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Molecular characterization of field strains of *Mycoplasma gallisepticum* in Malaysia through *pMGA* and *pVPA* genes sequencing

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Abstract: To characterize field strains of *Mycoplasma gallisepticum* (MG) in Malaysia, selected gene target-specific sequences to MG, hemagglutinin protein A (*pMGA*) and phase variable putative adhesion protein A (*pvpA*) genes, were amplified from positive MG samples using conventional polymerase chain reaction (PCR). A total of 25 MG-positive samples out of 94 field samples were sequenced with primer targeting *pMGA* and *pvpA* genes. Sequencing and phylogenetic analysis were conducted using bioinformatics softwares (Bioedit and MEGA 5) and genetic variation patterns were evaluated based on partial nucleotide sequencing of *pMGA* and *pvpA* genes. In case of *pMGA*, all 25 field strains showed similar gene size patterns with strains of pathogenic reference (MGS6) and vaccine (MG F), which were different from the PCR product size of less pathogenic vaccine strain (TS 11). However, *pvpA* partial nucleotide sequences of MG local strains showed that 20 out of 25 strains possessed similar gene size pattern with MG F, which were different from MGS6 and TS 11. In conclusion, based on the molecular characterization from phylogenetic analysis of *pMGA* and *pvpA* partial nucleotide sequences, it was found that Malaysian MG strains were different from strains reported in other countries.

ABOUT THE AUTHORS



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Farhana Yasmin is a dynamic, enthusiastic and hardworking veterinary professional who practices on ruminants and birds. She has done DVM and MS from the Hajee Mohammad Danesh Science and Technology University (HSTU), Bangladesh and Universiti Putra Malaysia (UPM), Malaysia respectively. In her master's programme, she worked in the field of molecular biology and technically, she has gained skills on sample collection and preservation, culture, DNA extraction, DNA cloning, conventional PCR and sequencing with phylogenetic analysis and real-time PCR with efficiency and specificity for the thesis concerning "Development of a Diagnostic Real Time PCR Assay, Molecular Detection and Characterization of *Mycoplasma gallisepticum*". This paper was culled from the MS thesis supervised by Aini Ideris, Abdul Rahman Omar and Mohd Hair-Bejo. Tan Sheau Wei, Tan Ching Giap, Rakibul Islam and Kartini Ahmad were her labmates and research colleagues who helped to collect the data and write the manuscript.

PUBLIC INTEREST STATEMENT

Mycoplasma gallisepticum is a primary pathogen that can cause chronic respiratory disease (CRD) which leads to severe economic losses to the commercial poultry industry through reduced feed efficiency, decreased egg production and increased mortality. Although the farmers follow strict biosecurity management, vaccination programme and medicinal treatment in the poultry farm to prevent this disease, recurrent infection and outbreaks are still present. Therefore, the control of mycoplasmosis is very important in order to increase the output of poultry farm. MG genes are related with pathogenic, antigenic and immune evasion properties. It has been reported that molecular characterization enables epidemiological tracing and classification of new field MG strains. So, this study will be a core for further efficient evaluation to develop a potential strategy for prevention and control of MG infection.

Subjects: Microbiology; Molecular Biology; Immunology; Medicine

Keywords: *Mycoplasma gallisepticum*; pMGA and pvpA genes; gene size pattern; phylogenetic analysis; Malaysia

1. Introduction

Mycoplasma are members of the Mycoplasmataceae family, which belongs to the Mycoplasmatales order in the Mollicutes class (Sasaki et al., 2002); and there are more than 200 species in this class. One of the species is *Mycoplasma gallisepticum* (MG), which has a worldwide distribution and is the most economically important pathogenic avian *Mycoplasma*. MG is a primary pathogen that can cause chronic respiratory disease (CRD) which leads to severe economic losses to the commercial poultry industry (Liu, Garcia, Levisohn, Yogev, & Kleven, 2001).

Sequencing methods have been introduced to study molecular epidemiology of bacterial pathogens (Enright & Spratt, 1999). To differentiate MG strains, the improvement on the molecular biology of MG (Razin, Yogev, & Naot, 1998) and MG complete genome sequence availability have been used to evaluate gene-targeted sequencing as a typing tool (Papazisi et al., 2003). DNA polymorphisms are the basis for the application of PCR based methods to detect and differentiate MG strains. Nascimento, Yamamoto, Herrick, and Tait (1991) showed that, gene-targeted sequencing analysis might help in the synthesis of specific strain primers and can be used to detect MG strains.

The adherence of MG can be affected by the changes in expression of both *pMGA* and cytoadhesin genes (*mgc1* and *mgc3*) but *pMGA* genes play a big potential role for antigenic variants generation (Bencina, 2002). Basically, *pMGA* and *pvpA* are two gene families of MG that encode major surface proteins having pathogenic, antigenic and immune evasion properties (Markham, Glew, Whithear, & Walker, 1993). The *pMGA* gene family involves a minimal of 16% of R strain and 7.7% of F strain (Baseggio, Glew, Markham, Whithear, & Browning, 1996) which are powerful genomic engagements to antigenic variability, and hypothetically play a key role in immune evasion (Markham et al., 1993). The *pMGA* gene family also accommodate a system for prompt and reversible switches in its expression of proteins (antigenic switching) in response to antibodies or other environmental clues (Glew, Browning, Markham, & Walker, 2000). The *pvpA* is integral membrane proteins of MG having size variation that possess high-frequency phase and antigenic capability (Levisohn, Rosengarten, & Yogev, 1995). In this study, genes used for sequencing were *pMGA* and *pvpA* which had size polymorphism (Tan, 2008).

In Malaysia, MG field strains showed gene size polymorphism in *pMGA* and *pvpA* genes with a significant pattern in comparison with reference pathogenic and vaccine strains (Zahraa, Aini, Omar, Hair-Bejo, & Tan, 2011). However, there is very limited information on molecular characterization; regarding the comparison of Malaysian local strains with international strains, upon analysis of phylogenetic tree based on *pMGA* and *pvpA* genes. Therefore, the objectives of the study were to sequence *pMGA* and *pvpA* genes of MG field strains and also to characterize these local strains based on *pMGA* and *pvpA* phylogenetic analysis.

2. Materials and methods

2.1. *Mycoplasma gallisepticum* samples

A total of 300 swab samples were obtained from different states of poultry farms in Malaysia where 94 were MG-positives field strains by real-time PCR using gapA 5F + 6R primer set of *gapA* gene (Yasmin et al., 2014). Ninety-four positive samples were also amplified by conventional PCR using AUTS11 F + ATTS11 R and pvpA 1F + pvpA 2R primer sets of *pMGA* and *pvpA* genes accordingly. Based on the brightness and thickness of bands, 25 field strains out of 94 samples were selected and amplified again by conventional PCR for molecular characterization. The sampling size, location, vaccination, farming type and number of positive samples are summarized in Table 1.

Table 1. Sampling size, sampling location and vaccination history of layer breeder, broiler breeder, broiler chicken and village chicken used in this study

Farm No.	Farm ID	Farming type	Vaccination	State	No. of samples	pMGA	pvpA
1	SMF1A	Layer breeder	Yes	Melaka	15	13	9
2	SMF1B	Layer breeder	Yes	Melaka	15	5	11
3	SMF5B	Layer breeder	Yes	Melaka	15	3	3
4	SMF2C	Layer breeder	Yes	Melaka	15		
5	TTJF1	Broiler chicken	Yes	Melaka	15	1	
6	TTJF2	Broiler chicken	Yes	Melaka	15		
7	SM4	Broiler chicken	Yes	Melaka	15		
8	SM5	Broiler chicken	Yes	Melaka	15		
9	G5	Layer breeder	Yes	Melaka	20		
10	SMF3	Layer breeder	Yes	Melaka	15		
11	JSF1	Broiler chicken	Yes	Johor	30	2	
12	JSF2	Broiler chicken	Yes	Johor	30		
13	HL	Village chicken	No	Selangor	10	1	1
14	Tanj	Broiler chicken	Yes	Selangor	30		1
15	RNS	Village chicken	No	Negeri Sembilan	24		
16	KS	Broiler breeder	Yes	Selangor	21		
Total					300	25	25

2.2. Extraction of genomic DNA

An ethanol-cleaned forceps was used each time to remove cotton swab from the phosphate-buffered saline (PBS) under sterile condition. Conventional salt-based method was used to extract genomic DNA with some modifications. In the first step of DNA extraction, 100 μ l of 10% sodium dodecyl sulphate (SDS) was used to lyse cells and tissues as well as to remove lipid membrane. Then, cellular and histone proteins bound to DNA were removed by adding 2 μ l of 50 μ g/ μ l Proteinase K (Epicentre® Biotechnologies, USA). The pellet was resuspended by vortexing the mixture. The mixture was then incubated at 65°C for 30 min and thoroughly shaken in every 5 min. Then, all the tubes were left at -20°C for 8 min or at 4°C for 30 min to 1 h.

Protein and peptide were precipitated by adding 300 μ l of ammonium acetate and vortex for 10 s to mix it properly. A micropipette was used to resuspend the pellet if it stuck at the bottom of tube and the pellet was centrifuged at 14,000 rpm for 15 min. Then, a volume of 850 μ l from the supernatant was saved by transferring it into another new microcentrifuge tube. For precipitation of DNA, a volume of 550 μ l of isopropanol was added into the supernatant and the tube was inverted about 40 times. The mixture was then centrifuged at 14,000 rpm for 15 min at 4°C. The isopropanol was then poured off carefully to avoid DNA pellet dislodging. Resultant DNA pellet was washed twice in 1 ml of 75% (v/v) ethanol by centrifugation at 14,000 rpm for 15 min at 4°C. The ethanol was poured off carefully so that no dislodge of DNA pellet. The DNA samples were then placed as inverted position in the laminar air flow chamber for 2 h until the evaporation of ethanol. At last, these samples were resuspended in 30 μ l of nuclease free water, then kept at 4°C for 30 min and finally stored at -20°C until to be used for PCR.

Table 2. PCR product size and primer sequences of *pMGA* and *pvpA* genes used for MG characterization

Primer	Gene	Sequence 5' to 3'	PCR Product Size (bp)	Optimal Annealing Temp.°C
AUTS11 F	<i>pMGA</i> or	TCTTCTTCGAAAA- CAAAAGG		
ATTS11R	<i>vlhA</i> genes family	GTTTGGAGTTGGTG- TATAGTTAG	~329	52
pvpA 1F	<i>pvpA</i>	GCCAMTCCAAC TCAA- CAAGCTGA		
pvpA2R		GGACGTS GTCCTGGCTGGT- TAGC	~702	58

Reference: Zahraa et al. (2011) and Kartini (2012).

2.3. PCR assay targeting *pMGA* and *pvpA* genes

For molecular characterization, field strains of positive MG with reference strains were amplified by conventional PCR for gene sequencing. Primers used in this study (Table 2), were synthesized by First Base Laboratories Sdn. Bhd., Selangor, Malaysia. The PCR protocol was optimized by varying conditions, viz. DNA concentration, amplification cycle, amplification time and primer annealing temperature. An automatic thermal cycler (MyCycler, BioRad, USA) was used to amplify fragmented DNA and the reaction set-up was consisted of an initial denaturation step at 94°C for 4 min, followed by 40 cycles of denaturation at 95°C for 15 s, annealing at 52°C (for *pMGA* gene) and 58°C (for *pvpA* gene) for 60 s, extension at 72°C for 15 s and final extension at 72°C for 5 min. The reaction volume was set up in a 25- μ l reaction mixture where 12.5 μ l (2X) (Bioline, Malaysia) of 2X MyFiMix, 0.4 μ l (25 pmol) of forward primer, 0.4 μ l (25 pmol) of reverse primer, 2 μ l (200 ng) of DNA template and 9.7 μ l of distilled water. The gel electrophoresis system at 1.5% agarose gel was used to view the result of amplified DNA in 1X TAE performed at 90 V for 40 min, stained with ethidium bromide (10 mg/ml). Bands were visualized on UV light and the photograph was taken with analyser (Syngene, Gene Genius Bioimaging System). PCR products were compared with 100-bp plus DNA ladder (Vivantis®, Malaysia) and the presence of band indicated a positive result, whereas the absence of band indicated a negative result.

2.4. DNA purification, DNA sequencing and sequence analysis

Positive bands were selected for DNA purification which was performed using the PCR SV kit (Gene All® Expin™ PCR SV). Purified DNA of PCR products were sent for sequencing to Macrogen Inc, Seoul, Korea. Nucleotide sequences of *pMGA* and *pvpA* genes were identified through direct sequencing from both directions using primers as shown in Table 2. The Automatic Sequencer-ABI 3730XL DNA Analyzer using the ABI PRISM® BigDye™ Terminator Cycle Sequencing Kits with AmpliTaq DNA Polymerase (FS enzyme) (Applied Biosystem, CA, USA) was used to perform sequencing reactions. Purified DNAs of reference pathogenic (MGS6, MG R and PG 31) and vaccine (TS 11, MG F and 6/85) strains were also sent for sequencing. The web-based pair-wise BLAST (Basic Local Alignment Search Tool) search programs under NCBI (National Centre for Biotechnology Information) was used for the primary determination and comparison among different strains of the raw sequence data. The Bioedit Sequence Alignment Editor Program with 7.0.9.0 version (Hall, 1999); based on Clustal W Multiple alignment (Thompson, Higgins, Gibson, & Clustal, 1994); was used for the assembling and analysing of sequence data. This program was also used to find sequence identity matrix. Multiple sequence alignment of local field strains, reference strains and international sequences were also analysed using the MEGA with version 5 software program (Tamura et al., 2011). MG international sequences were collected from Gene Bank (Table 3).

Table 3. MG International sequences used for sequence alignment and phylogenetic analysis

Isolate	Origin	Gene with Gene bank accession No.		Reference
		<i>pMGA</i>	<i>pvpA</i>	
80083	Australia	FJ 844438		Ghorashi, Noormohammadi, and Markham (2010)
87081	Australia	FJ 654141		Ghorashi et al. (2010)
B40/95	Australia	GU 166688		Ghorashi et al. (2010)
VR-5	Australia	GU 133052		Ghorashi et al. (2010)
K2101	USA	FJ 654136		Ghorashi et al. (2010)
K1453	USA	FJ 654137		Ghorashi et al. (2010)
K5792D	USA	GU 133051		Ghorashi et al. (2010)
Au94043CK94	Australia		AY 556377	Ferguson et al. (2005)
Au80083	Australia		AY 556376	Ferguson et al. (2005)
Au99169CK99	Australia		AY 556380	Ferguson et al. (2005)
Au97019	Australia		AY 556379	Ferguson et al. (2005)
Au96022CK96	Australia		AY 556378	Ferguson et al. (2005)
IRHB09CK06	Iran		EF 188268	Hosseini, Bozorgmehrfard, Peighambari, Pourbakhsh, and Razzazian (2006)
KS2	Israel		AY556370	Ferguson et al. (2005)
MK8	Israel		AY 556372	Ferguson et al. (2005)
BRT14	Israel		AY 556368	Ferguson et al. (2005)
DSD6	Israel		AY 556369	Ferguson et al. (2005)
OR2	Israel		AY 556373	Ferguson et al. (2005)
Ts8	China		EU 216075	Jiang, Chen, Yan, and Zeng (2009)
NE12	China		EU 216077	Jiang et al. (2009)
MG080508	Russia		FJ 965823	Sprygin et al. (2011)
MG140905	Russia		FJ 965808	Sprygin et al. (2011)
MG310807	Russia		FJ 965819	Sprygin et al. (2011)

Figure 1. Picture of PCR products of MG reference and field strains (12 field samples) by *pMGA* gene using primer AUTS11 F + ATTS11 R. 100-bp plus DNA ladder (lane 1), MGS6 (lane 2), MG F (lane 3), TS 11 (lane 4), 6/85 (lane 5), MG R (lane 6), PG 31 (lane 7), A5969 (lane 8), MG-positive field strains (lane 9–20), Negative control (lane 21).



2.5. Phylogenetic analysis

To fulfil the phylogenetic analysis, MG field strains together with reference strains were compared with strains of other countries as shown in Table 3. The neighbor joining tree of the MEGA with version 5 software was used to do phylogenetic analysis (Tamura et al., 2011). The accuracy of clustering of the neighbor-joining tree was assessed using 1000 bootstrap replicates.

Figure 2. Picture of PCR products of MG reference strains and local strains (13 field samples) by *pMGA* gene using primer AUTS11 F + ATTS11 R. 100-bp plus DNA ladder (lane 1), MGS6 (lane 2), MG F (lane 3), TS 11 (lane 4), 6/85 (lane 5), MG R (lane 6), PG 31 (lane 7), A5969 (lane 8), MG-positive field strains (lane 9–21), Negative control (lane 22).

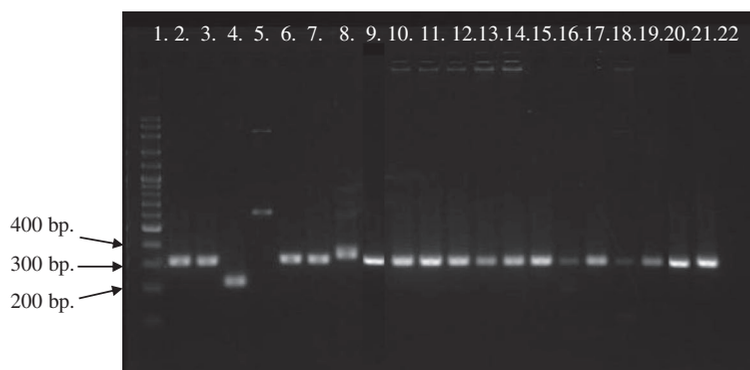
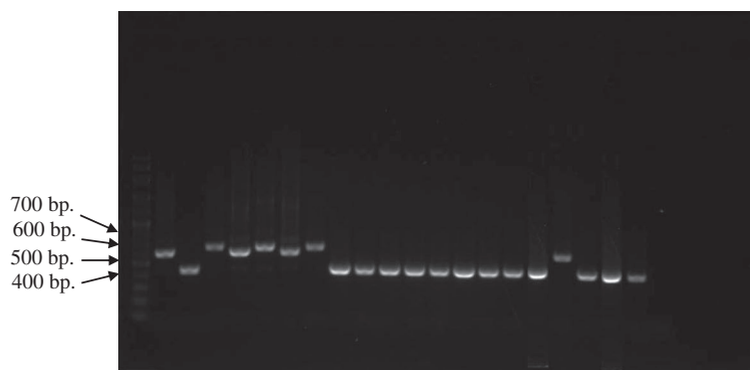


Figure 3. Picture of PCR products of MG reference strains and local strains (12 field samples) by *pvpA* gene using primer *pvpA* 1F + *pvpA* 2R. 100-bp plus DNA ladder (lane 1), MGS6 (lane 2), MG F (lane 3), TS 11 (lane 4), 6/85 (lane 5), MG R (lane 6), PG 31 (lane 7), A5969 (lane 8), MG-positive field strains (lane 9–20), Negative control (lane 21).



Figure 4. Picture of PCR products of MG reference strains and local strains (13 field strains) by *pvpA* gene using primer *pvpA* 1F + *pvpA* 2R. 100-bp plus DNA ladder (lane 1), MGS6 (lane 2), MG F (lane 3), TS 11 (lane 4), 6/85 (lane 5), MG R (lane 6), PG 31 (lane 7), A5969 (lane 8), MG-positive field strains (lane 9–21), Negative control (lane 22).



3. Results

3.1. Amplification of MG field strains by conventional PCR

For sequencing, 25 positive samples were selected for each *pMGA* and *pvpA* gene, which were based on the brightness and thickness of bands (Figures 1–4). Both *pMGA* and *pvpA* genes had size polymorphism on specific target sequence. Specific primer targeting *pMGA* and *pvpA* genes could detect the specific gene on the MG polymerized reaction with the primer-specific annealing temperature, and thus were able to detect the presence of MG in field samples.

Table 4. Analysis of *pMGA* and *pvpA* partial nucleotide sequences

Gene	<i>pMGA</i>	<i>pvpA</i>
PCR product size	226–336	434–677
Polymorphism	Polymorphic	Polymorphic
Insertion	Position 139–156 nt in two field strains	Position 1–65 nt in one field strain
Deletion	None	Position 573 nt—rest in 24 strains
G+C content	29–34%	48–51%
Sequence similarity	23 field strains = 98–100% MGS6	1 field strain = 100% MGS6
	2 field strains = 97–100% TS 11	14 field strains = 98–99% TS 11
		9 field strains = 99% A5969
		1 field strain = 98% PG 31

3.2. PCR result of MG field strains by *pMGA* gene

Analysis of MG PCR products of the *pMGA* gene (or M9 gene) by 1.5% agarose gel electrophoresis indicated that the location of bands were within 200–400 bp (Figures 1 and 2). Gene size variations were observed among PCR products of the *pMGA* gene both local field and reference strains. PCR product sizes of field strains used in this study were varied in between 315 and 336 bp, whereas product sizes of reference strains were ranged in between 226–329 bp. So, the overall range of PCR product sizes were 226–336 bp when were amplified by AUTS11 F + ATTS11 R primer set of *pMGA* gene. The smallest product length observed in vaccine strain (TS 11) was 226 bp, whereas the largest product found in MM33 field strain was 336 bp. Moreover, rest of 24 field strains showed similar gene size pattern with reference pathogenic (MGS6) and virulence vaccine (MG F) strains which were different from the less virulence vaccine strain TS 11 (PCR product 226 bp) and the reference strain A5969 (PCR product 329 bp). However, although all strains showed single band, the vaccine strain (6/85) showed multiple bands indicating nonspecific to the primer set of *pMGA* gene.

3.3. PCR result of MG field strains by *pvpA* Gene

Analysis of MG PCR product size of *pvpA* gene by 1.5% agarose gel electrophoresis indicated that the location of bands were within 400–700 bp (Figures 3 and 4). Gene size variations were observed within PCR products of the *pvpA* gene among field and reference strains. This study also showed the gene size polymorphism among PCR products of MG local strains. PCR product sizes of local and reference strains were varied in between 434–665 bp and 446–677 bp, respectively. So, the overall range of PCR products were ranged in between 434 and 677 bp when were amplified by *pvpA* 1F + *pvpA* 2R primer set of *pvpA* gene. The smallest product length was 434 bp, which was observed in local strains of MM8, MM10, MM12, MM14 and MM19, whereas the largest product size was 677 bp found in the reference strain MG R.

3.4. Sequence analysis

Partial nucleotide (nt) sequence analysis of *pMGA* and *pvpA* genes from MG locals with reference strains were successfully accomplished using Bioedit and MEGA softwares and the summary of findings are shown in Table 4. Result from the analysis of sequence identity matrix and PCR product sizes of *pMGA* and *pvpA* genes are stated below.

3.4.1. *pMGA* gene

Upon the analysis of *pMGA* partial nucleotide sequences from field strains and five reference strains (MGS6, MG F, MG R, PG 31 and TS 11), results indicated that there were nucleotide variation and gene size polymorphism ranged between 226 and 336 bp having 29–34% of GC content. Although no nucleotide deletion was observed, nucleotide insertion of two field strains were found in the position of 139–156 nt. A total of 23 and 2 field strains showed 98–100% and 97–100% sequence similarity with MGS6 and TS 11 accordingly. Local field strains also showed some dissimilarity with strains of other countries regarding nucleotide variation. From the analysis of sequence identity matrix, the

highest similarity was 0.987, which was observed in between MM14 and MM31 field strains. The lowest similarity was found in a field strain MS1 and an international strain 80,083 Australia. Overall, Malaysian local field strains showed maximum identity with the virulence vaccine strain MG F. Mention may be made that the similarity percentage of local field strains were very low to international strains indicated that Malaysian local strains were different from strains of other countries.

3.4.2. *pvpA* gene

Upon the analysis of *pvpA* partial nucleotide sequence from field strains and six reference strains (MGS6, 6/85, MG F, MG R, PG 31 and TS 11), results indicated that there were nucleotide variation and gene size polymorphism ranged between 434 and 677 bp having 48–51% of GC content. There was a nucleotide insertion in one field strain in the position of from 1 to 65 nt and the nucleotide deletion was observed in the position of 573 nt to rest in 24 field strains. A total of 1, 14, 9 and 1 field strains showed 100%, 98–99, 99 and 98% sequence similarity with MGS6, TS 11, A5969 and PG 31, respectively. Local field strains also showed some dissimilarity with international strains regarding nucleotide variation. From the analysis of sequence identity matrix, the highest similarity value was 0.966, which was found in between field strains of MM19 with MM21 and MM14 with MM31. The lowest similarity value was 0.121, which was found in between field strains MM10 with MS2 and MM10 field strain with IRHB09CK06/Iran international strain. Mention also be made that the similarity value was very low between field strains and international strains, which indicated that Malaysian strains were different from strains of other countries.

3.5. Phylogenetic tree analysis

3.5.1. Phylogenetic analysis on the basis of *pMGA* partial nucleotide sequence

A phylogenetic tree was created on the basis of 226–336-bp partial nucleotide sequences of *pMGA* gene (Figure 5). Upon the analysis of phylogenetic tree of *pMGA* partial nucleotide sequences from MG field, reference and international strains, results showed that a total of 24 out of 25 local strains were in the same cluster with the virulence vaccine strain MG F and the pathogenic reference strain MGS6. It also indicated that only the local strain MS1 was grouped with reference (MG R, PG 31 and TS 11) and international strains.

3.5.2. Phylogenetic analysis on the basis of *pvpA* partial nucleotide sequence

A phylogenetic tree was created on the basis of 434–677-bp partial nucleotide sequences of *pvpA* gene from local field, reference and international strains (Figure 6). Upon the analysis of the phylogenetic tree of *pvpA* partial nucleotide sequences It had been illustrated that a total of 23 out of 25 Malaysian local strains were in the same group with the reference of high virulence vaccine strain MG F. were in different group from the pathogenic reference strain, MGS6 and less pathogenic vaccine strain, TS11 and located in the same cluster with two international strains. Only the two field strains (MS1 and MS2) placed in same group with international strains (K5054 USA and IRHB09CK06 Iran) and other reference strains (MG R, TS 11, MGS6, 6/85 and PG 31). It also showed that other international strains were grouped separately as there was no similarity found to Malaysian field strains.

4. Discussion

Mycoplasma gallisepticum (MG) is an important and most pathogenic avian *Mycoplasma* which is responsible for the great economic losses worldwide as it transmits vertically and horizontally. In this study, a total of 25 MG local field strains with reference strains for both of *pMGA* and *pvpA* genes were selected for gene targeted sequence analysis amplified by using published primers on the basis of *pMGA* and *pvpA* partial nucleotide sequence, respectively. The present study indicates the presence of *pMGA* and *pvpA* gene sequences and size variations among MG field strains from Malaysia. This size variation or gene size polymorphism is controlled by *Mycoplasmas* genetic mechanisms which depend on the combination of appropriate gene subsets encoding surface proteins having short homo- or heteropolymorphic tracts, undergoing intermittent and transformable changes in nucleotide numbers (Meseguer, 2008).

Figure 5. Phylogenetic relationship among MG local strains, reference strains and international strains on the basis of *pMGA* partial nucleotide sequences.

Notes: ● indicates reference strains, ● indicates local field strains and ● indicates international strains.

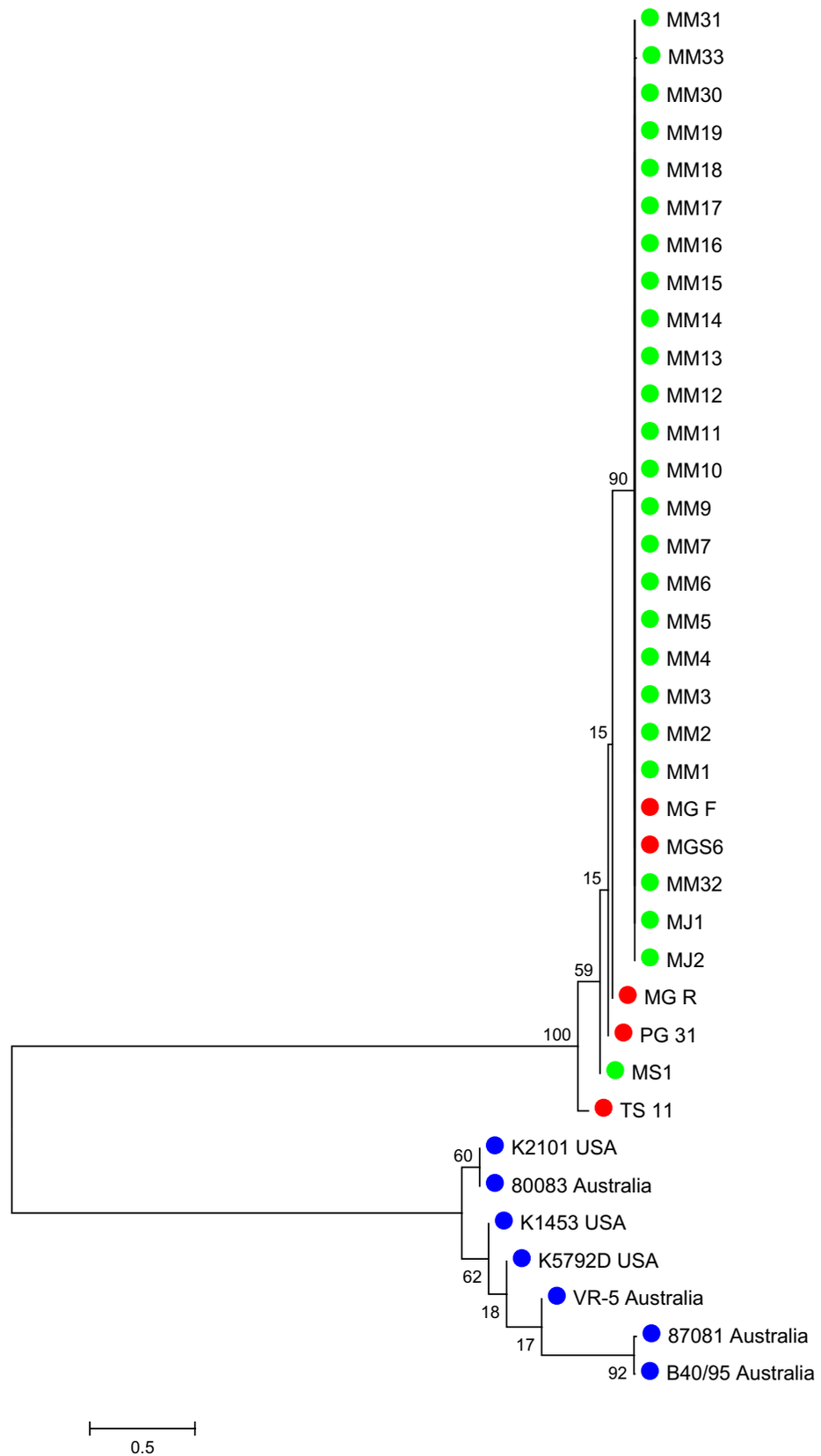
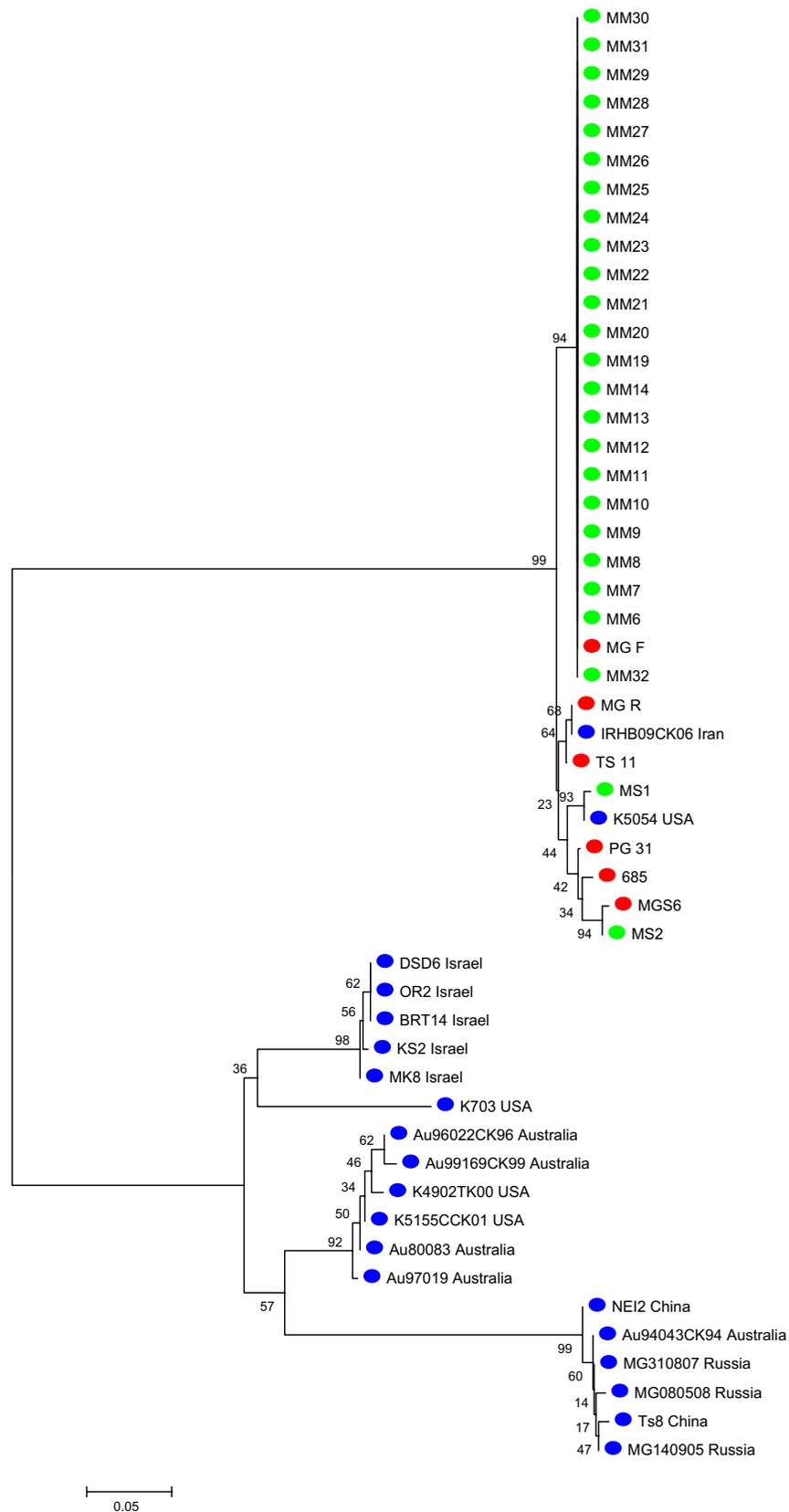


Figure 6. Phylogenetic relationship among MG local strains, reference strains and international strains on the basis of *pvpA* partial nucleotide sequences.

Notes: ● indicates reference strains, ● indicates local field strains and ● indicates international strains.



This finding concurred with other study where MG 6/85 strain was not amplified by AUTS11 F + ATTS11 R primer set of *pMGA* gene (Yasmin et al., 2014). Although these data were different from another study where one field strain from USA was amplified by AUAT TS 11 primer set showed similar gene size pattern with TS 11 and the vaccine strain 6/85 also amplified by this primer set but these two strains were different from each other due to the different gene size pattern (Kleven, 2002). The overall range of PCR product size was 434–677 bp when amplified by *pvpA* 1F + *pvpA* 2R primer set of *pvpA* gene which possessed some similarities with previous studies where *pvpA* gene showed size polymorphisms, with PCR products of 437, 578, 606 and 665 bp as detected among MG reference strains and local strains (Boguslavsky et al., 2000; Liu et al., 2001). Another researcher also showed the largest deletion found was that of a 230-bp fragment in vaccine strain F (Liu et al., 2001). Previous sequence analysis showed that MG reference strain R yield a 665-bp PCR product (Boguslavsky et al., 2000); which indicated similarity to the data found in this study where MG R reference strain yield 668-bp product size after amplification by *pvpA* 1F + *pvpA* 2R primer set of the gene *pvpA*. In this study, a total of 20 out of 25 local field strains showed 434–457-bp product size which is totally different from other study carried out in Malaysia, where it showed that, Malaysian local strains isolated from commercial chickens exhibited 567–611-bp PCR product when amplified by *pvpA* 1F + *pvpA* 2R primer set of *pvpA* gene (Zahraa et al., 2011). However, the findings of this study showed similarity with another study which stated that, Malaysian local strains exhibited gene size polymorphism at 437-bp PCR products (Kartini, 2012).

The partial nucleotide sequence analysis of *pMGA* and *pvpA* gene on MG local strains demonstrated that, there were nucleotide variation, size polymorphism, insertion and deletion, where *pMGA* showed similarities but *pvpA* gene showed dissimilarities with other study carried out in Malaysia, as in this study, in case of *pvpA* gene, most of the field strains showed 434–457-bp PCR product but Zahraa et al. (2011) found 567–611 bp. Nucleotide sequence data also demonstrated that, there were insertion when amplified by *pMGA* partial nucleotide sequence primer and deletion when amplified by *pvpA* partial nucleotide sequence primer set which was similar to the previous studies (Kartini, 2012; Zahraa et al., 2011).

Findings of this study showed similarity with other study which stated that, MG-positive samples were closely related with MG F and MGS6 reference strain and distant from TS 11 vaccine strain (Zahraa et al., 2011). However, another study demonstrated that, MG local strains were distant from the reference strain (Kartini, 2012). There was no similarity found between Malaysian local strains and international strains (Figures 5 and 6). These findings are different from the other study which stated that, MG-positive field samples were distinct from vaccine reference strain (Zahraa et al., 2011). However, Kartini (2012) showed that, MG-positive samples had similarity with MGS6 and had no similarity with MG F and MG R. Most of the international strains showed no similarities with Malaysian local strains.

5. Conclusion

Based on the phylogenetic tree analysis of *pMGA* and *pvpA* partial nucleotide sequences of MG local strains, reference strains and international strains, it can be concluded that Malaysian local strains are different from the international strains. This is an important information in relation to the types of vaccines to be used for the prevention of *Mycoplasma* infection. It would be necessary for further research on other different types of gene (*IGSR*) for characterization of MG field strains in Malaysia. A more efficacious vaccine may also be developed to control MG in Malaysia, based on the identified characteristics of MG field strains.

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Competing interests

The authors declare no competing interests.

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