Identification of the Key Differential Transcriptional Responses of Human Whole Blood Following TLR2 or TLR4 Ligation *In-Vitro*

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Abstract

The use of human whole blood for transcriptomic analysis has potential advantages over the use of isolated immune cells for studying the transcriptional response to pathogens and their products. Whole blood stimulation can be carried out in a laboratory without the expertise or equipment to isolate immune cells from blood, with the added advantage of being able to undertake experiments using very small volumes of blood. Toll like receptors (TLRs) are a family of pattern recognition receptors which recognise highly conserved microbial products. Using the TLR2 ligand (Pam3CSK4) and the TLR4 ligand (LPS), human whole blood was stimulated for 0, 1, 3, 6, 12 or 24 hours at which times mRNA was isolated and a comparative microarray was undertaken. A common NFκB transcriptional programme was identified following both TLR2 and TLR4 ligation which peaked at between 3 to 6 hours including upregulation of many of the NFκB family members. In contrast an interferon transcriptional response was observed following TLR4 but not TLR2 ligation as early as 1 hour post stimulation and peaking at 6 hours. These results recapitulate the findings observed in previously published studies using isolated murine and human myeloid cells indicating that *in vitro* stimulated human whole blood can be used to interrogate the early transcriptional kinetic response of innate cells to TLR ligands. Our study demonstrates that a transcriptomic analysis of mRNA isolated from human whole blood can delineate both the temporal response and the key transcriptional differences following TLR2 and TLR4 ligation.

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Introduction

Microarray analysis is increasingly being used to advance our understanding of the complex transcriptional responses generated downstream of an experimental perturbation or during a disease state [1,2]. Using modern microarray platforms it is possible to measure the expression of over 40,000 mRNA transcripts, encompassing all of the known functional human genome. By sequentially sampling over time, the temporal dynamics of the transcriptional response to a given event can be delineated and depending on the scale of the response many hundreds or thousands of significantly differentially expressed genes may be identified. To better understand and interpret this complex data bioinformatics tools have been developed that take advantage of known biological relationships which may influence gene expression. Using these tools it is possible to use an unbiased methodology to determine distinct classes of differentially regulated genes as well as potentially important transcriptional regulatory pathways or networks which may change when exposed to a given stimulus [2–4].

Toll-like receptors (TLR) are a family of pattern recognition receptors (PRR) which recognise highly conserved microbial products. In humans 10 functional TLRs have been described including TLR2 and TLR4 which are expressed on the cell surface and recognise conserved bacterial products [5]. TLR2 recognises lipoteichoic acids of Gram-positive bacteria and bacterial lipoproteins whereas TLR4 recognises lipopolysaccharides (LPS) of Gram-negative bacteria [5,6]. TLR2, TLR3 and TLR4 are expressed and functional in a wide variety of cells found in human whole blood including dendritic cells (DC) and monocytes [6]. Early in the immune response to a pathogen, ligation of TLRs induces gene transcription leading to inflammation, tissue repair and the initiation of adaptive immunity [6,7]. All TLRs utilise the MyD88 adaptor molecule except for TLR3 which uses the TRIF-TRAM adaptor molecules only. TLR2 and TLR4 both use the MyD88-TIRAP adaptor molecules. Additionally TLR4 also uses the TRIF-TRAM adaptor molecules [5–7]. Signalling via the
adaptor molecule MyD88 results in activation of transcription factors such as NFκB and AP-1 [5–7], whereas signalling via the TRIF-TRAM adaptors leads to activation of the interferon regulatory transcription factors (IRF) including IRF3 [6,9]. IRF3 is constitutively expressed and its activation by ligation of TLR4 or TLR3 results in an induction of IFNβ (amongst other cytokines), which via the IFNα/β receptor leads to positive feedback regulation of type 1 interferon (IFN) genes including the type 1 IFN inducible transcription factor IRF7 [8]. There is also a differential temporal transcriptional response following NFκB activation in macrophages, which can be broadly categorised into three phases: early primary response, late primary response and secondary response genes. This differential activation of sets of genes depends on their chromatin status and the potential need for remodelling of the target genes which can lead to the differential kinetics of induction [7,9]. Much of the knowledge we have gained of these signalling mechanisms are derived from studying TLR ligation of monocytes, macrophages or DCs from mice or humans.

A study collectively analysing data from multiple published papers from both human and murine macrophage transcriptional studies identified a group of genes upregulated following both TLR2 and TLR4 stimulation, which were predicted to be regulated by NFκB. Additionally a separate IFN-sensitive response element (ISRE) regulated set of genes expression was seen to be upregulated in the TLR4 (and TLR3) stimulations but not the TLR2 stimulation [10]. Importantly the temporal kinetics of these two groups of genes differed, with the genes predicted to be regulated by NFκB peaking in expression earlier than the predicted ISRE regulated genes. In addition, genes thought to be regulated by both ISRE and NFκB had a greater magnitude of induction though similar temporal kinetics to the genes regulated only by ISRE [10]. However this study had limited time points available for the TLR2 stimulations and so the majority of the temporal analysis was undertaken using LPS stimulated samples. Another comparative study including TLR2, TLR3 and TLR4 stimulation of murine primary DCs identified an “inflammatory programme” mediated by TLR2 ligation and an “anti-viral programme” mediated by TLR3 ligation. TLR4, owing to its use of both MyD88-TIRAP and TRIF-TRAM adaptor molecules, resulted in the activation of both “inflammatory” and “anti-viral” programmes [11]. The “inflammatory” programme was enriched for genes predicted to be regulated by NFκB (RELA, NFκB1, NFκB2 and MYD88) and the “anti-viral” programme was enriched for genes predicted to be regulated by the transcription factors STAT1, STAT2, STAT4, IRF3 and IRF9 [11]. Study of the temporal transcriptomic response following TLR4 ligation in human and murine macrophages has also been used to specifically analyse the expression of transcription factors [12–14]. For example in LPS stimulated human macrophages the majority of transcription factors whose expression was seen to change had done so by 2 hours. Groups of transcription factors peaked in expression at different times, coinciding with the transcriptional peaks of the genes they were predicted to regulate following TLR stimulation in both murine and human macrophages [12,15]. However studies in isolated cells may not reflect the overall host response to TLR ligation, since interaction will occur between different cell types, leading to a complex interplay of autocrine and paracrine signalling events resulting in differentiation, proliferation, cell trafficking and further chemokine/ cytokine production and feedback loops for positive and negative effects on gene regulation.

Transcriptomic studies following in vivo TLR4 ligand administration have also previously been undertaken [16–18]. From human whole blood leukocytes obtained 0, 2, 4, 6, 9 and 24 hours following in vivo administration of LPS, it was shown that mRNA expression of proinflammatory chemokines and cytokines (TNF, IL1A, IL1B, CXCL1, CXCL2, CCL2, CXCL8 and CXCL10) peaked at 2 to 4 hours after LPS administration, whereas the cytokine IL10 was maximal at 6 hours. In this study the transcription factors NFκB1, NFκB2, RELA and RELB were significantly expressed and seen to peak after the cytokines and chemokines. The peak time for transcription factors including the STAT (signal transducer and activator of transcription) and IRF genes was at 4 to 6 hours [16,19]. Analysis of mRNA isolated from circulating human neutrophils after 0, 2, 4 and 6 hours following LPS administration revealed significant upregulation of the TNF signalling pathway and NFκB genes such as NFκB1 and NFκB2 by 2 hours [18]. Although it is possible to undertake certain human in vivo experiments, studies are limited by ethical and practical considerations.

Whole blood comprises cells of both the innate and adaptive immune system. Therefore the use of whole blood for in vivo studies has the potential advantage over isolated cell populations as these different components may have a differential response to stimulation. Autocrine and paracrine signalling between the differing cell populations may result in a response of the whole system that potentially better reflects the in vivo response. Additionally whole blood potentially has advantages over PBMC, DCs or monocyte derived macrophages since it can be used in situations where it is not possible to obtain large volumes of blood to derive the isolated cell populations. Previously, In vitro human whole blood has been used as a model to study TLR ligation, predominantly with measurement of specific cytokine protein levels [20–23] or specific cytokine mRNA levels as the readout [24,25]. Human whole blood has also been used as a model to assess the whole genome transcriptional response to TLR4 [26] or TLR2 and TLR4 ligands [27], although these studies only looked at one time point with limited analysis.

The objective of our study was to undertake a detailed comparative analysis of the in vitro global temporal transcriptional response to TLR4 and TLR2 ligation in human whole blood. To better understand this gene transcriptional response we used a variety of bioinformatics approaches to delineate both the temporal response and the key transcriptional differences resulting from TLR2 and TLR4 ligation and demonstrate that in a whole blood system that the response to TLR stimulation can resemble that previously identified in isolated immune cells.

Materials and Methods

Ethics Statement

This study was approved by the Central London 3 Research Ethics Committee (09/H0716/41). All participants gave written informed consent.

Human Volunteers

Six healthy volunteers (self-reported questionnaire); three male, three female; aged 25–50 years old; of similar ethnic background were recruited into the study. Sixty ml of whole blood from each volunteer was taken at 9 am into 10 ml Vacutainers with sodium heparin 17 international units/ml (BD Vacutainer).

Whole Blood Cellular Composition

Measured by Celltac Automated Hematology Analyzer (MEK-6400)/K, Nikon Kohden) at 0 hour, volunteer’s results listed Table S1.
In vitro Whole Blood Stimulation
In 24 well plates (Costar 3524, Corning Incorporated), 1 ml of heparinised whole blood was stimulated either in the presence of a final concentration of 200 ng/ml of Pam3CSK4 (Invivogen), 1 ng/ml of LPS (from Salmonella Minnesota R595, Enzo Life Sciences) added in a volume of 100 µl with RPMI-1640 with GlutaMAX (Life Technologies). Media control samples were cultured with the addition of 100 µl of RPMI-1640 with GlutaMAX. Samples were incubated at 37°C, 5% CO₂ for 0, 1, 3, 6, 12 or 24 hours at which point the contents of the well were thoroughly mixed with 2 ml Tempus Solution (Applied Biosystems/Ambion) to lyse the cells and stabilise the RNA. Samples were stored at −80°C until RNA processing.

Endotoxin Testing
Reagents (excepting LPS and Pam3CSK4) including Sodium heparin Vacutainers were tested for endotoxin contamination by Limulus assay and were found to be endotoxin free (<0.03 EU/ml). Pam3CSK4 was tested by the manufacturer and confirmed to be endotoxin free (<0.001 EU/µg).

RNA Processing
RNA was isolated using the PerfectPure RNA Blood kit (5-PRIME) according to manufacturer's instructions. 2.5 µg of isolated RNA was globin RNA reduced using the GLOBINclear 96-well format kit (Applied Biosystems/Ambion) according to the manufacturer's instructions. Isolated and globin reduced RNA quantity was assessed using either NanoDrop 1000 or NanoDrop 8000 spectrophotometer (Nanodrop Products, Thermo Fisher Scientific), RNA quality was assessed using an Agilent 2100 Bioanalyzer (Range 6.5–9.5) (Agilent technologies). 200 ng of globin reduced RNA was amplified to generate biotinylated amplified antisense cRNA using the Illumina CustomPrep RNA amplification kit (Applied Biosystems/Ambion). 750 ng of cRNA was hybridized to Illumina Human HT-12 V4 BeadChip arrays (Illumina) and scanned on Illumina iScan. GenomeStudio (Illumina) was used to perform quality control and generate signal intensity. Two samples were excluded from further analysis at this stage as they failed quality control measures (0 hour media control x1, 6 hour media control x1).

Microarray Analysis
Raw background subtracted data was processed using GeneSpring V12.6 (Agilent Technologies) and the following principles were applied to all analyses. After background subtraction low signal values (<10) were then set to a threshold of 10, log2 transformed and per chip normalised using 75th percentile shift algorithm. Per-transcript normalisation was undertaken by normalisation to the median of a defined control group. Transcripts were then filtered out if they were not significantly (p<0.01) different in intensity value compared to the background in at least 10% of all the samples.

The resulting transcripts were then subjected to statistical filtering (either One-way ANOVA or 2-way ANOVA) with multiple testing correction (Benjamini-Hochberg p<0.01), followed by a further filtering of transcripts by fold change (FC) in which transcripts were filtered if less than 1.8 FC different between variables of interest. Expression heatmaps were generated within GeneSpring V12.6. Heatmap clustering was undertaken using Differential distance metric and Wards linkage rule, unless otherwise stated.

Media controls from 2 volunteers had evidence of activation of inflammatory genes by 3 hours of culture (Figure S1A). This activation persisted in all of the subsequent time points for these individuals (not shown) appears to be revealed in culture, is independent of the individual, and independent of the length of time in transport conditions (Figure S1B). These samples were excluded from the study.

The data discussed in this publication have been deposited in NCBI's Gene Expression Omnibus and are accessible through GEO Series accession number GSE55375.

k-means Clustering
Within Genespring 12.6 the normalised significant transcript lists were separately clustered by k-means clustering into 9 clusters using Euclidian distance metric. Number of clusters was chosen by the number of 3rd order branches in the dendrogram from the LPS expression heatmap. Clusters were compared across stimulations using Pearsons Correlation (within Graphpad Prism V6).

Ingenuity Pathway Analysis (IPA)
The canonical pathway, gene function annotation and upstream analyses were generated through the use of IPA (Ingenuity Systems, www.ingenuity.com). Significant transcripts identified from GEO microarray analysis were uploaded into IPA. For time point analyses these lists were filtered by mean FC (>1.8) compared to media control at the time point.

Type I IFN Regulated List
A list of human type I regulated genes was obtained from the Interferome V2 database (Accessed June 2013) [28].

Quantitative PCR
From the globin reduced RNA cDNA was synthesised using High Capacity cDNA Reverse Transcription kit (Applied Biosystems), according to the manufacturer's instructions followed by RNase H (Promega) treatment for 30 min at 37°C. IFNB1, IL1A, IL6, NFKB1, NFKB2, STAT1, STAT2 and IRF7 gene expression were quantified by real-time PCR (7900HT, Applied Biosystems) using the TaqMan system, and normalised to GAPDH mRNA. Primer probes used were IFNB1 (Hs01077958_s1); IL1A (Hs00174092_m1); IL6 (Hs00985639_m1); NFKB1 (Hs00765730_m1); NFKB2 (Hs01020890_s1); STAT1 (Hs01013996_m1); STAT2 (Hs01013123_m1); IRF7 (Hs01014809_g1); GAPDH (Hs02758991_g1) (all Applied Biosystems).

Results
Media Controls
Analysis of the media controls alone over time revealed 377 significantly expressed transcripts over time. The peak of this difference was at 24 hours with 281 transcripts more than 1.8FC different compared to the 0 hour samples (Figure S2A). These differentially expressed transcripts at 24 hours were enriched for inflammatory and metabolic function pathways (Figure S2B). For this reason all data for stimulations are filtered against the media controls from the time point of interest and fold changes calculated as relative to the media controls at the same time point.

Identification of Significantly Differentially Regulated Transcripts following LPS and Pam3CSK4 Stimulation
To gain insight into the differential temporal gene expression in response to TLR4 and TLR2 ligation we performed a comparative microarray analysis of LPS and Pam3CSK4 stimulated human whole blood and accompanying media controls over a
time course. From four healthy human volunteers 1 ml of heparinized whole blood was stimulated with either the TLR4 ligand LPS (1 ng/ml), the TLR2 ligand Pam3CSK4 (200 ng/ml), (concentrations shown to result in a plateau in previous studies within our laboratory), or incubated only with media as a control. RNA was isolated at 0, 1, 3, 6, 12 and 24 hours for each stimulus or control.

LPS (TLR4) stimulation induced a greater number of differentially regulated transcripts as compared to Pam3CSK4 (TLR2) and had a higher magnitude of response. For this reason the two stimulations were first analysed independently to generate the significant transcript lists compared to media control over the time course after stimulation. LPS stimulation resulted in the differential expression of 4777 transcripts (mapping to 3571 unique genes in IPA) whereas Pam3CSK4 stimulation resulted in only 1202 differentially regulated transcripts (mapping to 922 unique genes in IPA) (Figure 1A). Expression of these transcripts varied over time. Overall 90% of the 1202 significant Pam3CSK4 transcripts were shared with the LPS stimulation (Figure 1B). These shared 1093 transcripts when analysed by canonical pathway analysis (within IPA) were shown to contain TREM, TNF and NFκB signalling amongst the top 5 pathways (ranked by significance). Canonical Pathway analysis of the 3684 transcripts significantly expressed following LPS and not Pam3CSK4 stimulation revealed “IFN signalling” as the most significant pathway.

k-means Clustering Analysis Reveals Similarities and Differences between Stimulations

k-means clustering was applied to cluster the significant transcript lists based on their similarity in expression over time for each stimulation separately (full composition of the clusters listed in EXCEL Files S1 & S2). For each cluster the most significant canonical pathway (IPA) was determined, reflecting gene enrichment within each cluster. The clusters were then compared by their expression profile over time (Pearson correlation, Table S2) and the top canonical pathway to discover similar clusters between LPS (termed L) and Pam3CSK4 (termed P). Based on these criteria ten clusters in response to both stimulations were equivalent. Of the remaining clusters, six shared similar expression profiles but had different top canonical pathways and two clusters (P4 and P7) had no equivalent in the LPS stimulated clusters (Figure 2).

The similar clusters with “Granulocyte Adhesion and Diapedesis” as the top canonical pathway (L7 and P2) were small in terms of numbers of genes which were highly expressed by 1 hour after stimulation with both LPS and Pam3CSK4 and peaked in expression between 3 to 6 hours. 80% of the genes in the P2 cluster were found within the L7 cluster. These common genes were predominantly chemokines and cytokines: C15orf48, CCL2, CCL20, CCL3, CCL3L1, CCL3L3, CCL4L1, CCL4L2, CCL8, CCR1L2, CXCL2, EBI3, IL1A, IL6 and TNF.

The “Acute Phase response signalling” clusters (L4 and P8) were characterised by genes involved with inflammatory response. Approximately 60% of the genes in the P8 cluster were also found within the L4 cluster and these common genes included those involved with the inflammatory response: ORM1, ORM2, HAMP, IRAK2, PI3, PTGES, TNFAIP6, TNIP3. In addition, within the L4 cluster but not the P8 cluster were genes involved with IFN regulation: IFIH1, IFT1, IFT3, IFNG, OAS3, OASL.

The “Protein Ubiquitination pathway” clusters (L1 and P1) had 104 common genes between stimulations which included heat shock protein genes (HSP90AA1, HSP90AB1, HSPA5, HSPD1, HSPB1 and HSPH1) and Proteasome PA700/20S genes (PSMA1, PSMB3, PSMC4, PSMD1 and PSMD14).

The “Complement system” clusters (in both L3 and P5) were characterised by genes that were upregulated, had their highest expression at 24 hours and included the metallothionein genes (MT1G, MT1H, MT1E, MT1X, MT1M and MT1F) which were amongst the highest expressed genes at 24 hours following both LPS and Pam3CSK4 stimulations (Figure S3). Clusters L5 and P6 contained transcripts that were down regulated over time and were similarly enriched for the pathway “Pathway Biology Synthesis (Late stages)”.

Cluster L2, which peaked at 6 hours and was characterised by “Activation of IRF by cytosolic pattern recognition receptors” as the top pathway did not have a similar pathway in the Pam3CSK4 clusters. Examination of the genes within this pathway revealed that there were detected a number of IFN regulated/regulatory genes (DDX58, DDX58L2, IFIH1, IFT12, IFNB1, IRF7, STAT2). Clusters L3, L0, L6 and P4, P7 and P0 were all enriched for immune function pathways although there were no similarities between the stimulations in terms of top significant pathway and kinetic profile. Cluster P3 was enriched for the pathway “UDP-N-acetyl-D-galactosamine Biosynthesis II” and there was no similar cluster following LPS stimulation.

Common and Distinct Changes in Transcription at the Different Time Points

To better understand the temporal response we undertook analysis at each time point using the significant transcript lists. Significantly expressed transcripts at each time point were identified by comparing the mean expression of the differentially regulated transcripts in response to LPS and Pam3CSK4 stimulation at each time point to the media control at that time point and filtering those transcripts which were less than 1.8 FC different to the media control (full listings of identified transcripts given as EXCEL Files S3 & S4). From this it was observed that the peak of the transcriptional response compared to media control occurred at 6 hours for both LPS and Pam3CSK4 stimulations (Figure 3A). This differential transcriptional response was still evident at 24 hours following in vitro LPS stimulation of whole blood.

At each time point the per-time point transcript lists were analysed by canonical pathway analysis within IPA. The top 25 significant pathways (by mean –log p value of the LPS time point pathways and compared to Pam3CSK4) are shown (Figure 3B). This analysis revealed a large number of canonical pathways with similar kinetics of significance in both LPS and Pam3CSK4 stimulations, with the exception of the “IFN Signalling” pathway where there was a clear difference in significance between stimulations.

The pathways that were most significant at 1 hour and then diminished in significance over time were “Agranulocyte Adhesion and Diapedesis”, “Granulocyte Adhesion and Diapedesis”, “Differential Regulation of cytokines production in Epithelial Cells by IL-17A and IL-17F”, “Differential regulation of cytokines production in macrophages and T-Helper Cells by IL-17A” and the pathway “Role of Hypercytokinemia/hyperchemokinemia”. All the genes contributing to these pathways at this time point were chemokine and cytokine genes. “IL-10 signalling”, “Communication between Innate and Adaptive Immune cells” and “Atherosclerosis Signalling” pathways were also most significant at 1 hour and then diminished in significance over time. In addition to cytokine and chemokine genes the “Atherosclerosis Signalling” and “Communication between Innate and Adaptive Immune cells” pathways had CD40, ICAM1, ORM1 and ORM2 as being significantly expressed within the pathways at 1 hour following LPS and Pam3CSK4 stimulation. The “IL-10 signalling” pathway
in addition to chemokines and cytokines had the transcription factors, JUN, NFKB1, NFKB2, NFKBIA, NFKBIE as well as the gene SOCS3 within the pathway.

The pathways which were most significant at 3 hours following both LPS and Pam3CSK4 stimulation were “NFkB Signalling” and “IL-6 Signalling”. There were a number of pathways that were significant by 1 hour and remained significant following both LPS and Pam3CSK4 stimulations: “TNFR2 signalling”, “TREM1 signalling”, Altered T cell and B cell signalling in Rheumatoid Arthritis and “Dendritic cell maturation” pathways.

The “IFN signalling” pathway in response to LPS was only significant (p<0.01) from 3 hours onwards and was most significant at 6 hours (Figure 3B). There was a clear difference between the level of significance of the IFN signalling pathway following LPS and Pam3CSK4 stimulation, with the “IFN signalling” pathway not reaching the threshold of significance (p<0.01) at any time point following Pam3CSK4 stimulation.

The significantly expressed genes at 1 hour were analysed, as from the canonical pathway analysis we had identified enrichment of cytokines and chemokines genes at this time point. In addition this was the earliest measured time point following stimulation, when autocrine and paracrine signalling leading to induction of mRNA should be at its minimum and, 334 LPS and 165 Pam3CSK4 genes respectively were more than 1.8 FC different to media control at 1 hour. Cytokines/chemokines and transcriptional regulators (identified by IPA gene function annotation) combined accounted for approximately 20% of the significantly expressed genes at this time point. There was a large degree of

Figure 1. LPS or Pam3CSK4 stimulations results in a differential response in gene expression over time. (A) 1 ml of human whole blood from healthy volunteers (N = 4) was stimulated with either Pam3CSK4 (200 ng/ml), LPS (1 ng/ml) or media control for different lengths of time (0, 1, 3, 6, 12 and 24 hours). Stimulations were analysed independently: media control compared to Pam3CSK4 and media control compared to LPS revealed 1202 and 4777 significantly expressed transcripts respectively. Transcripts were identified by normalising expression values to the median of the 0 hour samples, filtering by detection from background, statistical filtering (2 way ANOVA with Benjamini Hochberg multiple testing correction p<0.01) and retaining transcripts whose expression was greater than 1.8 FC different between the media control and stimulation samples at one or more time point. (B) A Venn diagram of both significant transcript lists. Within the Venn for each subset the number of transcripts is given, with unique genes within IPA in brackets. For transcript lists the top 5 canonical pathways (IPA) are shown as well as a heat map of the normalised expression values of these transcripts for both stimulations over time.

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overlap between stimulations of the significant cytokines/chemokines at 1 hour, with all of the 17 significant cytokines/chemokines genes from the Pam3CSK4 list also being identified in the LPS list (Figure 3C). Following LPS but not Pam3CSK4 stimulation, of note IL12B, IFNB1, IFNG, IL23A and CXCL10 were also significantly expressed as early as 1 hour (Figure 3C). There were a number of significantly expressed transcriptional regulators by 1 hour, with the majority of significantly expressed transcriptional regulators identified in the Pam3CSK4 1 hour list shared with LPS, including those of the NFκB family (NFKB1, NFKB2, NFKBIA, NFKBIE, NFKBIZ and REL) and the AP-1/CREB family (JUN, JUNB, ATF3, BATF3). Transcriptional regulators involved in cell development, proliferation and differentiation (BTG2, EGR3, ETV3, HHEX, MAFF, SKIL) and the post-transcriptional regulator ZFP36 (also known as Tristetraprolin) were also identified (Figure 3C).

Upstream Analysis Identifies Potential Transcriptional Regulators

In order to identify which transcriptional regulators may be responsible for the differences observed in gene expression between LPS and Pam3CSK4 stimulation we undertook upstream analysis of the gene lists at each time point within IPA. Upstream analysis attempts to predict which transcriptional regulator may be responsible for the observed differential gene expression by comparing the genes known to be regulated by a transcriptional regulator (derived from the literature) to those significantly differentially expressed genes identified at each time point from this analysis. It can be seen that the top predicted transcription regulator for both stimulations was the NFκB complex and that it was predicted to be activated from 1 hour onwards (Figure 4A).

By comparison of the mean mRNA expression of the predicted transcriptional regulators compared to media control it can be seen that the magnitude and temporal response was similar for LPS and Pam3CSK4 stimulation for PPRC1 and SP1 (Figure 4B). The predicted transcriptional regulators NFκB1, CEBPA, EGR1 shared similar mRNA temporal profiles between stimulations; however the magnitude of the response was greater upon LPS stimulation. RELA, EZH2 and STAT3 were predicted to be activated in both stimulations however had different temporal profiles between stimulations. RELA, EZH2 and STAT3 were upregulated earlier and to a greater magnitude in LPS compared to Pam3CSK4 and this upregulation persisted over time in the LPS stimulation compared to Pam3CSK4 stimulation. The mRNA expression of STAT1 and IRF1, which were only predicted to be activated following LPS stimulation, had different profiles being induced earlier and to a greater magnitude in LPS compared to Pam3CSK4 stimulations.

The NFκB genes REL, RELA, RELB, NFKB1 and NFKB2 were all significantly expressed and present in both LPS 4777 transcript and Pam3CSK4 1202 transcript lists (Figure 4C). Transcriptional regulators involved in IFN regulation IRF1, IRF3 and STAT1 were predicted to be activated in the LPS stimulation but not the Pam3CSK4 stimulation (Figure 4A). IRF1, IRF2, IRF4, IRF7, IRF8, IRF9, STAT1, STAT2, STAT3, STAT4 and STAT5A were all identified in the 4777 LPS and not the Pam3CSK4 1202 transcript list as being significantly differentially expressed, IRF7 having the greatest FC induction compared to media control. IRF3, IRF5 and IRF6 expression was not significantly regulated following LPS or Pam3CSK4 stimulation compared to media control (Figure 4D). Overall the inducible IRF and STAT gene expression level peaked later (6 hours) in expression at 6 hours compared to that of the NFκB family of genes which peaked between 3 to 6 hours (Figure 4E). We validated this difference between LPS and Pam3CSK4 in terms of the magnitude of expression at the 6 hour time point for NFκB1, NFκB2, STAT1, STAT2 and IRF7 by quantitative real time PCR (Figure S5). In addition the FC relative to the media control was lower for STAT1, STAT2 and IRF7 compared to the NFκB1 and NFκB2 following Pam3CSK4 stimulation.

**Upstream Analysis Identifies Potentially Active Cytokines**

In order to attempt to identify the cytokines that were potentially involved in autocrine/paracrine signalling and subsequent gene expression regulation in response to LPS or Pam3CSK4 stimulation, we again undertook per-time point upstream analysis within IPA. TNF, IL1B and IL1A were predicted as potentially activated cytokines, and this predicted activation was early (by 1 hour) and sustained (Figure 5A). The mRNA expression of these cytokines compared to media control revealed them to be highly upregulated by 1 hour in both LPS and Pam3CSK4 stimulations. Although the magnitude was higher in LPS stimulation, they shared similar kinetic profile between stimulations (Figure 5B). IL1B was predicted to be activated upon LPS stimulation but not Pam3CSK4 and this difference was observed in the mean mRNA expression of the IL12B gene (encoding for IL12p40) which was upregulated in response to LPS compared to media control and not following Pam3CSK4 stimulation (Figure 5). IFN cytokine activation was also only predicted in the LPS stimulation (IFNA2, IFNL1 and IFNB1), in keeping with this the kinetic profile of IFNB1 can be seen to be different between LPS and Pam3CSK4 stimulations. This difference between LPS and Pam3CSK4 stimulation was validated by real time PCR at the peak of IFNB1 expression at 3 hours. IL1A and IL6 which also peaked at 3 hours were seen to have a robust response following both LPS and Pam3CSK4 stimulations, although with a much greater response following LPS (Figure S5). IFNA and IFNL1 genes were of low magnitude expression (<1.8 FC different to media control) under both conditions of stimulations (not shown).

**IFN Gene Expression is Dominant Following LPS but not Pam3CSK4 Stimulation**

Our data show that as early as 1 hour post stimulation IFN gene expression is seen to be upregulated following LPS but not Pam3CSK4 stimulation. This difference in IFN signalling was further emphasised by the k-means clustering, canonical pathway analysis and upstream analysis of both potential transcriptional regulators and cytokines highlighting a difference in IFN signalling following LPS and Pam3CSK4 stimulation. IRF and STAT genes had been identified by k-means clustering, predicted upstream analysis, and canonical pathway analysis as being differentially activated between LPS and Pam3CSK4 stimulations.

To test if this difference in expression of IRF and STAT genes resulted in differential expression of IFN regulated genes we used a list of human type 1 IFN regulated genes generated from the Interferome database v2.0. We compared the expression of these...
Figure 3. Transcript lists analysed at each time point. (A) A graph showing the number of genes from the respective significant transcript lists (4777 LPS and 1202 Pam3CSK4 lists) at each time point which are more than 1.8 FC different compared to the media control at that time point. (B) Transcriptional Response to TLR2 or TLR4 Ligation

**Cytokines and chemokines**

- LPS
  - CCL14, CXCL10, IFNB1, IFNG, IL12B, IL23A, IL36G, OSM
  - CCL2, CCL20, CCL23, CCL3, CCL3L1/CCL3L3, CCL4, CXCL1, CXCL2, EBI3, EDN1, IL1A, IL1B, IL1RN, IL6, IL8, TNF

**Transcriptional regulators**

- LPS
  - ATF3, BATF3, BTG2, CSRP1, EGR3, ELL2, ET3, HEX, HIVEP1, JUN, JUNB, MAF, MAF1, NF2L3, NFKB1, NFKB2, NFKBIA, NFKBIE, NFKBIZ, REL, SKIL, ZFP36

**Interferon Signalling**

**TNFR2 Signalling**

**TREM1 Signalling**

**Communication between Innate and Adaptive Immune Cells**

**Dendritic Cell Maturation**

**Role of Pattern Recognition Receptors in Recognition of Bacteria and Viruses**

**Death Receptor Signalling**

**Type 1 Diabetes Mellitus Signalling**

**Activation of IRF by Cytosolic Pattern Recognition Receptors**

**TNFRI Signalling**

**TWEAK Signalling**

**Agranulocyte Adhesion and Diapedesis**

**Granulocyte Adhesion and Diapedesis**

**Differential Regulation of Cytokine Production in Intestinal Epithelial Cells by IL-17A and IL-17F**

**Differential Regulation of Cytokine Production in Macrophages and T Helper Cells by IL-17A and IL-17F**

**Atherosclerosis Signalling**

**Role of Hypercytokinemia/hyperchemokinenia in the Pathogenesis of Influenza**

**Role of IL-17F in Allergic Inflammatory Airways Diseases**

**IL-6 Signalling**

**NF-κB Signalling**

**Role of Macrophages, Fibroblasts and Endothelial Cells in Rheumatoid Arthritis**

**IL-17A Signalling in Fibroblasts**

**Acute Phase Response Signalling**

**IL-10 Signalling**
The significantly expressed transcript lists (1202 Pam3CSK4 transcript list and 4777 LPS transcript list) were analysed in IPA. For each time point only genes whose expression were 1.8 FC different from the media control at that time point were taken into consideration. Shown is a heatmap of pathway significance of the top 25 IPA canonical pathways for each time point where significance criteria met (Fisher’s Exact test p < 0.01). The IPA canonical pathways were chosen by identifying from the LPS stimulation analyses the top 25 most significant pathways across the time points (mean –log p value) and then compared to Pam3CSK4. (C) Venn diagrams of cytokine/chemokines and transcriptional regulators identified using IPA gene functional classification from LPS 4777 and Pam3CSK4 1202 transcript lists with mean expression greater than 1.8 FC different to media control at 1 hour. Listed adjacent to the Venn diagrams are the genes from each subset.

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Discussion

A central goal of our study was to identify the key transcriptional differences between TLR4 and TLR2 ligation in a human whole blood system and demonstrate that in a whole blood system that the response to TLR stimulation can resemble that previously identified in isolated immune cells. We show that the magnitude of the transcriptional response both in terms of the number of differentially regulated genes as well as level of mRNA expression following LPS stimulation was greater and more sustained than the transcriptional response to Pam3CSK4. There was a common transcriptional response following stimulation with LPS and Pam3CSK4 which was dominated by NFkB regulated genes. In addition there was a separate IFN regulated transcriptional response (IFN and STAT mediated) seen mainly following LPS stimulation. This significant difference in the IFN response between TLR4 and TLR2 ligation could be seen as early as 1 hour post stimulation in our study, and reinforced in the later time points despite the potentially complex autocrine and paracrine signalling in the human whole blood system.

The common early transcriptional response following both LPS and Pam3CSK4 stimulations at 1 hour was characterised by highly expressed cytokines and chemokines (CCL2, CCL20, CCL23, CCL3, CCL3L1/CCL3L3, CCL4, CXCL1, CXCL2, EB13, EDN1, IL1A, IL1B, IL1RN, IL6, IL8 and TNF) as well as highly expressed cytokines and chemokines (CCL2, CCL20, CCL23, CCL3, CCL3L1/CCL3L3, CCL4, CXCL1, CXCL2, EB13, EDN1, IL1A, IL1B, IL1RN, IL6, IL8 and TNF) as well as the significant upregulation of transcriptional regulators including the NFkB family (NFKB1, NFKB2, NFKBIA, NFKBIE, NFKBIZ and REL) and AP-1/CREB (JUN, JUNB, ATF3, BATF3). The gene ZFP36 which encodes for the protein Tistreaprolin was also identified as significantly upregulated by 1 hour which has been shown to act in a post-transcriptional regulatory role by binding to the mRNA of some cytokines and promoting their degradation [29]. The peak of expression compared to media control of NFKB genes was 3 to 6 hours following both LPS and Pam3CSK4 stimulation. By 24 hours following Pam3CSK4 stimulation the expression of these NFKB genes was not significantly different.
Figure 4. Predicted transcriptional regulator identification and NFκB, IRF and STAT gene expression following LPS and Pam3CSK4 stimulation. (A) Predicted upstream transcriptional regulators from IPA; stimulations analysed independently using Pam3CSK4 1202 and LPS 4777 transcripts lists, at each time point only genes whose expression were 1.8 FC different from the media control at that time point were taken into
compared to media control, and similarly the expression of many proinflammatory cytokines (e.g. IL1A, IFNG and TNF) had diminished significantly by 24 hours compared to media control. This is in contrast to the expression following LPS stimulation where these NF\textsubscript{kB} genes and cytokines were still significantly upregulated compared to media control at 24 hours. This prolongation of the NF\textsubscript{kB} signalling seen following LPS stimulation maybe as a result of secondary induction of NF\textsubscript{kB} via the TRIF-TRAM adaptor molecules, or as a consequence of the IFN signalling [7,8]. We also identified a common group of metallo-

![Graph](image-url)

**Figure 5. Identifying potential cytokines involved in autocrine gene regulation by upstream analysis within IPA.** (A) Predicted activated cytokines from IPA upstream analysis; stimulations analysed independently using Pam3CSK4 1202 and LPS 4777 transcripts lists, at each time point only genes whose expression were 1.8 fold different from the media control at that time point were taken into consideration. Predicted upstream cytokines which met the criteria (p value <0.01 (Fishers Exact Test) and z-activation score >2.5) shown plotted by z-activation score only at the time points where significance criteria met. (B) Cytokine identified from either predicted upstream analysis or canonical pathway analysis (Fig. 3) mean mRNA expression plotted as log2 fold change (y axis) across time (x axis; 0, 1, 3, 6, 12 and 24 hours), fold change is relative to the media control at each time point, only those predicted cytokines whose mRNA expression is >1.8 fold upregulated relative to media control at one or more one time points are shown.

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Metallothionein genes (MT1G, MT1H, MT1E, MT1X, MT1M, MT1F) as being amongst the most up-regulated genes at 24 hours following both LPS and Pam3CSK4 stimulation. Metallothionein genes are known to have a key role in oxidative stress and heavy metal detoxification [30]. It has also been shown that they may be induced by cytokines such as IL-6 and they are thought to have a gene regulatory role in inflammation [31–33]. The late induction of these genes in our system suggests that they may be being induced by subsequent auto/paracrine signalling downstream of the MyD88 adaptor.

Following LPS stimulation a differential transcriptional response of IFN regulated/regulatory genes was observed as compared to Pam3CSK4. This difference was detectable as early as 1 hour post LPS stimulation with the significant upregulation of the cytokine genes IFNB1 and IFNG and the transcriptional regulator IRF8 following LPS and not Pam3CSK4 stimulation. The IRF and STAT (IRF1, IRF2, IRF4, IRF7, IRF9, IRF3, STAT1, STAT2, STAT3, STAT4 and STAT5A) genes were seen to be differentially regulated following LPS stimulation and the peak of their expression was at 6 hours. This difference in induction of IRF and STAT transcription regulators following TLR4 and not TLR2 ligation is consistent with previous studies which have compared the temporal transcriptional response following TLR2 and TLR4 ligation in murine DCs and macrophages [10,11]. The IFN signalling induced following TLR4 ligation and not TLR2 ligation can be accounted for by signalling through the MyD88 independent TRIF-TRAM, IRF3 pathway following TLR4 ligation resulting in induction of IFNB, which then leads to positive feedback of IFN gene induction through induction of IRF7 amongst other IRFs [8]. IRF3 is thought to be constitutively expressed and its expression is not thought to be induced by TLR ligation, or type I or II IFN. This may explain why IRF3 was predicted by upstream analysis to be activated and its mRNA expression not significantly induced following TLR4 or TLR2 ligation in our study [8]. IRF7, which is known to be induced by type I IFN subsequent to IRF3 activation by TLR4 ligation was seen to be the most induced of the IRFs following TLR4 ligation [8]. This stepwise induction of the IRFs may account for the later peak of expression at 6 hours of the inducible IRF and STAT genes compared to the NFkB genes which peaked at 3 hours in our study. This later peak in transcription factors associated with IFN regulation compared to NFkB genes is in agreement with studies in murine macrophages as well as ex-vivo human blood leukocytes. Slight differences in the timing of peaks observed in our study and these other studies could potentially be explained by the difference in time points sampled or differences in the systems used [10,34].
Transcriptional Response to TLR2 or TLR4 Ligation

A

In-vitro Blankley - LPS

In-vivo GSE3284 - LPS

4777 transcripts

3599 transcripts

In-vitro 3571 genes

In-vitro 2602 genes

2560

1011

1591

1. Crosstalk between DCs and NK cells
2. Communication between Innate and Adaptive cells
3. IL-17A Signaling in fibroblasts
4. PI3K signalling in B lymphocytes
5. TREM1 Signalling
6. Role of IL-17F in allergic inflammatory airway disease
7. B Cell receptor Signalling
8. TWEAK signalling
9. Ceramide Signalling
10. Differential regulation of cytokine production in Macrophages and T Helper cells by IL-17A and IL-17F

1. Interferon Signalling
2. Role of PRR in bacteria and viruses
3. TNFR1 Signalling
4. Type 1 DM Signalling
5. Activation of IRF by cytosolic PRR
6. TREM1 Signalling
7. Dendritic Cell maturation
8. Death receptor Signalling
9. NFκB Signalling
10. Tec Kinase Signalling

1. EIF2 Signalling
2. Regulation of eIF4 and p7056K Signalling
3. mTOR signalling
4. Mitochondrial Dysfunction
5. Natural Killer cell signalling
6. CD28 Signalling in T Helper cells
7. Huntington’s Disease Signalling
8. Molecular mechanisms of Cancer
9. RAN signalling
10. Role of NFAT in regulation of the Immune response

B

Log2 Fold change

Time (Hours)

In-vitro NFκB gene family

In-vitro IRF and STAT gene family

In-vivo NFκB gene family

In-vivo IRF and STAT gene family

C

IL1A

IL1B

TNF

IL15

IL32

In vitro LPS stimulation

In vivo LPS stimulation
Following LPS stimulation the peak of differential transcription compared to media control was at 6 hours in terms of the number of differentially regulated genes. At 24 hours 1443 genes were still significantly differentially regulated. This persistence of differential transcription at 24 hours is observed in human LPS stimulated in vitro transcriptional studies [12,13]. However this in contrast to that noted in human in vivo stimulation where transcriptional difference peaked between 4 to 6 hours and by 24 hours the transcriptional signature had returned to baseline [16]. Several cytokines/chemokines were not seen to be significantly induced in vivo compared to in vitro and these differences in expression and persistence of differential expression observed in vitro compared to in vivo could be explained by trafficking/removal of activated immune cells out of the circulation in vivo, which is not possible to be represented in an in vitro whole blood system.

The ability to utilise in vitro stimulated human whole blood for transcriptomic analysis of the early innate immune response has potential advantages over the use of isolated immune cells as whole blood stimulation can be carried out in a laboratory where the expertise or equipment to isolate immune cells from blood is lacking. In addition the volume of blood needed for stimulation is much less than that required to isolate immune cells, meaning experiments can be performed in populations where access to larger volumes of blood is not possible e.g. paediatrics or it could be possible to undertake more stimulations/time points with a given volume. The results however must be interpreted in the context of the complex autocrine/paracrine interactions which occur in a mixed cell culture and therefore stimulation of isolated immune cells will remain advantageous to interrogate in detail a specific transcriptional response.

We have shown that human whole blood can be used to study the early temporal transcriptional response following TLR2 and TLR4 ligation and that the results obtained are comparable to those from isolated murine and human immune cells.

Supporting Information

Figure S1 Activation of samples by culture conditions. (A) Heatmap of expression, clustered by transcripts, shows that 2 individuals (out of 6) media controls show activation (marked with arrows), the genes differentially expressed in these 2 media control samples are similar to LPS, but lower in magnitude. Transcripts identified by normalisation to 0 hour samples, filtering by detection from background, statistical filtering (2 way ANOVA with Benjamiini Hochberg multiple testing correction $p<0.01$) and retaining transcripts whose expression was greater than 1.8 FC from another condition. (B) Activation is not due to length of time in transport conditions, nor is it individual specific. The single individual shown here had not previously activated, and was included in final dataset. Blood was collected at time point 0 and left in sealed vacutainers, the vacutainers were opened at one hour intervals and 1 ml human whole blood was either immediately mixed with Tempus solution (labelled as Direct from vacutainer) or plated for 3 hours with either media control (RPMI-1640 with GlutaMAX) or LPS (1 ng/ml) and then mixed with Tempus solution. Reagents and containers (including vacutainers) are endotoxin free (undetectable by Limulus assay - sensitivity <0.03 EU/ml). Heatmap of expression (duplicate stimulations from the same individual shown, 2619 transcripts), clustered by transcripts shows that regardless of length of time in vacutainer activation occurred in all media control samples, and is not observed in the direct from vacutainer samples, implying that activation is dependent on culture conditions and is not a function of length of time ex vivo or length of time spent in the vacutainers. Transcripts identified by normalisation to median of “Direct from vacutainer” samples, filtering by detection from background, statistical filtering (ANOVA with Benjamiini Hochberg $p<0.01$) and then transcripts retained whose expression was >1.8 FC from another condition. (TIF)

Figure S2 Transcriptional changes in media controls over time. (A) Heatmap of normalised expression values of 377 transcripts which were identified to be significantly differentially expressed overtime (transcripts identified by normalisation to 0 hour samples, filtering by detection from background, statistical filtering (One way ANOVA with Benjamiini Hochberg $p<0.01$) and then transcripts retained whose expression was >1.8 FC from the 0 hour samples. Plotted above the heatmaps is the number of significantly expressed genes (mapped in IPA) that were differentially expressed at each time point by more than 1.8 FC compared to the 0 hour samples. (B) Top ten IPA canonical pathways of the significantly expressed genes at 24 hours, with the $-\log p$ value for the pathway and the significantly differentially expressed genes listed for each pathway. (TIF)

Figure S3 Metallothionein gene expression. (A) Heatmap of averaged Metallothionein mRNA expression over time following LPS or Pam3CSK4 stimulation, values normalised to the median of the 0 hour. Note asynchronous scale. (TIF)

Figure S4 Interferon regulated genes. Heatmap of averaged expression values of Type 1 Interferon regulated genes (List obtained from Interferome v2.0), normalised to the median of the 0 hour, genes retained if they were expressed greater than 1.8 FC from media control in at least one stimulation in one or more time points (resulting in 1105 genes). Graphed above heatmap is the mean absolute fold change of these Type 1 interferon regulated genes. (TIF)

Figure S5 Real time PCR. Real time PCR of selected genes following LPS and Pam3CSK4 stimulations and media controls, normalised to GAPDH expression. Mean fold change calculated between media controls and stimulations.
Table S1 Volunteer whole blood composition measured by Celltac Automated Hematology Analyzer (MEK-6400) at time point 0 hour.

Table S2 Pearson correlations for k-means derived clusters from Figure 2.

File S1 Listings of transcripts, LPS k-means clusters from Figure 2.

File S2 Listings of transcripts, Pam3CSK4 k-means clusters from Figure 2.

File S3 LPS time course data. Listings of transcripts from 4777 significant transcript list whose mean expression was 1.8 FC different to media control at each time point from Figure 3.

File S4 Pam3CSK4 time course data. Listings of transcripts from 1202 significant transcript list whose mean expression was 1.8 FC different to media control at each time point from Figure 3.

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Author Contributions

Conceived and designed the experiments: SB CMG CIB MPRB VP ML AOG. Performed the experiments: SB CMG CIB MPRB. Analyzed the data: SAOG. Contributed reagents/materials/analysis tools: DC AOG. Wrote the paper: SB CMG AH CIB MP VP ML AOG.