Expression and immunogenicity of six putative variable surface proteins in *Mycoplasma mycoides* subsp. *mycoides* SC

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Variable surface protein Vmm and five Vmm-type proteins from *Mycoplasma mycoides* subsp. *mycoides* SC were analysed to determine whether these proteins are expressed in vivo in animals affected by contagious bovine pleuropneumonia (CBPP) and in vitro. Recombinant versions of these proteins were constructed and expressed in *Escherichia coli* after mutation of the TGA Trp codons to TGG. These proteins were then analysed by dot and Western blotting with sera from CBPP-affected cattle. Furthermore, affinity-purified polyclonal antibodies to the recombinant proteins were used in Western and colony blotting to look for expression of the putative Vmm-type proteins in cultured *M. mycoides* SC. This study demonstrates that immunoglobulins in CBPP sera recognize all putative Vmm-type proteins tested, indicating that these proteins or their homologues are expressed by mycoplasmas during natural infections. Vmm and one of the putative Vmm-type proteins showed variable expression in vitro.

INTRODUCTION

*Mycoplasma mycoides* subsp. *mycoides* small colony type (*M. mycoides* SC) belongs to the class *Mollicutes* and is the causative agent of contagious bovine pleuropneumonia (CBPP), a severe disease affecting cattle. It is a notifiable disease that has to be reported to the World Organization for Animal Health (OIE). CBPP is a vast problem in Africa, where 18 countries reported 168 outbreaks in 2005 according to the OIE. The last outbreaks in Europe were in Italy (1993), Spain (1994) and Portugal (1999). The occurrence of CBPP in Asia has recently been reported from several countries, the latest being Kuwait in 2006 (World Animal Health Information Database, OIE).

The use of antibiotics, such as tylosin and tetracyclines, for treatment of CBPP is a much-debated subject. It has been considered ineffective and a potential promoter of silent carriers of the disease (Provost, 1996). However, newer antibiotics, such as danafloxacin, have been shown to reduce the spread of CBPP (Hubschle et al., 2006). The slaughter of infected herds still appears to be the most effective means to eradicate CBPP (Windsor & Wood, 1998). This strategy has been used in Europe, but the disease has re-emerged in every decade of the 20th century (Nicholas et al., 2000). CBPP is endemic in much of sub-Saharan Africa, and eradication by mass slaughter would be too expensive and have severe consequences (Kusiluka & Sudi, 2003; Windsor, 2000). In Botswana, CBPP was successfully eradicated in 1995 by a stamping-out strategy (Windsor & Wood, 1998) which turned out to be directly correlated with increased malnutrition in children (Boonstra et al., 2001). Extensive vaccination campaigns seem to be the best option for Africa (March, 2004; Windsor, 2000). However, recent vaccination campaigns have given poor results due to weak vaccine effects (Nicholas et al., 2000; Thiaucourt et al., 1998, 2000). The vaccines currently in use are live strains with reduced pathogenicity, and the vaccine currently recommended by the OIE, strain T1-44, has disadvantages such as lack of long-term immunity (Kusiluka & Sudi, 2003), late protection (1–3 months for primo-vaccination).

**Abbreviations:** ABP, albumin binding protein; CBPP, contagious bovine pleuropneumonia; FBS, fetal bovine serum; HRP, horseradish peroxidase; IMAC, immobilized metal affinity chromatography; pAb, polyclonal antibody.

A supplementary table listing the primers used in PCR, mutagenesis and DNA sequencing is available with the online version of this paper.
(Thiaucourt et al., 1998) and poor protection in vaccine trials (Thiaucourt et al., 2000). Pathogenicity of T1-44 has also been shown (Mbulu et al., 2004).

Little is known of the pathogenicity of M. mycoides SC. No secreted toxins or surface receptors mediating adhesion or cellular responses in host tissues have been reported. There are factors associated with pathogenesis, although their precise functions are unclear. The galactan capsule, an oligosaccharide layer surrounding the cell membrane, appears to promote binding to host tissues, convey resistance to phagocytosis and have toxic effects, and may cause autoimmune responses by structural similarities to bovine pneumogalactan (Buttery et al., 1976; Nicholas & Bashiruddin, 1995). Oxidative damage to the host tissue from hydrogen peroxide (\(H_2O_2\)) produced by glycerol metabolism in M. mycoides SC (Miles et al., 1991) may contribute to CBPP lesions and has been described for other mycoplasma species (Tyrion & Baseman, 1992). Five immunogenic lipoproteins of M. mycoides SC, LpA (Cheng et al., 1996; Monnerat et al., 1999), LpB (Vilei et al., 2000), LpC (Pilo et al., 2003), LpQ (Abdo et al., 2000) and Vmm (Persson et al., 2002), have recently been described; however, their biological functions or possible roles in pathogenicity are, to our knowledge, still unknown.

Many mycoplasmas express surface proteins that undergo reversible changes to alter the antigenic repertoire at the cell and population levels (Citti & Rosengarten, 1997; Razin et al., 1998), as an adaptation to evade the host immune response. Some variable surface proteins are involved in adhesion and immunomodulation (Le Grand et al., 1996; Sachse et al., 2000; Washburn et al., 1993). Only two variable surface proteins are known in M. mycoides SC (Gaurivaud et al., 2004; Persson et al., 2002). One of them, Vmm, is a small lipoprotein of 17 kDa whose expression is regulated at the transcriptional level by the number of \((TA)_n\) repeats in the promoter spacer. Fourteen putative proteins with similar promoters are present in the M. mycoides SC genome (Westberg et al., 2004), and are referred to as Vmm-type proteins in this study.

The primary aim of this study was to investigate six of the Vmm-type proteins, to detect whether humoral immune responses are raised against them in vivo, if they are expressed in vitro and if the expression in vitro is variable. An additional aim was the development of an optimized scheme for high-throughput production of recombinant M. mycoides SC proteins in Escherichia coli. This would enable the screening of a large number of proteins for possible selection as target antigens for the development of diagnostic tests and recombinant vaccines.

**METHODS**

**In silico analysis.** Since several of the 14 Vmm-type proteins have high similarity to each other, a subset of six of them with low similarity to each other [determined by BLAST (Altschul et al., 1997)] were selected for this investigation. This subset included Vmm. Their peptide sequences were retrieved from the genome sequence at EMBL/GenBank/DDBJ entry BX293980 and analysed with TMHMM (Krogh et al., 2001) and SignalP (Nielsen et al., 1997; Nielsen & Krogh, 1998) to identify transmembrane regions and signal peptide sequences, respectively, LipoP (Juncker et al., 2003) to predict lipoproteins, and InterProScan (Quevillon et al., 2005; Zdobnov & Apweiler, 2001) to identify protein motifs.

** Primer design and expression vector.** PCR primers (see Supplementary Table S1 available with the online version of this paper) were designed to amplify the whole genes except the signal peptide sequences. NotI or AscI restriction sites were added to the 5' end of the primers to allow directed insertion into the vector pAff8c (Larsson et al., 2000). The reverse primer was biotinylated to enable solid-phase cloning, and a 3C protease cleavage site (not used in this study) was included in the forward primer handle to enable removal of the His_{6}-albumin binding protein (ABP) fusion tag of the recombinant protein. The His_{6} moiety of the fusion tag allows purification by immobilized metal affinity chromatography (IMAC), while the ABP (Nygren et al., 1988) enhances solubility and is immunopotentiating (Libon et al., 1999; Sjolander et al., 1997). Names of the recombinant proteins were derived from the corresponding ORF names (Westberg et al., 2004) from EMBL/GenBank/DDBJ accession number BX293980.

** M. mycoides SC strains.** The M. mycoides SC type strain PG1\(^T\) was grown in F medium (Bolske, 1988). Genomic DNA was prepared and purified by proteinase K lysis and phenol:chloroform extraction. Total RNA was isolated with Trizol reagent (Life Technologies) from 200 ml culture. For colony immunoblotting, M. mycoides SC strains PG1\(^T\) and M223/90, a pathogenic strain from Tanzania (Bolske et al., 1995), were grown on F medium agar plates.

** Cloning.** PCR was performed with 20 ng template DNA using AmpliTaq DNA polymerase (Roche). The biotinylated PCR products were immobilized on Dynabeads M280-streptavidin paramagnetic beads (Dynal Biotech), and the bound PCR fragments were washed and cleaved with NotI (New England Biolabs). The buffer was replaced, and amplicons were released from the beads by AscI (New England Biolabs) digestion. Cleaved fragments were thereafter ligated into AscI/NotI-digested pAff8c plasmid using T4 DNA ligase (Fermentas) and the constructs were heat-shock transformed into E. coli strain BL21(DE3) cells (Novagen). Correct clone sequences were verified by sequencing. Single-stranded DNA template was generated using TempliPhi (GE Healthcare) prior to cycle sequencing with BigDye Terminator chemistry (Applied Biosystems). The sequencing reactions were analysed on an ABI PRISM 3700 sequencer (Applied Biosystems) and the data analysed using Sequencer software (Gene Codes). To substitute the mycoplasma TGA tryptophan codon with the E. coli TGG tryptophan codon, the QuikChange Multi Site-Directed Mutagenesis kit (Stratagene) was used. Plasmid preparations were made with the QIAprep Spin Miniprep kit (Qiagen).

** RT-PCR.** For expression analysis, RT-PCR was performed with the SuperScriptIII One-Step RT-PCR system with a Platinum Taq kit (Invitrogen) in 40 cycles. First-strand synthesis was performed with 1 \(\mu\)g RNA and 15 pmol reverse primer at 50 °C for 30 min.

**Protein expression and purification.** The recombinant proteins were expressed in E. coli BL21(DE3) and were purified by IMAC (Porath et al., 1975), as previously described (Stein et al., 2006). Purified samples were diluted from 6 to 1 M urea with PBS (2 mM NaH2PO4, 8 mM Na2HPO4, 150 mM NaCl) and were subsequently concentrated on a Vivapore 10/20 concentrator (Vivascience) to a final volume of 1.5 ml. The protein concentrations were estimated...
with the bicinchoninic acid (BCA) assay, and the protein-50 assay (Agilent Technologies) was used to measure protein purity.

**Antibodies and sera used in immunoblotting.** Sera from 15 CBPP cases were used in this study. Four were from an outbreak in Botswana in the middle of the 1990s (G1–4), eight were from an outbreak in Namibia in May 2004 (1MUK15A–17MUK15A), one was an experimental infection from Namibia in 2001 (Exp. Inf.), one serum was from an outbreak in Tanzania in 1997 (PW227), and the final serum was from Kenya in 1998 (C102). For information on CBPP diagnostic test results, see Table 1. Fetal bovine serum (FBS; Gibco) and sera from five healthy Swedish cattle were used as negative controls. The sera were diluted 1:400 and 1:5000; both concentrations were used for each serum. They were blocked with an *E. coli* lysate and purified His$_6$–ABP to prevent false signals due to interaction with residual *E. coli* proteins in the recombinant protein samples or to the His$_6$–ABP tag which originates from streptococcal protein G. For the pre-adsorption experiments, 500 µg of the six recombinant Vmm-type proteins was added to separate 1 ml aliquots of serum 15MUK15A, prepared as described above.

**Immunoblotting.** The recombinant proteins were dot-blotted on nitrocellulose membranes (Bio-Rad Laboratories). The proteins (2 µg of each) as well as His$_6$–ABP and a PG1$^T$ lysate for negative and positive controls were applied to the membrane. The membranes were blocked in PBST [PBS, 0.1 % (v/v) Tween 20] containing 25 % (v/v) horse serum (Gibco), followed by incubation with the bovine sera for 60 min at room temperature. Bound antibodies were detected with secondary goat anti-bovine IgG conjugated with horseradish peroxidase (HRP; 4 ng ml$^{-1}$, Jackson Immunoresearch) and a chemiluminescent HRP substrate (Pierce Biotechnology). The membranes were developed in a ChemiDoc XRS detection system (Bio-Rad Laboratories). For Western blotting, 3 µg recombinant proteins and/or a PG1$^T$ lysate were first separated by SDS-PAGE in a 10–20 % Tris/HCl Criterion Precast gel (Bio-Rad Laboratories) and then electrotransferred onto a nitrocellulose membrane (Bio-Rad Laboratories). Membranes were washed for 5 min in Tris-buffered saline (TBS; 1 mM Tris, 15 mM NaCl, pH 7.5) prior to staining with Ponceau S (Sigma-Aldrich; 1:20 in MilliQ water). Blocking and serum screening were performed as described for dot blotting.

**Immunizations and affinity purification of antibodies.** Rabbits were immunized with the purified recombinant Vmm-type proteins in accordance with national guidelines (Swedish permit A84-02). The protocols for immunization and subsequent affinity purification of antibodies have been described previously (Nilsson et al., 2005). Briefly, the serum was depleted of antibodies reactive to the His$_6$–ABP fusion tag, followed by enrichment of Vmm-type-protein-specific antibodies on an affinity column containing the recombinant proteins, and removal of all other antibodies. Collected antibody fractions were diluted 1:1 with 87 % (v/v) glycerol and 0.02 % (v/v) NaN$_3$, and stored at −20 °C. Antibodies were named after the corresponding recombinant Vmm-type protein, e.g. polyclonal antibody (pAb) A117 was produced from recombinant protein R117, etc.

**Colony immunostaining.** Freshly grown mycoplasma colonies were transferred to nitrocellulose membranes and blocked in TBST [TBS, 0.05 % (v/v) Tween 20] containing 10 % (v/v) horse serum, followed by incubation with the affinity-purified pAbs (diluted 1:50, to 1.5–6.6 µg ml$^{-1}$) for 1.5 h at room temperature. The Vmm-specific mAb 5G1 (9 µg ml$^{-1}$) (Brocchi et al., 1993; Persson et al., 2002) was used as a positive control. Bound antibodies were detected with HRP-conjugated swine anti-mouse or goat anti-rabbit IgG (DakoCytometry) at a final concentration of 0.3 µg ml$^{-1}$ using 4-chloro-1-naphthol as substrate. The membranes were also stained with Ponceau S in order to identify negative colonies.

**RESULTS**

**Selection and *in silico* analysis of Vmm-type proteins**

The following protein genes were selected from the group of 15 Vmm-type proteins: MSC$_0117$, MSC$_0364$,...

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### Table 1. Information on the sera used in immunoblotting experiments

<table>
<thead>
<tr>
<th>Serum</th>
<th>Origin</th>
<th>Year</th>
<th>CBPP status</th>
<th>Retested</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1–4</td>
<td>Botswana</td>
<td>1990s</td>
<td>Positive</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>1MUK15A</td>
<td>Namibia</td>
<td>2004</td>
<td>CFT positive*</td>
<td>LAT positive†</td>
<td></td>
</tr>
<tr>
<td>2MUK15A</td>
<td>Namibia</td>
<td>2004</td>
<td>CFT negative*</td>
<td>LAT positive†</td>
<td></td>
</tr>
<tr>
<td>4MUK15A</td>
<td>Namibia</td>
<td>2004</td>
<td>CFT positive*</td>
<td>LAT positive†</td>
<td></td>
</tr>
<tr>
<td>6MUK15D</td>
<td>Namibia</td>
<td>2004</td>
<td>CFT negative*</td>
<td>LAT positive†</td>
<td></td>
</tr>
<tr>
<td>13MUK15A</td>
<td>Namibia</td>
<td>2004</td>
<td>CFT negative*</td>
<td>LAT positive†</td>
<td></td>
</tr>
<tr>
<td>15MUK15A</td>
<td>Namibia</td>
<td>2004</td>
<td>CFT positive*</td>
<td>LAT positive†</td>
<td></td>
</tr>
<tr>
<td>16MUK15A</td>
<td>Namibia</td>
<td>2004</td>
<td>CFT positive*</td>
<td>LAT positive†</td>
<td></td>
</tr>
<tr>
<td>17MUK15A</td>
<td>Namibia</td>
<td>2004</td>
<td>CFT positive*</td>
<td>LAT positive†</td>
<td></td>
</tr>
<tr>
<td>Exp. Inf.</td>
<td>Namibia</td>
<td>2001</td>
<td>CFT positive*</td>
<td>LAT positive†</td>
<td>Intubated with local strain</td>
</tr>
<tr>
<td>PW277</td>
<td>Tanzania</td>
<td>1997</td>
<td>CFT positive‡</td>
<td>ND</td>
<td>CFT 1:40</td>
</tr>
<tr>
<td>C102</td>
<td>Kenya</td>
<td>1998</td>
<td>ELISA positive‡</td>
<td>LAT positive‡</td>
<td></td>
</tr>
<tr>
<td>Swe 1–5</td>
<td>Sweden</td>
<td>2007</td>
<td>Negative</td>
<td>ND</td>
<td>Healthy Swedish cattle</td>
</tr>
<tr>
<td>FBS</td>
<td>France</td>
<td>–</td>
<td>Negative</td>
<td>ND</td>
<td>Supplied by Gibco, tested mycoplasma-free</td>
</tr>
</tbody>
</table>

MSC_0816, MSC_0847, MSC_1033 and vmm MSC_0390. Two proteins have an amino acid sequence similar to that of other proteins in M. mycoides SC according to BLAST comparisons in which the cut off was set to 80% identity for at least 80% of the sequence length. MSC_0816 is similar to MSC_0812, MSC_0815, MSC_0818 and MSC_0817, and MSC_1033 is homologous to MSC_1058. Sequence analysis with SignalP and TMHMM revealed a signal peptide for each protein, but no transmembrane regions were identified. Analysis with Lipop identified five proteins (excluding MSC_0816) as lipoproteins, which was consistent with the genomic annotation. Further analysis with InterProScan revealed no additional signature/pattern/motif information except for MSC_0816, which had InterPro motifs IPR005046 and IPR011889 in one region. Both motifs exist in families of proteins with unknown functions. Many of these proteins contain a tandem peptide repeat sequence of 25 or 26 amino acid residues (26 aa in MSC_0816, repeated twice) found in predicted surface proteins, often lipoproteins. The InterPro database contains 85 mollicute proteins that match the IPR005046 motif, of which several belong to the M. mycoides cluster. See Table 2 for a summary of in silico analysis.

Cloning, in vitro mutagenesis and recombinant protein production

In order to mimic the protein that is presented to the host immune system, recombinants were designed based on the full-length Vmm-type proteins, the signal peptide excepted. The average size of the signal peptide sequences that were excluded from each protein was 24 aa. Plasmids containing the gene fragment fused to a His6–ABP tag were constructed, and as many as nine TGA tryptophan codons were successfully substituted within a gene by using up to four codon substitutions per reaction. All six recombinant proteins were successfully expressed and purified, yielding protein concentrations of 0.5–4.9 mg ml\(^{-1}\) from 100 ml culture.

Humoral immune responses to the Vmm-type proteins in CBPP-diseased bovines

In order to analyse whether native Vmm-type proteins are expressed in mycoplasmas during infection, the corresponding recombinant Vmm-type proteins were subjected to dot and Western blotting against 15 bovine sera from four CBPP outbreaks, as well as five sera from healthy Swedish cattle and FBS as negative controls. Dot blotting was performed as an initial screening to allow analysis under non-denaturing conditions. An M. mycoides SC strain PG1\(^T\) lysate was used as a positive control, and the recombinant fusion partner His\(_6\)–ABP and an E. coli whole-cell lysate were used as negative controls. The results (Fig. 1, Table 3) showed that all CBPP-positive sera contained antibodies that bound several recombinant proteins. Interestingly, the signal intensities varied between the sera. Generally, R816 was the predominant spot, while Vmm (R390) was absent or indistinguishable from signals seen in some of the negative controls, which is also demonstrated in the blot detected with a pool of all disease sera. There were no detectable false-positive signals due to the His\(_6\)–ABP tag; however, a weak reaction to the E. coli lysate was seen for some sera. The five control sera from healthy bovines and FBS showed that there is no general cross-reactivity of serum IgGs to the recombinant proteins. A faint positive signal, considered to be noise, did occur for four of the control sera.

Additional dot-blotting experiments were performed to confirm the specificity of the serum antibodies. Serum 15MU/K15A was aliquoted into seven fractions, of which six were pre-adsorbed with individual recombinant proteins prior to the dot-blotting analysis, and the remainder were used untreated. An example of the results with R816 is given in Fig. 2. The protein-specific signals were significantly reduced in the pre-adsorbed sera compared to untreated sera, while all other proteins acted as controls and reacted with equal signal intensities, indicating specific binding of the serum antibodies to the recombinant proteins. Since a more concentrated PG1\(^T\)

### Table 2. Summary of bioinformatic analyses of the selected Vmm-type proteins

<table>
<thead>
<tr>
<th>ORF</th>
<th>Promoter repeat</th>
<th>Lipobox</th>
<th>Conserved domain</th>
<th>Molecular mass (kDa)*</th>
<th>Recombinant protein ID</th>
<th>Molecular mass, recombinant protein (kDa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSC_0117</td>
<td>poly(TA)</td>
<td>VVA ↓ C</td>
<td></td>
<td>24.1</td>
<td>R117</td>
<td>42.6</td>
</tr>
<tr>
<td>MSC_0364</td>
<td>poly(TA)</td>
<td>TAS ↓ C</td>
<td></td>
<td>15.6</td>
<td>R364</td>
<td>33.9</td>
</tr>
<tr>
<td>MSC_0390</td>
<td>poly(TA)</td>
<td>VVA ↓ C</td>
<td></td>
<td>3.7</td>
<td>R390</td>
<td>22.2</td>
</tr>
<tr>
<td>MSC_0816</td>
<td>poly(A)</td>
<td>–</td>
<td>IPR011889</td>
<td>43.5</td>
<td>R816</td>
<td>62.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>IPR005046</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MSC_0847</td>
<td>poly(A)</td>
<td>TVS ↓ C</td>
<td></td>
<td>26.8</td>
<td>R847</td>
<td>44.8</td>
</tr>
<tr>
<td>MSC_1033</td>
<td>poly(TA)</td>
<td>VIA ↓ C</td>
<td></td>
<td>6.3</td>
<td>R1033</td>
<td>25.6</td>
</tr>
</tbody>
</table>

* Molecular mass calculated using the ProtParam tool (www.expasy.org), excluding signal peptides.

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lysate was used, the PG1T signal is stronger in comparison to Fig. 1.

To verify protein size and protein purity, and to identify the recombinant proteins, Western blot analyses were used. Each membrane was Ponceau S-stained to visualize total protein content followed by specific detection with one of the 15 CBPP sera, a His6-ABP-directed antibody, control sera or secondary HRP-conjugated antibodies alone. A selection of dots is shown in Fig. 3 and all results are compiled in Table 3. The His6-ABP-antibody staining showed clear and strong bands at the expected sizes of the recombinant proteins and for the His6-ABP control. Protein R117 had a second weak band of smaller size, indicating partially degraded protein or proteolytic fragments, which was further substantiated by the serum-detected blots that also showed bands of smaller sizes. A comparison of His6-ABP and Ponceau S stains showed that only small amounts of contaminating proteins were present, although protein R364 had several bands corresponding to E. coli proteins.

Fig. 1. Screening for immunoreactivity of the six Vmm-type proteins, here shown as dot blots of recombinant proteins and detected with 15 different bovine sera from CBPP-affected cattle, a pool of these sera, and six negative controls, including five healthy Swedish cattle and FBS. A representative blot from a total of three experiments is shown for each serum. Each membrane strip contained whole-cell lysate of M. mycoides SC strain PG1T as a positive control, the recombinant proteins R117, R364, R390 (Vmm), R816, R847 and R1033, and the negative controls His6-ABP (Tag) and E. coli whole-cell lysate. Binding of serum antibodies to a protein is indicated by a dark spot. Blots with sera C102, pooled sera and Swe 1–5 were made with a more concentrated PG1T lysate than the other blots.

<table>
<thead>
<tr>
<th>Serum</th>
<th>Method</th>
<th>Protein</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>R117</td>
</tr>
<tr>
<td>1MUK15A</td>
<td>DB</td>
<td>+/−</td>
</tr>
<tr>
<td>2MUK15A</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td>4MUK15A</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td>6MUK15D</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td>13MUK15A</td>
<td>DB</td>
<td>+/−</td>
</tr>
<tr>
<td>15MUK15A</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td>16MUK15A</td>
<td>DB</td>
<td>+/−</td>
</tr>
<tr>
<td>17MUK15A</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td>Exp. Inf.</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td>G1</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td>G2</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td>G3</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td>G4</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td>PW227</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td>C102</td>
<td>DB</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3. Summary of dot blot (DB) and Western blot (WB) results from screening the recombinant proteins R117, R364, R390, R816, R847 and R1033 with 15 CBPP-diseased bovine sera

Signal intensities in dots and bands: +++, strongly positive; +, positive; +/−, weak or inconclusive; −, negative. The results were judged from three separate sets of dot blot and two separate Western blot experiments performed under standardized conditions with two different dilutions of sera.
M. mycoides SC lysate varied among the sera even with identical aliquots of the same lysate. Blots with the control sera from healthy bovines and secondary antibodies were blank or showed rare, extremely faint bands; hence, the noise levels in the disease serum blots were considered to be negligible.

Overall, the dot and Western blots gave similar results. MSC_816 and its homologues raised a strong humoral response for all tested CBPP sera, while IgGs binding Vmm were rare. Antibody levels to the other Vmm-type proteins showed more variation.

**Generation of Vmm-type protein-specific antibodies**

pAbs to the five recombinant proteins were generated in rabbits. In addition, the mouse mAb 5G1, which targets Vmm (Persson et al., 2002), was used. The rabbit sera were affinity-purified using the recombinant proteins as ligands to obtain monospecific polyclonal antisera. The cross-reactivity of the polyclonal antisera was tested by dot blotting in duplicates. The six recombinant proteins, an M. mycoides SC strain PG1T lysate and four recombinant proteins of human origin were analysed with the five

**Fig. 2.** Dot blots made for evaluation of specificity of the serum antibodies to the recombinant Vmm-type proteins. The specific signal for a Vmm-type protein was blocked without affecting the other signals, demonstrated here with and without pre-adsorption with protein R816. The PG1 signal is stronger in comparison to Fig. 1, as a more concentrated PG1T lysate was used.

**Fig. 3.** Western blots of the recombinant proteins R117, R364, R390 (Vmm), R816, R847 and R1033 to screen for immunoreactivity of the corresponding native proteins in M. mycoides SC infections. The blots were analysed with bovine sera from CBPP-affected cattle (1MUK15A, 6MUK15D, Exp. Inf., G2, G4, C102 and PW227). His6–ABP protein (Tag) and a whole-cell lysate of M. mycoides SC strain PG1T were included as negative and positive controls, respectively. The last two blots were stained with Ponceau S in order to visualize the total protein content on the membrane and with a Tag-specific antibody that identifies all recombinant proteins.
affinity-purified pAbs and the 5G1 mAb (Fig. 4). All five antibodies against the recombinant proteins generated signals with their respective partner at 1:10 000 dilutions. A117, A816 and A847 generated weak cross-reactivity with each other. None of the specific pAbs stained the PG1 T lysate. The 5G1 staining showed strong specific staining of the recombinant Vmm (R390) and weak staining of the PG1 T lysate. Furthermore, there was no cross-reactivity with the His6–ABP tag or the recombinant proteins of human origin produced using the same vector and methodology, which were used as negative controls. It is noteworthy that antibodies A816 and A1033 will probably bind all the homologues to proteins MSC_0816 and MSC_1033 mentioned in the ‘Selection and in silico analysis of Vmm-type proteins’ section above.

Expression of Vmm-type proteins on in vitro cultured M. mycoides SC

Colonies of M. mycoides SC strains PG1 T and M223/90 were analysed with the five affinity-purified antibodies and mAb 5G1. Colonies that were positively stained entirely or in sectors were obtained for both strains with antibody A364 (Fig. 5), which confirms the bioinformatic prediction that protein MSC_0364 has variable expression. Approximately 20% of the colonies in M223/90 were stained, but only 5% in PG1 T. Antibody A847 gave an inconclusive result in that one colony was positive among thousands of negatives in strain M223/90 only; however, this may be a result of very rare expression of MSC_0847 under in vitro conditions. Expression of MSC_0117, MSC_0816 and MSC_1033 could not be detected with antibodies A117, A816 or A1033 by colony blotting. The control experiment with mAb 5G1 showed a high frequency of variation in the expression of Vmm in PG1 T, in accordance with previous results (Persson et al., 2002).

To investigate whether the negative results of colony blotting were caused by lack of expression, epitope masking or failure of the antibodies, Western blotting was performed on PG1 T lysates (Fig. 6). Antibody A364 generated a strong band of the correct size and two smaller bands that were considered to be noise, thus supporting the previous colony blotting results. Antibodies A117 and A847 gave a single band that would support the hypothesis that MSC_0117 and MSC_0847 were expressed, but the protein sizes were different from the theoretical sizes of the native proteins (see Table 2). Two distinct protein bands were obtained with A1033, indicating that both MSC_1033 and its homologue MSC_1058 were expressed. Even though the lower band corresponds to the theoretical size of MSC_1058 (15.7 kDa), it is tempting to assume that both proteins have retarded migration. No bands were obtained for antibody A816 despite the fact that this antibody should recognize five protein homologues. It is noteworthy that these Western blots contained a more concentrated PG1 T lysate than the dot blots.

To further investigate the expression data, RT-PCR was performed on RNA extracted from a PG1 T culture. Amplicons were obtained for all the Vmm-type proteins by RT-PCR but were not obtained by regular PCR on the same RNA template, showing that the transcripts were present in the cultured PG1 T. The presence of transcripts supported the findings for Vmm-type protein MSC_0364. For proteins MSC_0117 and MSC_0847, we conclude that the transcript had been detected, and it is likely that the
bands obtained in the Western blots were the correct proteins; however, they had an atypical migration in SDS-PAGE, which may be accounted for by imprecise bioinformatic size predictions. The transcript for protein MSC_1033 and/or MSC_1058 was amplified; however, we have no method to differentiate them. This is also true for protein MSC_0816 and its homologues, although no protein expression at all could be detected despite multiple immunoblottings being performed under different conditions with cultured PG1 T and M223/90. Interestingly, this protein often gave the strongest signals when the recombinant proteins were immunoblotted and screened with bovine sera.

**DISCUSSION**

In this investigation, five Vmm-type proteins of *M. mycoides* SC and Vmm itself were analysed. The bioinformatic predictions identified signal peptides in all the proteins (five of which were prolipoproteins), suggesting surface location, and also identified promoter structures that indicate variable expression. A scheme for high-throughput cloning, mutagenesis, expression in *E. coli* and purification of recombinant *M. mycoides* SC proteins was developed as a tool to perform studies that may support and supplement the bioinformatic predictions. This scheme will be beneficial for further research on *M. mycoides* SC and can be used in studies on other mycoplasma species. This scheme yielded sufficient recombinant Vmm-type protein from 100 ml of culture to perform Western and dot blotting experiments to assess the immunogenicity of the native Vmm-type proteins, to make antigens for antibody generation, and to make columns for affinity purification of the antisera.

When sera from natural CBPP infections were analysed for the presence of antibodies that target Vmm-type proteins, it was found that all sera contained antibodies to some or all of the Vmm-type proteins or their homologues. However, the sera generated different dot and Western blot patterns (Figs 1 and 3, Table 3), suggesting that the immune responses in the individual bovines were triggered by different proteins, either because this is the nature of the immune system or because the protein composition on the mycoplasma surface differed due to variable expression of each Vmm-type protein. For most sera, dot and Western blotting gave similar results, but we could observe differences in signal strength. This is expected due to the conformational differences of the proteins in the assays. Surprisingly, the previously studied Vmm (MSC_0390), which is thought to be immunodominant, generated the poorest IgG response and no response at all in most sera. Protein MSC_0816 and its homologues generated the strongest response overall. Interestingly, the differences in immunoreactivity between individual sera from one outbreak were as pronounced as the deviations seen between sera from different outbreaks in vastly separated geographical regions. This is clearly seen with the Botswana serum G2, which has a particular protein profile, whereas most Namibian and Botswana sera follow a more uniform pattern. The sera from Tanzania and Kenya gave distinct protein profiles; however, more sera must be analysed to find out if this is representative of the region or just specific for the individual bovines.

The dot and Western blotting results presented were repeated three times and twice, respectively, with high reproducibility. Standardized experimental conditions were used for each set of blots, with identical protein batches and volumes, and fixed serum dilutions. Signal intensities in Table 3 were annotated by judging all available replicates of the blots. It is not possible, however, to directly quantify the amounts of bound IgG antibodies by measuring the signal intensity when using an enhanced-chemiluminescence detection system, since subtle changes will influence the signal strength, such as the amount of protein in the blot, the time for photo acquisition and the enzymatic luminescence reaction. Clearly the light emitted from a strong positive signal will suppress the detection of a weak signal on the same blot. An example is the dot blot with serum PW227 in Fig. 1, where responses to the recombinant proteins are weak compared to the PG1 T lysate. A weak signal on the same blot. An example is the dot blot with serum PW227 in Fig. 1, where responses to the recombinant proteins are weak compared to the PG1 T lysate response, which is why the PG1 T signal appears more intense than for blots in which one of the recombinants gives the predominant signal. Similarly, a negative control often generated a weak signal when no positive signal was present. For this reason signals were categorized as inconclusive when only weak signals were obtained compared to the controls, or if the higher dilution of a serum lacked the corresponding signal of a lower dilution. Furthermore, we do not know the effects of storage, transport, cycles of freezing and thawing, age, lyophilization, etc. on the immunoglobulins. Naturally, the quality of the disease sera
will affect the general signal strength in an experiment, but should have less effect on the relative amounts of protein-specific antibodies that cause the protein profile within one blotting membrane. The only fresh samples used in this study were from the five healthy control cattle, and it is possible that these five samples had a higher general IgG titre than those of the stored CBPP-positive sera.

Having concluded that the Vmm-type proteins or their homologues are expressed in natural infections and generate a humoral immune response, variable protein expression in vitro was also examined. All results for MSC_0364 were congruent and showed that this protein is expressed in a culture medium environment, and the colony immunostaining confirmed that its expression is variable. Transcripts of all Vmm-type proteins were identified by RT-PCR in total RNA of strain PG1\textsuperscript{1}, although this method is very sensitive and does not indicate transcription levels. Colony blots were essentially negative and therefore inconclusive. Western blots of PG1\textsuperscript{1} lysates provided some support for expression of the Vmm-type proteins, with one distinct band for A117 and A847, and two distinct bands for A1033, as expected. There is still some uncertainty regarding these results, since the detected bands were of sizes that differed from the calculated theoretical protein sizes. It is well known, however, that proteins and especially lipoproteins can display unpredictable migration in SDS-PAGE (Banker & Cotman, 1972; Miyake et al., 1978; Simons & Helenius, 1970). At this point one can only speculate whether the failure to detect proteins MSC_0117, MSC_847 and MSC_1033 in colony blots is due to epitope masking or a protein conformation problem that makes the antibodies unsuitable for assays with native, folded proteins, even if they worked well in Western blot applications and dot blotting of recombinant proteins. Using these techniques we were unable to determine if these three proteins are variably expressed. The last protein, MSC_0816, and its homologues could not be detected using any of the immunoassays.

An interesting aspect of the analyses was the deviation between our experimental observations and the expected expression profile of the Vmm-type proteins, as judged from the promoter sequences. Looking at the genomic sequence of *M. mycoides* SC strain PG1\textsuperscript{1}, most of these Vmm-type proteins would be expected to be silent in this strain according to the length of the promoter spacer. Only one of the expressed Vmm-type proteins, MSC_117, had a promoter spacer length of 17 bp and should theoretically be expressed. Colony blotting indicated that MSC_364 and MSC_390 were expressed, while MSC_117 was not. One explanation would be that the genome sequence and the colony blots were made from different passages of PG1\textsuperscript{1}. Furthermore, transcripts were detected for all the Vmm-type proteins, but the sensitivity of PCR would detect a very small fraction of expressed genes that may be undetectable with the other methods. Generally, comparisons between experiments and batches of cultures are unreliable when working with variable proteins.

It may be argued that the antigen repertoire of in vitro-cultured strains of *M. mycoides* SC differs from those of strains in natural infections, since vaccination with non-viable in vitro cultivated strains often gives insufficient protection. Therefore, one hypothesis is that vaccine candidates should be searched for within the set of proteins that are expressed in the infected host but not in the laboratory. Theoretically, a vaccine of lysed whole cells spiked with a blend of all Vmm-type proteins should match all expression combinations in infected hosts.

Following the systematic analysis of surface proteins by recombinant technology we have presented results for six Vmm-type proteins that generated immune responses in vivo. We have also shown that the IgG response targets different Vmm-type proteins in individual sera. Further work using serial bleeds during a natural infection would increase our understanding of the natural variability in the expression of these proteins and possible infection-stage-dependent variability. It is appealing to consider these Vmm-type proteins as potential components of a recombinant protein vaccine, although their use in a vaccine needs to be further evaluated, since a humoral immune response does not necessarily mean that these proteins raise a cell-mediated immune response or a protective immune response. A recombinant vaccine to *M. mycoides* SC would have several advantages over live attenuated vaccines, which include durability in storage and transport, fewer undesirable side effects, since it would be more defined, and most importantly in this case, there are more options to modulate a recombinant vaccine. For example, one can readily add or remove fusion partners to enhance host responses or make di- and multimers of the recombinant proteins in the vaccine.

The recombinant proteins and their corresponding specific antibodies produced in this study have the potential to be powerful reagents for future protein investigations such as ELISA development, immunohistochemistry and studies of protein interactions.

**ACKNOWLEDGEMENTS**

We would like to thank Gerrit Viljoen, International Atomic Energy Agency, Vienna, Baptiste Dungu, Onderstepoort Biological Products, Onderstepoort, South Africa, Otto Hübsche, Ministry of Agriculture, Water and Fishery, Windhoek, Namibia, and John B. March, BigDNA Ltd, Roslin, Scotland, for providing the bovine sera used in this study. We would also like to thank our colleagues Stefan Jernstedt, Agneta Stillesjo and Virginia Meys at the National Veterinary Institute (SVA) in Uppsala, Sweden, for assisting with the colony blotting. The study was funded by the Swedish International Development Cooperation Agency (SIDA).

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Edited by: C. Citti