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# ORIGINAL ARTICLE

# Identification of syntrophic acetate-oxidizing bacteria in anaerobic digesters by combined protein-based stable isotope probing and metagenomics

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Inhibition of anaerobic digestion through accumulation of volatile fatty acids occasionally occurs as the result of unbalanced growth between acidogenic bacteria and methanogens. A fast recovery is a prerequisite for establishing an economical production of biogas. However, very little is known about the microorganisms facilitating this recovery. In this study, we investigated the organisms involved by a novel approach of mapping protein-stable isotope probing (protein-SIP) onto a binned metagenome. Under simulation of acetate accumulation conditions, formations of <sup>13</sup>C-labeled CO<sub>2</sub> and CH<sub>4</sub> were detected immediately following incubation with [U-<sup>13</sup>C]acetate, indicating high turnover rate of acetate. The identified <sup>13</sup>C-labeled peptides were mapped onto a binned metagenome for improved identification of the organisms involved. The results revealed that *Methanosarcina* and *Methanoculleus* were actively involved in acetate turnover, as were five subspecies of *Clostridia*. The acetate-consuming organisms affiliating with *Clostridia* all contained the FTFHS gene for formyltetrahydrofolate synthetase, a key enzyme for reductive acetogenesis, indicating that these organisms are possible syntrophic acetate-oxidizing (SAO) bacteria that can facilitate acetate consumption via SAO, coupled with hydrogenotrophic methanogenesis (SAO-HM). This study represents the first study applying protein-SIP for analysis of complex biogas samples, a promising method for identifying key microorganisms utilizing specific pathways.

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#### Introduction

Anaerobic digestion (AD) of organic materials to biogas has several environmental benefits with the methane-rich biogas as a source of renewable energy, as a wastewater treatment technology for the removal of pathogens, as well as odor and pollution reductions from agricultural, industrial and municipal wastes.

The AD of organic matter to biogas involves the coordinated activity of diverse subgroups of highly specialized microbial organisms, hence the stability of AD is highly dependent on the microbial community structure and interactions within the community. Failure to maintain the balance between these groups can lead to reactor inhibitions and breakdown (Demirel and Yenigün, 2002). Accumulation of volatile fatty acids (VFAs) may cause acidification and result in reduced process performance or worst-case scenario: complete reactor failure (Angelidaki and Ahring, 1993; Chen et al., 2008; Krakat et al., 2011). Accumulation of acetate, an important intermediate VFAs in the anaerobic decomposition of organic matter, has been observed under various operational conditions, and the effects on the microbial community have been addressed in several studies (Palatsi et al., 2011; Fotidis et al., 2013; Lü et al., 2013; Rajagopal et al., 2013; Labatut et al., 2014). Although methanogenesis from acetate (aceticlastic methanogenesis) is fairly well described, less is known about the organisms and metabolisms involved in oxidation of acetate to hydrogen and carbon dioxide (syntrophic acetate oxidation (SAO)) catalyzed by SAO bacteria. The formyltetrahydrofolate synthetase-encoding gene, fthfs, is key in the reductive acetogenesis (acetyl-CoA pathway). The enzyme encoded by this gene is also able to catalyze the reverse reaction, oxidizing acetate into H<sub>2</sub> and CO<sub>2</sub> (Xu et al., 2009; Hori et al., 2011). The gene

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has therefore been widely used as a biomarker for acetogenesis, although also present in a variety of non-acetogenic bacteria (Lovell and Leaphart, 2005).

a few SAO bacterial species have been isolated, and only their genomic potential has been determined (Manzoor et al., 2013; Müller et al., 2013). Isolated synthrophic acetate-oxidizing bacteria (SAOB) cover several bacterial phyla and represent both thermophiles (Thermacetogenium phaeum (Hattori et al., 2000); and Thermotoga lettingae (Balk et al., 2002)), thermotolerant (Tepidanaerobacter acetatoxydans (Westerholm et al., 2011)); and mesophiles (Clostridium ultunense (Schnürer et al., 1996); and Synthrophaceticus schinkii (Westerholm et al., 2010)). The oxidation of VFAs by syntrophic bacteria is not a thermodynamically favorable process and requires that they are closely associated with methanogens or with non-methanogenic hydrogenotrophs.

Culture-independent approaches using stable isotopes under high levels of acetate revealed high utilization activity of *Methanosarcina* from the archaeal population and putative SAO bacteria affiliating with the *Clostridia* (Hao *et al.*, 2014). Other studies have identified the *Synergistes* group 4 as a major group of acetate-utilizing bacteria in anaerobic sludge batch reactors fed with 2.5–10 mm acetate (Ito *et al.*, 2011).

Protein-stable isotope probing (protein-SIP) can be used to determine not only the identity, but also the activity of active key microorganisms (Jehmlich *et al.*, 2010). Although protein-SIP has been used on other less complex systems (Jehmlich *et al.*, 2008; Bastida *et al.*, 2010; Taubert *et al.*, 2011, 2012; Herbst *et al.*, 2013), it has, as far as we are aware, not been applied on complex environmental samples such as those present in an AD.

In this study, a combination of metagenomics, amplicon sequencing and protein-SIP was used for characterizing acetate-consuming communities in AD. The proteins from actively incorporating organisms were mapped onto a binned metagenome for identification of the bacteria involved. The results obtained in this study indicate SAO bacterial species are important factors in the recovery after acid accumulation, demonstrating the potential of the combined use of metagenomics and protein-SIP as a tool for linking identity and function, as well as providing an understanding of the underlying biology in AD.

# Materials and methods

Sources of inoculum

Inoculum was obtained from a commercial full-scale biogas digester at Research Centre Foulum, Denmark. The digester works with a mixture of pig and cattle manure, maize silage and deep litter manure. It is operated under thermophilic conditions (approximately 52 °C). The total solid, volatile solid, pH value and total ammonia nitrogen of the inoculum were 50.2, 40.2, 7.64 and 1.54 g l<sup>-1</sup>, respectively. The inoculum was starved under anaerobic conditions at 52 °C for 2 weeks before the main tracer experiment to reduce the background contribution of carbon dioxide and methane from the original substrates. A digestate sample was collected at Lynggård biogas plant (11-01-2012), Denmark. Lynggård is operated at 52 °C and runs with pig manure, and grass and maize silage.

# Operation of AD

Glass serum bottles (500 ml) were used for preparing the anaerobic incubation assay. Aliquots (195 ml) of inoculum were transferred into the 500 ml serum bottles, which were then sealed with butyl rubber stoppers and aluminum crimps. Five treatments were prepared with [U-13C] and [12C] acetate; detailed information of the reactor setup is summarized in Table 1. For the high-acetate concentration condition, substrate was added at the beginning of the experiment, whereas for the low-concentration condition, substrate was added on a daily basis. All experiments were carried out in triplicate, with the exception of the controls (blank reactors), which were carried out in duplicate. The experiment was run under static incubation conditions for 9 days at 52 °C using strict anaerobic techniques.

#### Basic analytical methods

The volume of a produced biogas was measured using an acidified water displacement method at room temperature and atmospheric pressure. Headspace biogas was collected using a gas-tight syringe with a needle through a septum and transferred into a 20 ml headspace vial. The compositions of CH<sub>4</sub> and CO<sub>2</sub> in the biogas samples were analyzed by a gas chromatograph 7890A (Agilent Technologies, Horsholm, Denmark) equipped with

Table 1 Description of batch reactors setup

Substrate	Feeding rate
195 ml Inoculum+5 ml water 195 ml Inoculum+5 ml concentrated CH <sub>3</sub> COONa to a final of 4 mm acetate 195 ml Inoculum+5 ml concentrated CH <sub>3</sub> COONa to a final of 100 mm acetate 195 ml Inoculum+5 ml concentrated [U- <sup>13</sup> C] acetate to a final of 100 mm acetate 195 ml Inoculum+5 ml concentrated [U- <sup>13</sup> C]acetate to a final of 4 mm acetate	Once at the beginning Daily Once at the beginning Daily

a thermal conductivity detector and a GC sampler 80 (Agilent Technologies).

Liquid samples for VFA analysis were collected periodically and the concentration of VFA determined on a 7890A gas chromatograph (Agilent Technologies), equipped with flame ionization detector. For details of the analysis, refer to Supplementary Methods.

# Membrane introduction mass spectrometry measurement

Membrane introduction mass spectrometry was used to monitor the incorporation of <sup>13</sup>C into the produced methane and carbon dioxide during the degradation of <sup>13</sup>C fully labeled acetate as described elsewhere (Mulat *et al.*, 2014). The membrane introduction mass spectrometry data are reported in terms of atom percent as follows:

$$13X(atom\%) = {^{13}X/(^{13}X + ^{12}X)}, \tag{1}$$

where  $^{13}X$  represents  $^{13}CO_2$  or  $^{13}CH_4$ , and  $^{12}X$  represents  $^{12}CO_2$  or  $^{12}CH_4$ .

#### Protein-SIP and amplicon analysis

Liquid samples (5 ml) for protein-SIP and amplicon analysis were collected from all reactors periodically and stored at  $-20\,^{\circ}\text{C}$  until analysis. Sampling was performed at 8, 24, 32, 48, 96, 144 and 192 h following the beginning of the experiment. Protein extraction was performed on samples collected at 8, 24, 48 and 192 h using a protocol as previously described (Hansen *et al.*, 2014). Cell debris was removed by centrifugation at 14 500 xg for 10 min at 4  $^{\circ}\text{C}$ .

Proteins were acetone precipitated as previously described (Botelho *et al.*, 2010). Precipitated proteins were resuspended in sodium dodecyl sulfate—polyacrylamide gel electrophoresis sample buffer, supplemented with dithiothreitol to a final concentration of 40 mm and denatured by boiling at 95 °C for 10 min before being loaded onto a pre-cast 4–15% gradient sodium dodecyl sulfate-gel (Bio-Rad, Sundbyberg, Sweden) and separated for 10 min at 160 V.

Liquid chromatography-tandem mass spectrometry In-gel digestion of proteins was performed as previously described (Shevchenko et al., 2006) and tryptic peptides were analyzed by an automated liquid chromatograph-electrospray ionization tandem mass spectrometer, consisting of an UltiMate 3000 RSLCnano system (Thermo Scientific, Bremen, Germany) coupled to a Q Exactive mass spectrometer (Thermo Scientific) via a Nanospray Flex ion source (Thermo Scientific). The analytical conditions were as previously described (Kjeldal et al., 2014), with the following modifications; analytes were eluted during a 120 min linear gradient, ranging from 12 to 40% (V/V) of solvent B (100% acetonitrile

supplemented with 0.1% (V/V) formic acid) followed by a final step gradient to 90% solvent B which was maintained for 20 min.

#### Protein analysis

A six-frame translation (every region of DNA has six possible reading frames) and prediction of open reading frames in the in-house constructed metagenome of the anaerobic reactor of Foulum was carried out in MaxQuant v. 1.5.1.2 (Martinsried, Germany). A two-search strategy as previously described was utilized (Seifert *et al.*, 2013). Briefly, an initial survey search was performed, searching against the non-redundant NCBI database, restricting the taxonomy to prokaryota. Sequences that were identified in the survey search were exported from NCBI and merged with the six-frame translation of the metagenome, and the merged database was used in the subsequent main search. For details, refer to Supplementary Methods.

Raw files were analyzed using OpenMS (https://www.openms.de) and the TOPP tools (Kohlbacher et al., 2007; Sturm et al., 2008). RIA and protein LR were, as described elsewhere (Kleindienst et al., 2014). determined using the opensource software OpenMS and the MetaproSIP tool (Sachsenberg et al., 2015).

#### DNA extraction

DNA extraction was conducted on the liquid biological triplicates collected at 8, 32 and 192 h of incubation periods, using the FastDNA Spin Kit for Soil (MP Biomedicals, Taastrup, Denmark) with minor modifications (for details, refer to Supplementary Methods).

#### Metagenome preparation

Two metagenomes were prepared from samples collected at the Foulum (13-12-2011) and Lynggård (11-01-2012) biogas plants. DNA was extracted, following a cetyltrimethylammonium bromide and enzyme-based method as described elsewhere (Klocke *et al.*, 2007). DNA was paired-end sequenced (2×150 bp) on a Illumina HiSeq2000 (Illumina, Carlsbad, CA, USA), and metagenome reads following analyzed in CLC Genomics Workbench v. 7.03 (CLC Bio, Aarhus, Denmark). For details, refer to Supplementary Methods.

The analysis and binning were performed exactly as described elsewhere (Albertsen *et al.*, 2013). Briefly, the binning of scaffolds to population genomes was performed by plotting the coverage estimates of the metagenomes of Foulum and Lynggard against each other for all scaffolds. Completeness and contamination of six genome bins was evaluated through calculation of GC content, tetranucleotide frequency and by the identification of conserved essential single-copy marker genes.

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A detailed description of the approach has been published elsewhere (Albertsen *et al.*, 2013).

Core genes of the Wood–Ljungdahl pathway were searched against the metagenome using Hidden Markov model profiles of these genes downloaded from protein family domains (Pfams) homepage (http://pdam.xfam.org/; Finn et al., 2014) and HMMER3 package (http://hmmer.janelia.org/). For details, refer to Supplementary Methods.

Amplicon sequencing

Materials and methods of amplicon sequencing are provided in Supplementary Methods.

Data availability

All amplicon data are available at European Nucleotide Archive (ENA) under accession number PRJEB10871. Metagenomes and assemblies, and bin genomes are available at ENA under accession number PRJEB10932. The mass spectrometry proteomics data have been deposited at ProteomeX-change Consortium (Vizcaíno et al., 2014) via the PRIDE partner repository with the data set identifier PXD002996.

#### Results

Degradation of acetate and methane production Batch reactors inoculated with anaerobic digester sludge were fed two different concentrations of labeled acetate, and the concentration of acetate as a function of time was monitored (Figure 1). The degradation of acetate in the blank reactor (inoculum only) and in the reactors fed with low (4 mm) and high (100 mm) concentrations of [U-¹³C]acetate were significantly different; the background concentration of acetate in the blank reactor remained at ~ 2 mm for the first 48 h and declined to below the detection limit between 48 and 72 h. No acetate accumulation was observed in reactors fed with low acetate concentration, and the total turnover of acetate was

approximately  $2.95\,\mathrm{g}\,\mathrm{l}^{-1}$  during the 9 days of incubation.

A continuous turnover of acetate occurred in the high acetate concentration-fed reactor, and the total turnover of acetate was around  $8.2\,\mathrm{g}\,\mathrm{l}^{-1}$  (acetate was spiked in at the beginning of the experiment only). The degradation of acetate in the high concentration-fed reactor started immediately and followed a linear trend during the first 120 h (corresponding to a consumption of ~83% of the amended [U- $^{13}$ C] acetate). After 120 h, the turnover of [U- $^{13}$ C] acetate decreased and finally reached a stationary level during the last 68 h of the incubation. Following 9 days of incubation, the concentration of acetate reached ~1.6 mm.

Besides the degradation of acetate, the production of labeled and unlabeled methane and carbon dioxide, respectively, was also monitored (Figure 2). In the low acetate concentration-fed reactor, the proportion of the  $^{13}\mathrm{CO}_2$  produced to total carbon dioxide increased almost linearly from 6 atom% up to 32 atom% during the 9 days of incubation, whereas the proportion of the  $^{13}\mathrm{CH}_4$  produced to total methane increased gradually from 45 atom% up to 76 atom%.

In the high acetate concentration-fed reactor, the production of <sup>13</sup>CO<sub>2</sub> increased from 20 atom% to about 45 atom% at 96 h and later stabilized at this value. The production of <sup>13</sup>CH<sub>4</sub> reached the peak value (80 atom%) at 24 h and remained almost stable for 96 h before starting to decline. During the time at which the levels of <sup>13</sup>CH<sub>4</sub> appeared to remain constant (Figure 2), almost 83% of the acetate was degraded (Figure 1). The isotopic abundance of <sup>13</sup>C-labeled CO<sub>2</sub> and CH<sub>4</sub> at both low and high concentrations of <sup>13</sup>C-labeled acetate incubations indicate the uptake and turnover of the added <sup>13</sup>C-labeled acetate.

Phylogenetic microbial community composition of acetate-fed reactors

The microbial community compositions were evaluated by amplicon sequencing of triplicate reactors

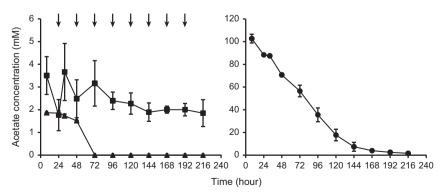
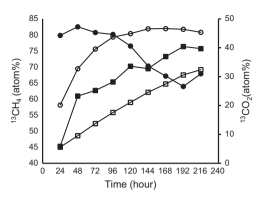


Figure 1 Temporal change of the residual acetate in (a) the blank reactors ( $\triangle$ ); reactors fed with low concentration (4 mm) of [U- $^{13}$ C] acetate ( $\blacksquare$ ); and (b) reactors fed with high concentrations (100 mm) of [U- $^{13}$ C] acetate ( $\blacksquare$ ). The lines represent mean values (n = 3), and error bars denote the standard deviation. Arrows indicate addition of acetate.



**Figure 2** Temporal change of the atom percent of  $^{13}\text{CO}_2$  and  $^{13}\text{CH}_4$  in the reactors fed with low concentrations of [U- $^{13}$ C]-acetate (■  $^{13}\text{CH}_4$ ;  $\square$   $^{13}\text{CO}_2$ ); and in the reactors fed with high concentrations of [U- $^{13}$ C]acetate (●  $^{13}\text{CH}_4$ ;  $\bigcirc$   $^{13}\text{CO}_2$ ). The lines represent mean values (n=3), and error bars denote the standard deviation.

fed with low and high concentrations of acetate, respectively, see Supplementary Information. Samples at three time points were chosen for amplicon sequencing (8, 32 and 192 h). At least 29 428 sequences per sample passed filtering. From the archaeal population, the most abundant genera were affiliated with Methanobacterium, Methanosarcina, Methanobrevibacter and Methanoculleus. However, Methanomassiliicoccus and Methanothermobacter were also detected. The five most abundant classes of bacteria belonged to Clostridia, Bacteroidia, Bacilli, Thermotogae and Anaerolineae (Supplementary Figure 1).

Among abundant genera (>0.1% of the total reads), several genera from the bacterial population were seen to either decrease or increase from 8 to 192 h (Supplementary Figure 1A). The most noticeable change in the microbial community seems to occur among the two orders of *Firmicutes*, *Bacillales* (increasing), and *Clostridiales* (decreasing). The differences in the microbial communities in the high and low acetate time series, as well as the three replicates are illustrated by non-metric multidimensional scaling analysis (Supplementary Figure 1B). The findings revealed high reproducibility of the biological replicates and changes between the time series of each experiment.

#### Protein-SIP analysis

Time-resolved protein-SIP analysis showed that <sup>13</sup>C was incorporated into peptides for the reactor fed with 100 mm [U-<sup>13</sup>C] acetate, starting from 48 h (Table 2, Figure 3). No <sup>13</sup>C labeling was detected in the reactor fed with 4 mm [U-<sup>13</sup>C]acetate, control (fed with unlabeled acetate) and blank reactors at any time point (data not shown).

In the reactor fed with high concentration of acetate, a total of five peptides incorporated  $^{13}\mathrm{C}$  at 48 h (Table 2) after the start of the incubation. From these  $^{13}\mathrm{C}$ -labeled peptides, one of them were

assigned to the domain of bacteria and the other two to the domain of archaea. The last two peptides could not be assigned to any sequence present in either of the two metagenomes. A functional annotation was given to three of the peptides. The two peptides belonging to *Archaea* were identified as different subunits of the methyl coenzyme M reductase, a key enzyme involved in methane formation from methanogens (Ermler, 1997). The third peptide, which was unassigned, was identified as flagellin subunit B and is thus related to motility.

Following 192 h, a total of 56 peptides could be identified that showed incorporation of <sup>13</sup>C (Figure 3 and Table 2). These peptides were dominated by bacterial species (*Clostridia*). Three labeled peptides originated from the domain of *Archaea* (two from the genus of *Methanoculleus* and one peptide from *Methanosarcina barkeri*). The proportion of the <sup>13</sup>C-labeled peptides, known as relative isotope abundance (RIA), as well as how much of a peptide population is labeled, LR, is noted in Table 2. An example of the development of labeling from 24 to 192 h is given in Figure 3b, for a methyl coenzyme M reductase (chain B) from *Methanosarcina barkeri*.

A functional annotation could be assigned for 6 out of the 56 peptides found at 192 h (Figure 3a, Table 2). The remaining five peptides came from proteins from *Clostridia* (four peptides) and an unassigned peptide and had household functions, for example, initiation of RNA synthesis, signal transduction and transport substrate binding. However, one peptide belonged to putative TetR family, which is associated with antibiotic resistance.

The two metagenomes from Foulum and Lynggård biogas plants were sequenced to yield a total of 130 190 434 and 79 999 986 reads, respectively. When assembled, this amounted to a size of 165 123 257 bp for the Foulum metagenome, whereas the Lynggård metagenome was 82 793 162 bp. Both of the metagenomes represent approximately 98% Bacteria and 2% Archaea (Supplementary Figure 2). The two metagenomes were binned, based on the coverage and scaffold length, and the phylogeny from BLAST of each scaffold was superimposed (Figure 4a). The two metagenomes are relatively similar and were used as a reference database for the identified and <sup>13</sup>C-labeled peptides (Figure 4b). The phylogenetic identification in the two metagenomes resembles the phylogenetic identification found by amplicon sequencing (Supplementary Figure 2). Protein-SIP data were superimposed onto the metagenome and all scaffolds belonging to a genome bin for which labeling of proteins were observed were extracted. The scaffolds of a total of six genome bins were extracted and completeness and potential contamination of each genome bin was evaluated (Table 3). The details of the assembly can be found in the Supplementary File 1.

Table 2 Peptides for which an increase in isotopic incorporation of <sup>13</sup>C was observed in the reactor fed with 100 mM [U-<sup>13</sup>C]acetate

1	,			1				
$Accession^{\mathrm{a}}$	$Peptide\ sequence^{ m b}$	$Description^\circ$	$Exp. m/z^{\rm d}$ 7	Theo. $m/z^{\circ} Z^{f}$ I	RIA 18 RIA	$2^{\mathrm{h}}$	LR <sup>i</sup> Taxonomy <sup>i</sup>	
48 h GII 219919005   3291:TRUE:2282 GII 11514434   2256:TRUE:10376 40545:TRUE:0	GPNEPGGLSFGHLSDIIQTSR LIGHGPFILDQYK SVAVNLAGIQGALASGK AGDDAAGLSISEK LTPEEFVSTFIPADLTWM (Oxidation)R	Methyl coenzyme M reductase subunit alpha Chain B, Methyl-Coenzyme M Reductase Flagellin, subunit protein B	728.0337 500.9459 778.4442 617.3017 757.0406	728.0344 3 500.9452 3 778.4438 2 617.3015 2 757.0452 3	1.2 57 0.6 28 0.6 57 0.6 90 1.1 1	51.1 25.1 57.6 90.3 9	7.4 Archaea 0.8 Clostridia (C2) (genus Gelria) 5.8 Methanosarcina barkeri 91.3 Unassigned 60.6 Unassigned	genus <i>Gelria)</i> barkeri
192 h 16:FALSE:5223 1400:FALSE:7184 18:FALSE:4093 2364:FALSE:3984 34:TRUE:27427 8772:TRUE:988 515:TRUE:2	AILPSPYGAFTR VDELLELGR FQVGFEEGVK DNPESIFVPLPVDPLVEER GLVGELFQEDLAALGIK ELLAEAGIEFGEISIR FVTVFEGIPVAK	ABC transporter substrate-binding protein	646.8548 522.2914 570.2911 797.7637 922.5129 848.9621	646.8535 2 522.2902 2 570.2902 2 592.5118 2 848.9594 2 692.3901 2	0.6 0.7 0.6 33 11.2 22 11.2 22 22 22 23 23 23 24 25 25 26 27 27 27 28 28 28 28 28 28 28 28 28 28 28 28 28	32.2 1 31.7 2 30.4 1 28.4 1 27.8 27.5	10.7 Clostridia (C1) ( 26.1 Clostridia (C1) ( 9.9 Clostridia (C1) ( 12.2 Clostridia (C1) ( 9.3 Clostridia (C1) ( 6.2 Clostridia (C1) ( 13.9 Clostridia (C1) (	genus Gelria) (genus Gelria) (genus Gelria) (genus Gelria) (genus Gelria) (genus Gelria)
888:FALSE:1680 448:TRUE:12186 515:TRUE:2 448:TRUE:12186 34:TRUE:12186	AAFLGVILR AAFLGVILR RGEEIGGTIR AIGINANPAFPDAGVYNDR IDEVIMI A AOP	Signal transduction histidine kinase-like protein	465.2752 771.4387 544.2969 987.9882	465.2744 2 771.4379 2 544.2964 2 987.9871 2			Clostridia (C1) Clostridia (C1) Clostridia (C1) Clostridia (C1) Clostridia (C1)	$genus \ Gelria$
34:1 RUE:2.44.7 515:TRUE:2 447:FALSE:10792 515:TRUE:2 73:FALSE:5845 34:TRUE:27427 9364:FALSE:5845	LIEVWLAAQK REEIGGTIR SGAQVLLSR AGELAFSASK DPPLYGAGDRTEDNLIK GOGLILVK CVMDFTFIN NYDOVIEK		544.2968 465.7724 490.7562 641.9862 549.8302	544.2964 2 465.772 2 490.756 2 641.9864 3 549.8295 2			Clostridia (C.1)	
2304:I7ALSE:3904 2074:TRUE:5302 54:FALSE:1185 73:TRUE:3399 109:TRUE:9954 515:TRUE:27427 109:TRUE:27427	GVIDED ITELATED TEACH TO THE A		10/3.043 648.6618 557.3177 868.4566 438.7331 593.3216 874.4877 807.6958	648.6618 3 557.323 4 438.7323 4 593.321 3 874.4851 2 807.6963 3 554.6407 3			Clostridia (C.1)	genus <i>Gelria</i> ) (genus <i>Gelria</i> )
139;FALSE:61670 49:TRUE:24338 515;TRUE:2 1400;FALSE:7184 515;TRUE:27427 73;FALSE:5845 448;TRUE:12186 448;TRUE:12186 2250;TRUE:7164 3291;TRUE:2282	1 LUEFYQLAR TLUEFYQLAGETPYNELQNEGR DLFVQAGLPTPNELQNEGR AVTGPLPPLVWASR GFVSNPYTGNYM(Oxidation)PHR EYEISEDGTEVTFYLR YNVEVEFKPVPR AIGINANPAFPDAGVYNDR DDNWWGNAVFGQPKPK FVAIEHVSADAAR QAADEAQLILAR		606.3202 1049.533 585.9351 732.4232 585.9357 975.9533 642.9338 658.9934 620.3004 462.5779 649.8583	606.319 2 1049.532 2 2585.9351 3 732.4221 2 585.9351 3 975.952 2 492.933 3 658.9938 3 658.9938 3 6620.3006 3 642.5772 3		222.3 3 221.7 2 221.7 2 221.7 2 200.7 200.7 115 115 113.6 11 3.6 3.8 3.8 3.8 3.8	38.3 Clostnida (C1) (22.9 Clostnida (C1) (21.6 Clostnida (C1) (13.4 Clostnida (C1) (14.7 Clostnida (C1) (12.9 Clostnida (C1) (11.4 Clostnida (C1) (11.4 Clostnida (C1) (35.0 Clostnida (C1) (39.0 Clostnida (C2) (39.0 Clostnida (C2) (39.0 Clostnida (C2) (39.0 Clostnidia (C2) (39.0 Clostni	genus <i>Gelria</i> ) (genus <i>Gelria</i> )
1633:FALSE:12343 3291:TRUE:2282 862:TRUE:2131 3291:TRUE:282 709:TRUE:15243 3291:TRUE:2282	LİJALQTSDK LIDEAGYTVDPATGIR IANVFATGGLGDK REPLADDVIR HFTNSIRPIR LIGHGPFILDQYK	Putative TetR family transcriptional regulator	551.3309 845.9382 631.3625 592.3258 414.2352 500.9458	551.3293 2 845.936 2 631.3612 2 592.3251 2 414.2352 3 500.9452 3	0.7 3.7 0.6 3.6 0.6 2.6 0.6 0.6 0.6 0.6 0.6 0.6 0.6 0.6 0.6 0	34.3 2 32.2 30.5 30.1 28.9 1 26.2 24.6		(genus Gelria)

Table 2 (Continued)

$Accession^{a}$	Peptide sequence <sup>b</sup>	$Description^{\circ}$	$Exp. m/z^d$	Theo. $m/z^e$ $Z^f$	RIA 18	$RIA~2^{ m h}$	$Exp.\ m/z^{ m d}$ Theo. $m/z^{ m e}\ Z^{ m f}\ RIA\ 1^{ m s}\ RIA\ 2^{ m h}\ LR^{ m i}\ Taxonomy^{ m i}$
709:TRUE:15243 1006:TRUE:9099	NLTFAEVGFR VVEAAIAAGK	Simple sugar transport system substrate-binding protein	577.3048 464.7768	577.3037 2 0.7 464.7767 2 0.7	0.7	17.3 30.9	8.8 Clostridia (C2) (genus Gelria) 12.6 Clostridia (C3)
1279:TRUE:16513	AIISVDANTQSHGVVIR	•	593.996	593.9952 3	1.3	27.2	17.6 Clostridia (C3)
1279:TRUE:16513	VAIGVEDLGHTSLAER		556.2979	556.2968 3	1.2	26.8	15.0 Clostridia (C3)
2799:FALSE:25402	IFTVDQISFIPK		704.3993	704.3978 2	9.0	21.4	9.5 Clostridia (C4)
2799:FALSE:25402	ALELSLEDSPR		615.323	615.3222 2	9.0	20.7	5.0 Clostridia (C4)
527:FALSE:43604	VLELALKDSVR		414.9192	414.9187 3	9.0	23.9	17.7 Clostridia (C5)
2628:TRUE:10590	ELDLDIVGNKDAVISK		576.9853	576.9841 3	1.2	51.7	2.8 Methanoculleus
2628:TRUE:10590	TIAVNLGGIEGALK		678.3993	678.3983 2	9.0	49.6	2.3 Methanoculleus
GI 11514434	SVAVNLAGIQGALASGK	Chain B, Methyl-Coenzyme M Reductase	778.445	778.4438 2	9.0	58.1	39.2 Methanosarcina barkeri
33816:FALSE:2321	ILDLLDSAPDLATAK		778.4337	778.4325 2	1.2	31.9	14.9 Unassigned
GI   652561026	EIALSLDLSPR	RNA polymerase sigma-70 factor, expansion family 1 607.3438	607.3438	607.343 2	0.7	24.4	15.4 Unassigned

Peptide sequence (with identified modification). Description of the identified protein. Experimentally determined mass-to-charge ratio of the observed sequence. "Theoretically determined mass-to-charge ratio of the observed sequence. Charge of the tryptic peptide. Relative isotope abundance of the naturally occurring isotopic cluster of the peptide. 1-Relative isotope abundance of isotopic cluster of the peptide showing incorporation of 1-8C. Labeling ratio. Highest possible taxonomic for the corresponding protein. <sup>a</sup>Protein identifier from metagenome or gene identifier (GI) classification

The Wood–Ljungdahl pathway has four marker genes, acsB (carbon monooxide dehydrogenase), the two subunits of the corrinoid iron sulfur protein (acsC and acsD), and ftfhs (formyltetrahydrofolate synthatase) (Can et al., 2014). Searching Hidden Markov model profiles of each of the functional domains of these marker proteins against the sixframe translated metagenome, revealed that all of the six Clostridia bin genomes contained the ftfhs gene, whereas genes acsC and acsD were only found in one of the bin genomes (Clostridia (C4). All of the six genome bins tested positive for all the genes of the Wood–Ljungdahl pathway (Supplementary File 2).

Eleven proteins from a syntrophic glutamate-oxidizing bacterium of the genus *Gelria* (C2) were labeled with <sup>13</sup>C during the SIP experiment (Table 2). The genus *Gelria* (phylum *Clostridia*) (C1) was the most heavily labeled in the proteome with 35 <sup>13</sup>C-labeled peptides.

Gelria belongs to the family Thermoanaerobacteraceae and is closely related to the genus of two of the known SAOB, namely Thermacetogenium phaeum and Tepidanaerobacter acetatoxydans, which are also part of the Thermoanaerobacteraceae family. Thermacetogenium had a relative read abundance of 0.01–0.11% in the amplicon data. Overall, the Thermoanaerobacteraceae family was found with an abundance of 0.8–2.0% in the different batch samples. Several ftfhs genes from Clostridia were found in the six-frame translated metagenome of the Foulum plant (Supplementary File 2).

Thermotoga lettingae is another known SAOB in the family Thermotogaceae. We did not detect any labeling in the proteins of this organism. A closely related genus, S1, was highly abundant in the amplicon data with a relative abundance of 2.1–7.2%. FTFHS was observed in Thermotogae in the six-frame translated metagenome (Supplementary Figure 1).

Six proteins from *Clostridia* were <sup>13</sup>C labeled, but not found in any of the bins and could not be identified at a higher taxonomic level. Amplicon sequencing revealed that the genus *Clostridium* had a relative abundance of 0.6–1.4%. However, the known SAOB, *Clostidium ultunese*, was not detected with <sup>13</sup>C incorporation, nor in the amplicon and metagenome data. Furthermore, *Syntrophaceticus schinkii*, from the family *Thermoanaerobacterales* Family III, order *Thermoanaerobacterales*, which is the last of the known SAOB, was not identified from the SIP experiment, nor was this family detected with amplicon sequencing.

#### **Discussion**

Our aim was to apply protein-SIP to describe the complex system of AD. Peptides showing incorporation of <sup>13</sup>C were mapped onto a binned metagenome in order to identify key microorganisms involved in

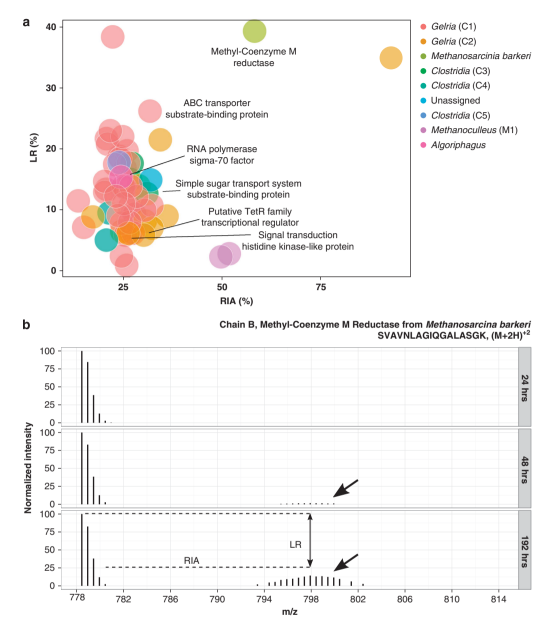


Figure 3 (a) RIA as a function of LR for peptides, showing incorporation of <sup>13</sup>C at 192 h in the reactor fed with 100 mm of [U-<sup>13</sup>C] acetate (colors represent highest possible taxonomic rank with names in brackets corresponding to extracted genome bins). Functionally annotated proteins are indicated in the figure. The peptide SVAVNLAGIQGALASKGK, which showed a high degree of <sup>13</sup>C incorporation, was identified belonging to *Methanosarcina barkeri*. (b) Time resolved analysis of the incorporation of <sup>13</sup>C in the peptide SVAVNLAGIQGALASKGK belonging to the methanogenic archaea *Methanosarcina barkeri*. Incorporation of <sup>13</sup>C in the peptide was evident after 48 h (the arrows indicate the <sup>13</sup>C-labeled isotopologue). The peptide was identified as a methyl coenzyme M reductase, which catalyzes anaerobic oxidation of methane. Besides the methyl coenzyme M reductase, proteins with functions related to transport of substrate and sugars, signal transduction, translation as well as a transcriptional repressor were also identified (Figure 3 and Table 2).

recovery of acetate accumulation under conditions simulating normal running conditions (low acetate levels) and stressed conditions with high levels of acetate.

The  $^{13}$ C isotopic profiling during the start of incubation in both the low and high concentrations of [U- $^{13}$ C]acetate showed that CO<sub>2</sub> and CH<sub>4</sub> are the main products of acetate degradation. As both the

methyl and carboxyl groups are  $^{13}$ C labeled in the [U- $^{13}$ C]acetate, it is difficult to estimate the proportion of methane produced through SAO-HM, using the values of atom percent of  $^{13}$ CO<sub>2</sub> and  $^{13}$ CH<sub>4</sub>. The  $^{13}$ CO<sub>2</sub> formed can be produced either through acetoclastic methanogenesis or through SAO during the degradation of [U- $^{13}$ C]acetate. In a previous study, performed on sludge from the same plant as

that analyzed in this study, methyl  $^{13}$ C-labeled acetate ([2- $^{13}$ C]acetate) was used to trace incorporation of  $^{13}$ C into CO<sub>2</sub> and CH<sub>4</sub> in real time during the

degradation of [2- $^{13}$ C]acetate. The results showed that SAO-HM had a key role in the conversion of acetate to methane as  $^{13}$ CO $_2$  can be produced from

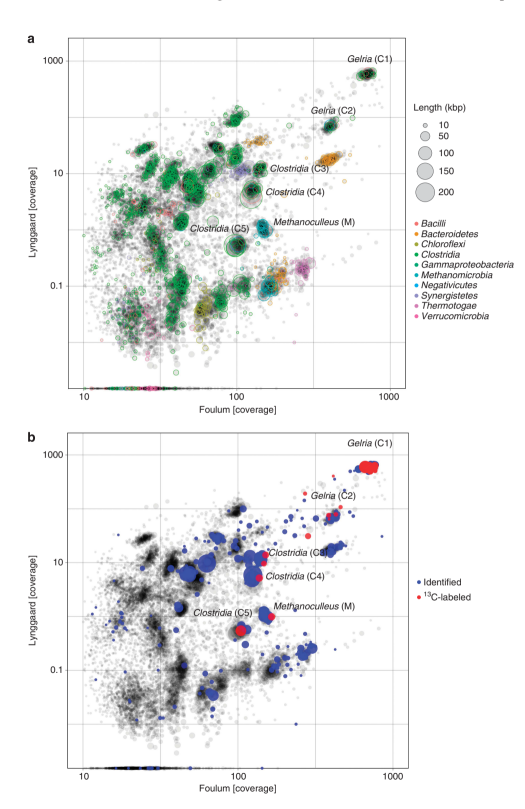


Figure 4 The scaffold coverage from two metagenomes are plotted. The dot sizes indicate the scaffold length. (a) Coloring is according to phylogeny. (b) Coloring is observed in the protein-SIP analysis according to identified and <sup>13</sup>C-labeled peptides. Labeling was seen in six clusters of scaffolds belonging to five subspecies of *Clostridia* and *Methanoculleus*.

Table 3 Assembly statistics of the six genome bins containing scaffolds to which proteins showing incorporation of <sup>13</sup>C mapped to in the protein-SIP analysis

Phylogenetic affiliation	No. of contigs	Total length (bp)	GC (%)	N50	Foulum coverage	Lynggaard coverage	No. essential unique genes	No. of total essential genes	Figure 4a
Clostridia (Gelria)	128	2 092 416	59.40	25 636	731.79	593.5	104	107	Gelria (C1)
Clostridia (Gelria)	306	1 833 135	49.50	11 477	405.00	71.02	94	95	Gelria (C2)
Clostridia	145	1 786 211	56.90	19 622	140.58	12.09	100	103	Clostridia (C3)
Clostridia	71	2 167 847	53.80	71 792	125.47	4.93	102	104	Clostridia (C4)
Clostridia	59	$2\ 188\ 512$	53.60	61 047	102.11	0.54	100	103	Clostridia (C5)
Methanomicrobia (Methanoculleus)	161	2 200 216	59.40	24 804	148.67	1.09	33	35	Methanoculleus (M)

The number of essential genes was evaluated using 107 Hidden Markov models of protein coding-essential single-copy genes.

the degradation of [2-<sup>13</sup>C]acetate through SAO-HM alone (Mulat *et al.*, 2014). Similar observations have been reported after amendment of [2-<sup>14</sup>C]acetate in batch incubations inoculated with manure and food waste (Karakashev *et al.*, 2006).

A qualitative interpretation of the measured atom percent of  $^{13}\text{CH}_4$  and  $\text{CO}_2$  during the degradation of  $[\text{U-}^{13}\text{C}]$  acetate revealed that the atom% of  $^{13}\text{CO}_2$  was generally lower than that of the  $^{13}\text{CH}_4$  in both the low and high concentrations of  $[\text{U-}^{13}\text{C}]$  acetate. This is most likely due to the high background pool of unlabeled  $\text{CO}_2$  in the system.

This is corroborated by the levels of <sup>13</sup>CH<sub>4</sub>, which reached a maximum of 75 atom% and 80 atom% in the low- and high-acetate-fed reactors, respectively. The production of unlabeled CH<sub>4</sub> is possibly caused by a reduction of the background pool of unlabeled CO<sub>2</sub> by hydrogenotrophs in syntrophy with SAOB. This assumption is in line with previous findings from the same AD, where SAO-HM was found to have a key role for the production of methane during the degradation of high concentrations (100 mm) of [2-13C]acetate (Mulat et al., 2014). Experimental conditions in this study and the previous study (Mulat et al., 2014) were very similar with the exception of the inoculum, which were a few months apart from the same full-scale biogas plant (Foulum). Although fluctuations in community dynamics are expected over time, the community structure between the time of the present and previous studies remained relatively stable (Mulat et al., 2014). The dominant microbial communities are almost similar in both studies (see the discussion below), indicating that SAO-HM had a significant role in the reactor fed with the high concentration of [U-13C]acetate.

The effect of the experimental design on the microbial community composition was investigated by amplicon sequencing of the 16S ribosomal RNA gene in both low- and high-acetate-fed reactors (see Supplementary Information). In general, the microbial communities observed in both batch reactors are in accordance with previous studies of AD communities (Lee et al., 2012; Wirth et al., 2012; Sundberg et al., 2013; St-Pierre and Wright, 2014). The majority of the organisms identified were

affiliated to Clostridia. Clostridia participate in various stages in AD, including hydrolysis of cellulosic plant biomass and acetate oxidation (Hattori, 2008; Schlüter et al., 2008; Wirth et al., 2012). The bacterial classes of *Bacteroidia*, *Bacilli*, Thermotogae, Anaerolineae, Synergistia, as well as several Proteobacteria were also highly abundant. This composition is also in accordance with our metagenome constructed from the Foulum and Lynggaard biogas plants. Methanobacterium, Methanosarcina, Methanobrevibacter and Methanoculleus, representing hydrogenotrophic and acetoclastic methanogens, were present in the four most abundant genera of Archaea detected in our reactors, which is in accordance with previous studies from similarly operated AD (Blume et al., 2010; Sasaki et al., 2011; Hagen et al., 2014; Tuan et al., 2014; Ziganshina et al., 2014).

The abundance of *Archaea* (1–2%) was relatively low, compared with Bacteria (98-99%). Several previous studies of the microbial diversity in AD are based on separate analysis for Archaea and Bacteria, and the abundance of the two is thus not directly comparable (Patil et al., 2010; Kim et al., 2011; Lee et al., 2012; Rodríguez et al., 2012). Although we applied a universal primer set targeting both kingdoms, the abundance count of Archaea can be biased because of extraction and PCR. The approach used in this study does not take account of the 16S ribosomal RNA gene copy number, which has previously been shown to be generally lower in Archaea than Bacteria (Lee et al., 2009). However, the abundance ratio of Archaea and Bacteria is in accordance with the metagenome.

Time, rather than the two different concentrations of acetate tested, was the cause of the observed fluctuations in the community structures. A significant increase in *Bacillales*, specifically the genera *Ureibacillus*, was observed. Although the abundance of the genera of *Ureibacillus* can potentially be attributed to the presence of acetate, no incorporation of <sup>13</sup>C from acetate was observed at protein level.

The abundances of *Methanobrevibacter* and *Methanoculleus* were constant during the incubation, the abundance of *Methanobacterium* decreased,

and the abundance of *Methanosarcina* increased. The small increase of *Methanosarcina* is interesting because this organism is capable of both acetoclastic (Liu *et al.*, 1985) and hydrogenotrophic (Thauer *et al.*, 2008) pathways. Members of *Methanosarcina* are favored at acetate concentrations higher than 1 mm (Karakashev *et al.*, 2006; Hori *et al.*, 2011), although the biochemical reasoning remains unknown.

The microorganisms directly involved in acetate turnover were investigated with protein-SIP, not only identifying the microorganisms but also determining their level of activity. <sup>13</sup>C-labeled peptides were detected only in the reactors fed with high concentrations of acetate; a reason for this could be a higher turnover rate of acetate of fewer specialized organisms and/or the relative high levels of labeling required for isotopical profiling. The focus hereafter will be on samples from the 100 mm reactor. <sup>13</sup>C isotopically labeled peptides of the microorganisms were observed after 48 h of incubation (five peptides). The number of identified labeled peptides increased over time (56 peptides after 192 h), indicating that an adaptation period, for example, protein synthesis, for the microorganisms toward a change in environmental conditions was required.

The number of <sup>13</sup>C-labeled peptides identified increased by using the Foulum metagenome as a reference database. Still, the number of <sup>13</sup>C-labeled peptides detected in our reactors is low and can be explained by the very dense nature of the samples and the fact that protein extractions are often contaminated with humic substances, which, in correlation with a bias of mass spectrometry toward the most abundant proteins, challenges subsequent isotopical profiling (Doherty and Beynon, 2006; Heyer et al., 2013). Furthermore, the microbial communities in the AD are very complex, hence difficult to analyze using mass spectrometry. However, we detected between 1000 and 2100 unlabeled peptides in our reactors, which is a much greater number than those labeled with <sup>13</sup>C, the incubation for a total of 9 days may have been too short for the slow-growing methanogens to adjust and synthesize new proteins, causing insufficient labeling.

Peptides from five subspecies of Clostridia (C1–C5), as well as Methanosarcina and Methan oculleus (M), incorporated <sup>13</sup>C from the reactors fed with high levels of 13C-labeled acetate. Methyl coenzyme M reductase from Methanosarcina barkeri was <sup>13</sup>C labeled, which is involved in methanogenesis, both acetoclastic and hydrogenotrophic. However, many of the peptides came from household proteins and were not directly involved in the SAO pathway. Formyltetrahydrofolate synthetase catalyzes the formation of acetate from  $H_2$  and  $CO_2$  (Xu et al., 2009; Hori et al., 2011), but has previously been associated with SAO in AD (Hori et al., 2011). The gene encoding the enzyme formyltetrahydrofolate synthetase, fthfs, is an ecological biomarker for reductive acetogenesis (Xu et al., 2009; Hori et al., 2011). Thus, we searched for *fthfs* in the six-frame translated Foulum metagenome, which was present in all of the *Bacteria* detected with the <sup>13</sup>C-labeling approach. However, only two of the remaining marker genes, *acsC* and *acsD*, encoding CO dehydrogenase/acetyl-CoA synthase subunits.

Despite the absence of these genes, protein-SIP data suggest that these phylotypes are responsible for SAO. Most of the <sup>13</sup>C-labeled proteins affiliated with the genus *Gelria* (c1). The enzymes of the Wood–Ljungdahl pathway are generally poorly understood for SAOB, and it has previously been suggested that other metabolic strategies than the Wood–Ljungdahl are being used by SAOB (Müller *et al.*, 2013).

Most of the <sup>13</sup>C-labeled peptides had a RIA (proportion of the peptide <sup>13</sup>C labeled) of around 13–36%, however, four peptides had higher RIA values of 49.6% and 51.7% (*Methanoculleus*), 58.1% (*Methanosarcina*), and 93.8% (genus *Gelria* in the class *Clostridia*), respectively. The LR (how much of a peptide population is labeled) of these four peptides were 2.3% and 2.8% (*Methanoculleus*), and 39.2% (*Methanosarcina*) and 35.0% (*Gelria*). These data clearly confirm that these microorganisms are part of acetate degradation.

We detected unlabeled peptides from several methanogens, both aceticlastic and hydrogenotrophic. Among the methanogens, Methanosarcina had the highest <sup>13</sup>C labeling with a RIA of 58.1% and an LR of 39.2%. The labeled peptide came from the methyl coenzyme M reductase  $\beta$  subunit, which is a common intermediate reaction of all metabolic pathways leading to methane formation (Grabarse et al., 2001). Thus, methyl coenzyme M reductase takes part in both methanogenic pathways (Rademacher et al., 2012). The highest <sup>13</sup>C labeling of Methanosarcina is in accordance with the measured acetate consumption rate and the observed increase in the relative abundance of Methanosarcina during the course of incubation, as shown by the amplicon sequencing analysis. This methanogen participates in acid recovery in ADs by transformation of acetate in accordance with previous studies (Petersen, 1991). Methanosarcina was also highly abundant in the reactor fed with 100 mm [2-13C] acetate presented in our previous study, and the results of the isotope analysis demonstrated the key role of SAO-HM in the degradation of acetate (Mulat

Adjusting the parameters of the AD to favor the presence and activity, *Methanosarcina* can possibly improve the stability of the reactor as this organism is able to carry out both acetoclastic and hydrogenotrophic methanogenesis and is more tolerant of several inhibitors such as ammonium, low retention times and high organic loading rates (De Vrieze *et al.*, 2012). *Methanosaeta*, which are strictly acetoclastic methanogens, were not detected with amplicon sequencing. This could point in the direction of SAO-HM being the dominant process. This pathway

et al., 2014).

is similarly dominant at thermophilic temperatures and in plants with increased levels of acetate (De Vrieze *et al.*, 2012). However, as *Methanosarcina* is capable of both methanogenesis pathways, it is not possible to define the dominant pathway in the digester under high acetate levels.

Methanoculleus was the other methanogen found with labeled peptides. This particular methanogen is hydrogenotrophic and grows on H<sub>2</sub> and CO<sub>2</sub> (Mikucki et al., 2003). Applying [1-<sup>13</sup>C]acetate and [2-<sup>13</sup>C]acetate would allow to differentiate between acetoclastic or hydrogenotrophic methanogenesis. Furthermore, the use of primers targeting the fthfs gene would allow us to identify the specific SAOB present in our reactors. As Methanoculleus incorporated <sup>13</sup>C into their peptides while growing on 100 mm [2-<sup>13</sup>C]acetate, they likely use the SAO-HM pathway in syntrophy with SAOB.

# Conclusion

In this study, we showed that protein-SIP is a method that can be used to detect the active microorganisms incorporating <sup>13</sup>C into their proteins in complex samples from AD batch reactors. We conclude that the mapping of protein-SIP onto a binned metagenome is highly applicable for identifying members of functional groups in complex microbial ecosystems. Peptides from Clostridia, Methanosarcina and Methanoculleus were labeled with <sup>13</sup>C and therefore confirmed that these microorganisms were involved in the recovery after inhibitory events with high levels of acetate. The <sup>13</sup>C-labeled and identified *Clostridia* are most likely oxidizing acetate as part of a synthrophy as they all contain the fthfs gene coding for formyltetrahydrofolate synthetase, a key enzyme in reductive acetogenesis. The findings therefore strongly indicate that these cells are new SAOB that can facilitate acetate consumption via SAO, coupled with hydrogenotrophic methanogenesis (SAO-HM). As Methanosarcina is a mixotrophic methanogen, its exact role as acetoclastic or hydrogenotrophic methanogenesis was not verified. Methanoculleus are hydrogenotrophic and thus likely involved in SAO-HM pathway under conditions with high concentrations of acetate.

# **Conflict of Interest**

The authors declare no conflict of interest.

# **Acknowledgements**

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