Gluconacetobacter entanii sp. nov., isolated from submerged high-acid industrial vinegar fermentations

Gerd Schüller, Christian Hertel and Walter P. Hammes

Acetic acid bacteria have been isolated from submerged high-acid spirit vinegar fermentations in the Southern part of Germany. Four strains (LTH 4560T, LTH 4341, LTH 4551 and LTH 4637) were characterized in more detail and it was revealed that they have in common certain properties such as requirement of acetic acid, ethanol and glucose for growth, and no over-oxidation of acetate. Growth occurs only at total concentrations (sum of acetic acid and ethanol) exceeding 60%. A method for their preservation was developed. Comparative analysis of the 16S rRNA revealed sequence similarities of >99% between strain LTH 4560T and the type strains of the related species Gluconacetobacter hansenii. However, low levels of DNA relatedness (<41%) were determined in DNA–DNA similarity studies. In addition, specific physiological characteristics permitted a clear identification of the strains within established species of acetic acid bacteria. The strains could also be differentiated on the basis of the distribution of IS element 1031 C within the chromosome. Based on these results, the new species Gluconacetobacter entanii sp. nov. is proposed for strain LTH 4560T (DSM 13536T). A 16S-rRNA-targeted oligonucleotide probe was constructed that was specific for G. entanii, and the phylogenetic position of the new species was derived from a 16S-rRNA-based tree.

Keywords: Gluconacetobacter entanii sp. nov., acetic acid bacteria, spirit vinegar fermentation

INTRODUCTION

The production of high-acid vinegar has been enabled by the use of submerged fermentation processes in so-called acetators. Depending on the strain of acetic acid bacteria and the nature of the fermentation substrate, final acetic acid concentrations of up to 15–16% can be achieved. In contrast to the high standard of the fermentation technology, the microbiology of vinegar fermentation is well below the standard of that of dairy or meat fermentation, for example. When vinegar fermentations have to be started or a breakdown of a fermentation process requires the restart of an acetator, a microbiologically undefined inoculum is commonly used (Sokollek & Hammes, 1997). The lack of defined pure starter cultures is due to problems in isolation and conservation of acetic acid bacteria involved in high-acid vinegar production (Sievers & Teuber, 1995). After the introduction of a double-layer agar technique by Entani et al. (1985), the first isolations of acetic acid bacteria from that environment were achieved. ‘Acetobacter polyoxogenes’ was described as a new species; however, the strain was not deposited in a culture collection. Later on, Sievers et al. (1992) described Gluconacetobacter entanii sp. nov. as the dominant species of acetic acid bacteria in industrial vinegar fermenters in central Europe. The authors reported that with the exception of the type strain of G. europaeus, the acetic acid bacteria isolated from high-acid submerged vinegar fermentations could not be preserved. Recently, preservation methods were described to improve the handling of the acetic acid bacteria outside of the fermentation process (Sokollek et al., 1998a). In these studies, strains were isolated from running spirit vinegar fermentations which could...
not be allotted to any species of the acetic acid bacteria, thus indicating that in addition to *G. europaeus* other species are involved in the fermentation process. It is the purpose of this communication to describe the isolation and characterization of a new *Gluconacetobacter* species involved in high-acid spirit vinegar fermentations.

**METHODS**

**Bacterial strains and growth conditions.** The strains investigated were isolates from running acetators. Isolates LTH 4341 and LTH 4551 were obtained by Sokolke *et al.* (1998a). Strains LTH 4560 and LTH 4637 were isolated from spirit vinegar fermentations at total concentrations [i.e. percentage (w/v) acetic acid plus percentage (v/v) ethanol] of 14.6 and 13.4%, respectively. The type strains of the acetic acid bacteria were obtained from the DSMZ (Deutsche Sammlung von Mikroorganismen und Zellkulturen, Braunschweig, Germany). *Gluconacetobacter xylanus* was grown in YPM medium containing (l–1): 5 g yeast extract, 3 g peptone and 25 g mannitol. *Gluconacetobacter hansenii* and *Gluconacetobacter oboediens* (formerly *Acetobacter oboediens* (Yamada, 2000)] were propagated in RAE medium (4a, Sokollek & Hammes, 1997). AE medium (4a/3e) was used as growth medium for isolates from high-acid spirit vinegar fermentations. Shaking cultures on agar plates were incubated at 30 °C on a rotary shaker at 200 r.p.m. in 1 l triple-baffled Erlenmeyer flasks containing 200 ml medium. Surface cultures on agar plates were incubated at 30 °C at >95% relative humidity. Viable cell counts were determined by surface plating on AE (4a/3e) broth for the serial dilutions. Total cell counts were determined with the aid of a Thoma haemocytometer.

**Preservation of the strains.** For preservation, 200 ml AE broth (4a/3e) was inoculated with a 7-d-old colony taken from AE agar (4a/3e) and incubated on a rotary shaker for at least 5 d. At mid-exponential growth (at an OD$_{660}$ of 0.6–0.7), the concentration of acetic acid was determined by titration with 0.1 M NaOH. For each mole of acetic acid, 0.5 mol CaCO$_3$ was added to the culture broth and the incubation was continued for 2 min. The cells were immediately harvested by centrifugation at 3000 g for 4 min at 0 °C. After resuspension in 2 ml ice-cold 20% malt extract solution (Sokollek & Hammes, 1997), the suspension was poured drop wise in liquid nitrogen. The frozen culture was lyophilized and stored at −20 °C.

**Physiological and biochemical characterization.** Growth was investigated in AE broth containing ethanol and acetic acid at concentrations of 0, 1, 2, 3, 4 and 5% (v/v) and 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11 and 12% (w/v), respectively. The cultures were grown under aeration at 30 °C for 14 d. Growth on acetate at pH 2.5, utilization of lactate, and growth in the presence of 30% glucose were studied in bouillon as described by Sievers *et al.* (1992). The utilization of carbon sources was investigated in AE broth (4a/3e) by replacing glucose, ethanol or acetic acid with the substrates presented in Results. The substrates were added at a concentration of 20 g l$^{-1}$ or 20 ml l$^{-1}$, respectively. Over-oxidation and formation of 2- and 5-ketogluconic acid as well as gluconic acid were investigated in AE broth (4a/3e) containing 2% glucose at 30 °C. After 2, 7 and 14 d incubation, concentrations of 2- and 5-ketogluconic acid, glucose, gluconic acid and acetic acid were determined by HPLC as described by Stolz *et al.* (1993) with the following modifications. An Aminex cartridge cation H$^+$ and a HPX-87-H (Bio-Rad) were applied as pre-column and column, respectively. The mobile phase consisted of H$_2$SO$_4$ (0.05 M) and the flow rate was adjusted to 0.6 ml min$^{-1}$ at 60 °C. Formation of cellulose was studied in AE broth (4a/3e) at 30 °C. Quinones were determined as described previously (Sokolke *et al.*, 1998b).

**Fermentation in a pilot acetator.** The growth of the isolates was studied in a pilot acetator (Frings) as described previously (Sokollek & Hammes, 1997) with the following modification: the mash consisted of ethanol, acetic acid, tap water and 1.5 g nutrient concentrate Acetozym DS (Frings) l$^{-1}$. The total concentration was adjusted to 13.4% for strain LTH 4637 or 14.6% for LTH 4560.

**DNA isolation.** Chromosomal DNA was prepared according to the CTAB method as described by Ausubel (1994). Plasmid DNA was isolated with the aid of Nucleobond AX PC 200 (Macherey-Nagel) and Qiagen-tip 100 columns.

**Hybridization techniques.** For hybridization with oligonucleotide probes, 5 µg chromosomal DNA was denatured in 0.2 M NaOH/2 × SSC at 37 °C for 20 min and transferred to a nylon membrane (Qiabrande; Qiagen) using a dot blot apparatus (Schleicher & Schuell). Oligonucleotides were labelled with the 3’ digoxigenin oligo-labelling kit (Boehringer Mannheim). Hybridizations were carried out as recommended by the supplier. Stringent washing was performed twice in 2 × SSC/0.1% SDS for 10 min at probe-specific temperatures. The accessibility of immobilized DNA was determined by hybridization with the 16S rDNA bacterial probe EUB 338 (Amann *et al.*, 1990). The probe-specific hybridization and stringent washing temperatures were 44 and 48 °C, respectively. Detection of hybrids was performed by using a DIG luminescent detection kit (Boehringer Mannheim).

For Southern hybridization, 2 µg chromosomal DNA was digested with *Hind*III and subjected to agarose gel electrophoresis. Blotting was performed on nylon membranes (Quibrande; Qiagen) with the aid of Vaku-Blot Apparatus (Pharmacia). An 888 bp fragment of IS 1031C was labelled with DIG-dUTP by using a PCR DIG probe synthesis kit (Boehringer Mannheim), plasmid pDCB29 (Coucheron, 1993) as the template and primers IS1031AF (5’-TTTGA-ACAAATTACCCAGCMTG-3’) and IS1031BR (5’-SMYTGA-CAATGCCCTTG-3’). Hybridization was performed at 69 °C as recommended by the supplier. The membranes were washed three times for 20 min in 2 × SSC/0.1% SDS at 70 °C. Hybrids were detected with the aid of a DIG luminescent detection kit (Boehringer Mannheim).

For determination of DNA similarities, the hybridization was performed as described previously (Sokolke *et al.*, 1998b).

**Sequencing and analysis of 16S rDNA.** Amplification and sequencing of 16S rDNA were performed as described previously (Probst *et al.*, 1998). The sequence data were added to alignments of deposited complete primary structures of 16S rRNA. The phylogenetic relationship was determined by applying maximum-parsimony and maximum-likelihood approaches on data sets varying with respect to the selection of reference sequences as well as sequence positions. The respective tools of the ARB program package (Springer *et al.*, 1992) were used for alignment, selection of positions according to variability, calculation of similarities as well as tree reconstruction, evaluation and drawing.
**DNA base composition.** The DNA G+C content was determined by direct analysis of the nucleosides by HPLC according to the method of Mesbah *et al.* (1989). All enzymes for nucleoside preparations were purchased from Boehringer Mannheim. Non-methylated lambda DNA (Sigma) was used as the standard.

**RESULTS**

**Isolation of *Gluconacetobacter* strains**

Strains were isolated from running industrial spirit-vinegar fermentations in Southern Germany. The technique described by Entani *et al.* (1985) was employed. Small, round and light-grey to light-brownish colonies were obtained on AE agar (4a/3e) within 5–10 d upon incubation at 30 °C in an atmosphere of > 95% relative humidity. Each fermenter contained one predominant strain, as revealed by comparison of the plasmid profiles and IS element fingerprints. The strains could not be allotted to the type strains of acetic acid bacteria applying 16S-rRNA-targeted probes (Sokollek *et al.*, 1998a).

**Morphological and cultural characteristics**

Microscopically, the cells appeared as non-motile, and non-flagellated rods, preferentially occurring in pairs. In AE broth, the strains grew at temperatures ranging from 20 to 34 °C, with an optimum of 30 °C, and at pH values ranging between 2.2 and 3.0, with an optimum of pH 2.7. In this medium, the total concentration (acetic acid plus ethanol) was kept > 6.0 with a minimum of 4% acetic acid and a maximum of 4% ethanol. In addition, the strains could be cultivated in yeast nitrogen base (Difco) supplemented with 0.5% glucose, 3% ethanol and 4% acetic acid.

**Genotypic differentiation of the strains**

Plasmids were isolated from the strains and subjected to gel electrophoresis. Identical plasmid profiles were found for isolates LTH 4341, LTH 4551 and LTH 4637. A slightly different profile was obtained for strain LTH 4560<sup>T</sup> (Fig. 1a). The plasmid profiles were compared with those obtained from the cultures in the acetators used as the source of the isolates. Identical profiles indicated that the predominant strain of the spirit vinegar acetators had been isolated. Genomic DNA of the strains was hybridized with a poly-nucleotide probe specific for the IS element 1031C occurring in the genome of *G. xylinus* (Coucheron, 1993). As shown in Fig. 1(b), closely related IS profiles were obtained. These permitted unambiguous differentiation of the four strains.

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**Fig. 1.** (a) Electrophoretic patterns of plasmid DNAs isolated from two *G. entanii* strains. Lanes: 1, lambda DNA/HindIII marker; 2, LTH 4560<sup>T</sup>; 3, LTH 4637. (b) Distribution of IS 1031C in the genomes of four *G. entanii* strains. Total DNA was digested with HindIII, blotted and hybridized with a PCR-DIG-labelled probe of IS 1031C. Lanes: 1, LTH 4560<sup>T</sup>; 2, LTH 4637; 3, LTH 4341; 4, LTH 4551.
16S rDNA sequence analysis and probe construction

The sequence of the 16S rDNA of strain LTH 4560\textsuperscript{T} was determined and analysed by comparison with the 16S rRNA sequences of the type strains of acetic acid bacteria (Sievers \textit{et al.}, 1994a, b, 1995; Sokollek \textit{et al.}, 1998b; Boesch \textit{et al.}, 1998; Franke \textit{et al.}, 1999). A similarity of \(>99\%\) was found between the sequence of strain LTH 4560\textsuperscript{T} and that of \textit{G. hansenii}. Two single nucleotide deletions as well as 15 base changes were detected. Based on these sequence data, the phylogenetic tree depicted in Fig. 2 was constructed reflecting the position of the strains within the genera \textit{Acetobacter}, \textit{Acidomonas}, \textit{Gluconacetobacter} and \textit{Gluconobacter}. The results of the 16S rDNA sequence comparison permitted the construction of the oligonucleotide probe AI4560 (5'-GAAACCGAGGCTATAC-3') which is specific for strain LTH 4560\textsuperscript{T}. The specificity of the probe was evaluated by dot blot hybridization to genomic DNA, including that of strains LTH 4341, LTH 4551 and LTH 4637 and the closely related type strains of the \textit{Acetobacteraceae}. For hybridization and stringent washing, the probe-specific temperatures were defined as 48 and 50 °C, respectively. Probe AI4560 reacted exclusively with DNA of the four isolates.

DNA similarity

Quantitative DNA–DNA reassociation studies were performed with the chromosomal DNA of strains LTH 4560\textsuperscript{T}, LTH 4341, LTH 4551, LTH 4637, \textit{G. europaeus} DSM 6160\textsuperscript{T}, \textit{G. xylinus} DSM 6513\textsuperscript{T}, \textit{G. oboedienis} DSM 11826\textsuperscript{\textdagger} and \textit{G. hansenii} DSM 5602\textsuperscript{T}. These species were chosen on the basis of their close relatedness, which is reflected by the cluster formation as shown in Fig. 2. Hybridizations were performed using the labelled DNA of \textit{G. europaeus} DSM 6160\textsuperscript{T}, \textit{G. xylinus} DSM 6513\textsuperscript{T}, \textit{G. hansenii} DSM 5602\textsuperscript{T} and strain LTH 4560\textsuperscript{T} as probes. As shown in Table 1, the isolates exhibited levels of DNA relatedness with the type strains below 41%. The relatedness of the isolates among each other was \(\geq 89\%\).

Physiological and biochemical characterization

The taxonomically relevant physiological properties were investigated for strains LTH 4560\textsuperscript{T} and LTH 4637. As the two strains exhibited the same physiological properties, the results obtained for strain LTH 4560\textsuperscript{T} were compared with the characteristics of the type strains of the acetic acid bacteria (Table 1). Remarkably, the isolates did not grow in AE broth without acetic acid. Within the genus \textit{Gluconacetobacter}, this property has been described only for \textit{G. europaeus} and the unvalidated species ‘\textit{A. polyoxogenes}’. On the other hand, strain LTH 4560\textsuperscript{T} can be distinguished from the type strain of \textit{G. europaeus} by the inability to grow on acetate at any pH and on lactate. Furthermore, ubiquinones Q-10 and Q-9 were found to be the major and minor component of the ubiquinone system, respectively. The DNA G+C content of strain LTH 4560\textsuperscript{T} was determined to be 58 mol%.

For further physiological characterization of isolates LTH 4560\textsuperscript{T} and LTH 4637, the utilization of various carbon sources was studied in comparison with the type strains of \textit{G. europaeus} and \textit{G. hansenii}. AE broth was used as basic medium, and acetic acid, ethanol or glucose was exchanged as indicated in Table 2. The two isolates exhibited similar properties with regard to their capacity to utilize the various substrates. Strain LTH 4637 could only be distinguished from LTH 4560\textsuperscript{T} by its ability to utilize sorbitol and mannitol. None of the cultures grew on methanol, n-butanol or...
**Table 1.** Characteristics of strain LTH 4560\textsuperscript{T} permitting its differentiation from other species of the genera *Acetobacter*, *Acidomonas* and *Gluconacetobacter*

<table>
<thead>
<tr>
<th>Characteristic</th>
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<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
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<tbody>
<tr>
<td>Growth on 3% (v/v) ethanol in the presence of 4–8% (w/v) acetic acid</td>
<td>–</td>
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<td>Growth without acetic acid</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Growth on acetate at pH 2.5</td>
<td>–</td>
<td>ND</td>
<td>+</td>
<td>–</td>
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<td>–</td>
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<td>DNA–DNA homology with*:</td>
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<tr>
<td>LTH 4560\textsuperscript{T}</td>
<td>ND</td>
<td>ND</td>
<td>26</td>
<td>18</td>
<td>32</td>
<td>100</td>
<td>39</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
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<tr>
<td><em>G. xylinus</em></td>
<td>ND</td>
<td>ND</td>
<td>32</td>
<td>100</td>
<td>39</td>
<td>18</td>
<td>14</td>
<td>ND</td>
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<td><em>G. hansenii</em></td>
<td>ND</td>
<td>ND</td>
<td>16</td>
<td>14</td>
<td>17</td>
<td>41</td>
<td>100</td>
<td>ND</td>
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<tr>
<td><em>G. europaeus</em></td>
<td>ND</td>
<td>ND</td>
<td>41</td>
<td>36</td>
<td>100</td>
<td>28</td>
<td>11</td>
<td>ND</td>
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<td>Formation of ketogluconic acids from d-glucose</td>
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<tr>
<td>2-Ketogluconic acid</td>
<td>d</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>d</td>
<td>–</td>
<td>d</td>
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<td>d</td>
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<td>+</td>
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<tr>
<td>5-Ketogluconic acid</td>
<td>d</td>
<td>–</td>
<td>–</td>
<td>+</td>
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<td>d</td>
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<tr>
<td>Growth in the presence of:</td>
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<tr>
<td>Ethanol</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>(+)</td>
<td>d</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Lactate</td>
<td>+</td>
<td>(+)</td>
<td>+</td>
<td>+</td>
<td>d</td>
<td>–</td>
<td>–</td>
<td>d</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>30% (w/v) d-glucose</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>–</td>
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<td>–</td>
<td>–</td>
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<td>+</td>
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</table>

* Data obtained in this work; all other data were taken from Sievers *et al.* (1992) and Sokollek *et al.* (1998b).

**Table 2.** Utilization of carbon sources in AE (4a/3e) broth

<table>
<thead>
<tr>
<th>Carbon source</th>
<th>G. europaeus</th>
<th>G. hansenii</th>
<th>LTH 4560\textsuperscript{T}</th>
<th>LTH 4637</th>
</tr>
</thead>
<tbody>
<tr>
<td>AE (4a/3e) broth</td>
<td>+</td>
<td>w</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Without glucose</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>–</td>
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<tr>
<td>AE (4a) broth</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Without ethanol</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>–</td>
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<tr>
<td>With n-propanol</td>
<td>+</td>
<td>–</td>
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<td>–</td>
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<tr>
<td>AE (4a/3e) broth – glucose replaced by:</td>
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<tr>
<td>Fructose</td>
<td>+</td>
<td>–</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Maltose</td>
<td>+</td>
<td>w</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Sucrose</td>
<td>+</td>
<td>–</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Glycerol</td>
<td>+</td>
<td>–</td>
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<tr>
<td>Sorbitol</td>
<td>+</td>
<td>–</td>
<td>w</td>
<td>+</td>
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<tr>
<td>Mannitol</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>w</td>
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<tr>
<td>Gluconate</td>
<td>+</td>
<td>–</td>
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<td>–</td>
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<tr>
<td>Lactate</td>
<td>+</td>
<td>–</td>
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<tr>
<td>AE (3e) broth without acetic acid</td>
<td>–</td>
<td>+</td>
<td>–</td>
<td>–</td>
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<tr>
<td>AE (3e) broth – acetic acid replaced by:</td>
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<tr>
<td>Lactate</td>
<td>+</td>
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<tr>
<td>Gluconate</td>
<td>–</td>
<td>+</td>
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</table>

\(d\)-ribose. Finally, no over-oxidation of acetic acid was observed for strains LTH 4560\textsuperscript{T} and LTH 4637 in AE broth (4a/3e) containing 2% glucose. Under these conditions, the glucose was not metabolized and no gluconic acid was formed, and no growth was observed in AE broth (4a/3e) without glucose.
Performance of the isolates in pilot acetic acid fermentation

In pilot acetic acid fermentations, the strains proved highly adapted to the semi-continuous fermentation conditions and achieved productivities (η) (Sokollek & Hammes, 1997) of > 6 at total concentrations ranging between 7 and 11.0%. However, the strains isolated from four independently running spirit vinegar fermenters in Southern Germany showed different degrees of productivity (η) at high total concentrations of ethanol and acetic acid, and differed in their tolerance to high total concentrations. Strains LTH 4560T, LTH 4551, LTH 4341 and LTH 4637 achieved productivities of 4-5% acetic acid per 24 h at total concentrations of 14.6, 14.4, 14.4 and 13.4%, respectively. The fermentation characteristics (productivity and tolerance to high total concentration) were found to be identical to those of the industrial fermentations from which the strains were isolated.

Preservation of isolates from high-acid submerged fermentation

It was observed that the total counts in the fermenting spirit vinegar mash at total concentrations ranging between 7 and 14.5% did not exceed 2-4 x 10^8 cells ml⁻¹. However, the determination of the viable counts by surface plating on AE (4a/3e) agar yielded only < 1 x 10⁴ c.f.u. ml⁻¹. After the addition of CaCO₃ to the culture broth at primary isolation from the mash (total concentration 13-2%), viable counts of 5 x 10⁴ c.f.u. ml⁻¹ were achieved, for example with strain LTH 4560T. The protective effect of CaCO₃ at primary isolation is even stronger when the mash has a reduced total concentration. For example, the total cell count of an acetator culture of strain LTH 4560T running at a total concentration of 7.6% was 2-4 x 10⁸ c.f.u. ml⁻¹ and the viable count was 2.7 x 10⁷ c.f.u. ml⁻¹ (survival 22.5%). The strains were finally subjected to lyophilization as described by Sokollek & Hammes (1997) and were found to be viable upon storage for 2 months at -18 °C.

DISCUSSION

Based on the culture techniques for acetic acid bacteria described by Entani et al. (1985), it was possible to study in more detail the fermentation flora involved in high-acid spirit vinegar fermentations. With these techniques, Entani et al. (1985) characterized an organism ("Acetobacter polyoxogenes") involved in submerged vinegar fermentations in Japan. In our investigations, a homogeneity in the fermentation flora became apparent in fermenters run with spirit alcohol. The isolates were characterized on the genomic and physiological level and it was observed that they have defined properties in common: (i) growth occurs only in the presence of acetic acid, ethanol and glucose; (ii) growth occurs only at total concentrations (sum of acetic acid and ethanol) exceeding 60%; (iii) the strains do not over-oxidize acetic acid. These characteristics reveal a striking similarity between the properties of the isolates and those described for "A. polyoxogenes" (Entani et al., 1985). Sievers et al. (1992) isolated a further species from high-acid fermentation, G. europaeus. This organism can be differentiated from our isolates, which do not grow on acetic acid at pH 2.5 and do not utilize glycerol, gluconate or lactate. Furthermore, G. europaeus can over-oxidize acetic acid and grows at total concentrations below 6% (Sokollek et al., 1998a).

The study of the 16S rRNA revealed a sequence similarity of > 99% between strain LTH 4560T and G. hansenii. This level of phylogenetic relatedness, derived from the comparative rRNA sequence analysis, can only provide differentiating information and is not sufficient to be used for the definition of a species. On the other hand, the superior method for phylogenetic investigations at and below the species level is the quantitative hybridization analysis of genomic DNA (Stackebrand & Goebl, 1994). The DNA-DNA hybridization data clearly indicated that the isolates are new species in the family Acetobacteraceae. The relatedness was 28% between strain LTH 4560T and G. europaeus, and 41% between strain LTH 4560T and G. hansenii. The strains had a DNA homology of > 89% among themselves. From the 16S-rRNA-based tree of relatedness of acetic acid bacteria (Fig. 2) it is evident that the isolates cluster in the G. xylinus/hansenii group. The use of the genus description Gluconacetobacter instead of Acetobacter is consistent with the validated proposal of Yamada et al. (1997, 1998). The detected sequence differences permitted the construction of a specific probe to clearly identify Gluconacetobacter entanii. The data obtained reveal that the strains belong to a new species. We propose the name Gluconacetobacter entanii, with strain LTH 4560T as the type strain.

The four isolates exhibit a very high degree of similarity; for example, the strains revealed identical plasmid profiles and even the profile of strain LTH 4560T shows only a slight modification in this property. A clear differentiation of the strains is possible when the distribution patterns of insertion element 1031C (Coucheron, 1993) are taken into consideration. These fingerprints are closely related but are sufficiently different to permit a strain identification. These findings can be explained on the basis of the practice of the production of high-acid vinegars in acetators. The primary inocula may be obtained from one supplier, and it is a common practice that the producers support each other by making available inocula from running acetators. Therefore, it appears likely that one competitive strain of this species (G. entanii) had developed which may have changed its pattern of IS elements during continuous propagation in a defined factory. Depending on the specific conditions employed in different factories, adaptation takes place by random mutation and selection, which is enhanced by the high content of IS elements. This conclusion is supported by the observation that the various strains exhibited...
Gluconacetobacter entanii sp. nov.

Gluconacetobacter entanii (en.ta’.ni. i. L. gen. entanii) of Entani, in honour of Etsuzu Entani, a Japanese microbiologist who performed the crucial experiments on isolation and propagation of spirit vinegar bacteria.

Cells are Gram-negative, ellipsoidal to rod-shaped, straight or slightly curved, 0.8–1.2 μm by 1.3–1.6 μm, non-motile, occurring singly, in chains and mainly in pairs. Endospores are not formed. Metabolism is respiratory, never fermentative. Colonies are round, regular, umbonate, soft, glossy and with a diameter of 1–2 mm on AE agar; catalase-positive, oxidase-negative. Growth only in the presence of acetic acid, ethanol and glucose; the total concentration (sum of acetic acid and ethanol) has to exceed 6% and does not oxidize acetic acid. Growth on 3% (v/v) ethanol in the presence of up to 11% acetic acid (v/v). Gluconate, glycerol and lactate are not assimilated. No formation of ketogluconic acid and ethanol decreases the rate of survival cells.

Description of Gluconacetobacter entanii sp. nov.

Gluconacetobacter entanii is a type strain, NCTC 19784 (ATCC 43444). Its DNA G+C content is 70.4 mol%, which is lower than the previously mentioned species. The type strain was isolated from a culture broth taken at the time of cell harvest, and it does not over-oxidize acetic acid. The type strain can only grow on 3% (v/v) ethanol in the presence of acetic acid (v/v). Gluconate, glycerol and lactate are not assimilated. No formation of ketogluconic acid and ethanol reduces the rate of survival cells.

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