Mycoplasma feliaucium, a New Species Isolated from the Respiratory Tract of Pumas

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Mycoplasmas isolated from the throats of three pumas (Felis concolor) were each cloned and examined in detail. All three were serologically and biologically indistinguishable from each other, and were serologically distinct from 83 recognized Mycoplasma and Acholeplasma species. They were designated as a new species, Mycoplasma feliaucium, with strain PU (NCTC 11703) as the type strain.

INTRODUCTION

Mycoplasma spp. have been isolated from a wide variety of animals, birds and plants. Most of the wild animals have not been examined for their mycoplasmal flora, but two reports have been published on mycoplasma isolation from captive large felines, including lions, lynx, pumas and tigers (Heyward et al., 1969; Hill, 1975). The tiger isolates and some of those recovered from lions and pumas were typed as Mycoplasma arginini (Tully et al., 1972; Hill, 1975). The other strains from lions have not been fully characterized but are serologically distinct from the puma isolates.

The puma strains cultivated by Hill (1975) had been isolated during a survey carried out to investigate the mycoplasmal flora of large felines, by examining the upper respiratory tract of pumas, when they were anaesthetized for surgical or other procedures. This paper is concerned with their further characterization.

METHODS

Mycoplasma strains. Mycoplasmas were isolated from the throats of three pumas (Felis concolor) from the Zoological Society of London, Regents Park, London, UK (Hill, 1975). Two distinct strains were isolated from one animal (one strain was identified as M. arginini) and one strain isolated from each of the other two pumas. The mycoplasma from each puma was cloned to produce a pure culture, by an initial filtration of a broth culture through a 220 nm membrane filter, culturing of the filtrate on solid medium, transfer of a single resulting colony to a further agar plate, the subsequent growth being inoculated into broth and the whole procedure repeated a further three times, thereby cloning four times in total (Subcommittee on the Taxonomy of Mycoplasmatales, 1972). One such cloned strain was designated PU and the other two, PU1 and PU2.

Mycoplasma species. Mycoplasma type strains of 83 Mycoplasma and Acholeplasma spp. (Table 1) were obtained from the National Collection of Type Cultures, Colindale, UK and Drs H. Atobe (University of Tokyo, Japan), M. F. Barile (National Institutes of Health, Bethesda, Md., USA), J. M. Bradbury (Liverpool University, UK), R. J. Fallon (Ruchill Hospital, Glasgow, UK), E. A. Freundt (University of Aarhus, Aarhus, Denmark), J. T. Heywood (National Communicable Disease Center, Atlanta, Ga., USA), F. T. W. Jordan (Liverpool University, UK), D. E. Jasper (University of California, Davis, Calif., USA), H. Kirchoff (Tierarztliche Hochschule, Hannover, FRG), R. H. Leach (PHLS Mycoplasma Reference Laboratory, Norwich, UK), G. Smith (Zoological Society of London, London, UK), D. Taylor-Robinson (Clinical Research Centre, Harrow, UK) and J. G. Tully (National Institutes of Health, Bethesda, Md., USA).

Medium and growth conditions. The basal culture media were those described by Taylor-Robinson et al. (1968) and Hill (1971). Liquid media contained either 1% (w/v) glucose (pH 7.8) or 1% (w/v) arginine (pH 7.3) depending...
Table 1. **Serological tests comparing strain PU with other, named species**

<table>
<thead>
<tr>
<th>Antigen</th>
<th>PU</th>
<th>Metabolism inhibition (titre)</th>
<th>Immunoperoxidase (titre)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Antiserum to:</td>
<td>Zone width (mm)</td>
<td>Antiserum to:</td>
</tr>
<tr>
<td>PU</td>
<td>M. muris</td>
<td>3-4</td>
<td>&lt;32</td>
</tr>
<tr>
<td>M. muris (RII14)</td>
<td>0</td>
<td>&lt;32</td>
<td>100</td>
</tr>
<tr>
<td>All other strains*</td>
<td>0</td>
<td>&lt;32</td>
<td>&lt;60</td>
</tr>
</tbody>
</table>

* M. agalactiae (PG2), M. alkalescens (PG51), M. altus (Illyes), M. anatis (1340), M. arginini (G230), M. arthritidis (PG6), M. bovigenitalium (PG11), M. bovirhinis (PG43), M. bovis (PG45), M. bovoculi (M165/69), M. buccale (CH-20247), M. californiae (S7-6), M. canadense (275C), M. canis (PG14), M. capricolum (California kid), M. caseae (G122), M. caseipharyngis (117C), M. citelli (RG-2C), M. cloacae (383), M. Collins (588), M. columbiculare (694), M. coprotherium (MMP1), M. columborale (MMP4), M. conjunctivae (HRC581), M. cricetuli (CH), M. cyon (H831), M. dispar (462/2), M. edwardii (PG24), M. equigenitalium (T37), M. equihinis (M432/72), M. fastidiosum (4822), M. fauclum (DC333), M. feliniunctum (Ben), M. felis (CO), M. fermentans (PG18), M. flocculare (MS42), M. gallinae (DO), M. gallinarum (PG16), M. gallisepticum (PG31), M. gallopavonis (WR1), M. gatae (CS), M. genitalium (G-37), M. glycolphilum (486), M. hominis (PG21), M. hypovacuolima (J), M. hyorhinis (BST7), M. hyosynoviae (S16), M. iners (PG30), M. iowae (695), M. lipofaciens (R171), M. lipophilum (MaBY), M. maculosum (PG15), M. meleagris (17529), M. mośski (MK405), M. molare (HS42), M. mustelae (MX9), M. mycoides subsp. capri (PG3), M. mycoides subsp. mycoides (PG1), M. neurolyticum (Type A), M. opalecens (M5408), M. orale (CH-19299), M. ovine (Y89), M. pneumoniae (FH), M. primatum (HRC292), M. pullorum (CKK), M. pulmonis (G34ASH), M. purificans (KS-1), M. salinarum (PG20), M. spumans (PG13), M. suavi (Mayfield strain (Clone B)), M. subdolium (1B), M. synoviae (WVU 1853), M. verecundum (107), A. axanther (S743), M. equifelata (N93), A. granularum (BTS39), A. hippikoni (C1), A. laiti (PG8), A. moditicum (PG49), A. morum (72-043), A. ovatum (19L), A. parvum (H23M).

† R, Reduced numbers.

upon the biochemical activity of the mycoplasmas cultivated. Agar cultures were incubated at 35–37 °C either in a humid chamber or under anaerobic conditions in a Gaspak system. Liquid cultures were stored at −70 °C in ampoules.

**Growth requirements and characteristics.** The cloned test strains were subcultured onto solid medium and grown for 1 week under both aerobic and Gaspak anaerobic conditions at 35–37 °C. Their sensitivity to methylene blue was investigated by adding 0.002% (w/v) methylene blue to the basal solid medium and comparing the growth of inoculated mycoplasmas with those cultivated on medium without methylene blue. Lipolytic activity was tested by inoculating the mycoplasmas onto basal medium enriched with 10% (v/v) egg yolk emulsion (Fabricant & Freundt, 1967). The incubated plates were then examined at 3, 7 and 14 d for visual evidence of lipolysis (clearing) or 'film' production.

**Absence of reversion.** The three test strains were each subcultured, by five passages on both solid and liquid media that contained no microbial inhibitors, in order to determine whether they would revert to a bacterial form. In addition, the colonies were treated with Dienes' stain, which differentiates mycoplasma colonies from bacterial L-forms (Timms, 1967).

**Morphological studies.** Mycoplasma colonies grown on agar were examined microscopically at ×100 magnification after 2, 7 and 14 d incubation. Some colonies were transferred to slides and stained with Giemsa stain (Fallon & Whittlestone, 1969). Liquid cultures were observed under dark-field microscopy and the organisms were also examined after staining by the Giemsa method (Fallon & Whittlestone, 1969). Organisms grown in broth were harvested by centrifugation and the resulting pellet was fixed at room temperature in 2% (v/v) glutaraldehyde for 2 h and post-fixed in 1% (w/v) osmium tetroxide for 1 h. After embedding, thin sections were cut and stained with uranyl acetate and lead citrate (Venable & Coggshall, 1965) and examined by electron microscopy.

**DNA base composition.** DNA was extracted from centrifuged broth culture deposits by the method of Kirby (1959), and the G + C content determined from its thermal denaturation temperature (Marmur & Doty, 1962). DNA extracted from Clostridium perfringens and Mycoplasma gallisepticum, which have known G + C contents, were included as controls.

**Filtration studies.** Cultures (24 h incubation) were diluted 1/10 in liquid medium, and then separate volumes of the diluted culture were filtered through membrane filters (Millipore) with pore diameters of 100, 220, 300, 450, 650 and 800 nm. The viable count (c.f.u. ml⁻¹) for each filtrate was determined and compared with that of the unfiltered culture dilution.
Mycoplasma felifaucium, a new sp. from pumas

Sterol dependence. Plates were prepared from serum-free medium supplemented with 0.5% bovine serum albumin (BSA), 0.5% glucose and 10 μg palmitic acid ml⁻¹ (Razin & Tully, 1970), to which cholesterol, dissolved in Tween 80, was added to give concentrations of 20, 10, 5 and 1 μg ml⁻¹. The albumin/glucose/palmitic acid plates without cholesterol, but with and without 20% (v/v) horse serum, were also included in the tests. Single mycoplasma colonies taken from a serum-agar plate were subcultured on the test plates and examined for growth during 14 d incubation (35 °C). Where growth occurred on any of the serum-free media, the single colonies were passaged similarly three successive times on the same medium (and also on serum-free medium without any supplements) to ensure that growth had not been due to a carry-over of serum from the original culture, giving misleading results (Razin & Tully, 1970; Edward, 1971).

The isolates were tested indirectly for sterol dependence by a paper disc inhibition method (Freundt et al., 1973), using either dried discs originally containing 0.02 ml of a 1:5% (v/v) ethanolic solution of digitonin (Sigma) or wet discs containing 0.02 ml of a 20% (w/v) aqueous solution of sodium polyanetholsulphonate (Koch-Light). The width of the zone of growth inhibition was recorded.

Biochemical activity. The three cloned strains were examined for: carbohydrate metabolism; hydrolysis of aesculin, arginine and urea (Williams & Wittler, 1971; Leach, 1976); reduction of methylene blue, resazurin, tetrazolium and tellurite; and phosphatase activity (Tully, 1965; Aluotto et al., 1970; Barber & Fabricant, 1971).

Erythrocyte techniques. The test strains were examined for haemolytic activity (Aluotto et al., 1970), haemadsorption and haemagglutination (Manchee & Taylor-Robinson, 1968) with fowl, guinea pig, human and sheep erythrocytes.

Polyacrylamide gel electrophoresis. This was done as described by Razin & Rotten (1967). Strains PU, PU1 and PU2 were compared with M. arginini, which has also been isolated from pumas (Hill, 1975). Centrifuged cell suspensions of the mycoplasmas were dried and 10 mg of each suspension was dissolved in 1 ml phenol/acetic acid/water (2:1:0.5, by vol.). The gels were loaded with 0.025 ml of the cell sample. After electrophoresis at room temperature for 3.5 h at a constant current of 5 mA per tube, the gels were stained with 1% (w/v) naphthol blue black.

Serological studies. Antisera to mycoplasmas were prepared in rabbits as described by Morton & Roberts (1967) and Hill (1971). The following three serological methods were used, each test being done in duplicate: growth inhibition tests with antiserum-impregnated sterile paper discs (Clyde, 1964), metabolism inhibition tests in microtitre plates (Purcell et al., 1966a, b; Taylor-Robinson et al., 1966) and immunoperoxidase tests on colonies grown on agar (Polak-Vogelzang et al., 1978).

RESULTS AND DISCUSSION

Colonies of strains PU, PU1 and PU2 were visible microscopically in subcultures after 2 d of incubation under both aerobic and anaerobic conditions. Primary isolates took 3–5 d to produce visible colonies. Individual colonies (Fig. 1) had a typical ‘fried egg’ appearance and growth was accompanied by a ‘film’ after prolonged incubation. Each strain produced a ‘film’ and clearing when cultured on egg yolk agar but the strains could not be cultured on agar containing methylene blue. Growth of the mycoplasmas was not retarded by the presence of penicillin in the medium. None of the isolates reverted to a bacterial form when serially subcultured on medium without bacterial inhibitors. Their colonies rapidly became stained with Dienes’ reagent, tending to confirm these isolates as true mycoplasmas (Boatman, 1979). The ultrastructure of the organisms in thin sections also proved typical of mycoplasma morphology (Boatman, 1979), the cells having a characteristic internal structure and being bounded by a single membrane (Fig. 2).

The DNA from strain PU was denatured at a mid-point temperature of 82.1 to 82.3 °C (corrected for thermal expansion) over several tests. The G + C content of 31 mol% was determined by the equation

\[ T_m = 69.3 + 0.41 (G + C), \]

where \( T_m \) is the thermal denaturation temperature (Marmur & Doty, 1962). This value is within the known range for the Mollicutes. The G + C contents of the controls, C. perfringens and M. gallisepticum, were the same as previously described (Marmur & Doty, 1962).

Filtration of the liquid cultures of each test mycoplasma (4 x 10⁸ c.f.u. ml⁻¹) showed that each passed in reduced numbers through membranes with 300 nm pore size (2 x 10⁷ c.f.u. ml⁻¹) and 220 nm pore size (6 x 10³ c.f.u. ml⁻¹). No organisms were detected after passage through the 100 nm membrane.
The three strains each required cholesterol for growth. They could not be cultured on the serum-free medium supplemented with BSA, glucose and palmitic acid without cholesterol, or with cholesterol at 1 µg ml⁻¹, but grew and were passaged on this medium when it contained 5 µg cholesterol ml⁻¹. In conformity with their sterol-dependence, they were susceptible to
digitonin and sodium polyanetholsulphonate in the disc tests, with large zones of growth inhibition (≥ 5 mm) produced by both reagents where growth was otherwise semi-confluent.

Each strain showed identical biochemical activity. Arginine was hydrolysed but neither urea nor any carbohydrates were metabolized. Resazurin, and tellurite under anaerobic but not aerobic conditions, were reduced and phosphatase activity was demonstrated. Aesculin was not hydrolysed and nor were methylene blue or tetracezolium reduced. Casein was not digested. In appropriate tests, all three strains were haemolytic for all four types of erythrocyte tested, haemadsorbing for sheep and human cells and haemagglutinating for sheep cells only.

The three cloned strains gave similar electrophoretic protein patterns and these were distinct from that of M. arginini, the only identified Mycoplasma sp. yet recovered from pumas.

Each of the serological techniques showed identical levels of cross-reaction between the three cloned puma isolates. The type strain PU was reacted with antiserum prepared against the species listed in Table 1. Antiserum to PU was tested with the named Mycoplasma and Acholeplasma species. No significant cross-reactions were detected, except for a slight one-way cross reaction with M. muris (Table 1) in growth inhibition tests, but this was not confirmed by the other two methods.

*M. pirum* was reported too recently to be included in the tests, but differs from strains PU, PU1 and PU2 in its inability to grow under aerobic conditions, in its biochemical characteristics (glucose metabolism and lack of phosphatase production), and in its lower (25-5) G + C mol %. (*A. florum* was also omitted, having not yet been obtained by the author, but since it is a sterol-independent Acholeplasma sp. it is clearly taxonomically distinct from the sterol dependent puma isolates.)

Strains PU, PU1 and PU2 belong to the *Mollicutes* because of their main properties, including absence of cell walls, filterability, lack of reversion to bacterial forms when grown in antibiotic-free media, penicillin resistance and production of typical morphology on agar. They belong to the genus *Mycoplasma*, because they are not strict anaerobes (cf. *Anaeroplasma*), are non-helical (cf. *Spiroplasma*), depend upon sterol for growth (cf. *Acholeplasma*), and because urease could not be demonstrated (cf. *Ureaplasma*).

Strains PU, PU1 and PU2 belong to the same species, as they had identical biological characteristics and showed serological and protein-pattern identity. Strain PU gave no cross-reaction with any existing named species, except for a minimal one of doubtful significance with *M. muris*, which in any case has some quite distinct biological characteristics. This strain, therefore, clearly represents a new *Mycoplasma* species, which also contains strains PU1 and PU2. I propose that this species be named *Mycoplasma feliafaucium* and that the type strain is PU (NCTC 11703). *M. feliafaucium* may be a common inhabitant of the upper respiratory tract of pumas, as it was recovered from each of three animals examined. There is no evidence as to its pathogenicity, but the infected pumas showed no signs of respiratory distress.

*Main characteristics of Mycoplasma feliafaucium*

feliafaucium. feles (or felis), L. n. cat; fauces, pl. L. n. throat; feliafaucium, of the feline throat.

*Habitat.* Throat of a puma.

*Morphology, physical and colony characteristics.* Pleomorphic cells bounded by a single unit membrane. Colonies on agar typical ‘fried egg’ appearance. Filterable through 220 nm membrane. Resistant to penicillin.

*DNA base composition.* G + C content 31 mol %.

*Growth characteristics.* Aerobic and anaerobic growth. Requires sterol for growth; growth is inhibited by digitonin and sodium polyanetholsulphonate (SPS).

*Optimal temperature* 35–37 °C.

*Metabolic characteristics.* Arginine hydrolysed. No metabolism of carbohydrates. Phosphatase produced and tetracezolium reduction negative. ‘Film’ positive.

*Serological characteristics.* Distinct from all other *Mycoplasma* species.

*Type strain.* PU (NCTC 11703).

I would like to thank J. Bird and C. Colhoun for the electron micrographs.
REFERENCES


