**Aminomonas paucivorans** gen. nov., sp. nov.,
a mesophilic, anaerobic, amino-acid-utilizing bacterium


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A novel, asaccharolytic, amino-acid-degrading bacterium, designated strain GLU-3^T, was isolated from an anaerobic lagoon of a dairy wastewater treatment plant. Strain GLU-3^T stained Gram-negative and was an obligately anaerobic, non-spore-forming, slightly curved, rod-shaped bacterium (0.3 x 4.0-6.0 μm) which existed singly or in pairs. The DNA G+C content was 43 mol%. Optimum growth occurred at 35 °C and pH 7.5 on arginine with a generation time of 16 h. Good growth was obtained on arginine, histidine, threonine and glycine. Acetate was the end-product formed from all these substrates, but in addition, a trace of formate was detected from arginine and histidine, and ornithine was produced from arginine. Strain GLU-3^T grew slowly on glutamate and produced acetate, carbon dioxide, formate, hydrogen and traces of propionate as the end-products. In syntrophic association with *Methanobacterium formicicuim*, strain GLU-3^T oxidized arginine, histidine and glutamate to give propionate as the major product; acetate, carbon dioxide and methane were also produced. Strain GLU-3^T did not degrade alanine and the branched-chain amino acids valine, leucine and isoleucine either in pure culture or in association with *M. formicicuim*. The nearest phylogenetic relative of strain GLU-3^T was the thermophile *Selenomonas acidaminovorans* (similarity value of 89.5%). As strain GLU-3^T^T is phylogenetically, physiologically and genotypically different from other amino-acid-degrading genera, it is proposed that it should be designated a new species of a new genus *Aminomonas paucivorans* gen. nov., sp. nov. (DSM 12260T).

**Keywords:** *Aminomonas paucivorans*, anaerobic bacterium, amino acid degradation, mixed culture, *Methanobacterium formicicuim*

**INTRODUCTION**

In nature, proteins are hydrolysed to peptides and amino acids and converted to methane and CO₂ by mixed bacterial populations under anaerobic conditions. A large number of peptides and amino acids can serve as carbon and energy sources for saccharolytic and asaccharolytic anaerobic bacteria (Smith & MacFarlane, 1997) and for many years, it was assumed that saccharolytic bacteria were primarily responsible for amino acid and peptide degradation (McInerney, 1988). However, studies on the rumen ecosystem have provided evidence of the importance of asaccharolytic micro-organisms in the turnover of amino acids (Chen & Russell, 1988, 1989). The descriptions of new species of obligate amino acid degraders isolated from a wide range of other ecosystems clearly indicate that their importance may have been largely unnoticed. Examples of asaccharolytic peptide/amino-acid-degrading bacteria include *Clostridium aminophilum* from the rumen (Paster et al., 1993), *Eubacterium acidaminophilum* from black anaerobic mud of a waste water ditch (Zindel et al., 1988), *Acidaminobacter hydrogenoformans* from Ems-Dollard estuary mud (Stams & Hansen, 1984), *‘Selenomonas acidaminophila’* from an anaerobic purification plant of dairy wastewater, and *Aminomonas paucivorans* from a dairy wastewater lagoon.
peptides and amino acids, we have isolated different 
(Nanninga et al., 1987) and Dethiosulfovibrio peptide-
vorans from an oilfield (Magot et al., 1997).
As part of our investigations on the degradation of 
peptides and amino acids, we have isolated different 
asaccharolytic amino-acid-degrading anaerobic bac-
teria from a treatment plant of a dairy wastewater 
aerobic lagoon of a dairy wastewater treatment plant (SantaFe de 
Bogota, Colombia) by completely filling a sterile bottle with 
a sediment slurry. The in situ temperature at the sample 
collection point was 24 °C and the pH was 6.8. Samples were 
transported to the laboratory and maintained at 4 °C until 
used.

Source of strains. Methanobacterium formicicum DSM 1535, 
S. acidaminovorans DSM 6589T and S. acidaminiphila 
DSM 3853T were purchased from DSMZ (Deutsche 
Sammlung von Mikroorganismen und Zellkulturen, 
Braunschweig, Germany).

Media and cultivation conditions. The Hungate technique 
(Hungate, 1969) was used throughout these studies. The 
basal medium used for isolation and cultivation of strain 
GLU-3T contained (1-l): 0.2 g KH2PO4; 0.3 g NH4Cl; 1 g 
NaCl; 0.4 g MgCl2; 6H2O; 0.5 g KCl; 0.15 g CaCl2; 2H2O; 
1 ml trace element solution (Imhoff-Stucke & Pfenning, 
1983); 10 ml 0.1% resazurin. The pH was adjusted to 7.2 
with 10 M KOH. The medium was boiled under a stream of 
O2-free N2 gas and cooled to room temperature. Aliquots 
(5 ml) of the medium were distributed into Hungate tubes 
under a stream of O2-free N2 gas. The gas phase was 
subsequently replaced with N2/CO2 (80:20). Prior to in-
oculation, 0.15 ml 2% Na2S·9H2O; 0.25 ml 10% NaHCO3 
and 0.05 ml Balch vitamin solution (Balch et al., 1979) 
were injected into each tube. Amino acids were added from sterile 
(heat or filter-sterilized) anaerobic stock solutions. The 
ability of the strain to grow aerobically was tested in the 
same growth medium but that was prepared aerobically and 
lacked Na2S·9H2O.

Isolation and growth. For isolation, the sludge sample was 
pulverized in an anaerobic chamber using a hand-held 
homogenizer. A tenfold serial dilution was prepared and 
inoculated into the Hungate tubes containing 5 ml basal 
medium with glutamate and yeast extract at final concen-
trations of 10 mM and 0.2%, respectively. Incubations were 
performed at 37 °C for up to 3-4 weeks, after which serial 
dilutions were prepared from the dilution which showed 
growth and inoculated into Hungate roll tubes containing 
basal medium, yeast extract (0.2%), glutamate (10 mM) and 
agar (2%). Axenic cultures were obtained by the repeated 
use of the Hungate roll-tube method.

Microscopy. Cell morphology and motility were determined 
using a Nikon phase-contrast microscope. Cells from 
cultures grown under different conditions and with different 
substrates were examined for the presence of spores. Thin 
sections and negative-staining of exponentially growing cells 
were prepared and examined as previously described 
(Fardeau et al., 1997).

Determination of pH, temperature and NaCl requirements 
for growth. All experiments were conducted in duplicate in 
basal medium containing 10 mM arginine and 0.2% yeast 
extract unless otherwise indicated. The isolate was sub-
cultured at least once under the same experimental con-
ditions. For pH studies, the medium was adjusted with 
sterile anaerobically prepared stock solutions of HCl (1M), 
NaHCO3 (10%) or Na2CO3 (10%) to give a pH of 5.7-9.2. 
The temperature range for growth was determined at 
18-55 °C. For studies on NaCl requirements, NaCl was 
weighed directly into tubes for concentrations higher than 
1% NaCl and the medium was dispensed into the tubes as 
described above. For concentrations lower than 1%, 
different amounts of a stock solution of NaCl (10%) were 
inhaled into predispensed medium to give the required 
concentrations.

Substrate utilization tests. The isolate was subcultured at 
least once under the same experimental conditions. For 
substrate utilization studies, 0.2% yeast extract was added 
to the basal medium. l-Amino acids, organic acids and 
sugars were tested at a final concentration of 10 mM. 
Mesaconate, β-methylaspartate, oxaloacetate, glycerol 
and ethanol were added to a final concentration of 5 mM. 
Biotypase, peptone, Casamino acids, gelatin and casein 
were tested at a final concentration of 0.5% (w/v). The 
electron acceptors tested were: thiosulfate (10 mM), sulfate 
(10 mM), elemental sulfur (2%), sulfate (2 mM), sodium 
fumarate (20 mM) and nitrate (10 mM).

Mixed culture experiments. To establish whether strain 
GLU-3T could benefit in the presence of hydrogen 
scavenging bacteria, a mixed culture with the hydrogeno-

trophic methanogen, M. formicicum was prepared. M. 
formicicum was grown under a H2/CO2 (80:20) atmosphere 
in basal medium containing 0.1% yeast extract, 10 mM 
acetate and 30 mM formate. Strain GLU-3T was grown in 
the basal medium containing 10 mM arginine and 0.2% 
yeast extract. A 10% inoculum from each of the cultures 
was injected into tubes containing 5 ml basal medium, 0.2% 

yeast extract and one of a selected amino acid (alanine, 
glutamate, valine, isoleucine, leucine, glycine, histidine, 

threonine or arginine) or organic acid (pyruvate, fumarate, 
succinate, α-ketoglutarate or malate) at a concentration of 
10 mM. At the end of the incubation period of 2-3 weeks at 
37 °C, the concentrations of amino acids utilized and the 
end-products formed were determined. Basal medium 
amended with only 0.2% yeast extract was used as a control.

Growth of strain GLU-3T under an atmosphere. 

The effect of a hydrogen atmosphere (H2/CO2; 80:20) on growth of 
strain GLU-3T was tested using basal medium containing 
0.2% yeast extract and 10 mM arginine, glutamate, threo-
inine, histidine or glycine.

Analytical techniques. Growth was measured by inserting 
growth tubes directly into a model UV-160A spectro-
photometer (Shimadzu) and measuring OD590. The fer-
mentation end-products were determined as described by 
Fardeau et al. (1993). Amino acid concentrations were 
determined by liquid chromatography (Moore et al., 1958). 
Growth and product formation were analysed after 2-3 
weeks incubation at 37 °C. Heat resistance was determined 
by incubating the culture at 80 °C for up to 20 min and at 
90 °C for up to 10 min followed by subculturing in fresh 
growth medium.

Determination of G+C content. The G+C content was 
determined by the DSMZ. The DNA was isolated and 
purified by chromatography on hydroxyapatite. The G+C 
content was determined by HPLC as described by Mesbah et al. 
(1989).
DNA extraction and amplification of 16S rRNA. *S. acidaminovorans* (Guangsheng et al., 1992) and 'S. acidaminophila' (Nanninga et al., 1987) were cultured as described previously and strain GLU-3 was grown on the basal medium containing 0.2% yeast extract and 10 mM glutamate. The DNA was extracted from the strains as described previously (Andrews & Patel, 1996; Redburn & Patel, 1993). The universal primers Fd1 and Rd1 were used to obtain a PCR product of approximately 1.5 kb, corresponding to base positions 8-1542 based on *Escherichia coli* numbering of the 16S rDNA (Van de Peer & De Wachter, 1993). A 50 µl reaction mix contained 1-20 ng genomic DNA, 1 µM each primer, 5 µl 10× buffer, 200 µM dNTP, 3.5 mM MgCl₂, and 2.5 U Taq polymerase (Promega). PCR was carried out by an initial denaturation at 94 °C for 7 min, then 29 cycles of annealing at 55 °C for 2 min, extension at 72 °C for 4 min, denaturation at 94 °C for 1 min, and finally an extension cycle of 55 °C for 2 min and 72 °C for 20 min.

**Direct sequencing of PCR products.** PCR products were purified using the QIAquick kit (Qiagen). DNA concentration of purified PCR products was estimated by comparison with the Low Mass Ladder (Gibco-BRL) on an agarose gel containing ethidium bromide. PCR products were sequenced with the ABI PRISM Dye Terminator Cycle Sequencing kit with AmpliTaq FS DNA polymerase and an ABI 373A sequencer. A 10 µl reaction mix contained 35 ng PCR product, 4 µl cycle sequencing reaction mix, 3.2 pmol primer (Andrews & Patel, 1996) and 2.5 µg BSA. Thermal cycling was carried out using a Rapid Cycler (Idaho Technology) at a temperature transition slope of 2, an initial denaturation of 94 °C for 15 s, then 25 cycles of denaturation at 94 °C for 0 s, annealing at 50 °C for 10 s and extension at 60 °C for 3 min.

**Sequence alignments and phylogenetic inferences.** The new sequence data that were generated were aligned and checked for accuracy manually with the alignment editor ae2 (Maidak et al., 1996). The program BLAST (Altschul et al., 1997) was initially used against the GenBank database to determine if any newly released, closely related sequences existed in the database. For analysis, the sequences of *Anaerobaculum thermoterrenum* and *D. peptidovorans* were extracted from GenBank (nos U5071 and U52817, respectively) and manually aligned with the prealigned sequences obtained from the Ribosomal Database Project (Maidak et al., 1996). Pairwise evolutionary distances based on 1151 unambiguous nucleotides were computed using DNADIST (Jukes & Cantor option) and neighbour-joining programs that form part of the PHYLIP package (Felsenstein, 1993). TREECON was used extensively for bootstrap analysis (500 data sets) (Van de Peer & De Wachter, 1993).

**RESULTS**

**Isolation**

The highest dilution showing growth on glutamate after 3-4 weeks incubation at 37 °C was used for further isolation. This positive culture was serially diluted and inoculated into roll tubes. Only one type of colony (small, round, whitish with smooth edges) developed after 4 weeks incubation at 37 °C. Several colonies were picked and a pure culture derived from one colony was selected, designated strain GLU-3T and characterized further. Strain GLU-3T grew slowly in medium containing glutamate and yeast extract, a medium used to initiate isolations. Subsequent experiments revealed that strain GLU-3T grew much better with arginine as the energy source, and therefore this...
### Table 1. Fermentation of substrates that supported growth of strain GLU-3T in pure culture

Results were recorded after 3 weeks incubation at 37 °C. The basal medium contained 0.2% yeast extract.

<table>
<thead>
<tr>
<th>Substrate*</th>
<th>Amino acid degraded (mM)</th>
<th>Products formed (mM)†</th>
<th>ΔOD₅₆₀</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Acetate</td>
<td>Propionate</td>
<td>Formate</td>
</tr>
<tr>
<td>Arginine</td>
<td>6·1</td>
<td>1·0</td>
<td>0·0</td>
</tr>
<tr>
<td>Histidine</td>
<td>ND</td>
<td>9·2</td>
<td>0·0</td>
</tr>
<tr>
<td>Glutamate</td>
<td>7·7</td>
<td>7·5</td>
<td>2·0</td>
</tr>
<tr>
<td>Threonine</td>
<td>ND</td>
<td>9·4</td>
<td>0·0</td>
</tr>
<tr>
<td>Glycine</td>
<td>3·0</td>
<td>4·5</td>
<td>0·0</td>
</tr>
</tbody>
</table>

* Poor growth with Casamino acids, peptone and cysteine was observed and acetate levels above those of the control were present.
† Tubes containing basal medium with 0.2% yeast extract but lacking substrates were used as control. All values were corrected for a small amount of acetate (2 mM) formed in the control tubes. ND, Not determined.

### Table 2. Amino acids used by strain GLU-3T in mixed culture and under a hydrogen atmosphere

H₂/CO₂ (80:20) was used at a pressure of 2 bar. Results were recorded after 3 weeks incubation at 37 °C. The basal medium contained 0.2% yeast extract and substrates at concentrations indicated.

<table>
<thead>
<tr>
<th>Culture conditions</th>
<th>Amino acid degraded (mM)</th>
<th>Products formed*</th>
<th>ΔOD₅₆₀</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Acetate</td>
<td>Propionate</td>
<td>Formate</td>
</tr>
<tr>
<td>Arginine + M. formicicum</td>
<td>8·9</td>
<td>2·2</td>
<td>6·0</td>
</tr>
<tr>
<td>Arginine + H₂</td>
<td>ND</td>
<td>1·0</td>
<td>0·0</td>
</tr>
<tr>
<td>Histidine + M. formicicum</td>
<td>6·2</td>
<td>4·0</td>
<td>3·3</td>
</tr>
<tr>
<td>Histidine + H₂</td>
<td>ND</td>
<td>5·0</td>
<td>0·0</td>
</tr>
<tr>
<td>Glutamate + M. formicicum</td>
<td>7·0</td>
<td>1·5</td>
<td>7·0</td>
</tr>
<tr>
<td>Glutamate + H₂</td>
<td>6·0</td>
<td>6·5</td>
<td>0·0</td>
</tr>
</tbody>
</table>

* Tubes containing basal medium with 0.2% yeast extract but lacking substrates were used as control. All values were corrected for the small amounts of acetate (2 mM) formed in the control tubes. Acetate, propionate, formate, ornithine and H₂ were determined in mM and CH₄ was determined in mmol.
ND, Not determined.

Amino acid was used for determining optimal growth conditions for strain GLU-3T.

**Morphology**

Cells of strain GLU-3T stained Gram-negative. They were slightly curved rods and measured 4·0–6·0 by 0·3–0·4 μm when grown on a medium containing arginine and yeast extract (Fig. 1a, b). Motility was not observed and electron microscopy of negatively stained cells revealed the absence of flagella. Spores were not observed from cells grown under various conditions and cells were not heat resistant. Ultrathin sections of strain GLU-3T revealed a cytoplasmic membrane and a complex cell wall layer characteristic of a Gram-negative cell (Fig. 1c).

**Growth, metabolic properties and G + C content**

Strain GLU-3T was a strictly anaerobic, chemoheterotrophic bacterium. The optimal growth temperature was 35 °C and the temperature range for

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growth was between 20 and 40 °C; no growth was observed at 18 and 42 °C. Strain GLU-3T did not require NaCl for growth, but tolerated up to 2.0% NaCl, with optimum growth occurring in the presence of 0.05–0.50% NaCl. The pH range for growth was 6.7–8.3 with an optimum around 7.5.

The G+C content of strain GLU-3T was 43 mol%.

**Substrates used for growth**

Strain GLU-3T did not grow in basal medium without yeast extract and 0.2% yeast extract was routinely used for substrate utilization tests. Strain GLU-3T fermented arginine, histidine, glutamate, threonine and glycine (Table 1). Arginine was fermented to ornithine, acetate and formate, histidine was fermented to acetate and formate, and glutamate was fermented to acetate and, to a minor extent, propionate and formate. Threonine and glycine were degraded to acetate. Acetate was also the major end-product of metabolism from Casamino acids, peptone and cysteine, but they were poorly used. As we have used 0.2% yeast extract to test for the utilization of amino acids, we cannot rule out the possibility that degradation occurs via the Stickland reaction, especially in the case of glycine. The following substrates were not used by strain GLU-3T: leucine, isoleucine, valine, alanine, lysine, proline, serine, methionine, asparagine, tyrosine, phenylalanine, aspartate, gelatin,
casein, glucose, saccharose, ribose, xylose, cellobiose, melibiose, maltose, galactose, mannose, arabinose, rhamnose, lactose, sorbose, mannitol, acetate, propionate, butyrate, valerate, fumarate, maleate, succinate, pyruvate, lactate, citrate, α-ketoglutarate, mesaconate, β-methylaspartate, oxaloacetate, glycerol and ethanol. Sulfate, thiosulfate, elemental sulfur, sulfite, nitrate and fumarate were not utilized as electron acceptors.

**Amino acid degradation in mixed cultures**

A mixed culture of strain GLU-3T and *M. formicicum* was unable to extend the range of utilisable substrates. However, compared to pure cultures, association of strain GLU-3T with *M. formicicum* resulted in a shift in end-product formation during degradation of arginine, histidine and glutamate with methane and propionate as the major fermentation products (Table 2). In contrast to the pure culture, ornithine did not accumulate during arginine degradation by the mixed culture. Methane was not detectable in mixed culture with glycine and threonine, and the same end-product profile as for a pure culture was observed (data not shown).

**Growth under hydrogen atmosphere**

Cultivation of strain GLU-3T under a H₂/CO₂ (80:20) atmosphere did not affect the fermentation of arginine, histidine, glutamate, threonine or glycine; the end-products were similar to those found without a hydrogen atmosphere (Table 2 and data not shown).

**16S rRNA sequence analysis**

An almost complete 16S rRNA gene sequence (1504, 1426 and 1530 nucleotides) of strain GLU-3T, *S. acidaminovorans* and *S. acidaminophila* corresponding to positions 17–1539, 14–1540 and 17–1539, respectively ([*E. coli* numbering; according to Winker & Woese (1991)]) were determined. Initial sequence comparisons indicated that strain GLU-3T was a member of the low G+C, Gram-positive branch. Several phylogenetic trees were generated using numerous combinations of sequences from the members of this branch and strain GLU-3T was consistently placed as an independent branch of descent adjacent to members of cluster V. This cluster currently comprises *Thermoanaerobacter* species, *D. peptidovorans* and *S. acidaminovorans* (Fig. 2). The closest relative of strain GLU-3T was *S. acidaminovorans* (similarity of 89.5%), which also forms an independent lineage in the vicinity of cluster V. Bootstrap analysis indicates a robust relationship between strain GLU-3T and *S. acidaminovorans* (100%). Comparison of the sequence of the 16S rRNA gene of *S. acidaminophila* with that of members of the low G+C, Gram-positive branch indicated that it clustered with *Clostridium quercicolum*, a member of cluster IX (Collins et al., 1994), with a similarity of 90%, rather than in the vicinity of cluster V, and was therefore not related to strain GLU-3T.

**DISCUSSION**

Strain GLU-3T is a Gram-negative, slightly curved, non-spore-forming, obligate amino-acid-degrading anaerobe and therefore cannot be assigned to the aminolytic members of the genus *Clostridium* which include *Clostridium sticklandii* and *Clostridium aminoophilum* (Paster et al., 1993), *Clostridium litorale*, *Clostridium pascui* and *Clostridium hydroxybenzoicum* (Fendrich et al., 1990; Wilde et al., 1997; Zhang et al., 1994) and *Clostridium acetireducens* (Örlygsson et al., 1996). These species are Gram-positive, spore-forming, straight rods. This conclusion is confirmed by phylogenetic analysis which places strain GLU-3T in the low-G+C-containing phylum in the vicinity of *Thermoanaerobacter* (cluster V), *S. acidaminovorans* and *D. peptidovorans*, with *S. acidaminovorans* being its closest relative. Like strain GLU-3T, *Thermoanaerobacter* species, *S. acidaminovorans* and *D. peptidovorans* use amino acids. However, *Thermoanaerobacter* species and *S. acidaminovorans* are thermophiles which also utilize carbohydrates, but in addition, *D. peptidovorans*, uses thiosulfate or elemental sulfur as the electron acceptor and strain GLU-3T does not.

Very little is understood about amino-acid-degrading pathways operating in obligately aminolytic anaerobes. Strain GLU-3T fermented histidine to acetate and formate. This fermentative pathway, albeit poorly studied, is similar to that used by aerobic bacteria as suggested by McSweeny et al. (1993) and may also be operating in strain GLU-3T. Strain GLU-3T degraded arginine to ornithine and this characteristic is similar to that reported for *S. acidaminovorans* and *Synergistes jonesii* strain 78-1 (McSweeny et al., 1993). This suggests that the arginine deaminase pathway may be operational in strain GLU-3T.

In pure culture, strain GLU-3T fermented glutamate to acetate, formate and trace amounts of propionate, a property that closely resembles *A. hydrogenoformans* in pure culture (Stams & Hansen, 1984). This trait differentiates it from other amino-acid-degrading members of the genera *Acidaminococcus*, *Peptostreptococcus*, *Fusobacterium* and *Clostridium* (Rogosa, 1969; Barker, 1981; Wilde et al., 1997), which produce acetate, butyrate, carbon dioxide, ammonium and hydrogen. In contrast to strain GLU-3T, *S. acidaminovorans* (Guangsheng et al., 1992) and *S. acidaminophila* (Nanninga et al., 1987) produce propionate as a major end-product from glutamate fermentation. Acetate was the only fatty acid produced from threonine and glycine fermentation. In the case of threonine degradation, this end-product is unusual as most species that degrade threonine, with the exception of a few examples such as *C. sticklandii* and *Clostridium subterminale* (Barker, 1981), are known to produce acetate and propionate.
Degradation of amino acids in mixed culture

An increase in the range of amino acids utilized was not observed in mixed culture with *M. formicicum*, in contrast to the situation observed for *A. hydrogenoformans*, *S. acidaminovorans* and *E. acidaminophilum* (Zindel et al., 1988). However, the metabolism of all the amino acids used by strain GLU-3T, except threonine and glycine, is clearly influenced by the presence of the hydrogenotrophic methanogen. No increase in acetate production, no change in the end-product profile, and the lack of methane production in the presence of the hydrogen scavenger suggest that threonine and glycine degradation occurs via a reductive rather than an oxidative process. In addition, a hydrogen gas phase did not alter the growth of strain GLU-3T on these two substrates further strengthening our hypothesis.

Co-culture of strain GLU-3T with *M. formicicum* shifted the metabolic end-products of arginine, histidine and glutamate degradation from acetate to propionate as the major end-product. This trait is similar to that of *A. hydrogenoformans* when it is co-cultured on glutamate and histidine with a hydrogen scavenger (Stams & Hansen, 1984). This may occur because the partial pressure of H₂ is reduced due to consumption by the hydrogen scavenger, with propionate production becoming thermodynamically more favourable (McInerney, 1988).

In spite of observed similarities between strain GLU-3T and *A. hydrogenoformans* with respect to glutamate metabolism, the latter is inhibited by a hydrogen atmosphere when grown on this amino acid. In addition, there are marked differences in the amino acids utilized. For example, *A. hydrogenoformans* utilized arginine and threonine only in mixed culture with hydrogenotrophic bacteria (Stams & Hansen, 1984), whereas strain GLU-3T could ferment both these amino acids in pure culture. In addition, unlike *A. hydrogenoformans*, strain GLU-3T did not utilize alanine, valine, leucine or isoleucine in the presence of hydrogen scavenging bacteria.

Because of the marked phenotypic and phylogenetic differences of strain GLU-3T from other amino acid degraders, we propose that it be assigned as a new species of a new genus *Aminomonas paucivorans* gen. nov., sp. nov.

Description of *Aminomonas paucivorans* sp. nov.

*Aminomonas paucivorans* (pau'ci.vor'ans. L. adj. paucus few, little; L. pres. part. vorans devouring, digesting; L. paucipers digesting little).

Slightly curved, rod-shaped bacterium (0.3 by 40–60 μm) which occurs singly or in pairs. Acetogenic, non-spore-forming. Strictly anaerobic. Colonies (up to 1.0 mm) are round, smooth and white. Mesophilic. Optimal growth temperature 35 ºC, range for growth 20–40 ºC; pH range of 6.7–8.3, optimum pH 7.5; does not require NaCl for growth but tolerates up to 2.0% with optimum growth occurring between 0.05–0.50% NaCl. Yeast extract is required for growth. Ferments arginine, threonine, glutamate, histidine and glycine. In mixed culture with *M. formicicum* on glutamate, arginine and histidine, a shift of end-products from acetate to propionate as the major end-product occurs. No growth was observed on carbohydrates, gelatin, casein, pyruvate, succinate, malate, fumarate, α-ketoglutarate, mesaconate, β-methylaspartate, oxaloacetate, glycerol, ethanol, acetate, propionate, butyrate, lactate, citrate, leucine, lysine, alanine, valine, proline, serine, methionine, asparagine, phenylalanine and aspartate. Sulfate, thiosulfate, elemental sulfur, sulfate, nitrate and fumarate were not utilized as electron acceptors. The G + C content of the DNA is 43 mol%.

ACKNOWLEDGEMENTS

We thank N. Zylber (BIP-CNRS) for amino acid analysis. This work was supported by grants from French Foreign Office (PCP) and Instituto Colombiano para el Desarrollo de la Ciencia y la Tecnología (COLCIENCIAS) (to S. B.) and in part from the Australian Research Council (to B. K. C. P.). We would like to thank Professor Trüper for having brought the bacterial nomenclature rules to our attention via the peer review process.

REFERENCES


