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Biogas production from hydrothermal liquefaction wastewater (HTLWW): Focusing on the microbial communities as revealed by high-throughput sequencing of full-length 16S rRNA genes

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Wastewater from hydrothermal liquefaction of rice straw

Microbial community analysis
by high-throughput sequencing of Full-Length 16S rRNA genes

Theoretical methane yield

Organic degradation by LT-C
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After petroleum ether extraction
Biogas production from hydrothermal liquefaction wastewater (HTLWW): Focusing on the microbial communities as revealed by high-throughput sequencing of full-length 16S rRNA genes

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Abstract

Hydrothermal liquefaction (HTL) is an emerging and promising technology for the conversion of wet biomass into bio-crude, however, little attention has been paid to the utilization of hydrothermal liquefaction wastewater (HTLWW) with high concentration of organics. The present study investigated biogas production from wastewater obtained from HTL of straw for bio-crude production, with focuses on the analysis of the microbial communities and characterization of the organics. Batch experiments showed the methane yield of HTLWW (R-HTLWW) was 184 mL/g COD, while HTLWW after petroleum ether extraction (PE-HTLWW), to extract additional bio-crude, had higher methane yield (235 mL/g COD) due to the extraction of recalcitrant organic compounds. Sequential batch experiments further demonstrated the higher methane yield of PE-HTLWW. LC-TOF-MS, HPLC and gel filtration chromatography showed organics with molecular weight (MW)<1000 were well degraded. Results from the high-throughput sequencing of full-length 16S rRNA genes analysis showed similar microbial community compositions were obtained for the reactors fed with either R-HTLWW or PE-HTLWW. The degradation of fatty acids were related with *Mesotoga infera*, *Syntrophomonas wolfei* et al. by species level identification. However, the species related to the degradation of other compounds (e.g. phenols) were not found, which could be due to the presence of uncharacterized microorganisms. It was also found previously proposed criteria (97 % and 98.65 % similarity) for species identification of 16S rRNA genes were not suitable for a fraction of 16S rRNA genes.

**Key Words:** hydrothermal liquefaction wastewater; biogas production; degradation of
1. Introduction

Hydrothermal liquefaction (HTL) is an attractive mean to generate renewable bio-energy from biomass. The organic components of biomass are converted into bio-crude under certain temperatures (200-350 °C) and pressures (4-22 MPa), and at the same time a large amount of wastewater containing various organic compounds is produced in the process (Fig 1) (Gai et al. 2015). Previous studies mainly focused on the characterization and potential utilization of the bio-crude (Davis et al. 2011, Xu and Lad 2008), and little attention was paid to the utilization of hydrothermal liquefaction wastewater (HTLWW) even though a significant fraction (20-50 %) of the organics in the biomass was converted and entered into HTLWW (Panisko et al. 2015, Tommaso et al. 2015, Xu and Lad 2008). HTLWW may contain cyclopentenones, phenols, acids et al. depending on the feedstocks and reaction conditions (Cheng et al. 2016, Panisko et al. 2015, Villadsen et al. 2012). Inappropriate disposal of HTLWW would result in the environmental pollution considering its high organic contents. The utilization of HTLWW is therefore important in order to achieve overall environmental and economical sustainability of the HTL process (Nelson et al. 2013).

Anaerobic digestion is widely used in the treatment of organic wastes, which can reduce the environmental pollution of organic wastes and at the same time produce energy in the form of biogas. Only one previous study investigated the biogas potential of HTLWW, which was obtained from the HTL of algae (Tommaso et al. 2015). It was reported that around 44 %-61 % of the COD was removed and converted to biogas after anaerobic digestion, which indicated that part of the organics in HTLWW was not bio-degradable or even toxic to the
microorganisms. Therefore, the characterization of the organic components in HTLWW and elucidation of their degradation during anaerobic digestion are crucial in order to make full utilization of HTLWW. Lignocellulosic materials, different from algae, are abundant in the world, and their utilization via HTL has been studied before (Gan 2012, Kumagai et al. 2007, Tekin et al. 2014), however, the HTLWW remains to be investigated. Since the organic components of HTLWW strongly depend on the feedstocks, it is necessary to investigate the biogas potential of HTLWW obtained from the HTL of lignocellulosic materials. In addition, several previous studies not only extract the bio-crude from solid phase as shown in Fig 1, but also from HTLWW (Leng et al. 2015, Shuping et al. 2010, Yin et al. 2010). The organic solvents could extract some compounds like ketones, phenols and aloxyphenolic according to Yang’s research (Yang et al. 2014), which were recalcitrant or inhibitory molecules for anaerobic digestion. Therefore, the extraction of HTLWW by organic solvents before anaerobic digestion could potentially increase the biogas yield (Cheng et al. 2016, Mottu et al. 2000). However, the effects of organic solvents extraction on the subsequent biogas production from HTLWW was still unknown. Organic solvents, including tetrahydrofuran, toluene, ethyl acetate, acetone, ether, methylene chloride, methanol petroleum ether and n-hexane, are organic solvents that can be used to extract bio-oil from HTL mixture products and their extraction properties were main determine by their polarity (Yang et al. 2014). Some organic solvents are highly toxic to human and therefore only four solvents with different polarities and less toxicity to human were chosen (Semenov 1986).”

Anaerobic digestion involves various microorganisms for the degradation of organic compounds (Luo et al. 2016b). Considering the complex organics in HTLWW, it is necessary
to reveal the microbial communities responsible for the degradation of organics in HTLWW, what would provide in-depth understanding of anaerobic digestion of HTLWW. The rapid development of next-generation sequencing technologies makes it possible to reveal the diversity and structure of the microbial community, with high sequencing depth (Luo et al. 2013). However, currently 16S rRNA genes analysis were mainly based on the second generation sequencing (e.g. 454 GS Junior (Roche), Miseq (Illumina), and Ion Torrent PGM (Life Technologies)), which could only make sequencing on short sequences (< 600bp) and were not able to provide reliable taxonomic information down to genus and species level (Loman et al. 2012, Mosher et al. 2013). Single molecule, real time sequencing (i.e. third generation sequencing) by PacBio RS SMRT chip can generate longer sequences than the second generation sequencing, and is possible to make high-throughput sequencing of the full-length 16S rRNA genes (Mosher et al. 2013, Mosher et al. 2014). A previous study demonstrated the sequences obtained from high-throughput sequencing of full-length 16S rRNA genes of *Shewanella oneidensis MR1* by Pacific Biosciences RS II sequencer can be accurately assigned to the species level (>99 % accuracy) (Mosher et al. 2014). However, high-throughput sequencing of full-length 16S rRNA genes has not been used for the microbial community analysis in mixed cultures (e.g. anaerobic digestion) until now.

Based on the above considerations, the present study aimed to elucidate the mechanisms involved in biogas production from HTLWW obtained from HTL of rice straw. The biogas production potentials from HTLWW extracted by various commonly used organic solvents were investigated, the organics and their removal during anaerobic digestion were characterized, and the microbial community involved in the anaerobic digestion of HTLWW
were revealed by high-throughput sequencing of full-length 16S rRNA genes using Pacific Biosciences RS II sequencer for the first time.

2. Material and methods

2.1. HTLWW

The HTLWW was obtained from a pilot-scale hydrothermal reactor with a volume of 80 L. 3.0 kg of minced rice straw mixed with 47 kg of water were added into the reactor and then heated to 280 °C at 12.0 MPa for 30 min (Chen et al. 2015). The mixture was filtered by a 300-mesh screen after HTL, and the filtrate was HTLWW.

HTLWW was then extracted by petroleum ether (PE), cyclohexane (CH), dichloromethane (DM) and ethyl acetate (EA) to separate parts of the organic components (Duan and Savage 2011, Yang et al. 2014), and they were named as PE-HTLWW, CH-HTLWW, DM-HTLWW and EA-HTLWW, respectively. The raw HTLWW was named as R-HTLWW. For the extraction, 125 mL organic solvent was added to a 500 mL bottle, and 250 mL HTLWW was also added. The bottles were then capped tightly and shaken with the speed of 120 rpm for 10 min by a shaker (Duan and Savage 2011). The mixture was then transferred to a funnel for the separation of organic solvents and HTLWW. The above procedure was repeated for the separated HTLWW for the second time extraction. The four samples PE-HTLWW, CH-HTLWW, DM-HTLWW and EA-HTLWW were then obtained. They were all placed in a refrigerator at -20 °C for further usage. Table 1 presents the COD values of the HTLWW samples and the saturated organic solvents in water.

2.2. Biogas production potentials of HTLWW

Batch experiments were conducted to determine the biogas potentials of HTLWW extracted
by various organic solvents. 118 mL serum bottles were used. 15 mL inoculum and 45 mL BA medium containing a certain amount of HTLWW were added to each bottle. The initial COD value of all the bottles were 0.75 g/L by adding different amounts of HTLWW to the BA medium. The pH value was adjusted to 7.5. All the bottles were flushed with N\textsubscript{2} for 5 min to remove oxygen, and then sealed with butyl rubber stoppers and aluminum screw caps. All the bottles were placed in an incubator with constant temperature 37 °C. The inoculum was obtained from an anaerobic reactor treating cassava stillage in an ethanol plant (Taicang, Suzhou, China). The bottles with only inoculum were used as control. All the experiments were done in triplicates.

2.3. Semi-continuous experiments

Based on the batch experiments, R-HTLWW and PE-HTLWW were used for the anaerobic sequencing batch reactors (ASBR) to determine the long-term biogas production performances, the degradation of organics, and the microbial community involved in the degradation of organics. ASBR has been widely used in previous studies for the treatment of organic wastewater (Angenent et al. 2002, Timur and Özturk 1999). Two 800 mL ASBR were used with working volume 400 mL. The reactors were fed every two days. The reactors were settled for 2 hours before discharging the supernatant, and new substrates were then fed to the reactors. The hydraulic retention time was controlled at 5 days and sludge retention time was controlled at 40 days by discharging excess sludge periodically for each reactor. Initially, 10 g/L glucose was used as the substrate to ensure both reactors had comparable performances. Then reactor R was fed with R-HTLWW, and reactor PE was fed with PE-HTLWW. For reactor R, R-HTLWW was diluted to the same COD concentration as PE-HTLWW in order
to have the same organic loading rate as reactor PE.

2.4 High-throughput sequencing of full-length 16S rRNA genes and bioinformatic analysis

Samples were obtained during the steady-states of both reactors. Total genomic DNA was extracted from each sample using QIAamp DNA Stool Mini Kit (QIAGEN, 51504). The quantity and purity of the extracted DNA were checked by Nanodrop 2000. PCR was then conducted with the primers 27F (AGAGTTTGATCCTGGCTCAG) and 1492R (GGTTACCTTGTTACGACTT) for bacteria and the primers 20F (TTCCGGTTGATCCYCCGCRG) and 1492R for archaea (DeLong 1992). All PCR amplifications were performed using the Taq PCR Core Kit (QIAGEN) with 1 uL template DNA and 20 pmol of each primer. The PCR conditions for bacteria were: 95 °C for 5 min, 28 cycles of three steps: 95 °C for 45 s, 55 °C for 1 min, and 68 °C for 2 min, followed by a final step at 68 °C for 7 min. The PCR conditions for archaea were: 95 °C for 2 min, 27 cycles of three steps: 94 °C for 45 s, 54 °C for 45 s, and 72 °C for 1.5 min, followed by a final step at 72 °C for 7 min. The samples were sent out for sequencing in one cell of the Pacific Biosciences RS II platform combined with the P4/C2 chemistry. The obtained sequences were deposited into the European Nucleotide Archive (ENA) with accession number PRJEB14373. The onboard software provided on the Pacific Biosciences RS II sequencer was used to eliminate CCS (circular consensus sequences) with <99 % predicted accuracy. The low-quality sequences (no exact matches to the forward and reverse primers, and length <1300 bp) and chimeras were removed from the raw sequencing data by MOTHUR program. The numbers of high quality sequences were 7911 (R) and 9099 (PE) for bacteria with
average length of 1390 bp, 1667 (R) and 1905 (PE) for archaea with average length of 1450 bp. The numbers of sequences were normalized to the same sequencing depths (7911 sequences for bacteria and 1667 sequences for archaea) to facilitate the comparison between different samples. The sequences were clustered into operational taxonomic units (OTU) with cutoff 0.03. Rarefaction curves, Shannon diversity index, coverage were also analyzed by MOTHUR program. The sequences were phylogenetically assigned to taxonomic classifications by RDP Classifier with a confidence threshold of 80 %. RDP could only assign the sequences into genus level. In order to get species classification, all the sequences were aligned using BLASTN against NCBI 16S rRNA database with strict criteria (percentage identity at both 97 % and 98.65 %, and alignment length>1300 bp). Both 97 % and 98.65 % of percentage identity were proposed in previous studies for species identification (Kim et al. 2014, Stackebrandt and Goebel 1994, Tindall et al. 2010). MEGAN software was then used to assign the sequences down to species level based on the BLASTN results (Huson et al. 2007). The volumes of gases reported in the present study were at standard temperature and pressure.

2.5. Analytical methods
COD was measured according to APHA (APHA 1995). Gas produced during the anaerobic digestion was detected by GC with thermal conductivity detector. Helium was used as the carrier gas (Liu et al. 2016). GC–MS was used to characterize the chemical compositions of organics extracted from HTLWW by different organic solvents. Gas chromatography was performed on a 30 m HP-INNOWax quartz capillary column with 0.25 mm inner diameter (I.D.) and 0.25 μm film thickness with injection temperature of 250 °C. The column was
initially held at 60 °C for 2 min and heated to 250 °C and held there for 10 min. Helium was used as the carrier gas (1.0 mL/min). A NIST Mass Spectral Database was used for compound identification. HPLC was used to measure the organic acids in the HTLWW samples as described previously (Chen et al. 2015). LC-TOF-MS was used to provide a detailed overview of the organic compounds in the HTLWW samples. It was performed on a Waters ACQUITY UPLC system equipped with a binary solvent delivery manager and a sample manager, coupled with a Waters Micromas Q-TOF Premier Mass Spectrometer equipped with an electrospray interface. Acquity BEH C18 column (100 mm×2.1 mm i.d., 1.7 µm; Waters, Milford, USA) was maintained at 45 °C and eluted with gradient solvent from A:B (99:1) to A:B (0:100) at a flow rate of 0.40 mL/min, where B was acetonitrile (0.1 % (v/v) formic acid) and A was aqueous formic acid (0.1% (v/v) formic acid). The wavelength was 280 nm and the injection volume 5.00 ul, column temperature was 50.0 °C. The source and desolvation temperature were 115 °C and 350 °C respectively. The UV–Vis spectrum was studied using absorptions at 254 wavelength, and the analysis was carried out using a double-beam UV–Vis spectrophotometer from Shimadzu (UV-1800). The molecular weight distributions of HTLWW before and after anaerobic digestion were determined by a GFC analyzer (LC-10ADVP, Shimadzu) according to a previous study (Wen et al. 2012).

3. Results and discussion

3.1. Biogas production potentials of HTLWW extracted by different organic solvents

Fig 2(A) presents the cumulative methane yields of HTLWW extracted by different organic solvents during the biogas potential tests. The methane yields increased fast in the first 10 days for the samples not including DM-HTLWW, which could be related with the
degradation of easy biodegradable organics. Slight increase of methane yields were observed after 10 days for R-HTLWW, PE-HTLWW and CH-HTLWW. Fig 2(B) shows the methane yields of the five HTLWW after 27 days digestion. The methane yield of R-HTLWW (184 mL/g COD) was much lower than the theoretical value (350 mL/g COD), and it indicated there were organics which were difficult to be biodegraded. However, the methane yield of HTLWW was increased after extraction by the organic solvents except DM, which showed that proper organic solvents could improve the anaerobic biodegradability of HTLWW. Further study was conducted to characterize the different HTLWW in order to understand how the organic solvents extraction affected its biodegradability.

3.2 Characterization of HTLWW extracted by different organic solvents
The high methane yield of EA-HTLWW was mainly attributed to the degradation of EA rather than the organics in the HTLWW since EA contributed to more than 98% of the COD in EA-HTLWW (Table 1), and the high methane yield of EA itself was shown in Fig S1. The negligible methane yield of DM-HTLWW was due to the toxicity of DM to the methanogens since no methane was produced when DM alone was used (Fig S1), and the toxicity of DM to methanogens was also reported in previous studies (Kim et al. 1996, McBride and Wolfe 1971). The above results showed that both EA and DM were not suitable as organic solvents since they would increase the difficulty of the subsequent utilization of HTLWW. Both PE and CH had low solubility as demonstrated by their contribution to the total COD in Table 1. Higher methane yield was obtained from PE-HTLWW compared to CH-HTLWW and R-HTLWW, which indicated that PE might have extracted more organics that are difficult to be biodegraded and thereby improved the biodegradability of HTLWW.
GC-MS was conducted to characterize the organics that extracted by different solvents Fig S2, and the relative amounts of major compounds extracted by four different organic solvents were summarized in Table S1. For PE and CH, the two weak polar solvents extracted weak polar components including furans, ketones and phenols. In general, more organics were extracted by PE compared to CH, which might result in the increased methane yield of PE-HTLWW since furans, ketones and phenols were recalcitrant or inhibitory molecules for anaerobic digestion (Speece 1983). Compounds detected from DM and EA organic phases had higher response values than those from PE and CH organic phases in terms of both quantities and types, which was consistent with their higher extracting yields (Table 1), and the results were also agreed with Yang’s (Yang et al. 2014) study where DM and EA with higher polarity were found to extract more organic acids, alcohol, ketones and phenols since many polar organic can be produced in HTL process. As GC-MS in our study only detected the extracted compounds by organic solvents, the organic acids in the HTLWW, which was shown to be dominant in the HTLWW in a previous study (Panisko et al. 2015), were further analyzed by HPLC, and the results were show in Table 2. The concentrations of residual organic acids in HTLWW after extraction decreased with the increase of solvent polarity. Lactic acid, acetic acid and propionic acid, which were easy to be converted to methane (Jeris and McCarty 1965, Vandenberg et al. 1976), were not extracted by PE and CH.

3.3 Biogas production from R-HTLWW and PE-HTLWW in ASBR

The two reactors were operated for around 100 days until steady-states were achieved (Fig S3). The methane yield (153 mL/g COD) of R-HTLWW was significantly higher than that (218 mL/g COD) of PE-HTLWW (P<0.01, ANOVA). The higher methane yield from
PE-HTLWW compared to R-HTLWW was consistent with the batch experiments. However, the methane yields from both PE-HTLWW and R-HTLWW were relatively lower than that from batch experiments, which could be due to the short HRT since the batch experiments allowed the full conversion of biodegradable organics. The above results further demonstrated that PE extraction improved the biodegradability of HTLWW in a certain extent.

3.4 Degradation of organic compounds in ASBR

The UV-VIS (Fig S4) of R-effluent and PE-effluent spectral absorption decreased compared to R-HTLWW and PE-HTLWW, respectively, which was related with the degradation of organic compounds in the anaerobic reactors. However, the absorption between 210-250 nm and 260-300 nm of R-effluent and PE-effluent suggested that ketones and phenols were not fully degraded in the anaerobic reactors (Cheng et al. 2016), which could resulted in the lower methane yield of both R-HTLWW and PE-HTLWW compared to the theoretical value (350 mL/gCOD). LC-TOF-MS identified 785 organic compounds from the four samples. As shown in Fig 3, the dominant organic compounds were well degraded, and the detected organic compounds in R-effluent and PE-effluent were less compared to R-HTLWW and PE-HTLWW, further indicating that most of the organic compounds were degraded in anaerobic reactors. NMDS analysis based on LC-TOF-MS results also showed a clear separation of the samples of influent and effluent. The main organic compounds as determined by GC-MS and HPLC were also identified from LC-TOF-MS as shown in Table S2. It is obvious that most of the organic acids, ketones and about half of the phenols were degraded in the anaerobic reactors.
and the organics left in the effluent were mainly phenols. The full degradation of organic acids were also demonstrated by HPLC analysis as shown in Table 2. Organic acids are preferable substrates for biogas production and therefore they could be fully degraded. Although phenols were reported to be biodegradable under mesophilic conditions (Agarry et al. 2008, Karlsson et al. 2000, Knoll and Winter 1989), there were various types of phenols detected in HTLWW (Table S1), which might result in the partly degradation of the phenols.

Since LC-TOF-MS only detected compounds with molecular weight (MW) less than 1000 in our study as shown in Table S2, the MW distributions of compounds in the samples were further measured by GFC. As shown in Fig 4, there were two peaks for R-HTLWW and PE-HTLWW, which corresponded to the MW 1798 and 180. The results indicated that a considerable amount of organics in the HTLWW were higher MW compounds, and it might be the polymers of HTL intermediate like carbohydrates, cellulose, hemicellulose, lignin and repolymerization compounds (Zhu et al. 2015). After anaerobic digestion, most of the compounds with MW less than 1000 were degraded, which was consistent with the LC-TOF-MS and HPLC results. However, one peak corresponding to MW 9300 was still observed for samples R-effluent and PE-effluent, which suggested that the organics with MW higher than 1000 were not well degraded. Therefore, the lower methane yields of both R-HTLWW and PE-HTLWW compared to the theoretical value (350 mL/gCOD) could be mainly attributed to the presence of MW higher than 1000 in the HTLWW. Furthermore, a small peak with MW around 180 was observed for both R-HTLWW and PE-HTLWW, which might relate with the organics which were not fully biodegraded as mentioned before.
3.5 Microbial community compositions as revealed by high-throughput sequencing of full-length 16S rRNA genes

The samples obtained from the continuous reactors were then used for microbial community analysis. The rarefaction curves of all the samples at 0.03 distance is shown in Fig S5. The curves of bacteria and archaea were overlapped for both samples, and it indicated samples R and PE had similar microbial richness, which was also reflected by the similar OTU numbers (Bacteria, around 1500 for both samples; Archaea, around 210 for both samples) (Table S3). The results showed that PE extraction of HTLWW did not have obvious effects on the microbial community richness. It should be noted that the sequencing depths for both bacteria (7911) and archaea (1167) were still not enough to cover the whole microbial diversity since plateaus were not achieved for all the rarefaction curves. However, the coverage values for bacteria (>86%) and archaea (>90%) indicated that most common OTUs were detected. The coverage values were relatively lower compared to previous studies (e.g. coverage value 97.4% with sequencing depth 50000 for bacteria (Luo et al. 2013), coverage value 98.7% with sequencing depth 63699 for bacteria (Pan et al. 2015)), which was mainly due to the sequencing depths was relatively lower in our study. However, it should be noted all the above mentioned studies were based on high-throughput sequencing of partial 16S rRNA genes (less than 500 bp). The Shannon diversity index provides both species richness and the evenness of the species in the microbial community (Lu et al. 2012). Similar with the microbial richness, the microbial diversities were not affected by PE extraction of HTLWW for both bacteria (around 5.44) and archaea (around 3.3). The higher OTU numbers and Shannon diversity of bacteria compared to archaea were consistent with previous studies.
(Luo et al. 2013, Zhang et al. 2009), further showing bacteria were more diverse than archaea.

The taxonomic classification of bacterial sequences by RDP classifier is shown in Fig 5(A). The similar taxonomic distribution in phylum, class and genus levels were observed for R and PE, further indicating PE extraction did not affect the bacterial communities. It could be due to that PE might only extract unbiodegradable organic compounds and therefore the degraded organic compounds in both reactors R and PE were similar. *Firmicutes, Synergistetes, Chloroflexi*, and *Bacteroidetes* were dominant phyla, and their dominance in mesophilic anaerobic reactors were also reported previously (Luo et al. 2016a, Sundberg et al. 2013). Although *Thermotogae* had high relative abundance, its dominance was mainly found in thermophilic anaerobic reactors (Shi et al. 2013). Genus level identification indicated *Thermotogae* were mainly composed of *Mesotoga*, which was recently reported to be the only mesophilic genus (Nesbø et al. 2012). *Mesotoga* was reported to use lactic acid and its dominance might be related with the degradation of lactic acid as seen in Table 2. *Clostridia* and *Synergistia* were the dominant classes in phylum *Firmicutes* and *Synergistetes*, respectively, and they were known as syntrophic partners together with hydrogenotrophic methanogens for the efficient degradation of lactic acid and VFAs (Li et al. 2016). Their dominances were most probably related with the high concentrations of lactic acid and VFAs in HTLWW (Table 2). The relative abundances of *Anaerolineae* and *Bacteroidia* were between 7-9 % in both samples, and they were capable of hydrolysis and fermentation of carbohydrates to VFAs (Narihiro and Sekiguchi 2007, Robert et al. 2007), however, the carbohydrates were not detected in our study (data not shown), which indicated that their
presence might be related with the degradation of other organics. The genus level classification showed that higher percentages (around 40%) of sequences were unclassified, which was consistent with previous studies (Lu et al. 2012, Luo et al. 2013), and it could be attributed to that most of biogas reactor’s communities are still uncharacterized (Bassani et al. 2015). The dominant genus were \textit{Syntrophobotulus}, \textit{Mesotoga}, and \textit{T78}. \textit{Syntrophobotulus glycolicus} is currently the only known member of the genus \textit{Syntrophobotulus}, however, it can only degrade glyoxylate (Yin et al. 2010), which was not detected in our study. Further species level identification did not detected \textit{Syntrophobotulus glycolicus} (Table 3), and it indicated the genus \textit{Syntrophobotulus} might contain unknown species with different metabolic potentials, which deserves further investigation. The role of \textit{Mesotoga} was mentioned previously for the utilization of lactic acid, while the exact role of \textit{T78} was still unknown (Goux et al. 2015).

Species level identification of full-length 16S rRNA gene sequences would provide more information on the microbial compositions and their metabolic potentials. Table 3 summarized the identified bacterial species. At 97\% similarity, the sequences assigned to species level were 5.6\% and 5.1\% of the total sequences for R and PE, respectively. However, increased sequences (9.9\% for R and 8.8\% for PE) assigned to species level were obtained at 98.65\% similarity. It would be expected less sequences would be assigned to species level with more critical criteria. The higher sequences assigned at 98.65\% similarity was attributed to the algorithm (lowest common ancestor) used by MEGAN (Huson et al. 2007). For instance, one sequence might match two or more species in NCBI 16S rRNA genes database at 97\% similarity, therefore MEGAN could not assign the sequence to
species level. However, the matched species might decrease to one at 98.65% similarity, and therefore it could be assigned to species level. Fig S6 shows that 550 sequences were assigned to the genus *Mesotoga*, however, only 172 sequences were further assigned to species *Mesotoga infera* and *Mesotoga prima* at 97% similarity, while 488 sequences were assigned to the genus *Mesotoga* at 98.65% similarity and all of the sequences were further assigned to species level (Fig S8). The above results indicated that 97% similarity was not enough to make species level identification. Although more sequences were assigned to species level at 98.65%, still the genus *Trichococcus* was not further assigned to species level (Fig S8 and S12). The sequences belonging to *Trichococcus* (Fig S12) were also extracted, and it was found that all the sequences had more than one match to the species in NCBI 16S rRNA genes database at 98.65% similarity (Table S4). 98.65% was previously proposed as the threshold for differentiating two species based on the analysis of 6787 genomes belonging to 1738 species (Kim et al. 2014). However, 98.65% was not the optimal value in our study since microbial community in anaerobic reactor was more diverse. It should be noted that 98.65% was still suitable for the species level identification of sequences belonging to most genus except *Trichococcus* (Fig S8 and S12). As shown in Table 2, lactic acid and VFAs were well degraded during anaerobic digestion, and their degradation could be correlated with the several known species as shown in Table 3. *Mesotoga infera*, *Mesotoga prima*, and *Petrimonas sulfuriphila* were reported to use lactic acid as carbon source (Ben Hania et al. 2015, Grabowski et al. 2005). *Syntrophobacter sulfatireducens* were known as propionate-oxidizing bacteria (Chen et al. 2005). *Syntrophomonas wolfei*, *Syntrophus aciditrophicus* and *Syntrophus buswellii* were demonstrated to be able to degrade
saturated four to eight fatty acids (Jackson et al. 1999, McInerney et al. 1981, Wallrabenstein and Schink 1994). Both *Syntrophus aciditrophicus* and *Syntrophus buswellii* could also degrade benzoate, which is the intermediate during phenol degradation (Na et al. 2016). However, the species for the degradation of phenols, ketones and alkenes were not detected, which were major organic compounds in HTLWW and were degraded in different extents during anaerobic digestion (Table S2). There were two reasons. First and most important, only a fraction of the bacterial species were recognized and characterized until now (Bassani et al. 2015, Schloss and Handelsman 2005), and therefore many new species remained to be explored, which was reflected by the large numbers of “not assigned” and “no hits” sequences as seen in Fig S6-S13. Second, the sequences had high similarity to several known species, and therefore they were not assigned to the species as discussed before.

Fig 5(B) shows the taxonomic classification of archaea sequences by RDP classifier, and the similar taxonomic distribution in order and genus levels for R and PE also suggested PE extraction did not affect the archaea communities. The order *Methanosarcinales* was dominant in both samples, and it was composed by the genus *Methanosaeta* and *Methanosarcina*. The microorganisms belonging to *Methanosaeta* were strict aceticlastic methanogens, and the higher percentage of *Methanosaeta* compared to *Methanosarcina* was due to the low acetic acid concentration in biogas reactors as seen in Table 2 (Karakashev et al. 2005). All the rest sequences were assigned to the orders *Methanomicrobiales* and *Methanobacteriales*, mediating hydrogenotrophic methanogenesis, which was consistent with the syntrophic degradation of fatty acids and the detected syntrophic species as described before. The genus *Methanoculleus* (Order *Methanomicrobiales*) was the main
hydrogenotrophic methanogenesis genus, which was also found to be dominant in other biogas reactors (Jaenicke et al. 2011, Krause et al. 2008). The species level identification by MEGAN showed that 40.9% and 47.9% of the sequences were assigned to species level at 98.65% similarity, which was much higher than that (<10%) for bacteria. It could be due to the higher diversity of bacteria compared to archaea as seen in Table S3 and as reported in previous studies (Luo et al. 2013, Zhang et al. 2009), which resulted in the more uncharacterized species in bacteria than that in archaea. The overwhelming majority of the sequences were assigned to *Methanosaeta concilii*. Although more than 200 sequences were assigned to the genus *Methanosarcina* (Fig S9 and S13), only very few sequences were further classified down to species level. The results further indicated that the 98.65% threshold for differentiating two species was not fully appropriate for all the archaea genus.

### 3.6 Outlook

The present study showed that HTLWW contains relatively higher amount of unbiodegradable organic compounds (e.g. phenols and other high MW (>1000) organic compounds), which were still left in HTLWW after anaerobic digestion. Therefore, further studies via aerobic biodegradation or chemical oxidation should be conducted to remove the residual organic compounds before discharging to the environment (Jang et al. 2015, Moreira et al. 2015). In addition, the usage of catalysis and changes of the HTL conditions also deserves further investigation in order to decrease the formation of unbiodegradable organic compounds without affecting the bio-crude production (Anastasakis and Ross 2011, Tekin and Karagöz 2013). For the first time, the third generation sequencing by PacBio RS SMRT was applied for the high-throughput sequencing of full-length 16S rRNA genes of mixed
cultures. The present study showed that the previously proposed thresholds (97% and 98.65% similarity) for species identification of 16S rRNA genes were not suitable for a fraction of 16S rRNA genes since different species might have high similarity (>98.65%) (Table S4). Therefore, the species level identification of 16S rRNA genes based on similarity is still challenging and remains further investigation. In addition, high percentages of “not assigned” and “no hits” sequences for bacteria sequences were observed, which could be related with the uncharacterized bacteria, and it could be solved with the gradually increased numbers of characterized species in 16S rRNA gene database. Recently, there were studies focusing on the identification of the genomes of microorganisms from mixed cultures by metagenomic analysis, which is independent of traditional cultivation methods, and thereby it might expand the sequences in 16S rRNA gene database (Bassani et al. 2015, Campanaro et al. 2016).

4. Conclusions

The present study showed that the methane yield of HTLWW (R-HTLWW) was 184 mL/g COD, while HTLWW after petroleum ether extraction had higher methane yield (235 mL/g COD) due to the extraction of recalcitrant organic compounds. The higher methane yields of PE-HTLWW (225 mL/gCOD) compared to R-HTLWW (160 mL/gCOD) was also demonstrated in the continuous experiments. Further study showed that organics with molecular weight (MW)<1000 were well degraded by LC-TOF-MS, HPLC and gel filtration chromatography analysis. The results from high-throughput sequencing of full-length 16S rRNA genes showed that similar microbial community compositions were obtained for the reactors fed with either R-HTLWW or PE-HTLWW, and the degradation of fatty acids were related with *Mesotoga infera*, *Syntrophomonas wolfei* et al. by species level identification.
However, the species related to the degradation of other compounds (e.g. phenols) were not found, and it could be due to the presence of uncharacterized microorganisms. The study also showed that previously proposed criteria (97 % and 98.65 % similarity) for species identification of 16S rRNA genes were not suitable for a fraction of 16S rRNA genes.

Acknowledgements

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Supporting Information

Supporting Information includes Tables and Figures as noted in the text.

References


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346-351.


Stackebrandt, E. and Goebel, B.M. (1994) Taxonomic note: A place for DNA-DNA reassociation and 16S rRNA sequence analysis in the present species definition in


Table 1 COD values of HTLWW and organic solvents

<table>
<thead>
<tr>
<th>Organic solvents</th>
<th>Agent polarity (gCOD/L)</th>
<th>Saturated solvent (gCOD/L)</th>
<th>HTLWW COD (gCOD/L)</th>
<th>COD contributed by COD extracting percent (%)</th>
<th>COD extracting percent (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>-</td>
<td>-</td>
<td>-</td>
<td>20.74</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>PE</td>
<td>0.01</td>
<td>0.09</td>
<td>15.99</td>
<td>0.56</td>
<td>23.33</td>
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<tr>
<td>CH</td>
<td>0.1</td>
<td>0.22</td>
<td>17.63</td>
<td>1.25</td>
<td>16.06</td>
</tr>
<tr>
<td>DM</td>
<td>3.4</td>
<td>7.95</td>
<td>13.28</td>
<td>59.86</td>
<td>74.30</td>
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<tr>
<td>EA</td>
<td>4.3</td>
<td>146.35</td>
<td>148.90</td>
<td>98.29</td>
<td>87.70</td>
</tr>
<tr>
<td>Name</td>
<td>R-HTLWW</td>
<td>PE-HTLW</td>
<td>CH-HTL</td>
<td>DM-HTLW</td>
<td>EA-HTL</td>
</tr>
<tr>
<td>--------------</td>
<td>---------</td>
<td>---------</td>
<td>--------</td>
<td>---------</td>
<td>--------</td>
</tr>
<tr>
<td>Lactic acid</td>
<td>3722</td>
<td>3708</td>
<td>3698</td>
<td>1628</td>
<td>592</td>
</tr>
<tr>
<td>Acetic acid</td>
<td>1802</td>
<td>1792</td>
<td>1782</td>
<td>740</td>
<td>–</td>
</tr>
<tr>
<td>Propionic acid</td>
<td>680</td>
<td>657</td>
<td>657</td>
<td>399</td>
<td>–</td>
</tr>
<tr>
<td>N-butyric acid</td>
<td>281</td>
<td>289</td>
<td>260</td>
<td>59</td>
<td>–</td>
</tr>
<tr>
<td>Isovaleric acid</td>
<td>146</td>
<td>142</td>
<td>133</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>
Table 3 Species level identification of the full-length 16S rRNA sequences

<table>
<thead>
<tr>
<th>Species/Genus</th>
<th>Number of sequences (97% Similarity)</th>
<th>Number of sequences (98.65% Similarity)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>R</td>
<td>PE</td>
</tr>
<tr>
<td><strong>Bacteria</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Acinetobacter seohaensis</em></td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td><em>Advenella faeciporci</em></td>
<td>9</td>
<td>2</td>
</tr>
<tr>
<td><em>Alkalibacter saccharofermentans</em></td>
<td>9</td>
<td>5</td>
</tr>
<tr>
<td><em>Aminivibrio pyruvatophilus</em></td>
<td>10</td>
<td>12</td>
</tr>
<tr>
<td><em>Halothiobacillus neapolitanus</em></td>
<td>15</td>
<td>6</td>
</tr>
<tr>
<td><em>Mesotoga infera</em></td>
<td>169</td>
<td>185</td>
</tr>
<tr>
<td><em>Mesotoga prima</em></td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td><em>Ornatilinea apprima</em></td>
<td>14</td>
<td>8</td>
</tr>
<tr>
<td><em>Parasporobacterium paucivorans</em></td>
<td>100</td>
<td>93</td>
</tr>
<tr>
<td><em>Petrimonas sulfuriphila</em></td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td><em>Pseudomonas caeni</em></td>
<td>4</td>
<td>7</td>
</tr>
<tr>
<td><em>Pseudomonas stutzeri</em></td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td><em>Syntrophobacter sulfatireducens</em></td>
<td>25</td>
<td>2</td>
</tr>
<tr>
<td><em>Syntrophomonas wolfei</em></td>
<td>28</td>
<td>37</td>
</tr>
<tr>
<td><em>Syntrophus aciditrophicus</em></td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td><em>Youngiibacter fragilis</em></td>
<td>9</td>
<td>6</td>
</tr>
<tr>
<td>Others* (28 species)</td>
<td>30</td>
<td>26</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>440 (5.6%)</strong></td>
<td><strong>404 (5.1%)</strong></td>
</tr>
<tr>
<td><strong>Archaea</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Methanosaeta concilii</em></td>
<td>746</td>
<td>896</td>
</tr>
<tr>
<td><em>Methanoculleus palmolei</em></td>
<td>90</td>
<td>109</td>
</tr>
<tr>
<td><em>Methanomassiliicoccus luminyensis</em></td>
<td>30</td>
<td>32</td>
</tr>
<tr>
<td>Others* (11 species)</td>
<td>16</td>
<td>13</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>882</strong></td>
<td><strong>1050</strong></td>
</tr>
</tbody>
</table>

*Others* are the species with numbers of sequences less than 5
Fig 1 Hydrothermal liquefaction process
Fig 2 Biogas production potentials of HTLWW (A) time courses of methane production (B) methane yields
Fig 3 Heatmap (A) and NMDS (B) analysis of the samples based on LC-TOF-MS results
Fig 4 GFC analysis of the samples
Fig 5 Taxonomic classification of bacteria (A) and archaea (B) sequences based on the high-throughput sequencing of full-length 16S rRNA genes
Highlights:

- The methane yield of HTLWW was increased after petroleum ether extraction
- Organics in HTLWW with molecular weight (MW) < 1000 were well degraded
- High-throughput sequencing of full-length 16S rRNA genes was applied
- Microbial community compositions were analyzed down to the species level