Two Clusters among *Mycoplasma haemomuris* Strains, Defined by the 16S-23S rRNA Intergenic Transcribed Spacer Sequences

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**ABSTRACT.** *Mycoplasma haemomuris* is a causative organism of infectious anemia or splenomegaly in rodents. Here, we report two distinct genetic groups among *M. haemomuris* strains detected from rats and mice, respectively, by examining the nucleotide sequences of the 16S-23S rRNA intergenic transcribed spacer region that has been shown to be a stable genetic marker for mycoplasma species. Our results may reveal host-tropism of each cluster of *M. haemomuris* strains, and suggest an idea to distinguish *M. haemomuris* into two different genetic clusters.

**KEYWORDS:** hemoplasma, mycoplasma, rRNA

NOTE  
Bacteriology

Hemotropic mycoplasmas also called hemoplasmas are causative of infectious anemia in various mammalian animals [15]. Hemoplasma strains have been isolated as an anemic pathogen from rodents including mice, rats and hamsters and had once been identified by only microscopic observation of blood smears [18]. Hemoplasma infections in laboratory rodents have been concerned to undermine the validity of animal experiments [1, 13]. They are often unrecognized, because of clinically silent infections. Such latent infections have been reported in Sprague-Dawley and Wistar rats [2, 3]. Currently, only one hemoplasma species *Mycoplasma haemomuris* Mayer 1921 formerly *Bartonella muris* or *Haemobartonella muris*, is established in rodents [16, 17]. Nucleotide sequence of the 16S rRNA gene of *M. haemomuris* has been determined on the Shizuoka strain that was the only strain maintained in vivo at that time [19]. Subsequently, nucleotide sequence of the 16S-23S rRNA intergenic transcribed spacer (ITS) region of the same strain was defined [10]. However, genetic variation in the 16S rRNA gene or ITS region remains unexplored, because no other rodent hemoplasma strains except for the Shizuoka strain have been available. Here, we report two genetic clusters in *M. haemomuris* strains by examining nucleotide sequence of ITS region as well as the 16S rRNA gene.

Anti-coagulated blood or spleen homogenates were obtained from black rats with splenomegaly trapped in Okinawa Prefecture, Japan. Detail of these samples examined is given in Table 1. Blood smears were prepared for Giemsa staining. Total DNA was extracted from 200 µl of the whole blood or spleen homogenate by using the QIAamp DNA Blood Mini Kit (QIAGEN, Hilden, Germany) according to the manufacturer’s instructions, eluting into 200 µl of buffer AE, and stored at −20°C until examination in the PCR assay.

Seven DNA samples were subjected to PCR to amplify entire region of the 16S rRNA gene and ITS region. The PCR was carried out with 50-µl reaction mixtures containing 1 µl of DNA solution, 0.8 µl of Tks Gflex™ DNA polymerase (5 units/µl), 25 µl of 2X Gflex PCR Buffer, 0.2 µl of relevant forward and reverse primers and water to a final volume of 50 µl. The forward (5’-AGAGTTT-GATCCTGGCTCAG-3’), equivalent to nucleotide numbers 11 to 30 of *M. wenyonii* (AY946266), or 5’-ATATCCCTAC-GGGAAACGCAGC-3’, equivalent to nucleotide numbers 328 to 347 of *M. wenyonii*, and reverse (5’-ACCGCAGCT-GCTGGCACATA), or 5’-ATACCTTGTTACGACTTAACT-3’, equivalent to nucleotide numbers 1446 to 1465 of *M. wenyonii*) (50 pmol/µl each) primers were used to amplify the 16S rRNA gene. On the other hand, ITS region was amplified by using forward primer Hemo16-23S-F (5’-GTTCGCGAGGTCTTCTTCGACATA-3’) and reverse primer Hemo16-23S-R1 (5’-CAGTACTTGTTCACCTTGTA-3’) as described previously [6]. After initial denaturation at 94°C for 5 min, the reaction cycle was carried out 30 times with denaturation at 98°C for 10 sec, annealing at 55°C for 60 sec and extension at 68°C for 30 sec in a thermal cycler. The PCR products were fractionated on horizontal, submerged 1.0% SeaKem ME agarose gels (FMC Bioproducts, Rockland, ME, U.S.A.) in TAE (40 mM Tris, pH8.0, 5 mM...
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sodium acetate and 1 mM disodium ethylenediaminetetraacetate) buffer at 50 volts for 60 min. After electrophoresis, the gels were stained in ethidium bromide solution (0.4 µg/ml) for 15 min and visualized under UV transilluminator. DNA in a clearly visible band was extracted by using NucleoSpin Extract II kit (Macherey-Nagel, Düren, Germany) and was subjected to direct sequencing in a 3500 Genetic Analyzer (Applied Biosystems, Foster City, CA, U.S.A.).

Almost entire nucleotide sequences of the 16S rRNA gene and ITS region of the seven strains were successfully determined. Nucleotide sequences of the 16S rRNA gene of these samples were almost identical (99% homology) and also showed 99% homology to those of M. haemomuris Shizuka strain (accession number U82963) isolated from a small field mouse in Japan [19]. This allowed us to classify the seven strains as M. haemomuris, though the Shizuka strain used as a reference has been lost and unavailable (Rikihisa, personal communication). Hemoplasma species has provisionally been classified or identified by only nucleotide sequence of the 16S rRNA gene because of uncultivable trait [16, 17].

The nucleotide sequences of ITS of the seven isolates were compared with those of authentic rodent mycoplasma species in a phylogenetic tree that was generated with the neighbor-joining method [20] from a distance matrix corrected for nucleotide substitutions by the Kimura two-parameter model [14]. Phylogenetic analysis indicated that the seven isolates were divided into two clusters A and B (Fig. 1). Nucleotide sequence similarity between these two clusters was 84.9%. This variation can be used for a genetic marker of M. haemomuris strains. Next, we examined primary and secondary structures of the ITS region of the isolates. Nucleotide sequences of ITS region from the seven isolates were compared with six other hemoplasma sequences in an alignment created by
Of the seven strains, ITS sequences of the five strains consisting of S151-2, S152-2–4, S152-5–7 and S159-11–13 were distinct from the three other strains, S154, Ikemajima 5–1 and Ikemajima 14–1 (Fig. 2). ITS sequences of these five strains were identical to those of *M. haemomuris* Shuzuoka strain. ITS region has been used for a comple-

![Fig. 2. Nucleotide variations appeared in alignment of the 13 ITS sequences from different hemoplasma strains. The nucleotide sequence numbers are given from a consensus sequence. Homologous nucleotides are shown as inverted characters. Dashes indicate nucleotide gaps between adjacent nucleotides introduced for the alignment. Ikema Is 5-1, Ikema Is14-1, CM haemobos1 and CM haemobos2 represent Ikemajima 5-1, Ikemajima 14-1 and *Candidatus* *M. haemobos* type 1 and type 2 [22] strains, respectively.](image-url)
mentary genetic marker for identification and classification of mycoplasmas, since this region is characteristic to each *Mycoplasma* species [7–9, 11, 21, 25]. Thus, difference in nucleotide sequence of ITS region may raise a hypothesis that *M. haemomuris* species contains at least two distinct clusters. We also found the ITS region of seven strains lacked spacer tRNA gene, but contained box A and box B motifs previously identified in other mycoplasma species [12]. These motifs are obvious in the sequence alignment of ITS regions of these hemoplasmas.

The secondary structures of the ITS were predicted according to the algorithm of Zuker and Stiegler [26]. Six stem-loop domains were allocated in ITS region of the clusters A and B (Fig. 3). Domains V and VI were common between the clusters. Domain II was well conservative, despite a single nucleotide substitution at loop region. Stem region of domain IV was well conserved between the clusters. Secondary structures in ITS region have sometime provided a key character to distinguish closely related species of mycoplasmas [8, 9, 22]. Thus, in addition to our previous illustration of ITS for *M. haemomuris* [10], the present analysis revealed existence of two genetically distinct clusters among *M. haemomuris* strains.

Collective analyses on the primary and secondary structures of ITS indicated *M. haemomuris* strains were divided into two clusters. This variation may not attribute to a geographical difference of locations where the hemoplasmas isolated, because cluster A included strains from Fukushima, Aomori and Shizuoka Prefectures, and cluster B included Fukushima and Okinawa Prefectures. Thus, this variation is most likely to depend on difference of natural host of hemoplasmas. In fact, cluster A strains have been isolated...
from small field mice, while cluster B was from black rats. *Mycoplasma haemomuris*, that was first observed in the blood of rats and named *Bartonella muris ratti* by Martin Mayer in 1921, was confirmed to be the causative agent of infectious anemia in albino rats following splenectomy [4]. Subsequently, another type of *Bartonella* morphologically distinct from *B. muris ratti* was found in the blood of albino mice by Schilling, and he called this variant as *B. muris musculi* [23]. Taken together, it turns out that the scientific designation, *M. haemomuris*, is composite of *B. muris* subsp. *ratti* in rat and *B. muris* subsp. *musculi* in mouse, despite possible cross-transfer between these animal species by experimental infection. Therefore, this raises a hypothesis that *M. haemomuris* Shizuoka strain isolated from a mouse may correspond to formerly *B. muris musculi*. In our microscopic observation, hemoplasma strains from each cluster appeared as tiny round bodies sometimes in chain within the red blood cells, though some of those from small filed mice might appear as projections from the red blood cell surface (Fig. 4). However, this minor difference may not be sufficient for differentiation of these two clusters on blood smears.

In conclusion, two genetic clusters of *M. haemomuris* were demonstrated by analyzing the primary and secondary structures of ITS region of *M. haemomuris* strains. Besides, our findings support the hypothesis that the cluster including *M. haemomuris* Shizuoka strain represents *M. haemomuris* subsp. *musculi*, and the other cluster corresponds to *M. haemomuris* subsp. *ratti*. This may provide a clue to elucidate differences in severity of anemia in rodent, since virulence of these two clusters is currently unknown. This variation can also be used for a genetic marker for monitoring of *M. haemomuris* infections in laboratory rodents. The GenBank/EMBL/DDBJ accession numbers appeared for the first time in this study are AB758434 through AB758440.

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REFERENCES

plasma haemomuris, previously classified as ‘Haemobartonella muris’. J. Vet. Med. Sci. 64: 1161–1164. [Medline] [CrossRef]


