ORIGINAL ARTICLE

Meta-Analysis of Human Molecular Responses to *Staphylococcus aureus*

Sidra Younis¹, Farha Deeba², Qamar Javed³, Miroslav Blumenberg⁴

ABSTRACT

**Objective:** To compare the local and systematic transcriptional responses of human body to *S. aureus* or its components.

**Study Design:** Meta-analysis of microarray data.

**Place and Duration of Study:** The study was conducted at R.O. Perelman Department of Dermatology, The NYU Cancer Institute, NYU Langone Medical Center, New York, USA, from March 2015 to May 2015.

**Materials and Methods:** Public repository “GEO Datasets” was searched using key term “*Staphylococcus aureus*” for data sets covering effects of *S. aureus* infection in *Homo sapiens* cells. The microarray data for immune cell responses to *S. aureus* was analyzed using Rank Prod, RMA Express and DAVID software.

**Results:** The analysis has shown that *S. aureus* infection was responsible for inducing immunity, platelet activation, vasodilation, MyD88 dependent gene expression and cell cycle. It suppressed gene expression of normal cell processes, protein catabolism and apoptosis. Heat-inactivated *S. aureus* challenged the cell induced immunity, cell cycle, growth regulators, anti-apoptosis and anticoagulant genes, while suppressed the genes for adaptive immunity, carbohydrate synthesis and Myd88 dependent pathway. Furthermore, in the *S. aureus*-infected patients the genes for defense, innate immunity, solute receptors and anti-apoptotic processes were upregulated, whereas adaptive immunity and positive regulators of apoptosis were downregulated.

**Conclusion:** MyD88 signaling pathway, ubiquitin mediated protein catabolism and IFNγ mediated cell death processes can be targeted for treatment against virulent *S. aureus* infections.

**Key Words:** Heat-killed *S. aureus*, Immunity, Live *S. aureus*, Meta-Analysis, *Staphylococcus aureus*.

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Introduction

*Staphylococcus aureus* (*S. aureus*) is a gram-positive bacterium found as part of normal flora in 50% of population. It is a common cause of skin infections including impetigo, folliculitis, cellulitis, infected ulcers, wounds, respiratory diseases and food poisoning. The severity of *S. aureus* infections can range from mild local to severe deeper soft tissue infections requiring hospitalization. Many antibiotics have been designed to target the bacterial wall synthesis. However, antibiotic-resistant forms of pathogenic *S. aureus*, e.g. methicillin-resistant *S. aureus* (MRSA), have become a widespread clinical problem. *S. aureus* produce enzymes that protect it from degradation and have acquired two types of strategies to avoid killing by host immune cells. *S. aureus* infection activates cell growth and metabolic pathways, which provides a rich growth environment. It activates anti-apoptotic protein transcription to decrease apoptosis of the infected cells.¹

*S. aureus* and associated toxins are recognized by a
number of surface and sub-cellular receptors in host cells, such as Toll-like receptor (TLR), retinoic acid-inducible gene-I-like receptors (RIG) and nucleotide-binding oligomerization domain-like receptors (NOD) like receptors. The activation of these receptors and co-stimulatory molecules results in cascades of reactions which lead to activation of nuclear factor kappa-light-chain-enhancer of activated B cells (NFkB) or interferon gamma (IFNG) through Myeloid differentiation primary response 88 (MyD88)-dependent or independent pathways. Subsequent production of cytokines, chemokines and complement molecules activate either innate or adaptive immune system, based on the triggering factors. Additionally, pathogen containing phagosomes are processed by endoplasmic reticulum and presented as peptides on the surface of antigen presenting cells as a complex with major histocompatibility molecules (MHC I or MHC II) or heat shock proteins. Some heat shock proteins can process pathogen specific peptides independent of endoplasmic reticulum and Golgi apparatus, thus giving extra advantages in antigen presentation and activation of innate immune specific proteins and pathways.

Although considerable information is available about S. aureus virulence, key pathways or signaling molecules are still obscure. To explore this problem, we carried out a meta-analysis of freely available data in public repositories to compare the local and systematic transcriptional responses of human body to S. aureus or its components.

**Materials and Methods**

We have compared transcription differences between untreated, control cells versus those challenged with active S. aureus, inactive S. aureus and S. aureus infected patients' blood.

**Search Strategy**

Public repository “GEO Datasets” was searched using key term “Staphylococcus aureus” for data sets covering effects of S. aureus infection in Homo sapiens cells. In these studies, effects of live heat- or UV-inactivated S. aureus cultures were studied in epithelial or immune cells. The cells were treated for different time periods. In 3 data sets differential expression in blood drawn from infected patients and healthy controls was studied. (Fig 1)

**Analysis of differential expression in cells infected with live S. aureus**

Data from 6 studies, 9 datasets, 214 microarrays and 21,632 genes were combined. In these studies, S. aureus infection responses were observed in macrophages, neutrophils, leukocytes, human umbilical vein endothelial cells and a hepatocellular carcinoma cell line. Affymetrix and Illumina platforms were used for differential expression analysis in infected and control cells. For this analysis, one data set was compromised; the first due to missing data and the second for fewer common genes with other data sets (Table 1 and Fig 2).

| Studies | Acc. No. | MA/C/T | Cell type | Infection | Platform |
|---------|----------|--------|-----------|-----------|----------|
| 1       | GSE39889 | 4-4    | Neutrophils | S. aureus | GPL570 [HG-U133_Plus_2] Affymetrix |
| 2       | GSE13670 | 15-15  | Macrophages | S. aureus | GPL570 [HG-U133_Plus_2] Affymetrix |
| 3a      | GSE13736 | 1-1    | Endothelial cells | S. aureus | GPL570 [HG-U133_Plus_2] Affymetrix |
| 3b      | GSE13736 | 1-1    | Endothelial cells | S. aureus | GPL570 [HG-U133_Plus_2] Affymetrix |
| 3c      | GSE13736 | 1-1    | Endothelial cells | S. aureus | GPL570 [HG-U133_Plus_2] Affymetrix |
| 4       | GSE16837 | 20-78  | PMNL | S. aureus | GPL570 [HG-U133_Plus_2] Affymetrix |
| 5       | GSE16129 | 10-42  | PBMC | S. aureus | GPL10575 [HG-U133A_2] Illumina |
| 6a      | GSE46943 | 3-3    | HCCC | S. aureus | [HuGene-1_0-0] Affymetrix Human |
| 6b      | GSE46943 | 3-3    | HCCC | S. aureus | [HuGene-1_0-0] Affymetrix Human |
| Total   | GSE16837 | 20-78  | PMNL | S. aureus | GPL570 [HG-U133_Plus_2] Affymetrix |
| GSE, Data series; MA, Microarray; C, Control; T, Treated; PMNL, Polymorphonuclear cells; PBMCs, Peripheral blood mononuclear cell, HCCC, Hepatocellular carcinoma cell line; S. aureus, Staphylococcus aureus; GPL, Gene platform |
respectively. This analysis included 2 studies, 3 data sets, 65 microarrays and 19,679 genes (Table 2).

### Differential expression analysis of *S. aureus* infected patients’ blood

Three studies containing 3 data sets were found comparing differential gene expression in blood of *S. aureus*-infected patients and healthy controls. These studies used Affymetrix (GPL570, GPL571) and Illumina (GPL6947) platforms. After combining the data from three different studies, 279 microarrays and 15,001 common genes were obtained. In this analysis the study using Illumina (GPL6947) platform and 143 microarrays was modified before combining. The empty spaces in data were replaced with "1" to use in RankProd software, as described in Mimoso *et al.*, 2014 (Table 3).

### Meta-analysis procedure

The data for transcriptional profiling were deposited in respective gene expression series as CEL or TXT files. These were processed for further analysis using RankProd software. RankProd software was used to identify the genes differentially expressed in *S. aureus*-challenged cells with *p*-values less than $10^{-5}$. RankProd analysis for each study group produced a table representing induced or suppressed genes in challenged cells. The DAVID software was used to get tables, charts and clusters for the induced or suppressed genes with *p*-value less than $10^{-4}$ obtained from RankProd output tables (Fig 1).

### Comparison of live *S. aureus* infected vs. control cells

*S. aureus* can cause skin and systemic infections. We analyzed differential expression in endothelial cells, immune cells and hepatocellular carcinoma cell lines challenged with live *S. aureus*. The results for the comparison of ontological categories induced and suppressed in *S. aureus* infected cells are given in Table 4. We found that extracellular proteins, membrane receptors and intracellular genes that are involved in defense response were induced. Whereas intracellular processes including replication, gene expression, cell death and ubiquitin mediated protease pathway were suppressed. The clusters containing overlapping ontological categories of genes induced in *S. aureus*-infected cells are presented in Table 4. In the top regulated cluster the ontological categories including “Extracellular region part” and “extracellular space” were found (ES 14.45). Extracellular region part mainly included pattern recognition receptors (PRRs), cytokines, chemokines, hematopoietic, plasminogen and thrombospondins. In addition, genes involved in blood coagulation and thrombogenesis, such as inflammatory cytokines and chemokines, were found in this cluster. Interestingly, in cells challenged with live *S. aureus*, interleukin 1 receptor (IL-1R) and TLR specific MyD88-dependent pathway signaling molecules
including mitogen-activated protein kinase (MAPK) and its regulators, which activate multiple processes in cells as gene expression, metabolism, cell division, cell morphology and cell survival, were largely induced. Anti-apoptosis genes were also upregulated.

In the 2nd cluster ontological categories “plasma membrane part” and “intrinsic to plasma membrane” were found (ES 11.47). This cluster presented genes for receptors, specifically G-protein coupled receptors (GPCRs) that activate ligand specific secondary messengers for example, phosphatidylinositol-calcium, adenylyl cyclase 3 (ADCY3) and voltage-dependent potassium channels. In the 3rd cluster ontological categories “response to wounding”, “inflammatory response” and “defense responses” were found (ES 8.65). The genes present in this cluster were largely overlapping with the first cluster. We found that genes involved in activation of prothrombin complex following stable fibril formation, were uniquely expressed in this cluster. We also found genes for cell surface receptors including glycoprotein, voltage gated channels, tumor necrosis factor (TNF) receptor family, integrins and importantly TLR2. In the 4th cluster (ES 6.42) ontological category “Positive regulation of biosynthesis” and “positive regulation of gene expression” are presented that included genes involved in the gene expression and replication.

In the 5th cluster ontological categories “cell-cell signaling” and “synaptic transmission” were upregulated (ES 6.31). In the 6th cluster (ES 5.72), we found gene ontologies “cell adhesion” and “cell-cell adhesion”. In the 7th cluster ontological categories “behavior” and “chemotaxis” were found (ES 5.59). We found the genes involved in cell to cell communication for example, calcium ion dependent transmembrane cell to cell junction, cadherin and protocadherins proteins in this cluster. Additionally, the genes for the proteins which bind to the cytoskeleton to carry cellular functions, including cell movement, cell division, endocytosis, and movement of organelles, were induced. The 8th cluster upregulated in S. aureus-infected cells contained ontological categories “vasculature development” and “blood vessel development” (ES 5.45). We found that genes for transcription factors, enzymes and signaling molecules were upregulated, they participate in transcription, replication and vasculature development processes. In the 9th cluster the ontological categories “cell migration” and “localization of cell” were found (ES 5.06). Detailed study of these ontologies revealed genes contributing in the process of cytoskeleton interactions which play important function in mitosis and receptor-ligand interaction were upregulated. Similarly, downstream effector molecules that participate in the transmission of signals from tyrosine kinase receptors and small GTPases to the actin cytoskeleton were induced. In the 10th cluster ontological category “regulation of myeloid cell differentiation” and “positive regulation of myeloid cell differentiation” were found (ES 5.00). The histone cluster and transcriptional factors were upregulated that specifically modulate the myeloid
cell differentiation, possibly participating in formation of immune cells which provide defense response against external stimuli. The expression of proteins for normal cell processes, such as DNA repair, replication, cell division, development, might point the ability of S. aureus to decrease apoptosis of infected cells by upregulating cell processes. Moreover, processes e.g., apoptosis, cytoskeleton development to improve immunocytes taxis, receptors for bacterial components and reduced energy expenditure represented cells defense mechanisms against bacterial infection.

In table 4 the clusters of gene ontologies suppressed by live S. aureus infection in human cells are presented. In the top cluster (ES 11.62), gene ontologies such as “intracellular organelle lumen” and “nuclear lumen” were found. In S. aureus-infected cells defense, thrombus formation, immunity and chemotaxis and the gene expression processes were particularly suppressed. The genes found in clusters 1, 2, 3, 5, 6, 9 and 10 were overlapping and involved in the processes confined to nucleus, mitochondria and endoplasmic reticulum. These processes were related to the gene expression, protein processing and translocation.

Nuclear receptor-mediated transcription was also repressed. We also found that the proteins for DNA modification, including histones, telomeres maintenance and DNA repair processes, were downregulated. The genes present in this cluster were completely different from the genes present in ontological category “positive regulation of gene expression” (cluster 4) in Table 5. The 2nd (ES 3.41), 3rd (ES 2.85), 5th (ES 2.65), 6th (ES 2.59) and 10th clusters (ES 1.77) again included genes that are involved in RNA polymerase II promoter dependent transcription process. In the 4th cluster gene ontologies such as “macromolecule catabolic process” and “protein catabolic process” were found (ES 2.80). This cluster presented genes for protein catabolism via ubiquitin-proteasome pathway. Similarly, in the 9th cluster we found proteins involved in ubiquitin-proteasome specifically involving ring finger proteins mediated proteolysis (ES 1.89). The 7th cluster represented gene ontologies such as “cell death” and “positive regulation of programmed cell death” (ES 2.55). These ontologies included genes such as caspasas that are necessary for the positive induction of cell death. Interestingly, we found the genes that modulate myeloid cells apoptosis. The proteins involved in mediating IFNγ induced cell death were also suppressed in the S. aureus-infected cells. In the 8th cluster cytoskeleton specific proteins were present in gene ontologies “non-membrane bound organelle” and “cytoskeleton” (ES 2.10). We found the proteins specific for microtubule cytoskeleton formation and regulation were suppressed. We have also found that proteins which may play an important role in integrin β-1 or B cell receptor (BCR) mediated signaling in B- and T-cells were also suppressed.

Comparison of heat inactivated S. aureus challenged vs. control cells

The live S. aureus and inactivated S. aureus challenged cells present different sets of differentially expressed genes. Therefore, we have compared differential expression of heat inactivated

| Table 5: Defense response genes induced in humans by Inactive S. aureus |
|---------------------------------------------------------------|
| Sr | Gene Ontologies | P Value | Sr | Gene Ontologies | P Value |
|---|-----------------|---------|---|-----------------|---------|
| 1 | response to wounding inflammatory response | 5.57E-30 | 2 | response to wounding inflammatory response | 2.56E-25 |
|   | defense response | 1.49E-23 |   | defense response | 1.39E-23 |
| 2 | R. of apoptosis | 1.85E-14 | 3 | R. of apoptosis | 1.85E-14 |
|   | Neg. R. of apoptosis | 2.83E-10 |   | lysozome | 4.26E-05 |
| 3 | R. of FAS | 2.07E-14 | 4 | R. of FAS | 3.07E-14 |
|   | cytokine activity extracellular region part | 1.42E-11 |   | cytokine activity extracellular region part | 1.29E-10 |
| 5 | cytokine activity Signaling by GPCR | 9.95E-06 | 6 | R. of RNA binding R. of transcription factor activity | 1.31E-03 |
|   | negative regulation of RNA binding R. of transcription factor activity | 1.31E-03 |   | hexose biosynthetic process | 1.74E-03 |
| 7 | response to laminin response to LPS response to cytokine stimulus | 3.54E-07 | 8 | response to laminin response to LPS response to cytokine stimulus | 3.54E-07 |
|   | R. of apoptosis | 1.85E-14 |   | R. of apoptosis | 1.85E-14 |
|   | Pos. R. of apoptosis | 3.24E-07 |   | Pos. R. of apoptosis | 3.24E-07 |
| 8 | R. of cytokine production | 4.28E-10 | 9 | R. of cytokine production | 4.33E-07 |
|   | extracellular stimulus | 1.56E-10 |   | extracellular stimulus | 1.56E-10 |
|   | response to nutrient | 1.56E-10 |   | response to nutrient | 1.56E-10 |
| 9 | response to organic substance Response to steroid hormone | 2.15E-15 | 10 | response to organic substance Response to steroid hormone | 2.15E-15 |
|   | glutathione metabolic process | 1.57E-09 |   | glutathione metabolic process | 1.57E-09 |
|   | coenzyme metabolic process | 1.57E-09 |   | coenzyme metabolic process | 1.57E-09 |
| 10 | R. of cytokine production | 4.28E-10 |   | R. of cytokine production | 1.83E-06 |
|    | Intestinal IgA production | 2.50E-03 |   | Intestinal IgA production | 2.50E-03 |
|    | antigen processing and presentation | 2.72E-03 |   | antigen processing and presentation | 2.72E-03 |

R. Regulation; Pos. R., Positive regulation; Neg. R., Negative regulation; LPS, Lipopolysaccharide; PCD, Programmed cell death
S. aureus challenged versus control cells using Meta-analysis.

The clusters of gene ontologies induced in heat-inactivated S. aureus challenged cells are given in Table 5. We found that three major processes were induced including defense response, regulation of apoptosis and cytokine production. We found that the 1st (ES 25.56), 6th (ES 8.39) and 9th (ES 7.44) clusters presented genes for defense mechanisms against bacterium. The ontological categories such as “response to wounding”, “inflammatory response” and “defense response” are present in the top regulated cluster (ES 25.56) as observed in Table 3 and 4. Like previous comparisons in S. aureus-challenged vs. control cells, we found that innate immunity specific Gram positive bacteria receptor present on cell surface, TLR2, or general bacterial receptor TLR8 found in intracellular endosomes were upregulated. Interestingly, proteins for MyD88-dependent TLR response pathway were induced. In contrast to live S. aureus challenged cells where innate immunity specific chemoattractant i.e. myeloid cells activators were overrepresented, in inactive S. aureus chemoattractant for innate as well as adaptive immunity specific cells were induced. Specially, a number TNF-α complex mediated macrophages-specific cytokines and co-stimulatory cluster of differentiation (CD) molecules, that share immune response by macrophages internalization, cell adhesion and T cell receptor (TCR) binding, were induced. In wounding response platelet derived growth factors and gene clusters were upregulated. Proteins involved in receptor-mediated endocytosis and regulators of apoptosis were also found in this cluster. The 6th cluster presented the signaling mechanisms in “response to bacterium” and “response to molecules of bacterial origin” (ES 8.39).

The innate immune receptor TLR2 and innate immune response genes including chemokines (C-C motif) cytokines including IL-1β, IL-6, IL-10, and Interleukin-1 receptor-associated kinase 3 (IRAK3), cytokines regulators were upregulated. Additional transcription and nuclear factors (FOS, JUN and NFKB1A) that participate in extrinsic cell death processes were also upregulated. In 9th cluster, genes for the response systems to endogenous peptide hormone stimulus were induced (ES 7.44). We observed that most of the calcium stimulated pathways involved in metabolism and signaling to peptide hormone leading to initiation of cell death pathways were induced.

In 2nd (ES 11.01), 3rd (ES 10.95) and 7th (ES 8.13) clusters; negative regulators of apoptotic pathways, that lead to increased cell survival were observed. Specifically, downstream signaling molecules for NLRs were induced. The major genes complexes which regulate anti-cell death processes were induced in this cluster including TNF-α and associated proteins, IILs (1, 6 and 10), DNA repair proteins, nuclear factors, oncogenes, transcription activators and serine threonine kinases. Cell cycle regulation proteins stimulated by MAPK pathways were also found in this cluster. Cytoskeletal proteins involved in cell communication, attachment and locomotion were also induced.

In the 4th (ES 10.50), 5th cluster (ES 9.18), 8th (ES 7.52) and 10th (ES 6.88) clusters genes involved in cytokine production and regulation were found. In chemokines and cytokines involved in defense against extracellular stimulus dominating the adaptive immune cells were found. In addition, growth regulators and proteins involved in local inflammatory process were induced. In addition, signaling pathways which use G protein-coupled receptors (GPCR) after activation by prostaglandins, adrenomedullin and immune cells were induced. The clusters representing the gene ontologies for suppressed genes in S. aureus challenged human cells are given in Table 5. The top cluster included ontological categories such as “plasma membrane part” and “integral to plasma membrane”. In this cluster the genes for adaptors and receptors largely associated with adaptive immune system, including lymphocytes and natural killer cells were found. A large number of the antigen-presenting proteins (CD molecules) were also suppressed. Additionally, adaptive immunity specific antigen processing proteins for MHCII family were downregulated. Adhesion molecules, dendritic cells specific genes were downregulated. Th1 regulators were also suppressed. ATP dependent constitutive or basal glucose transporters and some nucleoside transporters were downregulated. Calcium-mediated signaling molecules, G protein receptors, adaptors such as ADCY, and chemokines...
participating in cell proliferation, were found in this cluster. The receptors for Gram negative bacteria and immunoglobulin E (IgE) Fc fragments were also suppressed.

In the 2nd cluster (ES 3.73) the ontological categories “defense”, and “inflammation” related processes were presented. In detail, study showed that signaling pathways linked to Gram negative bacteria specific cell receptors (TLR4, TLR5) were presented. Lymphocyte specific antigen proteins, some myeloid and neutrophil specific proteins that bind TLR4 receptor to mediate downstream signaling were downregulated. In addition, MYD88 that acts as an adaptor downstream of TLR2 and TLR4 receptors was repressed. In the 3rd cluster proteins associated with “lysosome” involved in degradation of protein were represented (ES 3.69). These included CD proteins, MHCII complex which specify antigen presenting cells recognition and major enzymes important in degradation of the targeted molecule such as cathepsins, catalases, lipases and peptidases. The the 4th cluster presented ontological categories such as “membrane invagination” and “endocytosis” (ES 2.66). Interestingly, we found that the genes present in this cluster were related to endocytosis and intracellular processing of clathrin coated vesicle, which suggested that genes involved in phagocytosis of foreign bodies were downregulated. In 5th (ES 1.86) and 6th (ES 1.79) clusters of S. aureus challenge downregulated genes, the genes for “negative regulation of DNA binding” i.e. transcription process were principally found. The repressed genes were mostly DNA damage-induced transcriptional regulator and cytokine production genes. Surprisingly, we found that carbohydrate metabolism genes were downregulated.

In the 7th cluster (ES 1.60), we found that GTPase-Ras mediated signaling pathway proteins, that participate in endocytosis after binding with cations e.g., zinc, were downregulated. In the 8th cluster ontological category “response to extracellular stimuli” and “nutrient” was found (ES 1.56). These ontological categories included organic substance response genes that lead to activation of stress response, wound healing proteins. In addition, proteins for cell death regulation were also downregulated. In the 9th cluster the “glutathione metabolic process” and “coenzyme metabolic process” ontologies were represented (ES 1.53). Study of the genes in these clusters showed that enzymes processing glutathione were downregulated. In the 10th cluster antibody production and antigen presentation and processing activity is represented (ES 1.48). We found that in this cluster MHCII antigen presenting and processing proteins as well as adhesion proteins were presented.

**Comparison of transcriptional profiles of blood from S. aureus infected patients vs. healthy controls**

*S. aureus* is an important human pathogen which causes local as well as systemic infections. The study of infected blood transcriptome analysis should provide valuable information about systemic changes in molecular expression of leukocytes during infection that can be used for rapid diagnosis and therapeutical purposes. In present study, the transcriptional microarray data from blood of infected neonates and adults was combined and analyzed to identify the differentially expressed genes in *S. aureus* infected versus healthy controls through Meta-analysis (Table 6).

The clusters of gene ontologies induced in *S. aureus* infected human blood are given in Table 6. Principally, we found that genes for bacterial components recognition receptors and downstream processing pathways including internalization, processing and inflammatory responses were induced.

In the comparison of differential expression of genes in blood of *S. aureus* infected patients and controls, we found that the ontological categories “defense response”, “response to wounding” and “inflammation” are present in the top regulated cluster as observed in previous comparisons (ES 20.21). Innate immunity specific PRR's for Gram positive bacteria including TLRs and NLRs were upregulated in blood of infected patients. Acute inflammatory response genes including complement components, co-stimulatory molecules as CD55 and CD163, macrophages specific phagocytic receptor and associated enzymes were induced. The chemoattractant released from innate immunocytes and specific for adaptive immunocytes were upregulated. IL-18 released from antigen presenting cells and cytokines which affect thymus as well as proliferation of T-cells were also found. In addition,
IL-25 was induced which may lead to MyD88-dependent activation of NF-κB and also required for cytokine production and response mediation. Innate immunity specific phagocytosis, immune cells activators and thrombospondin, important in increasing movement of vesicle inside the cell, were also upregulated.

The 2nd (ES 10.57), 3rd (ES 7.80), 4th (ES 6.86), 6th (ES 5.29) and 8th (ES 5.11) clusters included genes for PRRs, TLRs (1, 2, 5, 6 and 8), sodium-chloride dependent neurotransmitter symporters.

The 2nd (ES 10.57) cluster represented gene ontologies such as "hemostasis" and "complement and coagulation cascade" were upregulated (ES 5.79). The genes included in this cluster were mostly involved in hemostasis process, concentrating immune cells density at site of injury. The upregulated gene complexes included coagulation factors, platelet glycoproteins, plasminogen activators and thrombomodulins. In addition, calcium binding proteins which accelerate ATP mediated fibrin formation process were induced. In 7th (ES 5.12) and 9th (ES 5.29) clusters genes for transport of lipoprotein particle by membrane invagination and vesicle formation were induced. Cytoskeleton and calcium transport genes involved in the process of translocation were also induced.

The 10th cluster presented gene ontologies such as "protein dimerization activity" and "homodimerization activity" (ES 4.49). We found that proteins involved in process of gene expression were largely upregulated; principally RNA polymerase II-dependent transcription was induced. In addition, transcription regulators as well as apoptosis mediators were induced. Similarly, cofactors (Ca^{2+} and Zn^{2+}) mediated guanylate cyclase and GTP derived energy process involved in processing of pathogen containing cells were induced.

The top cluster (ES 9.98) of the genes downregulated in S. aureus-infected patients included ontological categories "translational elongation" and "gene expression" (Table 6). These categories revealed that cytosolic and nucleic ribosomal proteins were the major suppressed genes in the top cluster. In addition, some translation elongation factors were also suppressed. Surprisingly, in contrast to
transcription of immunity specific genes induced in the infected patients, the ribosomal proteins transcription was largely suppressed. Similarly, the 3rd cluster (ES 6.28) presented the organelle lumen and nuclear lumen. The proteins for cell cycle progression, adaptive immune cells activation genes, glucose metabolism and positive regulation of cell death genes were also downregulated. The the 8th cluster (ES 3.89) again presented genes involved in nuclear processes such as formation of ribosomal RNA.

In the 2nd (ES 7.92), 4th (ES 5.66) and 10th (ES 3.50) clusters adaptive immunity specific genes involved in lymphocytes activation, proliferation, differentiation and regulation, involving both B- and T-lymphocytes were found. Co-stimulatory and adhesion molecules and accessory proteins present downstream in T-cell receptor signaling pathways which subsequently activate the two processes were found in suppressed genes: 1) expression of certain cytokines that enhance proliferation, differentiation and immune response 2) and downregulation of ubiquitin mediated proteolysis. In summary, The critical genes for the activation of helper T cell, Th1 (IFNγ) and Th2 (IL-4) lymphocytes, such as CD40LG and IL4, were also repressed.

In the 5th cluster ontological categories such as “graft versus host disease”, “cell adhesion molecules” and “antigen processing and presentation” were presented (ES 5.21). The genes involved in maturation of MHCII molecules that leads to survival of natural killer (NK) cells in response to infection-mediated activation were found in this cluster. In 6th cluster gene ontologies including “T-cell receptor complex” and “Regulation of calcium signaling” were found (ES 4.81). Proteins which have probable role in assembly and expression of the TCR complex as well as signal transduction upon antigen triggering were also downregulated.

The the 7th cluster presented gene ontologies such as “plasma membrane part” and “intrinsic to plasma membrane part” (ES 4.22). The genes present in these categories were plasma membrane integral proteins involved in adaptive immunity activation. Furthermore, a large number of the genes and processes related to adaptive immunity including IL-2 regulators, lymphocytes attractants, IL receptors (1, 2, 5, 7) stimulated Jak-stat signaling molecules, metal ion transporters (Ca²⁺, Na⁺, K⁺), active T-cells survival proteins, T-cell mediated mitogen-activated protein kinase (MAPK) pathway gens, and genes for IFNγ synthesis were suppressed. In 9th cluster (ES 3.76) ontological categories “positive regulation of apoptosis” and “Positive regulation of programmed cell death” were presented. The positive regulator of apoptosis, apoptosis-related signaling pathways and NKs activated formation of toxicity proteins were found here.

Discussion

The live S. aureus induced genes for both innate and adaptive immunity, where platelet function and vasodilation dominated the immune response. Moreover, MyD88-dependent signaling pathways were induced. Interestingly, unlike previous analyses live S. aureus did not modulate the apoptotic process as a cytoprotective strategy; but upregulated genes for normal cellular processes for example gene expression, neuron signaling and importantly mitotic cell cycle. This survival strategy was already reported in obligatory bacteria, which survive either by stimulating cell processes thus providing infected human cells resistance to the apoptosis stimulus, or directly regulate apoptosis machinery. The downregulated processes included gene expression, ubiquitin mediated protein catabolism and IFNγ mediated cell death processes. Furthermore, transcription process of innate immune response genes was induced whereas this process was suppressed for other cell processes.

The differential expression in the cells challenged with inactivated S. aureus was slightly different from live S. aureus challenged ones as revealed from the Meta-analysis. Similar to live S. aureus, positive induction of cellular processes such as cell cycle and growth regulation was also seen in inactivated S. aureus challenged cells proposing the comparable cytoprotective mechanism. Importantly, in inactivated S. aureus challenged cells carbohydrate biosynthesis process was downregulated whereas this process was upregulated in live S. aureus infected cells. Further MyD88 signaling pathway was induced by live S. aureus infection while suppressed by inactivated S. aureus treatment thus defining different cellular defense strategies against live and inactive S. aureus (Table 5). Interestingly, the results for Meta-analysis of

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differential expression in the blood of infected patients were marginally different from live- and inactivated- \textit{S. aureus} challenged cells. MyD88-independent signaling pathway was induced unlike the active \textit{S. aureus} mediated responses where MyD88-dependent signaling pathway was more prominent. Moreover, we observed that membrane receptors including symporters, aquaporin's and solute carriers were over-represented in this analysis. Similar to the live- and inactivated-\textit{S. aureus} analysis, innate immunity and negative regulators of cell death were induced while adaptive immunity and positive regulators of cell death were downregulated (Table 6).

In the live \textit{S. aureus} challenged cells, innate immunity, adaptive immunity and wounding process genes were induced whereas transcription of genes for normal cell processes, protein catabolism and apoptosis were suppressed by the \textit{S. aureus} infection. Similarly, Koziel and co-workers found a significant increase in the expression of antiapoptotic genes by transcriptome analysis of human monocyte-derived macrophages during \textit{S. aureus} infection.\textsuperscript{9,14,15} We also observed that in the \textit{S. aureus}-infected patients the genes for defense against bacterial infection, innate immunity, solute receptors and anti-apoptotic processes were upregulated, whereas gene expression, adaptive immunity and positive regulators of apoptosis were downregulated. In contrast, Smith et al, reported that in infected neonates, innate immune-negative feedback opposes innate immune activation while suppression of T-cell co-stimulation is coincident with selective upregulation of CD85 co-inhibitory pathways.\textsuperscript{15} Like activated \textit{S. aureus}, in heat-inactivated \textit{S. aureus} challenged cells innate and adaptive immune responses to molecules of bacterial origin, cell cycle, growth regulators, negative regulators of cell death as well as genes for blood thinning were induced. Further, among the suppressed one's genes for adaptive immunity, antibody production, antigen presentation and processing were found. However, for apoptosis process nuclear receptors for intrinsic cell death and calcium mediated signaling and apoptosis were induced. Glucose metabolism process was induced in live \textit{S. aureus} challenged cells it was suppressed by inactivated \textit{S. aureus}. Further MyD88 signaling pathway was induced by live \textit{S. aureus} infection whereas suppressed by inactivated \textit{S. aureus} treatment. Mayer et al., found that regulation of the TLR expression by host cells serves as a mechanism for adjusting sensitivity of microbial recognition in different compartments thus limiting infection. In contrast, Smith et al., found that \textit{S. aureus} infection develop an elevated set-point of myeloid regulatory signaling and sugar-lipid metabolism with concomitant inhibition of lymphoid responses in host cells.\textsuperscript{15}

\textbf{Conclusion}

Meta-analysis of microarray data from \textit{S. aureus}-challenged cells indicated that innate immune process genes were induced largely while adaptive immunity genes were suppressed. In addition, this Meta-analysis revealed cytoprotective strategies adopted by \textit{S. aureus} to evade host immune system mediated bactericidal activity. In this context, live \textit{S. aureus} and inactivated \textit{S. aureus} induced cellular processes and suppressed apoptosis genes. Similarly, apoptosis was suppressed in the infected patient's blood by upregulating anti-apoptotic genes. This review increased our understanding of the human molecular responses to infection and cytoprotective strategies adopted by the \textit{S. aureus} to evade host immune systems.

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