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**Diversity and metabolism of xylose and glucose fermenting
microbial communities in sequencing batch or continuous
culturing**

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Abstract

A mechanistic understanding of microbial community establishment and product
formation in open fermentative systems can aid the development of bioprocesses
utilising organic waste. Kinetically, a single rate-limiting substrate is expected to
result in one dominant species. Four enrichment cultures were operated to ferment
either xylose or glucose in a sequencing batch reactor (SBR) or a continuous-flow

stirred tank reactor (CSTR) mode. The combination of 16S rRNA gene-based analysis and fluorescence *in situ* hybridization revealed no complete dominance of one species in the community. The glucose-fed and xylose-fed SBR enrichments were dominated >80% by one species. *Enterobacteriaceae* dominated the SBRs enrichments, with *Citrobacter freundii* dominant for xylose and *Enterobacter cloacae* for glucose. *Clostridium*, *Enterobacteriaceae* and *Lachnospiraceae* affiliates dominated the CSTRs enrichments. Independent of substrate, SBR communities displayed 2-3 times higher biomass specific rate of substrate uptake ($q_s^{\max}$) and 50% lower biomass yield on ATP, to CSTR communities. Butyrate production was linked to dominance of *Clostridium* and low $q_s^{\max}$ ($1.06 \text{ Cmol}_s \text{ Cmol}_x^{-1} \text{ h}^{-1}$), while acetate and ethanol production was linked to dominance of *Enterobacteriaceae* and *Lachnospiraceae* and high $q_s^{\max}$ ($1.72 \text{ Cmol}_s \text{ Cmol}_x^{-1} \text{ h}^{-1}$ and higher). Overall, more diversity than expected through competition was observed, indicating mutualistic mechanisms might shape microbial diversity.

Keywords: Mixed culture fermentation – Bioreactor operation – Microbial diversity– r/K selection – Product spectrum – Kinetics

Introduction

The global aim of most societies to develop more circular economies (Ghisellini, Cialani and Ulgiati 2016) urges for a better use of organic waste as a resource. Until now, anaerobic digestion is the most common technology used to valorise this waste in the form of biogas. Several novel bio-based options that provide extra value to resource recovery are arising such as the production of polyhydroxyalkanoates

(Kleerebezem and van Loosdrecht 2007), alginate-like exopolymers (Lin *et al.* 2010), or medium chain length fatty acids (Spirito *et al.* 2014). The first step in these production routes consists of the conversion of polymeric carbohydrates into volatile fatty acids (VFAs) in a mixed-culture fermentative process (Marshall, LaBelle and May 2013). The alignment of VFA production to subsequent processing requires the identification of factors that drive product formation in microbial communities as function of process conditions. First attempts to describe steady-state patterns of mixed culture fermentation as function of an environmental parameter have provided incomplete insights in the product formation pathways established (Rodriguez *et al.* 2006; González-Cabaleiro, Lema and Rodríguez 2015). Observed product spectra at neutral pH could not be simulated properly using these models oriented to ATP production maximisation, indicating incomplete model assumptions. To aid model-based developments there is a need for experimental studies giving a more comprehensive insight into fermentation of specific carbohydrates into VFAs.

Xylose and glucose are the most abundant monomers found in lignocellulosic biomass (Anwar, Gulfraz and Irshad 2014). Fermentation of glucose or xylose can lead to different products, such as lactic acid, ethanol, hydrogen, and VFAs (Figure 1). Xylose can be fermented through the pentose phosphate pathway (PPP) or the phosphoketolase pathway (PKP), resulting in a different stoichiometry. Using the PKP, 40% of the carbon is directly converted to acetate, while the remaining carbon enters into glycolysis. In PPP, all carbon is converted to intermediates for glycolysis, thereby bringing all carbon to pyruvate first (Figure 1). In the first part of glycolysis, one glucose is converted to pyruvate producing four electrons that can be transferred to NADH. If one acetate is produced, a net amount of one NADH is produced. These

electrons cannot be transferred from NADH to hydrogen, as NADH does not possess sufficient energy to drive this reaction (-320 mV and -414 mV for NADH and hydrogen respectively, Buckel and Thauer 2013). Hydrogen is produced through ferredoxin (-400 mV), which is produced when oxidising pyruvate to Acetyl-CoA (Figure 1). The NADH surplus is oxidised by other fermentative pathways, e.g. ethanol production, thereby stoichiometrically coupling acetate and ethanol formation. Recently, electron bifurcation has been proposed as a metabolic strategy in *Clostridium pasteurianum* (Buckel and Thauer 2013) used to conserve energy in fermentation by directly coupling acetate and butyrate formation (Li *et al.* 2008). This mechanism has been successfully incorporated in balancing of NADH of product spectra over a range of pH values (Regueira *et al.* 2018).

Microbial enrichment cultures offer a powerful way of studying the establishment of a specific microbial niche (Beijerinck 1901), depending on the ecological conditions applied, such as pH, temperature, redox couple supplied, nutrients among others. Glucose fermentation has been relatively widely studied, including impacts of pH (Fang and Liu 2002; Temudo, Kleerebezem and van Loosdrecht 2007), temperature (Zoetemeyer *et al.* 1982), solid retention time (SRT) (Chunfeng *et al.* 2009), redox potential (Ren *et al.* 2007), inoculum type (Rafrafi *et al.* 2013), or hydrogen partial pressure (de Kok *et al.* 2013). Xylose is much less studied but its fermentation has been compared to glucose fermentation previously (Temudo *et al.* 2009).

Most studies have been conducted in continuous-flow stirred tank reactors (CSTR), under which regime one substrate is continuously limiting (*i.e.*, operation at low residual concentration). In CSTR systems, affinity dictates the selection: organisms

establishing the lowest residual substrate concentration (C_s) will dominate the enrichment (Kuenen 2014). Affinity is governed by both the maximum biomass specific growth rate ($\mu^{\max}$) and the affinity constant for substrate (K_s). Organisms competing for a substrate in a CSTR environment can, besides optimising their $\mu^{\max}$, optimise their K_s value to actively take up the substrate and dominate the microbial community.

In a sequencing batch reactor (SBR) operation, substrate is supplied in a pulse, leading to a high concentration in the environment of the microorganisms during most of the time that substrate is taken up. Organisms with the highest $\mu^{\max}$ will eventually dominate when substrate uptake is directly coupled to growth. The batch selective environment is traditionally used in microbiology to enrich and isolate organisms, using the shake-flask approach in combination with dilution series. Consequently, fast-growing microorganisms are overrepresented in databases of pure cultures (Prakash *et al.* 2013).

For both CSTR and SBR environments, $\mu^{\max}$ is a selective force, which is a function of the biomass specific rate of substrate uptake ($q_s^{\max}$), the biomass yield on substrate ($Y_{x,s}$) and the maintenance rate on substrate (m_s) (Pirt 1965). From a kinetic point of view, the microorganism with the highest competitive advantage in the environment will eventually outcompete the other microorganisms, which is either the highest $\mu^{\max}$ (in SBR) or highest affinity (in CSTR) on glucose or xylose. Ultimately, we aim to investigate the hypothesis if limiting a single substrate in an enrichment culture leads to the enrichment of a single microbial species. From a competition point of view, one limiting substrate will select for the most competitive

microorganism. Given enough generations or SRTs, this microorganism will eventually dominate the enrichment culture.

Next to microbial competition on substrate, the different pathways for product formation are competing within microorganisms. Anabolism needs chemical energy in the form of ATP to synthesize biomass. Under similar anabolic efficiency, the catabolic pathway that yields more ATP per substrate ($Y_{ATP,S}$) leads to the highest $Y_{X,S}$. Harvested ATP can also be used for active substrate transport. Hereby, microorganisms lower their K_s and thereby create a lower C_s to sustain their selection in a CSTR environment. Fermentative microorganisms are known to choose between a high flux pathway (optimizing $q_s^{\max}$) or a high yield pathway (optimising $Y_{ATP,S}$), which is best described by lactate versus acetate and ethanol formation in *Lactobacillus casei* (De Vries *et al.* 1970). Under CSTR cultivation, at high dilution rates lactate is formed and at low dilution rates acetate, ethanol and formate are formed. Lactate formation yields 2 ATP from 1 glucose, while acetate and ethanol yield 3 ATP from 1 glucose. Thus lactate production is linked to high $q_s^{\max}$, while acetate and ethanol production is linked to high $Y_{ATP,S}$. Thus, a microorganism will preferentially involve a metabolic pathway that maximizes $Y_{ATP,S}$ and/or $q_s^{\max}$ in a SBR environment and $Y_{ATP,S}$, $q_s^{\max}$ and/or K_s in a CSTR environment.

Here, we investigated whether SBR or CSTR environments fermenting either xylose or glucose enrich for an equal microbial community composition and result in equivalent metabolism and kinetics. Three environmental settings were applied to enrich for fermentative microorganisms: (1) a mineral medium with only glucose or xylose as carbon source for fermentation; (2) a combination of temperature, pH, and SRT to select mainly for primary fermentative microorganisms; and (3) suspended

cell cultures. The experimental set up was replicated from Temudo et al. (2009) for a direct comparison of results. The catabolic products, $q_s^{\max}$, and $Y_{x,s}$ were measured for each enrichment in steady state in order to verify if a certain stoichiometry was linked to a certain metabolic strategy. In parallel, we analysed the microbial community compositions to test the microbial diversity hypothesis for enrichment on single substrates, and to link community structures to fermentative products and metabolic strategies.

Materials and methods

Enrichment

All enrichments were performed in 3-L jacketed bioreactors (Applikon, the Netherlands) with working volumes of 2 L. pH was maintained at 8.0 ± 0.1 using NaOH at 4 mol L^{-1} and HCl at 1 mol L^{-1} . Temperature was maintained at $30^\circ\text{C} \pm 0.1$ using a E300 thermostat (Lauda, Germany). The cultures were stirred constantly at 300 rpm. Anaerobic conditions were maintained by sparging the reactor with a flow of $576 \text{ mmol N}_2 \text{ h}^{-1}$ and off-gas was cooled to 5°C using a gas condenser. For the SBRs, a hydraulic retention time (HRT) of 8 h was maintained by removing 1 L of culture per cycle under a cycle time set to 4 h. For CSTRs, the HRT was directly linked to the dilution rate applied.

The synthetic cultivation medium was identical to the one used by Temudo et al. (2007) using 4 g of either xylose or glucose as carbon source per litre. The carbon source and the ammonium, phosphate and trace elements were fed separately from

12.5× concentrated stock solutions and diluted using N₂-sparged demineralized water. Connected to the base pump was a pump supplying 3% (v:v) antifoam C (Sigma Aldrich, Germany), which ensured a flow of 3-5 mL h⁻¹ or 14-17 mL cycle⁻¹. The glucose and xylose solutions were sterilized at 110°C for 20 min.

The inoculum was obtained from cow rumen through a butcher in Est, the Netherlands, and on the same day, transported to lab at room temperature and filtered on 200 µm and aliquoted in 50-mL portions, and frozen at -20°C using 10% glycerol. The seed biomass was then thawed on ice before adding 10 mL to the reactor to start each enrichment culture. When a full first batch was performed the CSTRs were set to continuous mode and the SBRs were set in cycle mode, gradually moving from 24-h to 12-h and 6-h in 3 days to the final desired 4-h cycles to maintain a HRT of 8 h. Steady state was assumed if during a period of at least 5 days no variation was in the product concentrations.

Analytical methods

Samples from the reactors were immediately filtered on 0.45 µm polyvinylidene fluoride membranes (Millipore, USA) and stored at -20°C until analysis. VFAs (formate to valerate), lactate, succinate, ethanol, glucose and xylose were analysed using high performance liquid chromatograph (HPLC) equipped with an Aminex HPX-87H column (BioRad, USA) maintained at 60 °C and coupled to ultraviolet (UV) and refraction index (RI) detectors (Waters, USA), using phosphoric acid at 0.01 mol L⁻¹ as eluent. For high butyrate concentrations above 1 mmol L⁻¹, samples were analysed using gas chromatography (GC), since butyrate overlapped with ethanol on

the RI detector of the HPLC. GC was performed using a Chrompack 9001 (Agilent, USA) equipped with an injector maintained at 180°C, a fused-silica capillary column of 15 m × 0.53 mm HP-INNOWax (Agilent, USA) equilibrated at 80°C for alcohols with helium as carrier gas, and a flame ionization detector set at 200°C. Glycerol was detected using an enzymatic assay relying on glycerokinase, pyruvate kinase and L-lactate dehydrogenase, measuring NADH depletion at 340 nm (Megazyme, Ireland).

The off-gases were monitored on-line for H₂ and CO₂ by a connection to a NGA 2000 MLT 1 Multicomponent analyser (Rosemount, USA). Data acquisition (base, H₂, CO₂) was made using a BBI systems MFCS/win 2.1 (Sartorius, Germany).

Biomass concentration was measured using a standard method which relies on centrifugation to separate the cells from the medium (APHA, 1998). This analysis was coupled to absorbance measurement at 660 nm to establish a correlation. Absorbance values were used to calculate the biomass concentration during the batch experiments.

Cycle analysis

To characterise one cycle in SBR mode, one full cycle was sampled and product and biomass concentrations were measured in parallel to H₂ and CO₂ in the off-gas. In the CSTRs, one litre of volume was removed and one litre of medium was added to finally obtain a concentration of 4 g L⁻¹ of either xylose or glucose together with a stoichiometric amount of other nutrients. Sampling and off-gas analysis were carried out as in the SBRs.

Microbial community analysis

Genomic DNA was extracted using the Ultra Clean Soil DNA extraction kit (MOBIO laboratories, USA) following manufacturer's instructions, with the exception of heating the samples for 5 minutes at 65°C prior to bead beating. Highly molecular DNA was obtained (>10 kb) with a concentration of 10 ng μL^{-1} or higher. Extracted DNA was stored at -20°C until further use.

Analysis of 16S rRNA gene-based amplicon sequencing was conducted to get an overview of the predominant populations in the enrichments in time. The extracted DNA was sent for amplification and sequencing at a commercial company (Novogene, China). Amplification was achieved using the universal primer set 341f / 806r targeting the V3-V4 region of the 16S rRNA gene (Table S1). All polymerase chain reactions (PCR) were carried out in 30 μL reactions with 15 μL of Phusion® High_fidelity PCR Master Mix (New England Biolabs, USA), 0.2 $\mu\text{mol L}^{-1}$ of forward and reverse primers and 10 ng template DNA. Thermal cycling started with an initial denaturation at 98°C for 10 s, annealing at 50°C for 30 s and elongation at 72°C for 60 s and ending with 72°C for 5 min. These pools of amplicon sequences were then sequenced using an IlluminaHiSeq2500 platform. The sequencing datasets were cleaned and trimmed according to Jia *et al.* (2016) and processed with Qiime (Caporaso *et al.* 2010) using UCLUST with a 97% stringency to yield operational taxonomic units (OTUs). OTUs were taxonomically classified using the RDP classifier (Wang *et al.* 2007) with 0.85 confidence interval against the Greengenes database release of August 2013 (DeSantis *et al.* 2006). Double check of OTUs identity factors

was then obtained by alignment against the NCBI RefSeq database using the basic alignment search tool for nucleotides (BLASTn) (Johnson *et al.* 2008).

Cloning-sequencing was conducted to obtain species level information. The near-complete 16S rRNA gene was amplified using the primers GM3f and GM4r (Table S1). The PCR products were purified using QIAquick PCR purification kit (QIAGEN, Germany), ligated, and transformed into competent *Escherichia coli* cells using the TOPO TA Cloning Kit (Invitrogen, USA). Transformed cells were plated on Luria-Bertani medium plates containing 50 µg kanamycin mL⁻¹. After overnight incubation at 37°C, clones were randomly selected for amplification of the 16S insert into the PCR4-TOPO vector using the M13f and M13r primers (Table S1). Depending on the diversity of the sample, 8 to 55 clones were sequenced using Sanger sequencing (Baseclear, the Netherlands). The first and last 100 bp were removed using CodonCode aligner, as sequence quality was insufficient in these regions. Qiime processing was performed on the sequences as described above using a similarity criterium >99% which is defined to be the minimum similarity between species (Janda and Abbott 2007). BLASTn was used to retrieve the identity of each species, and BLAST results with the same species but a different strain were grouped together for phylogenetic resolution at species level. The closest relates strain was then used to retrieve genomic information. Sequences obtained are deposited under the BioProject accession number PRJNA505600 (raw merged amplicon reads) and MK185473 – MK185614 (1450 bp 16S genes) in the NCBI database. Cell fixation and fluorescence *in situ* hybridisation (FISH) were carried out as described by Johnson *et al.* (2009) using the probes listed in table S2, except that hybridization was carried out overnight. Additionally, DAPI staining was used to stain all microbial

cells by incubating the multi-wells microscopy slides of fixed cells with 10 μL of a solution of 10 mg DAPI mL^{-1} per well for 15 min. The samples were analysed using an epifluorescence microscope (Axioplan 2, Zeiss, Germany). Digital images were acquired using a Zeiss MRM camera together with Zeiss imaging software (AxioVision version 4.7, Zeiss, Germany). The 1000x magnified images were improved by setting the 1x sharpening. Three images were taken at 400x and exported as TIFF and used for quantification of the cell surface using the QUIPS feature in Leica QWin V3 (Leica, Germany).

Modelling of the cycle analysis

To obtain the $q_s^{\max}$ and $\mu^{\max}$ for the CSTRs from the cycle analysis, a model was constructed. Herbert-Pirt relation for substrate uptake was simplified by neglecting maintenance, as maintenance is not measured and is assumed to be a small contribution compared to $q_s^{\max}$:

$$\mu = Y_{xs} \cdot q_s \quad (1.1)$$

Monod kinetics were used to describe the growth rate as a function of the substrate concentration at a value of 0.1 mmol L^{-1} of either xylose or glucose:

$$\mu = \mu^{\max} \cdot \frac{C_s}{C_s + K_s} \quad (1.2)$$

The model estimated C_s and C_x by varying the biomass and substrate concentration at the start of the cycle analysis ($C_{x,0}$, $C_{s,0}$) and Y_{xs} and $q_s^{\max}$ values giving the best

fit, and a boundary value of μ is zero was applied when C_s was zero. The modelled values were then optimised to the measured data with a minimisation of the sum-squared error, using the non-linear solver in Microsoft Excel (2010).

Analysis of on-line data collected from the bioreactors

For SBRs, the $\mu^{\max}$ was calculated per cycle using the recorded base dosage values. Microbial growth was directly correlated to the base consumption due to acid production in fermentation (Figure S3). A script was developed in Matlab (version 2014, USA), further explained in the supplementary information (SI) section.

COD and carbon balances

During steady state carbon and chemical oxygen demand (COD) balances were set up using the elemental matrix given in table S4. COD and carbon balances were set up by multiplying the values in the table 9 with the in- and outgoing rates in the reactor, while the NADH, ATP and Gibbs energy balances were set up by multiplying the values in table 9 with the yield on glucose. Data reconciliation was used to obtain closed balances for H, C, O, N and charge using the method described by van der Heijden *et al.* (1994). These balances were used to calculate the Gibbs energy of dissipation.

Carbon and COD balances were set up for the cycle analyses by subtracting the amount of carbon or COD in the compounds measured at a time in the cycle from the measured available carbon or COD at the start of the cycle.

Results

Xylose and glucose fermentation product spectra are similar in SBRs and different in CSTRs

Four different enrichment reactors were operated and analysed for their main products in liquid and gas phase after steady-state was established; this was obtained after 20 SRTs for all enrichments. The glucose SBR exhibited the largest shift in product spectrum during the adaptation, as initially acetate and propionate were the dominant products which changed to acetate and ethanol as dominant products after 18 SRTs. The product spectrum in the xylose and glucose SBR enrichments was very similar, dominated by a catabolic reaction producing ethanol and acetate (Figure 2A), coupled with hydrogen and formate production (Figure 1). Regarding the by-products formed, the xylose SBR enrichment produced more succinate, while the glucose SBR enrichment produced more propionate and lactate.

The xylose CSTR enrichment also had a product spectrum dominated by acetate and ethanol (Figure 2B), coupled to the production of hydrogen and formate. In the glucose CSTR, butyrate was a dominant product, followed by acetate and ethanol (Figure 2B). Both these catabolic pathways were coupled with hydrogen and formate production. Regarding the by-products, similar to the SBRs, the glucose CSTR enrichment produced more propionate and lactate, while the xylose CSTR enrichment produced more succinate, with a significant yield of succinate production in this enrichment of $0.09 \text{ Cmol Cmol}_S^{-1}$ succinate formed.

351

352 Summing up, the glucose SBR and the xylose SBR and CSTR enrichment displayed
353 similar product spectra dominated by acetate and ethanol, while the glucose CSTR
354 showed a mixed product spectrum of butyrate, acetate and ethanol. Glycerol was not
355 detected in a significant amount in any of the enrichments. which was detected up to
356 0.1 Cmol Cmol_S⁻¹ by Temudo et al. (2009).

357

358 **Carbon and COD balances were nearly closed in all enrichments**

359

360 For all enrichments the carbon and chemical oxygen demand (COD, *i.e.*, electron)
361 balances could be closed from the measured products at 95% and 105%,
362 respectively (Table S3). Only in the glucose SBR enrichment a significant amount of
363 10% of carbon and COD could not be recovered in the outflows of the reactor. A
364 characteristic peak at a retention time of 19.1 min was present on the HPLC UV
365 channel for the glucose SBR which could not be identified but was confirmed to be
366 neither 1,3-propanediol nor malate, fumarate, 2,3-butanediol, acetoin or
367 hydroxyvalerate.

368

369 **No storage response or sequential fermentation during cycle analysis**

370

371 For all four enrichments a pulse experiment was performed, in which the substrate
372 and products were measured in time and used to set up a carbon and COD balance
373 over the cycle. A typical storage response would show COD “disappearing” during
374 the initial fermentation phase until the substrate is depleted, while it reappears after
375 substrate depletion as formed products. No such response was observed in both the

CSTR and SBR enrichments (Figure S2) and no sequential conversion of intermediate fermentation products was detected in the cycle analysis in SBRs (Figure S3).

Fast kinetics for SBR enrichments and high biomass yield for CSTR enrichment

At steady state, the yield of biomass formation on substrate was determined in all four enrichments (Table 1). There was no significant difference in biomass yield between the glucose CSTR enrichment reported here and by Temudo *et al.* (2009). The xylose CSTR enrichment displayed a 43% lower biomass yield than the glucose CSTR, and a 25% lower value compared to the xylose CSTR enrichment reported by Temudo *et al.* 2009. The glucose SBR, the xylose SBR and the xylose CSTR enrichment showed similar biomass yield values.

Through analysis of the on-line fermentation data the $\mu^{\max}$ -value for each fermentation cycle could be determined for the SBR enrichments (see SI, figure S5 and S6). A cycle analysis in the CSTR enrichment cultures was used to estimate $q_s^{\max}$. The actual q_s -value in the xylose CSTR enrichment was $1.06 \text{ Cmol}_S \text{ Cmol}_X^{-1} \text{ h}^{-1}$, which was 38% lower than the measured $q_s^{\max}$. The actual $q_s^{\max}$ -value in the glucose CSTR enrichment was $0.55 \text{ Cmol}_S \text{ Cmol}_X^{-1} \text{ h}^{-1}$ which was 48% lower than the maximal rate of glucose uptake. The xylose CSTR enrichment exhibited a 62% higher $q_s^{\max}$ -value than the glucose CSTR enrichment. The $q_s^{\max}$ value found for the xylose SBR enrichment was statistically significantly lower (33%) than for the glucose SBR enrichment (Table 1, $p = 0.002$).

Microbial community analyses highlighted higher diversity with xylose

Amplicon sequencing of the V3-V4 region of the 16S rRNA gene was used to obtain a relative snapshot of the dynamics of the community over time. Then, FISH analysis with three different probes targeting the 16S rRNA of populations of the genus *Clostridium* and of the families of *Enterobacteriaceae* or *Lachnospiraceae* was used to analyse the microbial communities in the enrichments. Lastly, clone libraries were created of the full 16S gene to obtain species-level information of the communities. Microbial diversity was evaluated by the abundance and number of families or genera present.

The xylose SBR enrichment was dominated by *Enterobacteriaceae* (Figure 3, Table 2, figure S7) and a side population of *Lachnospiraceae* and *Clostridium* (Table 2). The 16S amplicon sequencing revealed that the *Enterobacteriaceae* were dominated by *Citrobacter* species (Figure 3), which was confirmed to be *Citrobacter freundii* using the clone library (Figure 4).

The glucose SBR enrichment was dominated by *Enterobacteriaceae* (Figure 3, Table 2, figure S7) with a side population of *Lachnospiraceae*. The 16S amplicon sequencing shows that the *Enterobacteriaceae* were dominated by *Enterobacter* species (Figure 3), which is confirmed to be *Enterobacter cloacae* by the clone library (Figure 4). Two other species also were confirmed using the clone library, *Raoultella ornithinolytica* and *Citrobacter freundii*. Thus, both SBR enrichments were dominated

by a single *Enterobacteriaceae* species, with side-populations of *Lachnospiraceae* in both SBRs, and *Clostridium* in the xylose SBR enrichment.

The glucose CSTR enrichment is dominated by *Clostridium* species (Figure 3, Table 2, figure S7) with a side population of *Enterobacteriaceae* (Table 2). The 16S amplicon sequencing gave two main OTUs, an *Enterobacter* sp. and *Clostridium* sp. (Figure 3), which are confirmed to be *Clostridium intestinale* and *Raoultella ornithinolytica*.

The xylose CSTR enrichment is dominated by *Lachnospiraceae* and *Enterobacteriaceae* species (Figure 3, Table 2, figure S7). The 16S amplicon sequencing is dominated by a *Citrobacter* sp., while two OTUs from the *Lachnospiraceae* are present. The clone library reveals that the *Citrobacter* OTU corresponds to *Citrobacter freundii*, while only one of the *Lachnospiraceae* OTUs can be confirmed up to family level, as it only shows 96% sequence similarity with the closest cultivated relative *Lachnotalea glycerinii* (Table S6).

Summing up, it can be argued that the glucose SBR and CSTR enrichment showed a similar level of diversity, with a dominant species and a small side-population. The xylose SBR enrichment was more diverse than the glucose enrichments, as the side population contains both *Clostridium* and *Lachnospiraceae* species. In the xylose CSTR the largest diversity was observed, as here *Citrobacter freundii*, an uncultivated *Lachnospiraceae* species and a *Muricomes* population dominated.

Discussion

Pathway analysis of the enrichments

Under slightly alkaline and mesophilic conditions acetate and ethanol were the dominant products under SBR conditions, while butyrate formation occurred significantly under CSTR conditions. Compared to the work of Temudo *et al.* (2009) we observe a similar product spectrum in the glucose CSTR enrichment, though we observe more ethanol and less butyrate. The xylose CSTR enrichment is dominated by acetate and ethanol, while the enrichment of Temudo *et al.* (2009) had produced primarily butyrate and acetate. Acetate and ethanol have been shown as the dominant products at pH 7.9 and 30°C (Zoetemeyer, van den Heuvel and Cohen 1982), while acetate and butyrate have been dominant products under at pH 7.0 and 36°C (Fang and Liu 2002).

The rate of the supply of inert N₂ gas in the reactor broth was the only difference in experimental procedures between the present study and the work of Temudo *et al.* (2009). This could potentially change the hydrogen and carbon dioxide gas partial pressures. The impact of the gas flow rate on the fermentation pattern was investigated, in order to investigate if the gas flow rate could explain the differences in product spectrum observed. Little effect was found on all product yields and hydrogen partial pressure (Figure S1); thus, we expect no major impact of the gas flow rate. Furthermore, the glucose CSTR enrichment was duplicated and the resulting product spectrum of both enrichments was identical (Figure S1) which confirms the reproducibility of the enrichments.

A NADH balance was set up using the generalised metabolic network (Figure 1, Table S4), and the derivatives from the pyruvate to acetyl-CoA pathway were summed as a yield. The NADH balance of the four enrichments shows that the glucose CSTR has a small net producing NADH balance, whereas the two SBRs and the xylose CSTR have a small net NADH consuming balance. Minor discrepancies from the NADH-balance can possibly be explained by succinate production through an NADH producing pathway, such as through the oxidative branch of the TCA cycle. Assuming no net NADH consumption for succinate production would bring the two SBRs and the xylose CSTR to a closed NADH balance.

Comparable values for the acetyl-CoA derivatives and H₂/formate production (Table 3) indicate that H₂/formate production is directly coupled to pyruvate conversion to acetyl-CoA in the metabolic network as in Figure 1. Only for the xylose CSTR enrichment there is significantly less formate and H₂ found than acetyl-CoA derivatives, which suggest that H₂ and formate are consumed through homoacetogenesis as proposed by (Regueira *et al.* 2018).

The stoichiometric data argues for the PPP to be active in the xylose SBR, as acetate and ethanol are present in equimolar amounts and there is no excess of acetyl-CoA derivatives compared to formate/H₂. If the PKP would have been active, more acetate compared to ethanol would have been expected and less acetyl-CoA derivatives compared to formate/H₂. In *Clostridium acetobutylicum* the PKP has been significantly expressed under batch cultivation (Liu *et al.* 2012), but here the PPP is assumed to be the only pathway active under SBR conditions.

Bioenergetics and the role of substrate uptake

Using the metabolic network (Figure 1) the amount of ATP produced was estimated from the different catabolic products ($Y_{ATP,s}$). Combining this yield with the biomass yield, the biomass yield on ATP ($Y_{x,ATP}$) was calculated. The $Y_{x,ATP}$ values for the xylose SBR and CSTR are very similar (Table 4), while the $Y_{x,ATP}$ values for the glucose SBR and CSTR enrichments are higher (Table 4). $Y_{x,ATP}$ values are confirmed by the dissipation energy, as the xylose SBR and CSTR enrichment show a similar value, while the value for the glucose SBR enrichment is higher and the highest value is reported for the glucose CSTR enrichment. This means the xylose enrichments have a considerably lower energetic efficiency than the glucose enrichments. The dissipation values obtained for glucose is in accordance with the average values for glucose ($-236 \text{ kJ Cmol}_x^{-1}$), while that of xylose is considerably higher than according to the correlation function ($-246 \text{ kJ Cmol}_x^{-1}$) (Heijnen, van Loosdrecht and Tijhuis 1992).

The higher dissipation in the xylose enrichments can be caused by the cost of transporting xylose over the cell membrane. Xylose can be taken up into the cell by two different mechanisms. XylE is an enzyme which uses the proton motive force to take up xylose from the surrounding medium, through the symport with one proton (Davis and Henderson 1987). When assuming a stoichiometry of 2.67 mol H^+ per mol ATP used, this means xylose uptake XylE costs 0.375 mol ATP per mol xylose. A second method for active xylose uptake is via XylFGH, an ATP-binding cassette (ABC) transporter which uses the direct dephosphorylation of ATP to import xylose (Sumiya *et al.* 1995). XylE is known to be a low affinity transporter, while XylFGH is a

high affinity transporter (Sumiya *et al.* 1995). In *E. coli* it has been demonstrated that in batch conditions XylE plays a minor role in xylose uptake (Hasona *et al.* 2004).

The genome of the strain with the highest similarity was assessed for the presence of transporters. *Citrobacter freundii* strain P10159 dominant in the xylose SBR enrichment (Table S6) contains the XylE gene and not the analogues XylF, XylG or XylH (accession number CP012554.1) This argues for the nature of XylE as a high-rate xylose transport enzyme. A different *Citrobacter freundii* strain FDAARGOS (accession number CP026056.1) was populating the xylose CSTR, which contained neither XylE nor XylF, XylG or XylH. This suggests novel ABC transporters might be present in the xylose CSTR population.

Glucose uptake can be more energy efficient. The phosphotransferase system (PTS) is an uptake mechanism which couples the transfer of a phosphate group from PEP to glucose to transport glucose over the membrane, thus there is no net ATP cost for importing glucose as glucose-phosphate is directly produced. This complex is assumed to be active in both SBR and CSTR as this is observed to be the main transport system under glucose excess (Steinsiek and Bettenbrock 2012) and under substrate limitation (Babu *et al.* 2005). The *Enterobacter cloacae* strain AA4 dominant in the glucose SBR enrichment and the *Clostridium intestinale* strain URNW dominant in the glucose CSTR enrichment both contain all five genes necessary to express the PTS complex in their genomes (accession number CP018785.1 and HM801879.1).

When incorporating this biochemical consideration for substrate uptake, the $Y_{X,ATP}$ value for xylose and glucose becomes similar (Table 4), while the 50% difference in $Y_{X,ATP}$ between SBR and CSTR enrichments remains.

Xylose uptake is slower than glucose uptake in SBR

When substrate is only used for growth and no storage products are formed, the competition in a SBR process is based on the μ^{max} of the competing microorganisms, which can be maximised through $Y_{X,S}$ or q_s^{max} . The SBR grown cultures described in this paper are optimized for q_s^{max} (Table 1). The q_s^{max} of the glucose SBR enrichment is 50% higher than the xylose SBR enrichment. The lower uptake rate for xylose can be explained by a kinetic bottleneck identified in the PPP. Gonzalez *et al.* (2017) have shown that in glycolysis *E. coli* metabolises glucose to fructose-6-phosphate at a rate of $90 \text{ mmol g}_{DW}^{-1} \text{ h}^{-1}$, while in the PPP rates to form fructose-6-phosphate did not exceed $37 \text{ mmol g}_{DW}^{-1} \text{ h}^{-1}$. The production of formate, acetate and ethanol exceeded these values for glucose, indicating the lower part of fermentation was not rate limiting.

Acetate and ethanol production as a kinetic advantage

The q_s^{max} and μ^{max} for the CSTR grown glucose enrichment producing butyrate is significantly lower than the acetate and ethanol producing enrichment (Table 1 and Temudo *et al.* 2009). Furthermore, the xylose CSTR enrichment of Temudo *et al.* (2009) and the glucose CSTR enrichment performed here, showed a similar q_s^{max} -value (Table 1) and both enrichments are producing a significant amount of butyrate.

On top of that, both SBRs produce dominantly acetate and ethanol, where $q_s^{\max}$ is a more important competitive advantage than in CSTR conditions. The kinetic difference between butyrate forming and acetate and ethanol forming microorganisms is observed in pure cultures. The $\mu^{\max}$ of *Clostridium tyrobutyricum*, a butyrate producer, is 0.12 h^{-1} (Liu and Yang 2006) and *Citrobacter* sp. CMC-1, an acetate and ethanol producer, is 0.21 h^{-1} (Mangayil, Santala and Karp 2011) grown under similar conditions. The fact that acetate and ethanol formation is related to higher $\mu^{\max}$ is also indirectly shown by the study of Zoetemeyer *et al.* (1982), as a μ of 0.25 h^{-1} was applied here at pH 7.9 and 30°C obtaining a product spectrum of acetate and ethanol, while Temudo *et al.* (2009) and this study obtain also butyrate production at a μ of 0.13 h^{-1} . This kinetic advantage seems to hold only for fermentations at pH higher than 6.25, as enrichments performed in CSTR mode at pH 5.5 above $\mu^{\max}$ have demonstrated to systemically yield a product spectrum dominated by acetate, butyrate, and lactate (Rafrafi *et al.* 2013). This kinetic effect can be incorporated into model-based evaluation of mixed culture fermentations to improve the prediction of butyrate, acetate and ethanol production at neutral and alkaline pH.

Butyrate production as an efficient pathway

If acetate and ethanol production obtains a higher $q_s^{\max}$ value than butyrate, and both pathways produce 3 mol ATP, there seems to be no advantage for butyrate production over acetate and ethanol production. Thermodynamically, butyrate formation yields more energy than acetate and ethanol production, (-264 kJ mol^{-1} and -226 kJ mol^{-1} respectively). This energy is available in the step from crotonyl-CoA to

butyryl-CoA, which is calculated to be -50 kJ/mol (González-Cabaleiro, Lema and Rodríguez 2015). A direct conversion of this energy into a proton motive force has been rejected (Herrmann *et al.* 2008). Part of the energy can be conserved by coupling this energy to the transfer of the electrons from NADH to ferredoxin and then oxidizing ferredoxin with NAD^+ to generate a sodium motive force using the Rnf enzyme (Herrmann *et al.* 2008). Two of the six subunits of this complex are found in the genome of the *Clostridium intestinale* strain URNW, indicating the possibility of this mechanism being active in the glucose CSTR enrichment.

Metabolic strategies in fermentation: r-organisms vs K-organisms

The CSTR enrichments, when corrected for substrate uptake, show about 50% higher $Y_{x,\text{ATP}}$ value than the SBR enrichments. The q_s^{max} -value on the other hand is 2-3 times higher for the SBR enrichments compared to the CSTR enrichments. These observations correspond with the general microbial theory proposed on r- vs K-organisms (Andrews and Harris 1986). The r-organisms are more adapted to a substrate-abundant environment and display high q_s^{max} and μ^{max} values. K-organisms are more adapted to crowded environment where substrate is limited and display high $Y_{x,\text{ATP}}$ and K_s values. The reason r-organisms dissipate more energy than K-organisms in their metabolism may rely on the fact that at increasing growth rate more erroneous proteins are produced due to a higher error rate made during proofreading at higher speed (Yamane *et al.* 1977). Thus, more non-functional proteins are produced at higher growth rate. As protein production is estimated to cost >80% of the ATP to synthesise a cell (Hespell and Bryant 1979), larger error

rates will cause increased ATP cost per cell assuming a similar functioning protein content.

The community data shows that *Enterobacteriaceae* dominate the SBR environments, thus the *Citrobacter freundii* and *Enterobacter cloacae* species can be classified as r-organisms. *Enterobacteriaceae* species such as *E. coli* are well known to exhibit high growth rates in anaerobic environments with carbohydrates (De Vrije and Claassen 2003). *Clostridium* species on the other hand are often dominating in substrate-limited environments such as anaerobic digesters (Burrell *et al.* 2004), where the rate of hydrolysis of cellulose and hemicellulose is an order of magnitude lower than typical fermentation rates, creating a substrate-limited environment. In the glucose CSTR we observe a dominance of *Clostridium intestinale*, which fits with these observations.

The microbial community composition and the effect of limiting a single substrate

First of all, it is noteworthy that the FISH imaging and the 16S rRNA gene amplicon sequencing data do not always correspond. In the glucose SBR, the dominance of *Enterobacteriaceae* on OTU-level is confirmed by the FISH analysis, but in the glucose CSTR enrichment the *Enterobacteriaceae* are observed to be a minor fraction on cell-level (FISH image), while 30% of the reads relate to *Enterobacteriaceae*. In the xylose CSTR a similar bias is observed, as 53% of the community is identified as *Lachnospiraceae* using FISH (Table 2), while only 15% of the reads relate to *Lachnospiraceae*. As we have corrected the data for copy

numbers, the bias is likely caused by DNA extraction and PCR biases, which are known to cause biases in amplicon sequencing data (Brooks *et al.* 2015). As proposed by Amann, Ludwig and Schleifer (1995), 16S rRNA gene sequencing and FISH analysis have to be used in parallel to obtain an accurate estimation of the microbial community structure, which is confirmed in the study here.

Here, populations of *Enterobacteriaceae*, *Lachnospiraceae* and *Clostridium* dominated the enrichments. *Clostridium* and *Enterobacteriaceae* populations have been reported in enrichments on mineral medium (Table 5), though for the first time *Lachnospiraceae* were enriched on xylose. We find that a significant presence of *Clostridium* was linked to butyrate production, as in the glucose CSTR, which is confirmed by other enrichment studies (Table 5). The butyryl-CoA dehydrogenase gene, which is responsible for the reduction of crotonyl-CoA to butyryl-CoA using NADH, is found in organisms in the *Clostridium* species, while neither in *Enterobacter* nor in *Citrobacter* species according to the NCBI Gene database.

The glucose enrichments seem to be dominated by a single species with one side populating family, which is *Enterobacter cloacae* in the glucose SBR and *Clostridium intestinale* in the glucose CSTR. It was expected that, when limiting a single substrate, one specialist will dominate the community after prolonged cultivation, displaying either the highest $\mu^{\max}$ or the highest affinity. For the xylose enrichments, the communities are more diverse. In the xylose SBR, *Citrobacter freundii* dominated the culture, with a side-population of both, *Lachnospiraceae* and *Clostridium*. The xylose CSTR is populated by two *Lachnospiraceae* OTUs (Figure 3), one of which is confirmed to be an uncultivated *Lachnospiraceae* species (Table S6) next to a

population of *Citrobacter freundii*. Thus, xylose fermentation results in more microbial diversity than glucose fermentation.

All four enrichments are populated by more than one species, with stabilizing OTUs over time (Figure 3). This indicates that species have a reason to coexist in these single substrate limited systems. It is possible that mutualistic relationships between these species are present, *e.g.*, in the form of a B-vitamin exchange between species (Magnúsdóttir *et al.* 2015), as these communities are cultivated on mineral medium. Overall, it remains an important ecological question why in many cases rather diverse communities remain in very selective conditions with one limiting substrate.

Overall, this study aimed to show the impact of sequencing batch and continuous culturing on microbial communities fermenting lignocellulosic sugars such as xylose and glucose. Butyrate formation was linked to slow uptake rate, while acetate and ethanol formation was linked to high uptake rates. This kinetic effect can be taken into account in modelling efforts. In SBR, xylose was fermented 33% slower than glucose. SBR communities maximised their $q_s^{\max}$, while CSTR communities maximised their $Y_{x,ATP}$. SBR communities were dominated by r-strategists like *Citrobacter freundii* and *Enterobacter cloacae*, and the CSTR communities by K-organisms like *Clostridium intestinale* and *Lachnospiraceae* species. No significant storage of either xylose or glucose was observed in the SBR enrichments. The glucose enrichments confirmed the hypothesis that limitation of a single substrate leads to domination of a single species. The xylose enrichments displayed more microbial diversity, with the xylose CSTR up to three dominant populations.

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846 Dissimilation of Glucose in an Anaerobic Digestor. *Water Res* 1982;**16**:303–311.

847

848 Table 1: $Y_{x,s}$ calculated on the basis of TSS/VSS measurements at steady state (n=3). For
849 the SBRs, $\mu^{\max}$ was obtained from on-line fermentation data according to appendix VI. For
850 the CSTRs, $q_s^{\max}$ was obtained from a substrate pulse experiment and subsequent fitting the
851 substrate concentration data, with R^2 values of 0.97 and 0.92 for xylose and glucose
852 respectively. For the SBR $\sigma_{q_{s\max}}$ is calculated using error propagation and the covariance of
853 the $\mu^{\max}$ and $Y_{x,s}$ values. For the CSTRs $\sigma_{q_{s\max}}$ is calculated using error propagation and the
854 covariance of the C_s and C_x measurement, while $\sigma_{\mu\max}$ is calculated using error propagation
855 and the covariance of $q_s^{\max}$ and $Y_{x,s}$.

	$Y_{x,s}$ [Cmol _x Cmol _s ⁻¹]	$q_s^{\max}$ [Cmol _s Cmol _x ⁻¹ h ⁻¹]	$\mu^{\max}$ [h ⁻¹]	Reference
Xylose SBR	0.12 ± 0.01	2.28 ± 0.10	0.28 ± 0.01	This study
Glucose SBR	0.13 ± 0.01	3.41 ± 0.24	0.45 ± 0.01	This study
Xylose CSTR	0.12 ± 0.01	1.72 ± 0.02	0.22 ± 0.01	This study
	0.16 ± 0.01	1.01	0.16	Temudo <i>et al.</i> (2009)
Glucose CSTR	0.21 ± 0.01	1.06 ± 0.02	0.22 ± 0.01	This study
	0.21 ± 0.01	NA	NA	Temudo <i>et al.</i> (2009)

856
857 Table 2: Result of the FISH quantification (n = 3), with percentages denoting relative
858 abundances calculated from the target-probe surface area compared to EUB338 probe
859 surface. Unidentified populations were calculated as the remaining percentage after
860 summing up the relative abundances of the known populations. The last column shows the
861 amount of surface probed by EUB338 compared to DAPI.

	Chis150 vs. EUB338	Lac435 vs. EUB338	Ent183 vs. EUB338	Unidentified vs. EUB338	EUB338 vs. DAPI
Xylose SBR	2% ± 2%	5% ± 1%	90% ± 3%	2%	96% ± 2%
Glucose SBR	ND	3% ± 2%	91% ± 3%	6%	100% ± 7%
Xylose CSTR	ND	53% ± 3%	44% ± 6%	3%	104% ± 14%
Glucose CSTR	89% ± 12%	ND	5% ± 0%	6%	89% ± 8%

862

Table 3: Net NADH balance calculated using table S4. Acetyl-CoA derivatives were calculated from butyrate, acetate and ethanol production through the pyruvate to acetyl-CoA pathway (Figure 1).

	Net NADH balance metabolism [mol _{NADH} Cmol _S ⁻¹]	Acetyl-CoA derivates [mol Cmol _S ⁻¹]	Formate + H ₂ [mol Cmol _S ⁻¹]
Xylose SBR	-0.03 ± 0.00	0.27 ± 0.00	0.26 ± 0.00
Glucose SBR	-0.03 ± 0.01	0.22 ± 0.00	0.23 ± 0.02
Xylose CSTR	-0.06 ± 0.01	0.27 ± 0.00	0.22 ± 0.01
Glucose CSTR	0.02 ± 0.01	0.24 ± 0.20	0.25 ± 0.01

Table 4: $Y_{x,ATP}$ is calculated by assuming ATP formation per product (Table S4), for the measured data and corrected for substrate uptake. Xylose uptake in the CSTR is assumed by the XylFGH complex and the XylE complex in the SBR. Gibbs energy of dissipation is calculated at 30°C and pH = 8 using the reconciled data.

	Y_{xS} [Cmol _X Cmol _S ⁻¹]	$Y_{ATP,S}$ [mol _{ATP} Cmol _S ⁻¹]	$Y_{x,ATP}$ observed [g _X mol ⁻¹ ATP]	$Y_{x,ATP}$ corrected [g _X mol ⁻¹ ATP]	Gibbs energy of dissipation [kJ Cmol _X ⁻¹]
Xylose SBR	0.12 ± 0.01	0.42 ± 0.01	7.2	8.7	-378
Glucose SBR	0.13 ± 0.01	0.40 ± 0.01	8.2 ¹	8.2 ¹	-285
Xylose CSTR	0.12 ± 0.01	0.42 ± 0.01	6.8	12.8	-386
Glucose CSTR	0.21 ± 0.01	0.49 ± 0.03	13.4	13.4	-236

¹Only 90% of glucose conversion is assumed here, as the COD and carbon balance only close for 90%

874 Table 5: Reported predominant bacterial species for fermentative microbial communities
 875 enriched on xylose or glucose as carbon sources in CSTR mode. Species were detected
 876 using PCR and denaturing gradient gel electrophoresis or PCR and single strand
 877 conformation polymorphism analysis

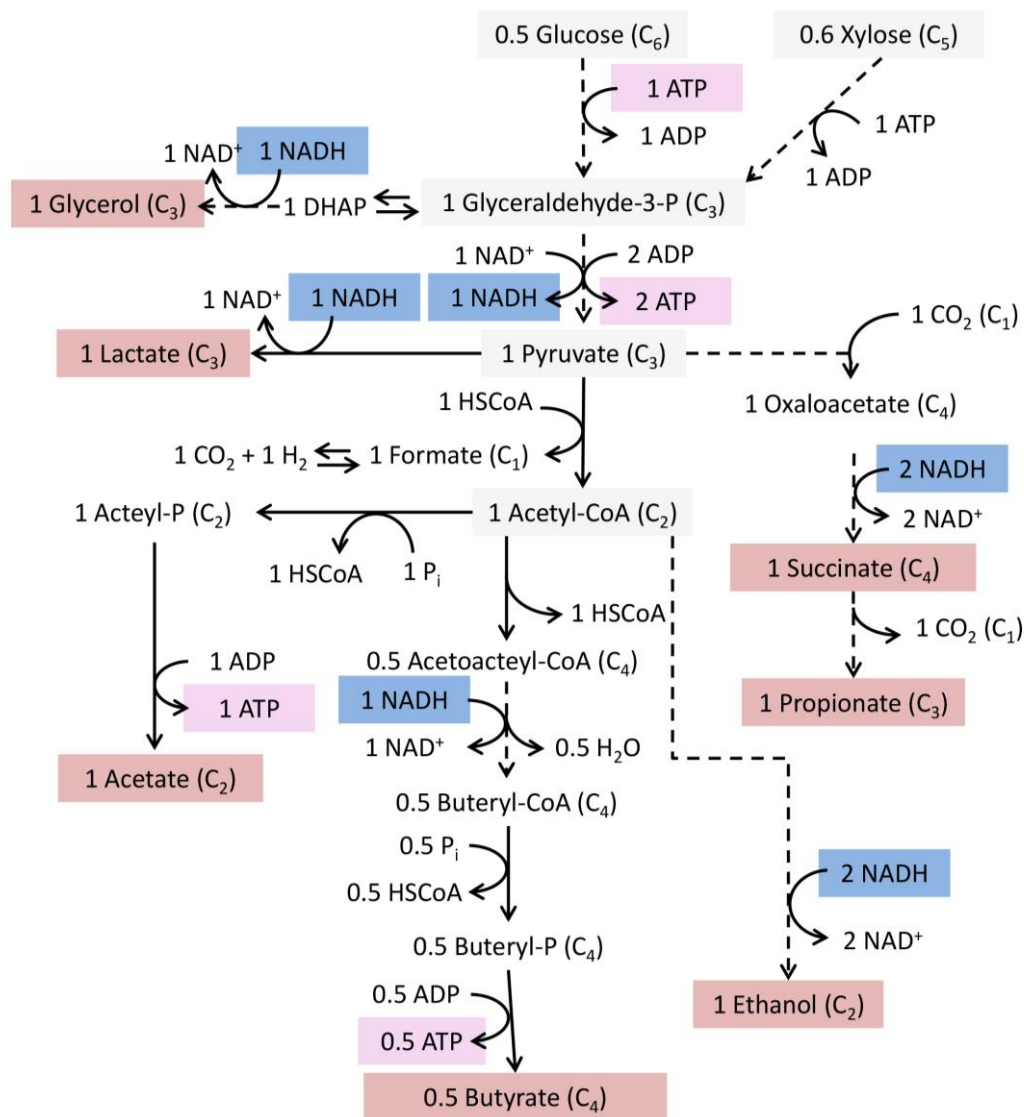
Substrate	Inoculum	T	pH range	Dominant carbon products	Organisms	Source
Xylose	Hot spring culture	45 °C	5.1	Acetate, butyrate	<i>Clostridium</i> <i>acetobutylicum</i> <i>Citrobacter freundii</i>	(Mäkinen, Nissilä and Puhakka 2012)
Xylose	Hot spring culture	37 °C	5.1	Acetate, butyrate, ethanol	<i>Clostridium</i> <i>acetobutylicum</i> <i>Clostridium tyrobutircum</i>	(Mäkinen, Nissilä and Puhakka 2012)
Glucose	Hot spring culture	37 °C	5.0	Acetate, butyrate	3 species of <i>Clostridium</i> 2 uncultured	(Karadag and Puhakka 2010)
Glucose	Activated sludge, cassava, rabbit droppings	37 °C	5.5	Butyrate, acetate, lactate*	<i>Clostridium</i> <i>pasteurianum</i> , <i>Clostridium beijerinckii</i> , <i>Lactobacillus paracasei</i>	(Rafrafi et al. 2013)
Xylose 4 g/L	Digestor sludge and acidification tank	30 °C	8.0	Acetate, butyrate	<i>Clostridium beijerinckii</i> , <i>Clostridium xylanovorans</i> , <i>Clostridium</i> sp. CCUG	(Temudo et al. 2008)
Xylose	Digestor	30 °C	8.0	Acetate,	<i>Citrobacter farmeri</i>	(Temudo et

11 g/L	sludge and acidification tank			butyrate, ethanol	<i>Clostridium intestinale</i> <i>Clostridium sp. CCUG</i>	<i>al.</i> 2008)
Glucose	Digestor sludge and acidification tank	30 °C	8.0	Acetate, butyrate, ethanol	<i>Clostridium quinii</i> **	(Temudo <i>et al.</i> 2008)

878 * 50% of the COD coming out of the reactor was glucose

879 ** two other bands are visible which are not mentioned

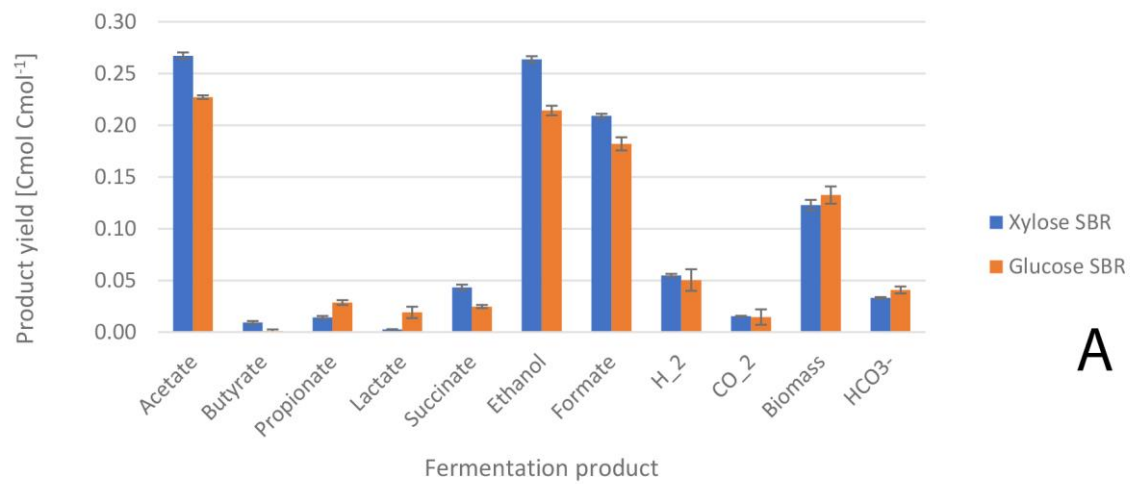
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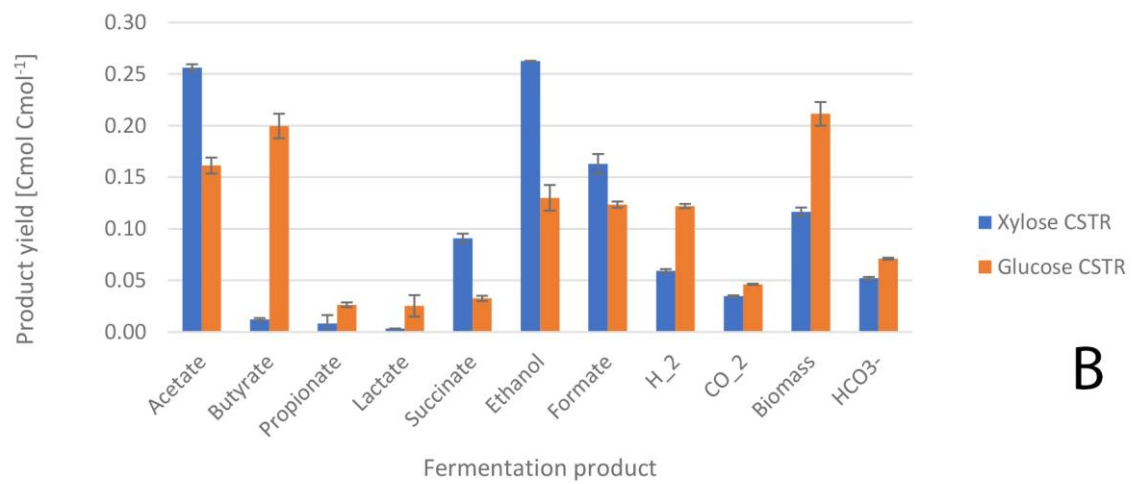
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882 Figure 1: Intracellular metabolic network for xylose and glucose fermentations. Dashed lines
 883 indicate lumped reactions, straight lines indicate single reactions. Xylose comes into the
 884 glycolysis through the synthesis of 2 fructose-6-phosphate and 1 glyceraldehyde-3-
 885 phosphate, through the PPP. The Emden-Meyerhof-Parnass pathway is used as this is the
 886 common type of glycolysis encountered in energy limited anaerobes (Flamholz *et al.* 2013).
 887 Figure is made on the basis of Madigan and Martinko (2006).

888



A



B

Figure 2: Product spectra of mixed culture fermentations of SBRs (A) and CSTRs (B) determined in steady state (n=3)

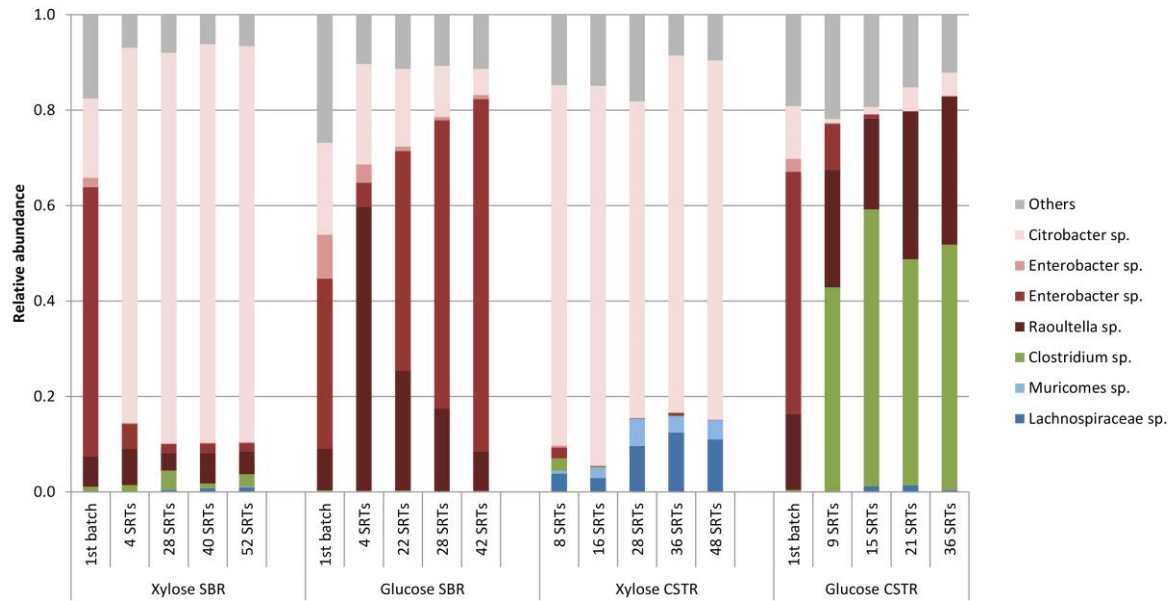
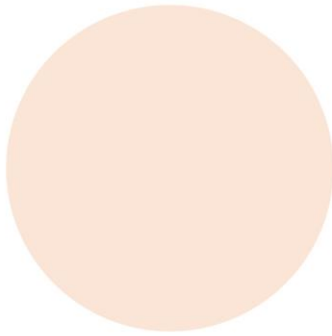
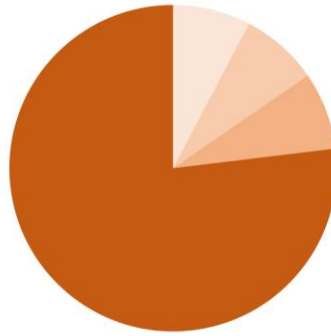


Figure 3: Overview of the amplicon results on the V3-V4 region of the 16S rRNA gene on OTU level. All OTUs that contribute to <1% of the reads are grouped into the others fraction (grey). In red OTUs belonging to the *Enterobacteriaceae* family are denoted, in green OTUs belonging to the *Clostridiaceae* family and in blue OTUs belonging to the *Lachnospiraceae* family. Closest related relatives found by BLAST used to characterize the OTU up to genus level (Appendix V). OTUs matched at <97% are presented as species from a family.

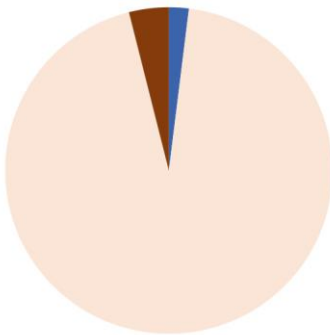
Xylose SBR



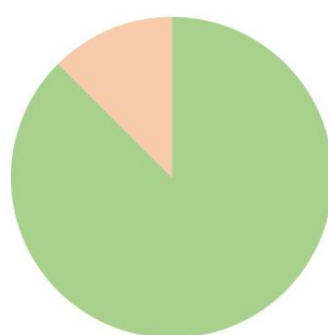
Glucose SBR



Xylose CSTR



Glucose CSTR



- Lachnospiraceae sp.
- Clostridium intestinale
- Citrobacter freundii
- Raoultella ornithinolytica
- Klebsiella sp. JT42
- Enterobacter cloacae
- Citrobacter pasteurii

903

904

905 Figure 4: Result of the clone library analysis in which strains that were found as closest

906 relative (Appendix VII) are grouped into species

907

908