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Characterization of a variant vlhA gene of *Mycoplasma synoviae*, strain WVU 1853, with a highly divergent haemagglutinin region

Awatef Béjaoui Khiari, Ibtissem Guériri, Radhia Ben Mohammed, Boutheina Ben Abdelmoumen Mardassi*

Abstract

**Background:** In *Mycoplasma synoviae*, type strain WVU 1853, a single member of the haemagglutinin vlhA gene family has been previously shown to be expressed. Variants of vlhA are expressed from the same unique vlhA promoter by recruiting pseudogene sequences via site-specific recombination events, thus generating antigenic variability. Using a bacterial stock of *M. synoviae* WVU 1853 that had been colony purified thrice and maintained in our laboratory at low passage level, we previously identified a vlhA gene-related partial coding sequence, referred to as MS2/28.1. The *E. coli*-expressed product of this partial coding sequence was found to be immunodominant, suggesting that it might be expressed.

**Results:** Reverse transcription-PCR amplification (RT-PCR), using a sense primer located at the 5′-end region of the expected vlhA transcript and a reverse primer located at the 3′ end of MS2/28.1 coding sequence, yielded a consistent amplification product showing that MS2/28.1 was indeed transcribed. Nucleotide sequence analysis of the RT-PCR product identified an 1815-nucleotide full-length open reading frame (ORF), immediately preceded by a nucleotide sequence identical to that previously reported for expressed *vlhA* genes. PCR amplifications using genomic DNA isolated from single colonies further confirmed that the full-length ORF of MS2/28.1 was located downstream of the unique *vlhA* promoter sequence. The deduced 604-amino acid (aa) sequence showed a perfect sequence identity to the previously reported *vlhA* expressed genes along the first 224 residues, then highly diverged with only 37.6% aa identity. Despite the fact that this *M. synoviae* clone expressed a highly divergent and considerably shorter C-terminal haemagglutinin product, it was found to be expressed at the surface of the bacterium and was able to haemagglutinate chicken erythrocytes. Importantly, the *E. coli*-expressed C-terminal highly divergent 60 residues of MS2/28.1 proved haemagglutination competent.

**Conclusions:** In contrast to the previously characterized *vlhA* expressed variants, MS2/28.1 displayed a highly divergent sequence, while still able to haemagglutinate erythrocytes. Overall, the data provide an indication as to which extent the *M. synoviae* *vlhA* gene could vary its antigenic repertoire.

**Background**

*Mycoplasma synoviae* is an economically important pathogen of poultry, causing synovitis, chronic respiratory tract disease, and retarded growth in chickens and turkeys [1,2]. *M. synoviae* is a member of the genus *Mycoplasma* of the class *Mollicutes*, a group of wall-less Gram-positive bacteria with genomes ranging from 1358 kb to as little as 580 kb [3]. The genome sequence of *M. synoviae* strain WVU 1853 has been determined and comparative analysis with *M. gallisepticum*, another major avian pathogen, provided evidence for horizontal gene transfer between the two species, though belonging to two distinct phylogenetic groups [4,5]. Among the genes that could have arisen by horizontal gene transfer are those encoding for haemagglutinins. In avian mycoplasmas, genes encoding for these immunogenic and surface exposed proteins are the subject of considerable antigenic variability [6]. By alternating the composition of their surface proteins, mycoplasmas are thought to colonize more efficiently mucosal surfaces and become...
more virulent [7,8]. Haemagglutinins account among the most important surface proteins involved in colonization and virulence of avian mycoplasmas [6,9].

In M. synoviae, haemagglutinins are encoded by related sequences of a multigene family referred to as vlhA genes [10-12]. The haemagglutinins of M. gallisepticum (pMGA) and M. imitans are also encoded by multigene families related to vlhA [13,14]. Both organization and control of expression of vlhA genes are quite different between M. gallisepticum and M. synoviae. In the former species, vlhA genes are located in five distinct genomic regions and each gene appears to be translationally competent [14,15]. By contrast, in M. synoviae, all vlhA sequences are confined to a restricted genomic region with a unique copy being expressed in a single strain [16,17].

The uniquely expressed vlhA gene of M. synoviae yields a product that is cleaved post-translationally into a N-terminal lipoprotein (MSPB) and a C-terminal haemagglutinin protein (MSPA) [11]. Cleavage was found to occur immediately after amino acid residue 344 [17]. Both MSPA and MSPB are surface-exposed proteins and exhibit high frequency antigenic variation, but only MSPA mediates binding to erythrocytes [10,11]. Sequence analysis of several M. synoviae strains suggested that MSPA was more antigenically variable than MSPB [6,10,11]. Consistently, in isogenically derived M. synoviae clones that have lost their haemadsorbing and/or haemagglutinating activity, MSPA was no more detectable by polyclonal antisera or monoclonal antibodies, suggesting extensive antigenic variation [12].

The molecular basis underlying the generation of antigenic variants of M. synoviae vlhA genes has been elegantly demonstrated in a study conducted by Noormohammadi et al. 2000 [17]. It resides in the ability of a single strain to undergo high frequency site-specific recombination, owing to the availability in the genome sequence of a significant pool of pseudogenes (vlhA-related partial sequences). Recombination between the single complete vlhA gene and one of the multiple pseudogene copies ensures the creation of a new vlhA gene variant. To date, three expressed vlhA gene variants (vlhA1, vlhA4, and vlhA5) have been characterized in M. synoviae strain WVU 1853 [17]. These genes are equally sized and show extensive sequence variability in a 400-bp DNA segment in the middle of the vlhA sequence, suggesting that the recombination event, though introduced sequence variations, tended to preserve the overall sequence length and composition. Although it has been concluded that the potential of vlhA genes to vary is considerable, there is no indication as to which extent a vlhA gene could diverge without losing its properties.

Previous studies from our laboratory have identified in M. synoviae strain WVU 1853, an immunodominant vlhA variant (termed MS2/28.1) [18] whose haemagglutinin region displayed a dramatic sequence shift and was considerably reduced in size, relative to the previously characterized expressed vlhA genes (vlhA1, vlhA4, and vlhA5) [17]. To better evaluate the extent of antigenic variation that could be tolerated by the M. synoviae haemagglutinin, we sought to know whether this highly divergent vlhA member was properly processed and whether it remained functionally competent. Our results provide evidence that the antigenic repertoire of M. synoviae vlhA genes might be wider than previously perceived.

Results
Isolation of the MS2/28.1 fragment
The complete nucleotide sequence of MS2/28, the λ phage-derived DNA fragment (GenBank accession number MSU66315) harbouring the immuno-reactive MS2/28.1 sequence, has been previously described [18]. It is 2657 bp long and contained two partial ORFs, referred herein to as, MS2/28.1 (5’ end) and MS2/28.2 (3’ end) (GenBank accession numbers ORF G2149016 and ORF G2149017, respectively). MS2/28.1 lacked its N-terminal sequence, whereas the C-terminal region of MS2/28.2 was incomplete. The two partial ORFs shared 71% and 61.27% identity at the nucleotide and deduced amino acids sequences, respectively, and were separated by a 481-nucleotide non-coding region (data not shown). Amino acid sequence comparisons between the two partial ORFs and the expressed vlhA1 gene have been previously described [11]. MS2/28.1 displayed 54.1% identity with vlhA1, along a 244-residue overlapping region, while MS2/28.2 showed 58.4% identity through a 495-amino acid overlapping sequence, starting at residue 260 of the vlhA1 gene sequence. Thus, the two partial open reading frames MS2/28.1 and MS2/28.2 are members of the vlhA gene family.

Evidence that MS2/28.1 was transcribed through the unique vlhA promoter
Immunoreactivity of the λ phage MS2/28 clone was associated with the 5’ end of the MS2/28.1 partial ORF. With regard to the vlhA1 sequence, the MS2/28.1 expressed region corresponded to the MSPA (haemagglutinin) sequence extending from residues 346 to 446, located immediately after the cleavage site.

Given the strong immuno-reactivity of the MS2/28.1 encoded product, we hypothesised that it might be expressed in the bacterium as a vlhA variant. Indeed, it has been well established that in M. synoviae, strain WVU 1853, there exist only a single copy of the vlhA promoter. New variant sequences, recruited from a pool of vlhA pseudogenes, are placed under the control of
this unique promoter via site-specific recombination [17]. Hence, we performed RT-PCR using a sense primer targeting the 5’-end of the expected vlhA promoter-derived transcript coupled to a reverse primer located at the 3’ end of MS2/28.1 partial coding sequence. As shown in Figure 1, RT-PCR reaction yielded a DNA fragment around 1.934 kb that could not be amplified when PCR was attempted without RT reaction. This result provides evidence that MS2/28.1 was transcribed as a vlhA variant.

Analysis of MS2/28.1 cDNA sequence

To further confirm the authenticity of the RT-PCR product and to complete the full-length coding sequence of the MS2/28.1, we subjected the RT-PCR product to nucleotide sequence analysis. As expected, the 5’-end region preceding the ATG initiation codon was identical to that reported for the previously reported vlhA expressed genes [17]. The MS2/28.1 full-length ORF consisted of 1815 nucleotides (GenBank accession number FJ890931). The deduced 604-amino acid sequence is predicted to encode a protein with an expected molecular mass of 64.3 kDa. Sequence alignments with vlhA1 (GenBank accession no. AF035624), as well as with the two full-length pseudogenes vlhA2 and vlhA3 (GenBank accession nos. AF085697 and AF085698, respectively) [17] (Figure 2), showed a perfect identity along the 670 5’-end first nucleotides, corresponding to the N-terminal first 223 amino acids. Downstream from this region, a very high divergence was observed with 37,6%, 37,8%, and 38,7% aa identity, respectively. Likewise, in this region, MS2/28.1 shared only 39,8% and 38,8% identity, respectively, with the two vlhA1 expressed variants, vlhA4 and vlhA5, previously identified in M. synoviae strain WVU 1853 (Figure 2). Overall, the haemagglutinin region of MS2/28.1 was found to be considerably reduced in size (148 aa less than in vlhA1) and displayed high level of sequence divergence in comparison to the previously reported vlhA expressed genes, namely vlhA1, vlhA4 (GenBank accession no. AF181033), and vlhA5 (GenBank accession no. AF181034) [17].

Demonstration that MS2/28.1 sequence is preceded by the vlhA1 promoter

To confirm that in our bacterial stock MS2/28.1 was located downstream of the unique vlhA promoter sequence, we performed PCR amplifications on single colonies using oligonucleotide primers placed in the vlhA promoter sequence with either vlhA1- or MS2/28.1-specific reverse primers. As shown in Figure 3, amplicons migrating at the expected mobility were obtained solely with MS2/28.1-specific reverse primers. Sequence analysis further confirmed that the upstream sequence is identical to that of the vlhA1 promoter, a result consistent with the finding that MS2/28.1 is transcriptionally active and that, in its transcript, the region preceding its ATG initiation codon was identical to that reported for vlhA1.

MS2/28.1 encoded full-length product is post-translationally cleaved with its C-terminal portion exposed at the bacterium’s surface

To characterize MS2/28.1 encoded product and to examine whether it was processed similarly as the vlhA1 product, we generated antisera towards four bacterially expressed distinct regions of the coding sequence. The reactivity of these antisera is shown in Figure 4. The antiserum raised against the N-terminal 323 residues (region A), which represent the vlhA1 MSPB protein, recognized three main protein bands (Figure 4, lane 1): a diffuse 25 to 30 kDa band similar to the previously described MSPC3 [10], a second diffuse band migrating at the range of 45 to 49 kDa, typical to the previously described MSPB2 and MSPB3 proteins, and a third less intense, high molecular mass protein of 97 kDa, consistent with the uncleaved vlhA1 gene product. All these protein bands were revealed by the rabbit polyclonal anti-M. synoviae serum (Figure 4, lane 5). The monospecific antiserum raised against the 19-amino acid peptide (region B) located immediately upstream of the putative cleavage site reacted essentially with a non diffuse single band of 45 kDa (Figure 4, lane 2), identical to the vlhA1
MSPB protein. Thus, MS2/28.1 product was properly cleaved. This was expected because, although MS2/28.1 diverged significantly from *vlhA1*, the sequence environment of the putative cleavage site was conserved along a 17-amino acid stretch (residues 339 to 355, relative to the *vlhA1* sequence). The monospecific antiserum to the highly reactive domain, located immediately downstream to the cleavage site (region C), reacted with only a doublet of 45 and 50 kDa (Figure 4, lane 3), similar to the two different sized bands previously described as size variants of the *vlhA1* MSPA protein [10]. Finally, the antiserum directed against the C-terminal portion of MS2/28.1 (region D) failed to recognize a distinguishable protein band (Figure 4, lane 4). By contrast, this antiserum strongly reacted in filter colony immunoblotting assay (Figure 5C), suggesting that this C-terminal

![Figure 2](image-url)  
**Figure 2** Comparison of the amino acid sequence predicted from *M. synoviae* MS2/28.1 gene with *vlhAs* 1 to 5. Alignment of the completed full-length MS2/28.1 deduced amino acid sequence with *vlhAs* 1 to 5 (GenBank accession numbers AF035624, AF085697, AF085698, AF181033, and AF181034, respectively). Identical aa regions are shaded in black while similar aa residues are shaded in grey.

![Figure 3](image-url)  
**Figure 3** Confirmation that MS2/28.1 is preceded by the unique *vlhA1* promoter sequence. Primer Expro, which anneals to the *vlhA1* promoter, was combined with either *vlhA1R* (lanes b) or with ORF5.1R (lanes c). No amplification from genomic DNA extracted from the four colonies was obtained with the *vlhA1*-specific reverse primer (lanes b). Expected amplicon was obtained with primers Expro/ORF5.1R (lanes c). PCR amplification of the full length MS2/28.1 was obtained with the primers pair Expoint and 2/28.1Rev (lanes d). As negative control, PCR was performed with no genomic *M. synoviae* DNA (lane a). Lane M, DNA size marker (1 kb).
region of MS2/28.1 protein is exposed at the cell surface. The C-terminal highly divergent region of MS2/28.1 encoded product was haemagglutination competent. Mycoplasma synoviae strain WVU 1853 antigen prepared from a single colony culture with an equivalent titer of 3 \times 10^3 CFU/ml showed haemagglutination of chicken red blood cells at a high dilution of 1:256, corresponding to a titer of 2 \times 10^4 CFU/ml. In addition, uniform hemadsorption of chicken erythrocytes to MS2/28.1 expressing colonies was demonstrated (additional file 1). Strikingly, the E. coli-expressed C-terminal 60 residues of MS2/28.1 showed an haemagglutination activity. Consistently, the antiserum raised against this C-terminal highly diverged region inhibited (at a 1/00 dilution) chicken erythrocytes haemagglutination. Collectively, these data demonstrate that the haemagglutinating activity of the vlhA variant MS2/28.1 maps to its surface-exposed and highly divergent C-terminal 60 residues.

Discussion

The molecular basis underlying the antigenic variability of M. synoviae vlhA protein, the abundant immunodominant surface haemagglutinin, has been attributed to site-specific recombination, where recruited vlhA pseudogene copies fuse with the unique expressed vlhA gene sequence [17]. Such a gene replacement mechanism, also known as gene conversion, allows a single strain of M. synoviae to generate a large number of variants by recruiting new sequences from a large pseudogene reservoir. This pseudogene reservoir was found to be confined to a restricted region of the genome [4,16], providing an optimal environment for site-specific recombination. The finding that MS2/28.1 gene sequence occurs in tandem with another vlhA related gene (MS2/28.2), suggests that it is part of this pseudogene reservoir.

Overall, the data point to the selection and clonal expansion of a WVU 1853 bacterial cell expressing a variant vlhA gene with an exceptionally highly divergent haemagglutinin region, comparatively to the expressed vlhA variant sequences described to date [17]. Indeed, all tested colonies contained an MS2/28.1 sequence located immediately downstream of the unique vlhA1 promoter. Comparative sequence analyses with the previously full-length vlhA genes, suggest that gene replacement could have occurred from aa residue 224 to the carboxy terminus. This finding adds a new 5’ recombination site to the previously identified three sites (codon for residues 136, 356, and 442) [17], thus increasing the potential to generate antigenic variability.

Selection of clones expressing other vlhA1-related genes from a culture of M. synoviae WVU 1853, led to the identification of two variant clones, referred to as vlhA4 and vlhA5 [17]. These expressed variants showed a predicted protein length close to that of vlhA1 and diverged in their amino acid sequence by only 15% and 25%, respectively, from residue 211 to the carboxy terminus. This limited sequence variability most likely allows maintaining proper vlhA processing, subcellular location, and haemagglutination activity, while providing sufficient antigenic variability. By contrast, the coding sequence of the full-length MS2/28.1 ORF is considerably shorter than vlhA1, from which it diverged by 64%.

The results showed that this highly variant sequence was properly processed, with its C-terminal highly divergent region exposed at the cell surface. In addition, the M. synoviae clone expressing MS2/28.1 efficiently haemagglutinated chicken erythrocytes, a property that was mapped to the surface-exposed and highly divergent C-terminal 60 residues of MS2/28.1. As far as could be ascertained, this is the first report mapping the haemagglutinating activity of a M. synoviae vlhA gene. The finding that the antiserum raised against this C-terminal region inhibited the haemagglutinating activity of the homologous M. synoviae culture, definitely confirmed that the surface exposed C-terminal 60 residues of MS2/28.1 is associated with haemagglutination. It remains to be seen whether other regions of MS2/28.1 contribute to haemagglutination.

Figure 4 Immunoblot of M. synoviae total antigens probed with antisera raised against regions A to D. Lanes 1 to 4 show immunostaining of M. synoviae whole-cell proteins with antisera raised against regions A to D respectively. Lane 5 shows the reactivity of the anti-M. synoviae polyclonal serum. Prestained broad range protein molecular mass markers are indicated in the left margin.
The results described above highlight the extent to which \textit{vlhA} genes could vary, in both the size and the sequence composition, without compromising their haemagglutination activity. Hence, comparing the expressed sequences from several naturally evolved haemagglutinin variant clones may help identifying critical residues involved in the haemagglutinating activity of \textit{vlhA}. These variations would enable the bacterium to expose an antigenically highly divergent product to better escape the immune system \cite{6,17}. Such a plausible consequence is presently under investigation. However, we anticipate that, during natural infection, in the face of the immune pressure, such an antigenic shift may occur frequently. It would thus be of interest to perform sequence comparisons between naturally derived \textit{vlhA} gene sequences by focusing on their variable haemagglutinin portion. Finally, because site-specific recombination events within \textit{vlhA} genes occur frequently through \textit{in vitro} culture passages, inter-laboratory variations in \textit{M. synoviae} stocks that had been colony purified are likely to exist.

**Conclusions**

The present study provided an indication of the extent to which the \textit{vlhA} haemagglutinin gene of \textit{M. synoviae} could vary without compromising the surface exposure and the haemagglutinating activity of its encoded product. We thus anticipate that the antigenic repertoire of \textit{M. synoviae} \textit{vlhA} gene could be much wider than previously thought.

**Methods**

**Bacterial strains, plasmids and culture conditions**

\textit{Mycoplasma synoviae} strain WVU 1853 was obtained from the American Type Cell Culture collection (ATCC 25204) and grown in Frey's medium \cite{19} supplemented with 15% (v/v) foetal calf serum. The strain was initially passaged in vitro at least 7 times before being subjected to three colony purification steps. A single colony was selected and grown. All mycoplasma cultures were then prepared from this primary stock and never exceeded two additional passages. Culture conditions and antigen preparation were performed as described elsewhere \cite{20,21}. The mycoplasma antigens were stored at -20°C until they were needed either for Western blot, RNA or DNA extraction protocols. The growth of \textit{E. coli} strains was carried out in LB or 2YT broths \cite{22}. Ampicillin was added to the media at the concentration of 100 \(\mu\)g/ml and Isopropyl\-B-D-thiogalacto-pyranoside (IPTG) was included for induction of protein expression at the concentrations recommended by the manufacturer (GE Healthcare) when the screening was performed in pGEX-4T-1 plasmid transformed into \textit{E. coli} BL21. Growth temperature were 37°C, except where indicated and growth rates were estimated by measuring the increase in OD\textsubscript{600}.

**Origin of the immunoreactive MS2/28 DNA fragment**

Isolation and characterization of the \textit{M. synoviae} DNA fragment MS2/28 [GenBank: MSU66315] was previously described \cite{18}. MS2/28 contains two partial ORFs, referred to as MS2/28.1 (5' end) and MS2/28.2 (3' end).

**Reverse transcription and polymerase chain reaction (RT-PCR)**

The total RNA of \textit{M. synoviae} strain WVU 1853 was isolated from a 24-h culture, using a protocol recommended for Gram-positive bacteria \cite{23}. Genomic \textit{M. synoviae} DNA was eliminated from the RNA preparation using DNAse I (2.5 mg/ml) digestion for 1-h period at 37°C. DNase I-treated total RNA of \textit{M. synoviae} was prepared as described above. Reverse transcription was performed at 55°C in a 20 \(\mu\)l reaction mixture containing 2 \(\mu\)g of total RNA, 4 \(\mu\)l of dNTP at 20 mM each, 12.5 \(\mu\)M of the reverse primer 2/28.1Rev (5'-GGGGCGGGCCTACACTGAGTACCTTGGCG-3'), 20 units of AMV reverse transcriptase and 2 \(\mu\)l of 10 \(\times\) buffer reaction (50 mM Tris-Cl, 8 mM MgCl\textsubscript{2}, 30 mM
KCl, 1 mM dithiotreitol, pH = 8). The first strand cDNA synthesis was allowed to proceed for 1 h followed by inactivation at 65°C during 10 min. PCR amplification was next performed using 2/28.1Rev coupled to the PromF primer (5'-GTCGACGAAATTAAGTATAAT-TATTAAG-3') which anneals to the 5' end region (-120 to -98) of the expected vlhA1-derived transcript. The amplification reaction consisted of 30 cycles of 94°C for 120 s, 55°C for 120 s and 72°C for 120 s, followed by an extension of 72°C for 7 min.

**Cloning and sequencing of the RT-PCR product**

The 1.934 kb RT-PCR product was purified and ligated into NotI/SalI-digested pBluescript II KS+ plasmid. The ligation product was used to transform E. coli HB101 cells and recombinant clones were screened using restriction analysis. Determination of the nucleotide sequence was performed with the Prism Ready Reaction Dye Deoxy Terminator Cycle sequencing Kit on an ABI PRISM 377 DNA sequencer (Applied Biosystems). The cloned amplicon was sequenced in both orientations from two different plasmid clones using sequence-specific internal and plasmid-anchored primers. The sequence data were edited and aligned using the software programs BioEdit [24] and ClustalW [25].

**Confirmation of the position of the completed MS2/28.1 gene sequence relative to the unique vlhA1 promoter**

Using genomic DNA extracted from single colonies as template, PCR amplifications were performed, combining EXpro (5’-CAAAAATTAGGATTTACATCTACTA-3’), a sense primer placed in the vlhA1 promoter region (-213 to -193), with either vlhA1 R (5’-TATTGTGTTCCGCTATTTGTCGTC-3′), a vlhA1-specific reverse primer, or ORF5.1R (5’-CGCTTCACCTTCATCTGGCTTACACTTGGC-3’), the MS2/28.1-specific reverse primer. To ensure that the full-length MS2/28.1 gene sequence was placed downstream of the vlhA1 promoter sequence, a second sense primer, EXpront (5’-ATTTACATTTTGTTACTCTTCACATT-3’), located in the vlhA1 promoter region (-172 to -151) was combined with 2/28.1Rev reverse primer (5’-GGGCGGCCGCTGAGCTGTTCATTGGAAT-3’), which anneals to the 3’-end sequence of MS2/28.1.

**Escherichia coli expression of distinct regions of the MS2/28.1 and purification of their products**

The coding sequences corresponding to amino acids 1 to 324 (the N-terminal region, region A), 326-344 (region B, 19-amino acid stretch lying immediately upstream of the putative cleavage site), 354-460 (region C, the region immediately downstream of the cleavage site), and 546-604 (the C-terminal 60 residues, region D) of the full-length MS2/28.1-associated ORF (referred to as MS2/28.1) were amplified by PCR using the three primer pairs 2/28.1For (5’-GGGATCCATGAAAATAAAAAATATTTAATTACATT-3′)-TGA1R (5’-GCGGCGCTTGAGCTGTTCATTGGAAT-3′), TGA1F(5’-GGATCCATTTCAAAGCTCAGCTCA-3′)-TG2AR (5’-GGGCGGCCGAGCTTTGGACTCAAGCTCTA-3′), and TGA6F (5’-GGATTCAATATTGGAATACAATAAATTTACCACA-3′)-2/28.1Rev (5’-GGGCGCCCTACCTTGACGTAGCTTGGC-3′) and cloned into BamHI/NotI-restricted pGEX-4T-1 expression vector, after being verified by nucleotide sequencing. The coding sequence of the region immediately downstream of the cleavage site (354-460, region C) was obtained from a plasmid containing the MS2/28.1 segment and subcloned in the EcoRI site of pGEX-4T-1. The recombinant plasmids encoding regions A, B, C, and D of MS2/28.1 were electroporated into competent E. coli strain BL21, to produce the GST-MS2/28.1A, GST-MS2/28.1B, GST-MS2/28.1C, and GST-MS2/28.1D fusion proteins, respectively. Briefly, overnight cultures of transformed bacteria were diluted 1:100 of 2YT medium containing ampicillin (100 µg/ml) and grown at 37°C C with shaking (250 rpm) to an A600 of 0.6. Protein expression was induced by the addition of 0.1 mM IPTG, and maintained for 4-h incubation at 30°C with vigorous agitation (250 rpm). The cells were then pelleted by centrifugation at 3000 rpm and resuspended in 1× PBS. The E. coli pellet was disrupted by sonication and solubilized with 1% Triton X-100 (Sigma) of 30 min. Both fusion proteins proved to be soluble and were readily purified by affinity chromatography on Glutathione Sepharose 4B Beads (GE Haelthcare), using the Bulk GST Purification Module, following the instructions of the manufacturer. The purified recombinant proteins were analysed by electrophoresis on sodium dodecyl sulfate (SDS)-12% polyacrylamide gels and allowed to react, in western immunoblotting, with a rabbit polyclonal anti-M. synoviae serum.

**Production of monospecific antiserum to MS2/28.1 regions A, B, C, and D**

Polyclonal monospecific antiserum to the purified fusion proteins GST-MS2/28.1A, GST-MS2/28.1B, GST-MS2/28.1C, and GST-MS2/28.1D were raised in female New Zealand White rabbits. Pre-immune serum was collected from each animal followed by intramuscular immunization on day 1 with 200 µg protein mixed with an equal volume of Freund’s complete adjuvant. Subsequent immunizations with 200 µg protein in Freund’s incomplete adjuvant were given at 2 week intervals. The produced antiseras were then used to identify, in M. synoviae, the proteins encoded by MS2/28.1.

A rabbit polyclonal anti-M. synoviae serum was generated as described above, using 200 µg of sonicated total M. synoviae antigen.

The immunization of rabbits and collection of sera were performed following the protocols approved by the Center for Biologics Evaluation and Research/Food and...
Drug Administration Institutional Animal Care and Use Committee.

Identification and Characterization of MS2/28.1 encoded proteins

*M. synoviae* total proteins were separated on SDS-polyacrylamide gels and electrophoretically transferred to nitrocellulose membranes (Bio-Rad). Rabbit antisera directed either against the fusion proteins or the whole *M. synoviae* antigen were then reacted with these membranes followed by incubation with a goat anti-rabbit immunoglobulin peroxidase conjugate (Sigma). The reactive protein bands were visualized using a substrate solution consisting of 0.05% 4-chloro-1-naphtol (Sigma) in PBS containing 20% (v/v) methanol.

Filter colony blotting

Fresh *M. synoviae* colonies growing on the surface of agar plates were transferred to nitrocellulose membrane discs (Bio-Rad). Discs were dried for 5 min at room temperature, then, they were incubated in blocking buffer (1 × PBS/5% BSA (Sigma)) for an hour. The discs were washed three times for 5 min in wash buffer (1 × PBS/0.1% BSA/0.05% Tween 20 (Bio-Rad)) and then incubated for 2 h with the primary antibody (diluted in wash buffer). After three brief washes, nitrocellulose discs were incubated for 1 h with peroxidase-conjugated secondary antibody against rabbit IgG (GE Healthcare) diluted at 1:3,000 in washbuffer. Colony blots were visualized, after washing steps, with substrate solution containing 4-chloro-naphtol (Bio-Rad) as chromogen and the reaction was stopped by washing blots in deionised water.

Haemagglutination and haemagglutination inhibition (HI) tests

Purified *M. synoviae* colonies were grown as described previously then harvested and adjusted to an equivalent in 50 dilutions of the mycoplasma cell suspension were made in rows of a U-bottomed microtiter plate duplicate serial twofold PBS to 1/50 of the original broth volume. In rows of a colony hemadsorption assay

Distinct colonies of *M. synoviae* strain WVU 1853 derived from a single clone expressing MS2/28.1 (2.5 × 10^7 CFU/ml), were observed on Frey’s agar after incubation for 48 h at 37°C, 5% CO₂. The colonies were covered with 15 ml of 0.5% chicken erythrocytes in PBS and incubated for 1 h at 37°C. Agar plate was then gently washed twice with PBS and examined at low magnification under a microscope for erythrocyte adherence to mycoplasma colonies.

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Authors’ contributions

ABK carried out major experimental work (PCR, RT-PCR, sequencing, sequence analysis, protein expression, production of polyclonal antisera, immunoblotting, filter colony blotting, haemagglutination and hemadsorption assays). Expression of the MS2/28.1C region and production of its monospecific antiserum were performed by GI. RBM carried out the amplification of MS2/28 5’-end cDNA and the completion of MS2/28 coding sequence. BBAM conceived, designed the study, and drafted the manuscript. All authors approved the final version of the manuscript.

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References

1. Keven SH: *Mycoplasma synoviae* infection. Diseases of Poultry Iowa State Press AmesSaif YM, Barnes HJ, Gilsson JR, Fady AR, McDougall LR, Swaye DE 2003, 756.
2. Feberwee A, de Wit JJ, Landman WJ: Induction of eggshell apex abnormalities by *Mycoplasma synoviae* field and experimental studies. Avian Pathol 2009, 38:187.
3. Calderon-Copete SP, Wigger G, Wunderlin C, Schmidheini T, Frey J, Quail MA, Falquet L: The Mycoplasma conjunctivae genome sequencing, annotation and analysis. BMC Bioinformatics 2009, 10:57.
4. Vascconcelos AT, Ferreira HB, Bizarro CV, Bonatto SI, Calvano NO, Pinto PM, Almeida DF, Almeida LG, Almeida R, Alves-Filho L, Assunção EN, Azevedo VA, Bogo MR, Bratado MW, Brocchi M, Burity HA, Camargo AA, Camargo SA, Carepo MS, Carraro DM, de Mattos Casciano JC, Castro LA, Cavaletti CA, Chemale G, Collevatti RG, Cunha CW, DallaGiovanna B, Dambrós DP, Dellagostin OA, Falcão C, Fantinatti-Carborgi F, Felipe MS, Fiorentin L, Franco GR, Freitas NS, Fries D, Grangeiro TB, Grisard EC, Guimarães CT, Hungria M, Jardim SN, Kreger MA, Laurino JP, Lima LF, Lopes MI, Loreto EL, Madeira HM, Manfrin GP, Maranhão AQA, Martinkovics CT, Medeiros SR, Moreira MA, Neiva M, Ramalho-Neto CE, Nicolás MF, Oliveira SC, Paiva RD, Pedroso FO, Penna SD, Pereira M, Pereira-Ferrari L, Pifferi I, Pinto LS, Potrich DP, Salim AC, Santos FR, Schmitt R, Schneider MP, Schraniik A, Schrank G, Schuck AF, Seuanez HN, Silva DW, Silva R, Silva SC, Soares CM, Souza KR, Souza RC, Staats CC, Steffens MB, Teixeira SM, Urményi TP, Vanneste MH, Zucchero LW, Simpson AJ, Zaha A: Swine and Poultry Pathogens: the Complete Genome Sequences of Two Strains of Mycoplasma hyopneumoniae and a Strain of Mycoplasma synoviae. J Bacteriol 2003, 185:5568-5577.
5. Sisani-Pugnet F, Larrique C, Marenza M, Jacob D, Baré A, Baré V, Schenowitz C, Mangelot S, Coulou L, Seguens B, de Daruvar A, Blanchard C, Citt C: Being pathogenic plastics, and sexual while living with a nearly minimal bacterial genome. PLoS Genet 2007, 3:744-758.
6. Bencina D: Haemagglutinins of pathogenic avian mycoplasmas. Avian Pathol 2002, 31:535-547.

Additional file 1: Hemadsorption of chicken erythrocytes on *M. synoviae* colonies. Adherence of chicken erythrocytes to colonies of *M. synoviae* expressing the viH4 variant MS2/28.1 cultured on Frey’s agar.
Phenotypic switching in mycoplasmas: Phase variation of diverse surface lipoproteins. Science 1990, 247:315-318.

Gorton TS, Geary SJ. Antibody-mediated selection of Mycoplasma gallisepticum phenotype expressing variable proteins. FEMS Microbiol Lett 1997, 155:31-38.

Naray M, Bencina D, Kleven SH, Habe F. The Haemagglutination-Positive Phenotype of Mycoplasma synoviae Induces Experimental Infectious Synovitis in Chickens More Frequently than Does the Haemagglutination-Negative Phenotype. Infect Immun 1998, 66:6004-6009.

Noormohammadi AH, Markham PF, Whithear KG, Walker ID, Gurevich VA, Ley OH, Browning GF. Mycoplasma synoviae has two distinct phase-variable major membraneantigens one of which is a putative haemagglutinin. Infect Immun 1997, 65:2542-2547.

Noormohammadi AH, Markham PF, Duffy MF, Whithear KG, Browning GF: Multigene families encoding the major haemagglutinins in phylogenetically distinct mycoplasmas. Infect Immun 1998, 66:3470-3475.

Bencina D, Naray M, Dave P, Drobvic-Valic M, Habe F, Kleven SH. The characterization of Mycoplasma synoviae EF-Tu protein and proteins involved in hemadherence and their N-terminal amino acid sequences. FEMS Microbiol Letters 1999, 173:85-94.

Markham PF, Grew MD, Sykes JE, Bowden TR, Pollocks TD, Browning GF, Whithear KG, Walker ID. The organisation of the multigene family which encodes the major cell surface protein, pMGA, of Mycoplasma gallisepticum. FEBS Lett 1994, 352:347-352.

Markham PF, Duffy MF, Grew MD, Browning GF: A gene family in Mycoplasma imitans closely related to the pMGA family of Mycoplasma gallisepticum. Microbiology 1999, 145:2095-2103.

Grew MD, Baseggio N, Markham PF, Browning GF, Walker ID: Expression of the pMGA genes of Mycoplasma gallisepticum is controlled by variation in the GAA trinucleotide repeat lengths within the 5′ non-coding regions. Infect Immun 1998, 66:5833-5841.

Allen JL, Noormohammadi AH, Browning GF: The vlhA loci of Mycoplasma synoviae are confined to a restricted region of the genome. Microbiology 2005, 151:935-940.

Noormohammadi AH, Markham PF, Kanci A, Whithear KG, Browning GF: A novel mechanism for control of antigenic variation in the haemagglutinin gene family of Mycoplasma synoviae. Mol Microbiol 2000, 35:911-923.

Ben Abdelmoumen B, Roy RS, Brousseau R: Cloning of Mycoplasma synoviae genes encoding specific antigens and their use as species-specific DNA probes. J Vet Diag Invest 1999, 11:162-169.

Frey ML, Hansen RP, Anderson DP: A medium for the isolation of avian mycoplasmas. Avian Pathol 1968, 29:2163-2171.

Ben Abdelmoumen B, Roy RS: An enzyme-linked immunosorbent assay for detection of avian mycoplasmas in culture. Avian Dis 1993, 39:251-262.

Sambrook J, Fritsch EF, Maniatis T: Molecular Cloning: A Laboratory Manual. Cold Spring Harbor, Cold Spring Harbor Press 1989.

Ausubel FM, Brent R, Kingston RE, Moore DD, Seidman JG, Smith JA, Struhl K. RNA extraction from gram positive bacteria. Current protocols in Molecular Biology 1997, 1, 4.4.3.

Hall TA: BioEdit: A user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. Nucleic Acids Symp 1999, 41:95-98.

Thompson JD, Desmond GH, Toby JG: ClustalW: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. Nucleic Acids Res 1994, 22:4673-4680.

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