



Methanogenic consortia from thermophilic molasses-fed structured-bed reactors: microbial characterization and responses to varying food-to-microorganism ratios

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Abstract

The heterogeneous character of fixed-film reactors may create highly specialized zones with a stratified distribution of microbial groups and varying capabilities to withstand high organic loads in anaerobic digestion (AD) systems. The microbial distribution and methane-producing potential of biomass from different regions (feeding zone and structured bed) of two second-stage thermophilic (55 °C) fixed-film reactors were assessed. Three levels of food-to-microorganism (F/M) ratio (0.4, 1.0 and 3.0 g-COD g⁻¹VS) using fermented (two-stage AD) and fresh (single-stage AD) sugarcane molasses were tested in batch reactors, simulating low to high organic loads. Specific methane production rates increased as the F/M increased when using fermented molasses, maintaining efficient methanogenesis at substrate availability levels threefold higher than single-stage schemes (3.0 vs. 1.0 g-COD g⁻¹VS). Success in methane production derived from the homogenous establishment (similar in both feeding zone and bed) of syntrophic associations between acetogens (*Pelotomaculum*, *Syntrophothermus*, *Syntrophomonas* and *Thermodesulfovibrio*), acetate oxidizers (*Thermoacetogenium*, *Mesotoga* and *Pseudothermotoga*) and hydrogenotrophic methogens (*Methanothermobacter* and *Methanoculleus*) replacing acetoclastic methanogens (*Methanosaeta*). Phase separation under thermophilic conditions was demonstrated to boost methane production from sugar-rich substrates, because the process depends on microbial groups (hydrogenotrophs) that grow faster and are less susceptible to low pH values compared to acetotrophs.

Keywords Two-stage biodigestion · Sugarcane molasses · 16S rRNA gene amplicon sequencing · Kinetic assessment · Syntrophic acetate oxidation + hydrogenotrophic methanogenesis

Abbreviations

AD	Anaerobic digestion	COD	Chemical oxygen demand
AnSTBR	Anaerobic structured-bed reactor	CODs	Soluble COD
ASV	Amplicon sequence variant(s)	CODt	Total COD
CHt	Total carbohydrates	F/M	Food-to-microorganism (ratio)
		FDZ	Feeding zone

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fMol	Fermented sugarcane molasses
GC	Gas chromatography
HM	Hydrogenotrophic methanogens
HRT	Hydraulic retention time
IA	Intermediate alkalinity
IN	Inoculum (sludge used to inoculate the continuous reactors)
k_1	First-order kinetic constant
k_1'	Specific first-order kinetic constant
Mol	Fresh sugarcane molasses
MPP	Methane production potential
NCBI	National Center for Biotechnology Information
OLR	Organic loading rate
PA	Partial alkalinity
P_{CH_4}	Potential methane production
PCoA	Principal coordinates analysis
PCR	Polymerase chain reaction
PheOH	Total phenols
PU	Polyurethane
PVC	Polyvinyl chloride
qPCR	Quantitative PCR
RA	Relative abundance
R_{CH_4}	Maximum methane production rate
R_{CH_4}'	Specific maximum methane production rate
RM1/RM2	Nomenclatures for the continuous reactors
SAOB	Syntrophic acetate-oxidizing bacteria
SO_4^{2-}	Sulfate
sOLR	Specific organic loading rate
STB	Structured bed
UASB	Upflow anaerobic sludge blanket (reactor)
VFA	Volatile fatty acids
VOA _{GC}	Volatile organic acids by gas chromatography
VOA _{tit}	Volatile organic acids by titration
VS	Volatile solids
λ	Lag phase

Introduction

Anaerobic digestion (AD) of high-strength wastewaters has been facilitated by developing high-rate systems, in which biomass retention mechanisms enabled uncoupling the hydraulic retention time (HRT) from the solid retention time in reactors (van Lier et al. 2015). Biomass granulation or the provision of stationary or moving media for cell attachment has created suitable conditions for the application of high organic loading rate (OLR) under low HRT, resulting in a series of benefits, such as achieving higher robustness and operating stability, requiring more compact units and increasing biogas production. Among the numerous configurations proposed are two that successfully achieved effective full-scale applications, namely, the sludge blanket and fixed-film

systems (Moletta 2005). The vulnerability of sludge granulation to compositional characteristics commonly found in numerous wastewaters, such as high suspended solid content and high salinity (van Lier et al. 2015), provides some advantages to using support materials in cell retention. Focus is given to the second group in this study, because recent research efforts have led to improvements on the conventional packed-bed systems through vertically arranging the support media to characterize the anaerobic structured-bed reactor (AnSTBR) (Aquino et al. 2017; Camiloti et al. 2014; Fuess et al. 2021a; Mockaitis et al. 2014).

AnSTBR systems simultaneously provide high void indices and adequate surface area levels for biomass attachment, which expand the applicability of fixed-film reactors. Compared to packed-bed systems, the higher bed porosity (> 90%) (Aquino et al. 2017; Mockaitis et al. 2014) minimizes bed clogging-related limitations, which enables both the processing of solid-rich substrates and the establishment of microbial populations with high growth yields (e.g. hydrolytic and fermentative bacteria). The attached growth itself also comprises an advantageous feature compared to other high-rate systems because less metabolic energy is required to establish high cell densities within the system (Aquino et al. 2017; Blanco et al. 2017). Consequently, the microbial community presents higher robustness when faced with adverse environmental conditions, such as organic overloading events (Chan et al. 2009).

Interestingly, fixed-film reactors do not behave as homogeneous systems, because the low-to-null cell mobility associated with the progressive conversion of the substrate may create highly specialized zones in the reactor. Following the flow direction, a “more acidogenic” microbial community should be expected in regions with high fresh substrate availability (e.g. feeding zones), whilst a “more acetogenic” and mainly methanogenic-like character should be observed in the subsequent compartments of the reactor. It is worth highlighting that the fermentability of the substrate may directly impact this microbial distribution, because the proposed specialization should be more evident in the processing of easily fermentable substrates, such as sugar-rich ones. Conversely, a relatively more homogeneous microbial distribution (or at least without the prevalence of fermentative groups near the feeding zone) is expected to occur in second-stage methanogenic reactors, in AD schemes where acidogenesis is carried out separately from methanogenesis.

Different microbial populations will have different growth rates (Mosey 1983), potentially leading to wide variations of the food-to-microorganism (F/M) ratio along fixed-film reactors. The amount of microorganisms is commonly measured as volatile suspended solids (VSS) or only volatile solids (VS), whilst the “food” is represented by the organic matter content measured as the chemical oxygen demand (COD). Recent studies associated optimal fermentation activity (targeting

biohydrogen production) with F/M ratios within the range of 4.0–6.0 g-COD g⁻¹VSS d⁻¹ (Blanco et al. 2017; Anzola-Rojas et al. 2015; Fuess et al. 2021b; Hafez et al. 2010), whilst stable methanogenesis has been conventionally pointed to occur under much lower F/M ratio values (<0.4 g-COD g⁻¹VSS d⁻¹ (Aquino et al. 2017; Barros et al. 2016; Chernicharo 2007), specifically in single-stage schemes. However, the compartmentalized assessment of a second-stage AnSTBR indicated the establishment of highly efficient substrate conversion into methane under F/M ratios as high as 3.0 g-COD g⁻¹VSS d⁻¹ (Fuess et al., 2021a) directly contradicting the expected behavior for methanogenic environments. In practical aspects, high F/M ratios indicate (or simulate, in the case of batch reactors) situations of organic overloading (or close to that), in which the activity of fermentative bacteria should prevail and inhibit methanogens by providing an excess amount of volatile fatty acids (VFA). Synonyms for the term F/M ratio in continuous reactors include both “biological loading rate” and “sludge loading rate” (Chernicharo 2007), as well as “specific organic loading rate” (Anzola-Rojas et al. 2015).

Some important questions arise from the stratified character of fixed-film reactors, regarding both the spatial distribution of specialized microbial groups and variations in the quality of the substrate (driven by the pre-fermentation intensity): [i] Does biomass specialization effectively occur as a result of differences in substrate quality (following the flow direction) or is a homogeneous distribution of all microbial groups participating in substrate conversion repeated along the entire reactor? [ii] More specifically, can methanogens withstand higher organic loads (or higher F/M ratios) when phase separation is used, i.e. when substrate fermentation is carried out (separately) prior to methanogenesis?

To answer these questions, the microbial distribution and methane-producing potential of biomass collected in different regions of two second-stage thermophilic AnSTBR systems were assessed. Samples collected from the feeding zone and the bed region were compared using 16S rRNA gene sequencing to understand performance discrepancies as a function of the prevailing microbial groups. Batch tests using biomass from the feeding zones were further conducted under different F/M ratio values using fermented and fresh (non-fermented) sugarcane molasses as the substrate to understand how efficiently methanogenic consortia are able to perform under low and excess substrate availability levels, potentially defining operating limits for anaerobic reactors.

Material and methods

AnSTBR systems and thermophilic methanogenic consortia

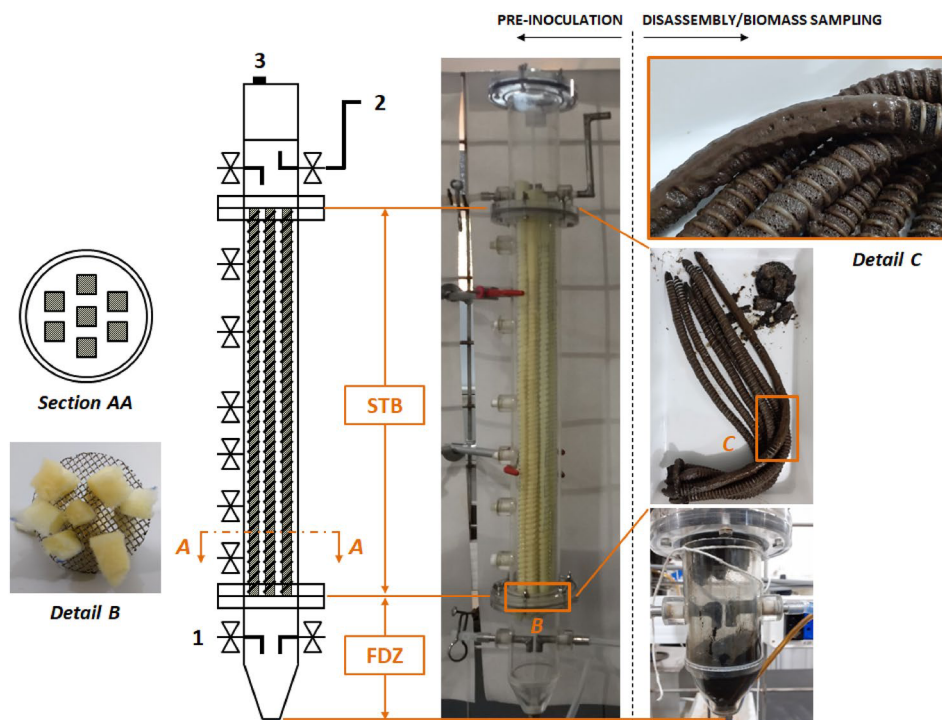
Two bench-scale (1.9 L each) AnSTBR systems were sampled to obtain the methanogenic consortia

microbiologically characterized and used as inocula in methane production potential (MPP) tests in this study. Polyurethane (PU) foam strips wrapped in spiral polyvinyl chloride (PVC) frames were used as the support material in both reactors, which were operated in upflow mode and fed with fermented sugarcane molasses for 250 (RM1) and 230 (RM2) days under thermophilic temperature conditions (55 °C). The reactors were subjected to different alkalization strategies: while sodium bicarbonate (NaHCO₃) dosing was carried out during the entire operating period of RM1, sodium hydroxide (NaOH) dosing coupled to effluent recirculation was initially used in RM2 (185 d), after which this approach was replaced by NaHCO₃ dosing. Increasing OLR levels were applied to both systems to define the operating limit conditions for each alkalization strategy. Complete details of the long-term operation of RM1 and RM2 were previously presented elsewhere (Fuess et al. 2021a). Despite the differences aforementioned, the operation of both reactors was finalized under equivalent conditions of OLR (10.0 kg-CODt m⁻³ d⁻¹; CODt = total chemical oxygen demand), HRT (24.0 h) and influent CODt (10 g L⁻¹), in addition to NaHCO₃ dosing. Biomass samples were further collected during reactor disassembling separately from the feeding zone (suspended cells in the bulk liquid) and from the bed region (PU strips—attached cells). Attached biomass was removed from the media by washing the strips with distilled water. Sample cleansing prior to DNA extraction was carried out immediately after collection, whilst the remaining amounts of biomass from the feeding zones were stored at 4 °C prior to use in MPP tests. Figure 1 shows constructive aspects of the AnSTBR and details of the biomass collection during the disassembling procedure.

Sugarcane molasses characterization

Fresh sugarcane molasses (Mol) was collected from a large scale annexed biorefinery (milling approximately 10 million tons of sugarcane per harvest) located in Pradópolis, São Paulo, Brazil. The samples were stored at 4 °C prior to use in both the feeding of the acidogenic reactor (see further description) and MPP tests. Fermented molasses (fMol) was collected from a bench-scale (2.0 L) thermophilic (55 °C) acidogenic AnSTBR operated under optimized conditions (OLR = 86.0 kg-CODt m⁻³ d⁻¹; HRT = 10.0 h) targeting enhanced biohydrogen production, as reported elsewhere (Fuess et al. 2021b). fMol samples were stored at -20 °C prior to use (feeding both RM1 and RM2 and as substrate in MPP tests) to interrupt the activity of cells washed-out from the acidogenic reactor and preserve the compositional aspects of the fermented

Fig. 1 Details of the AnSTBR systems (simplified sketch and experimental prior to inoculation) and biomass sampling from the feeding zone (FDZ) and structured bed (STB) during reactor disassembling. The same procedure was applied in RM1 and RM2. Legend: 1- reactor feeding, 2- effluent collection, 3- biogas collection



substrate. Detailed compositional aspects of Mol and fMol are presented in Table 1.

Methane production potential tests

MPP tests were carried out in 500 mL (nominal volume) Erlenmeyer flasks (with DIN standard PBT screw caps—GL45) filled with 100 mL of substrate and inoculum. Both Mol and fMol were diluted to obtain a COD of 10 g L^{-1} , simulating the terminal operating condition applied in the continuous reactors. Sugars were promptly solubilized once Mol was diluted. Meanwhile, fMol was centrifuged (9000 rpm, 5 min) prior to dilution as a strategy to remove cells (fermentative biomass) remaining from the acidogenic reactors, i.e. only the supernatant was used. In practical aspects, only the soluble organic content of both Mol and fMol was used as the substrate; however, the nomenclature “soluble COD (CODs)” was not used in this specific step because no $0.45 \mu\text{m}$ -filtration was carried out. Conversely, methanogenic sludge sources were centrifuged under similar conditions (9000 rpm, 10 min) to remove excess liquid phase, i.e. only the pellets were used, which were called wet sludge. In addition to biomass samples collected from the feeding zones of RM1 and RM2, the thermophilic methanogenic sludge used in the inoculation of both continuous reactors (identified as IN) was also assessed in MPP tests, following the same pre-centrifugation step. Assessing only biomass samples from the feeding zone was based on the performance of the continuous reactors, because the CODs was

Table 1 Compositional characterization of fresh and fermented molasses

Parameter	Sugarcane molasses	
	Fresh (Mol)	Fermented (fMol)
CODt (mg L^{-1})	$707 \pm 10^{\text{a,b}}$	$30,900 \pm 260$
CODs (mg L^{-1})	$707 \pm 10^{\text{a,b}}$	$28,203 \pm 927$
CHt (mg L^{-1})	$546 \pm 19^{\text{a}}$ (81.9%)	1717 ± 148 (6.5%)
Lactate (mg L^{-1})	—	$13,942 \pm 601$ (52.9%)
Ethanol (mg L^{-1})	—	329 ± 22 (2.4%)
Acetate (mg L^{-1})	—	1204 ± 160 (4.6%)
Propionate (mg L^{-1})	—	101 ± 4 (0.5%)
Butyrate (mg L^{-1})	—	2656 ± 168 (17.1%)
Caproate (mg L^{-1})	—	169 ± 31 (1.3%)
PheOH (mg L^{-1})	$13.0 \pm 0.1^{\text{a}}$ (2.1%)	667 ± 24 (2.7%)
SO_4^{2-} (mg L^{-1})	$2.4 \pm 0.1^{\text{a}}$	470 ± 5
CODs/ SO_4^{2-} (-)	299.6	60.0 ± 1.9
pH (-)	—	5.18 ± 0.07

CODt total chemical oxygen demand, CODs soluble chemical oxygen demand, CHt total carbohydrates, PheOH total phenols, SO_4^{2-} sulfate

^aValues in mg g^{-1}

^bCODt=CODs. Percentage values (%) indicate the relative participation of the compound in the CODs

majorly consumed ($> 70\%$) in this compartment in both reactors (Fuess et al. 2021a).

Once the volatile solids (VS) content was measured in each methanogenic consortium, the amount of wet sludge added to each flask was calculated according to three different F/M (or substrate-to-inoculum) ratios, namely, 0.4, 1.0 and 3.0 g-COD g⁻¹VS. The selected F/M ratios were based on the specific organic loading rate (sOLR) assessed in the continuous reactors, specifically in RM1 (Fuess et al. 2021a): the values 0.4 and 1.0 g-COD g⁻¹VS simulated the sOLR (obtained in terms of g-COD g⁻¹VSS d⁻¹) observed under stable operating conditions, using the total amount of biomass inside the reactor and the amount of biomass retained specifically in the feeding zone as the references, respectively. In turn, the value of 3.0 g-COD g⁻¹VS represented the maximum sOLR observed in the feeding chamber still associated with a stable operation. The three levels of the F/M ratio were assessed in fMol-fed reactors using all sludge sources (RM1, RM2 and IN). In the particular case of Mol, only sludge from RM2 was used. Finally, blank reactors were also monitored (duplicates) by replacing Mol or fMol with distilled water, using a fixed mass of 10 g of wet sludge per flask. For identification purposes, the nomenclature used to differentiate the conditions was formed by three parts: F/M ratio value-inoculum source(substrate type)—e.g. 0.4-RM1(fMol) and 3.0-RM2(Mol). The adopted experimental arrangement enabled comparing two- and single-stage AD schemes.

In addition to the definitions regarding both the F/M ratio and inoculum sources, reactor alkalization (including blanks) was based on NaHCO₃ dosing (0.25 g-NaHCO₃ g⁻¹COD), also using the continuous reactors as the reference (Fuess et al. 2021a). pH values in fMol and Mol after NaHCO₃ dosing reached ca. 6.70 and 7.60, respectively. After filling the flasks with alkalized substrate and sludge, the reactors were fluxed with ultra-high purity grade nitrogen for 5 min to establish conditions of anaerobiosis and sealed with rubber stoppers and plastic caps. The reactors were incubated in an orbital shaker (model Multitron PRO Incubator Shaker—Infors HT, Infors AG, Bottmingen-Basel, Switzerland) and monitored for periods ranging between 7 and 16 d, depending mainly on the temporal evolution of methane production. Temperature and agitation conditions were set respectively as 55 °C and 100 rpm.

Reactor monitoring was carried out through periodically sampling both the gas and liquid phases with 1 mL insulin syringes. In the specific case of biogas collection, the syringe was equipped with a Teflon body two-way valve (Supelco™ Analytical—Sigma Aldrich, Bellefonte, PA, USA). The sampling periodicity varied according to the response of the systems in terms of the cumulative methane production and COD decay. Experimental runs were finalized after observing coefficient of variation (CV) values lower than 5% for at least three consecutive points of the cumulative methane production, whilst the stabilization

of COD decay was used as a secondary reference. Liquid phase sampling was limited to 10% of the total substrate volume (10 mL). Biogas production was monitored by measuring the internal pressure of the flasks followed by the compositional analysis of the sample. Pressure measurements were carried out using a pressure gauge model TPR-16 (Desin Instruments, s.a., Barcelona, Spain), with an upper measurement limit of 500 mbar (50 kPa). Whenever required, pressure reliefs were conducted to prevent the occurrence of pressure values out of the measurement range.

Analytical methods, calculations and kinetic analysis

Overall liquid phase measurements included determining the pH, CODt, CODs, total carbohydrates (CHt), lactate, total phenols (PheOH), partial (PA) and intermediate (IA) alkalinity, volatile organic acids by titration (VOA_{tit}), volatile organic acids (VOA_{GC}; C2–C6) and solvents by gas chromatography (GC) and sulfate (SO₄²⁻). The periodic monitoring of the liquid phase in MPP tests included exclusively CODs measurements, whilst all remaining parameters were measured in non-diluted substrates (Table 1; except for PA, IA and VOA_{tit}) and by the end of the incubation periods (except for the CODt). COD (both total and soluble), pH and SO₄²⁻ measurements were based on the Standard Methods for the Examination of Water and Wastewater (APHA et al. 2012), whilst specific protocols were used for the remaining parameters: CHt (Dubois et al. 1956), lactate (Taylor 1996), PheOH (Buchanan and Nicell 1997), PA/IA (Ripley et al. 1986), VOA_{tit} (Kapp 1984) and VOA_{GC}/solvents (Adorno et al. 2014). Except for the case of CODt determination, samples were filtered in 0.45 µm syringe membranes (Chromafil GF/PET, Macherey–Nagel GmbH & Co. KG, Düren, Germany) prior to all remaining analyses.

Considering gas phase monitoring, biogas composition (nitrogen—N₂, methane—CH₄, carbon dioxide—CO₂ and hydrogen sulfide—H₂S) measurements were carried out in a GC equipped with a thermal conductivity detector (model GC-2014, Shimadzu Scientific Instruments, Japan) using hydrogen as the carrier gas (Lebrero et al. 2016). Specifically, the cumulative methane production up to time “n” (V_{CH₄,n}; NmL) was calculated as proposed elsewhere (Santos et al. 2019) using Eq. (1), in which the terms $iP_{pre-inj,n}$, $iP_{post-inj,n-1}$, $fCH_{4,n}$, $V_{headspace}$, T_0 , P_0 and T_{exp} , are, respectively, the internal pressure prior to sampling at time “n” (atm), the internal pressure after sampling at time “n—1” (atm), the methane fraction in biogas at time “n” (dimensionless), the volume of the headspace (L), the temperature at normal conditions (273.15 K), the pressure at normal conditions (1 atm) and the incubation temperature (55 °C = 328.15 K).

$$V_{\text{CH}_4,n} = \frac{\left\{ \sum_{i=0}^n \left[(iP_{\text{pre-inj},n} - iP_{\text{post-inj},n-1}) \times f_{\text{CH}_4,n} \right] \right\} \times V_{\text{headspace}} \times T_0}{P_0 \times T_{\text{exp}}} \quad (1)$$

The modified Gompertz model (Zwietering et al. 1990) (Eq. 2) was fitted to V_{CH_4} values to obtain the potential methane production (P_{CH_4} ; NmL), the maximum methane production rate (R_{CH_4} ; NmL h^{-1}) and the lag phase period (λ ; h). The maximum methane production rate was also calculated in specific terms (R_{CH_4}' ; NmL $\text{h}^{-1} \text{g}^{-1} \text{VS}$) by dividing R_{CH_4} by the amount of sludge (VS) used in the inoculation of each reactor. In Eq. (2), the terms $V_{\text{CH}_4}(t)$ and e are, respectively, the cumulative methane production (model) as a function of the incubation period (t) and the Euler's number (2.71828). Endogenous methane production (assessed in blank reactors) was discounted from methane production values obtained when processing fMol and Mol. Hence, V_{CH_4} values (both experimental and fitted) correspond to the net methane production. In parallel, kinetics of substrate consumption (CODs) was assessed through fitting the first-order decay model with residual (Rodrigues et al. 2003) (Eq. 3) to CODs temporal profiles, from which the residual CODs (CODs^R ; mg L^{-1}), the initial CODs (CODs^0 ; mg L^{-1}) and the first-order kinetic constant (k_1 ; h^{-1}) were estimated. The specific first-order kinetic constant (k_1' ; $\text{h}^{-1} \text{g}^{-1} \text{VS}$) was calculated analogously to R_{CH_4}' . Model fitting was carried out using the software Origin 2020 (OriginLab Corporation, Northampton, MA, USA) using the Levenberg–Marquardt algorithm. In the particular case of methane production, models were adjusted to experimental points collected up to fulfilling two criteria: [i] cumulative CODs removal efficiency (ER_{CODs} ; Eq. 4) higher than 70% and [ii] CODs removal efficiency between two consecutive points ($\text{ER}_{\text{CODs, interval}}$; Eq. 5) lower than 5%. In Eq. (4)–(5) the terms CODs_n , CODs_0 and CODs_{n-1} are the CODs (mg L^{-1}) measured at times “ n ”, 0 (zero) and “ $n-1$ ”, respectively.

$$V_{\text{CH}_4}(t) = P_{\text{CH}_4} \cdot \exp \left\{ -\exp \left[\frac{R_{\text{CH}_4} \cdot e}{P_{\text{CH}_4}} (\lambda - t) + 1 \right] \right\} \quad (2)$$

$$\text{CODs}(t) = \text{CODs}^R + (\text{CODs}^0 - \text{CODs}^R) \times \exp(-k_1 \times t) \quad (3)$$

$$\text{ER}_{\text{CODs},n} = 100 \times \left(\frac{\text{CODs}_n - \text{CODs}_0}{\text{CODs}_0} \right) \quad (4)$$

$$\text{ER}_{\text{CODs, interval}} = 100 \times \left(\frac{\text{CODs}_n - \text{CODs}_{n-1}}{\text{CODs}_0} \right) \quad (5)$$

Microbial community characterization by 16S rRNA gene sequencing

Biomass sampling and sample preparation.

Preliminary details of the microbial community sampling (targeting microbial characterization) were presented in Section “AnSTBR systems and thermophilic methanogenic consortia”. Samples from suspended cells in the feeding zone and attached cells from the bed region were obtained for both continuous methanogenic reactors (Fig. 1) during reactor disassembling and promptly stored at -20°C after cleansing, as described in the sequence. The thermophilic sludge (IN) used in the inoculation of RM1 and RM2 (collected from a thermophilic full-scale UASB processing sugarcane vinasse) was also characterized. Because the COVID-19 pandemic forced a relatively prolonged storage (4°C) of the remaining biomass samples from the feeding zones (ca. 6 months) and from IN (ca. 14 months) prior to MPP tests, aliquots of the stored samples were also analyzed (except for the case of RM1). Hence, a total of seven samples were processed and were identified by the sludge source (IN, RM1 or RM2) and the reactor region, i.e. feeding zone (FDZ) or structured bed (STB). The nomenclature “ps” (post-storage) was also used when pertinent. Table 2 brings details of the timeline followed to obtain biomass samples. All samples were subjected to 2–3 rounds of centrifugation (6000 rpm, 10 min) using a phosphate-buffered saline, which aimed to remove impurities (residual organic compounds). Finally, biomass pellets were stored at -20°C prior to further processing.

DNA extraction, 16S rRNA gene amplicon sequencing and bioinformatics

The procedures used in DNA extraction and 16S rRNA gene sequencing are presented elsewhere (Piffer et al. 2022)—refer to Supplementary material for details. Sequences were analyzed using quantitative insights into a microbial ecology’ pipeline (QIIME2 2021.4 release) (Bolyen et al. 2019). Demultiplexed paired-end sequencing reads were imported into the QIIME2, resulting in 358,837 joined sequences. Sequencing reads were ‘denoised’ using the ‘divisive amplicon denoising algorithm’ DADA2 (Callahan et al. 2016) plugin in QIIME2, which provided high resolution amplicon sequence variants (ASV) for downstream analysis by filtering out noises, correcting errors in marginal sequences, removing chimeric sequences and singletons and

Table 2 Timeline detailing biomass sampling periods. The text in bold refers to the nomenclature of the samples

Experiment	Timeline	Event	Description
Continuous reactors (Fuess et al., 2021a)	Before day 1	Inoculation	Reactor filling with macerated sludge
			Operation in closed cycle for 4 d
			Cleansing and storage (-20 °C) of one biomass sample from the inoculum (IN) for microbial characterization
			Storage (4 °C) of the remaining aliquot of the UASB-derived sludge
	Until day 230	Continuous operation of RM2	No biomass sampling
	Day 230	Disassembly of RM2	Reactor drainage, cleansing and storage (-20 °C) of biomass samples collected from the feeding zone (RM2-FDZ) and from the bed region (RM2-STB) of RM2
			Storage (4 °C) of the remaining aliquot of biomass collected from the feeding zone of RM2
	Until day 250	Continuous operation of RM1	No biomass sampling
	Day 250	Disassembly of RM1	Reactor drainage, cleansing and storage (-20 °C) of biomass samples collected from the feeding zone (RM1-FDZ) and from the bed region (RM1-STB) of RM1
			Storage (4 °C) of the remaining aliquot of biomass collected from the feeding zone of RM1
Activity interruption due to COVID-19 pandemic (~ 180 d)			
Batch tests (this study)	–	–	Cleansing and storage (-20 °C) of samples from the long-term stored inoculum (INps) and from the long-term stored biomass collected in the feeding zone of RM2 (RM2-FDZps) for microbial characterization ^a

^aAn unsuccessful preliminary round of batch tests using biomass collected from the feeding zone in RM1 decreased the availability of this sludge source, so that no sample was taken after the long-term storage in this particular case

dereplicating the resulting sequences. As a result, 313,539 sequences were obtained, ranging from 39,937 to 46,692 per sample ($n = 7$ in total), representing 719 ASV. The consensus sequences for the ASV were classified with a classify-sklearn classifier trained against the most recent SILVA 16S rRNA gene reference (release 138.1) database (Quast et al. 2013). In order to complete downstream diversity and composition analyses, sequences were rarefied to the lowest number of sequences per sample ($n = 39,937$ sequences). The sequences were submitted to the National Center for Biotechnology Information (NCBI; <http://ncbi.nlm.nih.gov>) under accession BioProject ID PRJNA778433.

Data analysis and visualization

All the statistical analyses were performed in R version 3.5.1 with R Studio environment, Version 1.3.1093. The biom file from QIIME2 was imported and analyzed through phyloseq-modified workflow (McMurdie and Holmes 2013). Taxon relative abundance bar charts were generated using custom R scripts and ggplot2 (v3.3.2) (Wickham 2016). A similarity matrix of weighted UniFrac distances was used for ordination by principal coordinates analysis (PCoA) of non-transformed relative abundance diversity datasets implemented in phyloseq and plotted using ggplot2 to visualize the

distribution of the microbial composition. Heat maps were generated using ampvis2 (v.2.6.5) (Andersen et al. 2018).

Results and discussion

Kinetics of methane production

The methane production potential of thermophilic microbial consortia processing sugarcane molasses was assessed under different F/M ratios, simulating different levels of substrate availability and also comparing single- and two-stage AD schemes. Temporal profiles for the cumulative methane production and CODs decay (experimental data) are depicted in Figs. 2 and 3, respectively. The kinetic parameters of the fitted models describing methane production and CODs decay are detailed in Tables 3 and 4, respectively. R_{CH_4} values followed an increasing pattern as the F/M ratio decreased (Table 3), regardless of the inoculum source and substrate type. This pattern characterizes an expected result, because higher biomass amounts (lower F/M ratios) will have the ability to promptly convert the available substrate. However, an inverse pattern was observed when considering R_{CH_4} values, which increased as the F/M ratio was increased for a given inoculum source (Table 3; considering specifically

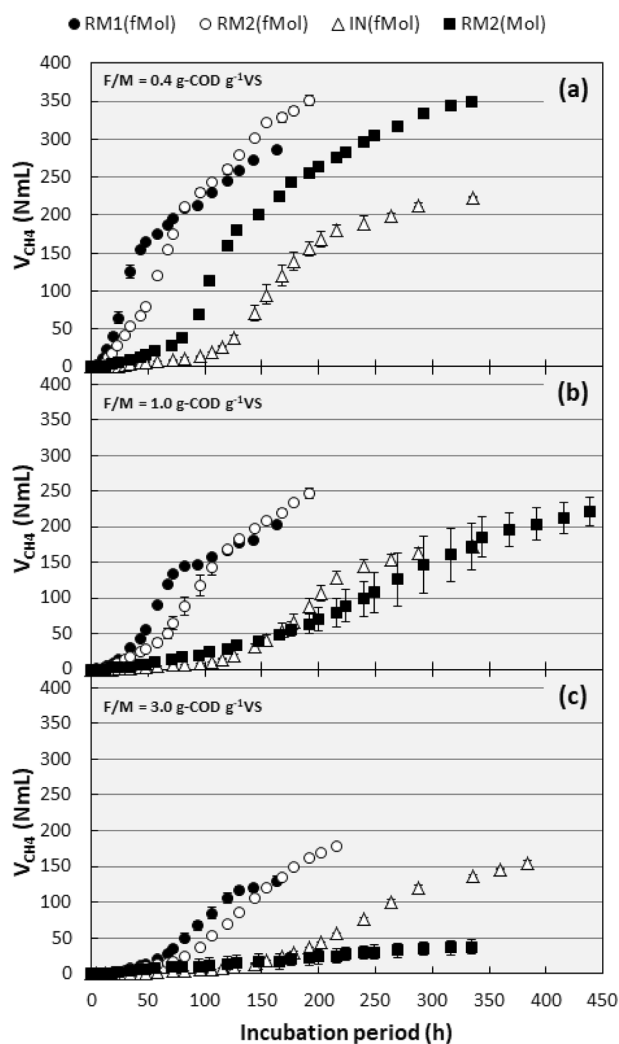


Fig. 2 Temporal evolution of methane production (V_{CH_4} , experimental data) for F/M ratios of **a** 0.4 g-COD g⁻¹VS, **b** 1.0 g-COD g⁻¹VS and **c** 3.0 g-COD g⁻¹VS

the use of fMol). This result indicates that lower amounts of biomass produced methane more efficiently, most likely due to the favoring of substrate conversion kinetics under excess substrate availability (prior to reaching the limit concentration, i.e. CODs, prior to inhibition). Under conditions of excess biomass (lower F/M ratios), a fraction of the cell most likely remained in a latency-like state, without effectively participating in methane evolution. From a kinetic perspective, lower substrate availability was the limiting factor for methanogens.

An integrated analysis including R_{CH_4} values obtained for RM1-derived sludge (this study) and specific methane production rate values in the different operating phases of RM1 (Fuess et al. 2021a) revealed a consistent variation pattern for R_{CH_4} (Fig. 4a–b), regardless of the operating mode, i.e. batch or continuous. The specific methane production

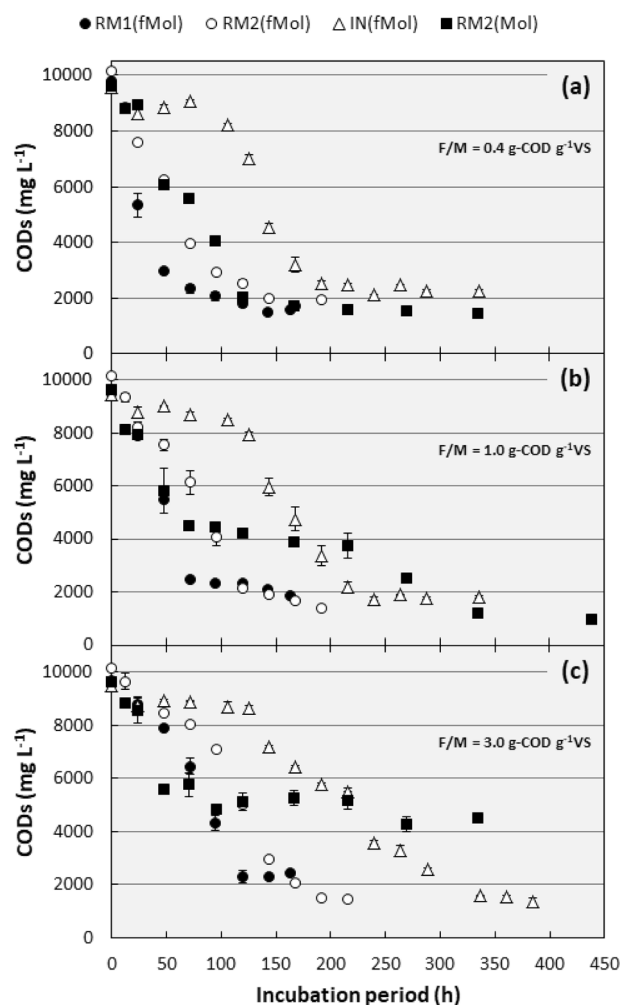


Fig. 3 Temporal profiles of CODs decay (experimental data) for F/M ratios of **a** 0.4 g-COD g⁻¹VS, **b** 1.0 g-COD g⁻¹VS and **c** 3.0 g-COD g⁻¹VS

rate in RM1 (at the different OLR levels) was calculated through re-arranging performance data previously presented, namely, the volumetric methane production rate and the biomass retention, providing a novel response not addressed in the base reference study (Fuess et al. 2021a). Details of this calculation are presented in the Supplementary data section. Despite the different measurement units for the F/M ratio (g-COD g⁻¹VS) and the sOLR (g-COD g⁻¹VSS d⁻¹), both parameters measure the amount of substrate available for conversion in the reactors on an equivalent basis, which supports the integrated comparison. R_{CH_4} varied linearly up to the maximum experimental sOLR measured in RM1 (~5.0 g-COD g⁻¹VSS d⁻¹; Fig. 4a), indicating that F/M ratio values assessed in batch tests (≤ 3.0 g-COD g⁻¹VS) characterized conditions of excess sludge compared to the amount of substrate, justifying the increasing trend previously highlighted. Differently from batch systems, in which the initial F/M is defined and fixed to a desired value, the

Table 3 Kinetic models fitted to experimental methane production profiles

Substrate	Sludge source	F/M ratio	Kinetic parameter				R ²
			P _{CH4}	R _{CH4}	R _{CH4} ^a	λ	
			(NmL)	(NmL h ⁻¹)	(NmL h ⁻¹ g ⁻¹ VS)	(h)	
Fermented molasses	RM1	0.4	207.11 ± 2.85	13.91 ± 0.55	5.68	10.84 ± 0.73	0.99
		1.0	173.38 ± 10.91	8.11 ± 0.60	7.70	26.63 ± 1.80	0.99
		3.0	165.48 ± 9.37	3.91 ± 0.11	11.91	46.92 ± 1.20	0.99
	RM2	0.4	301.30 ± 7.76	9.34 ± 0.24	3.69	21.39 ± 0.89	0.99
		1.0	279.24 ± 7.96	5.78 ± 0.14	5.70	40.62 ± 1.30	0.99
		3.0	234.71 ± 5.37	3.76 ± 0.05	11.13	68.27 ± 0.84	0.99
	IN	0.4	235.10 ± 14.86	4.94 ± 0.18	1.96	102.65 ± 1.90	0.99
		1.0	207.48 ± 13.61	3.43 ± 0.13	3.39	120.65 ± 2.38	0.99
		3.0	183.92 ± 5.58	2.16 ± 0.04	6.42	142.01 ± 1.67	0.99
Fresh molasses	RM2	0.4	263.81 ± 20.36	7.69 ± 0.60	3.08	65.05 ± 2.99	0.99
		1.0	298.75 ± 27.43	2.03 ± 0.08	2.00	102.50 ± 6.46	0.97
		3.0 ^b	—	—	—	—	—

P_{CH4} potential methane production, R_{CH4} maximum methane production rate, R_{CH4}^a specific maximum methane production rate, λ- lag phase period

^aCalculated using average values

^bNo model was fitted in this condition

Table 4 Kinetic models fitted to experimental CODs decay profiles

Substrate	Sludge source	F/M ratio	Incubation period (h)	Kinetic parameter				R ²
				CODs ⁰	CODs ^R	k ₁ (× 10 ⁻²)	k ₁ ^a (× 10 ⁻²) ^a	
				(mg L ⁻¹)	(mg L ⁻¹)	(h ⁻¹)	(h ⁻¹ g ⁻¹ VS)	
Fermented molasses	RM1	0.4	0–163	9817 ± 136	1566 ± 81	3.392 ± 0.155	1.384	0.99
		1.0	24–163 ^b	8050 ± 528	1764 ± 414	3.058 ± 0.787	2.904	0.95
		3.0	72–163 ^c	6525 ± 211	2027 ± 239	3.767 ± 0.687	11.478	0.95
	RM2	0.4	0–192	10,308 ± 182	996 ± 267	1.456 ± 0.121	0.575	0.99
		1.0	0–48	10,188 ± 152	6525 ± 806	2.766 ± 1.102	2.728	0.96
			48–192	7817 ± 316	349 ± 630	1.537 ± 0.319	1.516	0.96
		3.0	0–96	10,046 ± 197	5867 ± 1751	1.109 ± 0.759	3.282	0.91
			96–216	7209 ± 162	659 ± 289	2.014 ± 0.231	5.960	0.99
			216–384	5351 ± 164	1091 ± 225	1.620 ± 0.248	4.817	0.97
	IN	0.4	106–336 ^d	8566 ± 258	2066 ± 159	2.415 ± 0.254	0.956	0.97
		1.0	106–336 ^d	9061 ± 307	961 ± 382	1.357 ± 0.189	1.343	0.96
		3.0	125–216 ^d	8604 ± 78	5144 ± 166	2.489 ± 0.297	7.401	0.99
Fresh molasses	RM2		216–384	5351 ± 164	1091 ± 225	1.620 ± 0.248	4.817	0.97
		0.4	0–334	10,204 ± 504	888 ± 547	1.182 ± 0.221	0.473	0.96
		1.0	0–216	9723 ± 245	3632 ± 223	2.112 ± 0.274	2.083	0.96
			216–438	3737 ± 178	1031 ± 153	2.954 ± 0.997	2.995	0.96
		3.0	0–334	9945 ± 330	4678 ± 197	2.307 ± 0.386	6.827	0.91

CODs⁰ initial CODs, CODs^R residual CODs, k₁ first-order kinetic constant, k₁^a specific first-order kinetic constant

^aCalculated using average values

^bLinear decay between 0 and 24 h

^cLinear decay between 0 and 72 h: CODs(t) = (9796 ± 107) — (43.999 ± 2.381) × t (R² = 0.97)

^dThe CODs consumption was not significant until the lower limit of the interval was reached

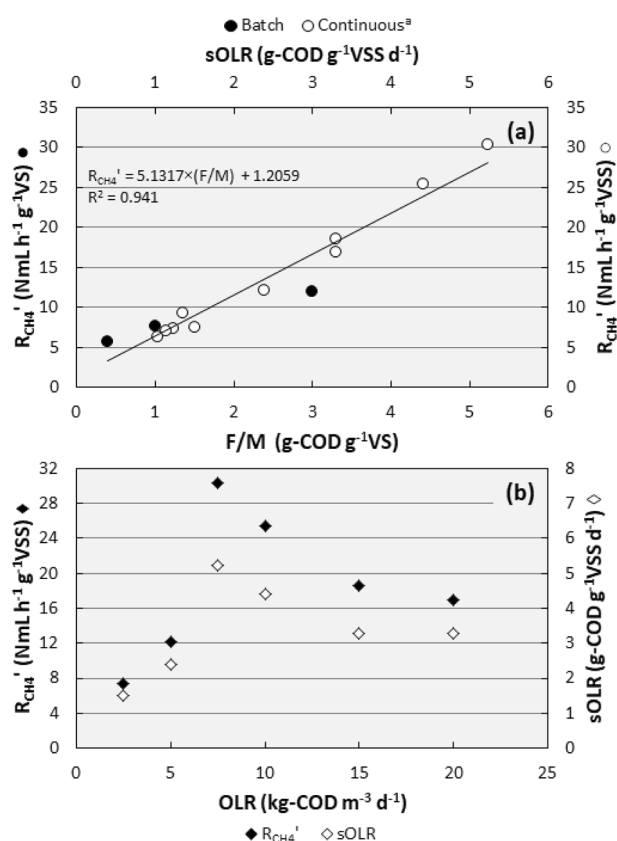


Fig. 4 Correlations between methane production and substrate availability: **a** maximum specific methane production rate (R_{CH_4}') as function of the F/M ratio (batch reactors) and specific organic loading rate (sOLR, continuous reactor) and **b** R_{CH_4}' as function of the organic loading rate (OLR) and sOLR (continuous reactor). Note: ^aValues calculated through re-arranging performance data collected from RM1 presented elsewhere (Fuess et al. 2021a)

sOLR varies as a function of both the applied OLR and the amount of retained biomass within continuous reactors (Fuess et al. 2021a; Anzola-Rojas et al. 2015), explaining the varying values in RM1. Interestingly, specific methane production was subjected to some degree of inhibition with the increase in the OLR (Fig. 4b), but not as a direct consequence of excess substrate availability. With the continuous retention of biomass within the continuous system, increasing the OLR (up to $20\ kg-COD\ m^{-3}\ d^{-1}$) was not sufficient to proportionally increase the sOLR (Fig. 4b), explaining the lower R_{CH_4}' values. Hence, the F/M ratios assessed in the batch tests may represent two operating situations of the continuous reactors: a short-term operation characterized by a low OLR under equally low biomass retention levels or a long-term operation with relatively high OLR and excess sludge retention.

An inverse pattern was observed when using fresh molasses as the substrate, in which lower R_{CH_4}' values were observed at higher F/M ratios (Table 3), until reaching a

marked inhibition of the methanogenic activity in the highest F/M ratio level ($3.0\ g-COD\ g^{-1}VS$) assessed (Fig. 2c). The carbohydrate-rich composition of Mol (> 80% of the CODs; Table 1) promptly favored the growth of fermentative bacteria, diverting the electron flow, i.e. CODs consumption, from methanogenesis towards cell synthesis. Similar results were observed in previous studies on the biodigestion of highly fermentable substrates, such as sugarcane juice and molasses (Fuess et al. 2021c; Vilela et al. 2021) and glycerol-supplemented sugarcane vinasse (Borges et al. 2022). In all cases, the impairment of the methanogenic activity was not impeditive for achieving relatively high substrate consumption levels, reinforcing the diversion towards cell synthesis. Comparing methane evolution patterns (and the kinetic parameters estimated) for the RM2(fMol) and RM2(Mol) conditions (Fig. 2 and Table 3) supports this hypothesis, because the lowest R_{CH_4} and R_{CH_4}' values were always associated with the non-fermented molasses (for a given F/M ratio). In practical aspects, this discrepancy demonstrated that phase separation increases the capability of anaerobic systems to withstand organic overloading conditions (simulated by high F/M ratios) far beyond the limits achieved in single-stage AD. The experimental arrangement indicated that two-stage schemes withstand at least a threefold higher organic load compared to single-stage ones, because methane production collapsed in condition 3.0-RM2(Mol)—despite the observation of methane production in this case (Fig. 2c), terminal pH values lower than 5.0 were observed, i.e. an evidence of VFA buildup. Hence, starting up a second-stage methanogenic reactor should require lower amounts of sludge and would demand a shorter period compared to single-stage schemes.

Shifting the analysis perspective, the comparison between the different biomass sources assessed (in a fixed F/M ratio) indicated that R_{CH_4}/R_{CH_4}' values always followed the order RM1 > RM2 > IN (Table 3), suggesting that biomass collected in RM1 was better adapted to the substrate and operation (incubation) conditions than the other inocula. Regardless of the differences observed in the long-term performance of RM1 and RM2, as a direct consequence of different alkalization strategies, CODs removal efficiency in the feeding zone of both reactors was equivalent, i.e. 74.9% (RM1) and 71.8% (RM2) (Fuess et al. 2021a), indicating that equivalent methane evolution patterns should be observed in MPP tests. Differences in the prevailing microbial communities (also resulting from the different alkalization approaches) may explain the observed patterns (Section “Microbial community composition”). It is worth highlighting the significant discrepancy in the performance of methanogenesis of IN relative to RM1/RM2, mainly with respect to much longer lag phase periods (Table 3), which resulted from the continuous adaptation of the thermophilic biomass to fermented molasses in RM1 and RM2 for periods

of at least 230 d. Besides the microbial differences (Section “[Microbial community composition](#)”), the OLR applied in the UASB from which IN was collected ($\sim 5.0 \text{ kg-CODt m}^{-3} \text{ d}^{-1}$) is half of that applied in RM1 and RM2 ($10 \text{ kg-CODt m}^{-3} \text{ d}^{-1}$) when microbial community samples were obtained, indicating a higher capability to withstand high substrate availability levels in the latter. However, the non-adaptation of the biomass to the incubation conditions was attenuated at high substrate availability: when comparing the different inocula (fMol-fed reactors), the higher the F/M ratio, the lower the variation coefficient calculated for R_{CH_4} (or R_{CH_4}') values, i.e. 30%, 40% and 50% respectively for the F/M ratios of 3.0, 1.0, 0.4 g-COD g⁻¹VS.

Kinetics of substrate consumption

Differently from methane production, CODs decay followed distinct kinetic patterns, according to both the sludge source and F/M ratio (Fig. 3 and Table 4). Single kinetic decays were observed only when assessing an F/M ratio of 0.4 g-COD g⁻¹VS in biomass samples from RM1 and RM2 (considering both fMol and Mol in the latter). The relative excess sludge amount in such cases favored a prompt substrate consumption, with a wide advantage for the case of RM1 ($k_1' = 1.384 \times 10^{-2} \text{ h}^{-1} \text{ g}^{-1}\text{VS}$; Table 4). Single decay patterns were also observed in the case of IN (F/M ratios of 0.4 and 1.0 g-COD g⁻¹VS); however, only after 106 h of experimental run (Fig. 3a-b), suggesting a progressive adaptation of the inoculum to the incubation conditions. In all remaining cases, substrate decay followed two distinct patterns, either with an initial linear decay (condition 3.0-RM1(fMol)) or with two sequential exponential decays (conditions 1.0-RM2(fMol), 3.0-RM2(fMol), 3.0-IN(fMol) and 1.0-RM2(Mol)) (Table 4). In these conditions, no trend repetition was observed, i.e. while k_1' in the first decay period exceed that of the second in 1.0-RM2(fMol) and 3.0-IN(fMol), the opposite was observed in 3.0-RM2(fMol) and 1.0-RM2(Mol).

For a given sludge source (fed with fMol), the higher the F/M ratio, the higher the value of k_1' (Table 4), repeating the behavior previously observed and described for R_{CH_4}' (Table 3). Complementing the hypothesis of higher metabolic efficiency towards methane production, substrate consumption also includes processes other than methanogenesis, such as cell synthesis. Hence, the high k_1' values estimated under lower biomass amounts (high F/M ratios) most likely resulted from both metabolic steps. In fact, relatively high biomass concentrations, i.e. $M > F$, are recommended when assessing the methanogenic activity of anaerobic sludge sources to minimize the negative effects of excess biomass synthesis over methane production (Aquino et al. 2007; Holliger et al. 2016). The establishment of enhanced cell growth as a significant electron (substrate) sink also justifies the

CODs consumption under unfavorable conditions to methanogenesis when using fresh molasses, mainly under high F/M ratios, i.e. 1.0 and 3.0 g-COD g⁻¹VS. These values are much higher than the ones previously recommended for efficient methane production from easily fermentable substrates, i.e. inoculum-to-substrate ratio higher than 4.0 (or F/M ratio lower than 0.25) (Holliger et al. 2016). Revisiting the parallel with reactor startup, efficient substrate conversion rates coupled to short latency periods will only be achieved when sufficient amounts of biomass are available (Haider et al. 2015). Sulfate reduction could have also contributed to the CODs decay, because sulfidogenesis can be associated with the complete oxidation of organic compounds. However, a significant deviation of electrons towards sulfate reduction should only be observed at high sulfate availability (COD/sulfate < 25.0) (Kiyuna et al. 2017), which was not observed for molasses, regardless of the pre-fermentation (COD/sulfate > 60.0; Table 1).

Similarly to the case of R_{CH_4}' , k_1' values followed the same order for a given F/M ratio, i.e. RM1 > RM2 > IN (Table 4), so that the discrepancy may be understood from an analogous perspective, focusing on microbial characterization aspects (Section “[Microbial community composition](#)”). It is worth highlighting that enhanced CODs decay in Mol-fed reactors at lower F/M ratios ($\leq 1.0 \text{ g-COD g}^{-1}\text{VS}$) was anticipated in comparison with IN (Fig. 3a-b), despite using fermented molasses in the latter. A similar response occurred in the case of methane evolution from fresh molasses at the lowest F/M ratio (0.4 g-COD g⁻¹VS; Fig. 2a), in which the exponential production phase anticipated the one from fermented molasses in reactors inoculated with IN. These results may be explained by the long-term ($\geq 230 \text{ d}$) adaptation of IN in the continuous reactors, which attenuated negative effects of enhanced VFA production over methanogens when processing Mol under higher sludge concentrations. Hence, the efficacy of phase separation in high-rate AD systems cannot be assessed separately from the degree of adaptation of the sludge to the desired operating conditions.

Breakdown of liquid phase constituents in biodigested molasses

Details of the liquid phase constituents at the end of the incubation periods are depicted in Fig. 5, including the terminal pH, the balance between bicarbonate alkalinity and VFA accumulation and the breakdown of the CODs. Different terminal pH values were observed according to the type of substrate, i.e. approximately 8.0 and 7.5 for fermented (regardless of the F/M ratio) and fresh (except for the F/M ratio of 3.0 g-COD g⁻¹VS) molasses (Fig. 5a). While a marked increase relative to the pH of fermented molasses supplied with NaHCO₃ prior to incubation (< 7.0) was observed in

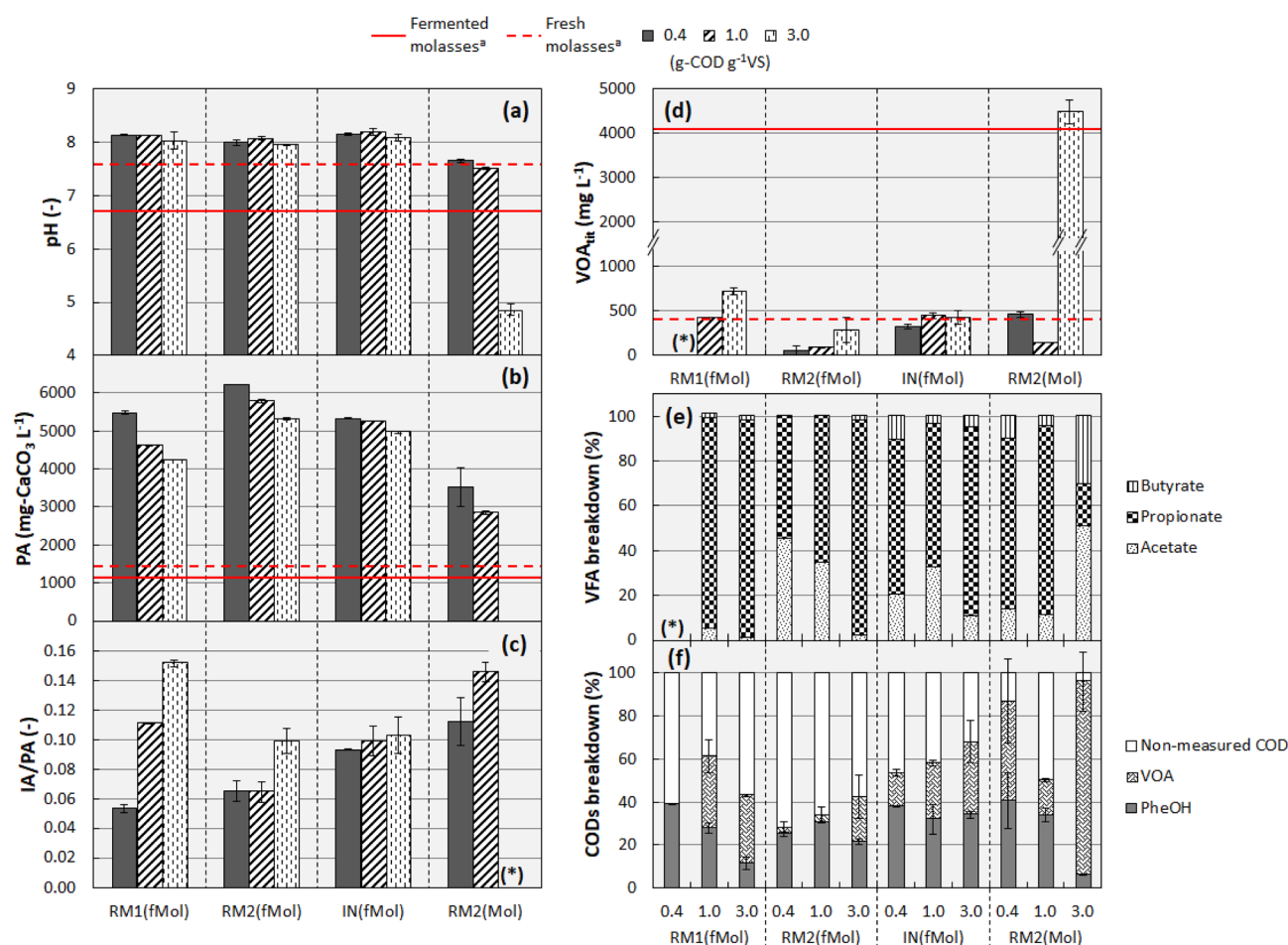


Fig. 5 Characterization of biodigested molasses at the end of the incubation periods: **a** pH, **b** partial alkalinity (PA), **c** intermediate-to-partial alkalinity (IA/PA) ratio, **d** concentration of volatile organic

acids by titration (VOA_{tit}), **e** breakdown of VFA (using gas chromatography) and **f** breakdown of the CODs. Legend: ^aAfter NaHCO₃ dosing, (*) indicates null values

fMol-fed reactors, initial and terminal pH values were equivalent (~7.5) when using fresh molasses (F/M ratios of 0.4 and 1.0 g-COD g⁻¹VS). In the first case, the non-occurrence of enhanced sugar fermentation positively impacted the balance of partial (bicarbonate) alkalinity (Fig. 5b), with values 1.5-to-2.0-fold higher than in reactors fed with fresh molasses (considering equivalent F/M ratios). These patterns (associated with the higher terminal pH values) indicate an excess of NaHCO₃ supply in second-stage methanogenic systems, suggesting the possibility to use lower doses without impairing the performance of methanogenesis. In fact, dosing 0.20 g-NaHCO₃ g⁻¹COD in RM1 (continuous reactor) did not negatively impact system performance (Fuess et al. 2021a). Conversely, removing NaHCO₃ supply triggered the inhibition of methanogenesis in a continuous reactor (also an AnSTBR) fed with fresh molasses, so that a minimum dose of 0.25 g-NaHCO₃ g⁻¹COD was defined as the limiting condition to maintain stable performance levels (Oliveira et al. 2020). As a side-effect remark

of the experiment, dosing equivalent amounts of NaHCO₃ (0.25 g g⁻¹COD) in methanogenic systems fed with fresh and fermented substrates results in excess bicarbonate alkalinity in the latter (up to 200% higher concentrations than in the first). In practical aspects, NaHCO₃ dosing in two-stage biodigestion systems have potential to be decreased to economically cost-competitive levels, improving mechanisms to ensure the maintenance of operating stability in industrial scale plants.

The intermediate-to-partial alkalinity (IA/PA) ratio measured at the end of the incubation periods reached values lower than 0.16 in all cases in which methane production was not inhibited (Fig. 5c), indicating much higher participation of bicarbonate-derived alkalinity than the one related to organic acids (despite the aforementioned discrepant patterns). Overall, the lower the F/M ratio, the lower the IA/PA value for a given source of inoculum (Fig. 5c), characterizing a relative accumulation of organic acids at high F/M ratios (VOA_{tit} usually below 500 mg L⁻¹; Fig. 5d). Residual

VOA_{tit} concentrations measured in reactors inoculated with biomass from RM1 (426 and 718 mg L⁻¹ in F/M ratios of 1.0 and 3.0 g-COD g⁻¹VS, respectively; Fig. 5d) were higher than those in reactors using RM2-derived sludge (<283 mg L⁻¹; Fig. 5d) most likely due to differences in the incubation periods. While methane production (CV < 5%; Section “Methane production potential tests”) and the associated CODs decay met the criteria for finalizing the experimental runs within 163 h in the case of RM1, reactors inoculated with sludge from RM2 required longer periods, which varied from 192 h (F/M = 0.4 and 1.0 g-COD g⁻¹VS) to 216 h (F/M = 3.0 g-COD g⁻¹VS). Enhanced acidification (VOA_{tit} = 4500 mg L⁻¹; Fig. 5d) was only observed in condition 3.0-RM2(Mol), in which the inhibition of methanogenesis (Fig. 2c) was associated with terminal pH values lower than 5.0 (Fig. 5a), as well as with the full consumption of the partial alkalinity (Fig. 5b).

The distribution of residual VFA concentrations (Fig. 5e) indicated distinct patterns according to the success in the establishment of methanogenesis. Propionate was the prevailing metabolite in conditions in which methane production evolved over time, regardless of differences in the kinetic patterns and type of substrate (Fig. 5e). In these cases, propionate oxidation into acetate and the subsequent oxidation of acetate into CO₂ and hydrogen most likely characterized the limiting steps, slowing down methane evolution via the hydrogenotrophic pathway, i.e. the primary methanogenic pathway established in all reactors, as supported by the microbial characterization of the inocula (Section “Microbial community composition”). Differently, acetate buildup was observed in condition 3.0-RM2(Mol), i.e. the only case in which methane production was impaired (Fig. 5e). In addition to the failure in acetate oxidation, which may have also limited the activity of hydrogenotrophic methanogens in this case, the activity of acetoclastic methanogens was equally inefficient to convert the excess acetate available, most likely due to the inhibition by the low pH (Hao et al. 2012).

Apart from the accumulation of VFA, the breakdown of the terminal CODs indicated consistent participation of total phenols (PheOH) in the group of non-converted organic matter, with most values within the range of 20 to 40% (Fig. 5f). PheOH usually accounts for a significant fraction of the residual COD in biodigested sugarcane-derived substrates, such as vinasse (Santos et al. 2019) and molasses (Fuess et al. 2021a). The presence of highly recalcitrant melanoidins (from the Maillard reaction between sugars and proteins) and colorants (except caramels) configures the main natural sources of phenols in these substrates (Chandra et al. 2018; Mohana et al. 2009), providing their characteristic dark brown color. It is noteworthy that the participation of PheOH in the CODs increased in all assessed conditions, i.e. from 2.1–2.7% (non-biodigested substrates; Table 1) to

(6.2–40.5% (biodigested substrates; Fig. 5e), with the highest terminal values usually observed at excess biomass concentrations (F/M = 0.4 and 1.0 g-COD g⁻¹VS). This pattern most likely resulted from a more intense hydrolytic activity, as previously observed in the dark fermentation of sugarcane vinasse (Piffer et al. 2021). The (partial) breakdown of complex structures (such as those found in melanoidins) implies the release of phenolic compounds to the soluble phase, explaining the high participation of PheOH in the residual CODs.

Microbial community composition

The composition of the microbial community samples collected at the beginning (inoculum) and at the end of the operation of the continuous methanogenic reactors (RM1 and RM2 in two different sampling points: FDZ and STB) and further used in MPP tests was determined using 16S rRNA gene amplicon sequencing. Prior to detailing the microbial composition of the samples and making inferences about the primary metabolic pathways, PCoA-based comparisons provided bases to understand differences and similarities among all samples from an overall perspective (Fig. 6), considering the interference of: [i] different regions (FDZ and STB) within a given reactor, [ii] different reactors (RM1 and RM2), [iii] the specialization of the sludge (IN) after long-term (≥ 230 d) operating periods, and [iv] the impacts of sample storage. It is worth highlighting that analyzing the impacts of sample storage was simply a parallel investigation of the study, in an effort to understand potential failures in MPP tests (which were not observed). PCoA results were interpreted considering both the total microbial community (Archaea and Bacteria domains; Fig. 6a) and exclusively the archaeal community (Fig. 6b).

Considering the total microbial community, while little differentiation was observed between different compartments of a given reactor (regardless of the storage in the case of RM2-FDZ), a marked distance was identified between RM1- and RM2-derived samples (Fig. 6a), which were grouped separately. Although the operation of both reactors was finalized under equivalent conditions of OLR (10.0 kg-CODt m⁻³ d⁻¹), HRT (24.0 h) and alkalization (NaHCO₃ dosing), RM2 was subjected to events of enhanced VFA accumulation when dosing NaOH in previous steps of the operation (Fuess et al. 2021a), which may explain the observed discrepancy. Total microbial communities from both reactors were also very different from IN, considering particularities of each AD system in which samples were collected (IN vs. RM1/RM2): reactor type (UASB vs. AnSTBR), substrate type (vinasse vs. fermented molasses) and OLR (5.0 vs. 10.0 kg-CODt m⁻³ d⁻¹). In the case of IN, the long-term storage (14 months) impacted the microbial composition to a higher extent than in the case of RM2-FDZ.

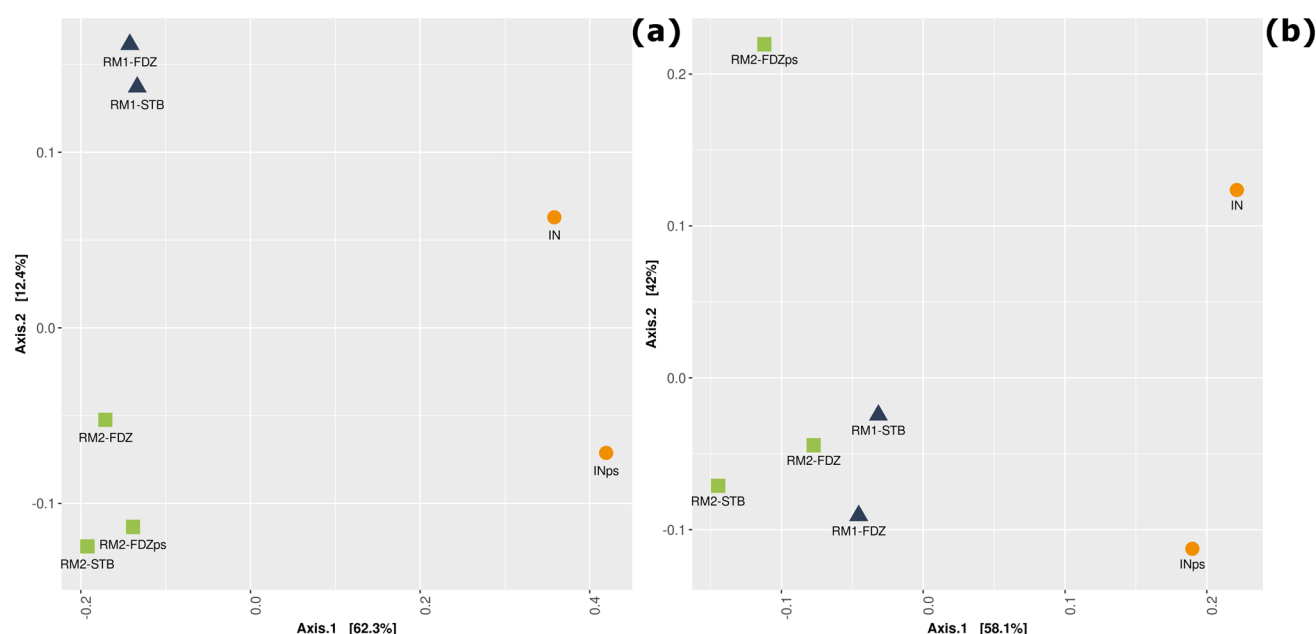


Fig. 6 Principal coordinates analysis (PCoA, UniFrac) plot for **a** total community and **b** archaeal community. Legend: samples related to the inoculum (filled circle), RM1 (filled triangle) and RM2 (filled square). “ps” indicates post-storage

Limiting the analysis to the archaeal community (Fig. 6b), samples collected from RM1 and RM2 were very similar, regardless of the compartment. This particular result confirms that variations in the bacterial community of RM2 triggered the discrepancy previously observed in the total microbial communities of both RM1 and RM2 (Fig. 6a). Marked difference was still observed between samples from RM1/RM2 and IN. Interestingly, long-term storage impacted the archaeal community to a much higher extent than the different growth conditions (suspended vs. attached) along the reactors, using the differences between samples RM2-FDZ and RM2-FDZps as the references. Hence, the relative similarity between microbial communities from different compartments of given reactor (regardless of considering only Archaea or both Archaea and Bacteria domains) answers one of the questions raised in Introduction, indicating that a more homogeneous microbial distribution is expected to occur in the biodigestion (methanogenesis) of pre-fermented substrates. However, the unbalanced conversion of organic matter observed in the continuous reactors, i.e. > 70% in the FDZ of both systems (Fuess et al. 2021a), leads to further questions: [i] How metabolically active are the microbial communities located in the bed region? [ii] Could the fixed bed simply be a “cell retention barrier” in the proposed reactor configuration?

Details of the microbial characterization of the biomass samples at the phylum and genus levels are depicted in Fig. 7, also considering the total microbial community (Archaea and Bacteria domains; Fig. 7a-b) and an isolated

analysis of the Archaea domain (Fig. 7c-d). Additionally, heat maps for ASV and genera are presented in the Supplementary data section for further reference. Overall, the Firmicutes phylum prevailed in all samples collected from RM1 and RM2 (Fig. 7a), characterizing a marked different pattern compared to IN and INps. While the phyla Campylobacterota, Firmicutes and Proteobacteria were identified relatively in equivalent abundances in IN, the Proteobacteria phylum prevailed in INps (Fig. 7a). The phylum Euryarchaeota, in which most methanogens are included (Lyu and Liu 2018), was the primary archaeal group identified in all samples (Fig. 7c; except for the case of INps, in which the phylum Halobacterota was also relevant). The *Methanothermobacter* genus was the most abundant methanogen identified in all samples collected from RM1 and RM2 (relative abundance—RA = 56.7–79.7% among Archaea; Fig. 7d), regardless of the growth condition, i.e. suspended (FDZ) or attached (STB). This result, in association with the identification of the *Methanosaeta* genus exclusively in samples from RM1 and RM2 (RA = 7.8–25.5% among Archaea; Fig. 7d), largely explain the similarity observed for archaeal communities (pre-storage) from both continuous reactors (Fig. 6b). The storage-dependent difference observed for samples RM2-FDZ and RM2-FDZps resulted from shifts in minor ASV associated with each genus identified. *Methanothermobacter* also prevailed in the thermophilic sludge (IN) used in the inoculation of RM1 and RM2 (RA = 64.7% among Archaea) and was followed by the *Methanoculleus* genus as the second most abundant archaeal group

(RA = 18.3%) (Fig. 7d). However, the latter (RA = 48.9%) surpassed the first (RA = 41.3%) after the long-term storage of IN (INps; Fig. 7d), which effectively characterized the primary distribution of methanogens during MPP tests. The significant presence of the *Methanoculleus* genus in both IN and INps samples (and the absence of the *Methanosaeta* genus) explains their difference relative to RM1 and RM2 (Fig. 6b), whilst the increase in the RA for *Methanoculleus* in INps explains the distance relative to IN. *Methanosarcina* (RA < 10.5%) and *Bathymarchaeia* (RA < 9.0%) genera were also identified in all biomass samples (Fig. 7d), but at lower RA levels compared to the previous groups.

Interestingly, the most abundant methanogens identified in all samples, i.e. *Methanothermobacter* (Hao et al. 2012; Cheng et al. 2011; Wasserfallen et al. 2000) and *Methanoculleus* (Dyksma et al. 2020; Manzoor et al. 2016; Shigematsu et al. 2004) genera, grow autotrophically on H₂ and CO₂, which may sound contradictory when considering two-stage AD schemes. Nearly 90% of the total carbohydrates found in fresh molasses were converted during dark fermentation (Fuess et al. 2021b) prior to feeding RM1 and RM2, markedly eliminating the prompt provision of both H₂ and CO₂ by acidogenic populations in the methanogenic units. However, the predominance of both genera in the inoculum (combined RA > 80% among Archaea; Fig. 7d) associated with the relatively low availability of acetate (4.6% of the CODs; Table 1) in fermented molasses, i.e. much lower levels than those of lactate (52.9%) and butyrate (17.1%) (Table 1), required the establishment of a strong association between acetogens, syntrophic acetate-oxidizing bacteria (SAOB) and hydrogenotrophic methanogens (HM) in both RM1 and RM2. In particular, the association between SAOB and HM has been systematically pointed to replace acetoclastic methanogenesis under thermophilic conditions (Hao et al. 2012; Cheng et al. 2011; Dyksma et al. 2020), which supports the occurrence of the metabolic sequence proposed. Compared to acetoclastic methanogenesis, methane production from the association between SAOB and HM is thermodynamically favored at high temperatures (Dolfing 2014). In addition, previous investigations indicated the prevalence of the latter pathway at relatively low acetate availability (Shigematsu et al. 2004; Petersen and Ahring 1991), which is in accordance with the metabolite distribution profile in fermented molasses (Table 1).

Differently from the participation of methanogens, notably characterized by a few representative genera (Fig. 7d), a higher diversity of acetogenic and acetate-oxidizing groups at much lower abundance levels was identified. The oxidation of fermentation metabolites into acetate was most likely carried out by the genera *Pelotomaculum* (RA = 0.9–6.6%, except in IN and INps), *Syntrophothermus* (RA = 0.6–3.8%, except in IN and INps), *Cloacimonadacea*

W5 (RA = 1.3–5.9%, except in IN and INps), *Syntrophomonas* (RA = 0.9–4.6%, except in IN and INps), and *Thermodesulfovibrio* (RA = 0.3–4.3%, all samples). Butyrate oxidation into acetate was previously associated with the *Syntrophothermus* (Sekiguchi et al. 2000) and *Syntrophomonas* (McInerney et al. 1981) genera, whilst both *Pelotomaculum* (Imachi et al. 2007) and *Cloacimonadacea* W5 (Dyksma and Gallert 2019) genera were involved in propionate oxidation. In all cases, syntrophic associations with methanogens were obligate to maintain low acetate levels, characterizing a different aspect of the cases reported in this study, because SAOB most likely consumed most of the available acetate. Interestingly, propionate availability in fermented molasses was much lower than all other fermentation metabolites (0.5% of the CODs; Table 1), so that an intermediate step releasing propionate was most likely established prior to acetogenesis. In particular lactate fermentation into acetate and propionate by species belonging to the *Clostridium* and *Veillonella* genera was identified as a relevant pathway in methanogenic environments (Seeliger et al. 2002; Stams et al. 1998), potentially fitting the proposed intermediate step. An ASV associated with the Veillonellales-Senomonadales order was identified primarily in biomass samples collected from the FDZ of both RM1 and RM2 (RA = 6.1–13.5%), which may support this hypothesis. Alternatively, *Thermodesulfovibrio* may have mediated the syntrophic degradation of lactate, which has been previously reported to occur in both the absence of sulfate and co-culture with hydrogenotrophic methanogens (Sekiguchi et al. 2008). The low sulfate availability in molasses (COD/sulfate > 60.0; Table 1) does not support the long-term persistence of such a sulfate-reducing group in the continuous reactors, suggesting the establishment of an alternative metabolic pathway to maintain cell growth.

Following the SAOB-HM pathway, genera responsible for mediating syntrophic acetate oxidation included *Thermoacetogenium* (RA = 1.4–2.3%, RM1-FDZ and RM1-STB), *Mesotoga* (RA = 3.4–7.8%, all samples) and *Pseudothermotoga* (RA = 0.2–2.1%, except in IN and INps). The *Thermoacetogenium* genus has been associated with high metabolic flexibility, according to the environmental conditions offered (Hattori et al. 2000). On one hand, acetate oxidation has been reported to occur in co-culture with hydrogenotrophic microorganisms or associated with sulfate reduction in pure culture. On the other hand, acetate production from alcohol fermentation or autotrophically from H₂ and CO₂ (homoacetogenesis) has also been observed. Acetate oxidation by *Pseudothermotoga* has been observed only in the presence of thiosulfate or methanogens (Balk et al. 2002), whilst in the case of *Mesotoga* the acetate-oxidizing pathway is still uncharacterized (Nobu et al. 2015). However, investigations using labelled carbon suggested that the latter

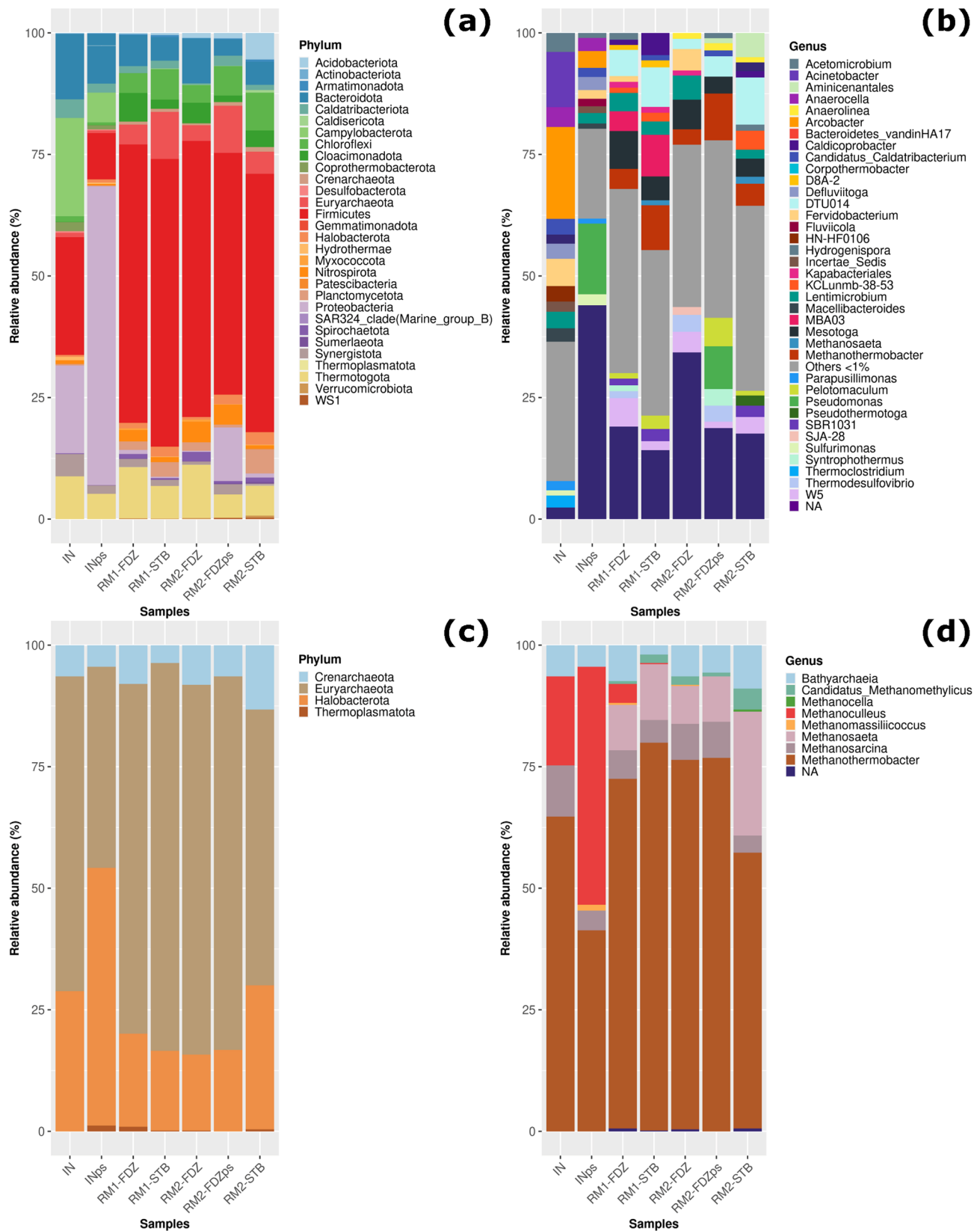


Fig. 7 Taxonomic distribution according to the 16S rRNA gene amplicon sequencing analysis of the total microbial community at phylum and genus level (a, b), and archaeal community at phylum and genus level (c, d). “ps” indicates post-storage

syntrophically oxidize acetate only in very low acetate concentrations (Wang et al. 2019).

Concurrently to SAOB, *Methanosaeta* most likely mediated secondary acetate utilization in methanogenesis via the acetoclastic pathway. Differently from *Methanosarcina*, which is able to mediate acetoclastic methanogenesis under high acetate availability (Petersen and Ahring 1991; Hori et al. 2006), the activity of *Methanosaeta* is favored under low acetate concentrations (Petersen and Ahring 1991; Karakashev et al. 2005; Suárez et al. 2018). Hence, efficient consumption of acetate by SAOB (maintained by the high activity of HM) made only a reduced proportion of acetate available for *Methanosaeta*, characterizing the secondary participation of this group in methane evolution. Interestingly, the highest RA levels observed for *Methanosaeta* were identified in the STB of both reactors, mainly in the case of RM2 (25.5%; Fig. 7d). Because of VFA accumulation events prior to the finalization of the operation, acetate concentrations higher than 1000 mg-COD L⁻¹ were measured in RM2 (Fuess et al. 2021a), suggesting higher participation of *Methanosaeta* in methane evolution. From a metabolic perspective, the SAOB-HM pathway was not able to efficiently metabolize excess acetate, requiring the establishment of parallel acetate-consuming pathways. High RA for the *Methanosarcina* genus was also identified in RM2 (7.4%, FDZ; Fig. 7d), which supports the hypothesis of favoring parallel acetate-consuming pathways. This group may also have participated as a HM, because it includes species capable of growing using any of the known methanogenic pathways, i.e. acetoclastic, hydrogenotrophic, carboxydutrophic (from carbon monoxide), methylotrophic (from methanol, methylamines and methylsulfides) and methyl respiration (methylated compounds + H₂) (Buan et al. 2011).

The microbial characterization also provided bases to understand the behavior of methane evolution in MPP tests. The long lag phase periods observed in reactors inoculated with INps (Fig. 2a–c) most likely resulted from non-identifying genera previously pointed out as key-players in acetogenesis (e.g. *Pelotomaculum*, *Syntrophothermus* and *Cloacimonadaceae* W5) and syntrophic acetate oxidation (e.g. *Mesotoga* and *Pseudothermotoga*) in the continuous reactors. Null RA values were equally identified for the *Methanosaeta* genus (Fig. 7d) in INps, suggesting an even stronger dependence of methanogenesis on the SAOB-HM pathway. This hypothesis is corroborated by the delay periods observed in CODs decay when assessing INps (Fig. 3a–c), because an effective participation of acetoclastic methanogenesis would promptly trigger CODs conversion. Because the *Thermodesulfovibrio* genus was present in INps (at very low initial RA, i.e. 0.3%), this group may have had an important participation in supplying the SAOB-HM pathway on the course of the incubation. Considering RM1- and RM2-derived samples, the minor availability of biomass samples

of RM1-FDZ was impeditive to effectively comparing the sludge samples used in MPP tests (post-storage). However, the comparison between RM1-FDZ and RM2-FDZ (both collected during the disassembling of the continuous reactors) indicated higher RA values for key acetogenic (*Cloacimonadaceae* W5, *Pelotomaculum*, *Syntrophothermus* and *Syntrophomonas*), SAOB (*Mesotoga*, *Thermoacetogenium* and *Pseudothermotoga*) and methanogenic (*Methanothermobacter*) genera in the first sample, explaining the better performances regarding methane evolution when using sludge from RM1. Moreover, the differences in MPP tests most likely did not reach higher levels because the participation of important groups increased in RM2-FDZps (relative to RM2-FDZ), such as the genera *Pelotomaculum*, *Syntrophothermus*, *Methanothermobacter* and *Methanosaeta*.

In any case, once the association between acetogens, SAOB and HM was efficiently established (regardless of the sludge source), both methane evolution and CODs decay proceeded more efficiently than in previous investigations with both fresh and fermented molasses under mesophilic temperature (30 °C) (Fuess et al., 2020). In these cases, the dependence of acetoclastic methanogenesis on the activity of acetogens markedly slowed down methane evolution, which occurred at linear (and not exponential) rates in periods longer than 100 h during the incubation (Fuess et al. 2020). Hence, whenever possible, using thermophilic conditions to intentionally select HM may also characterize an essential approach to increase the efficiency of methane production in two-stage AD, because HM present much lower doubling times (a few hours) compared to acetoclastic groups (a few days) (Mosey 1983). The capability of phase separation to increase the robustness of methane evolution relies more on this aspect than on “simply” eliminating enhanced fermentation from the methanogenic unit. Finally, sugar fermentation was most likely carried out by the genera *Acetomicrobium* (Hania et al. 2016), *Anaerolinea* (Sekiguchi et al. 2003), *Lentimicrobium* (Sun et al. 2016), *Fervidobacterium* (Cai et al. 2007) and *Caldicoproba* (Yokoyama et al. 2010) when using fresh molasses in RM2(Mol) tests, supplying sequential acetogenic and acetate-oxidizing groups.

Conclusions

The primary conclusions drawn from this study include:

- Phase separation under thermophilic conditions is the best approach to achieve efficient methane production from sugar-rich substrates. In addition to the consolidated idea of minimizing stressful conditions to methanogens by separating enhanced substrate fermentation, the use of high temperatures favor the participation of hydrogenotrophic methanogenesis. In practical aspects, methane

evolution will depend on microbial groups (e.g. *Methanothermobacter* and *Methanoculleus* genera) that grow faster and are less susceptible to low pH values compared to acetoclastic ones (e.g. *Methanosaeta* genus). Following the experimental results, the capability of second-stage methanogenic systems to withstand organic loads can be increased by at least threefold compared to single-stage schemes;

- The microbial community distribution tends to be less stratified in the methanogenic unit of the two-stage biogasification system, regardless of the differences in both the substrate availability (once the F/M ratio decreases as the liquid flows through the reactor) and the conditions provided for the cell growth (suspended in the bulk liquid or attached to the fixed media). In particular, homogeneity in the distribution of syntrophic acetate-oxidizing bacteria (e.g. *Mesotoga* and *Thermoacetogenium*) may be *sine qua non* for achieving all benefits previously associated with thermophilic two-stage biogasification (conclusion no. [i]); and,
- Further studies are still required to better understand the role of microbial communities attached to the fixed bed in such methanogenic reactors. In particular, the scale-up of the AnSTBR may be imperative to unravel the distribution of microbial communities in the different compartments of the system. The concentration of biomass in the feeding zone of bench-scale reactors tends to be proportionally high compared to that of the bed region, triggering a sharp substrate decay which limits defining the effective participation of the attached biomass in the overall substrate conversion.

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Declarations

Conflict of interest The authors declare that they have no known competing interests that could have appeared to influence the work reported in this paper.

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