Methylation and in vivo expression of the surface-exposed *Leptospira interrogans* outer-membrane protein OmpL32

Azad Eshghi,¹ Marija Pinne,²,³ David A. Haake,³,⁴ Richard L. Zuerner,⁵ Ami Frank⁵ and Caroline E. Cameron¹

¹Department of Biochemistry and Microbiology, University of Victoria, Victoria, BC, Canada
²Research Service, 151, Veterans Affairs Greater Los Angeles Healthcare System, Los Angeles, CA, USA
³Department of Medicine, David Geffen School of Medicine at University of California Los Angeles, Los Angeles, CA, USA
⁴Division of Infectious Diseases, 111F, Veterans Affairs Greater Los Angeles Healthcare System, Los Angeles, CA, USA
⁵Infectious Bacterial Diseases Research Unit, National Animal Disease Center (NADC), Agricultural Research Service (ARS), United States Department of Agriculture (USDA), Ames, IA, USA

Recent studies have revealed that bacterial protein methylation is a widespread post-translational modification that is required for virulence in selected pathogenic bacteria. In particular, altered methylation of outer-membrane proteins has been shown to modulate the effectiveness of the host immune response. In this study, 2D gel electrophoresis combined with MALDI-TOF MS identified a *Leptospira interrogans* serovar Copenhageni strain Fiocruz L1-130 protein, corresponding to ORF LIC11848, which undergoes extensive and differential methylation of glutamic acid residues. Immunofluorescence microscopy implicated LIC11848 as a surface-exposed outer-membrane protein, prompting the designation OmpL32. Indirect immunofluorescence microscopy of golden Syrian hamster liver and kidney sections revealed expression of OmpL32 during colonization of these organs. Identification of methylated surface-exposed outer-membrane proteins, such as OmpL32, provides a foundation for delineating the role of this post-translational modification in leptospiral virulence.

INTRODUCTION

Pathogenic *Leptospira* are the causative agents of the emerging zoonotic disease leptospirosis (Levett, 2001). Rats and house mice are asymptomatic carriers of *Leptospira* and serve as both reservoir (Bharti *et al.*, 2003; Ko *et al.*, 2009; Levett, 2001) and vector for the maintenance and transmission of bacteria to humans. Transmission to humans is thought to occur via contact with water contaminated by the urine of reservoir hosts (Levett, 2001). Leptospirosis is highly prevalent in developing nations with urban slums (Ko *et al.*, 2009; McBride *et al.*, 2005), and is endemic in settings where farmers that conduct their work manually are exposed to animals shedding leptospiral pathogens (Bharti *et al.*, 2003; Ko *et al.*, 2009; Levett, 2001). Disease symptoms in humans most commonly include uveitis (Verma *et al.*, 2008), and hepatic (Merien *et al.*, 1998) and renal (Seguro *et al.*, 1990) dysfunction. The most recent WHO statistics report more than 500 000 severe human leptospirosis cases per year with mortality rates greater than 10 % (WHO, 1999).

Understanding pathogenicity mechanisms surrounding the leptospiroid infection process has been hampered by the lack of targeted genetic tools available to leptospiral researchers, such as signature tagged mutagenesis (STM) (Hensel *et al.*, 1995) and transposon site hybridization (TraSH) (Sassetti *et al.*, 2001). Random transposon mutagenesis, while not as robust and efficient as STM and TraSH, has been successfully employed to identify virulence factor-encoding genes of leptospiral pathogens (Bourhy *et al.*, 2005). To date, five...
of six genes encoding virulence factors have been identified in this manner, including lat0222 in *Leptospira interrogans* serovar Lai, which encodes the outer-membrane lipoprotein Loa22 (Ristow et al., 2007), la2613 in serovar Lai, which encodes the flagella motor switch protein FliY (Liao et al., 2009), an orthologue of serovar Lai lb186 in *L. interrogans* serovar Manilae, which encodes the haem oxygenase HemO (Murray et al., 2009), and two genes which encode LPS biosynthesis proteins in serovar Manilae, with one being orthologous to serovar Lai la1641 and the other having no observed serovar Lai orthologue (Murray et al., 2010). Lastly, *invA*, a Nudix hydrolase specific for numerous dinucleoside oligophosphate substrates, has been directly inactivated using insertion mutagenesis and demonstrated to be required for virulence of serovar Lai in hamsters (Luo et al., 2011). In addition to these genetic approaches, various protein-based approaches have been utilized to identify potential leptospiral virulence factors. Such studies have elucidated the subcellular location for various proteins (Beck et al., 2009; Cullen et al., 2005; Haake & Matsunaga, 2002; Monahan et al., 2008; Pinne & Haake, 2009; Sakolvaree et al., 2007), identified immunoreactive proteins (Artiushin et al., 2004; Guerreiro et al., 2001; Sakolvaree et al., 2007), quantified absolute protein numbers per bacterium (Malmström et al., 2009), provided evidence for differential protein expression in response to changes in microenvironments (Cullen et al., 2002; Eshghi et al., 2009; Lo et al., 2009; Matsunaga et al., 2007; Nally et al., 2001a, b, 2007; Velineni et al., 2006), contributed to the identification of the first confirmed leptospiral virulence factor, Loa22 (Nally et al., 2007), and revealed diverse and extensive post-translational modifications (PTMs) (Cao et al., 2010).

Of particular interest, a recent global proteomic analysis of *L. interrogans* serovar Lai identified a total of 155 methylated proteins (Cao et al., 2010), suggesting a widespread role for methylation in *L. interrogans* protein function and/or regulation. In the context of virulence, methylation has been shown to be an essential PTM in various bacterial pathogens. Using genetic approaches, native methylation has been shown to be required for regulation of transcription and phase variation of outer-membrane components (Deitsch et al., 2009), and for maintenance of virulence (Giacomodonato et al., 2009). Methylation of glycolipids has been shown to be essential for *Mycobacterium avium* virulence in mice (Krzywinska et al., 2005), while methylation of proteins has been shown to alter the antigenicity of the outer-membrane protein OmpB from *Rickettsia typhi* (Chao et al., 2008), and to alter both the antigenicity of and the host T cell-mediated immune response against the heparin-binding haemagglutinin from *Mycobacterium tuberculosis* (Parra et al., 2004; Temmerman et al., 2004). In addition, methylation has been demonstrated to be essential for functional type III secretion, and thus virulence, in *Yersinia pseudotuberculosis* (Garbom et al., 2004, 2007), and methylation of the surface protein OmpB in *Rickettsia prowazeki* has been suggested to be central to the pathogenesis of that bacterium (Chao et al., 2004, 2007). Collectively, these investigations highlight the importance of methylation of outer-membrane surface components in bacterial virulence.

The present study describes the proteomic identification of the novel *L. interrogans* outer-membrane protein OmpL32 (LIC11848). Immunological analyses confirmed exposure of OmpL32 on the leptospiral surface and expression of this protein during the course of bacterial infection. Proteomic analysis revealed differential and extensive glutamic acid methylation of this protein under all the tested environmental conditions. The potential implications of this PTM are discussed within the context of leptospiral virulence.

**METHODS**

*Leptospira* and culture conditions. *L. interrogans* serovar Copenhageni strain Fiocruz L1-130 is a clinical isolate originating from Salvador, Brazil (Ko et al., 1999). *L. interrogans* serovar Pomona type kennedwicki strain RM-211 is an isolate from a swine abortion case (Thiermann et al., 1984). Cultures were maintained in Ellinghamausen and McCullough (Ellinghamausen & McCullough, 1965) as modified by Johnson and Harris (Johnson & Harris, 1967) (EMJH) medium at 29.5 °C. Bacterial enumeration and media shift experiments were performed as previously described (Eshghi et al., 2009). In these experiments, *L. interrogans* was grown under the following four conditions: (1) in EMJH medium at 37 °C; (2) in EMJH medium depleted of iron; (3) in EMJH medium supplemented with 10 % fetal bovine serum (FBS); or (4) in EMJH medium supplemented with 10 % FBS and depleted of iron.

2D gel electrophoresis (2DGE) and MALDI-TOF MS experiments. *L. interrogans* was harvested at 8500 g and washed twice with PBS supplemented with 5 mM MgCl₂. Cell pellets were lyophilized, weighed and resuspended to a concentration of 2.5 mg ml⁻¹. 2DGE, staining of gels with colloidal Coomassie blue and MALDI-TOF MS experiments were performed as previously described (Eshghi et al., 2009). Briefly, 2D gels (pH range 3–11) containing equal amounts of *Leptospira* by dry weight from each of the four different growth conditions were run in triplicate. Gels originating from the four different growth conditions were compared to detect proteins displaying altered intensity between the comparative conditions, while validation of protein spot intensity within a growth condition was obtained by comparison of matched triplicate gels.

Liquid chromatography-electrospray ionization-tandem MS (LC-ESI-MS/MS). Trypsin digests were performed with Genomic Solutions ProGest (Digilab). Briefly, gel slices were manually cut into 1 mm cubes and transferred to a Genomics Solutions ProGest perforated digestion tray. The gel pieces were de-stained (50:45:5, v/v, methanol/water/acetic acid) prior to reduction and alkylation with 10 mM DTT and 100 mM iodoacetamide, respectively (Sigma). Modified sequencing grade porcine trypsin solution (20 ng µl⁻¹, Promega) was added to the gel slices at an enzyme : protein ratio of 1:50. Proteins were digested for 5 h at 37 °C prior to collection of resulting peptides and acid extraction of the gel slices (50:40:10, v/v/v, acetonitrile/water/formic acid). Samples were then Speed Centrifuged to dry and stored at −20 °C until analysed by LC-ESI-MS/MS. An UltiMate Nano HPLC (LC Packings/Dionex) coupled to an Applied Biosystems/MDS Sciex QSTAR Pulsar I hybrid quadrupole-TOF LC-MS/MS mass spectrometer (AB Sciex) was used to perform the LC-MS/MS analyses. The nano-column was connected to a 20 µm internal diameter emitter tip (New Objective, Inc.)
positioned at the orifice plate of the mass spectrometer with spray established by applying a tip voltage of 1900 V to the pre-nano column via a platinum wire nano-tie high-voltage connection.

The lyophilized samples were rehydrated in 40 μl 2% (v/v) acetonitrile/0.1% formic acid, and 15 μl of each sample was injected using a FAMOS Autosampler (LC Packings/Dionex) for each analysis. The samples were concentrated/desalted using a SwitchOSII loading pump (LC Packings/Dionex) with 98% solvent A (2%, v/v, acetonitrile, 0.1% formic acid) at a rate of 30 μl min⁻¹ for 10 min over an Agilent Zorbax C18-SB trap column (5 x 0.3 mm) (Agilent Technologies) to protect the in-house-prepared nano-analytical column Magic C18 AQ resin, 150 mm x 75 μm column diameter (Michrom Bioresources). After the 10 min loading period, the trap column was switched in-line with the nano-column gradient flow. The following 1 HPLC analytical separation was performed at 300 nl min⁻¹ [33 min linear gradient from 5–60% solvent B (98%, v/v, acetonitrile, 0.1% formic acid), 2 min linear gradient from 60–75% solvent B, 5 % solvent B over 3 min and re-equilibrated with 5% solvent B for 12 min before the next injection].

Mass spectra analyses were acquired by collecting a 1 s TOFMS survey scan of 400–1200 m/2 followed by two 2.5 s product ion scans in the 100–1500 m/z mass range. MS/MS spectra were acquired in a data-dependent manner, selecting the top two most intense eluting ions in the 400–1200 m/z range with a 2⁺ to 4⁺ charge state greater than 20 counts. Following selection for MS/MS analysis, precursor ions were excluded from selection for MS/MS analysis for 180 s. A rolling collision energy for fragmentation was selected based on the precursor ion mass using the formula 0.05 m/z+5 V. Known keratin tryptic and trypsin autolysis product masses were excluded to prevent these contaminant ions from being selected for fragmentation.

Mass spectrometer parameters used were as follows: declustering potential setting of 50, focusing potential setting of 220, curtain gas setting of 25, and CAD gas setting of 5 with nitrogen in the collision cell.

**Bioinformatic analyses.** Peptide mass fingerprint (PMF) searches were conducted as previously described (Eshghi et al., 2009), with the following change: no variable modifications were selected. PMFs were further analysed for potential PTMs using the ExPASy Proteomics Server FindMod tool (Wilkins et al., 1999) (http://web.expasy.org/findmod/). Predictions were conducted with the following parameter settings: ion mode was set to Victoria-Genome BC Proteomics Centre. Search parameters used for prediction of subcellular localization, the LIC11848 amino acid sequence was analysed for the presence of a putative signal peptide sequence using the Liposcan 1.0 (Rahman et al., 2008) (http://www.cbs.dtu.dk/services/LipoP), SignalP (Emanuelsson et al., 2007) (http://www.cbs.dtu.dk/services/SignalP/) signal peptide prediction programs. Secondary structure analyses for prediction of β-sheets was conducted using GOR (Kloczkowski et al., 2002; Sen et al., 2005) (http://psib.ictbull.com/hfm/index.php), and homology predictions to Gram-negative outer-membrane proteins were conducted using OMPdb (Tsirigos et al., 2011) (http://aias.biol. uoa.gr/OMPdb/index.php). Transmembrane domain prediction was conducted using the hidden Markov model outer-membrane β-barrel protein prediction program PRED-TMBB (Bagos et al., 2004) (http://biophysics.biol.uoa.gr/PRED-TMBB/). B cell epitope predictions were made using the BCpreds B cell epitope prediction server (El-Manzalawy et al., 2008), using the search parameter ‘fixed length epitope prediction’ and epitope lengths of 12, 14, 20 or 22 aa (http://alab.cs.iastate.edu/bcpreds/predict.html).

**Recombinant protein expression and purification.** ORF LIC11848 was PCR-amplified from *L. interrogans* serovar Copenhageni strain Fiocruz L1-130 genomic DNA using the primer pair 5’-CTAGAGCCATATGTCGAGCCGATCCTAAACT-3’ and 5’-GTCACTCTGAGTTAGTAGGGAGGATTT-3’ (incorporated restriction sites indicated by underlined nucleotides). The LIC11848 amplicon (693 bp in length encompassing basepairs 70–762 of the ORF) was ligated first into the cloning vector pET1 (Clonejet, Fermentas) and digested with *Nde*I and *Xhol*, followed by ligation into a similarly digested PET28a expression vector (Novagen). The sequence and reading frame of the expression construct were verified by DNA sequencing with vector-specific primers. The LIC11848/PET28a construct was transformed into the *Escherichia coli* expression strain BL21 Star BL21(DE3) (Invitrogen). Bacteria were grown at 37 °C in a shaking incubator, and recombinant expression was induced at OD600 1.0 using a final concentration of 0.4 mM IPTG (Invitrogen). The incubation temperature was then reduced to 16 °C and bacteria were grown overnight.

Soluble recombinant protein was purified using the following methodology. Bacteria were harvested by centrifugation at 3000 g, resuspended in binding buffer (60 mM NaH2PO4, 500 mM NaCl and 20 mM imidazole, pH 8.0) and lysed via sonication (3 x 30 s) in the presence of a protease inhibitor cocktail [Protease Inhibitor Cocktail Set III; 100 mM 4-(2-aminoethyl) benzensulfonyl fluoride hydrochloride (AEBSF), 80 μM aprotinin, 5 mM bestatin, 1.5 mM E-64, 2 mM leupeptin, 1 mM pepstatin A (Calbiochem)]. The lysate was subsequently centrifuged at 20000 g at 4 °C for 30 min and the supernatant filtered through a 0.45 μm pore-size filter. Purification was performed on an AKTAprime Plus fast protein liquid chromatography system (GE Healthcare) fitted with a 1 ml HisTrap HP Column (GE Healthcare). All steps were performed at a flow rate of 1 ml/min⁻¹ and a pressure limit of 0.3 MPa. The run parameters included a 12 ml equilibration step with binding buffer, application of 30 ml of the filtered supernatant (protein sample), a 40 ml wash with binding buffer, and a gradient elution of 0–100% elution buffer (60 mM NaH2PO4, 500 mM NaCl and 500 mM imidazole, pH 8.0) over a 50 ml volume with fractions collected at 0.5 ml increments. Fractions containing recombinant OmpL32 (rOmpL32) (identified via *A*α*β* readings of fractions in combination with SDS-PAGE) were then combined and concentrated using a 10K Amicon Ultra-4 centrifugal filter unit (Millipore).

Desalting was achieved in HEPES buffered saline (20 mM HEPES, 150 mM NaCl, pH 7.4), offering 90 ml, via an AKTAPrime Plus fast
protein liquid chromatography apparatus fitted with a HiLoad 16/60 Superdex 75 prep grade gel filtration column (GE Healthcare). Purified rOmpL32 was quantified using a BCA Protein Assay (Pierce).

**Antibodies.** Polyclonal rabbit antiserum was prepared against recombinantly expressed LIC11848 (rLIC11848) by ProSci, Inc. Polyclonal rabbit antisera against OmpL36 (Eshghi et al., 2009), FlaA1 (Cullen et al., 2005) and LipL32 (Haake et al., 2000) have been described previously.

**Immunoblot analysis.** Approximately $1 \times 10^8$ *Leptospira* were harvested at 10,000 g and washed twice with wash buffer (PBS and 5 mM MgCl₂). The pellet was resuspended in SDS-PAGE loading buffer, boiled for 10 min and separated on a 15% SDS-polyacrylamide gel. Proteins were electrophoretically transferred to an Immobilon PVDF membrane (Millipore). Immunoblots were blocked with 2.5% milk powder in Tris-buffered saline, pH 7.4, with 0.05% Tween 20 (TBST). The membrane was washed with 1:1500 dilution of rLIC11848-specific rabbit antiserum or preimmune TBST, followed by incubation for 90 min at room temperature with a Kodak X-OMAT 2000A processor (Carestream Health). Membranes were exposed to X-ray film for 1 min and developed with a modification of incubating the membrane with the reagent for 1 min. The application as described by the manufacturer’s protocol with the modification of incubating the membrane with the reagent for 1 min. The membrane was exposed to X-ray film for 1 min and developed with a Kodak X-OMAT 2000A processor (Carestream Health).

**Immunofluorescence assay (IFA).** IFAs were conducted as previously described (Cullen et al., 2005; Pinne & Haake, 2011, 2009). Briefly, *L. interrogans* was harvested at 2000 g to maintain outer-membrane integrity, resuspended in PBS/5 mM MgCl₂ to a density of $5 \times 10^8$ cells ml⁻¹, applied in 1 ml aliquots to Nunc four-well Lab-Tek II Chamber Slides (Thermo Fisher Scientific) and incubated at 29.5 °C for 80 min for the purpose of adhering cells to the glass slides. All analyses were done in duplicate. Chamber slides were carefully aspirated, and bacteria were fixed using 1 ml per well of 2% (w/v) paraformaldehyde in PBS for 40 min at 29.5 °C. For the purpose of demonstrating an intact outer membrane, replicate chamber slides were permeabilized with 100% ice-cold methanol for 20 min at −20 °C. Chamber slides were then aspirated and blocked with 1 ml EMJH medium for 90 min at 29.5 °C. Rabbit antisera specific for OmpL36, rLIC11848 and FlaA1 were diluted 1:200, 1:100 and 1:600, respectively, and preimmune serum was diluted 1:100 and 1:200 dilutions, respectively. Normal goat sera block was used as a negative control (no primary antibody) on all sections from both infected and uninfected hamsters. Sections were washed with PBS to remove unbound antibody, and then incubated for 60 min at room temperature in the dark with a 1:5000 dilution of Alexa Fluor 488 F(ab’)₂ goat anti-rabbit secondary antibody and 0.4 μg DAPI ml⁻¹ (Invitrogen). Slides were mounted with Prolong Gold antifade reagent (Invitrogen). All images were captured on a Spot RT colour CCD camera mounted on a Nikon Eclipse E800. All immunohistochemistry images were captured under the same exposure conditions.

**RESULTS**

**L. interrogans** cells differentially express isoforms of a putative outer-membrane protein

To detect changes in protein expression resulting from exposure to varying environmental conditions, *L. interrogans* cells were grown at 37 °C in EMJH medium, in EMJH medium depleted of iron, in EMJH medium supplemented with 10% FBS or in EMJH medium supplemented with 10% FBS and depleted of iron. Total protein was separated by 2DGE, gels were stained with colloidal Coomassie stain, and protein spots that were observed to vary in intensity between the compared conditions (see below) were selected and subjected to trypsin digestion and subsequent MALDI-TOF MS. Comparative analysis of triplicate gels of *Leptospira* grown under these varied culture conditions, using conventional growth at 37 °C in normal EMJH medium as the comparator, revealed consistently altered expression of proteins corresponding to ORFs LIC10361 (electron transfer flavoprotein EtfB), LIC13050 (putative glycosyl hydrolase), LIC13166 (putative coagulase), LIC11336 (GroES), LIC12210 (IbpA-1) and LIC11890 (periplasmic flagellin FlaB1), as previously reported (Eshghi et al., 2009). An additional protein identified through this study, corresponding to the ORF LIC11848, was deemed to be of particular interest due to the identification of multiple LIC11848 isoforms that displayed altered intensity and differed in their respective isoelectric points over the pl range 3.5–7.5 (Fig. 1). Visual comparison of triplicate gels prepared from each of the comparative growth conditions confirmed that the individual LIC11848 protein spots remained constant between replicates of a particular growth condition (Supplementary Fig. S1), indicating that altered protein intensities did not result from gel-to-gel variation.

http://mic.sgmjournals.org

Downloaded from www.microbiologyresearch.org by IP: 54.70.40.11
On: Tue, 08 Jan 2019 13:28:16
Due to the complexity associated with analysis of the comparative 2DGE patterns, a brief description of how the different LIC11848 isoforms were identified is provided. A visual identification of similar LIC11848 isoforms was made by locating protein spots that migrated to the same location in each of the comparative growth conditions. This allowed for the designation of representative spots that constituted each of the identified isoforms and corresponded to spot 1 [deemed to be the same as spots 4, 8 and 11, indicated by red circles in Fig. 1, panels (a), (b), (c) and (d), respectively], spot 3 [deemed to be the same as spots 6, 10 and 13, indicated by green circles in Fig. 1, panels (a), (b), (c) and (d), respectively] and spot 2 [deemed to be the same as spots 5, 9 and 12, indicated by blue circles in Fig. 1, panels (a), (b), (c) and (d), respectively]. Representative spots 1, 3 and 7, as well as spots 6 and 13 to ensure accurate prediction of identical isoforms, were subjected to in-gel tryptic digestion, the resulting peptides were analysed via MALDI-TOF MS, and PMFs were used to perform MASCOT database searches to allow protein identification.

The mass to charge ($m/z$) ratios, peptide coverage and expect values obtained for each of the representative spots (1, 3, 6, 7 and 13) used for identification via PMF are summarized in Supplementary Table S1. A high degree of peptide mass coverage was obtained for the representative spots, allowing for a definitive identification of each of the five protein spots as LIC11848. Spot 2, which is representative of spots 5, 9 and 12, was definitively identified as a LIC11848 isoform through subsequent MS/MS analyses (see below).

**LIC11848 is differentially methylated on glutamic acid residues**

To identify a plausible cause for the observed pI shift in LIC11848, PMF data were further analysed using the ExPASy FindMod tool (Wilkins et al., 1999) to predict potential PTMs. This analysis predicted methylation of a total of 11 glutamic acid residues within LIC11848; specifically, methylation was predicted for four peptides for spot 1, five peptides for spot 3, three peptides for spot 6, six peptides for spot 7 and eight peptides for spot 13. The number of predicted methylated peptides was proportional to the number of $m/z$ ratios used to conduct searches, with a higher degree of peptide coverage correlating with the identification of an increased number of methyl-ester adducts. To test these *in silico* predictions, tryptic digests of...
LIC11848 protein from representative spots 1, 2, 6 and 7 (Fig. 1a–c) from replicate second-dimension gels were subjected to online high-performance LC-ESI-MS/MS. These analyses confirmed in silico predictions, and revealed that a total of 11 glutamic acid residues were methylated within LIC11848, with experimental confirmation of methylation of four peptides in spot 1 (Fig. 2a and Supplementary Fig. S2), seven peptides in spot 2 (Fig. 2b and Supplementary Fig. S3), five peptides in spot 7 (Fig. 2c and Supplementary Fig. S4) and seven peptides from spot 6 (Fig. 2d and Supplementary Fig. S5). The glutamic acid residue at amino acid residue position 54 showed methylation in all four spots (Fig. 2 and Supplementary Figs S2–S5). Similar to the PMF analyses, the number of methyl esters identified was proportional to the number of m/z values present in the MS/MS searches, with a higher peptide coverage correlating with identification of more extensive methylation. Of particular note was the observation that methyl group location varied among glutamic acid residues within an identical peptide identified between comparative growth conditions. The peptide spanning amino acids 105–116 (VEVYDFIRDEER) was represented in all isoforms, with three of four isoforms displaying methylation on the most N-terminal glutamic acid residue (residue 2, counting from the N terminus of the peptide; Fig. 2b–d) and one of four isoforms displaying methylation on the most C-terminal glutamic acid (residue 11, counting from the N terminus of the peptide; Fig. 2a).

**Correlation between the observed LIC11848 protein methylation pattern and predicted B cell epitope locations**

Comparison of the identified LIC11848 methylated peptides with B cell epitopes predicted using the BCpreds online prediction tool revealed all 11 methylated glutamic acid residues to be contained within regions of the protein predicted as possible B cell epitopes (Table 1). Further, four of the predicted B cell epitopes correlated with methylated peptides contained within all four of the protein spots subjected to methylation profile analysis (predicted epitopes 43–54, 89–110, 103–122 and 173–192). With the exception of two predicted B cell epitopes, the scores for these analyses were highly significant (≥0.926).

**LIC11848 is a surface-exposed protein**

To predict the cellular location of LIC11848, the amino acid sequence was analysed using the signal peptide prediction tools Lipop P 1.0 (Rahman et al., 2008) and SignalP 3.0 (Emanuelsson et al., 2007). Both tools predicted the presence of an N-terminal Sp1 signal peptide, with Lipop...
Table 1. Predicted B cell epitopes present within the LIC11848 protein (OmpL32) and correlation with the observed methylation profile

<table>
<thead>
<tr>
<th>Predicted B cell epitope*</th>
<th>Probability</th>
<th>Predicted epitope length (aa)</th>
<th>Amino acid range within LIC11848</th>
<th>Methylated amino acids (residue numbers)</th>
<th>Protein spots</th>
</tr>
</thead>
<tbody>
<tr>
<td>ETERKLDEKIFE</td>
<td>0.28</td>
<td>12</td>
<td>43–54</td>
<td>43, 45, 54</td>
<td>1, 2, 6, 7</td>
</tr>
<tr>
<td>CTPAINQEDPANNCIRVEVYDF</td>
<td>0.992</td>
<td>22</td>
<td>89–110</td>
<td>106</td>
<td>1, 2, 6, 7</td>
</tr>
<tr>
<td>IRVEVYDFIRDERGLKKNV</td>
<td>0.959</td>
<td>20</td>
<td>103–122</td>
<td>106, 115</td>
<td>1, 2, 6, 7</td>
</tr>
<tr>
<td>EVMDRGPNTQPSHNDKEVF</td>
<td>0.961</td>
<td>20</td>
<td>173–192</td>
<td>173, 190</td>
<td>1, 2, 6, 7</td>
</tr>
<tr>
<td>FKQDNYPEYGRPETPAEGVVK</td>
<td>0.996</td>
<td>20</td>
<td>195–214</td>
<td>200, 205</td>
<td>1, 2, 6</td>
</tr>
<tr>
<td>SFKKEYIKHLDQF</td>
<td>0.741</td>
<td>14</td>
<td>231–244</td>
<td>235</td>
<td>2, 6</td>
</tr>
<tr>
<td>TKIYMDQLQGNENYKENVDAL</td>
<td>0.926</td>
<td>22</td>
<td>249–270</td>
<td>261</td>
<td>2, 6, 7</td>
</tr>
</tbody>
</table>

*Bold type indicates methylated glutamic acid residues.

Predicting a potential cleavage site between serine 20 and glutamine 21, and SignalP predicting a potential cleavage site between glycine 24 and serine 25, a finding which is consistent with a potential outer-membrane location for this protein. Secondary structure analysis conducted using GOR predicted the secondary structure composition of the protein to be 15% β-sheet. To further assess the predicted location of LIC11848, the amino acid sequence was searched against the OMPdb database for Gram-negative bacterial location of LIC11848, the amino acid sequence was searched against the OMPdb database for Gram-negative bacterial location, a finding which is consistent with a potential outer-membrane location for this site between glycine 24 and serine 25, a finding which is consistent with a potential outer-membrane location for this protein. Secondary structure analysis conducted using GOR predicted the secondary structure composition of the protein to be 15% β-sheet. To further assess the predicted location of LIC11848, the amino acid sequence was searched against the OMPdb database for Gram-negative bacterial β-barrel-containing outer-membrane proteins (Tsirigos et al., 2011). This search identified significant homology between LIC11848 and the C-terminal residues 633–752 from the Bacteroides fragilis outer-membrane receptor (OMR-TonB-dependent receptor) family (27.13% sequence identity and 44.97% sequence similarity over 129 aa). Further evidence suggesting a membrane association for this protein was obtained via identification of nine transmembrane domains as predicted by PRED-TMBB analysis, although the generated score for this analysis of 3.020 was slightly higher than the outer-membrane protein threshold cut-off value of 2.965 for this prediction program.

To experimentally evaluate whether LIC11848 protein is present on the surface of L. interrogans, IFA (Cullen et al., 2005; Pinne & Haake, 2009, 2011) was conducted using intact and methanol-permeabilized L. interrogans with rLIC11848-specific serum, serum specific for the known surface-exposed protein OmpL36 (positive control) (Pinne & Haake, 2009) and serum specific for the periplasmic endoflagellar protein FlaA1 (negative control). Immunofluorescence microscopy revealed reactivity to intact L. interrogans with serum specific for rLIC11848 and OmpL36, but not with preimmune serum or serum specific for FlaA1 (Fig. 3a). Immunofluorescence was also detected in permeabilized Leptospira using serum specific for LIC11848, OmpL36 and FlaA1, but not preimmune serum (Fig. 3b). The specificity of the rLIC11848-specific antiserum was tested via immunoblot analysis. This analysis revealed reactivity to a protein band migrating at a molecular mass corresponding to the 32 kDa predicted molecular mass for LIC11848 (Supplementary Fig. S6).

Immunoblot analysis using preimmune serum showed an absence of reactivity.

**Leptospira express LIC11848 during colonization of hamster kidneys and liver**

To assess expression of LIC11848 during the infection process, immunofluorescence microscopy was conducted on kidney and liver sections of healthy uninfected and Leptospira-infected hamsters. Use of antiserum specific for LipL32 (positive control) or rLIC11848 in these experiments resulted in leptospiral fluorescence in both kidney and liver sections of hamsters infected with Leptospira (Fig. 4a–d). Although leptospires were observed throughout the tissue of infected hamsters, the level of LIC11848 protein expression was reduced in comparison with the highly conserved and abundant outer-membrane lipoprotein LipL32. This observation was in agreement with absolute protein numbers per Leptospira reported using selective reaction monitoring (Malmström et al., 2009), and more similar in protein expression levels to LipL46 (data not shown) in serially sectioned tissue. In kidney sections, the spirochaetes were mainly observed disseminated in the blood vessels and in the interstitium between tubules. In liver sections, the leptospires were observed within the intercellular spaces between hepatocytes. Use of antiserum specific for rLIC11848 on healthy (not infected with Leptospira) hamster kidney and liver sections did not result in fluorescence (Fig. 4e, f). Sections of Leptospira-infected hamster kidney and liver tissue incubated with secondary antibody alone did not display any fluorescence, confirming that the detected fluorescence was specific for the leptospiral proteins and did not result from non-specific staining from the fluorescent secondary antibody (Fig. 4g, h).

**DISCUSSION**

The results presented herein enhance our understanding of L. interrogans biology and serve as a platform for the study...
of leptospiral virulence. In this study, proteome analyses conducted on *L. interrogans* grown under the varying environmental conditions of media depleted in iron, media supplemented with 10% serum or media depleted of iron and containing 10% serum (*in vivo* growth conditions) identified the novel protein LIC11848. Immunofluorescence experiments revealed reactivity of serum specific for the 32 kDa LIC11848 protein product with intact *L. interrogans*, thus providing evidence for the exposure of this protein on the surface of viable *L. interrogans*. This finding, combined with the bioinformatic predictions that this protein possesses a cleavable SpI signal sequence and may have a propensity to form a β-barrel structure, prompted our designation of this protein as OmpL32. The present study also demonstrated by immunohistochemistry the expression of OmpL32 during leptospiral infection of the kidneys and liver of hamsters, correlating with a prior report of OmpL32 expression in virulent *Leptospira* freshly cultured from the liver and kidney of infected hamsters (Vieira et al., 2009). The level of OmpL32 expression compared with LipL32 in *in vivo* was in agreement with an earlier proteomic study that utilized selective reaction monitoring and demonstrated an 11-fold higher expression level of LipL32 (32,190 copies per cell) compared with OmpL32 (2,740 copies per cell) (Malmström et al., 2009) in *Leptospira* grown under laboratory conditions. Finally, and most importantly, proteomic analysis of multiple isoforms of OmpL32 revealed the interesting observation of differential glutamic acid methylation within this surface-exposed protein.

To date, glutamic acid carboxymethylation in bacteria has been discussed within the context of chemotaxis (Ahlgren & Ordal, 1983; Kehry et al., 1984; Kleene et al., 1977; Nishiyama et al., 1999). Specific carboxymethylation on glutamic acid residues serves to modulate protein function from an active to an inactive conformation or vice versa. Activation via methylation results in conformational changes of the methylated protein, leading to activation of downstream proteins that eventually activate flagella for directional motility (Clarke, 1993). The discovery that OmpL32 displays differential methylation suggests that a related methylation-induced regulation may occur with this leptospiral protein. However, two observations provide evidence suggesting that OmpL32 does not play a role in bacterial chemotaxis. First, the OmpL32 amino acid sequence does not reveal homology to proteins involved in either chemotaxis or motility, which in general are highly homologous to one another. And second, glutamic acid methylation of chemotaxis proteins has been demonstrated to occur on only a few targeted amino acid residues (Rice & Dahlquist, 1991), which is in distinct contrast to the widespread methylation pattern observed for OmpL32. These observations suggest that the glutamic acid-specific methylation pattern detected in OmpL32 has an alternative functional role.

Multiple studies have established a requirement for methylation in bacterial pathogenesis. A requirement for methylation has been demonstrated in *Y. pseudotuberculosis*, where an insertion mutant deficient in the VagH protein is avirulent in mice (Garbom et al., 2004). In *Y. pseudotuberculosis*, the VagH protein is homologous to the *E. coli* HemK protein (Garbom et al., 2007), an N2-methyltransferase (Heurgué-Hamard et al., 2002; Nakahigashi et al., 2002). The *Y. pseudotuberculosis* vagH insertion mutant demonstrated repression of secretion of the virulence determinant YopD via the type III secretion system (Garbom et al., 2007). Additionally, this mutant displayed characteristics similar to a type III secretion mutant during pathogenesis, suggesting a role for protein methylation in type III secretion function (Garbom et al., 2007). Within the context of outer-membrane protein methylation and bacterial pathogenesis, a link has been demonstrated in the bacterial pathogen *R. prowazekii*. The virulent Madrid E strain has a null mutation in the methyltransferase-encoding rp027 gene (Chao et al., 2007), and consequently shows hypomethylation of the immunodominant outer-membrane protein B (OmpB). In comparison, the virulent *R. prowazekii* Breinl strain, which expresses the Rp027 protein product, exhibits hypermethylation of OmpB (Chao et al., 2004), suggesting a role in virulence for outer-membrane protein methylation in *R. prowazekii* (Chao et al., 2007).

Methylation of bacterial outer-membrane proteins could contribute to virulence in at least two ways. First, methylation could potentially be used by bacteria to regulate protein function, in a manner analogous to that seen for chemotaxis proteins (Wadhams & Armitage, 2004) and the *Y. pseudotuberculosis* type III secretion system (Garbom et al., 2007). In this scenario, methylation could act to regulate the function of leptospiral virulence factors present on the spirochaetal surface by providing a post-translationally controlled switch between an active/inactive state, and in this way ensure optimal timing of virulence protein function within the infection process. Also, the presence or absence of protein methylation could be critical for real-time phase variation as a method of altering the antigenicity of bacterial outer-membrane proteins, thereby modulating the host immune response recognition of these proteins. Support for the existence of effects on antigenicity and their influence on cellular and humoral immune responses to bacterial pathogens comes from studies conducted with the pathogens *M. tuberculosis* and *R. typhi*. In *M. tuberculosis*, correlation of the level of protein methylation with host immune response modulation has been demonstrated (Temmerman et al., 2004), with native methylation of the surface-exposed *M. tuberculosis* heparin-binding haemaglutinin protein being required for an effective T cell-mediated immune response. The effect of protein methylation on the host humoral immune response has been studied in *R. typhi* using methylated and non-methylated protein fragments of the outer-membrane protein OmpB (Chao et al., 2008). In that study experiments conducted using patient sera in an ELISA-based format revealed higher titres to the methylated OmpB fragment when compared with the unmethylated fragment. Together these studies provide
evidence that methylation of surface-exposed proteins in bacteria directly affects pathogen recognition by the host immune response.

In order for OmpL32 to function in a methylation-dependent, immune evasion capacity within Leptospira, the protein must be present on the surface of the bacterium.

Immunofluorescence microscopy experiments confirmed surface exposure of this protein in L. interrogans serovar Copenhageni strain Fiocruz L1-130. The presentation of this protein on the leptospiral surface would facilitate direct recognition by the host immune response. The capacity of the bacterium to differentially methylate OmpL32 could contribute to the ability of this bacterium to persist in the

**Fig. 3.** Immunofluorescence microscopy identifies the LIC11848 protein as a surface-exposed, outer-membrane protein. Intact (a) and methanol-permeabilized (b; following page) L. interrogans was probed with rabbit preimmune serum or rabbit serum specific for OmpL36 (outer-membrane protein positive control), rLIC11848 or FlaA1 (periplasmic negative control protein). The observed fluorescence corresponds to either Alexa Fluor 568-labelled leptospires or DAPI-stained leptospiral DNA. Merged images reveal the overlap of DAPI-stained and Alexa Fluor 568-labelled Leptospira.
presence of the generated host immune response, and thus establish a chronic infection. In this scenario OmpL32 methylation could serve to modulate the level of the host immune response in a manner similar to that documented for M. tuberculosis (Temmerman et al., 2004) and R. typhi (Chao et al., 2008). In support of this we have observed different isoforms within an individual growth condition, suggesting that the extent of methylation differs between OmpL32 isoforms. Further, the direct identification of multiple different locations of methylation on amino acid residues within OmpL32 provides an additional indication that this process may represent a novel mechanism that would alter the surface of the bacterium. A definitive answer as to whether Leptospira cells alter the number of methylated residues within OmpL32 upon exposure to a host environment must await further investigation. However, in this study we have provided a foundation for future studies by showing that an outer-membrane protein is methylated and that the methylation pattern of this protein varies, resulting in multiple isoforms. An alternative question arising from these observations is whether the position of methylation alone rather than the absolute number of methylations leads to altered recognition by the host.
Fig. 4. *Leptospira* cells express LIC11848 protein during colonization of hamster kidneys and liver, as evidenced by immunofluorescence microscopy. *Leptospira*-infected tissue samples from golden Syrian hamsters were probed with LipL32 (positive control) or rLIC11848 antisera and viewed by indirect fluorescence microscopy (Matsunaga et al., 2006). Fluorescence (green areas) was detected using both LipL32 and rLIC11848 antisera with infected kidney [(a) and (c), respectively] and liver [(b) and (d), respectively] tissue sections but not in healthy uninfected kidney (e) or liver (f) tissue sections using rLIC11848 antiserum. Use of secondary antibody alone did not result in fluorescence in kidney (g) or liver (h) tissue sections.
Consistent with a purported functional role for differential methylation of OmpL32 in evasion of a host-generated humoral immune response, the entire repertoire of differentially methylated glutamic residues observed within OmpL32 in our study is located within predicted OmpL32-specific B cell epitopes. Also of note is the observation that the gene encoding OmpL32 (lic11848) resides in a putative three-component operon with a gene encoding a hypothetical protein (lic11849) and a gene encoding a putative methyltransferase (lic11850). Since co-expressed proteins are routinely functionally linked, this suggests that OmpL32 serves as a methylation substrate for the downstream putative methyltransferase. Combined, these observations suggest that all the required components for a rapid and effective immune evasion strategy focused around OmpL32 may be in place. In this model, exposure to altered environmental conditions would promote methylation of OmpL32 by the simultaneously expressed methyltransferase on residues contained within B cell epitopes, thus leading to an altered immune recognition of this protein on the leptospiral surface.

In summary, the current study has identified a novel L. interrogans protein designated OmpL32, which we have shown to be surface-exposed, expressed during the course of infection and differentially methylated on glutamic acid residues. Further investigation into the methylation status of leptospiral surface-exposed proteins and the potential role that this PTM plays in the leptospiral infection process is warranted, and may lead to the identification of a novel immune evasion strategy within Leptospira.

ACKNOWLEDGEMENTS

The authors would like to thank Dr Martin Boulanger, University of Victoria, Victoria, BC, Canada, for his gift of the pET28a vector and assistance with construct design, Anna von Rossum and Marcus Barron for assistance with recombinant protein production, and Dr Christoph Borchers, Derek Smith, Jen Proc, Leanne Oehlund, Darryl Hardie and the members of the University of Victoria-Genome BC Proteomics Centre for their support and guidance with the proteomic analyses. This work was supported by a University of Victoria Fellowship, Graduate Scholarship and the Pacific Century Graduate Scholarship (A.E.), a Discovery Grant from the Natural Sciences and Engineering Research Council of Canada (327186; C.E.C.), the British Columbia Proteomics Network (C.E.C.), the Canada Foundation for Innovation (C.E.C.), the British Columbia Knowledge Development Fund (C.E.C.), VA Medical Research Funds (D.A.H.), and Public Health Service grant AI-034431 (D.A.H.) from the National Institute of Allergy and Infectious Diseases. C.E.C. is a Canada Research Chair in Molecular Pathogenesis and a Michael Smith Foundation for Health Research Scholar.

REFERENCES


VagH, of *Yersinia pseudotuberculosis* reveals a tight link between VagH and the type III secretion system. *Microbiology* 153, 1464–1473.


Edited by: D. L. Gally