Succiniclasticum ruminis gen. nov., sp. nov. – A Ruminal Bacterium Converting Succinate to Propionate as the Sole Energy-Yielding Mechanism

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A gram-negative, anaerobic, non-motile, non-spore-forming, rod-shaped bacterium that fermented succinate quantitatively to propionate was isolated from a high dilution of rumen ingesta obtained from a dairy cow fed a production diet containing grass silage as the main roughage source. This organism did not grow on any of the following energy sources: 12 carbohydrates, pyruvate, lactate, 7 dicarboxylic acids, aspartate, citrate, and trans-aconitate. Both rumen fluid and yeast extract were necessary for good growth on succinate. The organism was negative for the following characteristics: production of propionate from threonine, protein digestion, sulfide production, nitrate reduction, catalase activity, and urease activity. There was no growth at 22°C and reduced growth at 45°C compared with growth at 39°C. The DNA base composition was 52 mol% G+C. The complete 16S rRNA sequence (EMBL accession number, X81137) was obtained, and the phylogenetic relationships of the organism were determined. The most closely related genera were the genera Aciduminococcus and Phascolarctobacterium. The name proposed for this bacterium is Succiniclasticum ruminis gen. nov., sp. nov.; the type strain is strain SE10 (= DSM 9236). Additional isolation attempts revealed that S. ruminis is a common inhabitant of the rumina of cows that are fed production diets and of cows on pasture.

MATERIALS AND METHODS

Media. All media were prepared under strictly anaerobic conditions (15), and the gas phase (except where indicated otherwise) was CO₂-free CO₂. Most of the medium components were heat sterilized; certain constituents were filter sterilized. The incubation temperature was usually 39°C.

The basal medium consisted of (per liter): 0.225 g of KH₂PO₄, 0.225 g of KH₁PO₄, 0.45 g of NaCl, 0.45 g of (NH₄)₂SO₄, 0.045 g of CaCl₂ (anhydrous), 0.09 g of MgSO₄·7H₂O, 0.36 g of NaHCO₃, 0.25 g of cysteine hydrochloride·H₂O, 0.25 g of Na₂S (hydrated), 0.005 g of indigo carmine, and 400 ml of rumen fluid (strained through cheesecloth and centrifuged at 15,000 × g for 30 min). Anaerobic dilution contained the same ingredients except that rumen fluid and sulfide were not present and the concentration of cysteine was doubled. This dilution was used for serial dilution and for carrying inocula.

The basal medium was modified as follows for different purposes (the succinate used in the media described below was diosuccinate·4H₂O). Enrichment medium contained 50 mM succinate. Bacteria from enriched medium produced colonies on medium containing (per liter) 10 g of succinate, 5 g of yeast extract (Difco), and 20 g of agar. The maintenance medium used for slopes contained (per liter) 10 g of succinate, 5 g of yeast extract, and 15 g of agar. Poorly buffered medium contained 40 mM succinate and 5 g of yeast extract per liter but no NaHCO₃; the Na of NaHCO₃ was replaced by the Na of NaCl, the pH was adjusted to 6.8 with HCl, and CO₂ was replaced with O₂-free N₂. Gelatin liquefaction medium contained (per liter) 10 g of succinate, 5 g of yeast extract, and 120 g of gelatin. Cascin digestion medium contained (per liter) 10 g of succinate, 5 g of yeast extract, 20 g of casein, and 20 g of agar but no cysteine or sulfide. Sulfide (H₂S) production medium contained (per liter) 10 g of succinate, 5 g of yeast extract, and 3 g of SIM medium (Difco); the cysteine concentration was doubled but no sulfide was added to this medium (a similar medium without rumen fluid was used for the same purpose). Nitrate reduction medium contained (per liter) 10 g of succinate and 5 g of yeast extract; a filter-sterilized KNO₃ solution was added to a final concentration of 1 g/liter after sterilization, and cysteine, sulfide, and indigo carmine were omitted. Urease production medium contained (per liter) 10 g of succinate and 5 g of yeast extract, and a filter-sterilized urea solution was added to a final concentration of 15 g/liter after sterilization. The urease test was also done in poorly buffered medium (see above) performed with similar additions. Tests to determine the effect of pH on growth were performed in a medium similar to that used for the urease test except that urea was omitted.

In a number of tests I assessed growth by measuring optical density. When the media were used for these tests were prepared, the rumen fluid was centrifuged at 50,000 × g for 30 min. Indigo carmine was omitted, and 5 g of yeast extract per liter was included (modified basal medium). Energy sources were added as concentrated, filter-sterilized solutions. The final concentrations of acid substrates (as their sodium salts), aspartic acid, and threonine were 40 mM, 40 mM, and 3 g/liter, respectively. When lactate was used, 20 mM t-lactate and 20 mM l-lactate were added. The succinate concentrations tested were 20, 40, 70, and 100 mM. Every other substrate (except where indicated otherwise) was added at a final concentration of 10 g/liter; glucose and mannose were also tested at a concentration of 2 g/liter. Susceptibility to oxygen was examined in modified basal medium containing 40 mM succinate that was prepared with and without cysteine and sulfide, as well as in the latter medium in 13-m tubes (containing 5 ml of medium) injected with 2 ml of sterile air. The effects of different temperatures and the effect of Tween 80 (0.2 g/liter) on growth were assessed in modified basal medium containing 40 mM succinate. Bacteria were grown in modified basal medium containing 70 mM succinate to determine the G+C content.

Isolation. A sample of whole rumen ingesta was obtained from a lactating, rumen-fistulated cow fed a production diet containing grass silage as the main roughage source and was collected and treated as described previously (15). Aliquots (1 ml) of a 10⁻³ dilution in 5 ml of enrichment medium were incubated, and portions of these preparations were transferred successively to fresh medium after 4, 3, and 1 days. Aliquots of diluted enriched cultures were grown in roll bottles containing succinate agar medium. After 3 days 10 colonies were picked. The strains obtained were subsequently reisolated and tested for growth in liquid succinate medium.

Characterization tests. For optical density measurements growth was monitored with a Novaspec II spectrophotometer (Pharmacia LKB) at 600 nm in glass tubes. Fermentation end products were determined by high-performance liquid chromatography (1), using pivalic acid as the internal standard. Except for the fermentation of the media, the tests to determine motility, catalase activity, urease activity, gelatin liquefaction, casein hydrolysis, H₂S production, production...
tion of propionate from threonine, and nitrate reduction (with sulfanilic acid and dimethyl-alpha-naphthylamine as reagents) were performed as recommended by Holdeman et al. (7).

G+C content was estimated as previously described (17). DNAs from Escherichia coli B, Micrococcus luteus (Micrococcus lysodeikticus), and calf thymus (all obtained from Sigma) were used as calibration standards.

All determinations were done at least in duplicate.

Inocula. Inocula (0.1 ml/5 ml of medium) for tests were grown on succinate maintenance slopes for about 68 h, and the organisms were tagged with dye by using a Taq Dye-Deoxy Terminator Cycle sequencing kit (Applied Biosystems, Weiterstadt, Hessen, Germany) as recommended by the manufacturer. Sequences were electrophoresed by using an Applied Biosystems model 373A DNA sequencer. The 16s ribosomal DNA sequence was manually aligned with representative sequences of members of the Clostridium-Bacillus line of descent.

Pairwise evolutionary distances were computed by using the correction of Jukes and Cantor (10). The least-squares distance method of DeSocio (5) was used to construct a phylogenetic dendrogram from distance matrix data.

Nucleotide sequence accession number. The nucleotide sequence determined in this study has been deposited in the EMBL database under accession no. X81137.

### RESULTS AND DISCUSSION

A strain isolated from enrichment medium that had been inoculated with a 10⁻⁸ dilution of whole rumen ingesta was able to grow on succinate as the sole energy source. This strain, named SE 10⁷ (T = type strain), was characterized. Strain SE 10⁷ cells were gram-negative, nonmotile, non-spore-forming, short rods that grew in the absence of oxygen at temperatures prevailing in the rumen. Of the substrates tested, only succinate supported growth, and propionate was the only end product detected. Negative results were obtained for a variety of tests (see below).

**Relatedness to other taxa.** Some of the characteristics that distinguish taxa which have phenotypic or genotypic properties similar to those of SE 10⁷ are summarized in Table 1. Strain SE 10⁷ differs from Propionigenium modestum (13) in its inability to grow on pyruvate, oxaloacetate, fumarate, malate, or aspartate; Propionigenium modestum ferments fumarate, malate, and aspartate only after lag periods of 4 to 8 days. Colonies of Propionigenium modestum are yellowish, while colonies of SE 10⁷ are colorless. Whereas Propionigenium modestum can grow at 15°C, strain SE 10⁷ cannot grow at 22°C. SE 10⁷ differs from Acidaminococcus fermentans (3) in that it cannot utilize amino acids and peptides as main energy sources or ferment citrate and trans-aconitate. Of the phenotypic characteristics that have been determined, the only one that distinguishes SE 10⁷ from Phascolarcobacterium faecium (4) is cell morphology. While Phascolarcobacterium faecium can produce long branched cells, only very limited branching (V shapes) was observed with SE 10⁷ under certain conditions (6).

Of the energy sources examined, only succinate is fermented by Phascolarcobacterium faecium and SE 10⁷, and fumarate inhibits the growth of both organisms. The dendrogram (Fig. 1), based on 16S rRNA sequence comparisons, shows that SE 10⁷ is most closely related to Phascolarcobacterium faecium and A. fermentans. The G+C content of SE 10⁷ is closer to that of A. fermentans than to that of Phascolarcobacterium faecium (Table 1). Considering the phenotypic similarity of Phascolarcobacterium faecium and SE 10⁷, differentiation of these taxa rests largely on the difference in G+C contents and on the level of 16S rRNA sequence similarity, which is not greater than 92.7%, as shown in Table 2. Therefore, I propose that strain SE 10⁷ should be placed in a new genus. **Description of Succiniclasticum gen. nov.** Succiniclasticum (Suc.cin.i.clas'ti.cum. L. adj. succinctus, succinie; Gr. v. kloan, to break; L. adj. clasticum, breaking; L. neut. n. Succiniclasticum, breaking or splitting succinic acid). The genus is characterized by not being able to ferment carbohydrates, amino acids, or mono-, di-, and tricarboxylic acids other than succinate, which is converted to propionate. Members of this genus are not proteolytic and do not produce catalase or urease or reduce nitrate. Cells are not extensively branched.

### TABLE 1. Some properties of genera resembling strain SE 10⁷ either phenotypically or genotypically

<table>
<thead>
<tr>
<th>Taxon</th>
<th>Morphology</th>
<th>G+C content of DNA (mol%)</th>
<th>Succinic acid fermented (propionate produced)</th>
<th>Amino acids and peptides used as energy sources</th>
</tr>
</thead>
<tbody>
<tr>
<td>SE 10⁷</td>
<td>Short rods, limited branching on some media</td>
<td>52</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>Phascolarcobacterium</td>
<td>Short and long branched rods</td>
<td>41–42</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>Acidaminococcus</td>
<td>Cocci</td>
<td>56</td>
<td>−</td>
<td>+</td>
</tr>
<tr>
<td>Propionigenium</td>
<td>Short rods</td>
<td>34</td>
<td>+</td>
<td>−</td>
</tr>
</tbody>
</table>

*None of the taxa is motile, produces spores, or ferments carbohydrates.*
Description of *Succiniclasticum ruminis* sp. nov. *Succiniclasticum ruminis* (ru'mi.nis. L. neut. gen. n. ruminis, of the rumen). Cells are gram-negative and non-motile, do not produce spores, and are short rods which are 0.3 to 0.5 µm wide and up to 1.8 µm long (Fig. 2). The thicker rods tend to have rounded ends, while the narrower ones often have one end, or occasionally both ends, somewhat tapered and can be slightly curved. Large clumps of cells are often observed in liquid media or in the water of syneresis of cultures grown on agar slopes. These clumps are easily dispersed by shaking. After incubation on rumen fluid-succinate agar slopes for about 18 h, surface colonies have diameters of up to 0.5 mm. They are somewhat irregularly round with smooth edges, convex, practically colorless, and slightly opaque with a glistening appearance. Submerged colonies that develop after inoculation into molten agar are lens-shaped discs.

**Growth characteristics.** The following substrates do not support growth: glucose, galactose, mannose, rhamnose, fructose, xylose, arabinose, maltose, cellobiose, lactose, sucrose, trehalose, mannitol, glycerol, aspartic acid, the sodium salts of DL-lactic and pyruvic acids, the disodium salts of oxalic, malonic, oxaloacetic, malic, fumaric, glutaric, and methylmalonic acids, the trisodium salts of citric and trans-aconitic acids, and yeast extract (which contains amino acids). Prolonged incubation (9 days) with fumarate, malate, aspartate, citrate, or trans-aconitate does not induce utilization. When acetate and butyrate are present in the rumen fluid of the medium, they are not assimilated to a measurable extent. Succinate is the only substrate tested that is fermented. Growth in the presence of 40 mM succinate plus 10 mM fumarate gives about the same maximal optical density as growth in the presence of succinate alone, but the maximal value is reached 1 to 2 h later. Very little growth occurs in medium containing 40 mM succinate and 40 mM fumarate. There is very little growth in medium from which either rumen fluid or yeast extract is omitted (Fig. 3). Succinate concentrations of 40, 70, and 100 mM result in progressively slower growth, but the final optical densities are correspondingly higher. Succinate is converted quantitatively to propionate, and no other end products are detected. CO₂ is probably released, but CO₂ release has not been measured. Growth in medium from which CO₂ and NaHCO₃ are omitted (poorly buffered medium with NaCl added) is appreciably less (the optical density is one-half as great) and slower than growth in medium containing CO₂ and NaHCO₃ (well-buffered medium). This could be due to pH. In well-buffered medium, which contains 40 mM succinate, the pH increases about 0.1 U, while the pH increases about 0.5 U in poorly buffered medium after growth has ceased. Growth in medium prepared under anoxic conditions is about the same irrespective of whether a reducing agent (cysteine or sulfide) is added. However, when air is added to medium without a reducing agent, little growth occurs. At 39°C the maximum optical density is about twice the optical density at 45°C after about the same period of incubation. There is no growth at 22°C. Tween 80 has no effect on growth.

![FIG. 2. Phase-contrast micrograph of *Succiniclasticum ruminis* SE 10⁷ cells on a slide covered with agar. Bar = 5 µm.](image-url)
ruminis

Deutsche Sammlung von Mikroorganismen und Zellkulturen

nate; A, 20 mM succinate; 0.40 mM succinate. OD₆₀₀, optical density at 600 nm.

Symbols: 0, no yeast extract, 40 mM succinate; O, 0.5% yeast extract, no succinate; ∆, 20 mM succinate; Ω, 40 mM succinate. OD₆₀₀, optical density at 600 nm.

Characterization tests and G+C content. Strain SE 10⁷ is not proteolytic; it neither liquefies gelatin nor hydrolyzes casein, and it does not produce H₂S. Propionate is not formed from threonine. It does not produce catalase or urease or reduce nitrate. The G+C content of the DNA is about 52 mol% (values for three separate determinations: 51.1, 52.0, and 52.2 mol%).

Strain SE 10⁷ is the type strain and has been deposited in the Deutsche Sammlung von Mikroorganismen und Zellkulturen as strain DSM 9236.

Isolation of other strains. Following the characterization of the unusual bacterium described above, I isolated bacteria having similar properties from the rumina of cows fed production diets containing silage or cows on pasture (16). I concluded that this organism is common in the rumen biota. Two of these bacterial strains are being characterized now. One of them was isolated from rumen ingesta (2 × 10⁻⁷ dilution) from a silage-fed cow different from the cow from which SE 10⁷ was isolated, while the other was isolated from a cow (10⁻⁸ dilution of ingesta) on pasture supplemented with concentrate.

Role in rumin digestion. The relevance of *Succinichlamidobacterium ruminis* clearly lies in the fact that this organism specializes in fermenting succinate and converting it quantitatively to propionate, the most important precursor of glucose in ruminants. The low energy yield when anaerobes convert succinate to propionate (14) should make this bacterium highly efficient in this conversion (there is a high rate of substrate turnover for a low rate of cell growth). Despite this, *Succinichlamidobacterium ruminis* appears to occur in high numbers (at least 10⁶ cells per g of ingesta) in the rumina of cows fed production diets containing grass silage (and usually about an equal amount of concentrate) on a dry matter basis) as the main roughage source and in the rumina of cows on pasture. This implies that the growth rate of this organism is high enough so that the organism can compete successfully for essential growth factors and may depend on limited competition for succinate from other bacteria. The level of succinate production in the rumina of cows fed grass silage diets is probably high because the level of *Prevotella ruminicola*, a succinate producer, tends to be greater than 50% (15) of the “total culturable” bacteria and that of cows on pasture tends to be greater than 40% (16). Because *Succinichlamidobacterium ruminis* has not been detected in ruminants fed other diets, it is possible that this organism typically occurs only in animals fed particular types of diets and perhaps even in particular animal species. This appears to be the case with *V. parvula* (formerly *V. gazogenes*) (9), a succinate and lactate fermenter that has not been found to be numerically important except in sheep rumina. Similarly, lactate-fermenting sarcinas have been found to be important only in cows fed production diets containing grass silage (18). More work will have to be done before it can be determined whether the presence of *Succinichlamidobacterium ruminis* in the rumen biota is related to diet.

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REFERENCES


FIG. 3. Effect of yeast extract and concentration of succinate on the growth of *Succinichlamidobacterium ruminis* SE 10⁷ in medium containing 40% rumen fluid. Symbols: 0, no yeast extract, 40 mM succinate; O, 0.5% yeast extract, no succinate; ∆, 20 mM succinate; Ω, 40 mM succinate. OD₆₀₀, optical density at 600 nm.