Affinity purification and characterization of a fibrinogen-binding protein complex which protects mice against lethal challenge with *Streptococcus equi* subsp. *equi*

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Cell-wall-associated proteins from *Streptococcus equi* subsp. *equi*, the causative agent of strangles, were analysed with a view to identifying a potential protective antigen. Preparations of these proteins, isolated from mutanolysin extracts of cell walls, were shown to contain one major high-M, protein species (apparent M, 220000 and 550000 when analysed by SDS-PAGE and gel-filtration chromatography, respectively). The high-M, protein bound horse fibrinogen and was purified under non-denaturing conditions using fibrinogen affinity chromatography. The fibrinogen-binding protein (FgBP) reacted with serum taken from horses recovering from strangles and protected mice against lethal challenge from *S. equi* subsp. *equi*. The sequence of the corresponding gene (*fbp*) was determined and shown to encode a mature protein (M, 54597) with predicted coiled-coil structure. An FgBP truncate, lacking the C-terminal cell wall/membrane anchor domain, was overexpressed in and purified from *Escherichia coli* and was shown to behave in an analogous fashion to the wild-type product in terms of M, estimation, fibrinogen binding and seroreactivity.

**Keywords**: *Streptococcus equi* subsp. *equi*, fibrinogen-binding protein, protective antigen

INTRODUCTION

The group C streptococcus *Streptococcus equi* subsp. *equi* is the causative agent of strangles, a highly contagious disease of the upper respiratory tract of the family *Equidae* (Timoney, 1993). The disease is initially characterized by nasal discharge and fever, followed by abscess formation in local lymph nodes. In severe cases, infection can become disseminated (bastard strangles) eventually leading to death. Strangles continues to be of major economic importance to the horse industry. However, existing vaccines against strangles seem to afford little protection (Yelle, 1987; Timoney, 1988) and there is a clear need for a more efficacious product.

In Gram-positive cocci, one group of proteins which have received much attention as important virulence factors and as potential protective antigens are those associated with the cell wall. These proteins share a number of structural and functional features, not least of which is their ability to bind host plasma and/or extracellular matrix proteins (reviewed by Kehoe, 1994; Patti et al., 1994; Goward et al., 1993). Amongst the best-studied wall-associated proteins are the M protein family of *Streptococcus pyogenes*. These dimeric, fibrillar molecules bind several host proteins including fibrinogen (Fg), albumin and IgG, are clearly major antiphagocytic virulence factors, and may have a role in adhesion (Kehoe, 1994; Hasty et al., 1992; Fischetti, 1991). There is also a large body of evidence to suggest that they are protective (Bronze et al., 1992, 1988; Bessen & Fischetti, 1990; D’Alessandri et al., 1978; Polly et al., 1975). Other protective cell-wall-associated proteins include the fibronectin (Fn)-binding protein of staphylococci (Schennings et al., 1993), the pneumococcal PspA protein (AlonsoDeVelasco et al., 1995) and the *Streptococcus mutans* surface protein antigen I/II (Katz et al., 1993; Okahashi et al., 1989).

**Abbreviations**: Fg, fibrinogen; FgBP, fibrinogen-binding protein; Fn, fibronectin; IDA, iminodiacetic acid; i.p., intraperitoneal(ly); MALDI-TOF, matrix-assisted laser desorption/ionization time-of-flight; s.c., subcutaneous(ly).

The GenBank accession number for the sequence reported in this paper is AFO12927.
A protective anti-phagocytic M-like protein has been extracted from whole cells of *S. equi* subsp. *equi* (Timoney, 1993; Timoney & Mukhtar, 1993; Boschwitz *et al.*, 1991; Jean-François *et al.*, 1991; Galán & Timoney, 1988). Several studies have shown that binding of Fg to *S. equi* subsp. *equi* cells confers resistance to phagocytosis, and Timoney and co-workers have additionally provided evidence that M-like protein may be involved in this binding (Boschwitz & Timoney, 1994). Here we report on the identification, purification and overexpression of a major cell-wall-associated protein, which we have called fibrinogen-binding protein (FgBP), of *S. equi* subsp. *equi* which binds horse Fg, and provide evidence for its protectively immunogenic potential. We also describe the cloning and sequencing of the gene (fbp) encoding FgBP, and the overexpression in *Escherichia coli* of an FgBP truncate possessing properties analogous to those of the native protein. During preparation of this manuscript, an independent study by Timoney *et al.* (1997) detailed the cloning and sequencing of two M-like proteins from *S. equi* subsp. *equi*. The relationship of these proteins to our own is discussed.

**METHODS**

**Bacterial strains, plasmids and growth conditions.** The virulent isolate, strain TW, of *S. equi* subsp. *equi* was obtained from the late Dr Paul Storm (Intervet International BV, The Netherlands). *Escherichia coli* strains DH5α (Life Technologies) and XLOLR (Stratagene) together with the plasmids pGEM7 (Promega) and pBK-CMV (Stratagene) were used for subcloning and sequencing. *E. coli* strains LE392 (Promega) and XL-1 Blue MRF' (Stratagene) were used for propagation of *λ* phages. *E. coli* XL-1 Blue (Stratagene) and plasmid pQE30 (Qiagen) was used for expression of the recombinant FgBP. *S. equi* subsp. *equi* was grown at 37 °C on Columbia base agar supplemented with 5% (v/v) horse blood or statically overnight in Todd–Hewitt broth supplemented with 0.2% (w/v) yeast extract. *E. coli* strains were grown at 37 °C in either LB medium or in 2x YT medium (Lech & Brent, 1997). Where appropriate, medium was supplemented with ampicillin (100 µg ml⁻¹), kanamycin (50 µg ml⁻¹), IPTG (1 mM) or 2% (w/v) glucose.

*S. equi* TW cells used in mouse experiments were harvested by centrifugation (16000 g, 15 min, 4 °C), washed twice in PBS (0.2 cultulture volume), resuspended in PBS (0.01 culture volume) and suspensions stored at −70 °C. On the day of challenge, an aliquot of frozen cells was thawed and diluted appropriately in PBS, and c.f.u. confirmed by plating appropriate dilutions in triplicate onto blood agar plates.

**Isolation of cell-wall-associated proteins from envelopes of** *S. equi*. *S. equi* cells from broth cultures were harvested by centrifugation (16000 g, 15 min, 4 °C), washed once in 10 mM Tris/HCl buffer pH 7.2 (Tris buffer), and finally resuspended in Tris buffer containing DNase (50 µg ml⁻¹), RNase (50 µg ml⁻¹) and protease inhibitors (2 mM PMSF, 2 mM benzamidine hydrochloride). Bacteria were then lysed by two passages through a French pressure cell (221 MPa) and unlysed cells were removed by centrifugation (3000 g, 10 min, 4 °C). Cell envelopes (membranes plus cell walls) were pelleted from the cleared lysate (45000 g, 1 h, 4 °C) and washed three times in Tris buffer. Cell-wall-associated proteins were isolated using either of two procedures (A or B).

**Purification of horse Fg.** Contaminating Fn was first removed from horse Fg (Sigma) by adsorption onto a gelatin affinity column (Mosher *et al.*, 1980) prepared by covalently linking gelatin to CNBr-activated Sepharose 4B. Horse Fg, eluted as unbound material, was then dialysed against 0.1 M sodium acetate (pH 5.0), and a precipitate of lipoproteins removed by centrifugation (16000 g, 15 min, 4 °C). Contaminating horse IgG was then adsorbed onto a protein G affinity column using the procedure recommended by the manufacturer (Amersham). Purified Fg (free of Fn and IgG) eluted as unbound material and was then dialysed against distilled water and lyophilized.

**Biochemical procedures.** SDS-PAGE was performed using either a 12.5% or a 7.5% (w/v) polyacrylamide separating gel and a 4.5% (w/v) polyacrylamide stacking gel (Laemmlli, 1970). Samples were routinely heated for 5 min at 100 °C in Laemmlli sample buffer (Laemmlli, 1970) prior to electrophoresis. Proteins were detected with Coomassie brilliant blue or by silver staining (McVeigh *et al.*, 1988). Molecular masses were determined from the relative mobilities of the following standard molecular mass marker proteins: α-2-macroglobulin (340 kDa), rabbit myosin (205 kDa), β-galactosidase (116 kDa), phosphorylase b (94 kDa), bovine serum albumin (66.2 kDa), catalase (61 kDa), glutamate dehydrogenase (55.4 kDa), fumarase (48.5 kDa), alcohol dehydrogenase (41 kDa), Omp F protein of *E. coli* (38.3 kDa), carbonic anhydrase (30 kDa), chymotrypsinogen (26.5 kDa), trypsin inhibitor (20.1 kDa), lysozyme (14.3 kDa). Defined peptide
fragments of FgBP were generated essentially as described by Cleveland et al. (1977). Excised SDS-PAGE slices containing purified FgBP were re-electrophoresed on a second SDS-PAGE gel (12.5 %, w/v, polyacrylamide) in the presence of staphylococcal V8 protease. Resultant peptides were then blotted onto Problot membranes and visualized with amido black. N-terminal amino acid sequence analysis was performed on excised bands using an Applied Biosystems 47A pulsed-liquid protein sequencer.

Gel filtration chromatography was performed at 4 °C using Sephacryl S-300 HR, an elution buffer consisting of 20 mM Tris/HCl, 200 mM NaCl and 0.2% (w/v) NaOEt (pH 7.4) and a flow rate of 5 ml h⁻¹. Molecular masses were estimated from a consideration of the elution profiles of the following standards: ferritin (450 kDa), catalase (240 kDa), aldolase (158 kDa), BSA (68 kDa), hen egg albumin (45 kDa), chymotrypsinogen (25 kDa) and cytochrome c (12.5 kDa).

Protein concentration was estimated by a modification (Dulley & Grieve, 1975) of the Lowry method using bovine serum albumin as standard. Matrix-assisted laser desorption/ ionization time-of-flight (MALDI-TOF) mass spectrometry with performance by Dr Len C. Packman (Department of Biochemistry, University of Cambridge, UK).

Immunological and affinity procedures. Western immunoblotting was performed as described by Caffery et al. (1988) using 5 % (w/v) dried skimmed milk as blocking reagent, peroxidase-labelled affinity-purified anti-mouse IgG (Sigma), anti-horse IgG H + L (ICN) or anti-rabbit IgG H + L (ICN) as localizing antibodies and 4-chloro-1-naphthol as developing reagent. The procedure for Fg affinity blotting was similar to that for Western immunoblotting except that 2 % (w/v) dried skimmed milk was employed and peroxidase-labelled horse Fg (0.1 mg ml⁻¹; prepared as described by Harlow & Lane, 1988) was used as a localizing reagent. The procedure for affinity blotting using recombinant FgBP as a probe was also performed essentially as described above using 0.1% (v/v) Tween 20 as blocking agent. Blots were probed with 5 µg FgBP ml⁻¹ and bound FgBP was detected with rabbit anti-FgBP antibodies followed by peroxidase-labelled affinity-purified anti-rabbit IgG, H + L antibodies (ICN). Rabbit anti-FgBP antibodies were generated by standard procedures (Owen, 1985).

Test-tube affinity-preparation experiments were performed using a modification of the immunoprecipitation technique described by Doherty et al. (1986). Fg (300–600 µg) in sodium barbital hydrochloride buffer (pH 8.6) was preincubated for 1 h at 25 °C with 10 mM EDTA, protease inhibitors and then centrifuged (13 000 g, 15 min, 20 °C). Triton X-100 (final concentration 2%, v/v) was added to the cleared supernatant. Mutanolysin extracts containing FgBP (50–100 µg) were also centrifuged as above to clear any sedimentable material. Solutions containing Fg and FgBP were then mixed and incubated for 4 h at 25 °C. Resultant affinity precipitates were harvested by centrifugation (13 000 g, 15 min, 20 °C), washed twice in 200 µl 0.1 M NaCl containing 0.5% (v/v) Triton X-100 and finally resuspended in Laemmli sample buffer. Affinity diffusion assays and analysis of excised affinity precipitates were conducted using established techniques (Owen, 1985, 1986).

ELISA to detect anti-FgBP IgG was carried out using standard procedures (Newell et al., 1988). Wells of microtitre plates were coated with 2.5 µg purified FgBP in 50 µl 0.1 M sodium carbonate buffer (pH 9.6), blocked in 1% (w/v) bovine serum albumin and incubated with serial twofold dilutions of mouse sera prediluted 1:250 in blocking solution. A 1:2000 dilution of peroxidase-labelled affinity-purified anti-mouse IgG was used as secondary antibody and 3,3',5,5'-tetramethylbenzidine as substrate.

Mouse protection experiments. All animal experiments were performed in compliance with EC directive 86/609/EC as implemented in Ireland under Statutory Instrument 17/94. Laca mice (8–12 weeks old) were immunized subcutaneously (s.c.) on days 0 and 28 with 30 µg purified FgBP emulsified in a 200 µl volume of MPL + S-TDCM Ribi adjuvant (active ingredients monophosphoryl lipid A and trehalose dimycolate; RIBI Immunochim Research). Control mice were immunized s.c. either with PBS or with RIBI adjuvant emulsified in PBS. On day 34, blood samples (100–200 µl) were taken by cardiac puncture from all mice except from the control group for whom the procedure proved too stressful. All mice were challenged on day 42 by intraperitoneal (i.p.) injection of 1 × 10⁷ c.f.u. S. equi TW cells in 200 µl volumes of PBS and were monitored for 32 d post challenge. For humane reasons, mice were killed if they were considered terminally ill with body temperatures below 32 °C.

Isolation of chromosomal, plasmid and bacteriophage DNA. Genomic DNA, isolated from S. equi TW by treatment with mutanolysin, lysozyme and Sarkosyl (Yu & Ferretti, 1989), was extracted twice with hexadecyltrimethylammonium bromide/chloroform followed by stepwise extraction with phenol/chloroform and chloroform/isooamyl alcohol (Wilson, 1997). DNA was finally precipitated with 0.6 vol. 2-propanol, and ethanol-washed pellets were resuspended in 10 mM Tris/HCl containing 10 mM EDTA (pH 8.0). Plasmid and phagemid DNA was purified from E. coli by a modified alkaline lysis method (Feliciello & Chinalli, 1993) and recombinant λ DNA was purified from phage lysates as detailed by Maniatis et al. (1982).

Construction of genomic libraries of S. equi subsp. equi TW. Two libraries were made in lambda using Sac3A-digested S. equi genomic DNA. The first was made in λgem11 and involved ligation to the left and right arms of the λgem11 (precleaved with Xhol) and in vitro packaging as outlined by the manufacturer (Promega). The second was made in λZAP Express, and involved ligation to vector arms (predigested with BamHI) and packaging using Gigapack III Gold packaging extract as described by the manufacturer (Stratagene).

Screening of S. equi genomic libraries and isolation of positive subclones. The degenerate oligonucleotide 5' CAR-AARGCNAARGAYGARCG 3' (R = A + G, N = A + G + C + T, Y = C + T), made from a knowledge of the amino acid sequence QKAKDER of FgBP, was 5' end-labelled with [32P]dATP. λgem11 recombinant phages were propagated on E. coli LE392 and hybridizations with labelled oligonucleotide were performed at 50 °C as outlined by O'Reilly et al. (1988). Blots were then washed at 37 °C for 15 min in 5 × saline sodium citrate (SSC), for 15 min in 2 × SSC and at 42 °C for 15 min in 1 × SSC and positive plaques were detected by autoradiography. One positive reacting plaque (S6E12) was identified, resuspended in phage buffer (20 mM Tris/HCl, pH 7.4, containing 0.1 M NaCl and 10 mM MgSO₄) and propagated to homogeneity. A 1.8 kb Sacl fragment from S6E12 was labelled with [32P]dATP and used as a hybridization probe with the S. equi TW genomic DNA.

Protective fibrinogen-binding protein from S. equi
resistance helper phage and transformed into *E. coli* XLORL according to the manufacturer’s instructions (Stratagene). One recombinant phagemid (termed pFBF700) was chosen for further study.

**DNA sequencing.** The sequencing strategy involved the generation of plasmid subclones containing overlapping deletions. The nucleotide sequence of template DNA was then determined using the ABI PRISM Dye Terminator Cycle Sequencing Ready Reaction Kit (Perkin Elmer), commercial primers and, where appropriate, specific primers synthesized from a knowledge of established DNA sequence.

**Expression and purification of a FgBP truncate.** The region of *fbp* encoding the first 434 amino acids of the mature protein was cloned into pQE30. The *fbp* fragment in question was first amplified by PCR under standard conditions using *pfu* polymerase (Stratagene), *S. equi* chromosome as template, and 5’ CGCCGGATCCAACTCTGGTATTGCTGAGC 3’ and 5’ TTATGGTCTCAAGCGATTGATTTC 3’ as forward and reverse primers, respectively (underlined nucleotides in the above primers correspond to complementary sequences in the *fbp* gene). The amplified product was digested with *BamHI* and ligated to the vector pQE30 (digested with *BamHI* and *SalI*). The recombinant plasmid was transformed into *E. coli* XL-1 Blue and the positive transformant *E. coli* MM21 isolated following growth on L-agar supplemented with ampicillin and glucose.

The hexahistidyl-tagged FgBP truncate was purified using metal chelate affinity chromatography as follows. *E. coli* MM21 was grown in 2 X YT medium, supplemented with glucose, to OD660 ~0.9. Following induction for 3 h in the presence of 1 mM IPTG, bacterial cells were harvested, washed, resuspended in binding buffer (20 mM Tris/HCl, 0.5 M NaCl, pH 7.9) and lysed by passage through a French pressure cell (221 MPa). The lysate was then cleared by centrifugation (45000 g, 1 h, 4°C), and the soluble fraction applied to a nickel-iminodiacetic acid (IDA) column which was then washed extensively with binding buffer until the A280 of the effluent reached zero. Hexahistidyl-tagged FgBP truncates were recovered using a 0-1 M gradient of imidazole in binding buffer.

**RESULTS**

**Isolation of cell-wall-associated proteins**

Cell-wall-associated proteins isolated from *S. equi* TW by mutanolysin digestion of purified wall material (see Methods, procedure A) showed the presence of several protein species following analysis by SDS-PAGE. The most dominant of these had an apparent M₅₀ of 220000 (Fig. 1, lane 2) and will be referred to as the high-M₅₀ protein. Other minor proteins of apparent M₅₀ 94000, 74000 and 56000 were also detected following mutanolysin digestion (Fig. 1, lane 2). In contrast, negligible amounts of these proteins were released from control undigested cell wall (Fig. 1, lane 3). Furthermore, the dominant high-M₅₀ protein was barely detectable in the SDS-PAGE profile of SDS-solubilized envelopes (Fig. 1, lane 1) but was a prominent feature of envelopes pretreated with mutanolysin (Fig. 2, lane 2). These data provide convincing evidence that these proteins are specifically anchored to the (SDS-insoluble) cell wall.

**Fg binding**

Several lines of evidence suggest that the high-M₅₀ protein binds horse Fg. Firstly, a single precipitate was clearly demonstrated following affinity diffusion experiments conducted with horse Fg and either mutanolysin extracts or purified high-M₅₀ protein (Fig. 1, inset). Secondly, SDS-PAGE analysis of washed affinity precipitates (Owen, 1986) obtained following either affinity diffusion or test-tube affinity precipitation experiments revealed the dominant components to be the high-M₅₀ protein and the three characteristic polypeptides (α, M₅₀ 76000; β, M₅₀ 61000; γ, M₅₀ 51000) of reduced horse Fg (Fig. 1, lanes 4 and 5). In contrast, neither protein formed a precipitate when incubated alone (data not shown). Finally, affinity blotting experiments performed on preparations of cell-wall-associated proteins showed that the high-M₅₀ species reacted strongly with labelled
Protective fibrinogen-binding protein from *S. equi*

**Fig. 2.** Purification of the high-M, FgBP from *S. equi* TW. Samples, analysed by SDS-PAGE using a 12.5% (w/v) polyacrylamide separating gel, were stained for protein with Coomassie brilliant blue. Lanes: 1, cell envelope fraction; 2, cell envelope fraction following incubation with mutanolysin; 3, supernatant fraction obtained following incubation of cell envelopes with mutanolysin; 4, proteins eluted unbound from the Fg-affinity column; 5, material purified by Fg-affinity chromatography. The position of the high-M, FgBP is indicated. Also indicated to the left of the SDS-PAGE are the positions to which standard molecular mass marker proteins (sizes shown in kDa) migrated.

(a) Protective effect of FgBP against lethal *S. equi* infection in mice. A group of 10 mice were immunized (s.c.) with 50 µg affinity-purified FgBP in Ribi adjuvant 42 and 14 d prior to challenge. Five control mice were immunized with PBS emulsified in Ribi adjuvant only and five mice were immunized with PBS alone. All mice were subsequently challenged (i.p.) with $1.5 \times 10^8$ c.f.u. of *S. equi* cells. ○, Controls; □, vaccinates.

(b) Serum IgG response to FgBP in vaccinated and unvaccinated mice, as monitored by Western immunoblotting. Purified FgBP (lanes 1–19) was analysed by SDS-PAGE using a 12.5% (w/v) polyacrylamide separating gel and was electroblotted onto nitrocellulose. Each lane was probed with an individual serum sample taken from a panel of ten vaccinated mice (lanes 1–10) and from a panel of eight unvaccinated (control) mice (lanes 11–18). Lane 19 was probed with anti-mouse IgG antibodies alone. The position of the FgBP is indicated.

Purification of the FgBP

The ability of the high-M, FgBP bound rabbit Fg but did not appear to bind human Fg, horse Fn, horse IgG or collagen (data not shown).

SDS-PAGE analysis of the purified protein revealed it to have an apparent $M_r$ (220000) which was unaffected by prolonged boiling (up to 30 min) in sample buffer. Following analysis on 7.5% (w/v) polyacrylamide gels (cf. 12.5%, w/v, polyacrylamide gels), the protein ran as a more diffuse band (apparent $M_r$ 160000–190000). During gel filtration chromatography on Sephacryl S300 HR the purified protein showed the elution characteristics ($V_e/V_o = 1:21$) of a molecule with an estimated $M_r$ ($\sim 500000$) somewhat larger than that of ferritin ($M_r$, 450000). MALDI-TOF mass spectrometry failed to give signals for the protein in the working range of the instrument.

Protection and Western immunoblotting experiments

The protectively immunogenic potential of the FgBP was tested in a mouse model in which mice, immunized s.c. with purified high-M, FgBP, were subsequently challenged with a lethal dose of virulent *S. equi* TW. All vaccinates were protected following challenge ($P < 0.01$; see Fig. 3a). Furthermore, at no stage did any vaccinated mouse show clinical signs of illness, nor could *S. equi* be isolated from their peritoneal fluid at the conclusion of the trial (day 32 post-challenge). In contrast, mortality in the control group of mice was 80% (Fig. 3a), and *S. equi* was isolated from peritoneal
Fig. 4. Serum IgG response to cell-wall-associated proteins of S. equi subsp. equi in horses recently recovered from strangles. Supernatant fractions obtained following incubation of cell envelopes with mutanolysin (lanes 1 and 3) and mutanolysin-extracted cell envelopes (2 and 4) were analysed by SDS-PAGE using a 12.5% (w/v) polyacrylamide separating gel and were electroblotted onto nitrocellulose. Lanes 1 and 2 were probed with serum taken from one of seven horses recently recovered from strangles and lanes 3 and 4 were probed with serum taken from a healthy foal. The profile of reactive proteins was similar for all convalescent horse sera tested. The positions and molecular masses (in kDa) of major reacting proteins are indicated at the left of the Western blots.

Cloning and sequencing of fbp

V8 proteolytic digestion of FgBP generated at least six peptides of M, <14 000 (data not shown). N-terminal amino acid sequence analysis yielded the following two distinct sequences (in single-letter code): (1) NSEVSRTATPRL... and (2) LQKADERQALTESFNTL5.

An S. equi genomic library, constructed in ZAP Express, was probed with a degenerate oligonucleotide made from a consideration of amino acid sequence (2) above. One positive clone (ZSE12) containing a 15 kb insert was identified. Subcloning and subsequent DNA sequence analysis of restriction fragments from ZSE12 revealed it to contain a truncated fbp gene encoding the first 488 amino acids of the precursor FgBP. A 1.8 kb Sac1 fragment from ZSE12 containing the promoter region plus the first 1307 bp of the fbp gene was then used as a probe to screen a second genomic library in λ ZAP Express. Several positive phagemid clones were identified and one (pFBP700), containing a 3.8 kb insert in pBK-CMV, was chosen for further experiments. DNA sequence analysis revealed the presence of an open reading frame of 1605 bp encoding the FgBP. It has been reported that cloning of S. equi M proteins can be frustrated by DNA rearrangements (Timoney et al., 1991). However, direct nucleotide sequence analysis of the fbp gene amplified by PCR from the chromosome confirmed the sequence of the above cloned fbp and provided good evidence that no such DNA rearrangements had occurred.

Sequence analysis of fbp and FgBP

The fbp open reading frame is preceded by sequences typical of promoter signals (−10, tataat; −35, tgtcat) and ribosome-binding sites (gagagg) and is followed by a sequence resembling a transcriptional terminator, suggesting that the gene product is translated from a monocistronic message. The translated product has the following features (Fig. 5a). The first 36 residues of the deduced amino acid sequence show features characteristic of a signal sequence (von Heijne, 1986). This region shows strong homology (46–62% identity) with signal sequences of other streptococcal proteins (see Goward et al., 1993). Immediately after the predicted signal cleavage site is a stretch of amino acids corresponding to one of the two sequences (no. 1) obtained by direct amino acid sequencing of V8 fragments of FgBP. The second sequence (no. 2) is located internally in FgBP. The fbp gene encodes a primary translation product of 534 amino acids (M, 58344) which is post-translationally processed to yield a putative mature protein of 498 amino acids (predicted M, 54597).

The sequence of the mature protein contains features characteristic of cell wall-associated proteins (see Fig. 5a). Firstly, it possesses a high content of alanine, glutamic acid, leucine and lysine as reported for the M proteins of S. pyogenes. In addition, the protein contains two blocks (A, residues 226–272; B, residues 357–405) of degenerate repeated sequences. The A1 and A2 repeats are 21 amino acids in length and differ only in one amino acid. A3 is a partial repeat of about five amino acids. The B repeats are 14 amino acids long and are repeated three times. The B1 and B2 repeats possess 64% identical residues and are separated by a short stretch of seven amino acids. The B2 and B3 repeats possess 57% identical residues. Finally, the C-terminal part of the protein contains features characteristic of cell wall/membrane-anchoring domains of cell wall proteins. Thus, a short region (482–499) showing a relative...
enrichment of Gly/Pro residues is followed by the consensus LPSTGE motif. This is followed almost directly by 21 hydrophobic amino acids (membrane-spanning region) and a short tail of five charged amino acid residues. The sequence of the membrane-anchoring domain (from and including LPSTGE) exhibits strong homology (68-74% residue identity) with corresponding regions in other streptococcal M proteins.

Secondary structure analysis according to the algorithm of Garnier et al. (1978) predicts that 70% of the FgBP is \( \alpha \)-helical. More refined structural predictions using the COILS program (Lupas, 1996; Lupas et al., 1991) suggest that about 60% of the mature protein possesses an extremely high probability (95-100%) of coiled-coil structure, with zero to very low probabilities of such structures in the first 147 residues of the mature protein and at the putative C-terminal wall/membrane anchor region (see Fig. 5b). Analysis by the MULTICOIL program (Wolf et al., 1997) indicates that FgBP most probably assumes a two-stranded coiled-coil structure (Fig. 5c) and that the likelihood of trimer formation is low (data not shown). Similar two-stranded coiled-coil structures have also been proposed for the M proteins from S. pyogenes (Nilson et al., 1995).

Computer-assisted searches of existing databases (Altschul et al., 1990) revealed that FgBP showed no significant homology, at the nucleotide and amino acid level (except in the signal sequence and wall/membrane-spanning regions), to sequences published for most surface proteins from group C streptococci. These include the M-like protein, the Fn-binding protein (FNZ) and the macroglobulin/albunin/lgG-binding (ZAG) protein of S. equi subsp. zooepidemicus (Lindmark et al., 1996; Jonsson et al., 1995; Timoney et al., 1995), and the Fg/albunin/lgG-binding protein (FAI) from a horse-derived group C streptococcus (Talay et al., 1996). Nor did the sequence show any homology to the partial sequences published earlier by Timoney and co-workers (Timoney et al., 1991; Galán et al., 1988) for an M-like protein from S. equi. However, comparison of the sequence of fbp with that recently published for SeM of S. equi subsp. equi (Timoney et al., 1997) reveals near identity – eleven differences in nucleotide sequence translating into six differences in deduced amino acid sequence (at residues 8, 58, 63, 143, 181 and 410 of the FgBP precursor protein).

### Expression, purification and properties of an FgBP truncate

The region of fbp encoding the first 434 amino acids of the mature protein (Fig. 5) was cloned in-frame into the pQE30 expression system. This vector places a hexa-histidine affinity tag at the N-terminus of the resultant fusion protein. SDS-PAGE analysis revealed that resultant transformants expressed two major proteins (with apparent \( M_r \) values of 200000 and 90000), which were not present in lysates of E. coli XL-1 Blue harbouring pQE30 alone and which could be purified by Ni\(^{2+}\)-IDA affinity chromatography. Of these, the species of higher \( M_r \) was the more dominant (Fig. 6). The recombinant truncate, which could be purified in yields of about 3-5 mg per litre of culture, retained many of the properties of native FgBP (Fig. 6, lane 3). Thus, it gave an estimated \( M_r \) following gel filtration of \( \sim 550000 \). In addition, the dominant species observed during SDS-PAGE (Fig. 6, lane 4) migrated with apparent \( M_r \) values several fold higher than that (\( M_r \) 49541) anticipated from consideration of sequence. Finally, both major species bound horse Fg (Fig. 6, lane 5), could be purified by Fg affinity chromatography (data not shown), reacted with antiserum to native FgBP (Fig. 6, lane 6) and gave identical profiles to those shown in Fig. 6 (lane 6) when tested in Western immunoblot experiments conducted with sera from vaccinated mice and from various convalescent horses (described in Figs 3 and 4). In the above Western blots, and to a lesser extent in Fg affinity blots, a third reacting band (apparent \( M_r \) 49000) could be observed for purified truncate preparations (Fig. 6, lanes 5 and 6). This protein, which may represent the monomeric form of the molecule, was present at low levels since it was barely detectable in SDS-gels stained with Coomassie blue (Fig. 6, lane 4). All three bands observed for the purified recombinant FgBP truncate reacted during Western blotting experiments with anti-

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**Fig. 5.** Schematic representation of the primary and secondary structure of FgBP. (a) The primary structure of FgBP showing positions of the signal sequence (Ss), the A and B repeats, the wall-spanning region (WSR) and membrane spanning region (MSR). (b) Probability of coiled-coil structure as predicted by the COILS web server (http://cole3. unil.ch/software/COILS_form.html) using the MultiCoil web server (http://ostrich.ics.mit. edu/cgi-bin/multicoil) using a window of 28 residues. (c) Probability of a coiled-coil dimer as predicted by the MultiCoil web server (http://ostrich.ics.mit. edu/cgi-bin/multicoil) using a window of 28 residues. The probability ranges applicable to (b) and (c) are given as a key at the bottom of the figure.
Although previous studies (Timoney et al., 1997; Boschwitz & Timoney, 1994) have suggested that surface proteins from *S. equi* subsp. *equi* may be involved in binding horse Fg, the present report gives the first description of the dominant molecular species likely to be involved in such reactions. Thus, it is clear from binding experiments and SDS-PAGE analysis of affinity precipitates and of mutanolysin extracts, that the (220 kDa) high-\(M_r\) protein is the major Fg-binding component and that it represents the major cell-wall-associated protein of the organism. Whereas the balance of evidence strongly suggests that the protein binds horse Fg (but not horse Fn, IgG and collagen), the possibility cannot be discounted that additional binding to other host components also occurs. We also report for the first time a simple scheme based on the ligand-binding properties of the molecule which facilitates its purification under non-denaturing conditions. Previous attempts to purify such M-like proteins have generally involved denaturing conditions (Timoney et al., 1997; Boschwitz et al., 1991; Timoney & Trachman, 1985), which have often resulted in preparations enriched in fragments of the native protein.

Timoney et al. (1997) described, for an American isolate (CF32) of *S. equi* subsp. *equi*, two distinct M-like proteins (termed SeM and SzPSe) which, on the basis of ELISA assays, appeared to bind equine Fg. Of the two, SeM was the more dominant opsonizing antigen for *S. equi*. It is clear from sequence comparison that FgBP is essentially identical to SeM, the few differences in amino acid sequence serving to further reinforce the remarkable uniformity previously noted amongst diverse *S. equi* isolates (Timoney et al., 1997; Jorm et al., 1994; Galán & Timoney, 1988). FgBP shows very little homology with SzPSe and affinity blotting experiments, SDS-PAGE analysis and N-terminal amino acid sequencing of V8 proteolytic fragments have failed to demonstrate the presence of a second Fg-binding component in our own purified preparations. The close similarity in behaviour between our purified high-\(M_r\) FgBP and a recombinant derivative (see below) also supports this conclusion.

It is clear that the \(M_r\) values of the molecule as estimated by SDS-PAGE (~ 220000) and gel filtration chromatography (~ 550000) are greatly at variance with that (\(M_r\) 54597) predicted from consideration of gene sequence. Several possible explanations could be invoked to account for these phenomena. These include (a) anomalous migration during SDS-PAGE and gel filtration, (b) formation of heat-stable oligomers, and (c) formation of a high-\(M_r\) aggregate composed of the \(M_r\) 54597 FgBP and other protein species and/or cell wall carbohydrate (Boschwitz et al., 1991).

In respect of (a) above, unusual behaviour during SDS-PAGE and gel filtration might be anticipated since FgBP is predicted to be a coiled-coil fibrous molecule (Fig. 5), and estimations of \(M_r\) in these systems are routinely based on the use of globular protein standards. Certainly, aberrant behaviour of other cell-wall-associated proteins during SDS-PAGE has been observed by several authors (Hartford et al., 1997; Jönsson et al., 1991; Hollingshead et al., 1986; Fahnestock et al., 1986). A relevant example is the 89 kDa Fg-binding protein (clumping factor, ClfA) from *Staphylococcus aureus* which migrates during SDS-PAGE with an apparent \(M_r\) (> 200000) more than twice that predicted (Hartford et
Neither for staphylococcal ClfA nor for S. equi FgBP can this phenomenon be attributed to the presence of the putative wall-spanning domain, since in both cases the ratio of apparent to predicted $M_r$ is largely unaffected for recombinant truncates in which this region has been deleted or reduced in size (Fig. 6; Hartford et al., 1997). In respect of (c) it seems unlikely that the high-$M_r$ FgBP observed during SDS-PAGE and gel filtration is a complex of more than one molecular species, since a recombinant truncate lacking the putative wall/membrane anchor domain and produced in a heterologous (Gram-negative) expression system behaves in an almost identical fashion to the wild-type product in terms of $M_r$ estimation, Fg binding and seroreactivity. This does not eliminate the possibility that the native FgBP isolated from S. equi by mutanolysin digestion contains small residual amounts of (covalently) bound peptidoglycan. A more likely explanation for the high-$M_r$ estimations is the existence of stable homooligomeric complex(es) which only partially dissociate in the presence of hot SDS, i.e. situation (b) above. Certainly, the apparent $M_r$ values and relative amounts of the three immuno/Fg-reactive bands observed for purified recombinant FgBP following SDS-PAGE are not incompatible with the presence in a largely tetramer population of low concentrations of monomer and dimer. The existence of homooligomeric structures is also supported by the prediction that coiled-coil dimers (but not trimers) occur with very high probability over an extended region of the molecule. Unfortunately, predictions for higher-stranded structures (i.e. $>3$) are compromised at present since available matrices on which such predictions might be based are compiled from comparatively small databases (Lupas, 1996). Given this and the problems in assessing $M_r$ values of fibrous molecules (see above), it is difficult to be definitive in regard to the oligomeric structure of native FgBP. Nevertheless, the results of SDS-PAGE, gel filtration and MALDI-TOF mass spectrometry are collectively not at variance with a dominant species composed of four or more FgBP monomers.

A major difference between our FgBP and the SeM of Timoney et al. (1997) is the behaviour during SDS-PAGE, SeM focusing as a doublet (apparent $M_r$ values 56000 and 58000; Timoney et al., 1997), FgBP as a multimer. Whereas the precise reasons for this discrepancy remain uncertain, it may reflect in part differences in concentrations and in the relative ease of extraction of monomeric and oligomeric forms of the molecule. In our hands, time-course experiments have indicated that maximal extraction of the high-$M_r$ FgBP is only achieved following prolonged incubation (18 h) of envelopes with mutanolysin in the presence of protease inhibitors, incubation for 1 h generating 10–20-fold less product. It should, however, be noted that mutanolysin extracts and preparations of recombinant FgBP often show the presence of a relatively minor protein species in the predicted monomer $M_r$ range (see Figs 1 and 6). Analogous extracts from the Timoney group are generally prepared by incubation of whole cells) with mutanolysin for 1 h (Galán & Timoney, 1987, 1988; Boschwitiz et al., 1991) and are analysed by selective immunoenhancing techniques (Western immunoblotting) rather than by direct protein staining. Some of these extracts additionally show the presence of a high-$M_r$ immunoreactive band (estimated $M_r$ 120000–140000) speculated to represent M-like protein bound to cell wall carbohydrate (Timoney & Mukhtar, 1993; Boschwitiz et al., 1991; Galán & Timoney, 1985). Thus, shorter incubation periods, perhaps dictated in part by a need to minimize protoplast lysis, may facilitate the release of the monomer form, but not of the more dominant multimeric form, the efficient extraction of which requires more extensive degradation of the peptidoglycan layer. Certainly, given the high probability of coiled-coil structure predicted from this study, one might anticipate that oligomeric form(s) would predominate in situ.

The results from experiments with small animals, shown in Fig. 3, represent the first clear demonstration of the protectively immunogenic potential of a defined and purified unadenatured protein from S. equi. Previous studies in the mouse model have utilized acid extracts of whole cells, culture supernatants, immune horse sera and monoclonal antibodies directed against acid-extracted protein (Jean-François et al., 1991; Timoney & Trachman, 1985). The possibility that protection is elicited by some minor component/contaminant within the high-$M_r$ FgBP complex seems highly unlikely given that a recombinant truncated FgBP expressed in E. coli reacts with immune sera in an analogous manner to native FgBP. Our observations that (a) the high-$M_r$ FgBP is protective in mice (and also in the target species; M. Meehan & P. Owen, unpublished data), and (b) a recombinant FgBP truncate can be readily purified from E. coli in yields 15–30-fold higher than that of the native streptococcal product, together with the complementary observations that SeM antiserum is strongly opsonogenic for S. equi (Timoney et al., 1997), give hope that an efficacious vaccine based on this product may soon be a reality.

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