Multi-Organ Transcriptome Dynamics in a Mouse Model of Cecal Ligation and Puncture-Induced Polymicrobial Sepsis

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Purpose: During sepsis, an excessive inflammatory immune reaction contributes to multi-organ dysfunction syndrome (MODS), a critical condition associated with high morbidity and mortality; however, the molecular mechanisms driving MODS remain elusive.

Methods: We used RNA sequencing to characterize transcriptional changes in the early phase of sepsis, at 6, 12, 24 hour time points in lung, kidney, liver, and heart tissues, in a cecal ligation and puncture (CLP)-induced polymicrobial sepsis murine model.

Results: The CLP surgery induced significant changes (adj. p-value<0.05) in expression of hundreds of transcripts in the four organs tested, with the highest number exceeding 2,000 differentially expressed genes (DEGs) in all organs at 12 hours post-CLP. Over-representation analysis by functional annotations of DEGs to the Reactome database revealed the immune system, hemostasis, lipid metabolism, signal transduction, and extracellular matrix remodeling biological processes as significantly altered in at least two organs, while metabolism of proteins and RNA were revealed as being liver tissue specific in the early phase of sepsis.

Conclusion: RNA sequencing across organs and time-points in the CLP murine model allowed us to study the trajectories of transcriptome changes demonstrating alterations common across multiple organs as well as biological pathways altered in an organ-specific manner. These findings could pave new directions in the research of sepsis-induced MODS and indicate new sepsis treatment strategies.

Keywords: sepsis, RNA-seq, reactome, murine model

Introduction

Sepsis is a clinical syndrome resulting from a systemic inflammatory response to infection, with a high mortality rate, which ranges between 10–50%, depending on the patient’s age and disease severity. 1 The molecular hallmark of sepsis is the release of pathogen-associated molecular patterns (PAMPs) from the infectious agent and host release of damage-associated molecular patterns (DAMPs or alarmins), recognized by Toll-like receptors (TLRs) which lead to a profound increase in cytokine production. 2 Dysregulated mechanisms of innate and adaptive immunity result in the early pro-inflammatory phase of sepsis characterized by high production of pro-inflammatory cytokines, followed by an anti-inflammatory phase that translates into functional immunosuppression. 3

In 2016, an international consensus conference defined sepsis as life-threatening organ dysfunction caused by dysregulated host response to infection. 4 As reviewed
recently, neither tissue hypoxia, significant cell death, nor adaptive strategy to overwhelming inflammatory injury could be defined as a predominant isolated mechanism of sepsis-related organ dysfunction. Instead, several other mechanisms are probably involved in the pathogenesis of sepsis, including resistance to pathogen load and tolerance to organ injury by infectious agents and the host’s immune response, as well as transient loss of function of the heart, lung, kidney, liver, and intestine (multiple organ dysfunction, MODS) resulting from the alterations in mitochondrial energy generation and cell cycle arrest induction. As a consequence, sepsis-related alterations activate cross talk between dysfunctional organs. However, the exact mechanisms underlying development of sepsis, and particularly MODS, remains incompletely understood.

Gene expression profiling is a well-recognized method of searching for the molecular basis of diseases. The human blood leukocyte response to acute systemic inflammation was associated with more than 3,700 differentially expressed genes, and systemic administration of LPS altered expression of 8,458 genes in the mouse adrenal gland identifying gene sets and pathways potentially involved in sepsis-induced organ dysfunction. While transcriptional signatures induced by LPS stimulation are conserved between humans and mice, substantial species-specific differences in sepsis response signatures were also recognized; these may represent, among others, differences in organization of molecular studies that can be conducted in septic patients and animal models. For ethical reasons, organ tissue sampling is impractical in critically ill patients, with the exception of post-mortem examinations, and, therefore, the majority of gene expression studies in septic patients have been conducted using only whole blood samples or isolated white blood cells. Although blood comes into contact with the cells, tissues, and organs of the entire organism, constituting a primary aspect of the immune defense system, comparing blood gene expression profiles from septic patients with organ transcriptomes from animal models can cause possible erroneous conclusions. Furthermore, sepsis is a dynamic process within a relatively narrow time window. In clinical studies blood samples are usually obtained at a single time point and, as a consequence, may represent different phases of sepsis, while in animal models organ tissue transcriptional alterations can be analyzed repeatedly at precise times to show a trajectory of molecular changes.

In this study using RNA sequencing we analyzed transcriptional changes in the lung, kidney, liver, and heart tissues, in a cecal ligation and puncture (CLP)-induced polymicrobial sepsis murine model. Our objective was to identify both organ specific alterations in gene expression profiles and to characterize pathway changes conserved across multiple organs in the early phase of sepsis.

**Materials and Methods**

### Mice

In this study, 9–12-week-old C57BL/6J male mice were housed in specific-pathogen free conditions with 14–10 hours of light and dark cycles and free access to food and water until the surgery.

### Institutional Review Board Statement

Mouse experiments were conducted in accordance with The American Association for Accreditation of Laboratory Animal Care guidelines under approval of The University of Washington Office of Animal Welfare (Protocol #4287-01) at the University of Washington South Lake Union medical research campus.

### Cecal Ligation and Puncture (CLP)-Induced Polymicrobial Sepsis

The CLP procedure was performed as previously described and according to the general guidelines. Briefly, under isoflurane anesthesia, mice were laid supine and a midline incision was made in the lower abdomen after shaving and sterilizing the abdominal skin. The cecum was identified and carefully extracted. Stool in the cecum was squeezed distally and a 3–0 suture was tied around the proximal portion of the cecum and secured tightly, interrupting continuity with the bowel and cutting off blood flow. A 21 g needle was used to make a single puncture in the proximal, antimesenteric aspect of the cecum and a small amount of stool was expressed. The cecum was placed back in the abdominal cavity and the abdominal wall closed with 6–0 suture. Skin was approximated and closed with Gluteus tissue adhesive. Immediately post op, mice were given 20 mL/kg normal saline solution and buprenorphine 0.05 mg/kg. Two hours later, mice were given imipenem 25 mg/kg. At 12 hours mice were dosed with normal saline solution 10 mL/kg, and the same doses of imipenem and buprenorphine. Mice were monitored using a clinical severity score every 12 hours, as previously reported. Tissue samples were collected at the specified time points. Anesthesia was induced again with isoflurane and blood collected via cardiac...
puncture. The chest was opened and heart, and lungs collected rapidly into cryovials which were immediately snap frozen in liquid nitrogen. Abdominal organs were collected in a similar fashion immediately thereafter.

**RNA Extraction**
Total RNA was extracted from tissue using MirVana™ PARIS™ Kit (Ambion) according to manufacturer’s instructions. The concentration of each sample was measured by NanoDrop 2000 (Thermo). Total RNA quality was assayed with the RNA 6000 Nano Kit on Agilent 2100 Bioanalyzer (Agilent). For RNA-Seq we analyzed RNA samples from 74 organs derived from 35 animals. If possible, the sets of four organs from the same animals were analyzed. Samples from 12 hours had RNA integrity number (RIN) >6.8, while all samples had <10% rRNA contamination following the poly (A) fraction. **(Table S1)**.

**Quantitative Reverse Transcription PCR (qRT-PCR)**
To synthesize cDNA, 400 ng total RNA was reverse-transcribed with 200 units of MMLV reverse transcriptase (Thermo) and oligo-dT primers in 20-µL reactions. RT reactions were diluted 100-fold with water prior to use in qPCR. GraphGrid Excel-based software was used to acquire, store, and analyze qRT-PCR data. Pair-wise t-test was used to measure statistically significant differences, which are depicted by the size of a circle for each comparison made: p<0.05 by a small circle, p<0.01 by a large circle. **(Table S2)**. Primers sequence is provided in **Table S2**.

**The Poly (A) Fraction Purification and Sequencing**
Messenger RNA (poly(A)-containing mRNA) was purified from the 1–8 µg of total RNA by magnetic beads coated with oligo (dT) using a Dynabeads® mRNA DIRECT™ Micro Kit (Thermo). To estimate the quality and quantity of mRNA samples we analyzed samples with the RNA 6000 Pico Kit with the 2100 Bioanalyzer (Agilent). The sequencing library of each RNA sample was prepared by using Ion Total RNA-Seq Kit v2 according to the manufacturer’s protocol (Thermo). The libraries were prepared from 1–15 ng of mRNA. Briefly, the mRNA was fragmented using RNaseIII and then purified. The fragmented RNA was hybridized and ligated with ion adapters. The RNA fragments were then reverse transcribed and amplified to double-stranded cDNA. Next, the cDNA was purified by magnetic bead based method. The molar concentration and size selection (50–1,000 bp) of each cDNA library was determined using DNA 1000 Kit on Bioanalyzer 2100 (Agilent). Each library was diluted to ~80 pM concentration prior to template preparation and up to four barcoded libraries were mixed in equal volume and used for automatic template preparation on an Ion Chef (Thermo) instrument using reagents from the Ion PI IC 200 Kit (Thermo) and Ion PI v3 Proton Chip. Samples were sequenced on the Ion Proton System (Thermo) according to the manufacturer’s instructions.

**RNA-Seq Data Analysis**
Raw reads were mapped to mm10 reference genome with Ion Torrent RNASeqAnalysis Plugin version 5.0 which utilizes STAR as the primary aligner, and bowtie2 for previously unaligned reads. Gene abundance was quantified with htseq-count (HTSeq framework version 0.6.1) using Ensembl Gene gtf file from UCSC as reference. Differential gene expression was performed with R package DESeq2 version 1.18, using Wald’s test and Benjamini-Hochberg procedure to control false discovery rate (FDR).

**Functional Analyses**
Overrepresentation of Reactome pathways in genes which were differentially expressed was performed with R package ClusterProfiler version 3.6, separately for up- and down-regulated genes. Heatmap visualization was conducted with ComplexHeatmap R package version 2.3.1, using only pathways present in at least two data points and with at least 5% of differential genes present in the pathway.

**Data Availability Statement**
The RNA-Seq data were deposited as the BAM files in the European Nucleotide Archive under the PRJEB20791 identifier.

**Results and Discussion**

**Cecal Ligation and Puncture Induces Sepsis of Moderate Severity**
The CLP model of sepsis can be highly variable in that severity with a specific injury in one lab does not necessarily translate to the same severity in another. We previously titrated the severity of our model to achieve 25–30% mortality at 7 days in C57BL/6J mice to mimic clinical mortality observed in human septic shock. This mortality phenotype with the specific injury described here was stable in our hands over multiple experiments for
years (Figure 1A). In this model, CLP-operated mice develop significantly elevated clinical severity scores compared to sham-operated mice (Figure 1B).

**RNA-Seq Survey Across Multiple Organs in Polymicrobial Sepsis Reveals Dynamics of Transcriptional Changes**

To establish multi-organ gene expression signatures of the early phase of sepsis polymicrobial sepsis was induced by CLP surgery in male mice and compared to sham-operated animals. At 6, 12, and 24 hours after surgery, three animals from each group were sacrificed and the lungs, kidneys, livers, and hearts were removed and frozen. At 12 hours an additional one CLP and sham-operated kidney tissue sample was included. The organs we have chosen are more frequently affected in the course of sepsis and extent of their dysfunction determines patients’ course from infection to recovery or death.\(^{22}\) Altogether, 74 Poly (A)-enriched mRNA samples, representing four organs removed from CLP and sham-operated animals, were sequenced on an Ion Proton sequencer. The principal component analysis showed that the samples clustered according to the tissue origin during the experiment (Figure 1S). The pairwise comparison between mice with CLP induced sepsis and control (sham-operated) mice identified 1,205, 489, 1,276, and 374 differentially expressed genes (DEGs) (an adjusted p-value<0.05) in liver, kidney, lung and heart samples, respectively, at 6 hours, 3,769, 2,457, 3,828, and 2,442 DEGs, respectively, at 12 hours, and 2,481, 1,470, 1,800, and 154 DEGs, respectively, at 24 hours (Figure 2A; Table S3).

The expression of 294/1,019/883, 124/861/48, 829/1,899/1,752, and 862/1,730/1,201 transcripts were found exclusively altered in CLP induced sepsis at least at one time point (6/12/24 hours) in the kidney, heart, liver, and lung, respectively. In contrast, 50, 266, and 20 DEGs were common to all four organ comparisons for 6, 12, and 24 hour time points, respectively (Figure 2B).

At the earliest time point, the number of significantly changed transcripts passed a 1,000 threshold in the lung and liver only, followed by a burst in expression alteration passing the 2,000 transcripts threshold in every organ at 12 hours. At the experiment endpoint (24 hours) the number of DEGs decreased across all the organs where the reduction of DEGs was most striking in the heart. Of note, the number of DEGs seen at all three time points in a given organ was 105, 31, 410, and 391 in the kidney, heart, liver, and lung, respectively (Figure 3).

Our RNA-Seq survey uncovered dynamic transcriptional changes across mouse organs with the most pronounced number of DEGs at the 12 hour time point during CLP-induced polymicrobial sepsis. To date, most of the transcriptome studies in sepsis have surveyed expression changes in peripheral blood mononuclear cells. A meta-analysis by Godde et al\(^8\) of 389 published transcriptomes studies from mouse and human immune cells induced by LPS or CLP procedures showed that expression profiles between humans and mouse models appeared to be highly conserved. However, no clinical study evaluated sepsis-related gene expression profiles in individual organs as the utility of the biopsies in the critically ill patients are not practical. In turn, only a small number of reports have described organ tissue transcriptional profiles in animal models of sepsis; most of these have studied single organs.\(^7\)\(^{23–29}\) Thus, our study, for
the first time, reveals septic transcriptional profiles simultaneous analysis across different organs over the time of early sepsis phase, using RNA-seq.

**RNA-Seq Data Significantly Correlate with qPCR Measured Abundances of Inflammatory and Endothelial Transcripts**

As a way to corroborate RNA-seq data we assayed in qPCR (Figure 4) the same RNA samples as in RNA-Seq survey and included at least three additional RNA samples per group. We used primers to inflammatory (Lcn2, Tnfa, Il-6, Il-1a, Ccl2) and endothelial (Angpt1, Angpt2, Kdr) genes previously studied in sepsis models. In sepsis endothelial dysfunction results from decreased expression of genes that otherwise maintain microvascular integrity. The qPCR confirmed significant alteration of surveyed transcripts across organs in at least one time point and the direction of changes in their expression was in line to other studies utilizing sepsis models. The Spearman correlation coefficient of qPCR and RNA-Seq data for those genes across the time point and organs is 0.72, and is statistically significant. Of note, the qPCR measured expression of Lcn2, that encodes neutrophil gelatinase-associated lipocalin (Ngal), remained significantly elevated irrespectively of the time-point and organ assayed, confirming previously observed sustained expression of this transcript in CLP-induced sepsis.

**Functional Analyses of DEGs Highlight Shared and Organ Specific Biological Pathways Altered in CLP-Induced Polymicrobial Sepsis**

To uncover organ and time point specific changes as well as alterations shared across organs and time, we performed functional analyses of significantly up and downregulated
transcripts (an adjusted $p$-value<0.05 and fold change; FC≥1.5) encompassing 12 types of samples using the Reactome database. The functional analysis yielded 26/0, 25/0, 45/3, and 20/8 of significantly enriched pathways for up/downregulated DEGs in heart, kidney, liver, and lungs, respectively, at 6 hours (Table S4), 55/4, 23/0, 49/31, and 11/43 pathways, respectively, at 12 hours (Table S5), and 5/0, 31/12, 43/38, and 23/25 pathways, respectively, at 24 hours (Table S6). To further generalize the results of functional analyses we defined pathways with a high contribution of changed genes by using the threshold of 5% genes altered in a given term while at the same time being altered in at least two separate types of samples, yielding 41 pathways recurrently altered in our sepsis model (Figure 5).

The CLP upregulated genes categorized under the Immune system consisted of 15 pathways. Pathways overrepresented in all organs for at least one time-point include: Cytokine Signaling in Immune system, Interleukin-1 family signaling, Interleukin-1 signaling, Signaling by Interleukins pathways, Antimicrobial peptides, and Neutrophil degranulation. In contrast, none of the significantly downregulated transcripts could be categorized into the Reactome Immune system category.

As previously reported, the common response at the early phase of polymicrobial sepsis includes activation of inflammatory mediators. Organisms exposed to bacterial endotoxins react with cell surface receptors, including the TLR family, which trigger the release of both proinflammatory and anti-inflammatory mediators through activation of several signal transduction pathways, including nuclear factor κB (NFκB), mitogen activated protein kinase (MAPK), janus kinase (JAK), and signaling transducer and activator of transcription (STAT). Both concurrent pro-inflammatory and anti-inflammatory phase are considered as the sepsis hallmark. Other pathways with a high level of significance were Hemostasis, Platelet activation, signaling and aggregation, Platelet degranulation, and Response to elevated platelet cytosolic Ca2+, which represented the Hemostasis cluster. While all four pathways revealed overrepresentation of upregulated genes at each time point for all organs, with the exception of the lung where, at 12 hours after surgery, they represented significantly downregulated transcripts. Platelets play an important role in sepsis pathophysiology and multi-organ dysfunction as well as in the resolution of inflammation.

Extensive metabolic dysregulation has been implicated in the pathogenesis of sepsis-induced organ failure, which itself is strongly associates with mortality of septic patients. The liver is the central organ for carbohydrate, protein, and fat metabolism and it contributes to the innate immune system. Among significantly downregulated genes, Reactome annotation analysis revealed overrepresentation of genes involved in the metabolism category and represented by seven pathways, mostly related to fatty acids metabolism and their oxidation (Figure 5). The metabolic pathways were already dysregulated at 6 hours post-surgery in the liver, with an increasing level of significance in subsequent time points. The significant signatures of metabolic alterations were also found in septic kidneys (at 12 hours post-surgery) and lungs (at 24 hours post-surgery), but not in the hearts. Interestingly, in the liver at 12 and 24 hour time points the metabolism of proteins related to the asparagine N-linked glycosylation pathway and proteins transport to the Golgi was prominently altered. These two pathways are closely related to other pathway branches that comprise vesicle-mediated transport and were altered at 12 and 24 hours in the liver. Previous studies using the CLP mouse model have highlighted the hepatic changes related to proteins and fatty acids metabolism.

Interestingly, at the 12 hour time-point a parallel increase in RNA metabolism and splicing was observed.
Figure 4 Quantitative polymerase chain reaction (qPCR) gene expression measurements of selected inflammatory (A) and endothelial-related (B) transcripts across multiple organs in CLP-induced polymicrobial sepsis. Total RNA from heart, lungs, kidneys, and livers was used in reverse transcription reactions with oligo-dT primers. cDNA was used in qPCR with gene-specific primers.14 Data are represented as mean±standard error of the mean (n=6 mice in each group). Statistical differences between two means (p-value) are shown by the size of the solid circles and additionally denoted with an asterisk (*): p<0.05 by a small circle, p<0.01 by a large circle.14
that could be attributed to the overall elevated transcription and protein translation in the liver. Indeed, the hepatic synthesis of acute phase proteins is a hallmark of infection and inflammation, and N-glycosylation is a main posttranslational modification of these proteins that regulates both their function and lifespan. Reactome annotation of significantly downregulated transcripts also revealed hepatic suppression of steroids metabolism, bile acid, and bile salt metabolism pathways (Tables S5 and S6) at 12 and 24 hours post-CLP. The impairment of bile formation in the liver during sepsis promotes hyperbilirubinemia associated with intrahepatic cholestasis. It has been shown before that LPS or pro-inflammatory cytokines alter bile acid transporters expression in oxidative stress that results in the inhibition of cAMP-dependent transport function, and by the alteration of hepatocytes cytoskeleton and liver tissue morphology, ultimately contributing to altered bile flow.

Signal transduction by the G protein-coupled receptors (GPCRs) was another major cluster comprised of seven significantly altered pathways. GPCR-related pathways, including WNT signaling, play an important role transducing signals that lead to the production of inflammatory cytokines, and regulation of communications associated with increased blood flow and increased vascular endothelial permeability. Therefore the GPCRs function to both control inflammation and its resolution. The expression changes in GPCR-related pathways were most prominent in lungs with five, three, and six out of seven pathways altered at 6, 12, and 24 hour time points, respectively. The pathways altered in the lung comprised both down and upregulated transcripts. On the other hand, in the kidney,
three and five GPCR-related pathways were significantly altered at 12 and 24 hour time points, and only represented by overexpressed genes. Signaling by WNT was one of the pathways overrepresented in lungs and negatively regulated at 6 hours while in the heart the downregulation occurred 12 hours post-CLP. Previous studies utilizing experimental mouse models of sepsis highlighted the WNT-pathway deregulation in the adrenal gland, heart, and lungs.

The small molecules transport pathway and its derivative term, the Solute Carrier (SLC)-mediated transmembrane transport, represented another major Reactome pathway with alterations in gene expression. The pathways comprised hepatic and pulmonary upregulated transcripts at the 6 hour time point. Conversely, at later time points downregulated transcripts contributed to the significant changes in either of these pathways in the kidney and liver.

Finally, the extracellular matrix (ECM) organization comprised both overexpressed and downexpressed gene sets found in livers and lungs, respectively, 24 hours after surgery. The ECM plays a significant role in the bacterial infection as its adhesive components, including collagens, fibronectin, and laminin, are common binding sites for the invading pathogens. Moreover, the host tissues and immune cells cross-talk as the tissue repair unfolds within the ECM microenvironment. In turn, the ECM undergoes an extensive remodeling both as a result of its degradation by pathogens released proteases and local immune cells response to the insult. Importantly, the endothelial cells associated ECM, the glycocalyx, undergoes extensive remodeling, ultimately leading to the disruption of the integrity of the endothelial barrier, a hallmark of sepsis. The functional analyses of our transcriptomic data pinpointed over-representation of altered transcripts associated with the hepatic and pulmonary ECM organization. In the liver, the upregulated genes were annotated to the significant terms related to collagen degradation and production at the same time. In the lungs, on the other hand, the downregulated genes were significantly over-represented in the term of collagen and elastin formation. Elastin is a component of a complex fiber structure distributed in compartments of the mammalian lung, including the pulmonary blood vessels. These results suggest ongoing tissue remodeling of these organs.

Overall, from the functional analyses it could be concluded that all four organs examined here manifest upregulation of transcripts related to inflammatory response and hemostasis during the early phase of a mouse sepsis model. In contrast, the downregulated genes were over-represented in metabolic pathways related to lipid metabolism, and were observed in the liver at all time points, and in the lungs and kidney at 12 and 24 hours post-CLP, respectively. The changes in proteins and RNAs metabolism were the functional categories that were significantly altered in the liver only.

Post-CLP Organs Tissues Contain DEGs Sets That are Consistent with the Transcriptome Signatures from Septic Peripheral Blood in Mouse Models and Human

In order to relate the DEGs detected in our experimental model across the organs to available transcriptomic sepsis signatures, we compared our dataset to a set of genes from a compendium of immune signatures derived from multiple sepsis human and mouse peripheral blood transcriptome studies. Specifically, we focused on the 667 genes set (Table S7), named the “leading edge” genes, enriched in peripheral blood transcriptome in both species under septic conditions. In our dataset (Table S7) grouped as “leading edge” 281 transcripts were altered in at least one sham-operated vs CLP comparison. Based on these transcripts the unsupervised hierarchical clustering of samples from a given organ nearly perfectly separated sham-operated from CLP samples, with an exception of two samples, one from heart and one from kidney tissues (Figure 6).

Conclusion

The CLP induced systemic inflammation evoked dynamic transcriptional changes across heart, kidney, liver, and lung tissues, with the highest number of transcripts altered at 12 hours post-CLP. Moreover, host response set of genes to sepsis was mediated by groups of co-expressed transcripts, regardless of the involved organ. Functional analyses with Reactome database of up- and downregulated DEGs yielded numerous significantly enriched pathways including the Immune system, Hemostasis, Metabolism, Signal transduction, and ECM organization that were shared between at least two organs. Thus, RNA sequencing allows us to study transcriptome dynamics and enables
a holistic view of sepsis-altered pathways, several of these previously unexplored, that could pave new directions in research of sepsis-induced organs dysfunction and provide clues for sepsis treatment strategies.

Abbreviations

MODS, multi-organ dysfunction syndrome; CLP, cecal ligation and puncture; DEGs, differentially expressed genes; ECM, extracellular matrix; GPCR, G protein-coupled receptor.

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Disclosure

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