Proteome and Antigen Profiling of *Coxiella burnetii* Developmental Forms

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A biphasic developmental cycle whereby highly resistant small-cell variants (SCVs) are generated from large-cell variants (LCVs) is considered fundamental to the virulence of *Coxiella burnetii*, the causative agent of human Q fever. In this study a proteome analysis of *C. burnetii* developmental forms was conducted to provide insight into their unique biological and immunological properties. Silver-stained gels of SCV and LCV lysates separated by two-dimensional (2-D) gel electrophoresis resolved over 675 proteins in both developmental forms. Forty-eight proteins were greater than twofold more abundant in SCVs than in LCVs, with six proteins greater than twofold more abundant in SCVs than in LCVs. Four and 15 upregulated proteins of SCVs and LCVs, respectively, were identified by mass spectrometry, and their predicted functional roles are consistent with a metabolically active LCV and a structurally resistant SCV. One-dimensional and 2-D immunoblots of cell form lysates probed with sera from infected/vaccinated guinea pigs and convalescent-phase serum from human patients who had recovered from acute Q fever, respectively, revealed both unique SCV/LCV antigens and common SCV/LCV antigens that were often differentially synthesized. Antigens recognized during human infection were identified by mass spectroscopy and included both previously described immunodominant proteins of *C. burnetii* and novel immunogenic proteins that may be important in the pathophysiology of clinical Q fever and/or the induction of protective immunity.

*Coxiella burnetii* is the etiologic agent of an acute, disabling influenza-like illness in humans termed Q fever. In rare instances chronic disease can occur, usually presenting as hepatitis or endocarditis. *C. burnetii* infects a wide range of vertebrate and invertebrate hosts and is found worldwide with the exception of New Zealand. Most human Q fever cases are acquired through inhalation of contaminated aerosols originating from products shed from chronically infected domestic livestock. Indeed, massive numbers of *coxiiella* (>10^10/g) are present in the products of parturition (reviewed in reference 19).

*C. burnetii* has an obligate requirement for replication in a eukaryotic host cell vacuole with lysosomal characteristics (15). Resistance to this harsh intracellular environment correlates with remarkable environmental stability, a trait that distinguishes *C. burnetii* from other obligate intracellular bacteria. The pathogen is highly resistant to osmotic shock, elevated temperature, desiccation, UV light, and various chemical disinfectants (21, 36). Environmental stability, along with aerosol transmission and an infectious dose approaching one organism, has resulted in classification of *C. burnetii* as a CDC category B biothreat. The spore-like characteristics of *C. burnetii* are attributed to a stable small-cell variant (SCV) that arises as part of a *C. burnetii* biphasic developmental cycle (22). SCVs are rod shaped and 0.2 to 0.5 μm in length and have an electron-dense condensed chromatin with an array of intracytoplasmic membranes. Large-cell variants (LCVs) can exceed 1 μm in length and are pleomorphic with a dispersed chromat in (12, 22). *C. burnetii* preparations enriched in LCVs are more metabolically active in vitro than those enriched in SCVs (21).

By infecting Vero cells with SCVs, we have recently established the kinetics of *C. burnetii* morphological development and defined the LCV as the replicative form of the pathogen (5). The *C. burnetii* growth cycle has defined lag, log, and stationary phases (5). SCV-to-LCV morphogenesis occurs during a lag phase lasting approximately 2 days. Exponential replication of LCVs ensues over the next 4 days with LCVs condensing to SCVs in significant numbers during a stationary phase that starts approximately 6 days postinfection (p.i.).

Elucidation of the molecular events that regulate *C. burnetii* development has been hampered by an inability to genetically manipulate the organism; however, some progress has been made in identifying developmental-form-specific proteins. McCaul et al. (20) identified a 29.5-kDa immunogenic protein termed P1 that is differentially synthesized by the LCV. The encoding gene was later cloned by Varghees et al. (33), who described P1 as having porin activity. Additional proteins that are dramatically enriched in the LCV include EF-Tu and EF-Ts (26) and RpoS (28). Two DNA-binding proteins, ScvA and Hq1, are preferentially expressed by the SCV (13, 14).

Intracellular morphological differentiation to generate environmentally stable SCV and replicatively active LCV develop-
mental forms is considered central to Coxiella virulence. However, the unique or upregulated proteins that confer the distinctive ultrastructural and biological characteristics of SCVs and LCVs are still largely unknown, as are the immunogenic constituents of developmental forms that are recognized during infection. The recent elucidation of the genome sequence of the Nine Mile isolate (RSA493) of C. burnetii (27) now allows a proteomic approach, i.e., two-dimensional (2-D) gel electrophoresis combined with peptide mass mapping and fingerprinting to address these problems. Using these approaches, we conducted an analysis of C. burnetii SCV and LCV protein composition and the developmental form antigens recognized during infection.

**MATERIALS AND METHODS**

**Cultivation and purification of C. burnetii.** C. burnetii Nine Mile Crazy (RSA514) was grown in African green monkey kidney (Vero) fibroblasts (CCL-81; American Type Culture Collection) cultivated in RPMI (Invitrogen) supplemented with 10% fetal bovine serum. This strain was isolated from the placenta of a guinea pig that had been chronically infected for 343 days with the reference strain (RSA439) (9). Bacteria were purified from infected cells 7 days p.i. by Renografin gradient centrifugation (37). SCVs and LCVs were fractionated from the Nine Mile phase I strain (RSA439) (23). Bacteria were purified from infected cells at 7 days p.i. by Renografin gradient centrifugation (37). SCVs and LCVs were separated by equilibrium density centrifugation in 32% cesium chloride (CsCl) as previously described (14). Cell variants were resuspended in K-36 (0.1 M KCl, 0.015 M NaCl, 0.05 M potassium phosphate, pH 7.0) and stored at –80°C.

**Immune sera.** A guinea pig was aerosol infected with 10^8 C. burnetii Nine Mile bacteria in phase I (RSA493) and bled for immune sera at 7, 14, and 28 days p.i. Guinea pig immune serum was also generated by vaccination and subsequent challenge with live organisms. Specifically, a guinea pig was vaccinated with 40 μg of formalin-fixed Nine Mile phase I bacteria mixed 1:1 in incomplete Freund’s adjuvant and boosted 14 days later. After 2 weeks vaccinated animals were aerosol challenged with 10^6 live Nine Mile phase I bacteria, and immune serum was harvested at 28 days p.i. (Protocols for production of guinea pig immune sera were approved by the Texas A&M University Institutional Animal Care and Use Committee.) Human convalescent-phase sera from two patients who had recovered from acute Q fever were obtained from an Australian serum collection recently acquired by J.E.S. Generation of anti-ScvA serum has been previously described (14).

**One-dimensional gel electrophoresis and immunoblotting.** Proteins of CscL2-purified SCVs and LCVs were separated by polyacrylamide gel electrophoresis as previously described (14) using NextGel polyacrylamide solution (Amresco, Solon, OH). For both one-dimensional and 2-D gels (below), equal amounts of SCV and LCV protein were separated as determined by the RC DC protein assay (Bio-Rad). Gels were stained with a SilverQuest silver staining Kit (Invitrogen) and destained with a solution of 40% acetonitrile, 0.1% trifluoroacetic acid. An 0.5-μl volume of porcine trypsin (Promega) at 20 μg/ml was incubated with the mixture at 37°C for 2 h. Five microliters of 5% trifuluroacetic acid was added to the modified peptide, and they were subsequently purified using C4 ZipTips (Millipore). Purified peptides were dried and then rehydrated with 1.5 μl of matrix solution consisting of 2.5 mg/ml α-cyano-4-hydroxycinnamic acid (Sigma) in a solution of 40% acetonitrile, 0.1% trifuluroacetic acid. An 0.5-μl volume of the peptide-matrix solution was spotted on the sample plate and allowed to air dry. Matrix-assisted laser desorption ionization-time of flight (TOF) mass spectra and peptide mass fingerprints were collected by using an Applied Biosystems/ MDS Sciex 4800 matrix-assisted laser desorption ionization TOF/TOF mass spectrometry workstation (Applied Biosystems, Framingham, MA). The resulting peptide peak list and peptide fragment list were submitted for a search of the CDS combined database (Celera Discovery System) by using the Mascot (Matrix Science) search software.

**Transmission electron microscopy.** Purified SCVs and LCVs were fixed overnight at 4°C with 2.5% glutaraldehyde-4% paraformaldehyde in 100 mM sodium cacodylate buffer (pH 7.2). Cells were postfixed with 0.5% osmium tetroxide in 100 mM sodium cacodylate buffer followed by 1% tannic acid in distilled water. Samples were stained overnight with 1% uranyl acetate, washed with distilled water, dehydrated with a graded ethanol series, and embedded in Spurr’s resin. Thin sections were stained with 1% uranyl acetate and Reynold’s lead citrate. Sections were viewed at 80 kV on a Philips CM-10 transmission electron microscope (PEL, Hillsboro, OR). Digital images were acquired with an AMT digital camera (AMT, Chazy, NY) and processed with Adobe Photoshop (version 7.0; Adobe Systems, Mountain View, CA).

**RESULTS**

**Purification of SCVs and LCVs.** Earlier electrophoresis studies revealed distinct protein heterogeneity between SCVs and LCVs (12, 14, 20). To more completely assess the degree of differential protein synthesis by SCVs and LCVs, we conducted a proteomic analysis using 2-D electrophoresis and silver staining. Highly pure preparations of sample are required for reproducible proteome surveys. In a previous study, we fractionated SCVs and LCVs of the Nine Mile phase I strain by exploiting their different buoyant densities in CsCl2 equilibrium gradients (35). While nearly homogeneous populations of developmental forms were obtained, yields were low due to the poor infectivity of phase I C. burnetii for tissue culture cells, a characteristic partly attributed to its hydrophilic, full-length lipopolysaccharide (LPS) (23, 37). The Nine Mile phase II variant is approximately 500 times more infectious for cultured cells (23); however, a severely truncated LPS makes this isolate highly hydrophobic, which causes it to aggregate in CsCl2 (37). We therefore investigated whether the Nine Mile Crazy variant, which produces an intermediate-length LPS and is roughly 20 times more infectious for cell culture than the Nine Mile phase I strain (23), could be separated by CsCl2 equilibrium density centrifugation. Clear separation of Nine Mile Crazy was achieved with LCVs banding at a higher density than SCVs due to their presumed permeability to CsCl2 (35) (Fig. 1A). Although C. burnetii was harvested at 7 days p.i., when the intracellular numbers of SCVs and LCVs are roughly equal (5), the yield of purified SCVs always greatly exceeded that of LCVs, a result attributed to the fragile nature of the LCV. Ultrastructural analysis by transmission electron
microscopy showed that nearly homogeneous SCVs and LCVs were isolated using this procedure (Fig. 1B). Prototypic SCVs were observed that were 0.2 to 0.5 μm in length and rod shaped and with an electron-dense condensed chromatin (Fig. 1B, left panel). The LCV ultrastructure was also typical with cell forms approximately 1 μm in length, pleomorphic, and displaying a granular appearance due to a dispersed chromatin (Fig. 1B, right panel). The homogeneity of SCV and LCV fractions was further confirmed by immunoblotting for ScvA, an SCV-specific protein (5, 14, 28) (Fig. 1C). ScvA was detected only in lysates of fractionated SCV.

**Comparison of SCV and LCV proteomes.** Lysates of purified cell forms were separated in the first dimension using immobilized pH gradient strips having a pH range of 4 to 7 or 7 to 10. This protocol was employed to better resolve the *C. burnetii* proteome and to facilitate detection of low-abundance proteins. In both pH ranges combined, over 675 protein spots were detected in 2-D gels of both SCV and LCV lysates (Fig. 2).

Based on densitometry, 48 proteins were greater than twofold more abundant in LCVs than in SCVs, with 32 and 16 of these proteins having isoelectric points between pH 4 and 7 and pH 7 and 10, respectively. Fifteen upregulated LCV proteins were identified by mass spectrometry, two of which were proteins of unknown function (Table 1). In general, identified proteins had predicted functions in general bacterial physiology including transcription (N utilization protein A [NusA]), translation (ribosomal protein S1 [RpsA], ribosomal protein L9 [RplI], and elongation factor EF-Tu), cell division (cell division protein FtsZ), chromosome partitioning (segregation and condensation protein B [ScpB] and chromosome partitioning protein ParB), riboflavin biosynthesis (6,7-dimethyl-8-ribityllumazine synthase [RibH]), and protein folding (60-kDa...
chaperone [GroEL] and chaperone protein HtpG), and their upregulation is consistent with the LCV being more metabolically and replicatively active than the SCV (5, 21). Upregulation of stringent starvation protein (SspA) (10), universal stress protein A (UspA) (24), and thioredoxin peroxidase (Bcp) (38) may reflect a stress response to the lysosomal environment by the LCV.

Six proteins were detected that were greater than twofold more abundant in SCVs than in LCVs, all having isoelectric points between 7 and 10 (Fig. 2). Four of these were identified by mass spectrometry as TolB, a protein involved in outer membrane stability (17); GTP binding protein Era homolog (Era) (3); cystathionine beta-lyase (MetC); and a hypothetical protein (Table 1).

C. burnetii genes encode an unusually high number of basic proteins with % having a predicted isoelectric point greater than 10 (27). Obvious

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**TABLE 1. Identification of differentially expressed SCV and LCV proteins**

<table>
<thead>
<tr>
<th>No.</th>
<th>SCV/LCV</th>
<th>Gene</th>
<th>Description</th>
<th>Predicted mass (kDa)</th>
<th>Predicted pl</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>LCV</td>
<td>CBU0528</td>
<td>Ribosomal protein S1 (RpsA)</td>
<td>62.1</td>
<td>5.0</td>
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<tr>
<td>2</td>
<td>LCV</td>
<td>CBU0867</td>
<td>Ribosomal protein L9 (RplI)</td>
<td>16.6</td>
<td>6.6</td>
</tr>
<tr>
<td>3</td>
<td>LCV</td>
<td>CBU1433</td>
<td>N utilization substance protein A (NusA)</td>
<td>56.2</td>
<td>4.3</td>
</tr>
<tr>
<td>4</td>
<td>LCV</td>
<td>CBU0236</td>
<td>Elongation factor EF-Tu</td>
<td>43.5</td>
<td>5.2</td>
</tr>
<tr>
<td>5</td>
<td>LCV</td>
<td>CBU1718</td>
<td>60-kDa chaperone (GroEL)</td>
<td>58.3</td>
<td>4.9</td>
</tr>
<tr>
<td>6</td>
<td>LCV</td>
<td>CBU0309</td>
<td>Chaperone protein HtpG (HtpG)</td>
<td>72.8</td>
<td>5.0</td>
</tr>
<tr>
<td>7</td>
<td>LCV</td>
<td>CBU1747</td>
<td>Stringent starvation protein A (SspA)</td>
<td>24.4</td>
<td>5.6</td>
</tr>
<tr>
<td>8</td>
<td>LCV</td>
<td>CBU0141</td>
<td>Cell division protein FtsZ (FtsZ)</td>
<td>40.7</td>
<td>4.3</td>
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<td>9</td>
<td>LCV</td>
<td>CBU1060</td>
<td>Segregation and condensation protein (SmpB)</td>
<td>23.7</td>
<td>4.4</td>
</tr>
<tr>
<td>10</td>
<td>LCV</td>
<td>CBU0658</td>
<td>Hypothetical protein</td>
<td>16.0</td>
<td>4.5</td>
</tr>
<tr>
<td>11</td>
<td>LCV</td>
<td>CBU1754</td>
<td>Hypothetical protein</td>
<td>22.3</td>
<td>4.7</td>
</tr>
<tr>
<td>12</td>
<td>LCV</td>
<td>CBU1927</td>
<td>Chromosome partitioning protein ParB (ParB)</td>
<td>32.0</td>
<td>9.4</td>
</tr>
<tr>
<td>13</td>
<td>LCV</td>
<td>CBU0963</td>
<td>Thioredoxin peroxidase (Bcp)</td>
<td>16.9</td>
<td>8.0</td>
</tr>
<tr>
<td>14</td>
<td>LCV</td>
<td>CBU0648</td>
<td>DMRL synthase (RibH)</td>
<td>16.6</td>
<td>7.3</td>
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<tr>
<td>15</td>
<td>LCV</td>
<td>CBU1916</td>
<td>Universal stress protein A (UspA)</td>
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<td>7.2</td>
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<tr>
<td>16</td>
<td>SCV</td>
<td>CBU0090</td>
<td>TolB protein (TolB)</td>
<td>47.8</td>
<td>8.9</td>
</tr>
<tr>
<td>17</td>
<td>SCV</td>
<td>CBU2025</td>
<td>Cystathionine beta-lyase (MetC)</td>
<td>42.7</td>
<td>8.3</td>
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<tr>
<td>18</td>
<td>SCV</td>
<td>CBU2079</td>
<td>Hypothetical</td>
<td>14.2</td>
<td>8.6</td>
</tr>
<tr>
<td>19</td>
<td>SCV</td>
<td>CBU1502</td>
<td>GTP binding protein Era homolog (Era)</td>
<td>33.9</td>
<td>9.4</td>
</tr>
</tbody>
</table>

*a Unique to or upregulated greater than twofold in the LCV or the SCV.

b DMRL, 6,7-dimethyl-8-ribityllumazine.
derived from a guinea pig vaccinated with formalin-fixed \textit{C. burnetii} in SCVs. Serum obtained at 28 days p.i. recognized 12.5-, 15-, and 30-kDa antigens in both SCV and LCV lysates. Serum reactivity was observed with the 12.5-, 15-, 30-, and 92-kDa common LCV/SCV antigens, and the 84-kDa LCV-specific antigen, that was recognized with 28-day-p.i. serum. Also detected was a novel 22-kDa antigen specific to the SCV. No SCV or LCV antigens were recognized by sera from naive guinea pigs (data not shown). Collectively, these results indicate that SCVs and LCVs express unique antigens. Moreover, based on immunoblot signal intensities, some shared antigens are associated with developmental forms in different amounts.

SCV and LCV antigens recognized by convalescent-phase sera from patients who recovered from acute Q fever. Convalescent-phase immune sera from two patients who had recovered from acute Q fever (2036 and 3004) were analyzed for immunoreactivity against SCV and LCV antigens separated by 2-D gel electrophoresis (Fig. 4). Serum 2036 recognized 20 SCV and 26 LCV antigens. Fifteen antigens were common to both cell variants, five were unique to SCV, and 11 were unique to LCV. By overlaying films of immunoblots with parallel silver-stained gels, we identified by mass spectrometry 10 strongly immunogenic antigens that were generally shared by SCVs and LCVs. Identified common cell variant antigens recognized by patient serum 2036 were GroEL, ribosomal protein L7/L12 (RplJ), acute disease antigen A (AdaA), trigger factor (Tig), and 3-oxoacyl-acyl carrier protein synthase II (FabF) (Table 2). Identified LCV-specific antigens included Bep and repressor protein C2. Serum 3004 recognized 24 SCV and 27 LCV antigens. Thirteen antigens were common to both developmental forms with 11 and 14 antigens unique to SCV and LCV, respectively. Identified antigens common to both cell variants included GroEL, elongation factor EF-Tu, RplJ, ATP synthase F-1 alpha subunit (AtpA), FabF, and isocitrate dehydrogenase (Icd) (Table 2). Tig was recognized only in LCV lysates. We were unable to identify SCV-specific antigens recognized by either serum.

**DISCUSSION**

We have previously demonstrated by gel electrophoresis a few differences in protein composition between \textit{C. burnetii} SCV and LCV developmental forms (12, 14); however, attempts to extensively identify developmental form-specific proteins were not undertaken due to low yields of purified cell variants coupled with the lack of sensitive methods for protein identification. Moreover, previous studies were hampered by poor solubilization of cell variants, with fewer than 200 SCV and LCV proteins resolved by 2-D gel electrophoresis and silver staining (12). In this study we exploited improved yields and solubilization of \textit{C. burnetii} developmental forms along with a dual-pH-range 2-D electrophoresis procedure to develop a more complete proteome map of the SCV and the LCV by more than tripling the number of proteins resolved by silver staining over a previous study (12). Forty-eight proteins...
are greater than twofold more abundant in LCVs than in SCVs, with six proteins greater than twofold more abundant in SCVs than in LCVs. Thus, the morphological differences observed between C. burnetii developmental forms clearly correlate with differences in protein composition.

Using mass spectroscopy, we identified 15 of the 48 proteins differentially synthesized by the LCV. These proteins are generally involved in cell division, RNA synthesis, and protein synthesis/processing. As the more metabolically active cell form of C. burnetii (5, 21), the LCV would be expected to have increased ribosome content. Consistent with this idea, we identified upregulation by the LCV of N utilization substance protein A (NusA), a protein required for robust transcription of rRNA (34), and the ribosomal proteins S1 and L9.

TABLE 2. Identification of immunoreactive SCV and LCV proteins

<table>
<thead>
<tr>
<th>No.</th>
<th>Gene</th>
<th>Description</th>
<th>Predicted mass (kDa)</th>
<th>Predicted pl</th>
<th>Serum 2036</th>
<th>Serum 3004</th>
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<td>Cell variant MS/MS identification</td>
<td>Spot in other cell variant lysate</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Cell variant MS/MS identification</td>
<td>Spot in other cell variant lysate</td>
</tr>
<tr>
<td>1</td>
<td>CBU1718</td>
<td>60-kDa chaperone (GroEL)</td>
<td>58.3</td>
<td>4.9</td>
<td>S</td>
<td>S+L</td>
</tr>
<tr>
<td>2</td>
<td>CBU0236</td>
<td>Elongation factor EF-Tu</td>
<td>43.5</td>
<td>5.2</td>
<td>S</td>
<td>S+L</td>
</tr>
<tr>
<td>3</td>
<td>CBU0229</td>
<td>Ribosomal protein L7/L12 (RplJ)</td>
<td>13.2</td>
<td>4.4</td>
<td>S</td>
<td>S+L</td>
</tr>
<tr>
<td>4</td>
<td>CBU1943</td>
<td>ATP synthase F-1 alpha subunit (AtpA)</td>
<td>58.8</td>
<td>6.0</td>
<td>NA</td>
<td>L</td>
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<tr>
<td>5</td>
<td>CBU1416</td>
<td>Repressor protein C2</td>
<td>24.3</td>
<td>7.6</td>
<td>L</td>
<td>L</td>
</tr>
<tr>
<td>6</td>
<td>CBU0952</td>
<td>Acute disease antigen A (AdaA)</td>
<td>26.0</td>
<td>9.3</td>
<td>L</td>
<td>L</td>
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<tr>
<td>7</td>
<td>CBU0963</td>
<td>Thioredoxin peroxidase (Bcp)</td>
<td>16.9</td>
<td>8.0</td>
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<td>L</td>
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<td>8</td>
<td>CBU0737</td>
<td>Trigger factor (Tig)</td>
<td>50.2</td>
<td>5.1</td>
<td>+</td>
<td>+</td>
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<tr>
<td>9</td>
<td>CBU0497</td>
<td>3-Oxoacyl-acyl carrier protein synthase II (FabF)</td>
<td>44.1</td>
<td>5.5</td>
<td>+</td>
<td>+</td>
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<tr>
<td>10</td>
<td>CBU1200</td>
<td>Isocitrate dehydrogenase (Icd)</td>
<td>46.6</td>
<td>6.4</td>
<td>NA</td>
<td>L</td>
</tr>
</tbody>
</table>

* Protein identified by mass spectrometry in lysates of the SCV (S), the LCV (L), or both (S+L).
* Corresponding spot present in immunoblot of lysates derived from other cell variants is indicated with a +.
* NA, not applicable.
* Immunoreactive spot at the same coordinates in both SCV and LCV 2-D immunoblots as those of the protein identified by serum 3004 in the LCV.
enhancing the translational machinery of the LCV is upregulation of EF-Tu, a finding in keeping with a previous study (26). We have recently confirmed that the LCV is the replicatively active cell form of the C. burnetii developmental cycle (5). As such, and in accordance with increased metabolic activity, the LCV upregulates FtsZ, required for bacterial binary fission, and two proteins involved in chromosome partitioning, ScepB and ParB (11, 30), relative to the SCV.

At first glance, upregulation of stringent starvation protein A (SspA) by the LCV seems counterintuitive as, among other effects, expression of this protein in Escherichia coli results in increased synthesis of the stationary-phase sigma factor RpoS (10). However, we (5) and others (28) have previously shown that RpoS is upregulated by the LCV during exponential growth, where it potentially induces genes involved in protecting C. burnetii against lysosomal stress. Indeed, RpoS induction of hydroperoxidase I in exponentially growing Vibrio vulniﬁcus is critical for survival of this organism during oxidative stress (25). In addition to regulating RpoS levels, SspA is thought to be a global regulator of bacterial virulence genes (10). Along with SspA, elevated levels of Bep, UspA, GroEL, and HtpG in the LCV also support the idea that this developmental form is responding to lysosomal stress. As a thioredoxin peroxidase, Bcp resides within the general family of thiol-speciﬁc antioxidant proteins termed peroxiredoxins. They are abundant proteins in bacteria and serve to detoxify various peroxide compounds (38). In E. coli, UspA is essential for survival of a variety of insults including exposure to hydrogen peroxide (24).

While the chaperones GroEL and HtpG are abundant in bacteria growing under normal conditions, they are also upregulated in response to a variety of adverse conditions (6, 29). Moreover, GroEL and HtpG are implicated in protein secretion (16, 32), and we have previously speculated that the LCV is the secretion-competent form of C. burnetii (5).

The most abundant LCV-upregulated protein that we identiﬁed is annotated as a hypothetical protein (CU90668). The Coxiella genome encodes a high proportion of hypothetical proteins (33.7% of coding sequences) (27). It is reasonable to suspect that these play critical roles in C. burnetii morphological development and pathogenesis; however, their functional assignments await future biochemical and genetic analysis. We did not exhaustively attempt to identify by mass spectrometry the remaining subset of LCV-upregulated proteins. However, it likely includes RpoS and EF-Ts, previously described as differentially synthesized by the LCV (35, 37), as LCV-upregulated proteins with molecular masses and pIs approximating those of these proteins are present on 2-D gels.

We identiﬁed four proteins differentially synthesized by the SCV with three, TolB, Era homolog, and MetC, having predicted functions. In E. coli, TolB interacts with other Tol proteins to stabilize the outer membrane (17). Elevated TolB levels in the SCV may partially explain this developmental form’s noted resistance to physical disruption. The E. coli protein Era is a member of the small regulatory GTPase superfamily (3). Among the proposed functions of Era and its homologs are regulation of the cell cycle and cellular differentiation (3). For example, Era homologs of Bacillus subtilis are required for normal cell division and spore formation (1, 4). Thus, this protein may be a key regulator of SCV morphological differentiation. Upregulation of MetC, a biosynthetic enzyme, by the SCV is puzzling. This enzyme catalyzes the last intermediate of the methionine biosynthetic pathway, converting cystathionine to l-homocysteine. In E. coli, homocysteine globally affects transcription and inhibits growth by perturbing the activity of threonine deaminase, which results in depletion of branched-chain amino acids (8, 31). Consequently, homocysteine may serve a signaling role in the SCV to downregulate metabolism. Alternatively, SCV may require more methionine for production of S-adenosylmethionine. Among the many uses of S-adenosylmethionine is production of spermine and spermidine (7). These cationic polyamines bind to DNA which stabilizes the molecule and facilitates chromatin condensation (1), a hallmark of the SCV. Two very basic SCV-speciﬁc DNA-binding proteins, ScvA and Hq1, have been previously described (13, 14) that also presumably play roles in chromatin condensation. Interestingly, all SCV-upregulated proteins were found in pH 7 to 10 2-D gels. Moreover, there is an abundance of unresolved SCV proteins that “stack up” at the basic end of pH 7 to 10 2-D gels relative to the LCV. In addition to ScvA and Hq1, these likely include additional SCV-upregulated proteins involved in chromatin condensation or other processes that contribute to the resistance properties of SCV. Resolution and identiﬁcation of these proteins will require an alternative method of protein fractionation. Similarly to the LCV, the most abundant SCV-upregulated protein based on densitometry is annotated as a hypothetical protein (CU90279).

P1 is the only surface antigen known to be differentially synthesized by C. burnetii developmental forms (20, 33). Here we extensively examined SCV and LCV antigens recognized in the context of both infection and vaccination with killed organisms. An infected guinea pig developed antibodies to an increasingly complex array of SCV and LCV antigens over the time course of infection. Although different levels of expression are apparent, all SCV antigens recognized at 28 days p.i. are also present in the LCV. A small subset of these common antigens is also recognized by vaccinated animals that were challenged with live organisms. The less complex humoral response elicited by vaccination and subsequent infection is not unexpected and has been reported elsewhere for C. burnetii (40). At least two antigens with approximate molecular masses of 42 and 84 kDa appear unique to the LCV. Like P1, these antigens may be surface exposed and involved in nutrient uptake by the LCV. Moreover, antigenic variation in vivo between the SCV and the LCV may play a role in the immune evasion and persistence of C. burnetii that lead to chronic infections (39).

An infected guinea pig ﬁrst generates antibodies to 12.5- and 15-kDa antigens with strong recognition of these proteins throughout the course of infection. Similar reactivities are observed with sera derived from vaccinated animals. Based on immunoblot signal intensities, these antigens appear more abundant in the SCV than in the LCV. Diagnosis of Q fever is usually by serological methods with the reference technique being microimmunofluorescence (19). This test employs inactive whole-cell C. burnetii as antigen. A serodiagnostic test based on recombinant antigen would eliminate the need for cultivation and puriﬁcation of C. burnetii and would likely be more speciﬁc. As the 12.5- and 15-kDa antigens are the predominant antigens recognized during the early stages of infec-
tion, they may have utility as serological diagnostic target antigens in the form of recombinant protein. At 28 days p.i., LPS comprises a portion of the 12.5-kDa-antigen reactivity. *C. burnetii* LPS is strongly immunogenic and also comprises the organism’s phase I antigen (41). Recognition of LPS only late in infection is in keeping with the serological phenomenon of phase variation whereby only late antisera (>20 days p.i.) of an infected animal binds to phase I antigen (i.e., LPS) (23). Consistent with a previous report (20), there appears to be more LPS associated with the SCV than with the LCV. To develop a more complete understanding of the human humoral response to *C. burnetii*, we identified immunogenic proteins recognized in the context of human acute Q fever. Similar to the reactivity observed with infected guinea pig sera, in LCV than SCV proteins. While a number of SCV- and LCV-specific proteins are recognized by convalescent-phase sera, in most cases these are weakly immunogenic. However, one human serum strongly recognized two LCV-specific proteins: Bep and repressor protein C2. C2 is an S24 peptidase family autoproteinase which is commonly involved in activation of RecA and repressor protein C2. C2 is an S24 peptidase family specific proteins are recognized by convalescent-phase sera, in most cases these are weakly immunogenic. However, one human serum strongly recognized two LCV-specific proteins: Bep and repressor protein C2. C2 is an S24 peptidase family autoproteinase which is commonly involved in activation of RecA (18). As discussed earlier, Bep may protect *C. burnetii* against oxidative stress. Selective recognition by the human immune system of both proteins only in the LCV is consistent with an elevated stress response by this developmental form. The remaining identified antigens are present in both the SCV and the LCV with GroEL, RplJ, and FabF recognized by both human sera in both cell forms. GroEL and the ribosomal protein L9 were recently identified as immunogens by screening a *C. burnetii* expression library with serum from *Coxiella*-infected mice (40). FabF, an enzyme involved in fatty acid biosynthesis; Icd, a Krebs’ cycle enzyme; and AtpA, a subunit of ATP synthase, are considered housekeeping enzymes. However, antigenic *C. burnetii* housekeeping proteins have been reported elsewhere (40), indicating that general metabolic enzymes of the organism are exposed to the host immune system.

AdA, EF-Tu, AtpA, and Icd were recognized in both SCV and LCV lysates by a single convalescent-phase serum. AdA was recently identified as an immunodominant outer membrane protein synthesized only by isolates acquired from patients with acute disease (42), and mice generate an antibody response to EF-Tu when vaccinated with formalin-fixed *C. burnetii* (26). One human serum recognized Tig, an abundant ribosome-associated chaperone (6), in both SCV and LCV lysates while the other human serum recognized Tig only in LCV lysates.

In summary, this study provides an extensive proteome analysis of *C. burnetii* developmental forms. Identification of cell-form-specific and common antigens provides new information for development of rationally designed subunit vaccines and new diagnostic tests. This is particularly relevant with respect to surface antigens of the SCV, the environmentally stable cell form.

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