

**Clostridium villosum** sp. nov. from Subcutaneous Abscesses in Cats

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*Clostridium villosum* sp. nov., isolated from subcutaneous abscesses of cats, is an obligately anaerobic, rod-shaped, sporeforming, monmotile organism which is gram positive in its early stages of growth and gram negative after 18 to 24 h. All strains of this species produce acetic, isobutyric, butyric, isovaleric, lactic, methyImalonic, and succinic acids from cooked meat-carbohydrate and peptone-yeast extract cultures supplemented with 5% horse serum. All of the strains produce ammonia and liquefy gelatin weakly but do not ferment acids or react in any other biochemical tests performed. The pattern of volatile acids produced, the lack of biochemical activity, the colonial morphology, and the late spore production distinguish *C. villosum* from other *Clostridium* species so far described. The type strain of *C. villosum* is VSB 3349; a culture of this strain has been deposited in the National Collection of Type Cultures under the number 11220.

From June to October 1978 we isolated from subcutaneous “fight wound” abscesses in cats 11 strains of an organism which does not appear to have been described previously. The organisms appeared in smears from purulent material as long, filamentous, gram-negative or gram-variable, rod-shaped bacteria. No spores were seen in the pus smears, but some of the rods appeared barred with nonstaining areas at intervals along the filaments. Purulent material which contained these morphological forms yielded an isolate with a distinctive colonial appearance.

This paper describes the characteristics of the above-mentioned organism, for which we propose the name *Clostridium villosum* sp. nov. (vil. los'um. L. adj. hairy, shaggy, rough haired). The type strain of *C. villosum* is Veterinary Pathology and Bacteriology Laboratory (VPB) 3349 (National Collection of Type Cultures strain 11220).

**MATERIALS AND METHODS**

**Bacterial strains.** The 11 strains studied (VPB) strains 523, 524, 3322, 3326, 3339, 3342, 3348, 3349, 3358, and 3359) were isolated from 11 cats, each with a solitary closed subcutaneous fight wound abscess. The organisms were part of a mixed flora of anaerobic and facultatively anaerobic bacteria found in these abscesses. Cultures of these strains were freeze-dried in skimmed milk after preliminary characterization; subsequently the cultures were revived and then fully characterized in parallel.

**Culture media and methods.** The organisms were isolated and grown initially on sheep blood agar plates and brain heart infusion agar plates (1), which were poured on the day of use, inoculated, and incubated anaerobically in a GasPak anaerobic system at 37°C.

Pure cultures of the isolates were grown in cooked meat plus peptic digest of meat (BVF) broth (4) (CMM) or cooked meat plus BVF broth supplemented with 0.4% glucose, 0.1% cellobiose, 0.1% maltose, and 0.1% starch (CMC). The basal media for fermentation studies were those of Holdeman et al. (1), and the fermentation of carbohydrates was determined by measuring after 5 days the pH of the cultures with carbohydrate and comparing it with the pH of the basal medium without carbohydrate incubated for the same length of time. For all biochemical tests, the liquid media except CMM and CMC were supplemented with 5% horse serum. For the biochemical tests, either GasPak jars in which catalysts had been changed after use and rejuvenated by heating at 160°C for 2 h or prerduced media (1) were used. Tubes of the latter were gassed with oxygen-free carbon dioxide at the time of inoculation.

The reagents and methods described previously (1) were used for the following tests: nitrate reduction; lecithinase, lipase, catalase, ammonia, and acetyl methylcarbinol production; esculin hydrolysis; and indole production in CMM. The medium for gelatin liquefaction was VF gelatin (BVF broth, 100 ml; glucose, 0.1 g; gelatin, 12 g; cysteine-HCl, 0.05 g; and vitamin K-heme solution, 1 ml [1]; the gelatin was clarified with beaten egg white and sterilized at 121°C for 15 min).

Fatty acids were extracted from CMC, peptone-yeast extract broth containing 5% horse serum (SPY), SPY-glucose, SPY-threonine, SPY-pyruvate, and SPY-lactate cultures as described previously (1) and were detected by gas-liquid chromatography on a Hewlett-Packard model 5830A gas chromatograph. A glass column 160 cm long and with an internal diameter of 2 mm was packed with 10% AT1200 plus 1% H3P04 on Chromosorb W-AW (80/100 mesh [AT-1200]) was used. The carrier gas was nitrogen at a flow rate of 30 ml/min, the oven temperature was 115°C, and the flame ionizer detector was run at 225°C. A Hewlett-Packard model 7671A automatic sampler was
used for all injections. The machine was computer
programmed to quantitate the products of fermention.
Gram stains were made on young cultures (8 to 12
h old) from cooked meat medium without carbohydrate and from blood agar to determine whether gram-
positive cells were present. Young cultures were fixed
with 2% glutaraldehyde in 0.1 M cacodylate buffer,
postfixed in 2% uranyl acetate, embedded, sectioned,
and examined in a transmission electron microscope
to determine cell wall structure. Cultures grown on
blood agar plates and CMC slants incubated anaerobically for up to 3 weeks were Gram stained and, if
spores or barred gram-negative cells were present,
samples were emulsified in CMC, and each was heated
by submerging in a water bath at 80°C for 10 min or
90°C for 5 min. The heated broth was then inoculated
into fresh CMC and incubated at 37°C to detect heat-
resistant cells. Usually two subcultures at 3-day inter-
vals were necessary to regain good growth of organ-
nisms. Samples from 3-week-old CMC slants were
fixed in 2% glutaraldehyde in 0.1 M cacodylate buffer,
postfixed in 2% uranyl acetate, embedded, sectioned,
and then examined in a transmission electron micro-
scope for spore formation and characteristics.
Repeated subcultures to blood plates and CMC
slants incubated aerobically or in a candle extinction
jar were performed to determine the oxygen relation-
ships of the organism. Growth at the bottom of tubes
of tryptose phosphate medium (yeast extract, 1 g;
tryptose, 20 g; NaCl, 5 g; glucose, 1 g; agar, 1 g; and
Na₂PO₄·12H₂O, 2.5 g; all dissolved in 1 liter of water
and sterilized at 121°C for 15 min) and movement
away from a central inoculation site in such tubes were
used to aid determination of motility, spore formation,
and anaerobiosis. Cultures at various stages of growth
in CMC were examined for motility in wet prepara-
tions and were examined for flagella by negative stain-
ing with 2% phosphotungstic acid and viewing in a
transmission electron microscope. The broth disk test
of Williams and Thiel (5) was used to determine sus-
ceptibility to antimicrobial agents.

RESULTS AND DISCUSSION

Morphology and Gram staining. All of the
strains studied had similar morphology and
Gram stain reactions. Cells were rod shaped with
parallel sides and rounded ends and measured
0.6 by 4.0 to 6.0 μm. Filaments reaching 24 to 30
μm in length were common in cultures more
than 24 to 48 h old. Thin-section electron micro-
scopy showed that the structure of the cell wall and the mode of division were consistent
with gram-positive organisms and that numerous mesosomes were produced. Individual cells
retained the Gram stain for up to 18 to 24 h in broth culture or on an agar surface but became
gram variable or gram negative after this time.
By 7 days most of the cells showed gram-nega-
tive staining, with nonstaining gaps at intervals
along the cell, thus giving the organisms a barred
or beaded appearance. Electron microscopy
showed many large cytoplasmic vacuoles in the cells.

Colonial characteristics. Agar colonies of
all of the strains were basically similar. Surface colonies on solid media measured 0.5 mm at 24
h (Fig. 1) and increased to approximately 5 mm
by 3 days. Mature colonies were differentiated
into central and peripheral portions (Fig. 2). The
central portion was raised convex or dome-
shaped, rough, dull, and whitish to yellowish in
color, whereas the peripheral portion varied
among strains but had a greater or lesser extent
of rhizoid or fimbriate edge (Fig. 2). All colonies
were strongly adherent to the underlying me-
dium, and colonies with less peripheral portions
had their central zones more deeply indented
into the agar. No hemolysis was noted on blood
agar plates. The colonies were very difficult to
emulsify, and rigorous trituration was required
to obtain even a granular suspension. Deep stab
colonies appeared like whisps of wool or cotton
with a dense granular center. There was no

![FIG. 1. Surface colony on blood agar at 24 h. Bar = 1 mm.](image1)

![FIG. 2. Surface colony on blood agar at 3 days. Bar = 1 mm.](image2)
growth on the surface of blood agar plates or CMC slants incubated aerobically or in a candle extinction jar.

**Cultural characteristics.** Organisms would not grow in prereduced or conventionally prepared, anaerobically incubated peptone-yeast extract without the addition of 5% horse serum. Tween 80 and formate-fumarate supplementation (1) did not stimulate growth. Cultures in SPY showed slight turbidity with membranous or clumpy growth (Fig. 3). Strains grew well in CMC whether the medium was (i) prereduced, (ii) conventionally prepared and incubated anaerobically, or (iii) steamed and cooled before inoculation and incubation. In CMC the organisms formed a delicate membrane near the surface of the broth, and throughout this membrane the bacteria clumped and appeared as discrete colonies resembling those formed in agar stabs. Late in growth, the membrane structure collapsed and settled into and around the meat particles (Fig. 4).

**Motility.** Extensive observations of broth cultures were made to try to detect motility. Motility was not observed in cultures varying from a few hours to 10 days of age when examined with a light microscope, and electron microscope examination of negatively stained preparations of these cultures failed to reveal flagella. Organisms seeded from CMC into the center of deep tubes of the semisolid tryptose phosphate medium failed also to provide evidence of motility in any of the strains. These rigorous observations were thought necessary, as the colonial appearance of the organisms suggested that they could be motile.

**Spore formation.** Spores were single, oval, and subterminal and caused a slight distention of the sporangium. Immature spores were smaller, retained the Gram stain, and appeared as small, bluish spots in the otherwise gram-negative rods. When the organisms were isolated, spores were not detected, possibly because the cultures were not incubated long enough. However, for all of the strains, anaerobically incubated CMC slant cultures were heat resistant (10 min at 80°C; 5 min at 90°C) after 2 weeks of incubation. Cells of these cultures were gram negative and had barred staining. Heat-resistant spores of strains 523, 3349, and 3857 were also produced on blood agar plates incubated anaerobically for 3 weeks. Heat-resistant cells were not detected in any strain grown for up to 3 weeks in either CMC or tryptose phosphate medium.

**Biochemical reactions.** All of the strains liquefied gelatin weakly, and ammonia was produced in CMM and SPY. None of the strains fermented any of the carbohydrates tested (esculin, fructose, glucose, lactose, maltose, mannitol, mannose, melibiose, ribose, sucrose, xylose), as determined by pH measurement. None
of the strains gave positive reactions in any of the other biochemical tests performed.

Fermentation products. Strains were grown for 2 days in CMC and for 5 days in SPY. The major fermentation acids detected from CMC cultures are listed in Table 1. Trace amounts of lactic, succinic, and methylmalonic acids were detected. As reflected in Table 1, the organisms grew better in CMC than in SPY media. However, organisms grew better in CMC incubated in GasPak jars or aerobically than in prereduced CMC. It seems, therefore, that CMC is beneficial with respect to both nutrition and pH. The fermentation products resulting from growth in SPY, SPY-glucose, SPY-threonine, SPY-pyruvate, and SPY-lactate can be seen in Table 1. Threonine was not converted to propionate, and lactate was not utilized, but greater amounts of butyric acid were produced in cultures which contained pyruvic acid than in those which did not. All strains produced fermentation products from SPY-based medium, but, with the exception of SPY-threonine, the amounts of products produced were considerably lower than those from CMC.

Resistance to antimicrobial agents. All strains were susceptible to penicillin (2 U/ml), amoxycillin (2.5 µg/ml), carbencillin (100 µg/ml), doxycycline (6 µg/ml), chloramphenicol (12 µg/ml), and erythromycin (3 µg/ml).

Distinguishing characteristics. The characteristics of *C. villosum* do not conform to those of any of the published descriptions of other members of the genus *Clostridium* (3), and no reference to this sort of organism could be found in a thorough search of the current literature.

Spores are not readily detected in this species, and the cells are usually observed to be gram-negative rods. It is possible, however, to distinguish it from gram-negative, rod-shaped species by its gram-positive or gram-variable reaction in young cells, by its strictly anaerobic growth, and by its distinctive colonial appearance. However, its fermentation products suggest that it may be confused with some members of the genus *Bacteroides* (2), but the fact that it does not produce propionic acid should help distinguish it from members of that genus.

Pathogenic significance. The organism described here was found as one of a number of anaerobic species present in each abscess examined. It was always present in large numbers, but it has yet to be determined whether these bacteria alone are able to produce disease. The indication is that the organism is present as part of the oral flora of cats and contributes to the purulent disease after gaining entry, along with other anaerobes, into an unusual site. It is perhaps significant that we have also isolated strains identical to those reported here from the pyothorax in cats. In that site, it is likely that the organism is pathogenic; and in the thoracic lesions it was the predominant isolate.

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**REPRINT REQUESTS**

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**LITERATURE CITED**


