Xylose anaerobic conversion by open-mixed cultures

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Abstract Xylose is, after glucose, the dominant sugar in agricultural wastes. In anaerobic environments, carbohydrates are converted into volatile fatty acids and alcohols. These can be used as building blocks in biotechnological or chemical processes, e.g., to produce bioplastics. In this study, xylose fermentation by mixed microbial cultures was investigated and compared with glucose under the same conditions. The product spectrum obtained with both substrates was comparable. It was observed that, in the case of xylose, a higher fraction of the carbon was converted into catabolic products (butyrate, acetate, and ethanol) and the biomass yield was approximately 20% lower than on glucose, 0.16 versus 0.21 Cmol X/Cmol S. This lower yield is likely related to the need of an extra ATP during xylose uptake. When submitted to a pulse of glucose, the population cultivated on xylose could instantaneously convert the glucose. No substrate preference was observed when glucose and xylose were fed simultaneously to the continuously operated bioreactor.

Keywords Xylose · Glucose · Fermentation · Bulk chemicals · Hydrogen · Mixed microbial population

Introduction

Lignocellulosic materials, such as wheat- or rice-straw, corn stovers, or sugar cane bagasse, are available in large amounts at a reasonable cost. Every year, about 73.9 Tg of dry wasted crops are generated in the world (Kim 2004). During the pre-treatment, the cellulose (35–45%) can be separated from the hemicellulose (25–40%). Hemicellulose is a heteropolymer of hexose and pentose sugars, with D-xylose as the major constituent. Xylose is the second most abundant carbohydrate monomer after glucose in agricultural wastes. Whereas glucose can be efficiently converted to ethanol and fine chemicals, an economical application for xylose needs to be developed.

Anaerobic digestion is a widely applied process for treatment of wastewater and organic solid waste aiming at methane-rich biogas production. An alternative anaerobic process could aim for the production of intermediates in the methanogenic process from complex substrates. This can be established by specific inhibition of methane production using operational measures like non-neutral pH-values or decreasing the solid retention time. Potentially interesting products that can be produced in this way include organic acids (acetate, butyrate, lactate, succinate), solvents (ethanol, butanol, acetone) and energy carriers like molecular hydrogen. Which mixtures of products are formed will depend on the substrate treated, the microbial population present and the operational
conditions (Kleerebezem and van Loosdrecht 2007; Lengeler et al. 1998). An extra advantage of anaerobic processes is its association with a low biomass yield, i.e., low waste production. Fermentation of hemicellulose and hemicellulose hydrolysates is a major subject of research, mostly aimed at the production of ethanol (Sommer et al. 2004; van Maris et al. 2006). One of the problems of this process is the inability of many microorganisms to metabolize efficiently all the carbohydrates derived from hemicellulose. Moreover, the presence of mixtures of sugars may lead to a sequential utilization of the substrates: diauxic growth.

Open-mixed culture biotechnology based on natural inocula with a high microbial diversity allows for operation of bioprocesses under non-sterile conditions with no risk of strain degeneration (Kleerebezem and van Loosdrecht 2007). As opposed to pure-culture processes, mixed cultures furthermore allow for the use of continuous processing, and are capable of dealing with mixtures of substrates of variable composition. All these aspects significantly reduce the costs of mixed culture processes compared to pure-culture-based industrial fermentations. The impact of the operational conditions on glucose fermentation by mixed cultures has been studied extensively (Hawkes et al. 2007; Temudo et al. 2007; Zoetemeeyer et al. 1982a; Zoetemeeyer et al. 1982b). However, until now, almost no attention has been given to xylose fermentation. Recently, xylose has been reported as a suitable substrate for biohydrogen production (Calli et al. 2008; Lin et al. 2008; Lin and Cheng 2006; Lo et al. 2008) or electricity production by microbial fuel cells (Huang and Angelidaki 2008; Huang et al. 2008), in both processes by open-mixed cultures. In order to optimize the above-mentioned processes, it is of high relevance to gain knowledge on the differences that exist when fermenting glucose or xylose with open-mixed cultures.

D-Xylose is not so readily utilized as D-glucose for the production of chemicals by microorganisms (Jeffries 1983). The reason may lie on the biochemical pathways used for pentose and hexose metabolism. In most fermentative anaerobic bacteria, D-xylose has to be converted into D-xylulose-5-phosphate (X5P). This metabolite is further metabolized through either pentose phosphate pathway (PPP), or through the phosphoketolase pathway (PKP; Fig. 1). The PPP concerns a series of reactions where D-xylulose-5-phosphate is rearranged and converted into glyceraldehyde-3-phosphate (GAP; Eq. 1), an intermediate of the glucose metabolism. The phosphoketolase cleaves X5P into GAP and acetyl phosphate, which is further converted into acetate and generates one ATP (Eq. 2; Tanaka et al. 2002). Despite the fact that the PPP is the most common pathway, many bacteria also have the capacity to catalyze the PKP pathway.

\[
\text{PPP : } 3 \text{ Xylose } \rightarrow 5 \text{ Pyruvate } + 5 \text{ NADH } + 5 \text{ ATP } \quad (1)
\]

\[
\text{PKP : } 3 \text{ Xylose } \rightarrow 3 \text{ Pyruvate } + 3 \text{ NADH } + 3 \text{ ATP }
\]

\[+ 3 \text{ Acetyl } \rightarrow \text{ P} \quad (2)
\]

The aim of this research is to investigate the product spectrum of xylose fermentation by mixed cultures and compare it to glucose fermentation under comparable process conditions. A relationship between the product spectrum and the type of substrate will be derived, based on generalized biochemical principles.

Materials and methods

Inoculum

The anaerobic mixed culture used as inoculum consisted of a mixture of two types of biomass, obtained from two different sources. The first inoculum came from a distillery wastewater treatment plant, Biothane in Delft, The Netherlands. This plant consists on an UASB operated at 30°C, pH 6.8–7.2, COD=2,500 mg/l. The second inoculum was a sludge solution from a potato starch processing acidification tank in Broek op Langedijk, Smiths Food, The Netherlands. Approximately 20 ml of each inoculum was added to the two reactors at the beginning of the experiment, and operated in batch conditions until biomass growth was observed.

Reactor operation

A continuously stirred reactor with a 3-l capacity (2-l working volume) was used (mechanical stirring 300 rpm). Water was recirculated through a water jacket to maintain a constant temperature of 30±1 °C. The reactor medium was sparged with nitrogen gas at a flow rate of 3.6 vvh in order to maintain anaerobic conditions. The pH was controlled (8±0.1) by automatic titration (ADI 1030 Bio controller) with 4 M NaOH and HCl solutions. To prevent excessive foaming, 3 ml/h of a 3% solution of silicone antifoaming agent (BDH, England) was dosed continuously. The dilution rate was 0.12 h⁻¹.

The reactor operated under the initial conditions (xylose 4 g/l, 0.133 Cmol/l) was run for 4 months, during which pulse experiments were performed. After this period, it was tested the conversion of an influent substrate concentration of 0.333 Cmol/l and co-fermentation of glucose and xylose (each at a final concentration of 0.067 Cmol/l). These experiments were run for 2 weeks at each state and before starting the last experiment, the reactor was reset.
to the initial concentration (0.133 Cmol/l). The ratio carbon/mineral solution was maintained during all the experiments.

Medium

Initially, the cultivation medium contained the following (in g/l): xylose 4.0 (0.133 Cmol/l); NH₄Cl 1.34; KH₂PO₄ 0.78; NaCl 0.292; Na₂SO₄·10H₂O 0.130; MgCl₂·6H₂O 0.120; FeSO₄·7H₂O 0.0031; CaCl₂ 0.0006; H₂BO₃ 0.0001; Na₂MoO₄·2H₂O 0.0001; ZnSO₄·7H₂O 0.0032; CoCl₂·6H₂O 0.0006; CuCl₂·2H₂O 0.0022; MnCl₂·4H₂O 0.0025; NiCl₂·6H₂O 0.0005; EDTA 0.050. The carbon and mineral solutions were prepared and fed separately. The carbon-containing solution was sterilized in an autoclave at 110 °C for 20 min.

Pulse experiments

These experiments were performed in the chemostat reactor while it was in pseudo-steady state. During each experiment, the inflow pumps were switched off and a pulse of a substrate (xylose, glucose, or a mixture of glucose and xylose) and minerals was given; samples were taken and the base addition to control the pH was monitored. In the first experiment, two pulses of xylose were supplied, 0.067 and 0.167 Cmol/l. The second pulse was supplied after xylose of the first pulse was fully converted (approximately 2 h). Additional pulse experiments were performed with glucose (0.083 Cmol/l), and with a mixture of xylose and glucose (both 0.067 Cmol/l). A period of at least 10 days was waited before giving another pulse to assure full recovery of the reactor performance.
System characterization

The reactor was run in continuous mode until pseudo-steady state was established, characterized by a stable product composition and biomass concentration. The off gas composition (H₂ and CO₂) and the base added for pH control were monitored on-line, allowing for direct analysis of the system stability. When these three rate measurements were stable or varied within a limited range (±10%), not tending to increase or decrease, a set of samples was taken during the following week. The samples were analyzed for the concentrations of substrate and soluble organic fermentation products and the biomass concentration.

Analytical methods

Reactor broth samples were immediately filtered (Millipore membrane of 0.45 μm). The substrate and end products were determined and quantified. Xylose, glucose, xylitol, volatile fatty acids (acetate, propionate, butyrate, iso-butyrate, valerate, iso-valerate, and caproate), lactate, succinic acid and, formic acid were determined by HPLC, using an Aminex HPX-87H column from Bio Rad (T=60 °C) coupled to a UV and a RI detector, while phosphoric acid 0.01 M was used as an eluent. The alcohols analyzed were ethanol, propanol, and butanol. These analyses were performed by gas chromatography, Chrompack 9001, equipped with a flame ionization detector (FID) and a fused-silica capillary column 15 m × 0.53 mm HP-INNOWAX. The column temperature was 80 °C for alcohols. The temperature of the injector and detector were 180 °C and 200 °C, respectively. Furthermore, helium was used as a carrier gas.

Measurements of H₂ and CO₂ in the off gas were performed on-line. A gas-detection system was coupled to the bioreactor outlet and a Rosemount Analytical NGA 2000 MLT 1 Multi-component analyzer (infrared detector). Data acquisition (pH and gas) was achieved with SCADA software (Sartorius BBI systems MFCS/win 2.1).

The total soluble and total chemical oxygen demand (COD) was measured in the influent and in the reactor liquid using the Dr Lange kit (1,000–10,000 mg COD/l). The total organic carbon (TOC) and total inorganic carbon (TIC) were measured using a Shimadzu TOC 5050A analyzer. For the total TOC and COD in samples with suspended solids, the samples were first homogenized through cell disruption in an ultrasound bath for 15 s. The inorganic carbon was also estimated based on the Henry coefficient of CO₂, the mass-transfer coefficient (kLa) of the system and carbonate equilibrium constants (Temudo et al. 2007).

The biomass dry weight was determined after filtration, according to standard methods (Greenberg et al. 1992).

The yields of the fermentation products were calculated per substrate consumed, and were corrected for the dilution factor due to dosage of base for pH-correction. The COD and carbon balances were established based on the number of electrons and carbon atoms per mole, while assuming a standard biomass composition of CH₁₈O₅N₀.₂ (Roels 1983).

Calculations

The specific substrate uptake rate (−qₛ in Cmol S/Cmol X h) was determined by adjusting a linear function to the experimental data of the substrate concentration plotted over time, considering the slope of the tangent and dividing the value obtained by the active biomass concentration at that point.

Results

Comparison between glucose and xylose conversion by mixed cultures

The bioreactor was freshly inoculated with 20 ml of each inoculum, and was left in batch mode with either glucose or xylose until growth and microbial activity was observed. Microbial activity could be detected by the online acquisition of the gas production (CO₂ and H₂) and the base added to maintain a constant pH. After this adaptation period, continuous feeding was initiated. The operational performance of the reactors was characterized when a pseudo-steady state was achieved. The pseudo-steady state as characterized by a constant product spectrum and biomass concentration was established after approximately 4 weeks of operation (80 volume changes).

Under substrate-limiting conditions, the product spectrum was largely comparable to glucose in the same conditions (Table 1). Compared to glucose, xylose fermentation produced more butyrate and less ethanol and the biomass yield per C mol substrate was 20% lower. Succinate and propionate yields are not shown in Table 1 because they were always lower than 0.02 Cmol/Cmol. The carbon and electron balances showed 100±10% recovery. Other possible fermentation products such as pyruvate, malate, fumarate, valerate, iso-valerate, caproate, propanol, butanol, 2,3-butanediol, acetoin, and methylglyoxal were below the detection limit.

The yield of the acetyl-CoA-derived products (sum of acetate, ethanol and butyrate) in both reactors was close to the combined yield of formate and molecular hydrogen. This indicates that formate and hydrogen are generated upon cleavage of pyruvate into acetyl-CoA. This suggests that both substrates have been metabolized through glycolysis; the Embden Meyerhof Parnas Pathway (EMP) in case of glucose, and the pentose phosphate pathway in the case of xylose (Eq. 1). In case of a significant contribution of the phosphoketolase pathway (PKP), a substantial amount of
acetate not accompanied by hydrogen or formate production (Eq. 2) should have been detected.

The ATP generated through catabolism was estimated by assuming that the ATP is only generated by substrate level phosphorylation (SLP) involved in each product formation pathway (Decker et al. 1970). This means, each pyruvate-derived product involves a net formation of one ATP from glycolysis as written in Eq. 1; additionally, per mole of acetate and butyrate formed one more ATP is generated also via SLP. This means that in total, acetate and butyrate involve the net formation of 2 and 3 ATP, respectively. The estimated catabolic ATP yields per carbon mole of substrate converted are also given in Table 1. These values suggested that xylose catabolism was more efficient than glucose, because more ATP was generated per carbon of xylose converted (0.54 versus 0.42 mol ATP/Cmol S). However, the measured biomass yield indicated that xylose-grown biomass had a significantly lower efficiency in ATP-utilization for biomass production compared to growth on glucose.

When the xylose concentration was increased to 0.333 Cmol/L, the product spectrum shifted to more ethanol and acetate at the expense of butyrate (Table 1). Additionally, the catabolic ATP yield decreased, but the biomass yield on ATP remained in the same range of the yield at lower substrate concentration. A similar effect of increased substrate/product concentration has been observed when glucose was the substrate (Temudo et al. 2008).

Characterization under substrate non-limiting conditions

To make a clearer comparison between xylose and glucose metabolism by the selected population, three pulse experiments were conducted in the xylose-fed chemostat. Results are shown in Fig. 2. While in the chemostat reactor, the rates (growth and substrate uptake) were set by the dilution rate; during these pulse experiments, substrate non-limiting conditions were established. The calculated kinetic values and the biomass yield are given in Table 2. The initial substrate uptake rates were comparable with the chemostat reactor (1.01, 0.78, and 0.97 compared with 0.89 Cmol/Cmol X h). The catabolic products were the same as obtained during continuous operation: mainly butyrate and acetate.

The operational performance of the bioreactor was largely unaffected by the first xylose pulse. However, when a second pulse with the double amount of substrate was supplied, the uptake rate decreased 26%. This suggests either substrate or product inhibition occurred. In terms of product concentrations, the total amount of acids increased more than 30% due to the first pulse supply of substrate. When a glucose pulse was supplied, the culture started taking up glucose immediately. Despite the lower substrate uptake rate, the culture could make a more efficient use of this substrate as indicated by the increase in the biomass yield (0.14 Cmol/Cmol S and 0.08 Cmol/Cmol S for glucose and xylose, respectively). Considering that the catabolic products were comparable, the difference in the biomass yield should be attributed to the substrate.

During the pulse with the two substrates, both substrates were simultaneously consumed in the beginning (first 2 h), Fig. 2. The glucose uptake rate was approximately 20% lower than the xylose uptake rate; this difference was also observed between the single-substrate pulse experiments. However, after this period, glucose consumption almost ceased and xylose conversion decelerated. The biomass yield was also affected and decreased by 20%. The product spectrum was not affected.

From these pulse experiments in the chemostat, it was confirmed that although no significant difference was observed in the catabolic products, the growth yield was enhanced when glucose was the substrate (Fig. 2). Addi-

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Table 1  Product yields of xylose and glucose fermentation in a chemostat by mixed cultures, at pH 8

|                          | Xylose 0.133 Cmol/L | Glucose 0.133 Cmol/L | Xylose 0.333 Cmol/L |
|--------------------------|---------------------|----------------------|---------------------|
| Dilution rate (h\(^{-1}\)) | 0.128±0.004         | 0.128±0.004          | 0.134±0.004         |
| Biomass yield (Cmol/Cmol) | 0.16±0.01           | 0.21±0.01            | 0.13±0.01           |
| Product yields (Cmol/Cmol S) |                    |                      |                     |
| Acetate                  | 0.18±0.01           | 0.16±0.02            | 0.22±0.01           |
| Butyrate                 | 0.46±0.02           | 0.30±0.04            | 0.07±0.01           |
| Ethanol                  | 0.017±0.004         | 0.08±0.01            | 0.23±0.01           |
| Formate                  | 0.20±0.02           | 0.22±0.01            | 0.18±0.01           |
| H\textsubscript{2} (mol/Cmol S) | 0.13±0.01         | 0.05±0.01            | 0.06±0.01           |
| H\textsubscript{2}+Formate (mol/Cmol S) | 0.33±0.02         | 0.27±0.01            | 0.24±0.01           |
| Acetyl-CoA derivatives\(^a\) (mol/Cmol S) | 0.33±0.02         | 0.27±0.01            | 0.26±0.01           |
| \(\gamma\textsubscript{ATP/S} (ATP/Cmol S)\(^b\)) | 0.54±0.01         | 0.42±0.01            | 0.41±0.01           |
| \(\gamma\textsubscript{X/ATP} (X/ATP (g/mol ATP))\) | 7.6                 | 12.3                 | 8.0                 |

The values presented are the average of the samples (three to four) taken in the time period of one week and the respective standard deviation.\(^a\) Sum of the acetyl-CoA derivatives: acetate, butyrate and ethanol in moles of acetyl-CoA per Cmol of substrate converted.\(^b\) The yield of ATP was estimated based on the catabolic products: one ATP per ethanol produced, two ATP per acetate and three ATP per butyrate.
tionally, glucose conversion capacity was constitutively present in the microbial community growing on xylose.

Co-fermentation of glucose and xylose by mixed cultures in continuous operation

In agricultural waste streams, a mixture of sugars is present. Since a slightly different product spectrum was obtained when glucose and xylose were fermented independently, co-fermentation of xylose and glucose by the culture grown on xylose was studied. The two substrates were fed together in equal carbon amounts (0.067 Cmol/l each; Fig. 3). During the first 24 h (three retention times), little difference could be detected. However, after 3 days, the ethanol yield increased and the butyrate yield decreased. This product spectrum seems to be an intermediate to the one obtained when the two substrates were fed separately. The yields of the other metabolic products and biomass remained almost constant.

The residual concentrations of the two substrates were below the detection limit (0.01 Cmmol/L) and no substrate preference could be detected when a xylose–glucose mixture was used.

Table 2  Kinetic parameters of the mixed microbial culture operated in continuous (substrate concentration of 0.133 Cmol/L, pH 8.0, dilution rate of 0.12 h\(^{-1}\) and temperature 30 °C) and when subjected to a pulse of a substrate (xylose, glucose and a mixture of the two substrates)

| Experiment          | Substrate          | Concentration (Cmol/L) | Specific substrate uptake rate \((-q_s\), Cmol/Cmol X L h) | Biomass yield (Cmol/Cmol) |
|---------------------|--------------------|------------------------|-----------------------------------------------------------|---------------------------|
| Continuous operation| Xylose\(^a\)        | 0.133                  | 0.89                                                      | 0.16                      |
|                     | Glucose            | 0.133                  | 0.62                                                      | 0.21                      |
| Pulse experiments   | Xylose 1st addition| 0.067                  | 1.01                                                      | 0.081                     |
|                     | x Xylose 2nd addition| 0.167                  | 0.55                                                      | 0.064                     |
|                     | Glucose            | 0.083                  | 0.78                                                      | 0.143                     |
|                     | Xylose and glucose | Phase 1\(^b\)          |                                                           |                           |
|                     |                    | Xylose+glucose: 0.167   | 0.97                                                      | 0.111                     |
|                     |                    | Xylose 0.083           | 0.56                                                      | n.a.                      |
|                     |                    | Glucose 0.083          | 0.41                                                      | n.a.                      |

\(^a\) Average of the values determined during three days before these experiments  
\(^b\) Phase 1 comprehends the first 3 h of this experiment  
n.a. Not available
Discussion

A mixed microbial population was selected from a rich inoculum, in a CSTR fed with xylose under substrate-limiting conditions. This culture was capable of converting xylose efficiently as the concentrations were kept lower than the detection limit (0.01 Cmmol/L), suggesting a high affinity for the substrate. The xylose-grown chemostat culture was also able to degrade glucose, not showing a preference for glucose or xylose when point doses of the two substrates are supplied to the bioreactor.

One of the major differences observed between mixed cultures grown on xylose and on glucose is the biomass yield on ATP ($Y_{X/ATP}$), see Table 1. The yield of catabolic products is higher for xylose, which would suggest a higher generation of ATP ($Y_{ATP/S}$); however, the biomass yield is in fact higher when glucose is the substrate ($Y_{X/S}$). This difference has also been observed in studies conducted with pure bacterial cultures, where the biomass yield on the catabolic ATP yield estimated ($Y_{ATP/S}$) is significantly higher for glucose than for xylose (Biesterveld et al. 1994; Heyndrickx et al. 1991; Ounine et al. 1985). Furthermore, in solvent-producing bacteria, not only growth is inhibited but also the xylose uptake is affected by the solvents (Ounine et al. 1985; Qureshi et al. 2006). This is in agreement with the lower energy provided during xylose catabolism. This might be a result of different transport systems that are present in the different species (Jeffries 1983). Facilitated diffusion is the most common mechanism in yeast, the glucose transporters can also catalyze xylose uptake, but have a much lower affinity (Hamacher et al. 2002). In bacteria, however, a number of xylose-specific transporters with different affinities have been identified (Strobel 1993a, b). Of particular interest is the high affinity transporter that consists of an ABC protein and therefore involves one ATP consumption for each xylose transported (Ahlem et al. 1982; Hasona et al. 2004). According to genome databases (http://www.genome.jp), this transporter is widespread over bacterial species able to metabolize xylose, e.g., in fermentative (gram-positive) microorganisms like C. acetobutylicum, but whether it is expressed remains unknown. In case this is the most active transporter in the microbial population selected, the catabolic ATP yield can be recalculated by considering the need of one ATP for xylose transport and one for activation inside the cell. As one molecule of pyruvate comes from 0.6 mol of xylose, this means that the ATP yield associated to pyruvate, as written in Eq. 1, will be affected by this value and becomes 0.4 mol ATP/mol pyruvate formed from xylose. Whereas glucose requires only one ATP for transport and activation, being the glycolytic efficiency estimated to be 1 mol ATP/mol pyruvate. Including these assumptions in our calculations, the $Y_{X/ATP}$ becomes 12.4 g VSS/mol ATP for xylose, which is highly comparable to glucose (12.2 g VSS/mol ATP). We therefore suggest that the lower biomass yield ($Y_{X/S}$) for growth on xylose compared to glucose, is related to the ATP consumption required for active xylose uptake. Such a difference has not been reported for modified yeast cells that were able to ferment xylose (Kuyper et al. 2005a), where the $Y_{ATP/S}$ remained in the same range for the two substrates.

In this study, a mixed microbial population was selected especially for xylose conversion. It was shown that even a culture that was cultivated only on xylose could immediately convert glucose (Figs. 2 and 3). Under non-limiting conditions (pulse experiment), however, the conversion of the two substrates showed some limitations. After a couple of hours, substrate conversion almost ceased. One possible explanation is that the selected population might be sensitive...
to the substrate concentration and/or increased product concentrations. Mixed substrate utilization in batch mode could be further improved by operating in a sequencing batch reactor on the two substrates (Kuyper et al. 2005b).

Use of real waste streams

During this study, all experiments were conducted with synthetic medium. The aim was to study the conversion of the two substrates under substrate-limiting and non-limiting conditions. For practical implementation of the process proposed here, experiments will need to be conducted with mixtures of substrates and real waste streams to verify the pure substrate results. In this respect, it is important to note that difficulties have been reported with the use of real waste streams; e.g., due to the presence of toxic compounds. These issues would have to be addressed before the application of a certain organic waste.

Another aspect of relevance is whether the waste streams are rich in readily biodegradable substrates or in polymers that still have to be hydrolyzed. In this situation, fermentation is not the limiting step and retention times may need to be extended, which may allow other bacteria to convert further the fermentation products by, e.g., methanogenic bacteria. This suggests the requirement to run the mixed culture fermentation process in conditions that are unfavorable for methanogens, e.g., by (periodical) lowering of the operational pH.

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