Spiroplasma velocicrescens sp. nov., from the Vespid Wasp Monobia quadridens

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Spiroplasma strain MQ-4T (T = type strain), which was isolated from the hemolymph of the vespid wasp Monobia quadridens, was serologically distinct from other spiroplasma species, groups, putative groups, and subgroups. Each strain MQ-4T cell was helical and motile and was surrounded by a single cytoplasmic membrane; there was no evidence of a cell wall. The strain grew well in 1% serum fraction medium, as well as in SM-1, M1D, and SP-4 liquid media, under both aerobic and anaerobic conditions. Strain MQ-4T grew at temperatures ranging from 10 to 41°C but did not grow at 43°C. The strain grew optimally at 37°C with a doubling time of 0.6 h, the shortest doubling time recorded for any spiroplasma. Strain MQ-4T catabolized glucose and arginine but did not hydrolyze urea. The guanine-plus-cytosine content of the DNA was about 27.5 ± 1 mol%. The genome size was 1,480 kbp (940 MDa). Strain MQ-4 (= ATCC 35262) is designated the type strain of a new species, Spiroplasma velocicrescens.

The genus Spiroplasma (31), which originally was considered a taxon that contains insect-borne pathogens of plant phloem, was later shown by Clark and his associates (6, 7) to be associated primarily with insects. As the true affinities of members of this genus were discovered, unique features that characterize each new group emerged. However, for some time, it appeared that the group XI spiroplasmas, represented by strain MQ-4T (T = type strain), which was isolated from the gut of the predaceous wasp Monobia quadridens, might have no unusual properties. This host had yielded several other spiroplasma taxa (6, 12), including the group VII organism Spiroplasma monobiae MQ-I (28), and it was felt that the spiroplasmas could have been acquired from prey insects (12). However, Konai et al. (14) have recently shown that strain MQ-4T is the fastest growing spiroplasma that has been assigned to a group.

In this paper we present the results of a taxonomic study of strain MQ-4T, in accordance with the proposed (13) minimal requirements for assignment of binominal names to mollicutes. We conclude that strain MQ-4, the type strain of group XI, represents a new species, Spiroplasma velocicrescens.

MATERIALS AND METHODS

Spiroplasma strains. Clark originally isolated strain MQ-4T by macerating the gut contents of vespid wasps (M. quadridens) collected in Maryland (6). Some of the genomic and serologic characteristics of this organism have been reported previously (3, 23, 27, 31). Strain MQ-4T was purified by classical filtration-cloning techniques (21). Representative strains of the 23 recognized groups and eight subgroups (11, 23), including the type strains of the 15 previously recognized species and seven new strains, each representing a putative new group (28, 29), were used in various parts of this study.

Culture media and cultivation techniques. Strain MQ-4T was selected from a population of helical cells growing in a primary culture in SM-1 liquid medium (25) at 30°C. After several early broth passages, the isolate was hypervirulent. Later, the dried culture was revived and passed twice in SM-1 broth at 30°C before triple cloning. After the last cloning, a strain derived from a single colony was designated MQ-4T and was used in characterization studies. The other culture media used included M1D and SP-4 media and serum fraction broth supplemented with 1% bovine serum fraction. A solid formulation of each of these media was prepared by adding Noble agar (Difco Laboratories, Detroit, Mich.) to a final concentration of 0.8%. Agar cultures were incubated at 30°C, either aerobically with 5% carbon dioxide (GasPak system; BBL Microbiology Systems, Cockeysville, Md.) or anaerobically (hydrogen GasPak system).

Temperature requirements. Temperature requirements for growth were assessed by preparing 10-fold dilutions of strain MQ-4T in M1D broth. One series of dilutions was incubated at each of 10 temperatures (5, 10, 15, 20, 25, 30, 32, 37, 41, and 45°C). Doubling times were computed by the method of Konai et al. (14).

Morphological and ultrastructural studies. Cells of strain MQ-4T in liquid culture in the early logarithmic phase were observed by dark-field microscopy at a magnification of ×1,250. For electron microscopy, the strain was grown in approximately 20 ml of broth and pelleted by centrifugation. The pellet was fixed for 2 h in 3% glutaraldehyde, postfixed in 1% osmium tetroxide for 1 h, dehydrated in acetone, embedded in Epon, sectioned, and stained with 1% aqueous uranyl acetate and Reynolds’s lead citrate (30).

Sterol requirement. The sterol requirement for growth of spiroplasma strain MQ-4T was demonstrated by the direct broth method (22) and a recently described dilution method (18). For the direct broth test we used one 100-ml bottle of regular serum-containing medium as a positive control, three bottles of serum-free medium for reagent controls, and four bottles of serum-free medium containing different amounts of cholesterol. All of the bottles were inoculated with a log-phase culture of strain MQ-4T and were incubated at 30°C until the media acidified. The cells were harvested by centrifugation at 15,000 rpm for 20 min, washed in phosphate-buffered saline (pH 7.5), and recentered, and the cell pellets were assayed to determine their protein contents by using the Bio-Rad DC protein assay (Bio-Rad Laboratories, Richmond, Calif.). The total amount of protein was used as an indication of spiroplasma growth in the various media.

Tests to determine biological and biochemical properties. The procedures used to study carbohydrate fermentation (1, 17), arginine catabolism (2), urea hydrolysis (16), hemadsorption (10), and film and spot production (9) have been described previously. Filtration characteristics were determined by serially filtering log-phase cultures in M1D broth, using techniques described previously (21).

Serological tests. Antiserum to strain MQ-4T was raised in rabbits as previously described (32). Hyperimmune antiserum to all previously described Spiroplasma species, groups, putative groups, and subgroups were obtained from the reference collections at the Beltsville Agricultural Research Center and the Mycoplasma Section of the National Institute of Allergy and Infectious Diseases laboratory in Frederick, Md. These antisera and strain MQ-4T were tested with standard disc gel growth inhibition tests (8, 27) by using M1D agar plates incubated aerobically at 30°C. For metabolism inhibition (20, 32) and deformation tests we also used previously described protocols (32).

Genomic analysis. One liter log-phase cultures of strain MQ-4T in M1D medium were harvested by centrifugation at 20,000 × g for 30 min. The pellets were
RESULTS AND DISCUSSION

Cultural and morphological properties. Strain MQ-4T grew well in liquid SM-1, M1D, and SP-4 media and on solid media prepared from these formulations. The strain also grew in conventional mycoplasma media containing horse serum (Edward formulation) or bovine serum fraction. Growth occurred in phosphate-buffered saline (pH 7.5), recentrifuged, and denatured with sodium dodecyl sulfate. Chromosomal DNA was purified further as described by Carle et al. (4). The guanine-plus-cytosine (G+C) content of purified strain MQ-4T DNA was determined by buoyant density, melting temperature, and high-performance liquid chromatography methods (5). Purified DNA from Spiroplasma cito R8A2 (G+C content, 26 ± 1 mol%) was used as a reference DNA in all procedures. The genome size was determined (3) by performing pulsed-field gel electrophoresis with a logarithmic-phase whole-cell culture of strain MQ-4T embedded in low-melting-point SeaPlaque agarose blocks. S. citri R8A2 processed and embedded in a similar way was used as a standard (genome size, approximately 1,820 kbp [1,156 MDa]).

Growth of strain MQ-4T on solid medium was observed on agar media under anaerobic conditions. Logarithmic-phase cultures of strain MQ-4T in GAM systems with carbon dioxide atmosphere and sodium dodecyl sulfate. Chromosomal DNA was purified further as described by Carle et al. (4). The guanine-plus-cytosine (G+C) content of purified strain MQ-4T DNA was determined by buoyant density, melting temperature, and high-performance liquid chromatography methods (5). Purified DNA from Spiroplasma cito R8A2 (G+C content, 26 ± 1 mol%) was used as a reference DNA in all procedures. The genome size was determined (3) by performing pulsed-field gel electrophoresis with a logarithmic-phase whole-cell culture of strain MQ-4T embedded in low-melting-point SeaPlaque agarose blocks. S. citri R8A2 processed and embedded in a similar way was used as a standard (genome size, approximately 1,820 kbp [1,156 MDa]).

Biochemical and biological properties. Strain MQ-4T produced acid from glucose and hydrolyzed arginine, but no evidence of urea hydrolysis was observed. Strain MQ-4T was positive for film and spot production, but colonies of the organism on an agar medium did not hemadsorb guinea pig erythrocytes. Passage of broth cultures of strain MQ-4T through 450- or 300-nm-pore-size membrane filters did not reduce the viable cell titer (10⁸ color-changing units/ml). The titer of a broth culture filtrate obtained after passage through a 220-nm-pore-size membrane was about 10-fold lower (10⁷ color-changing units/ml); a 100-nm-pore-size membrane filtrate did not contain viable cells.

Serological tests. The results of growth inhibition, metabolism inhibition, and spiroplasma deformation tests indicated that strain MQ-4T was not related serologically to representatives of previously described Spiroplasma groups or species or to seven ungrouped representatives of putative new groups (29).

Genome size and DNA base composition. The base compositions (G+C contents) of strain MQ-4T DNA were 27.5, 27.3, and 27.9 mol% as determined by the buoyant density, melting temperature, and high-pressure liquid chromatography techniques, respectively. The genome size was 1,480 kbp (940 MDa).

Habitat. The single strain described in this paper was isolated directly from the gut of the vespid wasp M. quadridens. Most mollicutes cultivated from insects have been isolated from guts (6, 12). Residence in the gut of a predaceous insect could simply reflect acquisition from prey. Some mollicutes that reside in insect hemolymph, such as Spiroplasma melliferum (7) and Spiroplasma apis (15), reduce the longevity of the host. Strain MQ-4T, like other insect gut spiroplasmas, is not known to be pathogenic to its insect host.

Diagnosis and significance. Like other Spiroplasma species (24, 31), S. velocecrecescens can be identified by serological tests. In metabolism inhibition and deformation tests, it exhibits, at most, minor one-way crosses with sera to all previously described Spiroplasma species, groups, subgroups, and putative groups (29). There is no known close relative; the group VII organism strain MQ-1 (S. monoblea), which was isolated from the same host, has a G+C content of 28 ± 1 mol% and a genome size of 657 MDa, values that differ from the strain MQ-4T values. The rapid growth of strain MQ-4T and its optimum temperature (37°C) are by no means unique. Unrelated strain HYOS-1 (29) grows almost as fast (doubling time at the optimum temperature [32°C], 0.7 h) and, strains of Spiroplasma floricola, S. apis, Spiroplasma chinense, and Spiroplasma culicicola and representatives of groups VIII and XII also grow very rapidly (14). The optimum temperature for 14 strains, representing nine groups of spiroplasmas, is 37°C (14).

Taxonomic placement. The properties described above for strain MQ-4T fulfill proposed criteria for species belonging to the class Mollicutes, including absence of a cell wall, filterability, and penicillin resistance. The growth requirement for sterol or serum, the inability to utilize urea, and the helicity and motility of strain MQ-4T place this organism in the family Spiroplasmataceae (19) and the genus Spiroplasma. Finally, a serologic comparison of strain MQ-4T with other Spiroplasma species and with other unclassified spiroplasma strains that were observed when various concentrations (1 to 17%) of fetal bovine serum were added to SP-4 base broth. Strain MQ-4T did not exhibit sustained growth when serial subcultures were made in serum-free broth or when 0.04% Tween 80 was added to this formulation (18). However, strain MQ-4T could be passed continuously in serum-containing medium, indicating that sterol was required for growth.
FIG. 2. Electron micrograph of a sectioned and stained cell pellet of strain MQ-4T. Sections were stained with 2% aqueous uranyl acetate and Reynolds's lead citrate. The arrows indicate the unit membrane.

represent putative species revealed the uniqueness of the new insect strain. Therefore, we propose the name Spiroplasma velocicrescens for this organism.

**Description of Spiroplasma velocicrescens sp. nov.** *Spiroplasma velocicrescens* (ve.lo.ci.cres'cens. L. adj. veloci, fast; L. part. crescens, growing; M. L. n. part. velocicrescens, fast growing). Cells are helical, motile filaments, vary from 200 to 300 nm in diameter, and lack cell walls. Colonies on solid medium containing 0.8% Noble agar are diffuse and never have a fried-egg appearance.

Chemoorganotroph. Acid is produced from glucose. Hydrolyzes arginine. Does not hydrolyze urea.

Film and spot reaction negative. Does not hemadsorb guinea pig erythrocytes.

Cholesterol or serum is required for growth.

The temperature range for growth is 10 to 41°C; the doubling time at 37°C, the optimum temperature, is 0.6 h.

Serologically distinct from previously described *Spiroplasma* species. Isolated from the gut of the vespid wasp *M. quadridens*. Pathogenicity for insects has not been determined.

The G+C content of the DNA is 27.5 ± 1 mol%. The genome size is 1,480 kbp (940 MDa).

The type strain is MQ-4 (= ATCC 35262).

**TABLE 1.** Growth responses of strain MQ-4T in the presence of different concentrations of cholesterol

<table>
<thead>
<tr>
<th>Medium</th>
<th>Cholesterol conc. (μg/ml)</th>
<th>Amt of protein (mg/100 ml)</th>
</tr>
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<tbody>
<tr>
<td>17% fetal bovine serum in SP-4 broth</td>
<td>0</td>
<td>2.16</td>
</tr>
<tr>
<td>Serum-free broth</td>
<td>0</td>
<td>I.G*</td>
</tr>
<tr>
<td>Serum-free broth containing palmitic acid (10 μg/ml) and albumin (1%)</td>
<td>0</td>
<td>0.42</td>
</tr>
<tr>
<td>Serum-free broth containing palmitic acid (10 μg/ml), albumin (1%), and Tween 80 (0.01%)</td>
<td>0</td>
<td>0.17</td>
</tr>
<tr>
<td>1.0</td>
<td>0.76</td>
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<td>5.0</td>
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<td>10.0</td>
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</tr>
<tr>
<td>20.0</td>
<td>1.80</td>
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* I.G, insufficient growth for detection.

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REFERENCES


