Reiterated repeat region variability in the ciliary adhesin gene of *Mycoplasma hyopneumoniae*

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*Mycoplasma hyopneumoniae* is a highly prevalent pathogen which colonizes the ciliated epithelial lining of the porcine respiratory tract. Expression libraries constructed from genomic DNA of the non-pathogenic strain *M. hyopneumoniae* J were screened with porcine hyperimmune antiserum against *M. hyopneumoniae*. One clone expressed a 28 kDa protein which was also reactive with monospecific antiserum raised against a putative *M. hyopneumoniae*-specific 94 kDa antigen derived from strain J. Trypsin digestion of whole *M. hyopneumoniae* cells showed the 94 kDa antigen to be surface-accessible. DNA sequence analysis of the gene encoding the 94 kDa antigen revealed greater than 90% homology to two adhesin genes, encoding P97 and Mhp1, cloned from pathogenic strain 232 and strain P5722 of *M. hyopneumoniae*, respectively. Two regions of repetitive DNA sequence were identified in the gene encoding the 94 kDa antigen. The first encoded the deduced amino acid sequence A(T)-K-P-E(V)-A(T) arranged as nine tandem repeats (RR1). The second region of repetitive DNA sequence encoded the deduced amino acid sequence G-A(E,S)-P-N(S)-Q-G-K-K-A-E arranged as five tandem repeats (RR2). Comparison of the three *M. hyopneumoniae* adhesin genes revealed that the genes encoding P97 and Mhp1, and the strain J gene encoding the 94 kDa antigen contained 15, 12 and 9 tandem repeats, respectively, in RR1, and 4, 5 and 5 tandem repeats, respectively, in RR2. Southern hybridization analysis of EcoRI-digested genomic DNA probed with an 820 bp fragment spanning RR1 and RR2 identified a strongly hybridizing fragment ranging in size from 2.15 to 2.30 kb among seven geographically diverse strains of *M. hyopneumoniae* but failed to hybridize with DNA from four strains of *Mycoplasma hyorhinis* or *Mycoplasma flocculare* strain Ms42. PCR primers flanking the DNA sequence encoding RR1 and RR2 were used to amplify DNA from the seven strains of *M. hyopneumoniae* and DNA sequence analysis of the amplification products showed that the number of tandem amino acid repeats in RR1 varied considerably between strains. RR1 from *M. hyopneumoniae* strains YZ, Beaufort, Sue, OMZ407 and C1735/2 comprised 11, 15, 12, 15 and 8 tandem copies, respectively, of the 5-aa repeat whilst RR2 comprised 4, 3, 4, 3 and 4 tandem copies, respectively, of the 10-aa repeat. Two putative integrin binding sites (GE-T and R-X-X-X-D) were identified in the 94 kDa ciliary adhesin. Variability in the number of amino acid repeats in RR1 amongst strains of *M. hyopneumoniae* may influence ciliary binding.

**Keywords**: *Mycoplasma hyopneumoniae*, pathogenesis, adhesin, antigenic size variation

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**Abbreviations**: i.m., intramuscular; PRR, proline-rich repeat.

The GenBank accession number for the sequence reported in this paper is AF001398.
INTRODUCTION

*Mycoplasma hyopneumoniae* attaches to the ciliated epithelial cells lining the porcine respiratory tract, inducing loss of ciliary function and a chronic inflammatory response which results in pneumonia and causes significant losses to swine production (Tajima & Yagishashi, 1982; Clark et al., 1991). Mycoplasma membranes are rich in protein, many of which stimulate a humoral immune response in the host (Razin, 1979). Although several immunoreactive *M. hyopneumoniae* antigens have been characterized and their genes cloned (Klinkert et al., 1985; Haldimann et al., 1993, Fagan et al., 1996), attempts to identify surface components which play a role in early colonization have only recently been reported. The development of microtitre plate adherence and TLC overlay assays has revealed that *M. hyopneumoniae* binds to sulfated glycolipid receptors present on porcine respiratory tract ciliated cells and that heparin, fucoidan, mucin and chondroitin interact with adhesive molecules on the surface of *M. hyopneumoniae* and are able to disrupt the adherence of the pathogen to intact ciliated cells (Zhang et al., 1994a, b). In addition, two mAbs (F2G5 and F1B6) which predominantly recognize a 97 kDa *M. hyopneumoniae* antigen inhibited adherence by up to 67% in a microtitre plate adherence assay (MPAA) (Zhang et al., 1995). mAb F2G5 recognized several antigens ranging in molecular mass from 65 to 125 kDa in immunoblotting studies of whole-cell lysates of different strains of *M. hyopneumoniae* and the immunoblot profiles varied among these strains (Zhang et al., 1995). In addition, low-passage strains of *M. hyopneumoniae* adhered strongly to porcine cilia in an MPAA whilst the high-passage strain J displayed significantly reduced adherence (Zhang et al., 1995).

Hsu et al. (1997) cloned, sequenced and expressed a gene encoding P97 from *M. hyopneumoniae* strain 232. The P97 coding region contained two regions of repetitive DNA sequence in the carboxy terminus. One of the repetitive regions encoded amino acids A(T)-K-P-E(V)-A(T) repeated 15 times (RR1) and the second region encoded amino acids G-A(E,S)-P-N(S)-Q-G-K-K-A repeated four times (RR2). The role of P97 in adherence to cilia was confirmed using transposon-induced mutations within the P97 coding sequence to eliminate reactivity to mAb F1B6. Tn1000 insertional mutagenesis studies mapped the mAb F1B6 binding epitope to a region between amino acid residues 704 and 1004 in the carboxy terminus of P97 which encompassed the 5-aa repeat region (RR1) but not the 10-aa repeat region (RR2). The insertion of Tn1000 in RR2 did not affect ciliary binding or the ability of mAb F1B6 to bind to P97, suggesting that RR2 is not involved in adhesion (Hsu et al., 1997). DNA sequence analysis revealed the presence of an ORF for a 124 kDa protein; however amino-terminal sequence analysis of P97 (Zhang et al., 1995) mapped 195 aa internal to the start codon, suggesting that the 124 kDa antigen was a precursor in *M. hyopneumoniae* that is subsequently processed to 102 kDa by proteolytic cleavage (Hsu et al., 1997). Preliminary DNA sequence analysis of the gene encoding a 28 kDa immunoreactive antigen from the avirulent strain *M. hyopneumoniae* J showed homology to the carboxy-terminal portion of a ciliary adhesin characterized from pathogenic strain 232 (accession no. U50901). The aim of this project was to clone, sequence and characterize the gene encoding the ciliary adhesin from the avirulent strain *M. hyopneumoniae* J and to investigate variations in gene sequence, particularly in the reiterated repeat regions, among geographically diverse strains of this pathogen. In addition, we investigated the surface accessibility of the adhesin and determined if pigs infected with *M. hyopneumoniae* elicited a serum antibody response which recognized this antigen.

METHODS

**Bacterial strains, plasmids and culture conditions.** Plasmid vectors pET23a, b and c (Novagen) in conjunction with *Escherichia coli* BL21(DE3)(pLysS) cells were used to produce the *M. hyopneumoniae* strain J library. *E. coli* strains XL-1 Blue (Stratagene) and BL21(DE3)(pLysS) (Novagen) were used to amplify and express pAS1, respectively. *E. coli* strains were grown in Luria–Bertani (LB) medium or on LB agar (Sambrook et al., 1989) supplemented with ampicillin (100 µg ml⁻¹) and chloramphenicol (32 µg ml⁻¹) where required.

**Mycoplasma strains and culture conditions.** *M. hyopneumoniae* strains J (NCTC 10110), Beaufort, Sue, C1735/2 and OMZ407, *M. hyorhinis* strains BTS-7, GDL, field strains 1 and 2 and *M. flocculare* type strain Ms42 were obtained from the Australian Mycoplasma Reference Collection (AMRC, University of South Australia). Australian isolates Beaufort, Sue, C1735/2 and OMZ407 were originally recovered from porcine lungs with lesions typical of enzootic pneumonia and are low-passage isolates (Scarnan et al., 1997). *M. hyopneumoniae* strains YZ (France) and 232 (USA) were obtained from M. Kobisch (Ploufragan, France) and T. Young (Iowa State University, USA), respectively. All strains were cultured as described previously (Scarnan et al., 1997). Briefly, each was grown in modified Friis medium at 37 °C and harvested in the pH range 6-9-72 as indicated by a colour change. Mycoplasmas were harvested by centrifugation at 7000 g for 20 min and the pellet washed three times with TBS (10 mM Tris/HCl, 0.9 % NaCl, pH 7-4). The identity of all strains of *M. hyopneumoniae* and *M. flocculare* strain Ms42 was confirmed using species-specific PCR primers which amplified portions of the 16S rRNA gene of these two species (Stemke et al., 1994).

**Preparation of antisera.** Preparation of porcine hyperimmune antiserum against *M. hyopneumoniae* has been described previously (Scarnan et al., 1997). Two pigs from a herd known to be free of *M. hyopneumoniae* were bled (prebleed) and the serum used to confirm the absence of antibodies against *M. hyopneumoniae* by immunoblotting and ELISA. A freshly grown culture of *M. hyopneumoniae* was harvested, washed and the final pellet (10 g) was resuspended in TBS. An aliquot (1 ml) containing approximately 250 mg of *M. hyopneumoniae* was emulsified with an equal volume of Freund's complete adjuvant and used to inoculate each pig intramuscularly (i.m.) in the neck and rump (1 ml in each site). One month later, 2 ml of a booster inoculum was prepared
using Freund's incomplete adjuvant and administered as before. Serum responses were monitored weekly thereafter until an anti-M. hyopneumoniae response was observed by immunoblotting and ELISA (Scarmar et al., 1997).

Rabbit hyperimmune serum was raised against recombinant 28 kDa (carboxy-terminal portion of the adhesin), 94 kDa (full adhesin) and 36 kDa (lactate dehydrogenase) M. hyopneumoniae antigens. The 28 kDa antigen was prepared as described by Djordjevic et al. (1994) with the following modifications. Polycrylamide gels (12%) prepared with a single sample well were loaded with 491 Prep Cell purified (Bio-Rad) 28 kDa antigen (approx. 50 mg). Gels were stained with Coomassie blue and the band containing the antigen excised, frozen in liquid nitrogen and ground with a mortar and pestle to a fine powder. Antigen was emulsified with Freund's complete adjuvant and then administered i.m. to two New Zealand White rabbits (delivering 0.75 ml to each of the hind legs and 0.25 ml to each front leg). Booster inoculations, delivered 1 month later, consisted of the same dose of the pulverized polycrylamide/antigen emulsified with Freund's incomplete adjuvant. Rabbit hyperimmune sera against 94 and 36 kDa M. hyopneumoniae-specific antigens were produced as described above except that each antigen did not require extraction from a gel. Column-purified (491 Prep Cell) 94 and 36 kDa antigens appeared as single bands when resolved by SDS-PAGE and stained with Coomassie blue or immunoblotted using anti-94 and anti-36 kDa antisera, respectively (data not shown).

Experimental infection of pigs with M. hyopneumoniae. Six pigs were experimentally infected with M. hyopneumoniae as described previously (Djordjevic et al., 1997). Briefly, pigs were challenged by intratracheal inoculation with a lung homogenate obtained from pigs previously infected with the virulent Beaufort strain of M. hyopneumoniae (Etheridge et al., 1979). The presence of M. hyopneumoniae in lung tissue routinely used for the preparation of lung homogenate was confirmed using a PCR assay incorporating primers specific for the 16s rRNA gene of this species (Stemke et al., 1994). Pigs experimentally infected with M. hyopneumoniae as described above were then blotted onto nitrocellulose, placed on Whatman 3MM filter paper separated with LB medium and 10 mM IPTG and incubated at 37 °C for 4 h. The blots were then treated with 50% (w/v) SDS to lyse the cells, gently washed with PBS, blocked by incubation in 5% (w/v) non-fat milk in TBS and immunostained using porcine hyperimmune M. hyopneumoniae antiserum which had been extensively adsorbed against E. coli proteins as described by Ro et al. (1994). Potential positive clones were recovered from master plates and the immunoreactivity of recombinant antigens was confirmed by Western blotting.

Protein expression and purification. Recombinant E. coli BL21(DE3)(pLysS) cells containing pAS1 were grown to mid-exponential phase in LB medium containing the appropriate antibiotics prior to the addition of 10 mM IPTG. After further growth for 4 h, cells were harvested, lysed by heating to 95 °C in Laemmli reducing mixture (Laemmli, 1970) and sonicated using a Branson Sonifier (10 × 30 s, 70% duty cycle over a 5 min period at 4 °C) to shear genomic DNA. Whole-cell lysate (2 ml loading vol., 0.5 g wet wt) was loaded onto a preparative 491 Prep Cell column (37 mm i.d.) containing a 12% polyacrylamide matrix with 4% polyacrylamide stacking gel and proteins separated by electrophoresis for 24 h at 40 mA constant current. Aliquots (15 μl) of fractions (5 ml) were analysed by immunoblotting using porcine hyperimmune antiserum against M. hyopneumoniae and fractions containing the immunoreactive 28 kDa antigen were pooled and concentrated by ultrafiltration (Amicon).

SDS-PAGE and Western blotting. SDS-PAGE and immunoblotting were performed essentially as described by Laemmli (1970) and Burnette (1981), respectively. After blocking in TBS containing 5% non-fat milk for 1 h and washing (3 × 10 min) in TBS containing 0.1% non-fat milk and 0.05% Tween 20, blots were incubated for 1:5 h with either porcine hyperimmune antiserum against M. hyopneumoniae (1:500) or rabbit anti-28-, anti-94- or anti-36 kDa sera (1:400) in TBS containing 0.5% non-fat milk. Blots were washed as above before being incubated with either goat anti-pig antibodies (Southern Biotechnologies) or sheep anti-rabbit antibodies (Silenus Laboratories) conjugated with horseradish peroxidase (1:2000). Blots were then washed (3 × 10 min) prior to development with freshly prepared DAB substrate (0.05% 3,3'-diaminobenzidine, 0.03% hydrogen peroxide in 100 mM Tris/HCl, pH 7.4).

ELISA for anti-28 kDa protein antibodies. Preparative SDS-PAGE-purified 28 kDa antigen (100 μl, 80 ng) was coated onto 96-well microtitre plates (Flow Laboratories) in carbonate coating buffer (0.15% Na₂CO₃, 0.29% NaHCO₃, pH 9.5) and plates were incubated overnight at room temperature in a humidified chamber. Plates were washed five times with PBS containing 0.05% Tween 20 using a SLT 96PW plate washer (Titetek). Primary antibody (sera from pigs experimentally infected with M. hyopneumoniae as described above) was diluted 1:200 in PBS containing 0.05% non-fat milk and 0.05% Tween 20, 100 μl was added to each well and the plates were incubated in a humidified chamber for 1:5 h at room temperature. Plates were washed as before, prior to the addition of 100 μl goat anti-pig horseradish peroxidase conjugate diluted 1:500 in PBS containing 20% non-fat milk and 0.05% Tween 20. Plates were then blotted onto nitrocellulose paper, treated with 50% (w/v) SDS to lyse the cells, gently washed with PBS, blocked by incubation in 5% (w/v) non-fat milk in TBS and immunostained using porcine hyperimmune M. hyopneumoniae antiserum which had been extensively adsorbed against E. coli proteins as described by Ro et al. (1994). Potential positive clones were recovered from master plates and the immunoreactivity of recombinant antigens was confirmed by Western blotting.

Cloning of the 3' end of the M. hyopneumoniae strain J adhesin gene. M. hyopneumoniae strain J genomic DNA was extracted as described previously (Fagan et al., 1996) and digested with EcoRI. DNA was size-fractionated through 10-40% sucrose gradients and fragments from 0.5 to 10 kb were ligated separately into EcoRI-digested pET23a, b and c vectors (Novagen) treated with calf alkaline phosphatase to prevent self-ligation. Ligation products were transformed into competent E. coli BL21(DE3)(pLysS) cells (Novagen). The cells were then inoculated into fresh LB medium and incubated for 2 h at 37 °C prior to plating onto LB agar containing chloramphenicol for overnight incubation at 37 °C. Colonies were then blotted onto nitrocellulose, placed on Whatman 3MM filter paper separated with LB medium and 1 mM IPTG and incubated at 37 °C for 4 h. The blots were then treated with 50% (w/v) SDS to lyse the cells, gently washed with PBS, blocked by incubation in 5% (w/v) non-fat milk in TBS and immunostained using porcine hyperimmune M. hyopneumoniae antiserum which had been extensively adsorbed against E. coli proteins as described by Ro et al. (1994). Potential positive clones were recovered from master plates and the immunoreactivity of recombinant antigens was confirmed by Western blotting.

Trypsin treatment of M. hyopneumoniae cells. M. hyopneumoniae cells (1 g) were resuspended in 5 ml sterile PBS and aliquots (500 μl) were introduced into sterile micro-centrifuge tubes. Cell suspensions of M. hyopneumoniae and freshly prepared stock solutions of trypsin were pre-incubated to 37 °C. Trypsin was added to cell suspensions of M.
**SDS-PAGE and immunoblotting using rabbit anti-28 and anti-94 kDa sera.**

DNA sequencing of the 3' end of the M. hyopneumoniae adhesin gene. DNA sequencing was performed using purified plasmid (pAS1) DNA (Qiagen) with synthetic oligonucleotide primers supplied commercially (Life Technologies) and the Taq DyeDeoxy Terminator Cycle Kit (Applied Biosystems). PCR was performed using a Perkin Elmer GeneAmp 9600 and DNA sequence reactions were analysed with an Applied Biosystems model 377 automated DNA sequencer. Comparison of the nucleotide and derived protein sequences with databases was performed using the package from the University of Wisconsin Genetics Computer Group (GCC) version 7, accessed via the Australian National Genomic Information Service (ANGIS, University of Sydney).

**N-terminal amino acid microsequencing.** The first 15 aa residues of the 28 kDa antigen purified by preparative SDS-PAGE were sequenced commercially using Edman chemistry with an Applied Biosystems Procise-HT Sequencer located at the Biomolecular Resource Facility, John Curtin School of Medical Research, Australian National University.

**Cloning and sequence analysis of the 5' end of the M. hyopneumoniae strain J adhesin gene.** Two homologous M. hyopneumoniae adhesin gene sequences (accession nos U27294 and U50901) were used to design primer 5'F (AGT TAA ATA AAT TTT TCA CT) and primer 5'R (CIT TAA TCT GAT TGT AAG GA) was designed using DNA sequence derived from pAS1. Primers 5' R and 5'R were used to amplify the 5' end of the M. hyopneumoniae strain J adhesin. A 50 μL reaction mixture comprised 200 μM deoxyribonucleotide triphosphates, oligonucleotide primers 5'F and 3'R and 3' (200 mM each), 2 units of AmpliTaq Gold DNA polymerase (Perkin Elmer), 16 mM (NH₄)₂SO₄, 67 mM Tris/HCl (pH 8.8) and 4 mM MgCl₂. The reaction mixtures were heated to 95-98°C (10 min). Cell lysates were analysed by SDS-PAGE and immunoblotting using rabbit anti-28 and anti-36 kDa sera.

**RESULTS**

**Immunoscreening of DNA libraries and immunological characterization of a 28 kDa M. hyopneumoniae strain J antigen.**

**DNA and protein sequence analysis of pAS1**

Digestion of pAS1 with EcoRI revealed the presence of a single 22 kb M. hyopneumoniae DNA insert (data not shown). DNA of pAS1 was used as a template to...
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Cloning and DNA sequence analysis of the 5' end of the ciliary adhesin gene

To facilitate a comparison of the gene encoding the 94 kDa *M. hyopneumoniae* antigen with genes encoding adhesins P97 (from strain 232) and Mhp1 (from strain P5722) the 5' end of the gene encoding the 94 kDa protein was amplified by PCR using primers 5'F and 3'R (Fig. 2a), cloned and sequenced three times. The predicted size of the 94 kDa antigen based on the complete sequence was 123 kDa which was similar to the calculated size reported for P97 and Mhp1 (124.9 and 124.2 kDa, respectively). Alignment of the predicted amino acid sequence of the 94 kDa antigen with the amino acid sequences of P97 and Mhp1 revealed differences in the number of tandem repeats among different strains of *M. hyopneumoniae*. The strain J adhesin contained nine copies of the 5-aa repeat and five copies of the 10-aa repeat whilst Mhp1 (strain P5722) had 12 copies of the 5-aa repeat and five copies of the 10-aa repeat (King et al., 1996). P97 (strain 232) had 15 copies of the 5-aa repeat and only four copies of the 10-aa repeat (Hsu et al., 1997). The first 32 aa of the 94 kDa antigen have features consistent with a prokaryotic signal sequence, including the presence of a cluster of positively charged amino acids (K) followed by a stretch of hydrophobic residues ending with a signal peptidase II cleavage site (data not shown). The absence of a cysteine residue immediately after the cleavage site or anywhere within the deduced amino acid sequence discounted the possibility that the 94 kDa antigen was a lipoprotein (Wieslander et al., 1992). Failure to detect sequence variation among three independent clones containing 5' adhesin gene sequences and the absence of more than a single strongly hybridizing fragment in Southern blot experiments (see Fig. 5b) suggested that only a single copy of the adhesin gene resides in *M. hyopneumoniae* strain J. Amino-terminal amino acid sequence data from purified P97 (Hsu et al., 1997) and Mhp1 (King et al., 1997) revealed that the first 195 aa had been removed and suggested that the ciliary adhesin was produced from a 124.5 kDa precursor which was subsequently processed to 102 kDa by proteolytic cleavage. Western blots of whole-cell lysates of different strains of *M. hyopneumoniae* reacted with anti-94 kDa serum (Fig. 3) identified a major immunogen ranging from 94 to 97 kDa (allowing for differences in the

sequence both strands of the 2.2 kb *M. hyopneumoniae* EcoRI fragment. The fragment was shown to be 2194 nt in length. Translation analysis revealed the presence of two ORFs (Fig. 2a). ORF1 was downstream of the T7 promoter of pET23c and the translated product, comprising 556 aa, showed strong homology to two translation products deposited in databases (U27294 and U50901, 92 and 88%, respectively); both sequences have been identified as *M. hyopneumoniae* adhesins (Hsu et al., 1997; King et al., 1996). From the deduced amino acid sequence, the predicted molecular mass of ORF1 was 62 kDa. The presence of two UGA codons at amino acid positions 546 and 766 in ORF1 precluded the expression of this protein in *E. coli* (Yamao et al., 1985). Amino acid analysis of the amino terminus of the 28 kDa antigen expressed by pAS1 identified the sequence A-F-S-Y-K-L-E-Y-T-D-E-N-K-L-S immediately following a methionine residue at position 881 (Fig. 2b). A fortuitous, putative Shine–Dalgarno sequence was also identified 10 bases upstream from the start codon, indicating that the 28 kDa antigen was expressed from this region of the sequence. A second ORF, ORF2, was identified immediately downstream from ORF1. No significant DNA or deduced protein sequence homology was identified using the ORF2 sequence in database searches. The presence of two regions of tandem repeats (RR1 and RR2) was also identified by sequence analysis of ORF1 (Fig. 2a). The first repeat region (RR1) consisted of repeating 15 bp units, encoding the amino acid sequence A(T)-K-P-E(V)-A(T), arranged as nine tandem repeats. The second (RR2) consisted of 30 bp units, encoding the amino acid sequence G-A(E,S)-P-N(S)-Q-G-K-K-A-E, arranged as five tandem repeats.

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**Fig. 1.** Immunological characterization of the 28 kDa antigen. Positions of molecular mass markers are given on the left. (a) Western blot of *E. coli* and *M. hyopneumoniae* whole-cell lysates separated in a 12% polyacrylamide gel reacted with anti-28 kDa serum. Lanes: 1 and 2, whole-cell lysates of *E. coli* BL21(DE3)(pLy5S)(pAS1) grown in the absence (1) and presence (2) of 10 mM IPTG; 3, whole-cell lysate of *M. hyopneumoniae* strain J. (b) Western blot of a 12% polyacrylamide gel of 491 Prep Cell-purified 94 kDa antigen (lane 1) and 28 kDa antigen (lane 2) reacted with anti-28 and anti-94 kDa sera, respectively.
number of repeats in RR1 and RR2), suggesting that this processing event occurs in all strains of M. 
hyopneumoniae so far examined.

Surface location of the 94 kDa antigen

Zhang et al. (1995) demonstrated the presence of P97 on the surface of M. hyopneumoniae using immunoelectron microscopy. To demonstrate the surface location of the 94 kDa antigen in M. hyopneumoniae strain J, whole 
hyopneumoniae cells were resuspended in PBS and digested with trypsin at concentrations of 0, 0.1, 0.3, 0.5, 1, 2, 3 and 5 µg ml⁻¹. Immunoblots of proteins of trypsin-treated whole M. hyopneumoniae cells probed with anti-28 kDa serum showed a steady decline in the detectable levels of the 94 kDa antigen at trypsin concentrations ranging from 0.1 to 1.0 µg ml⁻¹ (Fig. 4). The 94 kDa antigen was undetectable in M. hyopneumoniae treated with more than 2.0 µg ml⁻¹ trypsin for 15 min, suggesting that the 94 kDa antigen and the 28 kDa carboxy-terminal portion of this antigen reside on the surface of intact mycoplasmas. Cell integrity was demonstrated by reprobing the Western blot with rabbit antiserum raised to a purified preparation of recombinant lactate dehydrogenase from M. hyopneumoniae (A. L. Scarman, G. J. Eamens & S. P. Djordjevic, unpublished results). Lactate dehydrogenase (36 kDa) has been shown to reside in the cytosol in M. hyopneumoniae (Strasser et al., 1991). An immunoreactive antigen of equal staining intensity and with a molecular mass of 36 kDa was observed in all preparations of trypsin-treated whole M. hyopneumoniae cells confirming that trypsin did not cause cell lysis (Fig. 4).

Immunogenicity of the 28 kDa antigen

M. hyopneumoniae-free pigs were experimentally infected with the virulent Beaufort strain of M. hyopneumoniae as described previously (Fagan et al., 1996). Serum samples from six pigs, three taken prior to inoculation, were used to monitor the humoral immune response against the purified 28 kDa antigen. Although the temporal immune response varied between animals, antibodies which recognized the carboxy-terminal 28 kDa portion of the 94 kDa adhesin could be detected soon after challenge, becoming statistically significant (P < 0.05) 7 weeks after challenge (data not shown).

Strain variation in the adhesin antigen

Rabbit anti-28 and anti-94 kDa sera were used to probe identical Western blots of whole-cell lysates of seven M. hyopneumoniae strains from different geographic locations as well as four strains of M. hyorhinis and a single type strain of M. flocculare. The anti-28 kDa serum primarily reacted with a single antigen which varied in molecular mass between 94 and 97 kDa in all seven M. hyopneumoniae strains (Fig. 3a). The presence of more than one immunoreactive band can be observed for strains J (30 kDa as seen previously) and YZ (approx. 78 kDa). Strain Cl733/2 had the least immunoreactive band of all seven M. hyopneumoniae strains. Very weak cross-reactivity was observed with the whole-cell lysate of M. flocculare type strain Ms42, identifying two immunoreactive bands of 97 and 85 kDa (Fig. 3a). No cross-reactivity was observed with the whole-cell lysates of the four M. hyorhinis strains used in this study.
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Fig. 3. Western blot analysis of 12 porcine mycoplasma strains. Whole-cell lysates were resolved in a 12% polyacrylamide gel for reaction with (a) anti-28 and (b) anti-94 kDa sera. Positions of molecular mass markers are given on the left. Lanes: 1, *E. coli* BL21(DE3)(pLysS)(pASl) grown in the presence of 10 mM IPTG in (a) and column-purified 94 kDa antigen in (b); 2-8, *M. hyopneumoniae* strains J, 232, Beaufort, Sue, YZ, C1735/2 and OMZ407, respectively; 9-12, *M. hyorhinis* strains field strain 1, GDL, BTS-7 and field strain 2; 13, *M. flocculare* type strain Ms42.

Fig. 4. Western blot showing cell lysates of *M. hyopneumoniae* whole cells treated with 0, 0.1, 0.3, 0.5, 1, 2, 3 and 5 μg trypsin ml⁻¹ (lanes 1–8, respectively) separated in a 12% polyacrylamide gel and reacted with anti-28 and anti-36 kDa sera. Anti-36 kDa serum (raised against lactate dehydrogenase) failed to show a decrease in staining intensity at all trypsin concentrations, confirming cell integrity. Positions of molecular mass markers are given on the left.

In addition to detecting a 94–97 kDa antigen in whole-cell lysates of all strains of *M. hyopneumoniae* (except strain C1735/2), rabbit anti-94 kDa serum identified several other *M. hyopneumoniae* antigens ranging in molecular mass from 65 to 80 kDa (Fig. 3b). These antigen profiles are very similar to those described by Zhang *et al.* (1995) using mAbs raised against the P97 adhesin. Faintly reacting high molecular mass antigens were also observed in whole-cell lysates of two of the four *M. hyorhinis* strains (type strains GDL and BTS-7) and *M. flocculare* type strain Ms42 (Fig. 3b).

**PCR and Southern blot analysis**

PCR primers RRF and RRR (see Fig. 2a) amplified an 820 bp fragment spanning RR1 and RR2 using template DNA from *M. hyopneumoniae* strain J. When used to amplify the same fragment from genomic DNA isolated from six other strains of *M. hyopneumoniae*, a single amplification product ranging in size from 750 to 900 bp was observed (Fig. 5a). PCR products were not observed using genomic DNA isolated from four different strains of *M. hyorhinis* or a single type strain of *M. flocculare* (Fig. 5a), suggesting that the 94 kDa adhesin gene was not present in the genomes of these two phylogenetically related species which inhabit the respiratory tract of pigs (Stemke *et al.*, 1992). This result was confirmed by the failure of the 820 bp PCR fragment to hybridize under stringent conditions to EcoRI-restricted genomic DNA of these *Mycoplasma* species (Fig. 5b). A single, strongly reactive EcoRI fragment ranging in size from 2.15 to 2.30 kb (Fig. 5b) hybridized with the radiolabelled 820 bp PCR fragment, suggesting that different strains of *M. hyopneumoniae* may produce an adhesin with variable numbers of oligopeptide repeats within RR1 and RR2. Several weakly hybridizing EcoRI fragments were observed in Southern blot profiles of *M. hyopneumoniae* strains 232, YZ, Sue and C1735/2 and a single extremely weak fragment in the strain J profile (data not shown). No weakly hybridizing fragments were observed in the EcoRI profiles of *M. hyopneumoniae* strains Beaufort and OMZ407. Although whole-cell lysates of strain C1735/2 displayed a poorly immunoreactive antigen when reacted with anti-28 and 94 kDa sera, a strong hybridization signal (to a 2.15 kb EcoRI fragment) was observed using the 820 bp probe (Fig. 5b). The stronger hybridization signals with EcoRI fragments from 232 and C1735/2 were not correlated with a higher concentration of loaded DNA. This was determined by comparing the intensity of ethidium bromide staining in the agarose gel containing all seven strains of *M. hyopneumoniae* DNA used for Southern blotting (data not shown). The possibility that more than one copy of the ciliary adhesin gene may reside in these two strains would require that DNA sequences containing the EcoRI restriction site in ORF2 also be duplicated.

**Comparison of deduced amino acid sequences spanning RR1 and RR2 from geographically diverse strains of *M. hyopneumoniae***

DNA sequence analysis of cloned PCR fragments comprising a region of DNA encoding RR1 and RR2...
from geographically diverse strains of *M. hyopneumoniae* were compared (Fig. 6). The deduced amino acid sequences showed considerable variation, particularly in the number of tandem 5-aa repeats in RR1. Australian strains Beaufort and OMZ407 possessed 15 tandem copies of the 5-aa repeat in RR1 and three tandem copies of the 10-aa repeat in RR2. Australian strain Sue possessed 12 tandem copies of the 5-aa repeat in RR1 and four tandem copies of the 10-aa repeat in RR2. French strain YZ possessed 11 tandem copies of the 5-aa repeat and four tandem copies of the 10-aa repeat in RR1 and RR2, respectively. Finally, Australian isolate C1735/2 possessed the fewest number of tandem copies (eight) of the 5-aa repeat in RR1 and had four tandem copies of the 10-aa repeat in RR2 (Fig. 6).

**DISCUSSION**

Evidence that an immunoreactive 94 kDa *M. hyopneumoniae* strain J antigen is a ciliary adhesin is based on the following observations: (i) the gene encoding the 94 kDa antigen shows strong homology (greater than 90%) to two *M. hyopneumoniae* ciliary adhesin gene sequences deposited in GenBank, (ii) the 94 kDa antigen is surface-accessible and (iii) anti-94 kDa serum reacts with several antigens ranging predominantly from 65 to 97 kDa, displaying antigenic size variation patterns similar to those described by Zhang et al. (1995) using mAbs against the P97 adhesin. We have demonstrated that the 94 kDa antigen possesses repetitive proline-rich sequences in the carboxy terminus of the protein. The 94 kDa antigen from strain J had fewer 5-aa repeats in RR1 (nine repeats) compared with P97 which is expressed in a lower-passage strain (232) of *M. hyopneumoniae* (15 repeats). Mhp1 had 12 repeats (RR1) but is derived from *M. hyopneumoniae* strain P5722 of unknown virulence. Strain 232 also had one less copy of the 10-aa repeat of RR2. Except for the repeat regions and three deletions (aa 890–893, 1021–1030 and 1084) in strain 232, the deduced amino acid sequences of all three strains were very similar (93.3 and 95.7% homology, respectively).

The presence of a ciliary adhesin gene in the highly passaged, avirulent strain J of *M. hyopneumoniae* with significant homology to adhesin gene sequences from a low-passage, virulent strain of this pathogen encouraged us to investigate if differences in the number of reiterated repeat regions also existed amongst a panel of *M. hyopneumoniae* strains from different geographic localities. Whilst the number of repeats in RR2 remained between 3 and 4 copies, the number of repeats in RR1 ranged from 8 to 15. The presence of a transposon insertion (Tn1000) in RR2 (residue 1004) in P97 failed to abrogate either ciliary binding or diminish the ability of mAb F1B6 to bind to P97 (Hsu et al., 1997), strongly suggesting that RR2 is not essential for these functions. Although the exact nature of the epitope recognized by mAb F1B6 has not been determined, the ability of a Tn1000 insertion 109 residues upstream of the beginning of RR1 (residue 704) to abrogate ciliary and mAb binding defines the biologically important region(s) of the molecule between these two transposon insertion sites spanning 300 residues (aa 704–1004) in the carboxy terminus of P97. Although RR1 only comprises approximately 25% of the amino acids spanning the two transposon insertion sites, proline-rich repeats (PRRs, which reside in RR1 and RR2) are often highly immunogenic (Williamson, 1994), suggesting that RR1 may comprise an epitope(s) recognized by mAb F1B6 crucial for ciliary binding. Although RR2 does not appear to be essential for ciliary binding, it has been retained in all strains of *M. hyopneumoniae* so far examined and does not undergo much variation in the number of repeats (3–5 copies). Results described in this study clearly demonstrate that porcine antibodies are raised to RR2 during experimental infection with *M. hyopneumoniae*.

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**Fig. 5.** PCR and Southern blot analysis of genomic DNA from 12 porcine *Mycoplasma* strains. (a) PCR products (visualized on a 1.5% agarose gel with ethidium bromide staining) of an amplification using primers RRF and RRR which flank the area encoding repeat regions 1 and 2. (b) Southern blot of *EcoR*I-digested genomic DNA (transferred from a 1% agarose gel) probed with an 820 bp fragment spanning RR1 and RR2 amplified from *M. hyopneumoniae* strain J DNA using primers RRF and RRR. Lanes: 1–7, *M. hyopneumoniae* strains J, 232, Beaufort, Sue, YZ, C1735/2 and OMZ407, respectively; 8–11, *M. hyorhinis* strains field strain 1, GDL, BTS-7 and field strain 2; 12, *M. flocculare* type strain Ms42.
hyopneumoniae, tempting speculation that RR2 might serve as an immune decoy directing the host humoral immune response to a non-essential portion of the ciliary adhesin. Deletion or transposon mutagenic analyses rather than site-directed mutagenic approaches may be required to precisely map biologically essential portion(s) of the molecule since interactions involving PRRs are dependent upon multiple weak binding sites rather than lock and key binding (Williamson, 1994). Further studies are required to investigate the role of RR1 and RR2 in ciliary binding.

Studies by King et al. (1996) suggested that Mhp1 may be duplicated in the genome of M. hyopneumoniae strain P5722. Similarly, Hsu et al. (1997) suggested that P97 was duplicated in M. hyopneumoniae strain 232A. Using an 820 bp hybridization probe spanning RR1 and RR2, a single strongly hybridizing fragment ranging from 2.15 to 2.3 kb was observed on Southern blots of EcoRI digests of the seven strains of M. hyopneumoniae DNA used in this study (Fig. 5b). Although several weakly hybridizing bands were observed with DNA from strains 232, YZ, Sue and C1735/2, suggesting that multiple copies of RR1 and/or RR2 may be present, none was observed in Australian isolates Beaufort and OMZ407 and only one extremely faint band was observed in the EcoRI digest of M. hyopneumoniae strain J. If the adhesin gene is duplicated in the genomes of M. hyopneumoniae with differences in the number of repeats in RR1 and RR2, our PCR cloning strategy should have detected them. These data provide circumstantial evidence that ciliary adhesin genes with a
variable number of repeat regions do not exist as multiple copies in *M. hyopneumoniae*. However, it is feasible that less homologous DNA fragments, including RR1 and/or RR2, may be duplicated elsewhere in the genome.

The pattern of multiple antigens (65–97 kDa) on Western blots of whole-cell lysates of *M. hyopneumoniae* reacting with anti-94 kDa serum has similarity to immunoblot profiles described using mAbs raised against the P97 adhesin (Zhang *et al*., 1995; Hsu *et al*., 1997). Differences in the number of repeats in RR1 and RR2 is insufficient to account for this complex antigen profile. Anti-28 kDa serum predominantly recognized a single antigen of 94–97 kDa in whole-cell lysates of the seven *M. hyopneumoniae* strains which could be explained by differences in the number of repeats in RR1 and RR2. King *et al.* (1996) speculated that multiple immunoreactive antigens may be cleavage products of the mature protein. Asp-Pro bonds underlie cleavage under mildly acid conditions (Kyte, 1995) and cleavage occurs during the processing of samples for SDS-PAGE giving rise to a ladder pattern in immunoblots (Wästfelt *et al*., 1996). Several Asp-Pro peptide bonds (aa 107–108, 204–205, 1068–1069) are present in the deduced amino acid sequence of the 94 kDa protein. However, the molecular masses of such cleavage products are inconsistent with the immunoblot profile generated using anti-94 kDa serum and the predicted cleavage products should also be reactive with anti-28 kDa serum. Size heterogeneity may result from premature translation or proteolytic cleavage and the former of these two possibilities would be consistent with the immunoblot profiles observed with the anti-28 kDa serum. Interestingly, whole-cell lysates of *M. hyopneumoniae* strain C1735/2 were poorly immunoreactive with anti-94 and anti-28 kDa sera, although faintly reactive bands were observed. Whole-cell lysates of strain C1735/2 are also poorly immunoreactive with porcine hyperimmune antiserum raised against J and Beaufort strains of *M. hyopneumoniae* (Scarmam *et al*., 1997). Cryptic mycoplasmal epitopes which fail to react with mAbs have been described in several *Mycoplasma* species and can be explained by the ability of surface antigens to undergo phase variation (Rosengarten & Yoge, 1996). However, since hyperimmune sera typically recognize a plethora of epitopes it is unusual for whole-cell lysates to react poorly with homologous hyperimmune sera, although others have reported such a phenomenon (Avakian *et al*., 1991).

Like the extracellular porcine pathogen *M. hyopneumoniae*, *Mycoplasma pneumoniae* is an extracellular pathogen which specifically adheres to the ciliated epithelial lining of the human respiratory tract. It requires adhesion to host-cell receptors for colonization and subsequent development of disease. A surface protein of *M. pneumoniae* containing PRRs, displaying size polymorphism and possessing an R-G-D sequence has been identified and designated P65 (Proft *et al*., 1995). P65 has not yet been implicated in binding host cells but its exclusive localization in the Triton X-100-insoluble fraction makes it a likely candidate. The amino-terminal portion of P65 comprises two PRR regions that form a proline-rich acidic domain. The repeats (D-P-N-A-Y and D-P-N-Q-A-Y) are responsible for the observed size polymorphism in immunoblots of whole-cell lysates of two different strains of *M. pneumoniae*, one a clinical isolate and the other a high-passage laboratory strain. The high-passage strain FH of *M. pneumoniae* contained 12 D-P-N-A-Y repeats whilst the clinical isolate contained only nine copies.

Virulence in mycoplasmas has been shown to be attenuated by successive passage in laboratory media (Collier *et al*., 1985). Zhang *et al.* (1994a) demonstrated that *M. hyopneumoniae* strain 232 L127 showed decreased adherence to porcine cilia with increasing passage in Friis medium, noting a significant reduction (*P* < 0.01) in adherence after 50 passages. *M. hyopneumoniae* strains maintained in Friis medium with minimal passage bind relatively strongly to porcine cilia compared to the high-passage strain J (DeBey & Ross, 1994; Zhang *et al*., 1995). Furthermore, low-passage strains of *M. hyopneumoniae* are pathogenic for pigs whilst the high-passage strain J is unable to induce pneumonia or to be re-isolated from the lungs of animals after challenge (Zielinski & Ross, 1990; Zielinski *et al*., 1993). All strains of *M. hyopneumoniae* so far characterized (except C1735/2) revealed a greater number of repeats in RR1 compared to the high-passage, avirulent strain J. Although the ability of the four Australian strains (Beaufort, Sue, OM2407 and C1735/2) described in this study to bind to porcine cilia has not been determined, it is likely that they demonstrate binding activity since all were recovered from lung lesions and have undergone minimal passage in vitro.

Repetitive regions within DNA have been described in a wide range of organisms and are known to be active sites for homologous recombination or slipped-strand mispairing (Albertini *et al*., 1982; Petes & Hill, 1988; Wren, 1991; Dybvig & Voelker, 1996). Intragenic recombination amongst tandem repeats in a single gene in *Xanthomonas campestris pv. malvacearum* provide a mechanism for the evolution of new host specificities (Yang & Gabriel, 1993). Surface proteins displaying reiterated repeats are increasingly being reported in mycoplasmas (Proft *et al*., 1995; Zheng *et al*., 1995; Zhang & Wise 1996) and phylogenetically related Gram-positive bacteria (Wren, 1991; Wästfelt *et al*., 1996). The MB antigen from *Ureaplasma urealyticum* contains 42-5 repeats of a 6 aa motif G-K-E-Q-P-A which reside in the carboxy-terminal portion of the molecule, is antigenic and surface-exposed and is responsible for antigenic size variation. Furthermore, it has been postulated that different antigen sizes may be associated with a specific pathological manifestation (Zheng *et al*., 1995).

Proline is present in both repeats found in RR1 and RR2 in the *M. hyopneumoniae* adhesin. PRRs within proteins are often found as multiple tandem repeats of variable length in both prokaryotes and eukaryotes and are usually highly immunogenic (Williamson, 1994). Many
proline-rich sequences have been demonstrated to be involved in binding processes, a large number of which are located on the cell surface. In particular, bacterial outer-membrane proteins associated with transport functions, binding cytoskeletal proteins, binding peptidoglycan and intracellular signalling have been demonstrated to contain PRRs critical in performing such tasks (Williamson, 1994, and references therein). The involvement of PRRs in binding occurs in a non-stoichiometric, but functionally important way and has a unique ability to bind rapidly and tightly to receptor molecules (Williamson, 1994). The bulkiness of the proline side chain and the replacement of the amide proton by a methylene group disrupts α-helix and β-sheet formation. The relatively large proportion of charged amino acids in the ciliary adhesion and the propensity for proline-rich peptides to be highly soluble in water provides an explanation for the partitioning of the adhesin to the aqueous phase during extraction with Triton X-114 (Williamson, 1994; Zhang et al., 1995). An R-G-D sequence, implicated in integrin binding in a broad range of viruses, bacteria and eukaryote parasites, is often found in conjunction with PRRs (Lenninger et al., 1991). Although a search of the 94 kDa, P97 and Mhp1 adhesins failed to identify an R-G-D sequence, two integrin-binding L-E-T sequence motifs which recognize αβ integrins (Ruoslathi, 1996) were located at positions 618–620 and 630–632 in the deduced amino acid sequences of the 94 kDa, P97 and Mhp1 adhesins. In addition, two R-X-X-D sequence motifs, which recognize αβ integrins (Ruoslathi, 1996), were located at positions 221–225 and 791–795 in the deduced amino acid sequences of the 94 kDa, P97 and Mhp1 adhesins. Site-directed mutagenesis studies would be useful to determine if these sequence motifs are involved in ciliary adhesion.

Proline-rich sequences are often accompanied by a relatively high prevalence of glutamine. Glutamine is present in the reiterated repeats comprising RR2 in the reiterated repeats comprising RR2 in the deduced amino acid sequences of the 94 kDa, P97 and Mhp1 adhesins. In addition, 4 of the 5 aa residues immediately 5' to RR1 (P-Q-P-P-A) are either proline or glutamine moieties. Glutamine is the second most likely residue (proline being the first) to appear in a polyproline helix segment in globular proteins (Adzhubei & Sternberg, 1993). A sequence of four or more consecutive prolines adopts a conformation in solution known as a polyproline II helix making an extended conformationally restricted polypeptide chain (Adzhubei & Sternberg, 1993). However, polyproline II helices typically comprise repetitive short proline-rich sequences of 2–3 aa such as (X-P-Y)n. RR1 and RR2 comprise tandem repeats each containing 5 and 10 aa, respectively, and these sequences are typically involved in protein–protein binding. The restricted mobility afforded by such structures enhances binding ability by reducing the unfavourable entropy loss of peptides on binding (Williamson, 1994).

Earlier studies by Young et al. (1990, 1992) reported that P97 was expressed during M. hyopneumoniae infection and that an IgG and IgA immune response was elicited against it in the respiratory tract of contact-exposed pigs 35–60 d earlier than other M. hyopneumoniae antigens. Pigs experimentally infected with a virulent strain of M. hyopneumoniae mounted a humoral immune response against the 28 kDa recombinant antigen immediately after challenge and antibody concentrations continued to rise for at least 7 weeks. Whilst immunodominant epitopes appear to be present in the carboxy-terminal portion of the 94 kDa antigen, which encompasses RR1 and RR2, further studies are required to identify the crucial part(s) of the molecule involved in attachment and to determine which are capable of eliciting a protective immune response.

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REFERENCES


Variability in the *M. hyopneumoniae* ciliary adhesin gene


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