A unique *Coxiella burnetii* lipoprotein involved in metal binding (LimB)

James M. Battisti, Linda D. Hicks and Michael F. Minnick

Division of Biological Sciences, The University of Montana, Missoula, MT 59812, USA

*Coxiella burnetii* is the bacterial agent of Q fever in humans. Here, we describe a unique, ~7.2 kDa, surface-exposed lipoprotein involved in metal binding which we have termed LimB. LimB was initially identified as a potential metal-binding protein on far-Western (FW) blots containing whole-cell lysate proteins when probed with nickel-coated horseradish peroxidase (Ni-HRP) and developed with a chemiluminescent HRP substrate. The corresponding identity of LimB as CBU1224a was established by matrix-assisted laser desorption ionization-tandem time-of-flight mass spectrometry. BLAST analyses with CBU1224a showed no significant similarity to sequences outside strains of *C. burnetii*. Additional *in silico* analyses revealed a putative 20 residue signal sequence with the carboxyl end demarcated by a potential lipobox (LSGC) whose Cys residue is predicted to serve as the N-terminal, lipidated Cys of mature LimB. The second residue of mature LimB is predicted to be Ala, an uncharged envelope localization residue. These features suggest that CBU1224a is synthesized as a prolipoprotein which is subsequently lipidated, secreted and anchored in the outer membrane. Mature LimB is predicted to contain 45 aa, of which there are 10 His and 5 Cys; both amino acids are frequently involved in binding transition metal cations. Recombinant LimB (rLimB) was generated and its Ni-HRP-binding activity demonstrated on FW blots. Ni-HRP binding by rLimB was inhibited by >95% on FW blots done in the presence of EDTA, imidazole, Ni$^{2+}$ or Zn$^{2+}$, and roughly halved in the presence of Co$^{2+}$ or Fe$^{3+}$. The *limB* gene was maximally expressed at 3–7 days post-infection in *Coxiella*-infected Vero cells, coinciding with exponential phase growth. Two isoforms of LimB were detected on FW and Western blots, including a smaller (~7.2 kDa) species that was the predominant form in small cell variants and a larger isoform (~8.7 kDa) in large cell variants. LimB is Sarkosyl-insoluble, like many omps. The predicted surface location of LimB was verified by immunoelectron and immunofluorescence microscopy using anti-rLimB antibodies. Overall, the results suggest that LimB is a unique *Coxiella* lipoprotein that serves as a surface receptor for divalent metal cations and may play a role in acquiring at least one of these metals during intracellular growth.

**INTRODUCTION**

*Coxiella burnetii* is a gammaproteobacterium and the agent of Q fever in humans. *C. burnetii* is one of the most infectious pathogens known, with an ID$_{50}$ of 1–10 bacteria in the guinea pig model (Moos & Hackstadt, 1987). Human infections with *C. burnetii* are generally zoonoses acquired by inhalation of contaminated aerosols. Q fever typically presents as an acute, self-limiting flu-like illness accompanied by pneumonia or hepatitis.

In roughly 1% of cases, a severe chronic infection can occur, in which endocarditis is the predominant manifestation (Maurin & Raoult, 1999). These attributes and its past use as a biological weapon component (Regis, 1999) were grounds for classifying *Coxiella* as an ‘HHS Select Agent’.

In nature, *C. burnetii* is an obligate intracellular pathogen and undergoes a developmental cycle that exhibits two distinct cell morphotypes. The infectious small cell variant (SCV) has been described as 'spore-like', and it is extremely resistant to environmental stressors, including UV light and desiccation. Shortly after entry into an eukaryotic phagolysosome-like compartment called a parasitophorous vacuole (pH ~4.5), the SCV transforms into a 'vegetative' morphotype, termed a large cell variant (LCV) (McCaul & Williams, 1981). After 5–6 days of intracellular replication, the LCV transforms back to a SCV that is released from the host cell to complete the developmental cycle (Coleman...
**Methods**

**Bacterial strains, cell lines and growth conditions.** NM PII C. burnetii (strain RSA 439; clone 4) was propagated in African green monkey kidney (Vero) epithelial cells (CCL-81; American Type Culture Collection), as described previously (Raghavan et al., 2008). SCVs were prepared and purified from 28-day-old cultures as described previously (Raghavan et al., 2008). SCVs were used to inoculate Vero cell monolayers to produce synchronous infections (Coleman et al., 2004). Mixed-cell populations of LCVs and SCVs were harvested from synchronous cultures as for SCVs, but at 10 days post-infection. LCVs were prepared from 72 h synchronous cultures using a digitonin-based protocol (Cockrell et al., 2008). Whole-cell lysates of Coxiella were obtained by using a freeze-thaw method (Samoïlis et al., 2007).

**SDS-PAGE and far-Western (FW) blots.** Protein concentrations were determined by using a BCA protein assay kit (Thermo Scientific). Bacterial suspensions were solubilized with Laemmli sample buffer (Laemmli, 1970). Samples were typically boiled for 10 min and centrifuged (1 min, 10,000 g) to remove insoluble material; the resulting supernatant was separated on an SDS-PAGE gel (Laemmli, 1970). Protein bands were visualized by staining gels with Coomassie brilliant blue [CBB; 0.1 % (w/v) in 50 % methanol, 7 % (v/v) acetic acid].

**Inhibition studies.** Inhibition studies were done using recombinant LimB (rLimB) and FW blots were prepared as described above, except the probing solution contained a potential competitive inhibitor of binding, including EDTA (100 mM) or imidazole (500 mM). In some cases, HRP was used in place of Ni-HRP, at the same final concentration (1 μg ml⁻¹), to demonstrate that LimB binding activity was dependent on the nickel moiety of Ni-HRP. Finally, to qualitatively analyse the relative affinity of LimB for various metal ions, modified FW blots were performed based on a previously published protocol (Zhao & Waite, 2006). First, rLimB (5 μg protein) was separated by SDS-PAGE, transferred to nitrocellulose, blocked and washed as above. Second, blots were incubated (1 h; 25 °C) in TBST containing chloride salts of individual metal ions (FeCl₃·6H₂O, CoCl₂·6H₂O, NiCl₂·6H₂O or ZnCl₂) at a final concentration of 10 mM. Excess metal was subsequently removed by washing three times for 5 min with TBST containing 0.1 mM of the individual metal ion. Third, resulting FW blots were probed and washed as above except solutions contained 0.1 mM of the particular metal ion. Finally, blots were developed and visualized as above.

**Mass spectrometry (MS).** A C. burnetii whole-cell lysate (20 μg protein) was separated by using SDS-PAGE [10–15 % acrylamide (w/v)] and stained with CBB, as above. Bands corresponding to Ni-HRP-binding activity on FW blots were excised from gels and subjected to trypsin digestion and MALDI-TOF peptide mass fingerprinting and MALDI-tandem TOF (MALDI-TOF/TOF) peptide sequencing. All MS work and database searches (NRDB1 database using Mascot version 2.2.03 software) were performed by Alphalyse.

**Nucleic acid isolation and rLimB production.** Infected monolayers were harvested at specific time points (0, 3, 5, 6, 7 and 8 days) by replacing culture medium with 2.5 ml TRI reagent (Ambion), briefly incubating (5 min, 25 °C) and scraping. Harvested monolayers were stored (−80 °C) for subsequent nucleic acid isolation. RNA and DNA were isolated from each harvested monolayer sample essentially as

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*et al.*, 2004; McCaul & Williams, 1981). In addition to distinct developmental forms, two separate phase variants of the bacterium have also been described. First, wild-type *C. burnetii* is typically 'phase I', in reference to serological reactivity against its 'smooth', long-chain LPS, and must be manipulated in a bio-safety level (BSL)-3 facility. Second, repeated passage in tissue culture or embryonated hens' eggs results in the spontaneous generation of attenuated 'phase II' (PII) organisms that synthesize a 'rough' LPS that can be caused by a chromosomal deletion (Hoover et al., 2002; Vodkin & Williams, 1986), and may be handled in a BSL-2 laboratory. Finally, a number of *C. burnetii* strains [such as ‘Nine Mile’ (NM)] have been isolated from different parts of the world and have been grouped according to their association with acute or chronic disease manifestations of Q fever (Beare et al., 2009; Seshadri & Samuel, 2005).

Despite several fascinating attributes, little is known regarding Coxiella's virulence determinants. In fact, LPS is the only bona fide virulence factor described to date (Moos & Hackstadt, 1987), and it has been shown to function as an immune-evasion shield (Shannon et al., 2005). One potential set of virulence factors that has received relatively little attention is the pathogen's considerable assemblage of lipoproteins. Lipoproteins have long been recognized as virulence determinants in a variety of bacterial pathogens, and a number of them have been developed into vaccine immunogens (Fortney et al., 2002; Vodkin & Williams, 1986), and may be handled in a BSL-2 laboratory. Finally, a number of...
described previously (Raghavan et al., 2008). RNA was isolated from this extract by the addition of 300 µl 100% ethanol and use of a RiboPure kit (Ambion) according to the manufacturer’s instructions, and then stored at −20 °C. DNA and RNA were removed from these samples by DNase treatment (TURBO DNase kit; Ambion) and an RNA clean-up step (RNeasy kit; Qiagen). Purified RNA was quantified spectrophotometrically (260 nm), and 1 µg RNA from each sample was converted to cDNA with an iScript cDNA kit (Bio-Rad), as instructed. Finally, cDNA was used to analyse limB transcript profiles by quantitative real-time PCR (qRT-PCR).

DNA was isolated using TRI Reagent DNA/protein isolation protocol (Ambion). DNA-containing supernatant was transferred to another microcentrifuge tube and 40 µl HEPES (0.1 M) was added to adjust the pH to 8.2. Isolated DNA was cleaned using a QIAquick PCR purification kit (Qiagen) as instructed and stored (−20 °C) until needed.

A portion of Coxella limB encoding the predicted, mature LimB (i.e. Cys21 to His102) was cloned into pMAL-c5x (NEB) to create pJB202, a plasmid that encodes a LimB/maltose-binding protein (MBP) fusion. Briefly, primers (LimBF + XmnI, CTGGCAAGAA-GGCTTICAGGTGGGTCCCAGC; LimBR + BamHI, AAGGATTCTCCTTACAAATGATGATTACAATGACGATG) were designed to amplify the CBU1224a ORF from the codon for Cys 21 to the termination codon, and both primers contained restriction sites (underlined). PCR was done by using standard methods (Ausubel et al., 1995) and the resulting amplicon was purified (PCR Purification kit, Qiagen), digested with XmnI and BamHI, repurified, and ligated into the corresponding restriction sites of pMAL-c5x (NEB) using T4 DNA ligase. E. coli BL21 (DE3) was transformed with pJB202 (Chung et al., 1989) and the resulting transformants were selected by plating on Luria–Bertani plates containing ampicillin (100 µg ml−1). Plasmid constructs of selected clones were verified by automated sequence analysis. Amylene resin was used according to the manufacturer’s instructions (NEB) to isolate rLimB fusion protein from the supernatant. Fractions containing rLimB were identified by SDS-PAGE, pooled and dialysed against 20 mM Tris, 50 mM NaCl, 1 mM CaCl2 buffer (pH 6.5) using a Slide-a-Lyser cassette (Pierce; molecular weight cut-off, 3500 Da). rLimB was stored at −20 °C until needed.

qRT-PCR and quantitative PCR (qPCR). Primers were designed using Beacon Designer Software (Premier Biosoft). Primers were synthesized (Integrated DNA Technologies) and used to quantify limB mRNA over the course of a synchronous infection by normalizing to the number of C. burnetii genomes, as described previously (Coleman et al., 2004; Raghavan et al., 2008). First, the number of genomes in each purified DNA sample at each time point (0, 3, 5, 6, 7 and 8 days) was determined in triplicate qPCR reactions (each triplicate experiment was performed twice, thus resulting in six independent determinations). This was accomplished using an iCycler single-colour real-time PCR detection system (Bio-Rad), SYBR Green supermix (Bio-Rad) and a dotA (CBU_1648) primer set (DOTA-F, GATAGCGGTGGAGCCGTGAC; DOTA-R, TCTTTTGGCATCTGTGATT) by normalisation to a standard curve of 10-fold serially diluted (10−6 to 10−10) C. burnetii genomes. Second, cDNA from each time point was used to determine the starting quantity (SQ) mean, a value calculated by iCycler software (MyIQ, v1.0, Bio-Rad) normalizing limB transcript numbers with a limB primer set (LIMB-F, GGT-CCTGT GTTGATAAACTG; LIMB-R, GGTTTATGGTCTCCTT) to a standard curve generated by diluting 1:100 rLimB or rCom1 (Hicks et al., 2010) to both primary and booster injections. Rabbit anti-MBP polyclonal antibodies were purchased (NEB) and used as a negative control for the LimB fusion protein partner. Mouse monoclonal antibody against Coxella elongation factor-Ts (EF-Ts) was used as a negative control for intracellular proteins that may have inadvertently leaked from the bacterium during handling and preparation for microscopy (a generous gift from Robert Heinzen).

Immunoblots were done by transferring un-fixed SDS-PAGE gels to support nitrocellulose and probing overnight with anti-rLimB antibodies (1:5000 dilution) and peroxidase-conjugated goat anti-rabbit antibodies (Sigma), as described previously (Parrow et al., 2009), except ECL reagents were used during development according to the manufacturer’s instructions (Pierce), and signals were visualized with an LAS-3000 digital imaging system (Fujifilm).

Immunofluorescence microscopy to identify surface-exposed proteins was done by the general methods described by Ge & Rikihisa (2007). Bacteria were cultured and harvested from Vero cells at 5 days post-infection, as above, pelleted and washed in PBS containing 0.25 M sucrose (PBSS; pH 7.4). This, and all subsequent steps were done at 25 °C. Bacteria were fixed for 45 min in 2% (w/v) paraformaldehyde. After quenching in PBS containing 0.1 M glycine and subsequently washing in PBS, bacteria were incubated for 1 h with anti-LimB, anti-Com1, anti-EF-Ts, anti-MBP or pre-immune serum (PIS) at a 1:100 dilution in 0.2% (w/v) gelatin in PBS (PG buffer). After washing with PG buffer, bacteria were labelled for 1 h with Alexa Fluor 488 goat anti-rabbit or mouse (for anti-EF-Ts) immunoglobulin G (Invitrogen) at a dilution of 1:100 in PG buffer. Bacteria were washed in PG buffer, suspended in PBS and imaged with a BX51 fluorescence microscope and digital acquisition software (DP2-BSW) (Olympus).

To conduct immunoelectron microscopy, bacteria were cultured and harvested at 5 days post-infection as described above, and diluted in PBSS to a concentration of ~1×104 genomes µl−1. The suspension was combined with an equal volume (100 µl) of antiserum or PIS [diluted 1:100 in PBS-B (PBS containing 0.01 mg BSA ml−1)] and incubated for 1 h at 37 °C. Cells were then centrifuged (10 min at 10 000 g), the pellet was washed with 300 µl PBSS, and the pellet was suspended in 200 µl PBS-B. An aliquot (100 µl) of the cell suspensions was combined with an equal volume of protein A–gold conjugate [Electron Microscopy Sciences (EMS)] pre-diluted 1:20 in PBS-B. Following a brief incubation (50 min, 37 °C), cells were centrifuged and washed as above, and finally suspended in 30 µl PBS. Half the volume of these suspensions was placed on Formvar-coated TEM grids (EMS; 400 mesh) and incubated for 10 min at 23 °C to allow for adherence. Excess suspension was wiped from the grid surface by using blotting paper, replaced with 15 µl uranyl acetate [2% (w/v) in H2O, J. T. Baker, Phillipsburg, NJ], and incubated for 3 min at 23 °C. Excess protein A–gold was washed from grids by successive immersions in three beakers containing 2 l dH2O. Finally, grids were air-dried and analysed by using a Hitachi 7100 transmission electron microscope.

Sarkosyl fractionation. Omps of C. burnetii were enriched by Sarkosyl-fractionation of the bacterium. A whole-cell lysate from a mixed-cell population of Coxella was prepared and the insoluble cell-wall fraction was pelleted by centrifugation. Briefly, the pellet was suspended in 300 µl PBS (pH 7.4), to which 38 ml 0.1 M sodium bicarbonate (pH 11) was added, then incubated on ice for 1 h (with rocking). The mixture was centrifuged (141 000 g for 80 min, 4 °C) and the resulting pellet was extracted with 4 ml sodium N-lauroylsarcosine (Sarkosyl; Sigma-Aldrich) solution (2%; w/v). The extraction was done on ice for 30 min (with rocking). The Sarkosyl-
insoluble (omp-rich) fraction was isolated by centrifuging the mixture (138 000 g for 1 h, 4 °C). The Sarkosyl-soluble fraction (supernatant) was precipitated in 4 vols acetone for 1 h at −20 °C. The mixture was centrifuged (3000 g for 30 min at 4 °C) to obtain a pellet. Both fractions were analysed by SDS-PAGE and blotting.

**Sequencing, in silico analyses and statistics.** Sequence data were obtained with an automated DNA sequencer (ABI3130xl) and a BigDye terminator cycle sequencing ready reaction kit (ABI). GenBank searches were done by using BLAST (http://www.ncbi.nlm.nih.gov/BLAST/). In silico tools available at the database of bacterial lipoproteins (DOLOP; http://www.mrc-lmb.cam.ac.uk/genomes/dolop/; Babu et al., 2006) were used to determine if a predicted protein sequence contained a prolipoprotein signal peptide. MacVector genetic software (Version 11.1.1) was used for protein sequence analysis as well as sequence manipulations. Comparative densitometry of FW blot signals was done using ImageJ 1.42q software (http://rsbweb.nih.gov/ij/). At least three independent determinations were used to calculate means and standard deviations for all numerical data. Statistical and graphical analyses were done using InStat3 (GraphPad) and/or Excel (Microsoft) software. Statistical significance was determined using Student’s t test, where a P-value of <0.05 was considered significant.

**RESULTS**

**FW blots reveal a ~7.2 kDa component of *Coxiella* that binds Ni-HRP**

A prominent, chemiluminescent band of ~7.2 kDa was discovered on Ni-HRP-probed FW blots containing whole-cell lysate proteins from a mixed-cell population (10-day-old culture) of *C. burnetii* LCVs and SCVs (Fig. 1a and b). We were curious to determine whether the signal was a result of Ni-HRP binding to a *C. burnetii* constituent or whether it was due to an endogenous peroxidase that was directly acting on the chemiluminescent HRP substrate. To address this question, we performed FW blots as above but without Ni-HRP. The resulting FW blots showed that even with prolonged exposure (5 min) a chemiluminescent band was not detectable Fig. 1(b) (~7.2 kDa), suggesting that the FW blot signal was due to Ni-HRP binding to a *C. burnetii* component, and not due to an endogenous peroxidase. To ensure that Ni-HRP was not binding a contaminating host-cell constituent, Vero cells were cultivated and harvested for Ni-HRP FW blot analyses (Fig. 1c). The resulting FW blots, prepared as above, showed no detectable chemiluminescent signal in Vero host cell lysate proteins (Fig. 1d). Finally, FW blots of *C. burnetii* were prepared and probed with HRP; however, a chemiluminescent signal was not produced in the absence of Ni (data not shown). From these results, we hypothesized that the Ni-HRP was binding to a *C. burnetii* component, that binding was dependent on the nickel moiety of the probe, and that a small, His-rich protein might be responsible.

**MS identifies the Ni-HRP binding protein as CBU1224a**

To identify the *C. burnetii* protein responsible for Ni-HRP binding, MALDI-TOF peptide mass fingerprinting and MALDI-TOF/TOF peptide sequencing were employed; these identified two major peaks in the chromatogram that matched two of the tryptic peptide fragments calculated for GI-153209557, a *C. burnetii* putative lipoprotein. MS sequencing results represented 56% coverage of the predicted mature, 45 aa lipoprotein. These data allowed us to presumptively identify the Ni-HRP-binding ligand as the protein encoded by the CBU1224a locus of *C. burnetii* (strain RSA 493).

**In silico analysis of CBU1224a reveals a potential lipoprotein signal sequence, a ‘lipobox’ and a His-rich carboxyl-terminus**

Several in silico analyses were performed on CBU1224a and its encoded protein. BLAST was used to search the GenBank...
sequences of six *C. burnetii* strains (RSA 493, RSA 331, Q177, Q154, Q212 and 5J108-111) (Beare et al., 2009) (Seshadri & Samuel, 2005; Seshadri et al., 2003) possess ORFs that are 100 % identical to CBU1224a. In each strain, the 198 bp ORF is flanked by genes encoding haloalkane dehalogenase (111 bp upstream) and an RmuC family protein (105 bp downstream). RmuC (rearrangement mutator) family proteins are hypothesized to play a role in DNA structure/recombination (Slupska et al., 2000) and haloalkane dehalogenase (EC 3.8.1.5) is an enzyme involved in degradation of halocarbons. Based on considerable intergenic spacing, neither of the flanking genes appears to be transcriptionally linked to CBU1224a. Finally, BLAST searches of the non-redundant GenBank database using DNA, as well as the predicted 65 aa residue prolipoprotein and 45 aa mature lipoprotein sequences, did not yield significant similarities outside *C. burnetii*.

We also used *in silico* tools available on the DOLOP website (Babu et al., 2006) to investigate the putative lipoprotein encoded by CBU1224a. These analyses revealed a potential ‘lipobox’ and characteristic N, H and C prolipoprotein signal peptide regions, supporting the ‘lipoprotein’ annotation for the locus. Specifically, the characteristics of the lipobox are: (i) it consists of 4 aa with the consensus sequence [LV][AST][V][AS][G]=C (abbreviated as LXXC), (ii) it is located within the first 40 residues of the prolipoprotein, and (iii) it is also called the Cys region (Fig. 2).

Based on studies with other Gram-negative bacteria, the mature CBU1224a lipoprotein (unlipidated) is predicted to be 45 aa in length (~5.2 kDa), and would be located in the *C. burnetii* outer membrane. Several features and results support this hypothesis. First, the envelope localization residue is an Ala, and would therefore help direct the mature protein to the outer membrane (Yamaguchi et al., 1988). Second, two tryptic fragments identified by MALDI-TOF MS (Thr25-Lys41 and His53-Arg60) are peptides from the predicted mature lipoprotein and not the leader sequence.

The high-affinity binding of Ni-HRP observed in FW blots (Fig. 1) is probably explained by the high percentage of His residues located in the C-terminal portion of CBU1224a (Fig. 2). Specifically, the predicted, mature 45 aa lipoprotein contains ten His (22 % of the total amino acids), nine of which are located in the last 20 aa of the protein. Four Cys residues are also found in this region, an observation that is noteworthy for two reasons: (i) Cys residues could be partially responsible for Ni-HRP binding as both His and Cys are known to coordinate binding of transition metals, and (ii) intra- and/or inter-disulfide bonds could contribute to structure or multimerization of the protein, respectively.

**Recombinant protein encoded by CBU1224a binds Ni-HRP**

To unequivocally demonstrate that Ni-HRP binding on FW blots was due to the protein encoded by CBU1224a, a plasmid (pJB202) was constructed to produce a translational fusion with MBP. FW blots were prepared to compare Ni-HRP binding characteristics of *E. coli* BL21 (pJB202) versus *E. coli* BL21 (pMAL-c5X; empty cloning vector). Results clearly show that the recombinant protein binds Ni-HRP (Fig. 3, lanes 4 and 5), like the native protein (Fig. 3, lane 1). In addition, recombinant MBP (rLimB’s fusion partner) does not bind the probe (Fig. 3, lanes 2 and 3). Based upon these results and our *in silico* analyses, we designated CBU1224a as a lipoprotein involved in metal-binding (LimB).

**rLimB binds divalent cations, which involves His residues**

To further characterize Ni-HRP binding, rLimB was employed in FW blots done in the presence of inhibitors...
signal relative to untreated controls (Fig. 4, Ni \textsuperscript{2+}~95 % reduction in their Ni-HRP chemiluminescent signal of four metal ions. FW blots blocked with nickel showed FW blots by incubation in TBST containing chloride salts following transfer of proteins to nitrocellulose, we blocked particular metal to LimB. To this end, immediately 'block' the binding of Ni-HRP on FW blots. The relative competition capacity of free metal ions could therefore be analysed by FW blots. Where a reduction in chemiluminescent signal was not completely detectable relative to untreated controls. These data suggest that rLimB binds Ni\textsuperscript{2+} and Zn\textsuperscript{2+} with high affinity. In contrast, when FW blots were blocked with cobalt (Co\textsuperscript{2+}) or iron (Fe\textsuperscript{3+}), the chemiluminescent signal was roughly half of untreated controls, suggesting that these metals bind LimB with relatively lower affinity than Ni\textsuperscript{2+} or Zn\textsuperscript{2+}. Taken as a whole, the results demonstrate that LimB binds a variety of divalent metal cations with a higher affinity for Ni\textsuperscript{2+} and Zn\textsuperscript{2+} relative to Co\textsuperscript{2+} and Fe\textsuperscript{3+}.

**2-Mercaptoethanol (2-ME) treatment implicates the Cys residues of LimB in metal binding**

We were curious whether the Cys residues of LimB also play a role in Ni-HRP binding. As mentioned above, four Cys residues are located in the C-terminal portion of LimB and might bind nickel and/or confer inter- or intra-LimB disulfide bonds. To address these possibilities, cell lysate proteins of *C. burnetii* were prepared with and without 2-ME, and FW blots were done as described above. Resulting blots showed a marked reduction in the chemiluminescent signal of LimB when samples were prepared without 2-ME compared with those with 2-ME (data not shown). Increased binding of Ni-HRP in the presence of 2-ME may be due to increased exposure of His and/or increased availability of thiol ligands as a result of disulfide reduction. Multiple LimB bands that would indicate disulfide-mediated inter-LimB multimerization were never observed; however, it is still possible that multimers occur, especially considering the low Ni-HRP binding efficiency observed in the absence of 2-ME.

**limB expression is maximal during exponential phase**

To determine whether expression of *limB* changes during the developmental cycle of *C. burnetii*, synchronous infections were initiated and the transcript quantity was analysed over an 8 day time-course using qRT-PCR. Results showed that *limB* mRNA was significantly higher (*P*<0.05) at 3–7 days post-infection, relative to the initial day (day 0). The *limB* transcripts were highest at 5 days post-infection (~48-fold increase over the day 0 value). A corresponding analysis of *Coxiella* genomes in these samples shows that *limB* expression is highest during exponential phase growth of the bacterium (Fig. 5). Although transcription of *limB* is maximal during exponential phase growth (Fig. 5), the LimB protein can be detected in both LCVs and SCVs (Fig. 6). Interestingly, FW blots and immunoblots prepared from 12.5 % acrylamide gels revealed two distinct molecular mass forms of native LimB, with a larger (~8.7 kDa) LimB species dominant in LCVs and an ~7.2 kDa isofrom of LimB in SCVs (28 days post-infection) (Fig. 6). Not surprisingly, mixed-cell populations of the bacterium possess both LimB isoforms (Fig. 6a and b, lane 1). Although the two LimB isoforms can be distinguished as a doublet or smear on blots prepared from gradient gels (e.g.

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**Fig. 3.** rLimB binds Ni-HRP. FW blot prepared from a SDS-PAGE gel (12.5 % acrylamide; w/v) containing separated proteins (20 µg) of whole-cell lysates of *C. burnetii* (mixed-cell population; lane 1), *E. coli* (pMAL-c5x) (2 h, no IPTG; lane 2), *E. coli* (pMAL-c5x) (2 h, 1 mM IPTG; lane 3), *E. coli* (pJB202) (2 h, no IPTG; lane 4) and *E. coli* (pJB202) (2 h, 1 mM IPTG; lane 5). FW blot chemiluminescent signals from native LimB and rLimB binding of Ni-HRP are indicated by closed and open arrowheads, respectively. Molecular masses were determined from protein standards.
Fig. 1b), resolution of the bands was much clearer when blots were prepared from 12.5% acrylamide gels (Figs 6 and 7). The two LimB isoforms may reflect post-translational modification of LimB, and perhaps cleavage of LimB’s signal sequence and/or lipidation during the protein’s maturation.

**LimB is a surface-exposed omp**

The second residue of mature LimB is predicted to be Ala (Fig. 2). Previous work has shown that the second residue of mature lipoproteins plays an important role in coordinating transport of the protein to its final destination in the cell, thus it is referred to as an envelope localization residue (ELR) (Yamaguchi *et al.*, 1988). Since lipoproteins with an uncharged ELR are translocated to the outer membrane of Gram-negative bacteria, we hypothesized that LimB was surface-exposed in *C. burnetii.* To address the hypothesis, we used three different methods. First, we prepared a Sarkosyl-insoluble fraction of *Coxiella,* a method commonly used to purify omps of Gram-negative bacteria, previously employed for this purpose with *C. burnetii* (Zhang *et al.*, 2005). Analysis of Sarkosyl fractions of a mixed-cell population of *Coxiella* by FW blots and immunoblotting

![Fig. 4. Ni-HRP binding by rLimB is inhibited by EDTA, imidazole and divalent metal cations. (a) FW blots were done in parallel and prepared from the same SDS-PAGE gel (12.5% acrylamide; w/v) using whole-cell lysates of *C. burnetii* (mixed-cell population; 20 μg protein per lane). Blots were incubated in the presence of EDTA, imidazole (IMID) or divalent metal cations, as described in Methods. The resulting FW blots demonstrate a marked reduction in rLimB binding of Ni-HRP in the presence of Co²⁺ or Fe³⁺ (<50% of control), as shown by the reduced chemiluminescent signal (arrow). Binding signal was not detectable on blots incubated with EDTA or Zn²⁺ and was only slightly detectable (<5% of control) if imidazole (IMID) or Ni²⁺ were present. (b) Densitometric analysis of the blots in (a), shown as a percentage of the signal density of the untreated control (None). The assay is representative of two independent sets of experiments.](image)

![Fig. 5. *limB* expression is highest during exponential phase growth of *C. burnetii* in synchronous infections of cultured Vero cells. An analysis of *limB* transcript quantities (histogram) relative to the number of genomes (● and line) was determined by qRT-PCR and qPCR, respectively, over the course of an 8 day synchronous infection of Vero cells, as described in Methods. Values represent the means ± SD from six independent determinations. Asterisks denote a statistically significant increase in *limB* mRNA relative to 0 days (*P*<0.05).](image)
with anti-rLimB antibodies showed that: (i) both molecular mass forms of LimB were present in the insoluble fraction but absent in the Sarkosyl-soluble fraction, (ii) Ni-HRP was bound by both LimB isoforms and (iii) both protein isoforms were recognized by anti-LimB antibodies (Fig. 7). These data suggest that LimB is an omp.

Second, we employed immunoelectron microscopy with anti-rLimB or control antibodies and live bacteria. Resulting electron micrographs showed uniform protein A–gold binding on the surface of Coxiella treated with either anti-LimB or anti-Com1 (positive control) antibodies (Fig. 8a). In contrast, treatment with anti-MBP antibodies (a control for the fusion partner of the rLimB immunogen), anti-EF-Ts (a control for leaked intracellular proteins) or rabbit PIS from the same animal used to generate anti-rLimB antibodies, showed minor and inconsistent protein A–gold binding to the bacteria (Fig. 8a).

Finally, we employed these antibodies and PIS in immunofluorescence microscopy with Coxiella cells that had been fixed with paraformaldehyde. This procedure was previously employed to detect surface-exposed proteins of Ehrlichia and Anaplasma species (Ge & Rikihisa, 2007). Results of these experiments correlated with the immunoelectron microscopy data, and showed prominent immunofluorescence when Coxiella was treated with anti-rLimB or anti-Com1 antibodies, and minor, sporadic immunofluorescence when negative-control antibodies or PIS were used (Fig. 8b). Taken together, the Sarkosyl insolubility and detection of LimB on the surface of live cells by immunoelectron microscopy and on paraformaldehyde-fixed cells by immunofluorescence microscopy strongly suggest that LimB is a surface-exposed omp in Coxiella.

**Discussion**

In this study, we present data on the discovery and partial characterization of Coxiella LimB, a surface-exposed omp that binds divalent metal cations, including nickel and zinc with high affinity, and cobalt and iron with relatively lower affinity. Although assigning specific roles to LimB requires additional investigation regarding the protein’s structure and ligand-binding characteristics, the results of this study allow us to formulate reasonable hypotheses. First, maximal expression of the limB gene occurs during exponential phase growth, suggesting that LimB and its ligand(s) may be important for replication. Second, LimB’s surface location suggests that its metal ligand is obtained extracellularly. Third, because the replication of Coxiella in nature is restricted to an acidified, lysosome-like
compartment of the host cell, termed a parasitophorous vacuole (PV) (Voth & Heinzen, 2007), it is logical to assume that the ligand(s) is present in the PV niche. Fourth, it is plausible that LimB activity in the PV generates an active competition with the host cell, especially if the ligand is limiting in concentration. Based upon these results, we hypothesize that LimB is a lipoprotein that serves as a receptor for a divalent metal cation(s) at the surface of the pathogen during active replication within host cells. Considering that bio-reactive metal acquisition is often key to bacterial replication (Ratledge & Dover, 2000), LimB may play a role in the pathogenesis of C. burnetii.

Gram-negative bacteria employ a set of highly conserved proteins to process and transport lipoproteins (Babu et al., 2006; Hantke & Braun, 1973). Prolipoprotein precursors are synthesized in the cytosol and then translocated through the cytosolic membrane via a Sec translocon [or Tat for proteins with a type I signal sequence (Hutchings et al., 2009)]. Once across, a phosphatidylglycerol:proliprotein diacylglyceryl transferase (Lgt) catalyses addition of a DAG moiety to the thiol group of the lipobox Cys residue. Subsequently, a lipoprotein-specific signal peptidase (Lsp; also called signal peptidase II) cleaves the signal peptide at the N-terminal side of lipobox Cys. Finally, a phospholipid:apolipoprotein transacylase (Lnt) covalently attaches a fatty acid to the amide group of the N-terminal Cys to create the mature lipoprotein (Tokuda, 2009) (see Fig. 2). The ELR at position 2 of mature lipoproteins plays an important role in coordinating the protein’s transport to its final destination in the cell (Yamaguchi et al., 1988). Namely, mature lipoproteins that possess a charged ELR (especially Asp) are retained in the cytosolic membrane during transport, whereas those with an uncharged ELR (e.g. the Ala of LimB; Fig 2) are translocated to the outer membrane with the assistance of Lol sorting-system proteins in the periplasm (Tokuda, 2009; Yamaguchi et al., 1988). Based upon characteristics of the predicted LimB protein (Fig. 2) we hypothesize that pro-LimB follows a typical Sec-dependent (type II signal peptide) pathway wherein its 20-residue type II signal sequence would be cleaved by an LspA-like protein, its resulting N-terminal Cys is lipidated by Lgt and Lnt homologues, and the Ala ELR helps coordinate translocation of mature LimB to the outer membrane with the assistance of a Lol-like sorting-system. In fact, the surface location determined for LimB (Figs 7 and 8) is in keeping with the hypothesized outer membrane location of LimB, as predicted from its Ala ELR (Fig. 2).

Although the exact role of LimB in the biology of Coxiella remains to be determined, its demonstrated surface exposure and potentially chimeric nature (a protein with lipids) would be attractive features for a subunit vaccine candidate. In addition, the biosynthetic pathway of LimB might also provide an alternative strategy for therapeutic intervention during chronic Q fever; specifically, lipoprotein secretion could conceivably be inhibited by drugs like globomycin. Globomycin, a cyclic pentapeptide antibiotic, has been shown to inhibit secretion of lipoproteins in a wide variety of pathogens by inhibiting Lsp function (Geuken et al., 2006; Kiho et al., 2004; Rahman et al., 2007). Further investigation of LimB and other surface-exposed metal-binding proteins of Coxiella is expected to yield clues regarding the poorly characterized pathogenesis of this highly infectious agent. An illustration of LimB lipoprotein maturation as well as several hypothetical functions are provided in Supplementary Fig. S1, available with the online version of this paper.

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