The Pasteurella haemolytica 35 kDa iron-regulated protein is an FbpA homologue

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In a previous investigation, a 35 kDa iron-regulated protein was identified from total cellular proteins of Pasteurella haemolytica grown under iron-depleted conditions. This study reports identification of the gene (fbpA) encoding the 35 kDa protein based on complementation of an entA Escherichia coli strain transformed with a plasmid derived from a P. haemolytica lambda ZAP II library. Cross-reactivity was demonstrated between an anti-35 kDa mAb and a 35 kDa protein expressed in this strain. Furthermore, a translated ORF identified on the recombinant plasmid corresponded with the N-terminal amino acid sequence of the intact and a CNBr-cleaved fragment of the 35 kDa iron-regulated protein. Nucleotide sequence analysis of the gene encoding the 35 kDa protein demonstrated homology with the cluster 1 group of extracellular solute-binding proteins, especially to the iron-binding proteins of this family. Complete sequence analysis of the recombinant plasmid insert identified three other predominant ORFs, two of which appeared to be in an operonic organization with fbpA. These latter components (fbpB and fbpC) showed homology to the transmembrane and ATPase components of ATP-binding cassette (ABC)-type uptake systems, respectively. Based on amino acid/DNA sequencing, citrate competition assay of iron affinity and visible wavelength spectra, it was concluded that the P. haemolytica 35 kDa protein functions as an FbpA homologue (referred to as PFbpA) and that the gene encoding this protein is part of an operon comprising a member of the FbpABC family of iron uptake systems. Primary sequence analysis revealed rather surprisingly that PFbpA is more closely related to the intracellular Mn/Fe-binding protein IdIA found in cyanobacteria than to any of the homologous FbpA proteins currently known in commensal or pathogenic members of the Pasteurellaceae or Neisseriaceae.

Keywords: Pasteurella haemolytica, iron, FbpA, ABC transporter

INTRODUCTION

Iron is an essential and limiting resource in most environments. In the mammalian host this is largely due to the iron-sequestering effects of the mammalian iron-binding proteins transferrin (in sera) and lactoferrin (on mucosal surfaces). In response to the difficulties associated with acquiring iron, many microorganisms have developed high-affinity iron acquisition systems to obtain this nutrient (Mietzner & Morse, 1994). The expression of genes encoding proteins involved in the iron acquisition process is often coordinately regulated by the availability of iron. In bacterial pathogens, low levels of iron serve as a signal for expression of a number of bacterial genes involved in adaptation to the host, including genes responsible for the pathogenesis of infection. Thus, proteins expressed under iron-limited conditions may be of essential function for bacterial survival and may ultimately serve as immuno- or chemotherapeutic targets.

Pasteurella haemolytica is the causative agent of bovine...
pneumonic pasteurellosis (shipping fever), responsible for a significant economic loss to the cattle industry. Initial difficulties encountered in developing vaccines to *P. haemolytica* prompted investigators to identify and study proteins which were preferentially expressed under iron-limiting conditions. A number of such iron-regulated proteins have been identified, including proteins of molecular mass 100, 77, 71, 42, 35 and 31 kDa (Deneer & Potter, 1989; Murray et al., 1992; Tabatabai & Frank, 1997). The 100 kDa protein has been demonstrated to possess transferrin-binding activity (Ogunnariwo & Schryvers, 1990; Gray-Owen & Schryvers, 1996) and the genes encoding the 100 and 71 kDa proteins in *P. haemolytica* have since been cloned, sequenced and identified as the TbpA and TbpB transferrin-binding receptors (Ogunnariwo et al., 1997).

TbpA and TbpB form a host-specific outer-membrane receptor complex necessary for the removal of iron from transferrin at the cell surface (Gray-Owen & Schryvers, 1996). The TbpB receptor is a largely hydrophilic molecule, believed to be anchored to the cell surface through a lipitated acyl tail (Gray-Owen & Schryvers, 1996). The TbpA receptor is believed to function as an integral membrane gated porin through which iron is translocated after having been removed from transferrin. Upon removal from transferrin and translocation across the outer membrane, iron becomes associated with a periplasmic ferric-ion-binding protein (FbpA). FbpA has been demonstrated to function in complexing and transporting this iron through the periplasmic space (Chen et al., 1993; Kirby et al., 1997; Khun et al., 1998). FbpA is the periplasmic component of an iron ATP-binding cassette (ABC) transporter system (Tam & Saier, 1993), which also includes an inner-transmembrane protein, FpbB, and a cytoplasmic ATPase, FpbC (believed to be necessary for the transport of FbpA-bound iron across the inner membrane and mobilization of iron from FbpA, respectively). This integrated cascade of proteins results in transferrin-bound iron being mobilized from the cell surface, transported through the periplasmic space and deposited into the cytoplasm without the incorporation of the transferrin molecule into the bacterial cell.

In a previous investigation, a 35 kDa iron-regulated protein was identified by comparison of autoradiographic patterns between iodinated total cellular proteins of *P. haemolytica* grown in iron-replete and iron-depleted media (Lainson et al., 1991). An anti-35 kDa mAb demonstrated this protein to be localized in the periplasm. It was proposed that the 35 kDa protein, given its size, location and regulation, may function in the uptake or transport of chelated iron. In this investigation, we undertook to identify and characterize the gene encoding the 35 kDa protein using a *P. haemolytica* phagemid library to complement in trans an iron-utilization-deficient strain of *Escherichia coli*. Evidence is provided that the *P. haemolytica* fbpA gene encodes the 35 kDa protein. Biochemical properties of the *P. haemolytica* FbpA protein are examined and its relationship to other known FbpA proteins is presented.

**METHODS**

**Strains and growth media.** Bacterial strains, plasmids and deoxyoligonucleotides used in this study are shown in Table 1. *P. haemolytica* and *E. coli* strains were both stored in 30% glycerol at -70°C. Fresh *P. haemolytica* cultures were grown on BHI (brain-heart-infusion medium; Difco) plates incubated overnight at 37°C in a 5% CO₂ atmosphere. Fresh *E. coli* cultures were prepared on LA or LT plates [Luria agar (Gibco-BRL) supplemented with 100 µg ampicillin (Sigma) ml⁻¹ or 12.5 µg tetracycline (Sigma) ml⁻¹, respectively] and incubated overnight at 37°C. Liquid cultures of *P. haemolytica* were prepared in BHI broth. Liquid *E. coli* cultures were prepared in LA or LT broth [Luria broth base (Gibco-BRL) supplemented with 100 µg ampicillin ml⁻¹ or 12.5 µg tetracycline ml⁻¹, respectively]. Iron-restricted cultures of *P. haemolytica* were prepared by subculturing overnight BHI cultures into fresh BHI broth to an OD₆₀₀ of 0.2, at which point the cultures were supplemented to 100 µM with EDDHA (ethylene-diamine-di(o-hydroxyphenylacetic acid); Sigma). Whole cells were used for Western blot analysis after an additional 2 h growth at 37°C. NBDA agar [8% (w/v) nutrient broth (Difco), 5% (w/v) NaCl, 1.5% (w/v) agar (Difco), pH 7.0 supplemented to 0.2 mM with 2',2'-dipyridyl (Sigma) and 100 µg ampicillin ml⁻¹ (Eick-Helmerich et al., 1987) and 2x YT broth [16% (w/v) Bactotryptone, 10% (w/v) Bactoyeast extract, 5% NaCl, pH 7.0] were used in the library complementation experiment.

**N-terminal amino acid sequence analysis of the 35 kDa protein.** The 35 kDa protein was semi-purified by reversed-phase chromatography. It was then further purified by separation on SDS-PAGE gel, blotted onto Immobilon-P membrane (Millipore) and recovered for analysis. For internal sequencing, the 35 kDa protein was affinity purified as previously described (Murray et al., 1992). It was then cleaved with CNBr and a 14 kDa internal fragment was N-terminally sequenced. Sequence analyses were carried out on an Applied Biosystems 470 gas-phase protein sequencer at the Babraham Research Institute, Cambridge, UK.

**Complementation of e573 for growth on NBDA media using a *P. haemolytica* lambda ZAP II library.** Briefly, the following protocol [a hybrid of protocols described by Zimmermann et al. (1989) and Williamson & Slocum (1994)] was used to complement e573 (entA) for growth on NBDA media using a *P. haemolytica* h044 lambda ZAP II library (Stratagene). This library had been prepared as previously described for a *Haemophilus influenzae* library (Adhikari et al., 1995). e136 (SURE strain) and e573 (entA) cultures were grown in 50 ml LB broth supplemented with 0.2% maltose and 10 mM MgSO₄ overnight at 37°C in a shaking incubator. After overnight incubation the e573 culture was stored until needed at 4°C. Five microliters of a 10¹⁶ p.f.u. ml⁻¹ *P. haemolytica* lambda ZAP II library was diluted in 195 µl 2x YT broth. Aliquots (200 µl) of the diluted library were mixed with 200 µl of an overnight e136 culture and 10 µl of R408 helper phage (7.5 x 10¹⁶ p.f.u. ml⁻¹) and incubated at 37°C for 15 min. The mixture was added to 5 ml prewarmed (37°C) 2x YT broth in a 15 ml polypropylene tube and incubated for 3 h in a 37°C shaking incubator. The tube was then heated to 70°C for 20 min, centrifuged in a horizontal centrifuge at 4000 g for 15 min and the supernatant transferred to a fresh 15 ml polypropylene tube. The resulting infective ss-Bluescript phagemid was immediately rescued to ppBluescript plasmid by mixing 1 ml phagemid supernatant with 1 ml prewarmed e573 in a stationary 37°C incubator for 20 min. The 2 ml of culture were then added to 50 ml of 5:1 dilution of BHI/NBDA broth.
Table 1. Bacterial strains, plasmids and primers

<table>
<thead>
<tr>
<th>Strain, plasmid or primer</th>
<th>Description</th>
<th>Source</th>
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<tr>
<td><strong>Strains</strong></td>
<td></td>
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<tr>
<td>e136</td>
<td><em>E. coli</em> SURE strain (Tet&lt;sup&gt;R&lt;/sup&gt;)</td>
<td>Stratagene</td>
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<tr>
<td>e157</td>
<td><em>E. coli</em> DH5α F&lt;sup&gt;+&lt;/sup&gt; strain</td>
<td>Novagen</td>
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<tr>
<td>e201</td>
<td><em>E. coli</em> BL21(DE3) pLysS T7 expression strain</td>
<td>Liss (1987)</td>
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<tr>
<td>e573</td>
<td>Enterocin-deficient <em>E. coli</em> strain F&lt;sup&gt;+&lt;/sup&gt;, enA&lt;sup&gt;+&lt;/sup&gt; T7&lt;sup&gt;R&lt;/sup&gt;</td>
<td>R. Kadner, Charlottesville, USA</td>
</tr>
<tr>
<td>e638</td>
<td>e573 with plasmid pH1 integrated (pfbABC operon)</td>
<td>This study</td>
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<tr>
<td>e1103</td>
<td>e157 with plasmid pGEM PHFA</td>
<td>This study</td>
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<tr>
<td>e1124</td>
<td>e157 with plasmid pT7-7 PHFA</td>
<td>This study</td>
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<td>e201 with plasmid pT7-7 PHFA</td>
<td>This study</td>
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<tr>
<td>e1267</td>
<td>e201 with plasmid pT7-7</td>
<td>This study</td>
</tr>
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<td>h944</td>
<td>Serotype A1 <em>P. haemolytica</em> strain</td>
<td>S. Lundberg, Airdrie, Canada</td>
</tr>
<tr>
<td>h908</td>
<td>Serotype A2 <em>P. haemolytica</em> strain</td>
<td>ATCC 33366</td>
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<td><strong>Plasmids</strong></td>
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<tr>
<td>pBluescript SK II(+)</td>
<td>High-copy general cloning vector</td>
<td>Stratagene</td>
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<tr>
<td>pBSPh1</td>
<td>pBluescript SK II(+) vector with recombinant fbpABC insert from h044 lambda ZAP II genomic DNA library</td>
<td>This study</td>
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<td>T/A PCR cloning vector</td>
<td>Promega</td>
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<td>504-505 PHFA PCR product cloned into pGEM vector</td>
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<tr>
<td>pT7-7</td>
<td>T7 expression vector</td>
<td>Tabor (1994)</td>
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<td>pT7-7 PHFA</td>
<td>EcoR1/BamH1 PHFA cloned into similarly cut pGEM vector</td>
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<tr>
<td><strong>Primers</strong></td>
<td></td>
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<tr>
<td>Oligo 504</td>
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<td>This study</td>
</tr>
<tr>
<td>Oligo 505</td>
<td>5′ AGGATTCACAGGGTTAGGACGCAATAC 3′</td>
<td>This study</td>
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and incubated overnight at 37 °C in a shaking incubator. The cells were subcultured to solid media, to provide an environment selective for growth (and subsequent enrichment of the culture) of those clones capable of growing under moderately iron-restricted conditions. After overnight incubation, 200 μl of culture was plated in duplicate on LA agar (to determine the efficiency of the phagemid transfection) and NBDA agar (to select for clones capable of complementing the siderophore-deficient host). Pinpoint colonies on NBDA agar could be identified after 48–72 h incubation at 37 °C. Subsequent colonies were replated on NBDA agar. Colonies capable of regrowing on fresh NBDA agar were screened for siderophore production by plating on siderophore detection media (Schwyn & Neilands, 1987). Clones negative for siderophore production were further investigated.

**PCR and sequence analysis.** PCR reactions were performed using *Taq* thermostable DNA polymerase (BRL) on a Perkin Elmer model 480 thermocycler. The following profile was used to amplify PCR products: 1 × (94 °C for 2 min), 30 × (94 °C for 1 min, 50 °C for 1 min, 72 °C for 1 min) and 1 × (72 °C for 10 min). All other PCR parameters were performed as described by Innis & Gelfand (1990). Plasmids were prepared for sequencing using a double-stranded nested deletion kit (Pharmacia). Derivatives of plasmids were prepared using Qiagen plasmid miniprep columns and sequencing was performed using Thermosequenase fluorescent dye-primer cycle sequencing with 7-deaza-dGTP (Amersham) on an ABI PRISM 377 DNA sequencer.

**Subcloning, overexpression and purification of *P. haemolytica* FbpA (PFbpA).** Oligodeoxynucleotide primers were designed to PCR-amplify the fbpA gene, in order to clone and overexpress the gene product in pT7-7 vector (Tabor, 1994). Upstream primer 504 was designed to amplify the (predicted) native Shine–Dalgarno (underlined, Table 1) and coding (italicized ATG start codon, Table 1) regions of fbpA. It also possesses an EcoR1 site for subcloning purposes (bold region, Table 1). Primer 505 was designed to amplify the intergenic region between fbpA and fbpB, 144 bases downstream of the stop codon of fbpA, and possesses a BamH1 site for subcloning purposes (bold region, Table 1). A 504–505 PCR product amplified from genomic h044 DNA was cloned into pGEM T-vector, constructing pGEM PHFA and strain e1103. The EcoR1/BamH1 fbpA cassette was removed from pGEM PHFA and spliced into similarly cut pT7-7 vector, producing pT7-7 PHFA and strain e1124. pT7-7 PHFA was transformed into the lambda lysogen *E. coli* strain BL21(DE3), which directs expression of T7 RNA polymerase under the control of the IPTG-inducible promoter, lacUV5, constructing strain e1126. PFbpA was overexpressed and purified from strain e1126 using IPTG induction and osmotic shock procedures described for the production and isolation of periplasmic maltose-binding fusion proteins from pMal-P2 (Riggs, 1994). Osmotic shock fluid was lyophilized and resuspended in appropriate buffers for further analysis.

**Electrophoresis and Western blotting.** All SDS-PAGE gels were run with the buffer system of Laemmli (1970). Urea denaturing IEF gels were run using Dry-IEF gels reconstituted with 6 M urea/10 mM Tris (pH 7.2) on a Phast-gel electrophoresis system using carbamylate calibration standards (Pharmacia). TAE (Tris/acetate/EDTA) agarose gels, standard molecular biology techniques and immunological analy-
sis of electroblotted proteins were performed as previously described (Sambrook et al., 1989). Whole cells were applied to SDS-PAGE gels and a previously prepared murine mAb (Lainson et al., 1991) specific for the P. haemolytica 35 kDa protein was used to develop the blots using ECL detection reagent (Amersham).

Buffer exchange, preparation of holo- and apo-PFbpA, wavelength scans and citrate competition assay. To prepare protein for IEF, protein samples were dialysed overnight with one change of 6 M urea/10 mM Tris (pH 7.2). Holo-PFbpA was prepared by resuspending lyophilized protein in 0.1 M sodium citrate/0.1 M NaHCO₃ buffer (pH 8.6), adding 25-fold molar excess FeCl₂ and incubating at room temperature for 10 min. The solution was then either gel-filtered using Econo-Pac 10 DG columns (Bio-Rad) or dialysed against two overnight changes of 10 mM Tris/200 mM NaCl (pH 8.0). Apo-PFbpA was prepared by resuspending the protein with 0.1 M sodium acetate/0.1 M NaH₂PO₄ (pH 5.5), dialysing PFbpA against two overnight changes of 0.1 M sodium acetate/0.1 M NaH₂PO₄/25 mM Na₂EDTA/0.4% (w/v) Chelex 100 (pH 5.5) and one overnight change of 10 mM Tris/200 mM NaCl (pH 8.0). Protein samples were quantified using a Bio-Rad D_{2} Protein Assay and a Beckman DU-64 spectrophotometer. Wavelength scans of holo- and apo-PFbpA were performed on a Beckman DU-640 spectrophotometer. Citrate competition assays of PFbpA iron affinity were performed as described by Chen et al. (1993), except that visible absorbance was monitored at the peak Fe absorbance for PFbpA (419 nm).

**RESULTS**

**Isolation of a P. haemolytica fbpABC operon**

Our approach to cloning the fbpA gene was based on the premise that it would be part of an operon capable of mediating high-affinity iron transport in an E. coli mutant deficient in this process (Zimmerman et al., 1989). We used a siderophore-deficient (entA) strain as a host. This strain was incapable of growing on nutrient broth supplemented with the iron-chelating agent dipyridyl. The entA strain was transfected with a lambda ZAP II phagemid library generated from genomic DNA of a serotype A1 P. haemolytica strain (Ogunnarwio et al., 1997) and selected on the iron-limited medium. Upon selection, colonies were found which were negative for siderophore production, but positive for growth after replating on fresh iron-limited medium. One clone, designated e638, was randomly selected for further analysis.

Plasmid analysis of strain e638 yielded a recombinant plasmid, pBSPH1, with a 5.5 kb insert. Whole-cell Western blots were performed to determine if this recombinant strain expressed the previously identified 35 kDa protein (Lainson et al., 1991). A mAb reacting with the 35 kDa protein isolated from a serotype A2 strain of P. haemolytica was used to screen the electrobots. Extracts from iron-starved cells of both A1 and A2 serotypes were included in the analysis to ensure that the mAb was capable of detecting the 35 kDa protein from a serotype A1 strain (the source of the genomic library). A single protein of 35 kDa was detected in both serotypes of P. haemolytica, reaffirming the conserved antigenic nature of this protein between A serotypes of P. haemolytica (lanes 3 and 4, Fig. 1; Murray et al., 1992). In e638 two bands were detected, a predominant band at 35 kDa and a minor band at a slightly higher molecular mass (lane 1, Fig. 1).

**Sequence analysis of pBSPH1**

DNA sequence analysis of the 5327 bp insert in pBSPH1 identified four ORFs with homology to other known proteins (Fig. 2). Three of these ORFs appeared in tandem arrangement, while a fourth partial coding sequence was identified in opposite orientation to the former genes. The first ORF (fbpA) encoded a protein which bears homology to the iron-binding proteins belonging to the cluster 1 group of periplasmic binding proteins (Tam & Saier, 1993). A Shine–Dalgarno sequence immediately upstream of the fbpA coding sequence corresponds to 9/13 nucleotides of the E. coli consensus sequence (Shine & Dalgarno, 1974). A potential −10 region (TAAGAA) region was also identified upstream of fbpA. The recombinant gene insert ends immediately upstream of this −10 region; however the fusion of the recombinant insert to the pBluescript SK II(+) vector appears to have created an artificial −35 region (TTGATA), generating a functional promoter. Previous work has indicated that FbpA is an iron-regulated protein (Lainson et al., 1991), but unfortunately truncation of the promoter/operator regions in pBSPH1 precludes analysis of this region or the potential search for a fur consensus sequence.

An intergenic region of 187 bp with a number of potential regions of secondary structure separates fbpA from a potential TTG start codon of the next ORF. The second ORF (fbpB) encodes a highly hydrophobic protein with homology to the family of cytoplasmic membrane-spanning proteins believed to interact with the iron-binding cluster 1 periplasmic binding proteins. Based on this TTG start codon, the calculated molecular mass of FbpB would be 58.7 kDa. Two regions of FbpB match the permease consensus sequence EAA--G--------I-LP, common to the transmembrane component of these ABC transporter systems (Saurin et al., 1994; Adhikari et al., 1996). The third gene (fbpC), with a
calculated molecular mass of 37.7 kDa, has the typical signature sequences of an ATPase (Walker A and B motifs; Higgins, 1992) and begins 9 bp downstream of FbpB. The fourth truncated ORF (54.3) illustrates homology to a hypothetical protein of 54.3 kDa (Burland et al., 1995) identified in the genome of E. coli (51% identity with the available P. haemolytica sequence; ~65% of the gene sequenced, based on comparison with the E. coli gene). Two small potential ORFs (Fig. 2) immediately following FbpC [ORF5 of 389 bp (~15 kDa) and ORF 6 of 284 bp (~10 kDa)] show no significant homology to any known proteins, but are reminiscent of the small potential ORF (524 bp; ~19 kDa) following sfrC of Serratia marcescens (Angerer et al., 1990).

The fbpA gene encodes a 38 kDa protein containing a characteristic signal sequence (Fig. 2) which is cleaved from the mature protein following secretion. The predicted size of the mature PFbpA which would be exported to the periplasm is 35.8 kDa. Thus the higher molecular mass protein detected in Western blots of the strain containing pBSPH1 (Fig. 1) corresponds to the 38 kDa unprocessed precursor protein of PFbpA. As pBSPH1 is a high-copy plasmid, the presence of the higher molecular mass PFbpA in E. coli, but not in the native strains, may indicate a saturation of the E. coli periplasmic secretion apparatus. Two regions of the translated PFbpA protein correspond with the amino acid sequence derived from the 35 kDa protein (Table 2), confirming that the 35 kDa protein is encoded by the fbpA gene. The N-terminal amino acid sequence of the intact 35 kDa protein corresponds exactly to that of the translated DNA sequence. In contrast, there are several differences between the experimentally determined amino acid sequence of the 14 kDa polypeptide and the translated sequence from the fbpA gene. This may represent an artifact of DNA- or amino acid sequencing or perhaps is attributable to differences between PFbpA from the A1 and A2 serotypes.

The predicted amino acid sequence of the mature PFbpA
Table 2. N-terminal amino acid sequences of the 35 kDa PFbpA and CNBr-cleaved 14 kDa subfragment, and the corresponding translated DNA sequences of fbpA

Bracketed letters represent questionable amino acid identifications.

<table>
<thead>
<tr>
<th>Sequence source</th>
<th>Peptide sequence</th>
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<tbody>
<tr>
<td>N-terminal sequence</td>
<td>ANEVNVSYRQPYLIEPMLK</td>
</tr>
<tr>
<td>Translated DNA sequence</td>
<td>ANEVNVSYRQPYLIEPMLK</td>
</tr>
<tr>
<td>14 kDa fragment N-terminal sequence</td>
<td>MLDDEKQKS (C)AAAI(N) FPS</td>
</tr>
<tr>
<td>Translated DNA sequence</td>
<td>MLDDEKQKS W AAAC(A) N FPS</td>
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</table>

was aligned with the sequences of homologues found in other bacterial species. Surprisingly, there was greater identity with IdiA (iron-deficiency-induced Protein A) from the cyanobacterium _Synechococcus_ than with FbpA proteins from other pathogens, such as _H. influenzae_, belonging to the Pasteurellaceae (Fig. 3). The analysis also revealed that there are two short regions of identity which appear common to virtually all Fe-binding proteins of this class (TGIKV and PADV) and thus may serve as signatures of this protein family. It is pertinent to note that these do not represent any of the amino acids involved in liganding the metal ion (Bruns et al., 1997). A dendrogram was compiled to illustrate the similarities and relationships of the various iron-ligand-binding proteins relative to one another and to the larger family of cluster 1 extracellular solute-binding proteins (Tam & Saier, 1993), including the carbohydrate-binding proteins MalE, MalX, MsmE and UgpB (Fig. 4). The resulting non-rooted tree was compiled using primary protein sequence comparison and the neighbour-joining method of the CLUSTAL W 1.6 program (Higgins et al., 1996). Confidence in the reliability of the branch points in the dendrogram is illustrated by the boxed bootstrap value assigned at each branch point (out of 10000 bootstrap samples).

**Biochemical characterization of PFbpA**

The _P. haemolytica_ _fbpA_ gene was subcloned into the pT7-7 expression vector in order to provide substantial quantities of protein for further biochemical analysis. Simple overexpression of the recombinant FbpA and isolation of osmotic shock fluid provided a relatively pure preparation of protein. The process of iron loading and dialysis removed some minor contaminants such that the resulting protein was pure enough for bio-

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*Fig. 3.* Primary sequence alignment between the mature proteins of _P. haemolytica_ FbpA, _Synechococcus_ spp. PCC6301 IdiA and _H. influenzae_ FbpA. Boxed are the TGIKV and PADV sequences characteristic of the iron-binding protein subclass of cluster 1 periplasmic binding proteins. Outlined are the iron (I)-binding ligands and in bold are the phosphate (P)-binding ligands known from the _H. influenzae_ FbpA crystal structure (Bruns et al., 1997). Identical residues with reference to the PFbpA primary sequence are illustrated by dots; dashes represent spaces introduced to maximize alignment.
**Fig. 4.** Relationship of *Pasteurella haemolytica* FbpA to other members of the iron-binding protein subfamily and proteins of the cluster 1 group of extracellular solute-binding proteins (Tam & Saier, 1993). Analysis was performed using the Power Macintosh versions of CLUSTAL W 1.6 (Higgins et al., 1996). Illustrated are relative genetic distances between proteins (fractional numbers) and bootstrap branching confidence values (out of 10000 samples, in boxes).
Fig. 5. Coomassie-stained 12% SDS-PAGE gel of holo-PFbpA protein produced by IPTG induction and concentrated osmotic shock fractionation from strain e1126 (pT7-7 PHFA) (lane 1), and an equivalent amount of sample produced by IPTG induction and concentrated osmotic shock fractionation from the negative control strain e1267 (pT7-7) (lane 2). Lane M contains prestained SDS-7B molecular mass markers (Sigma).

Fig. 6. Wavelength spectral scan from 300 to 700 nm of holo-(-) and apo-(- - -) forms of PFbpA.

We initially attempted to isolate the gene encoding the 35 kDa protein using a PCR-based approach, designing oligonucleotides to amino acid sequences generated from the intact and a CNBr-cleaved fragment of the 35 kDa protein. However, we were unsuccessful at amplifying a specific PCR product from *P. haemolytica* chromosomal DNA during these experiments (data not shown). This may be in part attributable to the differences between the N-terminal amino acid and DNA sequences of the 14 kDa peptide (Table 2). We decided to pursue an alternative approach based on the method previously utilized by Zimmermann et al. (1989) to isolate the *sfuABC* operon from *S. marcescens*. We proceeded on the premise (i) that the 35 kDa protein may be an FbpA homologue organized in a three-gene operon similar to that of other *fipABC* operons, and (ii) these iron uptake systems may all share a similar affinity for iron, making it possible to clone the gene for the 35 kDa protein based on the ability of the associated operon to complement an iron uptake mutant of *E. coli*. Using a *P. haemolytica* A1 lambda ZAP I1 library we were successful in identifying a clone capable of complementing an *entA* *E. coli* strain on iron-restricted media. This clone was verified to be negative for siderophore production on the siderophore detection media of Schwyn & Neilands (1987). DNA sequencing using M13F universal primer and Western blotting using an anti-35 kDa mAb verified that the plasmid pBSPH1 possessed a translatable sequence which corresponded with the N-terminal amino acid sequence of the 35 kDa protein, and that the strain e638 produced a 35 kDa protein cross-reactive with the mAb. Overexpression and purification of this protein revealed reversible iron binding with a peak absorbance at 419 nm and a relative iron affinity similar to that reported for FbpAs from *H. influenzae* and *N. gonorrhoeae*.

Complete sequence analysis of pBSPH1 identified three genes in tandem arrangement with motifs characteristic of bacterial ABC importer systems (Higgins, 1992) and in particular, the three-component iron uptake operons within this family of transporters. Members of this ‘operon’ transporter family have been identified in a number of eubacterial species, including: *N. gonorrhoeae* (Berish et al., 1990b), *Neisseria meningitidis* (Berish et al., 1990a), *H. influenzae* (Sanders et al., 1994), *Actinobacillus actinomycetemcomitans* (GenBank accession number 2340838), *S. marcescens* (Angerer et al., 1990), *Yersinia enterocolitica*.

coloration previously reported for holo-*H. influenzae* FbpA, *Neisseria gonorrhoeae* FbpA and human transferrin (data not shown). Citrate competition assays confirmed a relative iron affinity similar to that reported for other FbpAs (approximate $K_d$ of $10^{19}$-$10^{20}$; data not shown). IEF of PFbpA revealed an isoelectric point slightly less than that of *H. influenzae* FbpA at $pI \sim 7.5$ (position $-4$ on a GADPH carbamylation train; Pharmacia) (data not shown).

DISCUSSION

We initially attempted to isolate the gene encoding the 35 kDa protein using a PCR-based approach, designing oligonucleotides to amino acid sequences generated from the intact and a CNBr-cleaved fragment of the 35 kDa protein. However, we were unsuccessful at amplifying a specific PCR product from *P. haemolytica* chromosomal DNA during these experiments (data not shown). This may be in part attributable to the differences between the N-terminal amino acid and DNA sequences of the 14 kDa peptide (Table 2). We decided to pursue an alternative approach based on the method previously utilized by Zimmermann et al. (1989) to isolate the *sfuABC* operon from *S. marcescens*. We proceeded on the premise (i) that the 35 kDa protein may be an FbpA homologue organized in a three-gene operon similar to that of other *fipABC* operons, and (ii) these iron uptake systems may all share a similar affinity for iron, making it possible to clone the gene for the 35 kDa protein based on the ability of the associated operon to complement an iron uptake mutant of *E. coli*. Using a *P. haemolytica* A1 lambda ZAP II library we were successful in identifying a clone capable of complementing an *entA* *E. coli* strain on iron-restricted media. This clone was verified to be negative for siderophore production on the siderophore detection media of Schwyn & Neilands (1987). DNA sequencing using M13F universal primer and Western blotting using an anti-35 kDa mAb verified that the plasmid pBSPH1 possessed a translatable sequence which corresponded with the N-terminal amino acid sequence of the 35 kDa protein, and that the strain e638 produced a 35 kDa protein cross-reactive with the mAb. Overexpression and purification of this protein revealed reversible iron binding with a peak absorbance at 419 nm and a relative iron affinity similar to that reported for FbpAs from *H. influenzae* and *N. gonorrhoeae*.

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(GenBank accession number Z47200) and Actinobacillus pleuropneumoniae (Chin et al., 1996). All of these species are obligate or opportunistic human or veterinary pathogens. It is interesting to note that the sequence similarities between the FbpA components of different species are strongly associated with host and environment, but not necessarily associated with taxonomic groups. Thus, there is little overall sequence similarity between FbpA proteins among members of the Pasteurellaceae [A. pleuropneumoniae (source of porcine pleuropneumonia), P. haemolytica (bovine shipping fever), A. actinomycetemcomitans (human periodontal disease) and H. influenzae (Haemophilus meningitis, otitis media)], while human mucosal pathogens [(A. actinomycetemcomitans, H. influenzae, N. gonorrhoeae (gonorrhea) and N. meningitidis (meningococcal meningitis)] and human enteric species [S. marcescens (urinary tract infection, septicaemia) and Y. enterocolitica (gastroenteritis)] form distinct clusters (Fig. 4).

Most surprising to us was the fact that PFbpA is more closely related to the 34 kDa protein IdiA (≈40% identity) from the free-living cyanobacteria Synechococcus and Synechocystis spp. than to any of the aforementioned FbpA homologues from human or veterinary pathogens (≈20–26% identity; Fig. 3; Klaus-Peter et al., 1996; Kaneko et al., 1995). IdiA has been demonstrated to be involved in the mobilization of both Fe and Mn necessary for growth and photosynthetic activity under Fe and/or Mn limitation in these species. To further compound our surprise, however, was the fact that IdiA is not a periplasmic protein, but rather is associated with thylakoid membranes (photosynthetic pseudoorganelles) within the cytoplasm of the cell. Both in this investigation (on the basis of an osmotic shock purification scheme and biochemical characterization) and in a previous study (Lainson et al., 1991) it has been demonstrated that PFbpA is a periplasmic protein that reversibly and with high affinity associates with iron. As this protein and its cognate inner-membrane receptor complex were cloned functionally on the basis of their ability to rescue a siderophore-deficient strain of E. coli for growth under iron-limited conditions, it can be concluded that the PFbpABC operon is an iron ABC uptake transporter system, despite the similarity of PFbpA to IdiA. It is an interesting example of biological conservation, however, that two highly related iron-binding proteins have been adapted to different roles and subcellular locations within different species of eubacteria.

The recent resolution of the crystal structure of the H. influenzae FbpA (HPfbpA) protein (Bruns et al., 1997) has provided some insight into the origins and function of this class of proteins. Structural data reveal that HFbpA is, in addition to being an iron-binding protein, a phosphate-binding protein. Complexed exogenous phosphate provides a necessary free O ligand for iron binding and is a structural analogue of the carbonate anion necessary for iron binding by transferrins and lactoferrins. Both transferrin N-lobe and HFbp bind iron with a similar spectrum of ligands (H249, Y95, Y188, D63 and two O ligands from exogenous CO3 in transferrin; H9, Y195, Y196, E57, one O ligand from exogenous PO4 and one O ligand from H2O in HFbp), however the source of these ligands is non-analogous regions of the primary structure of these proteins. There is also extremely low sequence identity between the N-lobe of transferrin and HFbp (≈10%). This, coupled with the fact that HFbp has more in common structurally with sugar- and anion-binding periplasmic proteins than transferrin, has led to the proposal that high-efficiency iron-binding has evolved independently multiple times over the course of evolution and, in the case of HFbp, has resulted from selective pressure for a phosphate-binding protein to associate avidly, but reversibly, with iron. A central a-helix implicated in anion binding appears to be the most structurally universal motif conserved between HFbp, sulfate- and phosphate-binding proteins.

Despite low sequence identity, two of the four amino acids involved in iron binding (Y198 and Y199) based on primary sequence alignment can be identified by alignment of PFbpA with HFbp (Fig. 3). In addition, three of the six amino acids implicated in PO4 binding in HFbp (Q58, N175, N193 and the a-helix A141) are also present in PFbp, including N196 and what appears to be a homologue of the central anion a-helix: S137, G138, K139. Currently ongoing crystallographic analysis of this protein should reveal whether it also binds phosphate or instead uses an alternative exogenous anionic ligand. Differences in the anion- or iron-binding ligands in PFbpA may be responsible for the blue-shifted absorbance spectra of this protein. Preliminary crystallographic diffraction data reveal that PFbpA cannot be exactly modelled on the HFbpA crystal structure, indicating that structural differences do exist between these two proteins (D. McRee, personal communication).

Early in our work we had anticipated a large degree of sequence conservation in the FbpA homologues between the various pathogenic members of the Pasteurellaceae, including the representative species P. haemolytica (bovine pathogen), A. pleuropneumoniae (porcine pathogen) and H. influenzae (human pathogen). This postulate was based on the proposal that FbpA represents a critical component of transferrin-receptor-mediated iron uptake pathways and, due to constraints of protein–protein interactions with other components of the uptake pathway (i.e. FbpB and possibly TbpA), would be restricted in the degree of permissible variation which would continue to allow this system to remain functional. Furthermore, as FbpA is a periplasmic entity it would not be expected to be under immunological pressure for variation and therefore should not vary more than any cellular ‘house keeping’ genes. It must be stated that it has been suggested that FbpA may be transiently surface-exposed (Gomez et al., 1996). However, the majority of the evidence, including that for PFbpA, suggests that this protein is predominately a periplasmic entity (Lainson et al., 1991; Berish et al.,
Within the cluster 1 group of periplasmic binding proteins, PFbpA and, to a greater extent, A. pleuropneumoniae FbpA, stand out as highly divergent from other members of this iron-binding protein subfamily (Fig. 4). It could be suggested that this divergence may have been the result of isolation of these pathogens from other members of this protein family (Schryvers & Gonzalez, 1990). However, the strong degree of identity between not only P. haemolytica fbpA gene and that of homologues identified within Synecococcus and Synecocystis spp., but also between P. haemolytica FbpB (~42% identity vs ~17–24% of other FbpBs) and P. haemolytica FbpC (44% identity vs ~32–38%) and homologues within these cyanobacteria is a curiosity, as obviously the cyanobacteria have diverged from the Pasteurellaceae long before the development of host specificity within the Pasteurellaceae. We cannot, however, give a rational explanation for how such a relationship could have developed, although the limitations of phylogenetic analysis and the complications of horizontal genetic transfer have been well-documented by others (Syvanen, 1994).

Although resistance to challenge infection by P. haemolytica serotype A1 in sheep has been demonstrated to be correlated with a rise in titre of anti-35 kDa serum antibodies (possibly anti-PFbpA) (Mosier et al., 1989) and the 35 kDa protein is highly immunogenic in sheep (Lainson et al., 1991), it is unlikely that the 35 kDa protein (PFbpA) alone could make an effective immunoprophylactic target given its lack of surface accessibility. However, it is unknown whether the strong humoral response to PFbpA could augment overall immune response to P. haemolytica infection. It has recently been suggested that FbpA could make a potential chemotherapeutic target, given the recent understanding of the mechanism of iron binding revealed by the crystal structure of HFbpA (Bruns et al., 1997). This is supported by the facts that iron acquisition is a critical component of bacterial pathogenesis in vivo and FbpA is a focal point for high-affinity iron uptake from both free iron sources and iron complexed by transferrin and lactoferrin within the host. Despite low overall sequence identity, it appeared that the mechanism of iron binding by FbpA was universally conserved between different species, as the core iron-binding ligands could be identified in all representative members of this protein; an attractive feature for a broad-spectrum target. However, the recent identification of ‘outliers’ for which these iron-binding ligands cannot be identified in primary sequence alignments of PFbpA and, to a larger extent, A. pleuropneumoniae FbpA underscores the question of whether alternative mechanisms of iron binding exist in these related pathogens. Structural information on these proteins will answer this question and may reveal whether FbpA should be further pursued as a target for antibacterial therapy.

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