**Dermatophilus chelonae** sp. nov., Isolated from Chelonids in Australia

A. M. MASTERS,* T. M. ELLIS, J. M. CARSON, S. S. SUTHERLAND, AND A. R. GREGORY

Animal Health Laboratories, Department of Agriculture, South Perth, Western Australia 6151, Australia

Three isolates of a previously undescribed *Dermatophilus* sp. obtained from chelonids (two strains obtained from turtles and one strain obtained from a tortoise) were compared with 30 *Dermatophilus congolensis* isolates obtained from Australian mammals. The microscopic appearance, the colony morphology, and most biochemical test results for the chelonid isolates were characteristic of the genus *Dermatophilus*. Our isolates differed from the mammalian *D. congolensis* isolates in a number of cultural characteristics, including faster growth at 27°C than at 37°C, formation of two hemolysis zones around colonies on blood agar at 37°C in the presence of 10% CO₂, poor motility, and production of a distinctive odor. The DNA restriction enzyme digestion and protein electrophoresis patterns of our strains were distinct. The electrophoretic mobilities of 11 enzymes differed from the mobilities observed with *D. congolensis* strains. A monoclonal antibody to a surface antigen of an ovine isolate did not react with zoospores or filaments of the chelonid isolates. Biochemical differences between our isolates and *D. congolensis* included the ability of the chelonid isolates to reduce nitrate to nitrite and the fact that the chelonid isolates exhibit collagenase activity in vitro. We propose that the chelonid isolates should be placed in a new species, *Dermatophilus chelonae*. Strain W16, which was isolated from a nose scab on a snapping turtle, is the type strain; a culture of this strain has been deposited in the American Type Culture Collection as strain ATCC 51576.

A skin disease caused by the actinomycete *Dermatophilus congolensis* affects a wide variety of animal species and occurs throughout the world. This disease is economically important in cattle in tropical regions and in sheep in high-rainfall areas. Dermatophilosis lesions have also been found in goats, horses, deer, gazelles, giraffes, foxes, chamois, Columbian ground squirrels, cottontail rabbits, owls, polar bears, seals, pigs, and humans (13). To date, there apparently have been no reports of isolations of *D. congolensis* from chelonids, although the organism has been isolated from lizards (2, 23, 30). It has been reported that there is considerable variation in the colony morphology and cultural characteristics of *D. congolensis* (12). Significant variations in serological, biochemical, and virulence characteristics within the species have also been documented (1, 8, 10, 12, 16, 18, 19, 21, 33). A study of the variations in the cultural, biochemical, and morphological properties of Australian mammalian isolates of *D. congolensis* was recently undertaken (9). In our study we compared the properties of three isolates obtained from chelonids with the properties of these mammalian *D. congolensis* isolates.

**MATERIALS AND METHODS**

**Bacterial isolates.** *Dermatophilus* sp. isolates were obtained from an abscess in a tortoise in 1980 (isolate W8), from skin lesions on a turtle in 1982 (isolate W11), and from a nose scab on a snapping turtle in 1987 (isolate W16) (T = type strain); all of these isolates were obtained at the Perth Zoo, in Perth, Western Australia. The isolates which we used for restriction enzyme analysis of bacterial DNAs included all of the isolates and reference strains listed by Trott et al. (34).

**Cultural characteristics.** The isolates were cultured on 9% bovine blood agar (in Oxoid Columbia agar base no. 2) and were observed after 2 to 3 days of incubation at 37°C in the presence of 10% CO₂ and at 27°C in the ambient atmosphere. The appearance of cultures in tryptone-phosphate–10% ovine serum broth inoculated with a dense suspension of zoospores was recorded after 3 days of incubation at 37°C and after additional incubation at room tempera-

ture. The volume of slime produced after 3 days of incubation at 37°C in thioglycolate broth (Oxoid) was determined and was expressed as a percentage of the total volume of the culture after centrifugation at 1,500 × g for 10 min. The annular radii of the hemolysis zones around streaks and colonies of the isolates were measured and compared after duplicate plates were incubated at 37 and 27°C for 2 to 3 days on bovine, ovine, and equine blood agar media. The isolates were also tested for their ability to grow on blood agar containing polymyxin B (1,000 IU/ml; Oxoid). Motility was determined after stab inoculation into semisolid motility medium (17). The amount of radial extension of haze from the stab line and the level of zoospore activity observed microscopically in wet mounts were measured after incubation at 37 and 27°C. Susceptibility to penicillin G (10 U), streptomycin (30 μg), tetracycline (30 μg), neomycin (30 μg), chloramphenicol (30 μg) and sulfadiazine (100 μg) was determined by the standard single-disc method (4, 5), with the following two modifications: the *Dermatophilus* isolates were tested on Mueller-Hinton agar supplemented with 9% bovine blood, and the annular radii of the zones obtained with the isolates were measured after 48 h of incubation. We also determined the antibiotic susceptibility of W8 and W11 at 27°C. Reference organisms *Escherichia coli* NCTC 10418 and *Staphylococcus aureus* NCTC 6571 were treated as they are treated in standard tests.

**Biochemical characteristics.** Zoospore suspensions in distilled water were prepared by using growth harvested from cultures on blood agar after 3 days of incubation at 37°C and ≥8 days of incubation at room temperature to promote caseous colonies. The zoospores were examined with a transmission electron microscope after negative staining with 2% phosphotungstic acid. Observations of flagella and other features were recorded, and representative zoospores were photographed for measurements. Morphological features of mature filaments after growth in thioglycolate broth were highlighted by negative staining with India ink (7), and samples were photographed with an Olympus Vanox microscope. Capsular thickness was measured by using the photographic prints and the final magnification of each print. The mean of two to seven measurements was recorded for each isolate. Air-dried, acetone-fixed smears of *D. congolensis* zoospores were stained by using an indirect fluorescent-antibody procedure and monoclonal antibody 2F₄α (11). This monoclonal antibody reacts with *D. congolensis* zoospore surface antigen. The smears were examined to determine the fluorescent staining pattern and intensity. The microscopic appearance after Gram staining and Zielh-Neelsen staining for acid fastness (7) was also examined.

**Catalase and urease tests.** Catalase and urease tests were performed as previously described (12). The proteolysis test media used were nutrient gelatin medium (128 g/liter; Oxoid) and litmus milk medium (10% skim milk powder, 1.2% litmus solution). Tests for gelatin hydrolysis (method 2), methyl red and Voges-Proskauer reactions, nitrate reduction, indole production (method 2), and acid production from sugars in cystine tryptic medium (Gibco) were performed as described by Cowan (7).

Isolates were screened for various enzyme activities by using a gel diffusion technique (9), as follows. Suspensions of zoospores in peptone water were
prepared from caseous or mucoid growth harvested from cultures on bovine blood agar after 24 days of incubation at 27°C (12, 24). The suspensions were incubated in 4-mm-diameter wells in 1% agarose gels containing substrates (see below). Gels were stained with substrates, which were dissolved at a concentration of 0.1 mg/ml in phosphate-buffered saline (pH 7.5) containing 1% Ultra Pure DNA grade agarose (Bio-Rad Laboratories, Richmond, Calif.), were incubated in 4-sulfate, chondroitin 6-sulfate (Sigma Chemical Co., St. Louis, Mo.), and sheep dermal derm/derm (kindly provided by the Animal Products Technology Branch of the Department of Agriculture, Western Australia). Duplicate plates were incubated at 37 and 27°C for 24 to 48 h. The gels were stained with 0.02% toluidine blue (Gurr Biological Stains, Chaddwell Heath, Essex, England) in 0.1 M acetic acid, and the sizes of cleared zones were measured after the preparations were destained with 0.1 M acetic acid. The hyaluronidase substrate was used at a concentration of 100 mg/ml in phosphate-buffered saline (pH 7.5), this collagen is soluble at 4°C, but it forms a gel at 37°C. The collagen gel was pipetted into sterile microtiter plates (Flow Laboratories), and the resulting preparations were incubated at 37°C overnight to allow the collagen to solidify. Suspensions of *Dematophilus* isolates in peptone water (10 μl per well; 4 × 10 × 2 × 10 CFU per well) were pipetted into the centers of the wells containing collagen. The collagen gels were incubated at 37°C and were observed daily for 10 days; the experiment was repeated three times. In the presence of collagenase, the collagen gel liquified. The liquified gel from positive wells was compared with collagen gel obtained from negative wells by electron microscopy. The samples were acidified with 4 volumes of 0.1 M acetic acid and dialyzed against 50 volumes of 0.4% ATP in 0.1 M acetic acid for 48 h at 4°C. Samples were dried on Formvar grids, negatively stained with 2% phosphotungstic acid, and examined by transmission electron microscopy for the presence of segment long-spacing aggregates (arrays of aligned collagen molecules), as described by Snowdon and Swann (31).

An elastase screening method, in which elastin-Congo red (catalog no. E-0052; Sigma) was used, was adapted for microtiter plates from the method of Shotton (29). An elastin-Congo red suspension (1 or 4 mg/ml) and a suspension of *Dematophilus* zoospores (1 × 10 CFU/ml) in broth were dispensed into each well. Dilutions of elastase (Boehringer Mannheim) in broth were used as the standards. The following five dyes for suspending elastin-Congo red were compared: 0.02 M borate buffer (pH 8.8), borate buffer-peptone water (1:1, volume/volume), phosphate-buffered saline, peptone water, and 0.067% sodium dodecyl sulfate (Sigma) (pH 7.5). Duplicate plates were incubated at room temperature and at 27°C and were observed daily for 7 to 14 days for release of Congo red from the elastin particles.

**Enzyme assay analysis of bacterial DNA.** DNA was extracted from the *Dematophilus* isolates by a method adapted from the method of Wilson (36). Restriction enzyme analysis with Apal, BamHI, and PvuII was performed as described by Sambrook et al. (27). These restriction enzymes were chosen because they differentiated groups of *D. congolensis* isolates in other studies (9, 34). Digests were electrophoresed in 0.7% Ultra Pure DNA grade agarose (Bio-Rad Laboratories) in Tris-borate-EDTA buffer at 2.7 V/cm for 4 h. The gels were stained with a solution containing 0.1 μg of ethidium bromide per ml and were photographed with Polaroid type 667 film under UV light.

**Bacterial protein characterization.** *Dematophilus* isolates were harvested from low-protein nutrient agar containing 1% tryptone, 0.5% NaCl, 1.75% casein hydrolysate, and 0.015% soluble starch after incubation for 4 to 5 days at 27°C. Suspended in normal saline, and heated at 56°C for 6 h to release protein from the cells. Native protein polycrylamide gel electrophoresis (PAGE) and sodium dodecyl sulfate (SDS)-PAGE were performed with a Phast System apparatus (Pharmacia) by using 8 to 20% gradient polyacrylamide gels at 150 V, 3 mA, 1.5 W, and 15°C (105 V-h for SDS-PAGE and 150 to 160 V-h for native protein PAGE). The gels were stained by using the Phast System silver staining procedures and photographed.

**Infectivity in sheep, guinea pigs, and rabbits.** Infectivity studies were conducted with approval of the Animal Ethics Committee of the Department of Agriculture, Western Australia.Suspensions of zoospores of each isolate containing 10, 10, 10, and 10 CFU/ml of peptone water and a peptone water control were inoculated onto white rabbits that weighed approximated 3.2 kg. The rabbits were 12 weeks old (ewe wethers). The back of each animal was casually clipped over an area extending on each side of the midline. The area was swabbed with a cotton wool swab soaked in soap water and then washed with ether to remove the surface lipids. Five sites (4 by 4 cm for sheep, 2.5 by 2.5 cm for rabbits, and 2.5 by 2 cm for guinea pigs) were marked on each back. Inoculation was performed by immersing a cotton wool swab in a zoospore suspension or peptone water control and rolling it across a site for 1 min. The swab was weighed before and after application to ascertain the volume of suspension applied to the site and hence the approximate number of *Dematophilus* zoospores. The sites were examined 3, 7, 10, 12, and 14 days after inoculation by a single operator and scored for severity as described by Sanders et al. (28), but on a scale of 0 to 5. The lesion score for each site was calculated as follows: lesion score = lesion severity × percentage of area affected.

For each isolate on each day of examination, a quadratic regression was fitted between the log_{10} number of zoospores applied and the lesion score in order to estimate adjusted lesion scores at the standard doses (10, 10, 10, and 10 CFU/ml). A nonhierarchical cluster analysis was carried out to group the isolates into virulence categories for each animal species, as described previously (9).

**RESULTS**

**Cultural characteristics.** All three chelonid *Dematophilus* isolates produced discrete colonies at both 37 and 27°C on bovine blood agar and polymyxin blood agar, but all three isolates grew faster at 27°C than at 37°C. At 37°C, they produced small, dry, adherent, very slightly raised, gray-white colonies after 3 days. After transfer to room temperature, the colonies were white, raised, and umbonate or annelliform and became sticky after several days and then caseous after prolonged storage. At 27°C, the colonies were adherent, raised, which was larger than the colonies produced at 37°C. Colonies changed from dry to sticky more rapidly when the initial growth was at 27°C. At 37°C, colonies of all three isolates were surrounded by zones (annular radii, <1 mm) of beta-hemolysis as defined by Cowan (7), with outer zones (annular radii, 5 to 6 mm) of greenish alpha-hemolysis. After growth at 27°C, the isolates exhibited only beta-hemolysis. On ovine blood agar, the isolates produced 0.5- to 1-mm zones of beta-hemolysis at 37°C and 1-mm zones at 27°C, while on equine blood agar they produced little or no hemolysis at 37°C and <0.5-mm hemolysis zones at 27°C. During prolonged incubation (3 to 6 weeks) at room temperature on the tryptone-casein hydrolysate agar used for bacterial protein characterization, proliferation continued, and raised, sticky, ivory-colored growth having a rubbery texture was produced. Strain W8, W11, and W16* cultured substrates, which produced a distinctive characteristic odor of skin, was kindly provided by the Animal Products Technology Branch of the Western Australia Department of Agriculture. After dialysis at 4°C against phosphate-buffered saline (pH 7.5), this collagen is soluble at 4°C, but it forms a gel at 37°C. The soluble collagen was pipetted at 2°C into sterile microtiter plates (Flow Laboratories), and the resulting preparations were incubated at 37°C and were observed daily for 10 days; the experiment was repeated three times. In the presence of collagenase, the collagen gel liquified. The liquified gel from positive wells was compared with collagen gel obtained from negative wells by electron microscopy. The samples were acidified with 4 volumes of 0.1 M acetic acid and dialyzed against 50 volumes of 0.4% ATP in 0.1 M acetic acid for 48 h at 4°C. Samples were dried on Formvar grids, negatively stained with 2% phosphotungstic acid, and examined by transmission electron microscopy for the presence of segment long-spacing aggregates (arrays of aligned collagen molecules), as described by Snowdon and Swann (31).

An elastase screening method, in which elastin-Congo red (catalog no. E-0052; Sigma) was used, was adapted for microtiter plates from the method of Shotton (29). An elastin-Congo red suspension (1 or 4 mg/ml) and a suspension of *Dematophilus* zoospores (1 × 10 CFU/ml) in broth were dispensed into each well. Dilutions of elastase (Boehringer Mannheim) in broth were used as the standards. The following five dyes for suspending elastin-Congo red were compared: 0.02 M borate buffer (pH 8.8), borate buffer-peptone water (1:1, volume/volume), phosphate-buffered saline, peptone water, and 0.067% sodium dodecyl sulfate (Sigma) (pH 7.5). Duplicate plates were incubated at room temperature and at 27°C and were observed daily for 7 to 14 days for release of Congo red from the elastin particles.

**Enzyme assay analysis of bacterial DNA.** DNA was extracted from the *Dematophilus* isolates by a method adapted from the method of Wilson (36). Restriction enzyme analysis with Apal, BamHI, and PvuII was performed as described by Sambrook et al. (27). These restriction enzymes were chosen because they differentiated groups of *D. congolensis* isolates in other studies (9, 34). Digests were electrophoresed in 0.7% Ultra Pure DNA grade agarose (Bio-Rad Laboratories) in Tris-borate-EDTA buffer at 2.7 V/cm for 4 h. The gels were stained with a solution containing 0.1 μg of ethidium bromide per ml and were photographed with Polaroid type 667 film under UV light.
producing branching mycelia. Septa formed initially transversely with a slight bulging of the filament on one side of the septum. After prolonged incubation when a small inoculum was used, some mycelia had transverse and longitudinal septa at regular intervals; this created an appearance of cuboidal packets of spores within the mature filaments (Fig. 1). With a large inoculum in tightly capped tubes, we observed no septa other than the transverse septa. The mean thicknesses of the capsules around the mature filaments were 0.13 \(\mu\)m for W8, 1.3 \(\mu\)m for W11, and 0.5 \(\mu\)m for W16\(^T\) (Fig. 1). Only three mammalian isolates had capsules that were \(\leq 2 \mu\)m thick. Fluorescent-antibody staining with monoclonal antibody 2F\(_a\) was negative for all three chelonid isolates. Gram-stained smears contained gram-positive coccoid forms, germ tubes, and branching filaments with transverse and longitudinal divisions. Transverse rows containing up to five zoospores were

FIG. 1. Capsules around mature filaments of *Dermatophilus* isolates. (a) Tortoise isolate W8. (b) Snapping turtle isolate W16\(^T\). (c to f) Mammalian *D. congolensis* strains W3, S2, MB, and W15. The cultures were negatively stained with India ink. On each scale bar 1 division = 10 \(\mu\)m.
Table 1. Comparison of chelonid Dermatophilus isolates with D. congolensis isolates

<table>
<thead>
<tr>
<th>Category</th>
<th>Characteristics common to chelonid isolates and D. congolensis isolates</th>
<th>Distinctive characteristics of chelonid isolates W8, W11, and W16&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Morphology</strong></td>
<td>Gram-positive branching filaments which divide by transverse and longitudinal septa to form zoospores, which form germ tubes, elongate into filaments, and repeat the cycle; zoospore size; not acid fast</td>
<td>Thin capsules around mature filaments; not reactive with monoclonal antibody 2F; few flagella per zoospore</td>
</tr>
<tr>
<td><strong>Cultural</strong></td>
<td>Growth on blood agar initially dry and adherent; later growth is moist, especially at ≤30°C; beta-hemolysis on bovine blood agar at 27°C; colony morphology; resistant to polymyxin B; susceptible to penicillin G, tetracycline, chloramphenicol, and sulfadiazine</td>
<td>Growth faster at 27°C than at 37°C; sticky (not mucoid) growth on agar media after prolonged incubation at ≤30°C; two hemolysis zones on bovine blood agar at 37°C in the presence of 10% CO₂ (inner beta-hemolysis zone and outer alpha-hemolysis zone); low motility; resistant to streptomycin and neomycin</td>
</tr>
</tbody>
</table>

**Genomic**

**Infectivity**

**Biochemical**<sup>a</sup> Catalase positive; gelatin and casein hydrolyzed; acid produced from glucose; no acid produced from sucrose, trehalose, dulcitol, xylose, lactose, mannitol, sorbitol, or salicin; indole not formed; chondroitinase activity against chondroitin 4-sulfate; no hyaluronidase or elastase activity detected; methyl red and Voges-Proskauer tests negative. There is nitrate weakly reduced to nitrite; collagenase activity (in vitro); native and SDS protein PAGE patterns; urease negative (strains W8 and W11); distinctive odor of cultures. In this paper we describe the first isolates of Dermatophilus sp. obtained from tortoises or turtles. The mode of transmission of the bacteria to the chelonids is not known. Demzatuphilus strains obtained from reptiles include reports of six isolates obtained from lizards (2, 6, 23, 30). After experimental inoculation, two of the lizard isolates were pathogenic, one for lizards and mice (2) and another for a sheep (30). The lesions on the sheep resulting from experimental infection with the lizard isolate were more severe and persistent than the lesions caused by the chelonid isolates in this study, although the inoculation procedures differed. Two of the lizard isolates were urease negative (2, 23); one of these isolates produced acid from mannitol (2), while the other fermented sucrose and was resistant to penicillin, lincomycin, and triple sulfonamides (23). A third lizard isolate was resistant to neomycin and sulfonamides (30). Other pre-

**DISCUSSION**

In this paper we describe the first isolates of Dermatophilus sp. obtained from tortoises or turtles. The mode of transmission of the bacteria to the chelonids is not known. Dermatophilus sp. was not isolated from water and soil samples collected on site at the same time that strain W16<sup>b</sup> was isolated (32). Other reports of Dermatophilus strains obtained from reptiles include reports of six isolates obtained from lizards (2, 6, 23, 30). After experimental inoculation, two of the lizard isolates were pathogenic, one for lizards and mice (2) and another for a sheep (30). The lesions on the sheep resulting from experimental infection with the lizard isolate were more severe and persistent than the lesions caused by the chelonid isolates in this study, although the inoculation procedures differed. Two of the lizard isolates were urease negative (2, 23); one of these isolates produced acid from mannitol (2), while the other fermented sucrose and was resistant to penicillin, lincomycin, and triple sulfonamides (23). A third lizard isolate was resistant to neomycin and sulfonamides (30). Other pre-
viously reported cultural, biochemical, and microscopic characteristics of the lizard isolates were similar to those of mammalian *D. congolensis* isolates. While the lizard isolates differed slightly from *D. congolensis*, they also differed from the chelonid isolates in cultural characteristics, in mannitol and sucrose test results, and in antibiotic resistance characteristics (except for neomycin resistance).

Many cultural, morphological, and biochemical characteristics of the chelonid *Dermatophilus* isolates and the mammalian *D. congolensis* strains were similar, but there were several important differences. Gordon (12) observed wide ranges of variation in colony morphology, pigmentation, acid production from maltose, proteolytic ability, degree of encapsulation, and hemolysis under anaerobic growth conditions among *D. congolensis* isolates. Roberts described the plasticity of *D. congolensis* under different culture conditions, including inhibition of sporulation with reduced aeration (24, 26). Variation in antibiotic susceptibility among *D. congolensis* isolates, including resistance to streptomycin and neomycin, has been described by Elad and Yeruham (8). The variation and plasticity described above were considered when we compared the characteristics of the three chelonid *Dermatophilus* isolates with the characteristics of the mammalian *D. congolensis* strains. The ability of the chelonid isolates to grow faster at 27°C than at 37°C may reflect adaptation to poikilothermic hosts in a cool aquatic environment, and their sticky slime may improve adhesion to the host’s skin in water. The difference in the hemolysis patterns may be linked to pathogenicity and awaits further investigation. The odor of cultures of the chelonid isolates differed from the odor of cultures of the mammalian *D. congolensis* strains, indicating that different metabolic product(s) may be present. The lack of fluorescent-antibody staining with monoclonal antibody 2F4 indicates that the chelonid isolates lack a characteristic *D. congolensis* cell wall antigen. The poor motility and low number of flagella on the zoospores of the chelonid *Dermatophilus* isolates which we observed contrasted with the characteristics of most mammalian *D. congolensis* strains examined (9). It is possible that the chelonid isolates express flagella and exhibit greater motility under culture conditions different from the conditions used in this study. In tryptose-phosphate-10% ovine serum broth *D. congolensis* isolates usually produce a layer of growth or a distinct pellicle at or near the surface, a suspended network of tiny colonies joined by fine filaments, and a flocculent deposit, whereas cultures of the chelonid isolates lack surface growth. This may indicate an absence of aerotaxis (25) or may reflect the poor motility of the zoospores in culture.

The rapid resolution of lesions (by 7 days) observed with the chelonid *Dermatophilus* isolates on three mammalian species differs from the resolution of lesions observed with mammalian *D. congolensis* strains, which produce lesions that are most severe at days 5 and 7 on guinea pigs and rabbits, respectively,
and at days 12 to 14 on sheep (9). This may be due to the slower growth of the chelonid isolates at higher temperatures and adaptation to chelonid hosts. Maung (22) demonstrated that the tortoise Testudo hera produced antibody against Brucella abortus very slowly compared with mammals.

Our restriction enzyme analysis of bacterial DNAs revealed that there is a genetic difference between the chelonid isolates and the mammalian strains of D. congolensis, but it is difficult to assess degrees of relatedness by this technique. Using multiocular enzyme electrophoresis, Trott et al. (34), compared allozymes of Dermatophilus isolates at 16 enzyme loci. At 11 of the loci investigated, the chelonid isolates had a different allele than any of the D. congolensis strains, including three American Type Culture Collection reference strains. The electrophoretic type of the chelonid isolates differed from the electrophoretic types of seven D. congolensis strains at an average of 13.6 loci, resulting in a calculated genetic distance of 0.852.

The high proportion of different allozymes is well within the range found previously to differentiate eukaryotic species within a genus (3, 15, 20).

The results of cultural, morphological, and biochemical studies, including restriction enzyme analysis of bacterial DNAs and analysis of the PAGE patterns of bacterial proteins (see above), together with the genetic distances obtained in the study of Trott et al. (34), indicate that the chelonid isolates and D. congolensis strains belong to different species. We propose the name Dermatophilus chelonae for the chelonid isolates.

**Description of Dermatophilus chelonae** sp. nov.

**Dermatophilus chelonae** (Kr. lo'nae. L. gen. n. chelonae, of a turtle or tortoise, the source of the first isolates). Cocccoid zoospores that are 0.7 by 0.8 μm (minimum diameter by maximum diameter) to 1.2 by 1.6 μm produce germ tubes (diameter, ~1 μm) which elongate into filaments, eventually producing branching mycelia. Initially, septa form transversely, and there is a slight bulging of the filament on one side of the septum. Longitudinal septa are produced later, creating mature filaments with transverse and longitudinal divisions at regular intervals; the filaments are up to five cuboidal segments (~4 μm) in diameter and are surrounded by a thin capsule (thickness, 0.13 to 1.3 μm). The segments separate, becoming zoospores with zero to six flagella; aflagellate zoospores are the most common zoospores. All forms (zoospores, germ tubes, filaments, and segments) are gram positive. Not acid fast. Negative for fluorescent-antibody staining with monoclonal antibody 2F, (11).

Colonies on bovine blood agar after 3 days of growth are initially dry and adherent; the colonies are small, very slightly raised, and grey-white and have two hemolysis zones (beta and alpha) at 37°C, and they are white, raised, and larger and have a beta-hemolysis zone at 27°C. After transfer to room temperature, growth continues, and colonies are adherent, white, raised, and umbonate or annelliform; the colonies become sticky after several days and caseous after prolonged storage. On ovine blood agar, colonies are beta-hemolytic; on equine blood agar, colonies are nonhemolytic at 37°C and weakly phosphate-10% ovine serum broth produce a flocculent de-}

Urease is not always produced. Weak reduction of nitrate to nitrite occurs. Indole is not formed; methyl red and Voges-Proskauer tests are negative. Chondroitinase activity occurs with chondroitin 4-sulfate but not with chondroitin 6-sulfate. Collagenase activity occurs with in vitro-assembled collagen.

Resistant to polymyxin B (1,000 IU/ml), streptomycin (10 μg), and neomycin (30 μg). Susceptible to penicillin G (10 U), tetracycline (30 μg), chloramphenicol (30 μg), and sulfadiazine (100 μg).

D. congolensis and D. torquatus are distinctive electrophoretic types of the chelonid isolates. D. congolensis, D. torquatus, and D. pachydermidis are distinctive electrophoretic types of seven D. congolensis strains at an average of 13.6 loci, resulting in a calculated genetic distance of 0.852.
32. Sutherland, S. S. (Department of Agriculture, Western Australia). 1987. Personal communication.