Ferrous iron-binding protein Omb of *Salmonella enterica* serovar Choleraesuis promotes resistance to hydrophobic antibiotics and contributes to its virulence

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*Salmonella enterica* serovar Choleraesuis (SC) is an important enteric pathogen that causes serious systemic infections in swine and humans. To identify the genes required for resistance to antimicrobial peptides, we constructed a bank of SC transposon mutants and screened them for hypersensitivity to the cationic peptide polymyxin B. Here we report one isolated polymyxin B-susceptible mutant that also exhibited increased sensitivity toward human neutrophil peptide alpha-defensin 1 (HNP-1) and hydrophobic antibiotics including erythromycin and novobiocin. The mutant had a mutation in an ORF identified as outer membrane \( \beta \)-barrel protein gene *omb*. The purified recombinant Omb protein was characterized as a ferrous iron-binding protein. The constructed *omb* isogenic mutant grew more slowly in iron-limiting conditions than the wild-type (WT) parent strain. In addition, compared with the WT strain, the *omb* mutant exhibited an increase in net negative charge upon the cell surface and was more easily killed by polymyxin B, HNP-1 and hydrophobic antibiotics. The *omb* gene was transcribed, regardless of the iron content within the growth medium, and the Omb protein appeared exclusively in the outer membrane fraction. Infection experiments demonstrated virulence attenuation when the mutant was administered orally or intraperitoneally to mice. This study indicates that Omb is a previously unrecognized ferrous iron-binding protein. *In vivo*, Omb may be involved in the acquisition of ferrous iron during the initial stages of SC infection and appears to be an important virulence factor for SC in mice.

**INTRODUCTION**

*Salmonella enterica* serovar Choleraesuis (SC) is a Gram-negative and facultative intracellular bacterium that causes serious systemic infections including typhoid disease, bacteraemia, pneumonia, septicaemia, enterocolitis, hepatitis, encephalitis and abortion in swine (Schwartz, 1999).
Although SC is highly host-adapted, the organism is also notorious for its extreme invasiveness and pathogenic nature in humans, frequently causing septicaemic disease with only marginal involvement of the intestinal tract (Chiu et al., 2006, 2005). In Taiwan, SC is the second most common of all Salmonella serotypes isolated and shows the greatest ability to cause extraintestinal infections (Wang et al., 2006). Recently, an increasing percentage of highly invasive strains of SC exhibiting high-level resistance to antimicrobial agents have become a serious problem that hampers the effective treatment of diseases caused by this pathogen.

The ability of bacterial pathogens to cause disease in mammals is dependent upon their ability to overcome the host immune defence system. Antimicrobial peptides (APs) are one of the important weapons of this defence system. Though structurally highly diverse, nearly all APs share the features of a net cationic charge and have the capacity to kill a broad spectrum of micro-organisms. Electrostatic interactions between the positively charged APs and the negatively charged bacterial cell envelope are critical for bacterial killing. It is therefore not surprising that bacterial resistance to APs is linked to charge-based modifications at the cell surface, which result in decreased binding of the cationic APs and afford the organism relative protection. Examples include the addition of L-lysine to phosphatidylglycerol (Kristian et al., 2003), the addition of aminoarabinose to the lipopolysaccharide (LPS) lipid A moiety (Tamayo et al., 2005), and the incorporation of D-alanine into Gram-positive cell wall teichoic acids (Kristian et al., 2005).

In an attempt to understand the mechanisms involved in the natural resistance of clinical isolates of SC to APs, we isolated a polymyxin B-susceptible Tn5 mutant that also exhibited increased sensitivity toward human neutrophil peptide alpha-defensin 1 (HNP-1) and hydrophobic antibiotics, including erythromycin and novobiocin. This paper reports the cloning of the Tn5-inactivated gene from this mutant and the characterization of its gene product.

METHODS

Bacterial strains and plasmids. The bacterial strains and plasmids used for this study are listed in Table 1. SC strain S280 is a clinical isolate from the blood of a patient at the National Cheng-Kung University Hospital, Taiwan, ROC. All strains were routinely grown in minimal medium (12.8 g Na2HPO4·7 H2O l−1, 3 g KH2PO4 l−1, 10 g NaCl l−1, 1 g NH4Cl l−1, 2 mM MgSO4·7H2O, 0.2 mM CaCl2, 4 g glucose l−1) or Luria–Bertani (LB) medium at 37 °C, with aeration. Antibiotics were used as follows: 100 μg ampicillin ml−1, 100 μg kanamycin ml−1 and 50 μg tetracycline ml−1 for both Escherichia coli and SC. MilliQ (Millipore) water was used for all growth media and chemical solutions. All iron-deficient media were prepared in plastic containers and filter-sterilized, and cells were grown in polypropylene and polystyrene vessels (Knight et al., 2005).

Molecular techniques. Standard techniques were used to construct recombinant plasmids (Sambrook et al., 1989). DNA fragments used in cloning were extracted from agarose gels by use of the Qiaex II kit (Qiagen). PCR was carried out according to the manufacturer’s recommendations by use of the Taq DNA polymerase kit (GE Healthcare Biosciences). Nucleotide sequences were determined with an autosequencer (ABI Prism 373 DNA sequencer; Applied Biosystems).

Construction of transposon mutant bank. The transposon insertional library of SC strain S280 was generated by a published method (Hensel et al., 1995), with minor modifications. E. coli S17-1 λpir carrying a transposon plasmid (Hensel et al., 1995) was delivered to the tetracycline-resistant SC strain S280 by conjugation. The transconjugants were selected by growth with kanamycin and tetracycline. The resultant mutants were further screened by the polymyxin B sensitivity assay described below.

AP and hydrophobic antibiotic sensitivity assay. An AP and hydrophobic antibiotic sensitivity assay was performed as described elsewhere (Kristian et al., 2005), with minor modifications. In sterile 96-well microtitre plates, exponential-phase S280 or transposon mutant culture was adjusted to 105 c.f.u. ml−1 in 100 μl minimal medium containing polymyxin B (0–10 μg ml−1) (Gibco BRL), HNP-1 (0–50 μg ml−1) (Sigma), erythromycin (0–50 μg ml−1) (Fluka), or novobiocin (0–2 mg ml−1) (Sigma), in serial dilutions. The MIC was defined as the lowest antimicrobial concentration yielding no detectable bacterial growth (OD600 measurement). All experiments were done with duplicate samples on three independent occasions.

Colony and Southern hybridization. Colony hybridization and Southern hybridization were performed as described elsewhere (Hensel et al., 1995). DNA probes were labelled with [α-32P]dCTP by use of a random priming kit (Megaprime DNA labelling system, GE Healthcare Biosciences), using either the PCR products or fragments excised from the recombinant plasmids as templates. The nylon membrane with DNA was prehybridized with hybridization buffer (ExpressHyb hybridization solution; Clontech Laboratories) for 30 min at 65 °C, hybridized for 2 h at 65 °C, washed, and visualized by autoradiography.

Analysis of transposon insertion sites. Chromosomal DNA from each Tn5 mutant was digested individually with BglII, EcoRI, KpnI, PstI and SalI (there are no recognition sites for these five restriction enzymes within the transposon). The presence of the Tn5 mutant was screened for with an α-32P-labelled kanamycin-resistance gene by Southern hybridization. The kanamycin probe was generated by excision from plasmid pUC4K to generate a 1.2 kb SalI fragment, which was then used as a template for the random priming kit.

Cloning of the SC omb gene. Chromosomal DNA from strain S280-1 was digested with EcoRI and inserted into identically digested pUC19. The ligation reaction mixtures were transformed into E. coli XL1Blue and then cells were selected for kanamycin resistance. Plasmid DNA was extracted, and the chromosomal DNA sequences flanking the transposon were obtained by DNA sequencing using primers P6 and P7 (Hensel et al., 1995). A genomic library of S280 was constructed in E. coli by using Sure3A.I partially digested S280 DNA ligated into the BamHI site of pBR322, as described previously (Chang et al., 1993). The complete coding sequence of omb was cloned from the genomic library of the S280 strain by colony hybridization with an α-32P-labelled DNA fragment of the omb gene. One positive clone was selected for DNA sequencing.

Overexpression of Omb in E. coli. A 663 bp fragment of the omb gene was amplified from plasmid pSC by PCR with primers F4 (\`TCTTCTGAGCAGGGTACCGCTGCGG-3\`) and R1 (\`TCTTCTGAGCAGGGTACCGCTGCGG-3\`). PCR products were digested with BamHI and Xhol and inserted into identically
digested pGEX-5X-3 (GE Healthcare) to generate plasmid pSC4. The GST–Omb fusion protein containing the amino acid region of Omb between Asp-34 and Leu-248 was expressed in E. coli BL21(DE3). The transformants were grown at 37 °C in LB medium containing 100 μg ampicillin ml⁻¹. When the culture reached OD₆₀₀ 0.5, IPTG was added to a final concentration of 1.0 mM. After a further 4 h of growth, the cultures were harvested by centrifugation at 4000 r.p.m., for 15 min at 4 °C. The pellet was resuspended in PBS containing 0.05% Tween-20 (PBST) in 1% of the original volume and lysed by sonication on ice, and then centrifuged at 10 000 r.p.m. for 20 min at 4 °C. The supernatant was applied to glutathione–Sepharose 4B beads (Pharmacia) and extensively washed with PBST. Factor Xa was injected into the column and the column was maintained at room temperature overnight. Using this method, the fusion protein was site-specifically cleaved and the recombinant Omb was eluted with 0.05% Tween-20 (PBST) in 1% of the original volume and lysed by sonication on ice, and then centrifuged at 10 000 r.p.m. for 20 min at 4 °C. The supernatant was applied to glutathione–Sepharose 4B beads (Pharmacia) and extensively washed with PBST. Factor Xa was injected into the column and the column was maintained at room temperature overnight. Using this method, the fusion protein was site-specifically cleaved and the recombinant Omb was eluted with 50 mM Tris/HCl (pH 7.7) containing 100 mM NaCl. The yield of the Omb by this method was estimated at about 4 mg from 1 l bacterial culture.

Iron-binding assay. An iron-binding assay was performed as described elsewhere (Petrat et al., 1999), with minor modifications. The fluorescent dye phen green SK (dipotassium salt) (PG SK) for iron detection was purchased from Invitrogen Molecular Probes. The fluorescent dye phen green SK (dipotassium salt) (PG SK) was used to determine the iron-binding capacity of the recombinant protein. The assay was performed by incubating the recombinant protein with a constant concentration of PG SK and varying concentrations of iron. The absorbance at 510 nm was measured and plotted against the iron concentration. The iron-binding capacity of the recombinant protein was determined by fitting the data to a linear regression model.

Preparation of polyclonal antisera. A 665 bp fragment of the omb gene was amplified from plasmid pSC by PCR with primers F1 (5'-TACGTTAGATGATGATCCACCGGCA-3') and R1 (5'-TTCTTCGACGACGGTACCGTGATTGGC-3'). PCR products were digested with Ndel and Xhol and inserted into identically digested pET21b (Novagen) to generate plasmid pSC1. His₆-tagged Omb containing the amino acid region of Omb between Met-33 and Leu-248 was expressed in E. coli BL21(DE3) and purified under denaturing conditions by using a nickel affinity column, according to the manufacturer’s instructions (Novagen). Purified His₆-tagged Omb at 100 μg (ml saline)⁻¹ was mixed with 1 ml Freund’s incomplete adjuvant. This mixture was then injected subcutaneously into an Elite New Zealand White rabbit. Two booster doses were administered at 2-week intervals, and the antiserum was collected after 6 weeks. The antiserum was purified through a protein A column according to the manufacturer’s instructions (GE Healthcare Biosciences).

Western blot analysis. Proteins separated by SDS-12% PAGE were electroblotted onto a nitrocellulose membrane (GE Healthcare Biosciences) and then incubated with a His₆-tagged Omb-specific rabbit polyclonal antibody as the primary antibody. The secondary antibody was a 1:15 000 dilution of goat anti-rabbit IgG conjugated to horseradish peroxidase (Sigma). The proteins were visualized by

<table>
<thead>
<tr>
<th>Strain or plasmid</th>
<th>Genotype or comments</th>
<th>Reference or source</th>
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</thead>
<tbody>
<tr>
<td><strong>SC strains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S280</td>
<td>WT clinical isolate</td>
<td>Hospital centre of Cheng-Kung University</td>
</tr>
<tr>
<td>S280-1</td>
<td>S280 omb::Tn</td>
<td>This study</td>
</tr>
<tr>
<td>S280-2</td>
<td>S280 omb::pCSC</td>
<td>This study</td>
</tr>
<tr>
<td>S280-3</td>
<td>S280-2 strain with plasmid pSC2</td>
<td>This study</td>
</tr>
<tr>
<td><strong>E. coli strains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BL21(DE3)</td>
<td>F’ ompT hsdSB (rK mC) gal dcm (DE3)</td>
<td>Novagen</td>
</tr>
<tr>
<td>S17-Lipir</td>
<td>Tp’ Sm’ recA pro hsdR’ M’ , RP4:2Tc::Mu: Km T7, ipir</td>
<td>Hensel et al. (1995)</td>
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<td>XL1Blue</td>
<td>F’::Tn10 pro A’ B’ lacD Δ(lacZ)M15 recA endA1 gyrA96(Nal’) thi hsdR17(rK mC) glnV44 relA1 lac</td>
<td>Stratagene</td>
</tr>
<tr>
<td><strong>Plasmids</strong></td>
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<tr>
<td>pBR322</td>
<td>Standard cloning vector; Ap’*</td>
<td>New England Biolabs</td>
</tr>
<tr>
<td>pCVD422</td>
<td>Vector for insertion disruption; Ap’</td>
<td>Donnenberg &amp; Kaper (1991)</td>
</tr>
<tr>
<td>pET21b</td>
<td>Expression vector; Ap’</td>
<td>Novagen</td>
</tr>
<tr>
<td>pGEX-5X-3</td>
<td>Expression vector; Ap’</td>
<td>GE Healthcare</td>
</tr>
<tr>
<td>pSC</td>
<td>Positive clone from S280 genomic library; ~3.0 kb chromosomal fragment in pBR322; Ap’</td>
<td>This study</td>
</tr>
<tr>
<td>pSC1</td>
<td>omb structure gene cloned into pET21b; Ap’</td>
<td>This study</td>
</tr>
<tr>
<td>pSC2</td>
<td>omb structure gene cloned into pBR322; Ap’</td>
<td>This study</td>
</tr>
<tr>
<td>pSC3</td>
<td>292 bp internal fragment from S280 chromosome omb gene cloned into pCVD422; Ap’</td>
<td>This study</td>
</tr>
<tr>
<td>pSC4</td>
<td>omb structure gene cloned into pGEX-5X-3; Ap’</td>
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</tr>
<tr>
<td>pUC4K</td>
<td>Source of kanamycin-resistance cassette; Ap’</td>
<td>Pharmacia</td>
</tr>
<tr>
<td>pUC19</td>
<td>Standard cloning vector; Ap’</td>
<td>New England Biolabs</td>
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</table>

chemiluminescence according to the manufacturer’s protocol (ECL Western Blotting Detection Reagent; GE Healthcare Biosciences).

**Cellular fractionation.** Extracellular, periplasmic and cytoplasmic cell fractions were isolated as described elsewhere (Cornelis et al., 1982). The sarcosine-insoluble outer membrane fraction of S280 was prepared as described elsewhere (Doig & Trust, 1994), with minor modifications. The cell pellets were disrupted by passage through a French pressure cell [20 000 p.s.i. (138 000 kPa)]. After removal of unbroken cells, the total membrane (pellet) and cytoplasmic (supernatant) fractions were collected by centrifugation (40 000 g, 30 min, 4 °C), resuspended in 20 mM Tris/HCl (pH 7.5) containing 2.0 % (w/v) sodium lauryl sarcosine, and incubated at room temperature for 30 min. Outer membrane proteins were collected by centrifugation (40 000 g, 30 min, 4 °C) and washed three times with distilled water.

**Effect of iron on the growth of the SC omb mutant.** Strains were grown at 37 °C in LB broth and harvested at their mid-exponential phase of growth. Bacteria were centrifuged for 5 min at 3000 g, washed twice with minimal medium, and resuspended in minimal medium to OD$_{500}$ 0.1. Bacterial suspensions were added at 1:10 to minimal medium containing different concentrations of the iron chelator 2,2’-dipyriddy1 (2,2-DPD) (Sigma Chemical) (5 × 10$^{-4}$ and 1 × 10$^{-3}$ μM) and incubated at 37 °C. The growth rate of the cells was monitored by OD$_{500}$ measurements. Samples were shaken gently, and aliquots removed at specified times were assayed for bacterial counts by plating out serial dilutions onto LB plates. The doubling times (g) for the number of bacteria in iron-limited medium were calculated with the following equation:

$$g = \frac{0.301t}{n} (\log N_t - \log N_0)$$

where $N_0$ is the initial population number, $N_t$ is the population number at time $t$, and $n$ is the number of generations in time $t$.

**Cytochrome c binding assay.** A cytochrome c binding assay was performed as described previously (Kristian et al., 2005). Bacteria were grown to early exponential phase, washed twice with MOPS buffer (20 mM, pH 7), adjusted to a final OD$_{500}$ of 7 in MOPS buffer plus 0.5 mg cytochrome c ml$^{-1}$ (Sigma-Aldrich), and incubated at room temperature. A plate counting assay was also performed to verify the numbers of viable cells. As a control, 0.5 mg cytochrome c ml$^{-1}$ was incubated in MOPS buffer under the same conditions but without bacteria. After 10 min, bacteria were removed by centrifugation (21 000 g, 3 min) and the cytochrome c content of the supernatants was quantified photometrically at 530 nm, the absorption maximum of the prosthetic group.

**Construction of the omb mutant.** A 292 bp internal fragment of the omb gene from pSC was generated by PCR with primers F2 (5’-CTGGCATGCGAATCCGACGGCT-3’) and R2 (5’-GGTGTCGACCGATATGCACCAT-3’). The PCR product was digested with SphI and SalI and inserted into identically digested pCVD422 (Donnenberg & Kaper, 1991). The resultant plasmid, pSC, was introduced into S280-2 by using a published method (Beliavskia et al., 2000). Transformants were selected by using ampicillin. The resultant strain was further verified by plasmid extraction and PCR.

**Virulence assay.** BALB/c mice (6–8 weeks old) purchased from the animal centre of the College of Medicine at National Cheng-Kung University were challenged by intraperitoneal (i.p.) injection or by feeding with a bacterial suspension. In feeding experiments, mice were starved for 6 h to clear the contents of the stomach before the oral challenge. A 0.1 ml volume of 100 mM NaHCO$_3$ was administered intragastrically to neutralize the stomach contents. A group of eight mice were challenged orally with 0.2 ml per mouse of a 10-fold serially diluted (in PBS) bacterial suspension by using a round-tipped stainless steel feeding needle, and mortality was recorded at 3–4 weeks post-infection. Each experiment was performed at least twice with similar results. The LD$_{50}$ for each strain was calculated by the method of Reed & Muench (1938).

**RESULTS**

**Isolation and characterization of polymyxin B-sensitive mutant S280-1**

To identify the genes required for resistance to APs, we constructed a bank of Tn5 mutants of a clinical strain of SC S280, and screened them for susceptibility to APs using AP sensitivity assays. One polymyxin B-sensitive clone, S280-1, was isolated and further investigated. Compared with the wild-type (WT) strain, this mutant not only exhibited clearly decreased MICs for polymyxin B and HNP-1, but also showed significantly enhanced susceptibility to hydrophobic antibiotics including erthyromycin and novobiocin (Table 2). However, the sensitivity of S280-1 to other antibiotics, including streptomycin, tetracycline, ampicillin, nalidixic acid and chloramphenicol, remained unchanged (data not shown). The S280-1 mutant had the same colony morphology as the WT strain, and both grew at similar rates in LB medium or minimal medium (data not shown), suggesting that S280-1 was not an auxotrophic mutant.

Southern blot analysis with the kanamycin gene as the probe revealed that the S280-1 mutant contained a transposon insertion site (data not shown). The DNA sequences immediately flanking the Tn5 integration sites in the S280-1 mutant were obtained as described in Methods. The resultant plasmid was sequenced, and through sequence analysis, 630 bp of S280-1 chromosomal DNA beyond the transposon was cloned.

**Characterization of the genetic loci**

For isolation of the complete coding sequence of the Tn5-interrupted gene, a genomic library of strain S280 DNA was subjected to colony hybridization, and a plasmid, pSC, obtained from a probe-reactive clone was selected for analysis. Three complete and one partial ORFs were identified. The DNA sequence of the first complete ORF (Fig. 1), where the mini-Tn5 was inserted in strain S280-1,
was designated *omb* (outer membrane β-barrel protein gene). The *omb* gene consists of a single transcriptional unit in which the coding sequence encodes a protein of 252 aa with a molecular mass of 27.8 kDa. The first 23 aa of the deduced Omb sequence have the characteristics of a typical bacterial signal sequence, followed by a typical Ala-X-Ala signal peptidase cleavage site (von Heijne, 1983) between Ala-23 and Ala-24. Cleavage of the signal sequence would give a mature protein with a molecular mass of 25 kDa, which is in reasonable agreement with the size derived from SDS-PAGE (Fig. 2). The putative amino acid sequence of intact Omb shared 99–100 % sequence identity with the putative outer membrane proteins of *Salmonella typhimurium* LT2 and CT18, and *Salmonella paratyphi* ATCC9150, as well as having high shared degrees of identity (>54 %) with the hypothetically extracytoplasmic proteins of several enteropathogenic *E. coli* strains and other pathogens, including *Shigella* spp., *Yersinia* spp., *Klebsiella pneumoniae*, *Erwinia tasmaniensis* and *Serratia proteamaculans* (data not shown). Only one of the aforementioned high-identity proteins has been characterized, the *E. coli* b1722 protein (Eb1722). Eb1722 has been classified as an outer membrane receptor (OMR) family protein by the β-barrel finder (BBF) program, although its amino acid sequence exhibits low similarity to known OMR family proteins (Zhai & Saier, 2002). Since Eb1722 shares 88 % sequence identity with Omb, it seems reasonable to propose that Omb is a member of the OMR family of proteins. The second ORF was heterologously positioned 48 bp downstream of *omb* and was predicted to encode a 319 residue protein with a molecular mass of ~34.8 kDa. Its amino acid sequence shared 71 % sequence identity with systemic factor protein A (SfpA) from *Yersinia enterocolitica* (Mildiner-Earley & Miller, 2006), and was therefore designated *sfpA* of SC. SfpA from *Y. enterocolitica* is a porin, located in the membrane, and is necessary for sustaining colonization of mesenteric lymph nodes; SfpA is conserved in other pathogenic bacteria that are involved in systemic disease (Mildiner-Earley & Miller, 2006). The third ORF was heterologously positioned 142 bp upstream of *sfpA* and was predicted to encode a 104 residue protein with a molecular mass of ~12.3 kDa. This amino acid sequence shared a high degree of sequence identity (96 %) with that of the putative inner membrane protein STM1329 [accession no. NP_460295 (gi: 16764680)] from *S. typhimurium* LT2, and was thus designated *imp* (putative inner membrane protein gene) from SC. The incomplete ORF was heterologously positioned 289 bp upstream of *omb* and potentially encoded the N-terminal 198 aa of a protein that shared significant homology (98 %) to 6-phosphofructokinase II from *E. coli* (Daldal, 1984) and was therefore designated *pfkB* from SC. The genetic organization and the transcriptional direction of the three genes, *omb*, *sfpA* and *imp* of SC strain S280, strongly suggest that the mutant S280-1 phenotype is attributable only to the *omb* insertional mutation, since no polar effects on downstream genes should be expected. Recently, the complete genome sequence of SC strain SC-B67 has become publicly available (http://www.ncbi.nlm.nih.gov/genomes/lproks.cgi?view=1). A comparison of the sequences of strain *S280* *omb* and *sfpA* with those of strain SC-B67 revealed that the respective sequences are the same at the nucleotide and amino acid levels. However, *imp* of strain *S280* is an ORF encoding 104 aa of a protein, while the *imp*-related gene of SC-B67 has been designated a pseudogene (SCPS16) (http://www.ncbi.nlm.nih.gov/genomes/lproks.cgi?view=1) which encodes 82 aa, and is 100 % identical with the N-terminal 82 aa sequence of *S280* Imp. Nevertheless, the ORF of *S280* *imp* is extended by an additional 22 aa at the C terminus.

**Table 2.** MICs for the APs and hydrophobic antibiotics used in this study

<table>
<thead>
<tr>
<th>SC strain</th>
<th>MIC (μg ml⁻¹)</th>
<th>APs</th>
<th>Hydrophobic antibiotics</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Polymyxin B</td>
<td>HNP-1</td>
</tr>
<tr>
<td>S280 (WT)</td>
<td>0.458 ± 0.014</td>
<td>3.17 ± 0.06</td>
<td>3.33 ± 0.06</td>
</tr>
<tr>
<td>S280-1 (Tn5 mutant)</td>
<td>0.150 ± 0.008*</td>
<td>0.67 ± 0.03*</td>
<td>0.83 ± 0.03*</td>
</tr>
<tr>
<td>S280-2 (isogenic mutant)</td>
<td>0.167 ± 0.006*</td>
<td>0.73 ± 0.03*</td>
<td>0.79 ± 0.03*</td>
</tr>
<tr>
<td>S280-3 (S280-2+pSC2)</td>
<td>0.471 ± 0.011</td>
<td>3.38 ± 0.09</td>
<td>3.58 ± 0.06</td>
</tr>
</tbody>
</table>

*Significantly lower MIC of AP/hydrophobic antibiotic for the mutant strain compared with the WT strain S280 (two-tailed t test, P<0.05).*

**Fig. 1.** Chromosomal organization of *omb* and the flanking region in SC S280. Arrows represent the direction of transcription and the ORFs of *omb*, *sfpA* and *imp*.
Identification of the Omb protein as a ferrous iron-binding protein of SC

Since Omb shared striking homology to Eb1722 (classified as an OMR family protein), and since several OMR proteins have been characterized as ferrichrome-iron receptors (Zhai & Saier, 2002), we suspected that Omb was also an iron-binding protein. To investigate this, we carried out fluorescence studies on purified recombinant Omb protein using the fluorescent iron indicator PG SK. The results showed that free PG SK in a simple buffered solution had an excitation maximum at 507 nm and an emission maximum at 532 nm. Addition of ferrous iron or ferric iron causes quenching of PG SK fluorescence. However, the addition of Omb with ferrous iron or ferric iron markedly decreased PG SK fluorescence quenching. Neither addition of purified Omb nor addition of iron (ferrous iron or ferric iron) led to a shift of the excitation or emission spectra. Loading PG SK to a final concentration of 4 μmol l⁻¹ to a simple buffered solution containing 10⁻⁶ M ferrous iron with various concentrations of Omb (10⁻⁶–10⁰ μg ml⁻¹) led to a rapid increase in fluorescence in a manner dependent upon the Omb protein dose (Table 3). The more Omb present, the stronger the fluorescence. However, this phenomenon was not observed when ferrous iron was replaced with ferric iron under the same conditions as described above. This suggests that the fluorescence of PG SK was already partly quenched by ferrous iron chelation under control conditions and that redistribution of ferrous iron from the fluorescence indicator to a chelator-like Omb abolished this quenching, i.e. ‘dequenched’ the fluorescence. These results indicated that the Omb protein specifically binds ferrous iron with high affinity, at least in vitro.

Gene expression and localization of Omb in S280

To investigate whether Omb could or could not be produced under iron-rich conditions, and to determine the cellular location of Omb, S280 cells were grown to exponential phase in LB medium, pelleted and fractionated, and then the production and distribution of Omb in each cellular fraction was analysed by Western blotting. The results showed that Omb could be produced in both iron-rich LB medium and minimal medium and migrated as a ~25 kDa polypeptide, which agreed well with the mass predicted from the putative amino acid sequence (Fig. 2a). In addition, Omb was found only in the outer membrane fraction (Fig. 2b). Most high-affinity iron acquisition systems are not expressed constitutively and their expression is upregulated in response to a decrease in the level of intracellular iron. To investigate the effect of iron on the

Table 3. Responsiveness of PG SK to various concentrations of Omb with Fe²⁺ and Fe³⁺

<table>
<thead>
<tr>
<th>Omb concentration (μg ml⁻¹)</th>
<th>PG SK fluorescence (percentage of baseline fluorescence)</th>
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<tbody>
<tr>
<td></td>
<td>Fe²⁺ (10⁻⁶ M)</td>
</tr>
<tr>
<td>0</td>
<td>6.6±0.3</td>
</tr>
<tr>
<td>10⁻⁶</td>
<td>9.7±0.5</td>
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<tr>
<td>10⁻⁵</td>
<td>18.4±0.4</td>
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<tr>
<td>10⁻⁴</td>
<td>27.3±0.5</td>
</tr>
<tr>
<td>10⁻³</td>
<td>41.5±0.9</td>
</tr>
<tr>
<td>10⁻²</td>
<td>56.8±0.7</td>
</tr>
<tr>
<td>10⁻¹</td>
<td>67.2±0.8</td>
</tr>
<tr>
<td>1</td>
<td>79.3±0.9</td>
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</table>
expression of Omb of SC, we performed RT-PCR and Western blot analyses under various culture conditions of iron restriction. The results from RT-PCR experiments indicated that the transcription levels of omb did not increase in response to a decrease in the level of iron (data not shown). Western blot analyses confirmed the results of the RT-PCR experiments, and indicated that the Omb protein level did not increase in response to a decrease in the level of iron, even at higher concentrations of the iron chelator 2,2-DPD (5 \times 10^{-4} \text{ and } 1 \times 10^{-3} \text{ µM}) (data not shown). These results indicate that in SC, Omb protein is located in the outer membrane, and that Omb expression is probably not regulated by the prevailing level of iron in the environment.

The absence of Omb renders SC sensitive to killing by hydrophobic antibiotics

To demonstrate that the observed AP-sensitive and hydrophobic antibiotic-sensitive phenotype was reproducibly linked to the transposon insertion at the omb locus, we reconstructed an omb isogenic mutant, S280-2. Inactivation of the WT omb with the insertional disruption of the omb gene in S280-2 was checked by PCR, using a pair of primers complementary to sequences located in the omb and ampicillin-resistance genes (data not shown). For confirmation of the lack of Omb protein in the mutant strain S280-2, Western blot analysis of the WT and mutant S280-2 strains was performed. The results of the Western blot analysis demonstrated that the S280-2 mutant does not produce any anti-Omb-reactive protein (Fig. 2a). The isogenic mutant S280-2 was examined for sensitivity to polymyxin B and HNP-1 in suspension at different concentrations. As shown in Table 2, the mutant strain S280-2 exhibited clearly decreased MICs for polymyxin B and HNP-1 compared with the WT strain. Similarly, S280-2 showed significantly enhanced susceptibilities to erythromycin and novobiocin compared with the WT strain (Table 2). These increased susceptibilities to polymyxin B, HNP-1, erythromycin and novobiocin are not significantly different from those of the Tn5 mutant S280-1. Furthermore, like the Tn5 mutant S280-1, sensitivity of S280-2 to other antibiotics, including streptomycin, tetracycline, ampicillin, nalidixic acid and chloramphenicol, remained unchanged.

The absence of Omb affects surface charge

Since the increased sensitivities of the Tn5 S280-1 and S280-2 mutants seemed to be restricted to cationic APs and hydrophobic antibiotics, and since the outer membrane protein Omb was identified as a ferrous iron-binding protein, we suspected that the lack of Omb caused alterations in the bacterial surface charge. To investigate this, the capacity of the WT and S280-2 mutant cells to bind a highly positively charged molecule, cytochrome c, was determined. As shown in Fig. 3, the mutant bound a higher amount of cytochrome c than the WT strain, suggesting that the cell surface is more highly negatively charged in the mutant than in the WT strain.

The absence of Omb leads to attenuated growth in iron-limited conditions

We further examined the growth of the WT strain and mutant strain S280-2 in media under different iron-limiting conditions. The results showed that both strains grew in minimal medium containing less than 5 \times 10^{-4} \text{ µM} of the iron chelator 2,2-DPD, and both grew with approximately identical growth rates (Fig. 4), indicating that the ability of the mutant to utilize available iron is similar to that of the WT strain with higher levels of readily available iron. However, in minimal medium containing 5 \times 10^{-4} \text{ and } 1 \times 10^{-3} \text{ µM} 2,2-DPD, which represents lower and the lowest levels of readily available iron tested, respectively, the mutant showed attenuated growth compared with the WT strain and the doubling times for the mutant were 36 and 280 min, respectively, whereas those for the WT strain were 18 and 190 min, respectively. The results revealed that the mutant showed attenuated growth in readily available iron-limiting conditions compared to the WT strain (Fig. 4).

Complementation of the S280-2 mutant

To confirm that (a) the attenuated growth of the S280-2 mutant in response to 5 \times 10^{-4} \text{ or } 1 \times 10^{-3} \text{ µM} 2,2-DPD, and (b) the AP- and hydrophobic antibiotic-sensitive phenotype of this mutant were indeed due to inactivation of the omb gene, we cloned the WT gene into plasmid pBR322; this construct (pSC2) was transformed into the S280-2 mutant strain. The results revealed that complementation of the S280-2 mutant with pSC2, but not introduction of a control plasmid, rescued the ability to exhibit WT growth rates in minimal medium containing 5 \times 10^{-4} \text{ or } 1 \times 10^{-3} \text{ µM} 2,2-DPD. Sensitivity measure-
incubated at 37 °C for 72 h in minimal medium containing the iron chelator 2,2-DPD at 0 μM (●, ○), 5×10⁻⁴ μM (▲, △) or 1×10⁻³ μM (■, □); the values presented are mean and SE (not shown where smaller than the symbols) of three independent experiments.

**Reduced virulence is associated with Omb**

The virulence of the isogenic mutant S280-2 and the WT strain S280 was studied by infecting BALB/c mice via either the oral or i.p. route. As shown in Table 4, the LD₅₀ values for mutant strain S280-2 in mice were 4.0×10⁵ and 3.5×10² at 3 weeks post-infection via the oral and i.p. routes, respectively. These values were 20- and 15-fold higher than the LD₅₀ values of 1.7×10⁴ and 2.0×10¹ for WT infection in mice, administered via the oral and i.p. routes, respectively.

**DISCUSSION**

The outer membrane of enteric bacteria functions as an effective permeability barrier to harmful hydrophobic molecules. Consequently, OM impermeability is generally considered to be a characteristic feature of enteropathogenic bacteria. In this report, we have identified an outer membrane protein (Omb) in SC that possesses a ferrous iron-binding capacity. We have demonstrated that the SC omb mutant is more sensitive to polymyxin B, HNP-1, erythromycin, and novobiocin than the WT strain, suggesting that Omb is responsible for the SC resistance to multiple hydrophobic antimicrobial agents. Furthermore, compared with the WT strain, the SC omb mutant also showed attenuated growth in iron-limiting medium and in infection experiments. We demonstrated virulence attenuation when this mutant was administered orally or intraperitoneally to mice. Thus, Omb represents a potential virulence factor that could contribute in multiple ways to the pathogenesis of SC infection.

Although bioinformatics analysis using BLAST revealed that the putative amino acid sequence of Omb shares high degrees of identity (>54%) with hypothetical extracytoplasmic proteins of various Gram-negative bacteria, nearly all of them are enteropathogenic bacteria. None of these Omb high-identity proteins has been characterized, except for Eb1722. In addition, because Eb1722 is classified as an OMR family protein according to bioinformatics, although without any experimental basis, the biological roles of these Omb-high-identity proteins in Gram-negative bacteria are unclear. Since we determined the location of Omb to be the SC outer membrane and demonstrated that purified recombinant Omb protein possesses a ferrous iron-binding ability, it is reasonable to propose that Omb may be part of an iron transport system upon which SC depends to scavenge ferrous iron from the environment. Nevertheless, Omb iron acquisition differs markedly from iron transport systems in Gram-negative bacteria that employ iron-regulated outer membrane receptors. Most Gram-negative bacteria have a Fur-like iron acquisition system whose related genes form an operon with a Fur box sequence that binds regulatory Fur repressor protein under iron-limiting conditions (Clarke et al., 2001; Wandersman & Delepelaire, 2004). However, no Fur box sequence was found upstream of omb. In addition, ambient ORFs were heterologously positioned downstream or upstream of omb, indicating that the omb gene consists of a single transcriptional unit and does not form an operon with other genes. Most importantly, no change in omb expression was observed for the SC strain grown under iron-rich or different iron-restricted conditions, indicating that the expression of omb is not regulated by iron. Thus, the Omb protein may represent a novel ferrous-iron transport system that promotes the acquisition of ferrous iron by SC from the environment.

Resistance to the bactericidal effect of APs is an important property of pathogenic bacteria. Most pathogenic bacteria...
have developed a variety of strategies to combat APs. These strategies include increasing the amount of capsule polysaccharide to act as a shield against APs (Campos et al., 2004), downregulation of bactericidal peptides by bacterial plasmid DNA (Islam et al., 2001), expelling APs through energy-dependent pumps (Bayer et al., 2006; Tzeng et al., 2005), decreased fluidity of the outer membrane outer leaflet through increased acylation of lipid A (Guo et al., 1998), cleaving APs with proteases (Nyberg et al., 2004), and the repulsion of APs by reducing the net negative charge of the bacterial cell envelope through covalent modification of anionic molecules. In this study, we found that Omb is a ferrous iron-binding protein and that the omb mutant is more sensitive to polymyxin B and HNP-1 than the WT strain. Because of the cationic nature of polymyxin B and HNP-1, the electrostatic attraction of these two APs to the negatively charged bacterial membranes prior to membrane insertion is likely to be critical for bacterial killing (Hancock & Chapple, 1999). In addition, the omb mutant showed significantly enhanced susceptibility to hydrophobic antibiotics (including erythromycin and novobiocin) when compared with the WT strain. Furthermore, sensitivity toward neutral antibiotics (including streptomycin, tetracycline, nalidixic acid and chloramphenicol) was the same for both the omb and WT strains. Because of the two factors above, we propose that the constitutive uptake of ferrous iron from the environment by the Omb outer membrane protein increases the positive charge at the bacterial surface sufficiently to decrease the affinity of APs and hydrophobic antibiotics for their cell wall targets, thereby affording the organism relative protection. Our finding that the omb mutant bound higher amounts of positively charged cytochrome c than the WT strain is consistent with this notion. Furthermore, it has been reported that polyionic LPS molecules of Gram-negative bacteria are electrostatically linked by divergent cations (e.g. Mg$^{2+}$ and Ca$^{2+}$) and form a ‘tiled-roof’ structure that functions as an effective permeability barrier against hydrophobic antibiotics, detergents, dyes and macromolecules [e.g. EDTA, polymyxin B nonapeptide (PMBN), lysine polymers and protamine] (Vaara, 1992). It may be reasoned that the binding of ferrous iron by outer membrane Omb protein molecules not only reduces the negative charge of the cell envelope, but also stabilizes and maintains the integrity of the outer lipopolysaccharide layer and cell wall, and thereby enhances the barrier.

The role of iron, as a regulator for virulence expression, was first established by Pappenheimer & Johnson (1936). Alteration of available iron levels, brought about by inherited disease or tissue injury, predisposes humans to infection with a variety of pathogens (Bullen et al., 2006). However, the level of free iron in biological fluids of healthy humans is very tightly limited and controlled (Weinberg, 1978). To obtain this unavailable iron, most pathogenic bacteria including Salmonella spp. have to lyse host cells or induce expression of haemophores and siderophores via the iron-responsive Fur regulon (Clarke et al., 2001; Wandersman & Delepelaire, 2004). We propose that the availability of iron via these routes is insufficient to meet the needs of SC in the early stages of infection. In this study, we identified a novel protein, Omb, in the SC outer membrane with ferrous iron-binding ability that may deliver sufficient additional iron to the bacterium. Since the results of the fluorescence analysis indicated that purified recombinant Omb is a high-affinity ferrous iron-binding protein, and since the expression of Omb in SC is independent of iron supply and appears to be constitutive, it would be reasonable to speculate that ferrous iron uptake by Omb in SC is unique and very important for early stage infection before other iron acquisition systems become induced. Bacterial pathogens contain several iron acquisition systems, so that mutation of one system can be compensated for by another system, thus protecting the organism from virulence attenuation (Nassif et al., 1987; Wu et al., 2002; Wyckoff et al., 2006).

In contrast, the virulence of our constructed omb isogenic mutant became attenuated in mice compared with the WT strain, which is consistent with the notion that Omb in SC is unique and very important for early stage infection. Although systems involved in ferrous iron uptake and usage in SC are beginning to be investigated, the precise mechanism by which these bacteria are able to bind and transport ferrous iron through their membranes is not well understood.

In summary, these data represent the first description, to our knowledge, of Omb as a novel outer membrane protein that is very important for scavenging ferrous iron. It is possible that the absence of Omb causes structural changes, with increased net negative charge on the surface, thereby facilitating interaction of APs and hydrophobic antibiotics with the bacterial surface. The BLAST search revealed that at least 20 different bacterial species contain a putative Omb protein (>54% identity), with only one such protein found in each of these species. Since all of these species are Gram-negative enteropathogenic bacteria, it would be interesting to investigate whether Omb is also a ferrous iron-binding protein involved in hydrophobic antibiotic resistance in any of these other bacteria. Such studies may provide a strategy for designing new drugs or novel combinations of drugs for the treatment of Gram-negative enteropathogenic infections including those with SC.

ACKNOWLEDGEMENTS

This work was supported by a grant (97-EC-17-A-10-S1-0013) from the Department of Industrial Technology, Ministry of Economic Affairs of Taiwan.

REFERENCES


Edited by: A. R. Walmsley