Differences in Host Innate Responses among Coccidioides Isolates in a Murine Model of Pulmonary Coccidioidomycosis

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Coccidioidomycosis, more commonly known as valley fever, is a fungal infection caused by inhaling either Coccidioides immitis or Coccidioides posadasii conidia, which are found in the soil in arid to semiarid regions of the Americas. The most common route of infection is by inhalation of airborne Coccidioides arthroconidia. Once a susceptible host inhales the conidia, a transition to mature endosporulated spherules can occur within the first 5 days of infection. For this study, we examined the host response in a murine model of coccidioidomycosis during a time period of infection that has not been well characterized. We collected lung tissue and bronchoalveolar lavage fluid (BALF) from BALB/c mice that were infected with a C. immitis pure strain, a C. immitis hybrid strain, or a C. posadasii strain as well as uninfected mice. We compared the host responses to the Coccidioides strains used in this study by assessing the level of transcription of selected cytokine genes in lung tissues and characterized host and fungal proteins present in BALF. Host response varied depending on the Coccidioides strain that was used and did not appear to be overly robust. This study provides a foundation to begin to dissect the host immune response early in infection, to detect abundant Coccidioides proteins, and to develop diagnostics that target these early time points of infection.

Coccidioidomycosis endemic to certain semiarid regions in the Americas. The most common route of infection is by inhalation of airborne Coccidioides arthroconidia. Once a susceptible host inhales the conidia, a transition to mature endosporulated spherules can occur within the first 5 days of infection. For this study, we examined the host response in a murine model of coccidioidomycosis during a time period of infection that has not been well characterized. We collected lung tissue and bronchoalveolar lavage fluid (BALF) from BALB/c mice that were infected with a C. immitis pure strain, a C. immitis hybrid strain, or a C. posadasii strain as well as uninfected mice. We compared the host responses to the Coccidioides strains used in this study by assessing the level of transcription of selected cytokine genes in lung tissues and characterized host and fungal proteins present in BALF. Host response varied depending on the Coccidioides strain that was used and did not appear to be overly robust. This study provides a foundation to begin to dissect the host immune response early in infection, to detect abundant Coccidioides proteins, and to develop diagnostics that target these early time points of infection.

Received 4 August 2015; Accepted 10 August 2015; Accepted manuscript posted online 14 August 2015

Citation: Lewis ERG, David VR, Doyle AI, Rajabi K, Kiefer JA, Pirrotte P, Barker BM. 2015. Differences in host innate responses among Coccidioides isolates in a murine model of pulmonary coccidioidomycosis. Eukaryot Cell 14:1043–1053. doi:10.1128/EC.00122-15

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Supplemental material for this article may be found at http://dx.doi.org/10.1128/EC.00122-15.

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doi:10.1128/EC.00122-15
on days 7 and 14 postinfection indicated nonprotective IL-10 and IL-4 responses to strain RS Coccidioides infections in susceptible C57BL/6 mice (8).

The cytokine response to challenge infections of other human fungal respiratory pathogens has also been examined in murine models of disease. For example, increased transcription of genes that encode IL-2, IL-4, IL-12, and IFN-γ was detected as early as day 3 postinfection in the lungs of C57BL/6 mice challenged with Histoplasma capsulatum (9). IFN-γ is necessary for control of H. capsulatum infection in mice (10). Elevated levels of TNF-α, IL-6, IL-1β, and macrophage inflammatory protein (MIP) were detected during the first 4 days postinfection in BALB/c mice infected with Paracoccidioides brasiliensis (11). The cytokines TNF-α and IL-1α have been shown to be important for leukocyte recruitment and control of infection of the pathogen Aspergillus fumigatus in mice (12, 13). Examination of Cryptococcus neoformans in a C57BL/6 mouse model of infection showed that IL-17α increase is mediated by leukocyte recruitment and activation and IFN-γ production after intratracheal challenge (14).

The goal of this study was to expand upon previous studies of the host innate immune response to pulmonary coccidioidomycosis by examining the first 5 days of infection in a BALB/c mouse model of infection, when the morphological shift from the saprobic to parasitic cycle occurs. Understanding the cytokine expression profile at the onset of infection could reveal new insights into infection dynamics of Coccidioides. For this study, we analyzed the immune responses of mice intranasally infected with arthroconidia from clinical isolates of Coccidioides: a C. immitis pure isolate (2006), a C. immitis hybrid isolate (RS), and a C. posadaij pure isolate (Silveira). The expression profiles of selected cytokines were assessed by real-time reverse transcription-PCR (RT-PCR) of RNA extracted from the mouse lung tissues and a multiplex bead array for proteins present in BALF. Finally, protein expression profiles from BALF during day 5 postinfection were determined using mass spectrometry to identify potential host biomarkers and fungal proteins expressed during pulmonary coccidioidomycosis.

**MATERIALS AND METHODS**

**Mouse inoculations.** Coccidioides immitis isolate 2006 (15), C. immitis hybrid isolate RS (16), and C. posadaij isolate Silveira (17, 18) were used in this study. Female 6- to 8-week-old BALB/c mice were anesthetized with ketamine-xylene and intranasally inoculated with 100,000 arthroconidia suspended in 30 μl phosphate-buffered saline (PBS) as described previously (19). Control mice were inoculated with PBS alone. The mice were housed according to NIH guidelines in a biosafety level 3 animal laboratory. All procedures were approved by the Institutional Animal Care and Use Committee for the University of Arizona. Three mice from each infection group (2006, RS, and Silveira) and two uninfected PBS-treated mice were sacrificed on day 1 through day 5. The lungs were rinsed with 2 ml of phosphate-buffered saline with 0.01 mM EDTA to collect BALF. The right lobe of the lung was harvested and flash frozen in liquid nitrogen. One milliliter of each BALF sample was formalin fixed and underwent a complete cell count using a hemocytometer. The remainder of each BALF sample was filtered using a 0.8-μm Millex-AA syringe filter unit (EMD Millipore, Billerica, MA), and a Thermo Scientific Pierce bicinechonic acid (BCA) protein assay (Life Technologies, Carlsbad, CA) was used to quantify protein concentrations.

**RNA extraction.** Modifications to a previously described protocol were applied to the RNA extraction procedure (20). The whole right lungs were lyophilized for 48 h. The lyophilized lungs were put in 2-ml tubes with lysing matrix D (MP Biomedicals, Santa Ana, CA) ceramic spheres and mixed with 1 ml of TRIzol (Bioline USA, Inc., Taunton, MA). Samples were subjected to bead beating for 2 min and then incubated for an additional 5 min. Chloroform (200 μl) was added, mixed well, and incubated 2 min at room temperature. Samples were centrifuged at maximum speed for 15 min at 4°C. The upper layers were collected and mixed with an equal volume of 80% ethanol (EtOH), and the entire volume was pipetted immediately to RNase spin columns (Qiagen, Inc., Valencia, CA). Columns were centrifuged for 1 min, the flowthrough was discarded and washed twice with kit-supplied RPE buffer (Qiagen), and columns were completely dried after the last wash. The spin columns were then transferred to new RNase-free 1.5-ml tubes, 200 μl of RNase free water was added to each spin column, and columns were incubated for 1 min. After that 1-min incubation, the tubes containing the spin columns were centrifuged to obtain nucleic acid. Samples were analyzed with NanoDrop ND-1000 (Thermo Fisher Scientific, Inc., Waltham, MA) and a 2100 Bioanalyzer (Agilent Technologies, Inc., Santa Clara, CA).

**Real-time RT-PCR.** The extracted RNA was treated with an Ambion Turbo DNA-free kit (Life Technologies Corporation, Carlsbad, CA) to remove genomic DNA. The RNA concentration was checked on a NanoDrop ND-1000 (Thermo Fisher Scientific, Inc., Waltham, MA). Conditions for real-time RT-PCR were 1 min of polymerase activation at 95°C followed by 40 cycles of 5 s at 95°C, 10 s at 60°C, and 20 s at 72°C. To determine fungal burden, oligomers designed to amplify the Coccidioides glyceraldehyde-3-phosphate dehydrogenase gene (gapdh) were used in a real-time RT-PCR assay (see Table S1 in the supplemental material). To examine cytokine gene expression, PrimeTime qPCR primer assays (Integrated DNA Technologies, Inc., Coralville, IA) of selected genes (see Table S1 in the supplemental material) were used for this study. Change (n-fold) was determined using the 2^ΔΔCT method (21). The mouse gapdh gene was selected as the internal control gene to calculate the relative expression of the tested mouse genes.

**Multiplex bead array assay.** The concentrations of selected cytokines (see Table S2 in the supplemental material) for the BALF were examined using a cytokine mouse magnetic 20-Plex panel (Life Technologies, Carlsbad, CA) on a Luminex LX200 system with Luminex xPONENT 3.0 software (Luminex Corporation, Austin, TX). A Thermo Scientific Pierce BCA protein assay (Life Technologies, Carlsbad, CA) was used to quantify protein concentrations. Cytokine concentrations were normalized to the amount of total protein quantified in each sample.

**Proteomic analysis.** Equal amounts of protein (20 μg each) were reduced and denatured by boiling for 10 min in loading buffer (Bio-Rad Laboratories, Inc., Hercules, CA) containing 5% β-mercaptoethanol. Each sample was then separated by gel electrophoresis on a 4 to 20% SDS-PAGE gel (Bio-Rad Laboratories, Inc., Hercules, CA) for 45 min at 120 V and stained using Coomassie Plus (Bio-Rad Laboratories, Inc., Hercules, CA). Gel lanes were fully excised into individual bands, destained, washed, dried, and further processed using a previously published method (22). Briefly, each lane fraction was reduced and alkylated using cyclic rehydration/dehydration of the acrylamide gel slabs. Samples were then digested using Trypsin Gold (Promega Corporation, Madison, WI) at 20 ng/ml overnight at 37°C. Peptides were extracted, concentrated to dryness under vacuum, and frozen at −20°C prior to mass spectrometry analysis.

Analyses were conducted on a nanoAcquity ultra performance liquid chromatography (UPLC) system (Waters Corporation, Milford, MA) coupled to a Thermo LTQ Orbitrap Velos mass spectrometer (Thermo Scientific, Waltham, MA). Each sample fraction was reconstituted in 0.1% formic acid for analysis by using online liquid chromatography coupled to mass spectrometry. Each fraction was reconstituted in 0.1% formic acid and loaded onto a 100-μm-diameter column packed with 100 μm with 3 μm Reprosil Pur C18 AQ resin eluted at 500 nl/minute. Solvents A and B
were 0.1% formic acid in water and acetonitrile, respectively. The gradient was 3% B to 40% B in 17 min followed by 40% B to 90% B in 0.5 min, then 90% B for 2 min, and re-equilibration for 10.5 min. The mass spectrometer operated in positive ion mode using a spray voltage of 1.8 kV and a capillary temperature of 200°C. Data were acquired in top-15, data-dependent acquisition mode using a collision voltage of 30 V. Raw mass spectrometry data were searched against a concatenated database from UniprotKB/Swissprot for *Mus musculus* and UniprotKB/Swissprot + Trembl for *C. posadasii* isolate Silveira and *C. immitis* isolate RS using Mascot (version 1.4.1.14; Matrix Science, London, United Kingdom) and X! Tandem (v2010.12.01.1; The GPM). Tandem mass spectra were extracted, and charge state was deconvoluted and deisotoped by Proteome Discoverer 1.4.1.14 (Thermo Fisher Scientific, Waltham, MA). Oxidation (Met), carbamidomethylation (Cys), and pyro-Glu formation (N-terminal Gln or Glu) were specified as variable modifications. A peptide mass tolerance of ±10 ppm, a fragment mass tolerance of ±0.8 Da, and a maximum of two trypsin missed cleavages were allowed. Peptide identifications were accepted at ≥98.0% as specified by the Peptide Prophet algorithm (23) and corrected using Scaffold m correction (Proteome Software Inc., Portland, OR).

**Biological concept enrichment analysis.** Biological concept enrichment analysis was performed on the four sets of proteins using the ClueGO v2.1.7 Cytoscape plugin (24). Enrichment was performed with the following Ontologies/Pathway gene sets: GO Biological Process, KEGG, Reactome, and WikiPathways. Advanced Term/Pathway selection options were set at Go Tree interval minimum level 3 and maximum level 8, GO term fusion, and minimum number of genes at five for uninfected and 10 for 2006, RS, and Silveira infected. The kappa score was set at 0.4. A two-sided hypergeometric test was used with Bonferroni step-down correction.

**ToppGene functional analysis.** We used the ToppGene functional analysis suite (25) to identify enriched terms in each of our four sets of proteins (uninfected and 2006, RS, and Silveira infected) (false discovery rate [FDR] correction; \( P < 0.05 \); gene limits, \( 1 \leq n \leq 2,000 \)). Enriched terms from the biological process ontology were ranked by \( P \) value, and only the five most significant terms were retained for interpretation of the host response to *Coccidioides* infection in BALF.

**Statistics.** Statistical analyses were performed using GraphPad Prism version 6.00 for Mac (GraphPad Software, La Jolla, CA) or Scaffold (Proteome Software Inc., Portland, OR) for mass spectrometry-derived data. \( P \) values were calculated either by using a two-way analysis of variance (ANOVA) with Tukey's posttest or Kruskal-Wallis test in the case of the proteomics data. A \( P \) value of \(< 0.05\) was considered statistically significant.

**RESULTS**

**BALB/c mouse infections.** Three mice from each infection group and two mice from the uninfected group were sacrificed on each day postinfection during this 5-day study. Real-time RT-PCR was used to detect *Coccidioides gapdh* mRNA extracted from the whole right lungs of each mouse to determine a relative fungal burden (Fig. 1A). *C. posadasii* isolate Silveira *gapdh* mRNA transcripts were detected on day 3 of infection and beyond (Fig. 1A), whereas *C. immitis* hybrid isolate RS *gapdh* mRNA transcripts were not detected until day 4 of infection (Fig. 1A). *C. immitis* isolate 2006 *gapdh* mRNA transcripts were not detected until day 5 of infection (Fig. 1A). No *Coccidioides gapdh* mRNA transcripts were detected in the uninfected mice (Fig. 1A).

A total leukocyte count was made from cells present in the BALF to confirm that the mice mounted an inflammatory response to infection (Fig. 1B; Table 1). Non-statistically significant increases were observed for all infection groups compared to the uninfected mice on the first day of infection (Fig. 1B). On day 2 of infection, the 2006 and RS infection groups had non-statistically significant increases of total leukocyte counts compared to the Silveira infection group and uninfected control group (Fig. 1B). On day 3 of infection, all three infection groups had leukocyte counts comparable to that of the uninfected control group. On day 4, however, the Silveira infection group had statistically significant higher number of total leukocytes than the 2006 infection group (\( P \leq 0.001 \)), RS infection group (\( P \leq 0.001 \)), and uninfected control group (\( P \approx 0.01 \)). On day 5, all infection groups had a higher number of leukocytes per ml of BALF than uninfected mice. In agreement with days 1, 3, and 4, the Silveira infection group had the highest number of leukocytes per ml of BALF on average. During necropsy, we observed that mice infected with *C. posadasii* isolate Silveira suffered from more severe lung damage starting at day 3 of infection than the mice infected with *C. immitis* isolate RS and *C. immitis* isolate 2006 (data not shown).

**Real-time RT-PCR analysis of mouse lung tissues.** The whole right lungs were harvested from each mouse and underwent real-time RT-PCR analysis to assess transcript levels of pro- and anti-inflammatory cytokines. The cytokine targets and primers used for this study are listed in Table S1 in the supplemental material. We detected a differential cytokine response among the *Coccidioides* isolates used in this study. Only the Silveira infection group showed a 20-fold increase in the proinflammatory cytokine gene *il-1β* (Fig. 2). A temporal trend toward an increase in levels of the proinflammatory cytokine gene *tnfα* varied among the infection
groups during the study (Fig. 2). There were also trends that all infection groups shared. On day 1, there was an initial increase of \textit{ifn\_y1} and \textit{il\_1r1} for all infection groups, with a significant decrease on days 4 and 5, respectively (Fig. 2). There were transcriptional increases for many of the cytokine genes, including the anti-inflammatory cytokine gene \textit{il-10} and the proinflammatory cytokine gene \textit{il-1\_a}, analyzed on day 5 of infection for all infection groups. Similar increases were seen in the expression of the proinflammatory cytokine gene \textit{il-17a} on day 5 for the mice infected with 2006 or Silveira (Fig. 2). No statistically significant increases were noticed for \textit{il-17a} before day 5 of infection (Fig. 2). We did observe non-statistically significant increases in changes for other transcripts examined, including \textit{il-2} and \textit{il-4}. Real-time RT-PCR results indicate a differential cytokine response dependent on which \textit{Coccidioides} isolate was used for infection.

**Multiplex bead array analysis of mouse BALF.** The 20 cytokines that were quantified are listed in Table S2 in the supplemental material. Few were detected above the assay’s lower limit, as only IFN-\(\gamma\) and IL-2 were detected in the BALF using this method (Fig. 3). The average values of protein concentration for IFN-\(\gamma\) and IL-2 in uninfected control animals are presented in Fig. 3. During the 5 days of infection, day 1 appeared to be the only day in which IFN-\(\gamma\) was at a higher, but not a statistically significantly higher, concentration in the 2006 group than in the uninfected group (Fig. 3A). On day 1 of infection, a non-statistically significant increase of IL-2 was present in the BALF of the 2006 and Silveira infection groups (Fig. 3B). Overall, we did not detect an abundant cytokine response.

**Proteomic analysis of mouse BALF.** Because we did not detect abundant cytokines in our samples, we employed proteomics analysis on remaining BALF. Overall, total protein concentrations remained below 100 \(\mu\text{g}/m\text{l}\) in all of the mouse groups until day 3 of infection (Fig. 4). On day 3 of infection, the Silveira-infected mice reached a mean total protein concentration of 300 \(\mu\text{g}/m\text{l}\), while other mouse groups remained below 100 \(\mu\text{g}/m\text{l}\) (Fig. 4). A marked increase in the mean total protein concentration within the BALF of the Silveira infection group on day 4 was statistically significant compared to the 2006 infection group \((P \leq 0.0001)\), RS infection group \((P \leq 0.0001)\), and uninfected mouse group \((P \leq 0.0001)\) (Fig. 4). On day 5, all infection groups had increased levels of total protein in BALF compared to the uninfected mouse group (Fig. 4). As the highest protein concentrations were observed at day 5, we performed an extensive downstream proteomic analysis on these samples.

We performed proteomic analysis on the BALF collected on day 5 of the experiment from the nine infected mice (three mice per infection group) and the two mice in the uninfected group. A total of 374 unique proteins were identified from the proteome of all BALF collected on day 5 of this experiment (Fig. 5; also, see Data Set S1 in the supplemental material). This number includes 566 unique proteins identified overall, as well as eight \textit{Coccidioides} proteins identified among the infected mice (Fig. 5). Data Set S1 lists the proteins identified in the BALF samples. Of the proteins that we identified, 33 mouse proteins displayed statistically significant differences between the four mouse groups examined in this study (see Data Set S1). Six mouse proteins were significantly differentially abundant in uninfected mice compared to all of the mouse infection groups (Table 2). Two of these, annexin A5 and aminopeptidase N, were increased in all uninfected compared to infected mice. Aminopeptidase N extracellular signaling
helps to reduce inflammation, and annexin A5 is a known inhibitor of protein kinase C, thus inhibiting apoptosis and degranulation in healthy mice (26, 27). This suggests activation of cytotoxic activity from mast cells and granulocytes and activation of inflammation and apoptosis in all infected mice. Four of the proteins, α-2-macroglobulin, complement C3, hemopexin, and proteasome subunit, were lower in uninfected mice, and these are associated with general lung damage and inflammation common in all the infected mice (28–31). Twelve proteins were significantly differentially expressed for mice infected with RS compared to the other mouse groups (Table 3). Interestingly, 10 of the 12 were more abundant, including chitinase and iron/heme-associated proteins. For the proteins that were lower in abundance, fibronectin and inter-α-trypsin inhibitor are both biomarkers associated with chronic obstructive pulmonary disease (COPD) and inflammatory processes when found in abundance, suggesting that RS does not induce inflammation via this mechanism (31, 32). Ten significantly differentially expressed proteins were found in mice infected with Silveira compared to the other groups (Table 4). Of the four that are less abundant, carbonyl reductase (NADPH) 2 suggest a nonprotective response to oxidative stress, whereas complement C5 and lactotransferrin suggest a lack of protective inflammatory response (29, 33–35). For proteins that were more highly expressed, all are associated with damage response, supporting the observation that Silveira causes significant damage to the lung. Interestingly, no unique proteins were differentially abundant for mice infected with isolate 2006 (see Data Set S1).

In order to determine representative biology associated with the four sets of proteins, ClueGO was used to perform biological concept enrichment analysis. ClueGO visualizes the functional enrichment results by grouping similar ontologies and networks into overlapping clusters based on similar function and gene membership. The four protein sets were analyzed with ClueGO using the GO Gene Ontology (GO) Biological Process, Kyoto Encyclopedia of Genes and Genomes (KEGG), Reactome, and

FIG 2 Real-time RT-PCR analysis of cytokine gene mRNA transcripts in the whole right lungs of BALB/c mice. Error bars indicate standard errors from biological triplicates. $P$ values were calculated using two-way ANOVA followed by Tukey’s multiple-comparison test. A $P$ value of <0.05 was considered statistically significant, and levels of significance were assigned as follows: *, $P \leq 0.05$; **, $P \leq 0.01$; ***, $P \leq 0.001$; ****, $P \leq 0.0001$. 

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October 2015 Volume 14 Number 10  Eukaryotic Cell
WikiPathways gene set categories. The enrichment network is visualized in Fig. 6. The red nodes represent ontologies or pathways that show preferential enrichment in the Silveira strain. The blue nodes are those enriched in the RS group of proteins. Interestingly, no category had preferential enrichment for proteins in the 2006 or uninfected protein set. The gray nodes did not have a preferential enrichment in any of the protein sets and represent common terms in all protein sets. The significant ontology/pathway nodes for the Silveira infection group protein set were categories such as innate immune system, proteolysis, cellular response to stress, lipid transport, EPH-ephrin signaling, and actin cytoskeleton organization (Fig. 6). The protein set from the RS infection group had significant node clusters, such as metabolism, phagosome, complement, and coagulation cascades, as well as glutathione metabolism, among others (Fig. 6). A tight cluster of nodes is centered on signaling pathways with a large component of proteasome-related genes (Fig. 6). A cluster of proteasome genes common across all protein sets drive enrichment in this cluster of pathways (Fig. 6).

ToppGene enrichment analysis was also performed to confirm the results of the protein clustering analysis (see Fig. S1 to S4 and Data Set S2 in the supplemental material). The host proteins identified align with the increase in \textit{Coccidioides} isolate fungal burden and infection dynamics (2006/RS/Silveira) we observed in other assays (Fig. 1 and 4). Total proteins assigned to these terms were notably lower in uninfected mice, possibly as a result of a baseline defense response to challenges at the endothelial interface between alveoli and capillaries (see Fig. S4 and Data Set S2). Key terms associated with inflammation were present in all \textit{Coccidioides} isolate groups (see Fig. S1 to S3 and Data Set S2). As expected, proteins associated with both innate and humoral re-

![FIG 3](image)

Concentration of cytokines IFN-γ (A) and IL-2 (B) in BALF. Cytokine concentrations were measured using a multiplex bead array and normalized to the amount of total protein quantified in each sample. The dashed line represents the mean for uninfected control animals. Error bars indicate standard errors from biological triplicates. \textit{P} values were calculated using two-way ANOVA followed by Tukey’s multiple-comparison test. A \textit{P} value of \(<0.05 (\text{*})\) was considered statistically significant.

![FIG 4](image)

Quantifying the concentration (µg/ml) of total proteins present in BALF for downstream proteomics analysis. Error bars indicate standard errors from biological replicates (\(n = 3\) for each infection group; \(n = 2\) for the uninfected group), \(P\) values were calculated using two-way ANOVA followed by Tukey’s multiple-comparison test (\(\text{*, } P \leq 0.05; \text{****, } P \leq 0.0001\)).

![FIG 5](image)

Venn diagram comparing and contrasting the number of unique proteins identified in the BALF collected from the day 5 groups using mass spectrometry analysis. A total of 374 unique proteins composed of 366 unique mouse proteins and eight unique \textit{Coccidioides} proteins were identified. In parentheses are the numbers of mouse proteins that were identified only in a particular group.

![TABLE 2](image)

| Protein                | Gene ID     | Abundance |
|------------------------|-------------|-----------|
| α-2-Macroglobulin      | A2M_MOUSE   | Less      |
| Aminopeptidase N       | AMPN_MOUSE  | More      |
| Annexin A5             | ANXA5_MOUSE | More      |
| Complement C3          | CO3_MOUSE   | Less      |
| Hemopexin              | HEMO_MOUSE  | Less      |
| Proteasome subunit beta type 6 | PSB6_MOUSE | Less      |
responses were prioritized for RS and Silveira (defense, inflammatory, and acute inflammatory responses). Proteins secreted in response to wounding were significant in all infection groups (see Fig. S1 to S3 and Data Set S2), confirming our findings on probable lung damage and tissue necrosis (Fig. 4). We detected an increase of platelet degranulation-associated proteins in the 2006 mouse infection group that was not as apparent in any of the other infection groups or uninfected control mice (see Fig. S1 and Data Set S2). Platelet degranulation is an antimicrobial host response to infection that has been observed for other human pathogens, including the fungus *Aspergillus fumigatus* and multiple bacterial species (36, 37). The high fungal burden of Silveira isolate also likely led to an increase in proteolytic host response (see Fig. S3 and Data Set S2).

**Coccidioides** proteins identified in BALF. We also identified *Coccidioides* proteins present in BALF (Table 3). Three proteins, a *Coccidioides* serine protease (CIMG_10287/CPSG_04717), a triosephosphate isomerase (CIMG_09361/CPSG_03911), and an uncharacterized protein (CIMG_09001/CPSG_01366), were identified in the BALF collected from all infection groups (Table 5). The *Coccidioides* serine protease we identified in all of the mouse infection groups is a member of an expanded subtilisin *N. ioides* implicated in host interactions but is currently uncharacterized (38). Orthologous serine proteases have been shown to be virulence factors expressed during infection by other human-pathogenic fungal and bacterial species (38).

Other *Coccidioides* proteins were found in only one or two infection groups (Table 5). As an example, endochitinase 1 (CIMG_02795/CPSG_08657) was identified from one mouse infected with RS and all three of the mice infected with Silveira (Table 5). It is a member of a well-characterized family of chitinase proteins in *Coccidioides* that have been shown to be upregulated during the parasitic life cycle (39, 40). CTS1 is also immunogenic and is used as a serodiagnostic antigen for disease (41).

A peroxisomal matrix protein (PMP1) (CIMG_05828/CPSG_04764) was identified in two of the mice infected with RS. A recombinant PMP1 was found previously to be reactive with serum from individuals with both acute and protracted coccidioidomycosis (42). The potential of this protein as a recombinant vaccine candidate was also examined; it evoked protection in two murine models of infection with *C. posadasii* (42). Thus, our approach was validated by the identification of previously characterized *Coccidioides* proteins that are known to be associated with the parasitic life cycle.

**DISCUSSION**

For this study, we sought to examine and characterize the early host response to *Coccidioides* infection in a BALB/c mouse model of pulmonary coccidioidomycosis at time points not previously characterized. Our focus on BALF was driven by the knowledge that this is a common clinical specimen and that current diagnostics are inadequate for detecting *Coccidioides* (43). We examined the host response to three different isolates of *Coccidioides* to assess whether the host responses to the isolates would be similar. We initially anticipated that the three *Coccidioides* species isolates would have comparable overall inflammatory responses and infection kinetics. In contrast to our expectation, there appeared to be significant differential responses among the isolates used in this study. We discovered that of the three *Coccidioides* isolates that were used in this study, the *C. posadasii* isolate Silveira appeared to cause the highest relative fungal burden in mouse lung tissue. This isolate also caused a significantly higher influx of leukocytes as early as day 4 of infection that was not observed in any of the other infection groups on that day. By day 5 of infection, a significant increase in leukocytes was present for all infection groups. Although we do not yet understand the mechanism of pathogenesis and increased fungal growth in the host, the Silveira isolate has been reported to be highly virulent in mice (17, 44).

A differential host response to the three *Coccidioides* isolates was also observed when host cytokine gene mRNA levels in lung tissue were examined. We discovered a differential and diverse cytokine gene expression profile associated with each *Coccidioides* isolate. Previous studies have shown that some cytokines, including IFN-γ, can be protective against *Coccidioides* infection in mice, whereas other cytokines, including IL-4 and IL-10, are not protective against infection (6, 7). Other cytokines, such as TNF-α, IL-1α, and IL-17α, have been shown to be protective in murine models of other mycoses (12–14). The differential cytokine responses involving both probable protective and nonprotective cytokines we observed in our study may associate with the inability of BALB/c mice to respond to *Coccidioides* infections. We detected few cytokines in the BALF using the multiplex bead array assay.

We quantified various amounts of total protein in BALF, with day 5 having the overall largest amounts of total protein present in infected mice compared to uninfected mice. Increases in total protein concentration and leukocytes are markers for lung injury with
other infections (45–47). By these assessments, Silveira appeared to be the most virulent isolate used in this study. Our observation of increasing levels of the total protein during the course of this study was probably due in part to lung damage caused by infection. Because few cytokines were detected, proteomic analysis on the samples collected on day 5 of infection allowed determination of other proteins present. We found differences in the proteomic profile that suggest that lung damage was not the only cause of increased protein concentration. Noninterleukin proteins involved in inflammation, immune signaling, and
differentiation.

**TABLE 5** *Coccidioides* proteins that were detected in the BALF collected from infected mice on day 5 of infection

| Protein                                      | Gene ORF\(^a\)                      | *Coccidioides* isolate\(^b\) (n°) |
|----------------------------------------------|-------------------------------------|-----------------------------------|
| 40S ribosomal protein S14                    | CIMG\_04348/CPSG\_09614             | RS (1)                            |
| 60S ribosomal protein L12                    | CIMG\_04811/CPSG\_07687             | Sil (2)                           |
| Endochitinase 1                               | CIMG\_02795/CPSG\_08657             | RS (1), Sil (3)                    |
| Gamma-glutamyltranspeptidase                 | CIMG\_05765/CPSG\_02828             | Sil (2)                           |
| Peroxisomal matrix protein                   | CIMG\_05828/CPSG\_04764             | RS (2)                            |
| Serine protease                              | CIMG\_10287/CPSG\_04717             | 2006 (3), RS (3), Sil (2)          |
| Triosephosphate isomerase                    | CIMG\_09361/CPSG\_03911             | 2006 (1), RS (1), Sil (1)          |
| Uncharacterized protein                      | CIMG\_09001/CPSG\_01366             | 2006 (2), RS (1), Sil (3)          |

\(^a\) Gene open reading frame (ORF) name for orthologous loci in both species, in the format *C. immitis* isolate RS/*C. posadasii* isolate Silveira.

\(^b\) 2006, *C. immitis* isolate 2006; RS, *C. immitis* isolate RS; Sil, *C. posadasii* Silveira.

\(^c\) Number of mice per infection group in which this protein was identified.
host defense response: proteins varied in abundance for all infection groups compared with uninfected mice. Similar to the cytokine response to infection, the proteome of the BALF varied depending on which isolate was used for infection. The observations that were made in this study will set the foundation for future studies characterizing the host response to different Coccidioides isolates. As well, some proteins and specific host response pathways may be useful host biomarkers for pulmonary coccidioidomycosis.

In addition to finding differential host protein expression profiles, we also identified at least three Coccidioides proteins that have been shown to be associated with the Coccidioides parasitic life cycle by other research groups, confirming that our method of proteomic analysis was able to detect expected proteins. Some of the Coccidioides proteins that we identified are uncharacterized proteins, including a protein (CIMG_09001/CPShG_01366) that was identified in all infection groups. Proteins associated with infections caused by both Coccidioides species are good candidates for further study, as these novel proteins may be necessary for the initiation of the parasitic life cycle and therefore potential targets for diagnostics and/or therapeutics.

Many Coccidioides isolates tested to date cause fatal disease within eight to 12 days with as few as 50 arthroconidia administered intranasally in immunocompetent BALB/c mice (17, 48). We administered 100,000 arthroconidia intranasally to each group and could have potentially overwhelmed the host innate immune response. The data suggest that this was not the case and that this dose is reasonable when dosages administered for other fungal pathogens are considered (49–51). As discussed above, we observed a differential cytokine response to Coccidioides isolates used in the study, and the early innate immune response did not seem overly robust. With regard to the infectious dose in humans, neither the average nor the minimum dosage is known (52). The dose administered to the mice in our study is biologically relevant, as a host could inhale 100,000 or more arthroconidia during an acute environmental exposure. In future studies, it will be worthwhile to assess host cytokine response in a dosage-dependent manner.

The lack of a strong cytokine response to early pulmonary coccidioidomycosis described in this study points to previous studies suggesting host immune system evasion. Circulating Coccidioides antigens have been shown to suppress cell-mediated immune response in BALB/c mice (6, 53, 54). More recently, the virulence factors spherule outer wall glycoprotein (SOWgp) and metalloproteinase 1 (Mep1) were identified and shown to evade the host immune response in a C57BL/6 mouse model of coccidioidomycosis (55–59). The most successful pathogens utilize one or several mechanisms to evade and/or counteract a protective immune response from the host (60–62). Because we did not observe these previously identified protein products among our data, we propose that Coccidioides may utilize additional host immune evasion mechanisms, and these may differ temporally as well as among isolates.

Previous studies compared the virulence of different Coccidioides isolates in a mouse model of infection and found differential virulence among the isolates tested (17, 44). To our knowledge, this is the first study that compares the early cytokine response of BALB/c mice to multiple Coccidioides isolates. All isolates that were used in this study are human clinical isolates with known infectivity. Our data suggest that BALB/c mice have a differential cytokine response dependent on the isolate used for infection. At this time, no differential disease phenotypes have been described for coccidioidomycosis caused by C. immitis compared to C. posadasii. This study does not provide enough evidence to support or negate that claim. However, our findings support the importance of performing additional comparative studies of clinical and even environmental isolates of Coccidioides species. Genomic sequencing has thus far revealed high levels of genetic diversity within and among species of Coccidioides (63). Performing comparative functional studies on a variety of these sequenced isolates could help define the genotypic and phenotypic diversity of the organisms and their relationship to clinical outcomes. As well, gaining a better understanding of the host innate immune response may lead to the development of more sensitive and specific diagnostic tools and effective treatment options.

ACKNOWLEDGMENTS

This work was supported by grant K22 AI104801 from the National Institute of Allergy and Infectious Diseases (B.M.B.), an Arizona Biomedical Research Corporation young investigator grant (B.M.B.), and Translational Genomics Research Institute (TGen) startup funds (B.M.B. and P.P.).

We truly appreciate the assistance provided by John Galgiani, Lisa Shubitiz, Hien Trinh, and Maria Lourdes Lewis with the mouse experiments and access to ABSL3 for this study. We thank Kylie Sage and Heather Mead for assistance with real-time RT-PCR experiments.

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