neonatal dried blood spots (DBS) because they represent an important archived, and readily accessible specimen to study factors of disease development. Indeed, DBS are usually collected at birth for the screening of rare genetic metabolic disorders, and are stored for several years (15). Previous studies have shown that quantitative RNA measurements, either with microarrays or RNA-seq, can be performed on neonatal DBS stored at room temperature for up to

9 years (14, 16-18). Additionally, since the transcriptional profiles of RNA derived from DBS in mice, stored for several months at room temperature, correlated with those from fresh whole blood (19), we assumed this may also be the case in humans. The transcriptome varies according to the cell types studied, and certain RNA markers are tissue-specific. Tissue-specific RNA molecules have been successfully extracted from blood and saliva stains, dried at room temperature for up to 16 years, and used for genome-wide expression analysis (20, 21). Since DBS are produced by spotting whole blood on filter paper, they were assumed to mainly reflect the neonatal immune system.

The aim of this exploratory study was to evaluate the feasibility of transcriptome analysis from archived neonatal DBS in relation to cCMV and LTI development. In particular, we wanted to determine whether the neonatal immune system at birth may be a determinant of LTI development at 6 years of age. This would provide insights into the immune regulation of cCMV, and, by identifying prognostic markers for clinical outcome, could provide the means to introduce the long-debated newborn screening program for CMV in DBS by defining subgroups of infants that would benefit from clinical and audiological follow-up, and possibly antiviral treatment (22). Our investigations revealed that transcriptome analysis of RNA from neonatal DBS stored at room temperature for 8 years of a nation-wide retrospective cohort of children with cCMV and controls is possible, and could potentially be used to unravel the pathogenesis of cCMV and CMV-related disease.

4.2. MATERIALS AND METHODS

4.2.1. Study population and clinical data

A previously described nationwide, retrospective cohort was used in this study (23). The cohort was derived from a total group of 31,484 children, born in 2008 in the Netherlands, which was retrospectively tested for cCMV by PCR of CMV DNA in neonatal DBS at five years of age. In total, 156 children (0.5%) were diagnosed with cCMV. Clinical data were retrieved from 133 congenitally CMV-infected children and from 274 non-infected children. Children were defined as symptomatic at birth if they had one or more of the following signs or symptoms in the neonatal period: prematurity, being small for gestational age, microcephaly, hepato- or splenomegaly, generalized petechiae or purpura, hypotonia, abnormal laboratory findings (elevated liver transaminases, hyperbilirubinemia, neutropenia or thrombocytopenia), cerebral ultrasound abnormalities, ophthalmologic abnormalities or neonatal hearing impairment. LTI was defined as the presence of impairment in one or more domain (hearing, visual, neurological, motor, cognitive and speech-language). The cCMV associated LTI in the original cohort has been described in detail (24). In brief, hearing impairment was defined as sensorineural hearing loss ≥ 40 dB; visual impairment was defined as a visual acuity below 0.3; neurological impairment included cerebral palsy, epilepsy, microcephaly, autism spectrum disorder and ADHD; motor developmental delay was based upon the physical therapist's report and if available on a score below the fifth centile in the Movement Assessment Battery for Children; cognitive developmental delay was defined as an intelligence quotient less than or equal to 70 if this was tested, or it was based on a diagnosis by a medical specialist; speech and language development were assessed by the speech therapist or speech and hearing centre. Additionally, the severity of the LTI was assessed by accumulating the number of domains affected and indicated as the presence of LTI in two or more domains. Since in this cohort

maternal seroimmunity to CMV before birth was unknown, it was assumed that cCMV infection could have resulted from either maternal primary or secondary infection. Due to the retrospective design of the study, there was no standardized clinical and laboratory assessment performed at birth. Therefore, we cannot exclude the possibility that we might have misclassified some newborns without clinically apparent disease or with mild and transient symptoms in the asymptomatic group. However, because of the Dutch child health care system, the chance of having missed major signs or symptoms can be considered negligible (23, 24).

For the study presented in this article, DBS were selected based on the clinical outcome of the infants, with a total of 6 CMV-negative without any clinical signs, 6 CMV-positive with LTI and 6 CMV-positive without LTI. This study was approved by the Medical Ethics Committee of the Leiden University Medical Center, and all the parents of the children included have given written informed consent for the use of clinical data and DBS.

4.2.2. DNA extraction from DBS and qPCR of CMV

After a first initial CMV PCR screening performed at the National Institute for Public Health and the Environment (RIVM), a second confirmatory PCR was performed at the Leiden University Medical Center (LUMC) (23). For this purpose, DNA was extracted from DBS by using the QIAamp DNA minikit according to the previously described protocol (25). For each test, one full DBS was punched by using an automated DBS puncher (1296-071, Perkin Elmer-Wallac, Zaventem, Belgium). CMV DNA amplification of a 126-bp fragment from the immediate-early antigen region was performed using an internally controlled quantitative real-time PCR, as described previously (26, 27), on the CFX96 Real-Time PCR Detection System (BioRad, Veenendaal, The Netherlands). The PCR was performed in triplicate, and the CMV viral load was expressed in IU/ml.

4.2.3. RNA extraction from DBS

One full DBS was punched using an automated DBS puncher (1296-071, Perkin Elmer-Wallac, Zaventem, Belgium). RNA was extracted from DBS by using the NucleoSpin miRNA kit (Macherey-Nagel, Duren, Germany), according to the manufacturer's instructions with a minor modification. This included pre-incubating the DBS with 300 μ l of lysis buffer ML for 30 min at 37°C with agitation (1000 rpm) (28). The supernatant was transferred to the NucleoSpin Filter, and the procedure was carried out according to the manufacturer's instruction. Small and large RNAs were purified in one fraction, without separation of small RNAs, and a DNase treatment was used to reduce DNA contamination. The RNA was eluted in 50 μ l of RNase-free H₂O, and RNA integrity was assessed using the RNA Nano 6000 Assay Kit on the Bioanalyzer 2100 system (Agilent Technologies, CA, USA). The RNA concentration was measured using a Qubit 2.0 fluorometer (Life Technologies, CA, USA).

4.2.4. Library preparation and sequencing

An average amount of 185 ng of RNA was used as input material for library preparation. Sequencing libraries were generated using the TruSeq Stranded Total RNA Sample preparation kit for Illumina (Illumina, Inc., San Diego, CA, USA) following the manufacturer's recommendations, and index

codes were added to attribute sequences to each sample. Briefly, rRNA was depleted from total RNA using rRNA removal magnetic beads (RRB). The remaining RNA was purified using RNAClean XP magnetic beads. As the RNA samples from DBS were already fragmented, the fragmentation step was skipped in order to avoid over-fragmentation. First strand cDNA was synthesized using random hexamer primers and SuperScript II reverse transcriptase. Second strand synthesis was performed using the polymerase provided with the kit. After adenylation of the 3' end of the blunt-ended DNA fragments, the RNA index adapters were ligated, and PCR was carried out using the PCR master mix and primer cocktail provided by Illumina to amplify the DNA in the library that had adapter molecules on both ends. Library quality was assessed using the DNA 1000 Assay kit for the Agilent Bioanalyzer 2100 system (Agilent Technologies, CA, USA), and the DNA amount was measured using a Qubit 2.0 fluorometer. Clustering of the index-coded samples was performed using the Illumina TruSeq PE Cluster Kit v3 (cBot-HS) according to the manufacturer's instructions. After cluster generation, the libraries were sequenced on the Illumina HiSeq 2000 platform (6 samples per lane), and 76 base paired-end reads were generated for the first batch (n=6, 2 of each group) and 50 base paired-end reads for the second batch (n=12, 4 of each group). All 76 base reads were trimmed to 50 bases to allow for uniform subsequent analysis across all samples, and the batch effect was accounted for in downstream analysis. Due to lack of resources it was not possible to sequence the whole cohort.

4.2.5. Read mapping to the reference genome

Sequence files were generated in FASTQ format, and all RNA sequence files were processed using the BIOPET Gentrapp pipeline version 0.7 developed at the LUMC (http://biopet-docs.readthedocs.io/en/latest/releasenotes/release_notes_0.7.0/). The BIOPET Gentrapp pipeline consists of FASTQ pre-processing (including quality control, quality trimming and adapter clipping), RNA-seq alignment, read and base quantification. FastQC version 0.11.2 was used for raw read quality control. Low quality read trimming was done using sickle version 1.33 with default settings. Cutadapt version 1.9.1 with default settings was used for adapter clipping based on the detected adapter sequences by FastQC toolkit. RNA-seq reads were aligned against human reference genome GRCh38 using RNA-seq aligner GSNAP version 2014-12-23 with settings "--npaths 1 --quiet-if-excessive". Ensembl human genome annotation version 87 was used for raw read counting. The gene read quantification step was performed using htseq-count version 0.6.1p1 with the setting "--stranded=reverse".

4.2.6. Differential expression analysis: individual genes

We identified significant gene expression differences between congenitally infected children (n=12) and controls (n=6), as well as between congenitally infected children that developed LTI (n=6) and congenitally infected children that did not develop LTI (n=6). Moreover, we also assessed gene expression differences in relation to logarithm of CMV viral load treated as continuous variable. Genes with low fragment counts were removed by requiring at least 2 fragments per million of aligned fragments to be observed in at least 2 samples. Library size normalization factors were obtained with the trimmed mean of M-values (TMM) method (29). Linear modelling using Bioconductor/R package 'limma' (30) was performed on read counts transformed to log-CPM values. Observational-

level weights obtained from the voom function were used to model mean-variance relationship. All three analyses were corrected for the batch effect in the design matrix. Multiple testing correction using false discovery rate control of Benjamini and Hochberg was performed at the threshold of 0.05.

4.2.7. Differential expression analysis: pathways

The Bioconductor/R package 'global test' designed by J. Goeman was used to evaluate differences in expression profiles of gene sets between the different groups (31). These were a group of congenitally infected children (n=12) and a group of controls (n=6). Within the group of congenitally infected children, those that developed LTI (n=6) and those that did not develop LTI (n=6). An additional analysis was performed to find gene set expression profiles dependent on CMV viral load as continuous variable. This method has been shown to have more power to detect gene sets with small effect size (29, 32, 33). We selected a limited number of candidate gene sets (pathways) for use in the global test, before inspecting the data using the QuickGO browser (34). The pathways were selected based on their putative role in the etiology of the disease. An additional selection criterion was the specimen, i.e. DBS, which derives from whole blood and therefore mainly reflects the neonatal immune system. These pathways were T-, B-, and NK-cell activation, innate immune response, and inflammatory response with its regulation. Each pathway contained from 17 to 435 genes. This analysis was performed on the voom-transformed data. Due to the exploratory nature of this study, and to the limited number of selected pathways, no multiple testing correction was applied.

Finally, an additional immune pathway that has emerged as one of the possible players in limiting the immune response during cCMV is the T cell exhaustion (7). However, this does not exist yet as a pathway in the QuickGo browser. Therefore, based on the transcriptional definition of exhaustion previously described (7, 35), and on our available data, a set of exhaustion genes was selected. An independent sample t-test was used to evaluate the difference in the square root of the reads per million (RPM) between the different categories. CMV+ vs CMV-, CMV+ without LTI vs CMV+ with LTI, CMV+ low load vs CMV+ high load. In the latter, the infected group was split in two according to the median log₂ viral load measured in DBS which was 10.2, namely low (< 10.2) and high (≥ 10.2) viral load groups. However, p-values were not reported because this analysis had the sole purpose of illustrating trends.

4.3. RESULTS

4.3.1. Study population and clinical data

The clinical data of the congenitally infected children included in this study, as well as of the non-infected controls, are listed in Table 1. A total of 12 children with cCMV, and 6 without cCMV, were included in order to assess the gene expression profile in relation to cCMV. Additionally, the 12 children with cCMV were selected in order to assess differences in gene expression in relation to LTI development. For this purpose, 6 infected children were selected, who did not have any symptoms at birth nor LTI at six years of age, whereas the other 6 had LTI in one or more of the following

Table 1. Study population and clinical outcome

	cCMV with LTI ¹ n = 6	cCMV no LTI ² n = 6	No cCMV ³ n = 6
Gender			
Male	4	3	3
Female	2	3	3
Gestational age (weeks)⁴	39 (36-40)	40 (37-41)	41 (37-41)
Birth weight (g)⁴	3040 (1890-4040)	3340 (2760-4240)	3298 (3070-4360)
CMV viral load⁵	3.1 (2.43-4.97)	3.1 (2.18-4.30)	-
Long term impairment			
Hearing impairment ⁶	0	0	0
Visual impairment ⁷	0	0	0
Neurological impairment ⁸	3	0	0
Motor impairment ⁹	6	0	0
Cognitive impairment ¹⁰	4	0	0
Speech/language problem ¹¹	4	0	0
More than one impairment¹²	5	0	0

¹ Congenitally infected children that develop LTI, 5 out of 6 had symptoms at birth including prematurity (n=1), dysmaturity (n=1), microcephaly (n=3); ² Congenitally infected children that did not develop LTI, none of them had symptoms at birth;

³ Non-infected controls, none of them had symptoms at birth nor LTI; ⁴ Values are medians with minimum and maximum; ⁵ CMV viral load measured on DBS, values are log (IU/ml) medians with minimum and maximum; ⁶ Sensorineural hearing loss \geq 40 decibels; ⁷ Optic nerve atrophy or cortical visual impairment; ⁸ Cerebral palsy (n=1), epilepsy (n=1), microcephaly (n=1), autism (n=2), ADHD (n=1); ⁹ Motor impairment (fine, gross or balance) based on test or diagnosis or sensory processing disorder or developmental coordination disorder (n=6); ¹⁰ Cognitive impairment based on test or diagnosis (n=4); ¹¹ Language impairment based on test or diagnosis, speech-impairment, oral motor skill difficulties or auditory processing disorder (n=4);

¹² Impairment in two or more domains of impairment: hearing, visual, neurologic, motor, cognitive, and speech-language.

domains of impairment: neurological, motor, cognitive and speech/language (Table 1). Five children out of those who developed LTI also had symptoms at birth. Importantly, none of the children in the control group had symptoms at birth nor developed LTI. Given the diversity of the specific symptoms at birth and impairments at the age of six, the subjects were selected in order to have a similar proportion of male and female across the groups. In this way, the influence of gender in the gene expression analysis was limited.

4.3.2. Library preparation and sequencing statistics

The average number of RNA-seq read pairs per sample was 38.5 million \pm 4.8 million, with 38.9 million \pm 5.4 million for the CMV- samples and 38.4 million \pm 4.7 million for the CMV+ samples. Within the CMV+ samples, those without LTI generated 37.6 million \pm 5.9 million paired-end reads, and those with LTI generated 39.1 million \pm 3.6 million paired-end reads. The mean RNA fragment size was 285 \pm 8 bp, and the mean DNA fragment size was 165 \pm 8 bp. On average, 92.25 % of bases exceeded Q30. The detailed information per sample is shown in Table 2.

Table 2. RNA-seq data per individual

ID ¹	cCMV ²	Gender ³	LTI ⁴	Input RNA (ng) ⁵	RNA fragment size (bp)	DNA fragment size (bp)	Total number of read pairs ⁶	Total bases ⁷	Raw bases Q10+ ⁸	Raw bases Q20+ ⁹	Raw bases Q30+ ¹⁰
1	CMV-	m	no	160	274	154	36559606	3623399872 (99.1%)	3582227412 (98.0%)	3398394726 (93.0%)	
2	CMV-	f	no	200	286	166	33157384	3286143611 (99.1%)	3247394425 (97.9%)	3076116879 (92.8%)	
3	CMV+	f	no	200	278	158	29540831	2927528373 (99.1%)	2892987291 (97.9%)	2740849118 (92.8%)	
4	CMV+	f	no	200	282	162	31300956	3102402121 (99.1%)	3066377738 (98.0%)	2905730026 (92.8%)	
5	CMV+	f	yes	120	281	161	33323826	3302791747 (99.1%)	3265370156 (98.0%)	3096491305 (92.9%)	
6	CMV+	m	yes	200	282	162	39311864	3897307546 (99.1%)	3853368879 (98.0%)	3657532714 (93.0%)	
7	CMV-	f	no	200	285	165	45571592	4499488134 (98.7%)	4432680661 (97.3%)	4219189147 (92.6%)	
8	CMV+	f	yes	200	278	158	42119123	4158027595 (98.7%)	4097388499 (97.3%)	3904204865 (92.7%)	
9	CMV+	m	no	140	286	166	43750889	4319612015 (98.7%)	4258366454 (97.3%)	4066273713 (92.9%)	
10	CMV-	f	no	200	282	162	35039051	3463528125 (98.8%)	3416670543 (97.5%)	3265849570 (93.2%)	
11	CMV+	m	yes	200	289	169	40100801	3951127179 (98.5%)	3878646166 (96.7%)	3587779279 (89.5%)	
12	CMV+	m	no	200	298	178	41322383	4081237277 (98.8%)	4023236869 (97.4%)	3838652400 (92.9%)	
13	CMV-	m	no	200	287	167	45568773	4486847437 (98.5%)	4416030829 (96.9%)	4204748423 (92.3%)	
14	CMV+	m	yes	200	289	169	36627608	3618784465 (98.8%)	3566791324 (97.4%)	3397972122 (92.8%)	
15	CMV+	f	no	200	302	182	41894051	4140976514 (98.8%)	4084677610 (97.5%)	3904733403 (93.2%)	
16	CMV-	m	no	200	271	151	37242565	3638553568 (97.7%)	3565521893 (95.7%)	3317637210 (89.1%)	
17	CMV+	m	yes	140	290	170	43235245	4228806228 (97.8%)	4149633476 (96.0%)	3947504971 (91.3%)	
18	CMV+	m	no	170	282	162	38075664	3753496088 (98.6%)	3696000143 (97.1%)	3467378795 (91.1%)	

¹ID child identification number; ²cCMV, congenital Cytomegalovirus infection; CMV+, congenitally infected children; CMV-, non-infected controls; ³f, female; m, male; ⁴LTI, long-term impairment at 6 years of age; ⁵Input RNA (ng) the amount of RNA used as input material for library preparation; ⁶Total number of paired-end reads, total number of paired-end reads that passed Illumina filter generated per sample; ⁷Total bases, total number of bases generated per sample; ⁸Raw bases Q10+, base calls with quality Q-scores of Q10+ (Q10 or higher) have an error probability of 0.1 (1 in 10) or less; ⁹Raw bases Q20+, base calls with Q20+ have an error probability of 0.01 (1 in 100) or less; ¹⁰Raw bases Q30+, base calls with Q30+ have an error probability of 0.001 (1 in 1,000) or less.

4.3.3. Differential expression: individual genes

Next, we determined whether any other gene could be associated with cCMV, LTI development at 6 years of age or CMV viral load. After low count features removal, ~25% of counts aligned on features and 18360 different genes were used in gene expression analysis. The R package LIMMA was used for the assessment of differential expression of individual genes between congenitally infected children (n=12) and non-infected controls (n=6). No statistically significant differences in gene expression were observed between the groups. We next assessed gene expression differences in relation to cCMV clinical outcome by comparing congenitally infected children that developed LTI at six years of age (n=6) to congenitally infected children that did not develop LTI (n=6). This analysis did not reveal any statistically significant differences between the groups. Finally, the differences in gene expression were assessed in relation to the logarithm of CMV viral load as continuous variable, and no statistically significant differences were observed.

4.3.4. Differential expression: pathways

In order to evaluate whether different biological mechanisms may underlie different clinical outcomes, a global test was performed on manually pre-selected pathways based on their putative role in the etiology of cCMV disease. The selected pathways for T-, B-, and NK-cell activation, innate immune response, and inflammatory response were assessed in relation to cCMV, LTI development at 6 years of age and CMV viral load. The results are shown in Table 3. This analysis revealed trend significant results in relation to CMV viral load and LTI development. In particular, the innate immune response (p=0.046, Fig 1) and the NK-cell activation (p=0.086) may be associated to CMV viral load; whereas the regulation of inflammatory response (p=0.077, Fig 2) to LTI development. In all cases, a small number of genes appeared to be responsible for these trends. Several antiviral genes were positively associated with CMV viral load, i.e. ISG15 and RSAD2, whereas the anti-inflammatory cytokine IL-4 was associated with the congenitally infected children that did not develop LTI.

Table 3. Global Test analysis

Pathways	cCMV ¹	LTI ²	CMV viral load ³
	p-values		
Innate immune response (435 genes)	0.706 ⁴	0.432	0.046
T cell activation (51 genes)	0.375	0.203	0.195
B cell activation (30 genes)	0.367	0.125	0.254
NK cell activation (17 genes)	0.717	0.499	0.086
Inflammatory response (381 genes)	0.725	0.341	0.232
Regulation of inflammatory response (68 genes)	0.577	0.077	0.567
Negative regulation of inflammatory response (78 genes)	0.633	0.339	0.133
Positive regulation of inflammatory response (74 genes)	0.444	0.652	0.791

¹ Gene sets expression differences between CMV- (n=6) and CMV+ (n=12); ² Gene sets expression differences between congenitally infected children with LTI (n=6) and without LTI (n=6); ³ Gene sets expression differences according to CMV viral load (continuous variable).

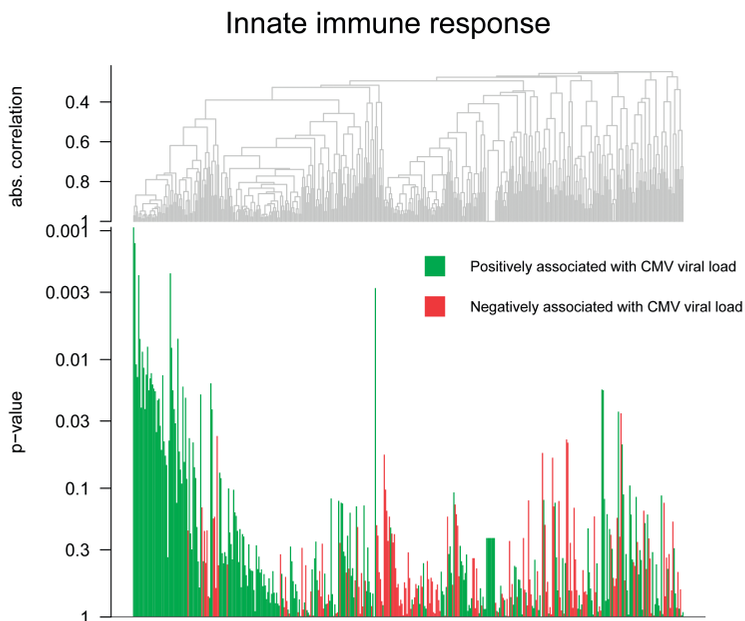


Figure 1. Global test: Innate immune response. Innate immune response in relation to CMV viral load as continuous variable measured on DBS, $p=0.046$. The gene names of x-axes are provided in supplementary S1 Table.

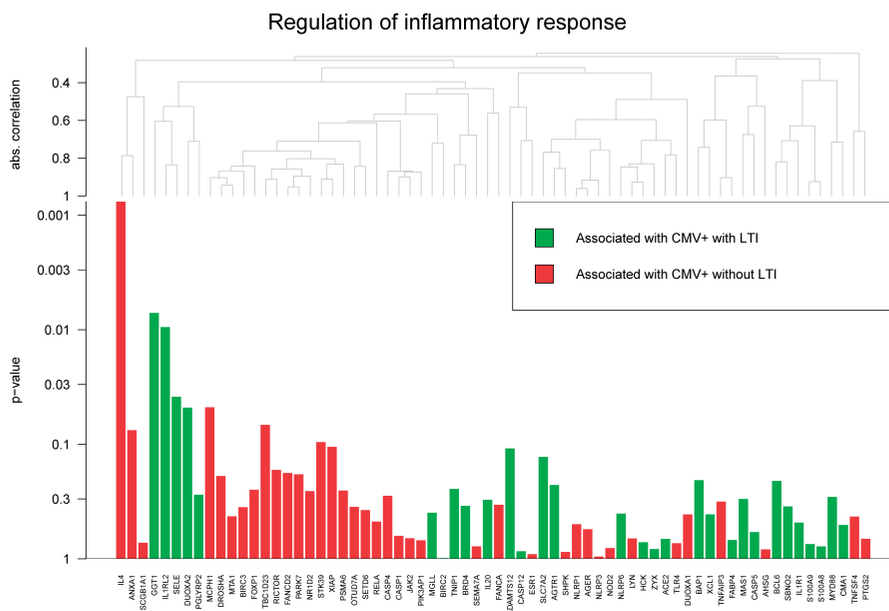


Figure 2. Global test: Regulation of inflammatory response. Regulation of inflammatory response in congenitally infected children that developed LTI at 6 years of age ($n=6$) and in congenitally children that did not develop LTI ($n=6$), $p=0.077$.

Finally, as previously shown by others, one of the possible mechanisms limiting the T cell response to CMV during early life is considered to be T cell exhaustion (7). Therefore, we wondered whether the same phenomenon could be observed in our cohort when comparing the CMV- group (n=6) to the CMV+ group (n=12). Additionally, this pathway was assessed in relation to CMV viral load and development of LTI at 6 years of age. For this purpose, based on the transcriptional definition of exhaustion previously described (7, 35), as well as on our available data, a set of genes was selected and reported in Table 4. Of these genes, the RPM were reported for each comparison in order to observe the trend to be further explored. A trend of increased expression of differentiation markers, mainly CD57 and transcription factor T-bet, and of increased effector markers, primarily IFN- and granzyme, was observed in the CMV+ group compared to the CMV- group (Fig 3 A-D). Furthermore, a trend of increased expression of inhibitory markers, mainly PD-1 and LAG-3, was observed in the CMV+ group (Fig 3 A-D). Next, the CMV+ group was split in two groups according to the median log₂ viral load measured in DBS which was 10.2, namely low and high viral load groups. Comparing the group with high viral load to the one with low viral load, the aforementioned observed trends relative to differentiation, effector and inhibitory markers were more pronounced than when comparing CMV+ to CMV-. Finally, when comparing the cCMV+ group that developed LTI to those who did not, no striking trends were observed (Fig 3 A-D).

Table 4. T cell markers

T cell markers
Differentiation and effectors¹
IFN γ
IL-2
MIP-1 β
TNF- α
Granzyme B
Perforin 1
CCR5
CD57
Transcription factors²
T-bet
Blimp-1
Inhibitory receptors
PD-1
LAG-3
FAS-L
Inhibitory cytokines
IL-10
TGF- β

¹ Markers defining a differentiation phenotype that leads to a functional response; ² Key transcription factors for T cell differentiation and exhaustion

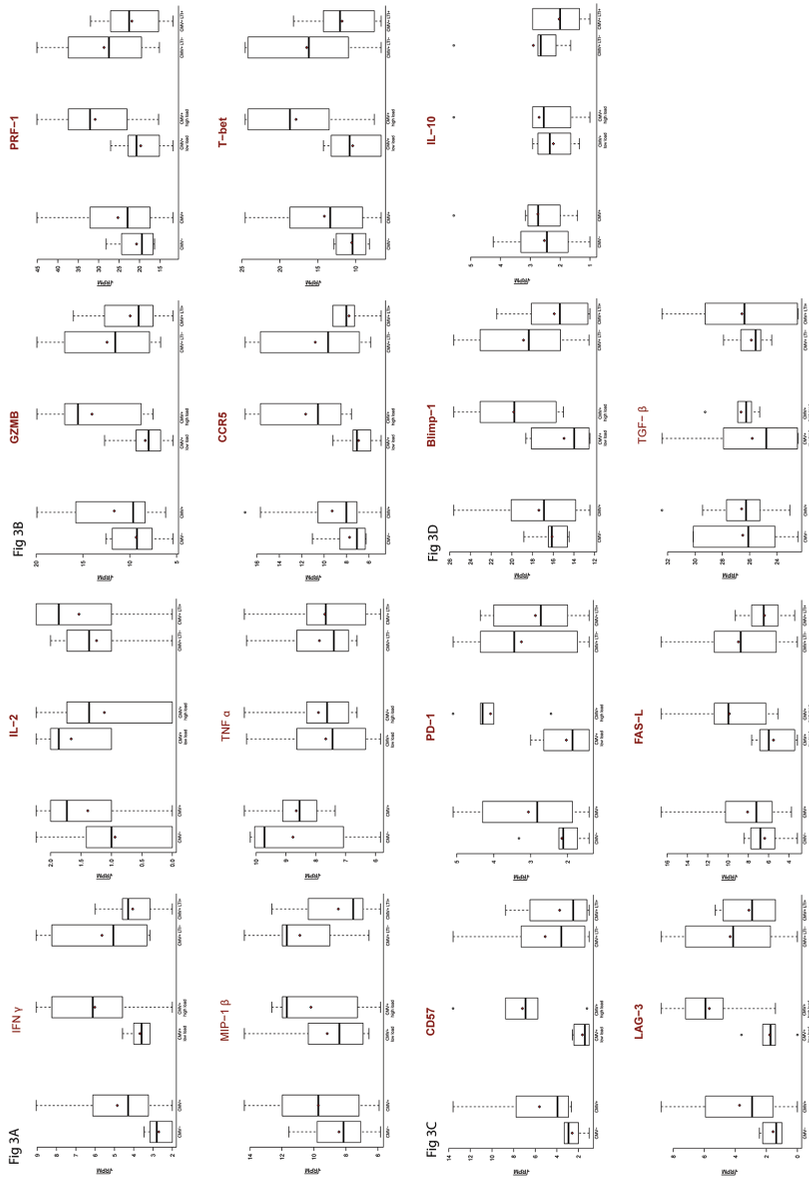


Figure 3. A-D. T cell exhaustion. T cell markers identifying the exhaustion phenotype in relation to cCMV, CMV viral load and LTI development at 6 years of age. *CMV+*, non-infected controls (n=6); *CMV+*, congenitally infected children (n=12); *CMV+ low load*, congenitally infected children with log₂ CMV viral load below the median measured in DBS which was 10.2 (n=6); *CMV+ high load*, congenitally infected children with log₂ CMV viral load equal to or above the median measured in DBS (n=6); *CMV+ LTI-*, congenitally infected children that did not develop LTI (n=6); *CMV+ LTI+*, congenitally infected children that developed LTI (n=6). Boxplot: bold line, median of square root of RPM; red dot, mean of square root of RPM.

4.4. DISCUSSION

This study aimed to evaluate whether transcriptome analysis by next generation RNA sequencing on DBS derived from a retrospective nation-wide cohort of children with cCMV and controls is feasible, and whether useful insights could be obtained on the etiology of different cCMV outcomes. This would allow the identification of potential biomarkers for long-term outcome, which could provide the means to introduce the long-debated newborn screening program for cCMV in DBS (22). Indeed, this would define subgroups of children benefitting from clinical, audiological follow-up, and possibly antiviral treatment.

The global test for differential expression of gene sets revealed, although only with trend significant results, an important feature of cCMV in relation to whole blood transcriptome, i.e. CMV viral load is the main factor to influence the pre-selected immune pathways, whereas CMV disease seems to be secondary. In our study, numerous antiviral genes were positively associated with CMV viral load, suggesting the involvement of the innate immune system in response to cCMV in the newborns, in particular with higher viral loads. The fact that no striking differences were observed when comparing CMV+ to CMV-, suggests that the high viral load is the main initiator of this expression pattern. Therefore, the presence of neonates with low viral load in the CMV+ group may have diluted the differences between CMV+ and CMV-. Congenitally infected children excrete CMV for several years after birth, whereas in adults this lasts only several months (36, 37), suggesting a deficient cell-mediated immune response in early life (38). Therefore, it is tempting to speculate that the activation of the innate immunity in the fetus may have an important role in controlling cCMV, however this is difficult to determine. One of the possible mechanism for this limited T cell response to CMV during early life is considered to be T cell exhaustion (7). In our cohort, also the exhaustion pathway was more pronounced in the high viral load group compared to the low viral load group, with PD-1 being the marker influenced the most, as previously shown (7). Therefore, also in this case the difference in exhaustion pathway between CMV+ and CMV- could have been diluted because of the presence of low viral load individuals in the CMV+ group. However, the exhaustion pathway analysis needs further confirmation as we only reported expression trends. T cell exhaustion is characterized by loss of T cell functions, and is induced by persistent infections (7, 35). Primary CMV infection induces functional T cell exhaustion in both adults and fetuses, though considerably more pronounced in the latter. As this phenomenon is associated with prolonged exposure with higher viral loads, the high viral loads reported in fetuses may be the cause of this effect (39-41). The exhaustion may contribute to the prolonged CMV viral excretion in the children (7). The influence of viral load in the immune responses has been shown before, both in humans and in the murine models of CMV infection. Here, the degree of CMV-specific memory CD8 T cells accumulation, as well as the phenotypic T cell profile, was influenced by the viral load (42, 43). However, the role of CMV viral load in the clinical outcome still remains controversial. Some studies have correlated CMV viral load, measured in blood, with clinical outcome (44, 45), whereas others have not (46-48). The neonatal viral load may differ depending on the trimester of vertical transmission, or whether it was a primary maternal infection. Indeed, earlier infections may lead to a more extensive cCMV. However, in our cohort this is impossible to establish (49). Additionally,

CMV viral load in whole blood may not correlate to CMV loads in other neonatal compartments, and therefore may not fully reflect viral replication in all affected organs and tissues.

The molecular mechanisms of LTI development are largely unknown, though the late-onset hearing loss is believed to be the result of a chronic productive infection throughout childhood (50, 51). In this context, a long-term dysfunctional immune response seems plausible, although it cannot be excluded that such dysfunction leads to a parallel uncontrolled inflammation that contributes to tissue damage. In studies of characterization of tissue damage in fetuses of 20-21 weeks of gestation with cCMV, an association between the degree of tissue damage in the brain, as well as in the inner ear, with viral load, inflammatory response and placental functionality was shown (52, 53). A dysfunctional immune response that leads to uncontrolled viral replication, and immune-mediated damage was suggested. Therefore, a similar pathogenesis may be assumed when such infection becomes chronic. The exhaustion pathway that was found in congenitally infected children, especially those with higher CMV viral load, did not seem to correlate to clinical outcome at 6 years of age. This suggests that other mechanisms are involved in the long-term immune dysfunction. In our cohort, when comparing congenitally infected children that developed LTI to those infected who did not, a role for the regulation of inflammatory responses seemed to partially contribute. Anti-inflammatory markers, such as the cytokine IL-4, were associated with congenitally infected children that did not develop LTI. The success of an immune response is the result of a balance between effector and regulatory mechanisms, therefore, the potential protective effect of IL-4 in those infected children that did not develop LTI may lie in its anti-inflammatory property. Interestingly, in a cohort of healthy CMV infected individuals, the CD4 T-cell response associated with a protective immunity involved cytokine production of IFN, and/or IL-17, in association with IL-4 (54). Similarly to IL-10, IL-4 has been shown to possess the capacity of down-regulating the production of pro-inflammatory mediators by microglia, both in humans and in mice (55-57), and its neuroprotective effect was associated with downregulation of brain inflammation in mice (58). When studying the regulation of the inflammatory response in children with cCMV and compare the group with LTI to that without LTI, we have to be aware that there may be other perinatal factors influencing the inflammatory pattern in DBS. Although we cannot fully exclude a role for non-cCMV related perinatal factors, there was no bacterial amniotic infection or neonatal sepsis in all children included in this study.

Several reasons may have contributed to the fact that we did not find a strong impact of cCMV on whole blood transcriptomes from DBS. First of all, one of the groups of congenitally infected children did not have symptoms at birth nor LTI, which is the case in most children with cCMV, and the clinical signs of symptoms associated with LTI are very diverse. Second of all, in our cohort, the fetal infection may have been the result of a primary or secondary CMV infection in the mother, and may have taken place at any time during pregnancy, especially in the asymptomatic children. Third of all, the small sample size of the groups may have led to a lack of power both in the gene expression analysis of individual genes, as well as in the pathway analysis. Lastly, the RNA degradation on these specimens, due to e.g. ribonucleases, pH, humidity or UV light, may have contributed to the lack of significant differences among the sample groups. The degradation of RNA from dried stains has been extensively studied in forensic studies for obvious reasons, and several RNAs have

been extracted from numerous conditions (59-64). From these studies, determinants for RNA stability appeared to be the specimen the RNA is extracted from, and the specific RNA molecule analyzed. In the former, the detection limit of blood-specific RNA has been shown to be lower than for other specimens (21). In the latter, some RNAs can be more stable in dried stains than others (21). Secreted RNAs, e.g. in fresh saliva, may be more susceptible to fast degradation by extracellular RNases, and therefore are not to be expected on dried stains (20). Importantly, for those RNAs detected on dried blood stored at room temperature, only few genes have been demonstrated to be differentially expressed during time (20). Therefore, we assumed that those markers detected on DBS in our study were less prone to degradation, and relatively stable for long periods of time. Furthermore, the influence of RNA contamination in the downstream analysis, e.g. from skin cells or external microorganisms, may be considered negligible as the most abundant RNAs species come from the host whole blood (65). Despite the fact that enough data were generated in our study for the downstream analysis, with comparable cDNA fragment size as shown in forensic studies (21, 59-64), we cannot exclude that fresh material may have revealed differences in expression patterns that we could not pick up.

Furthermore, due to the retrospective nature of the study, cCMV diagnosis was carried out by performing PCR of viral DNA on DBS, which in comparison with PCR on urine or saliva has been associated with limited and variable sensitivity (66). Therefore, a negative CMV PCR on DBS does not fully exclude cCMV. However, it is important to note that with the relatively high sensitivity of our CMV PCR on DBS (estimated > 85%), high specificity (> 99.9%) and the cCMV birth prevalence of 0.5%, the chance of a CMV false-negative result is 1/1000 (23). Therefore, it is very unlikely that a cCMV positive child ended up in our cCMV negative control group.

To the best of our knowledge, this is the first exploratory study assessing the feasibility of transcriptome sequencing using RNA isolated from archived neonatal DBS of children with cCMV, and non-infected controls, in relation to long-term outcome. Despite the lack of statistical power to detect individual gene expression differences, the pathway analysis suggested a potential differential gene expression in relation to CMV viral load and LTI. Therefore, this study represents a first step in unraveling the pathogenesis of cCMV, and in identifying prognostic markers for cCMV long-term outcome.

4.5. ACKNOWLEDGMENTS

The CROCUS study was initiated and supported by the National Institute of Public Health and the Environment (RIVM), we thank Marjolein Korndewal for the use of the CROCUS study clinical data. We thank Daniela Balzereit, Simon Dökel, Alexander Kovacovics and Matthias Linser (Otto Warburg Laboratory Gene Regulation and Systems Biology of Cancer, Max Planck Institute for Molecular Genetics, Berlin, Germany) for sequencing library preparation and Illumina HiSeq sequencing.

4.6. AUTHOR CONTRIBUTIONS

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4.7. DATA AVAILABILITY STATEMENT

The dataset underlying this study cannot be shared publicly, as it would violate restrictions imposed by the Medical Ethical Committee of Leiden University Medical Center (LUMC). Specifically, the authors must restrict the full genomic data of the participants of the study in order to protect participant privacy. A minimal underlying data set containing the read counts per gene value is available in the Supporting Information files. Interested and qualified researchers may request access to the full dataset by contacting Eric C.J. Claas, Molecular Biologist of the Medical Microbiology Department of LUMC, E.C.J.Claas@lumc.nl.

4.8. FUNDING

This work was supported by European Union Seventh Framework Programme FP7/2012-2016 under grant agreement number 316655 (VACTRAIN). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. HL is a founder and the chairman of Alacris Theranostics GmbH, Berlin, Germany. Alacris Theranostics GmbH provided support in the form of salary for author HL, but did not have any additional role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript. The specific role of this author is articulated in the 'author contributions' section.

4.9. COMPETING INTERESTS

We have the following interests: HL is a founder and the chairman of Alacris Theranostics GmbH, Berlin, Germany. There are no patents, products in development or marketed products to declare. This does not alter our adherence to all the PLOS ONE policies on sharing data and materials.

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SUPPORTING DATA

Table S1. Gene names. Gene names of x-axis of Fig 1 concerning the innate immune response in relation to CMV viral load as continuous variable. (XLSX)

Genes	p-value	Genes	p-value	Genes	p-value
ISG15	0,001	IFI16	0,043	CLEC5A	0,142
RSAD2	0,001	SPON2	0,044	PRKD1	0,149
GSDMD	0,002	GPBR1	0,044	CLEC10A	0,150
OASL	0,002	TNK1	0,046	TRIM14	0,154
IGHG1	0,003	DHX58	0,050	IGHG4	0,155
PRDM1	0,007	B2M	0,052	AIM2	0,159
OAS3	0,007	IL36RN	0,053	SIGLEC15	0,159
BST2	0,008	CD300LB	0,055	MST1R	0,160
NLRCS	0,008	TRIM21	0,056	CSK	0,160
IFIT3	0,009	NR1H4	0,058	TRIM26	0,164
PML	0,011	SLAMF7	0,062	IL36B	0,165
HERC5	0,012	MX2	0,066	LY86	0,166
IFIH1	0,013	KLRD1	0,069	SIRPB1	0,166
SERPING1	0,013	ELF4	0,072	CFI	0,167
ADAR	0,013	SARM1	0,076	IGHV3OR16-9	0,170
APOL1	0,014	MATK	0,083	IL1RL2	0,172
EIF2AK2	0,014	ZAP70	0,083	PGLYRP2	0,182
JAK3	0,015	APOBEC3D	0,084	SERINC3	0,186
IFIT5	0,016	TBK1	0,092	DEFB116	0,192
ZNF683	0,016	PTK2B	0,095	IFNA6	0,193
MX1	0,017	IRAK1	0,100	SRC	0,195
IFITM3	0,017	FCN1	0,102	HMGB2	0,200
FCGR1A	0,017	TRIM5	0,103	IFNL1	0,200
IRF7	0,017	IFNA21	0,108	FGA	0,202
DDX60	0,017	APCS	0,112	SRPK1	0,205
OAS2	0,018	UBC	0,114	IL27	0,207
APOBEC3F	0,019	SIGLEC14	0,117	TRIM11	0,208
APOBEC3G	0,020	IFNA13	0,120	NLRX1	0,213
IFITM1	0,020	IRF1	0,121	F12	0,213
IFIT1	0,021	APOBEC3C	0,123	MR1	0,215
ZBP1	0,024	IL23A	0,124	FES	0,215
ISG20	0,024	IGHG2	0,125	NFKB1	0,221
IFIT2	0,024	ANXA1	0,126	KLRG1	0,225
TBKBP1	0,024	TRIM15	0,129	CFP	0,225
NLRP2	0,025	IGHA1	0,130	GATA3	0,227
C4B	0,026	BPIFB1	0,130	NLRP3	0,227
ZC3HAV1	0,031	NOD1	0,130	FBXO9	0,230
DDX58	0,033	JCHAIN	0,131	BPIFB3	0,233
OAS1	0,037	NRROS	0,134	IGHV3-23	0,234
DEFB129	0,039	TNK2	0,136	KLRK1	0,238
SLAMF6	0,041	SLAMF1	0,138	TEC	0,238
DEFB114	0,042	VSIG4	0,138	IKBKG	0,243
TRIM25	0,043	FRK	0,141	DEFB105B	0,244

Table S1. (continued)

Genes	p-value	Genes	p-value	Genes	p-value
DEFB106A	0,244	CD84	0,350	APP	0,448
DEFB4B	0,244	DEFB107A	0,352	S100A9	0,449
DEFB103B	0,244	DEFB113	0,356	IFNA10	0,452
DEFB104B	0,244	TRIM4	0,357	TRIM13	0,453
RP11-330H6.5	0,244	CORO1A	0,358	SSC5D	0,454
DEFB103A	0,244	C4A	0,359	MAP3K5	0,456
C4BPA	0,245	GZMM	0,361	PPP1R14B	0,456
CARD9	0,257	IL1RAP	0,368	SH2D1B	0,460
IL23R	0,257	CASP4	0,369	ABL2	0,463
KLRC2	0,260	IFNL2	0,369	IRAK4	0,464
C8B	0,261	SUSD4	0,369	SLPI	0,467
PTK6	0,264	YES1	0,371	SH2D1A	0,474
APOBEC3B	0,266	COLEC12	0,374	TICAM1	0,478
DEFA3	0,266	TRIM56	0,375	TLR10	0,479
SAA1	0,270	C1R	0,380	CYBB	0,482
TRIM62	0,272	C5	0,381	FER	0,488
SIRT2	0,273	UNC93B1	0,382	MEFV	0,496
CIQBP	0,278	CD6	0,383	DEFB133	0,497
MB21D1	0,281	DEFB134	0,383	RNF135	0,498
SMPDL3B	0,283	MASP2	0,390	IPO7	0,500
TRIM28	0,283	DEFB115	0,391	DEFA1	0,503
CACTIN	0,288	CAPZA1	0,396	TREM1	0,508
LGALS3	0,290	KRT16	0,396	TREML1	0,508
IFNA5	0,292	ANKRD17	0,397	MAP4K2	0,511
APOBEC3A	0,295	PGLYRP4	0,397	TLR3	0,511
HMGB1	0,299	C9	0,401	MIF	0,515
CHID1	0,303	PGLYRP3	0,401	PIK3CG	0,515
BLK	0,311	FCN2	0,410	ITCH	0,518
POLR3H	0,313	CYLD	0,417	SRMS	0,519
RNASE7	0,313	CSF1R	0,417	CSF1	0,520
CLEC7A	0,315	IRGM	0,420	DEFB132	0,523
LCK	0,319	IFNL3	0,421	AKAP8	0,527
NCR2	0,324	TRIM35	0,423	C4BPB	0,530
DMBT1	0,324	IGHE	0,423	TLR6	0,531
CIQB	0,324	DEFB125	0,423	TRIM10	0,532
DEFA5	0,326	RPS27A	0,425	NLRP1	0,538
IFITM2	0,327	CD180	0,427	CD1D	0,553
CR2	0,336	OTULIN	0,427	PYCARD	0,554
POLR3D	0,337	PADI4	0,433	TRIL	0,554
C2	0,340	ITK	0,436	TMEM173	0,556
JAK2	0,344	DEFB126	0,439	TLR8	0,560
IFNA14	0,344	FYN	0,440	NLRC4	0,562
DAB2IP	0,345	DEFA4	0,440	TRIM38	0,569
ECSIT	0,349	ATG5	0,440	TRIM8	0,570
AXL	0,349	PIK3CD	0,445	LILRA5	0,571
ADAM15	0,349	FCN3	0,447	POLR3A	0,574

Table S1. (continued)

Genes	p-value	Genes	p-value	Genes	p-value
DEFB105A	0,575	PSTPIP1	0,672	TLR9	0,799
ATG12	0,576	CRISP3	0,677	MBL2	0,804
SYK	0,582	DEFB107B	0,678	BCL10	0,806
CIQC	0,583	ACOD1	0,691	IFNA4	0,809
F2RL1	0,584	VNN1	0,691	DEFB108B	0,814
CRCP	0,588	DEFA6	0,693	CLEC4M	0,814
APOBEC3H	0,590	CLEC4D	0,697	IFNA2	0,815
CD300E	0,591	BPIFA1	0,699	REL	0,817
LGR4	0,595	MASP1	0,701	IRF5	0,819
AGER	0,596	SFTPD	0,703	IGLL1	0,820
C1S	0,597	TIRAP	0,704	NOD2	0,822
CYBA	0,599	FADD	0,704	DEFB106B	0,823
TYK2	0,600	RELB	0,709	DEFB104A	0,826
BMX	0,602	SEC14L1	0,709	IL36A	0,828
CD209	0,610	RIPK2	0,715	DEFB4A	0,837
CLEC4E	0,614	LY9	0,715	TRIM27	0,841
NAIP	0,615	S100A7	0,715	STYK1	0,848
ANKHD1	0,618	TNFAIP8L2	0,718	MYD88	0,848
DEFB131	0,618	C1RL	0,723	DEFB118	0,850
DEFB123	0,618	ZBTB1	0,727	TRIM31	0,851
DEFB119	0,620	IL36G	0,727	NFKB2	0,853
CTD-2313N18.7	0,622	TKFC	0,730	ANG	0,854
DEFB110	0,622	TLR4	0,732	MID2	0,855
PCBP2	0,622	CNPY3	0,735	NLRP2B	0,859
PPARG	0,626	CFB	0,737	C7	0,862
DEFB135	0,629	CRI	0,738	FGR	0,862
AKIRIN2	0,630	TYROBP	0,740	TICAM2	0,863
TRAF3	0,630	CFD	0,744	CAPZA2	0,864
LCN2	0,631	S100A8	0,745	TOLLIP	0,867
BTK	0,634	C8A	0,749	FGF	0,867
MARCO	0,637	DEFB128	0,749	S100A12	0,873
PGLYRP1	0,640	POLR3E	0,750	POLR3F	0,876
IKKB	0,640	CLU	0,754	POLR3B	0,876
CLEC6A	0,642	TLR7	0,771	TREM2	0,879
DEFB127	0,645	NLRP6	0,775	CHGA	0,884
IFNA8	0,646	ABL1	0,777	IGLL5	0,886
HAVCR2	0,648	CHUK	0,781	IFNW1	0,890
JAK1	0,648	NCF2	0,782	CIQA	0,893
CFH	0,656	IFNA1	0,787	LY96	0,896
IRF3	0,659	SAMHD1	0,788	HMGB3	0,897
TRIM32	0,660	CLEC4A	0,790	HCK	0,897
POLR3G	0,661	CD14	0,791	TLR5	0,899
C6	0,665	CD244	0,794	MALT1	0,903
DDX3X	0,665	IL34	0,794	IFNA7	0,909
CAMP	0,670	TLR2	0,797	TLR1	0,917
DEFB1	0,671	DEFB124	0,798	IFNE	0,917

Table S1. (continued)

Genes	p-value	Genes	p-value	Genes	p-value
NCF1	0,918	CD46	0,942	FCER1G	0,971
STO0B	0,919	PTK2	0,944	LBP	0,973
POLR3C	0,920	ARHGEF2	0,947	C8G	0,973
ADARB1	0,925	CD55	0,949	MSRB1	0,978
PTX3	0,934	UBA52	0,953	SRPK2	0,978
NLRP10	0,936	C3	0,954	POLR3K	0,981
IFNA16	0,939	CLEC4C	0,957	IFNA17	0,984
UBB	0,940	DEFB112	0,957	SERINC5	0,985
LYN	0,940	MAVS	0,964	PYDC1	0,991
SDHAF4	0,941	DEFA1B	0,967	DEFB121	0,991

Table S2. Read counts per gene. (XLSX). Available online.

