Akkermansia muciniphila gen. nov., sp. nov., a human intestinal mucin-degrading bacterium

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The diversity of mucin-degrading bacteria in the human intestine was investigated by combining culture and 16S rRNA-dependent approaches. A dominant bacterium, strain Muc<sup>T</sup>, was isolated by dilution to extinction of faeces in anaerobic medium containing gastric mucin as the sole carbon and nitrogen source. A pure culture was obtained using the anaerobic soft agar technique. Strain Muc<sup>T</sup> was a Gram-negative, strictly anaerobic, non-motile, non-spore-forming, oval-shaped bacterium that could grow singly and in pairs. When grown on mucin medium, cells produced a capsule and were found to aggregate. Strain Muc<sup>T</sup> could grow on a limited number of sugars, including N-acetylglucosamine, N-acetylgalactosamine and glucose, but only when a protein source was provided and with a lower growth rate and final density than on mucin. The G+C content of DNA from strain Muc<sup>T</sup> was 47.6 mol%. 16S rRNA gene sequence analysis revealed that the isolate was part of the division Verrucomicrobia. The closest described relative of strain Muc<sup>T</sup> was Verrucomicrobium spinosum (92% sequence similarity). Remarkably, the 16S rRNA gene sequence of strain Muc<sup>T</sup> showed 99% similarity to three uncultured colonic bacteria. According to the data obtained in this work, strain Muc<sup>T</sup> represents a novel bacterium belonging to a new genus in subdivision 1 of the Verrucomicrobia; the name Akkermansia muciniphila gen. nov., sp. nov. is proposed; the type strain is Muc<sup>T</sup> (= ATCC BAA-835<sup>T</sup> = CIP 107961<sup>T</sup>).
The introduction of high-resolution molecular techniques has improved analyses of complex microbial ecosystems. The most important advance has been the use of the 16S rRNA gene as a molecular fingerprint to analyse microbial diversity. Molecular approaches have indicated that a lack of knowledge regarding cultivation conditions has hampered our view of the intestinal microbiota (Vaughan et al., 2000). As a consequence, a substantial proportion of the microbiota has not yet been cultured or described (Zoetendal et al., 1998; Suau et al., 1999); this may be due mainly to the lack of appropriate cultivation techniques. However, new, alternative and improved cultivation approaches are continuously being developed and recently a number of novel species and genera have been cultured from the GI tract: Roseburia intestinalis (Duncan et al., 2002), Campylobacter hominis (Lawson et al., 2001), Ruminococcus luti (Simmering et al., 2002), Anaerostipes cacaoe (Schwitz et al., 2002), Dorea longicatena (Taras et al., 2002) and Victivallis vadensis (Zoetendal et al., 2003).

In the present study, mucin-degrading bacteria from human faeces were enriched using a most probable number (MPN) approach in which the medium contained mucin as the sole carbon and energy source. The enrichments were analysed by denaturing gradient gel electrophoresis (DGGE) of PCR-amplified 16S rRNA gene sequences. A single DGGE type dominated all the positive MPN enrichments. The organism corresponding to the dominant DGGE type was isolated and characterized; it represents a novel intestinal bacterium, strain MucT, that is able to use gastric mucin in pure culture.

A faecal sample from a healthy adult volunteer was freshly collected in a polyethylene bag and 0.5 g was diluted into 9 ml sterile anaerobic Ringer’s solution containing 0.5 g cysteine l⁻¹. This suspension was thoroughly mixed and serially diluted (10-fold) in Ringer’s. Each dilution (1 ml) was inoculated in triplicate into 9 ml bicarbonate-buffered medium. This basal medium contained (l⁻¹): 0.4 g KH₂PO₄; 0.53 g Na₂HPO₄; 0.3 g NH₄Cl; 0.3 g NaCl; 0.1 g MgCl₂.6H₂O; 0.11 g CaCl₂; 1 ml alkaline trace element solution; 1 ml acid trace element solution; 1 ml vitamin solution; 0.5 mg resazurin; 4 g NaHCO₃; 0.25 g Na₂S.7–9H₂O. The trace element and vitamin solutions were as described previously (Stams et al., 1993). All compounds were autoclaved, except the vitamins, which were filter-sterilized. This basal medium was supplemented with 0.7 % (v/v) clarified, sterile rumen fluid and 0.25 % (v/v) commercial hog gastric mucin (Type III; Sigma), purified by ethanol precipitation as described previously (Miller & Hoskins, 1981). This medium is further referred to as mucin medium. Unless indicated, incubations were done in serum bottles sealed with butyl-rubber stoppers at 37 °C under anaerobic conditions provided by a gas phase of 182 kPa (1.8 atm) N₂/CO₂ (80:20, v/v). Enrichments were done in 30 ml serum bottles with 10 ml liquid volume. Negative controls comprised one series of mucin media that was not inoculated and another series that was inoculated, but not supplemented with mucin. Mucindegrading bacteria were quantified using the MPN technique (n = 3). The soft agar technique was used to isolate a pure culture as follows: the highest dilution where growth was observed was serially diluted in phosphate buffer (pH 7) until 10⁻⁹ dilution and the 10⁻⁶ to 10⁻⁹ dilutions were re-inoculated into the same medium containing 0.75 % agar (agar noble; Difco). Single colonies were picked, grown in mucin medium and re-inoculated in soft agar mucin medium. This step was repeated until purity.

Generation times were determined in mucin medium and growth was analysed in triplicate by measuring absorbance at 600 nm. The optimum pH and temperature were measured in triplicate on brain–heart infusion (BHI; Difco) supplemented with 1 mM Na₂S. Temperatures tested were 4–45 °C, at intervals of 5 °C; growth was determined at pH 5–9, at intervals of 0.5 pH units (adjusted with HCl or NaOH) at 37 °C. Cultures were incubated for at least 1 month.

Potential substrates for growth were tested at a concentration of 10 mM in the same liquid basal medium or in basal medium supplemented with peptone, tryptone, casitone and yeast extract at a concentration of 0.5 or 2 g l⁻¹. Cultures were incubated for up to 4 weeks. Human gastric mucin was isolated from HT-29 MTX human intestinal cell lines and this mucin was added to the basal liquid medium at a concentration of 0.05 %. Rich media BHI and Columbia broth (Difco) and 16 g Wilkens–Chalgren broth (WC broth; Oxoid) 1⁻¹ were also tested as growth substrates. To test the origin of the nitrogen source, the solution containing NH₄Cl was not added to the mucin medium.

Cell morphology, motility and spore formation were investigated using phase-contrast microscopy. The Gram reaction was assessed using Gram staining as described previously (Pluge et al., 2000). To test for the presence of a capsule, an Indian ink suspension was used.

For TEM of strain MucT, cells were fixed with 0.25 % glutaraldehyde. Negative staining was performed on 400 copper mesh grids with glow-discharged parlodion carbon support film. Micrographs were recorded at a magnification of 40000 x on a JEOL 1010 electron microscope operating at 80 kV. For SEM, droplets of strain MucT were put onto poly-L-lysine-coated Nucleopore polycarbonate membranes (Costar). These membranes were fixed for 1 h in 4 % glutaraldehyde in growing medium. Specimens were dehydrated in a graded series of ethanol and critical-point dried with carbon dioxide. The samples were glued onto a sample holder using carbon adhesive tabs. Samples were sputter-coated with 10 nm platinum in a dedicated preparation chamber (CT 1500 HF) and analysed with a field emission SEM (JEOL 6300 F) at 5 kV.

The G+C content of DNA of strain MucT was determined
at the DSMZ (Deutsche Sammlung von Mikroorganismen und Zellkulturen, Braunschweig, Germany) by HPLC (Mesbah et al., 1989).

To monitor the dynamics of the human faecal mucin-degrading population, DGGE analysis of 16S rRNA gene amplicons was performed. DNA was extracted from the faecal sample, enrichment cultures and a pure culture isolated from the highest dilution. DNA isolation and the amplification of the V6 to V8 regions of the 16S rRNA gene from these samples were performed as described previously (Zoetendal et al., 1998). PCR fragments were separated by DGGE consisting of 8 % (v/v) polyacrylamide (ratio of acrylamide to bisacrylamide, 37:5:1) and 0·5× Tris/acetate/EDTA (pH 8·0) (TAE) buffer; 100 % denaturing acrylamide was defined as 7 M urea and 40 % formamidé. Gradients of 38–48 % were used to separate products amplified with universal primers. After migration of the PCR products at 85 V for 16 h, the gels were stained with AgNO₃ as described previously (Sanguinetti et al., 1994).

PCR on the 16S rRNA gene of strain MucT was performed with universal primers 11f and 1510r (Lane, 1991). The following PCR programme was used: 94°C for 5 min; 40 cycles consisting of 94°C for 1 min 30 s, 48°C for 30 s, and 68°C for 1 min 30 s; and finally 68°C for 7 min. PCR products were purified and concentrated with the Qiaquick PCR purification kit (Qiagen) according to the manufacturer’s instructions. The purified 16S rRNA gene product was sequenced on both strands using infrared Dye 41-labelled primers 7f, 342r, 805f, 1100r and 1510r (Lane, 1991). One extra primer, Muc1 (5’-GGA AAC CCT GAT GGT GCG-3’), which targets a 339 bp specific region of the 16S rRNA gene sequence of strain MucT, was designed to obtain unambiguous results. Sequences were automatically analysed on a LI-COR DNA sequencer 4000L and corrected manually. Pairwise sequence alignment was performed with the program DNASTAR. The 16S rRNA gene sequence was compared to sequences from GenBank using the program BLASTN 2.0, available through the National Centre for Biotechnology Information (NCBI) website (http://www.ncbi.nlm.nih.gov/blast/). The ARB software package was used to align cloned sequences and 16S rRNA gene sequences of nearest relatives (Strunk & Ludwig, 1995). A phylogenetic tree was constructed with ARB using the neighbour-joining method. The distance matrix used in the neighbour-joining method included stretches of sequence corresponding to Escherichia coli positions 63–1491.

Substrates and fermentation product concentrations were determined in the culture before and after growth using HPLC and GC methods as described previously (Stams et al., 1993).

The use of serial dilution in an anaerobic medium containing mucin as energy source led to the isolation of a predominant mucin-degrading bacterium from the human faecal sample. As in all MPN studies, the low-dilution cultures presumably gave rise to the fastest-growing organisms under the given culture conditions, whereas the high-dilution cultures supported growth of the numerically dominant organisms. In our study, a single band generated by PCR-DGGE (Fig. 1) dominated all dilutions, indicating that there was one predominant mucin-degrading bacterium and that this was also the fastest-growing mucin-degrading organism. The MPN of mucin-degrading organisms present in this faecal sample was estimated at $2 \times 10^9$ (g faeces)$^{-1}$. No growth was observed in the uninoculated mucin medium, indicating the sterility of the mucin, nor in the medium inoculated with faecal cultures presumably gave rise to the fastest-growing organisms under the given culture conditions, whereas the high-dilution cultures supported growth of the numerically dominant organisms. In our study, a single band generated by PCR-DGGE (Fig. 1) dominated all dilutions, indicating that there was one predominant mucin-degrading bacterium and that this was also the fastest-growing mucin-degrading organism. The MPN of mucin-degrading organisms present in this faecal sample was estimated at $2 \times 10^9$ (g faeces)$^{-1}$. No growth was observed in the uninoculated mucin medium, indicating the sterility of the mucin, nor in the medium inoculated with faecal cultures but not supplemented with mucin. This indicates that growth occurred solely due to utilization of mucin by the faecal bacteria.

Microscopic analysis of the enrichment dilutions revealed that an oval-shaped organism was predominant in the first dilutions of the enrichment. The 16S rRNA gene amplicons from all the dilutions of the enrichment where growth was observed ($10^{-2}$ to $10^{-9}$) were analysed by DGGE (Fig. 1, lanes 3–10). In the first dilutions, many bands were detected, but one was predominant. This band became more intense with increasing dilution and, in the highest dilution where growth occurred (10$^{-9}$), this band was almost unique in the profile. The faecal sample profile showed a band at the same position, suggesting that the enriched micro-organism...
containing this 16S rRNA gene is the same as the one present in the faecal sample and represents at least 1% of the total intestinal bacterial community (Muyzer et al., 1993). The mucin-degrading bacterium, whose 16S rRNA gene corresponded to the major DGGE band present in the enrichment (lane 10), was cultured from the highest dilution with growth ($10^{-2}$) using the soft agar technique. After 6 days, the most predominant colony type, white, was grown in the mucin medium. White colonies were picked, diluted in the mucin medium and transferred into soft agar mucin medium. This purification step was repeated twice. Finally, a single type of white colony appeared. Phase contrast microscopy revealed only one morphotype, and the DGGE profile of the 16S rRNA gene amplicon of the strain isolated showed the presence of a unique band corresponding to the major band present in the enrichment (Fig. 1, lane 1). The pure culture was designated strain MucT.

An almost-complete 16S rRNA gene sequence of strain MucT was determined (1433 bp). The most similar 16S rRNA gene sequences, which were derived from studies of uncultured colonic bacteria [HuCA18 and HuCC13 (Hold et al., 2002) and L10-6 (Salzman et al., 2002)] were 99% identical to strain MucT. The cultured bacterium most closely related to strain MucT was Verrucomicrobi um spinosum, and this was only distantly related (92%). A phylogenetic dendrogram based on 16S rRNA gene sequences was constructed; it revealed that strain MucT is related to the genera Prosthecobacter and Verrucomicrobi um, which are members of the order Verrucomicrobiales. Thus, MucT belongs to the division Verrucomicrobia and the class Verrucomicrobi a (Fig. 2). The majority of the members of this new division are clones and only a few are cultivated bacteria: a single genus, Verrucomicrobi um (Schlesner, 1987), after which the division was named; four species of the genus Prosthecobacter (Prosthecobacter debontii, Prosthecobacter dejongei, Prosthecobacter fusiformis and Prosthecobacter vanneervenii) (Staley et al., 1976; Hedlund et al., 1997); Opitutus terrae (Chin et al., 2001) and other ultramicrobacteria (Janssen et al., 1997); and the recently described species Victivallis vadensis, the first member of the division Verrucomicrobi um to be isolated from the GI tract (Zoetendal et al., 2003). Members of the ‘Verrucomicrobi um’ group of bacteria have also been identified in low numbers in human faeces-derived 16S rRNA gene libraries (Wilson & Blitchington, 1996; Suau et al., 1999; Hold et al., 2002). The division Verrucomicrobi um is composed of five subdivisions (Hugenholtz et al., 1998) and the genera Prosthecobacter and Verrucomicrobi um are part of subdivision 1. Bacteria from these two genera were isolated from freshwater habitats and are both Gram-negative, aerobic and heavily fimbriated. Cells of Verrucomicrobi um have many prosthecae, whereas cells of Prosthecobacter have only a single prosthecae. Strain MucT shares some common characteristics; for example, it is Gram-negative and can grow without vitamins. However, strain MucT is distinct among the members of subdivision 1 in that it is strictly anaerobic and cells are oval-shaped in contrast to the other members (see Table 1). On the basis of a phylogenetic analysis, strain MucT does not belong to the Verrucomicrobi um or Prosthecobacter clusters and

![Fig. 2. Phylogenetic tree showing the position of strain MucT among selected clones or strains belonging to the division Verrucomicrobi um. The tree, which was rooted using Escherichia coli as the outgroup, was generated by the neighbour-joining method. The numbers before the interior branch points indicate the five major lineages within the division Verrucomicrobi um as proposed by Hugenholtz et al. (1998). Bar, 10% sequence divergence.](image-url)
Growth was determined by measuring OD600 in basal medium supplemented with the appropriate substrate (10 mM final concentration; see text). Strain MucT represents a novel species in a new genus belonging to subdivision 1 of the division Verrucomicrobia.

<table>
<thead>
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<th>Characteristic</th>
<th>1</th>
<th>2</th>
<th>3</th>
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<tr>
<td>Cell morphology</td>
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<td>Fusiform rod-shaped</td>
<td>Fusiform rod-shaped</td>
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<td>Cell size (µm)</td>
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<td>0.8–1.0 × 1.0–3.8</td>
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<tr>
<td>Tolerance to oxygen</td>
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<td>+</td>
<td>+</td>
</tr>
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<td>Temperature range for growth (°C)</td>
<td>20–40</td>
<td>26–34</td>
<td>1–40</td>
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<tr>
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<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Fimbriae</td>
<td>–</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Capsule</td>
<td>+</td>
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</tr>
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<td>Growth on*:</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Glucose</td>
<td>w†</td>
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<td>+</td>
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<td>Fructose</td>
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<td>+</td>
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<tr>
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<td>N-Acetylglucosamine</td>
<td>w†</td>
<td>+</td>
<td>+/–‡</td>
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<tr>
<td>N-Acetyl-galactosamine</td>
<td>w†</td>
<td>ND</td>
<td>ND</td>
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<tr>
<td>Mucin</td>
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<td>ND</td>
<td>ND</td>
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<td>DNA G+C content (mol%)</td>
<td>47.6</td>
<td>57.9–59.3</td>
<td>54.6–60.1</td>
</tr>
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</table>

*Growth was determined by measuring OD600 in basal medium supplemented with the appropriate substrate (10 mM final concentration; see text).
†When a protein source is provided (peptone, yeast extract, tryptone and casitone at final concentration of 2 g l−1 each).
‡Depends on species.

should be considered as a separate phylogenetic branch. It is therefore proposed that strain MucT represents a novel species in a new genus belonging to subdivision 1 of the Verrucomicrobia.

Strain MucT is an obligate chemo-organotroph. No growth was detected on basal medium supplemented with vitamins and purged with H2/CO2 (80:20). Rumen fluid and vitamins were not required for growth on mucin and, for further characterization of the strain, they were not added to the mucin medium. Growth was not observed in mucin medium in the absence of a reducing agent, as indicated by the pink colour of the medium, demonstrating the strict anaerobic nature of strain MucT. The isolate could grow between 20 and 40 °C, with optimum growth at 37 °C. The optimum pH for growth was 6-5. No growth was observed below pH 5.5 or above pH 8. The doubling time of the strain was approximately 1-5 h in mucin medium.

No growth was observed on glucose, cellobiose, lactose, galactose, xylose, fucose, rhamnose, maltose, succinate, acetate, fumarate, butyrate, lactate, casitone (0.5%), Case amino acids (0.5%), tryptophan (0.5%), peptone (0.5%), yeast extract (0.5%), proline, glycine, aspartate, serine, threonine, glutamate, alanine, N-acetylgalactosamine or N-acetyl-galactosamine after 4 weeks incubation. Gastric mucin isolated from human intestinal cell lines and adapted on 10−5 M methotrexate (HT-29 MTX) to produce a high amount of mucin (Lesuffleur et al., 1990) resulted in growth of strain MucT to the same density as with hog gastric mucin. Strain MucT could also grow on rich media, Columbia and BHI, but with a final optical density of half that of the mucin medium. No growth was observed on rich WC anaerobe broth. When peptone, yeast extract, tryptone and casitone (each at 0.5 g l−1) were added to the basal medium, growth was observed only when the sugars N-acetylgalactosamine, N-acetyl-galactosamine and glucose were added, although bacterial growth was less than a quarter of that on mucin medium. When the solution containing the nitrogen source was not added to the basal medium supplemented with mucin, strain MucT could grow to the same density, indicating that the isolate utilized mucin as both carbon and nitrogen source. Strain MucT did not produce H2, but acetate, propionate and ethanol were formed from mucin fermentation. No sulfides were produced. Sulfates were released during fermentation of mucin (0.71 mM), demonstrating sulfatase activity. It is presumed that the limited ability of strain MucT to grow on the many substrates tested may be due to the complex structure of the mucin, which is composed of both oligosaccharides and amino acids, and that strain MucT requires a combination of all these components to reach a high density. It is likely that strain MucT produces one or more

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appropriate glycosidases to degrade the N-acetylgalactosamine and N-acetylglucosamine components from mucin, which might be exposed in the terminal part, and to use them as growth substrates.

Cells of strain Muc\(^T\) were oval-shaped (Fig. 3a), showing a different size depending on the medium. In mucin medium, strain Muc\(^T\) was 640 nm in diameter and 690 nm in length and in BHI, strain Muc\(^T\) was 830 nm in diameter and 1 \(\mu\)m in length. Cells stained Gram-negative. Flagella were not seen on negatively stained EM preparations. Spore formation was never observed. In mucin medium, the organism could grow as single cells or in pairs, but rarely in chains; it often formed aggregates in which a translucent layer of material was observed between organisms. In BHI and Columbia media, this material was rarely, if ever, observed and cells occurred singly or in pairs, but rarely in groups. In basal medium, supplemented with N-acetylglucosamine or N-acetylgalactosamine, together with some sources of proteins (a combination of yeast extract, peptone, tryptone and casitone), cells occurred singly and sometimes in pairs. Cells of strain Muc\(^T\) grown in mucin medium could exclude Indian ink, which is characteristic of capsule-possessing bacteria. EM revealed the existence of filamentous structures on cells grown in mucin medium (Fig. 3b). It is assumed that these filaments are capsular polymers that are used to connect cells together. Since this aggregation is mainly observed in mucin medium, this capsule may aid in adhesion and colonization of mucin-secreting epithelia in the GI tract. On soft agar medium, colonies of strain Muc\(^T\) appeared white and were 0-7 mm in diameter.

Based on morphological, physiological and phylogenetic features of strain Muc\(^T\), a new genus, *Akkermansia*, with the type species *Akkermansia muciniphila* gen. nov., sp. nov. is proposed.

**Description of Akkermansia gen. nov.**

*Akkermansia* (Ak.ker.man’ si.a. N.L. fem. n. Akkermansia derived from Antoon Akkermans, a Dutch microbiologist recognized for his contribution to microbial ecology).


The type species is *Akkermansia muciniphila*.

**Description of Akkermansia muciniphila sp. nov.**

*Akkermansia muciniphila* (mu.ci.ni’ phi.la. N.L. neut. n. mucinum mucin; Gr. adj. philos loving; N.L. fem. adj. muciniphila mucin-loving).

Cells are oval-shaped, non-motile and stain Gram-negative. The long axis of single cells is 0.6–1.0 \(\mu\)m, depending on the substrate used. Cells occur singly, in pairs, in short chains and in aggregates. Growth occurs at 20–40 °C and pH 5.5–8.0, with optimum growth at 37 °C and pH 6.5. Strictly anaerobic. Able to grow on gastric mucin, brain–heart infusion and Columbia media, and on N-acetylglucosamine, N-acetylgalactosamine and glucose when these three sugars are in the presence of (each at 2 g l\(^{-1}\)) peptone, yeast extract, casitone and tryptone. Cellobiose, lactose, galactose, xylose, fucose, rhamnose, maltose, succinate, acetate, fumarate, lactate, casitone, Casamino acids, tryptone, peptone, yeast extract, proline, glycine, aspartate, serine, threonine and glutamate do not support growth. Capable of using mucin as carbon, energy and nitrogen source. Able to release sulfate in a free form from mucin fermentation. In mucin medium, cells are covered with filaments. Growth occurs without vitamins. Colonies appear white with a diameter of 0.7 mm in soft agar mucin medium.

The type strain is Muc\(^T\) (= ATCC BAA-835\(^T\) = CIP 107961\(^T\)), isolated from the human intestinal tract. Its DNA G+C content is 47.6 mol%.

**Acknowledgements**

The authors are grateful to E. Tosi-Couture (Institut Pasteur, Unité Toxines et Pathogénie bactériennes, Paris, France) for performing the TEM. Dr G. Huet (INSERM, U560, Lille, France) is gratefully acknowledged for the gift of the HT-29 MTX mucin. We thank E. G.
Zoetendal for discussion and for critically reading the manuscript and H. Smidt for help in the phylogenetic analysis. We thank Professor Dr H. G. Trüper for his help regarding the Latin nomenclature. This work has been carried out with the financial support of the European Community specific RTD programme ‘Quality of Life and Management of Living Resources’ research project EU & Microfunction (QLK1-2001-00135). C. M. P. was supported by the Research Council for Earth and Life Sciences (ALW) with financial aid from the Netherlands Organization for Scientific Research (NWO).

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