Optimization and microbial community analysis of anaerobic co-digestion of food waste and sewage sludge based on microwave pretreatment

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HIGHLIGHTS

- Methane production was enhanced by both co-digestion and MW pretreatment.
- The optimized ratio was 3:2 for anaerobic co-digestion based on MW pretreatment.
- Propionic accumulation led to lower methane production of MW-FW compared to MW-SS.
- Bacteroides only dominated in co-digestion, Methanosphaera only dominated in MW-FW.
- Methanosarcina was the predominant methanogens for mono-SS and MW-SS.

ABSTRACT

The effects of microwave pretreatment (MW) on co-digestion of food waste (FW) and sewage sludge (SS) have never been investigated. In this study, a series of mesophilic biochemical methane potential (BMP) tests were conducted to determine the optimized ratio of FW and SS based on MW, and the evolution of bacterial and archaeal community was investigated through high-throughput sequencing method. Results showed that the optimized ratio was 3:2 for co-digestion of FW and SS based on MW, and the methane production was 316.24 and 338.44 mLCH4/gVS added for MW-FW and MW-SS, respectively. The MW-SS was superior for methane production compared to MW-FW, in which accumulation of propionic acid led to the inhibition of methanogenesis. Proteiniborus and Parabacteroides were responsible for proteins and polysaccharides degradation for all, respectively, while Bacteroides only dominated in co-digestion. Methanosphaera dominated in MW-FW at the active methane production phase, while it was Methanosarcina in MW-SS and mono-SS.

1. Introduction

Sewage sludge (SS) production had an average annual growth of 13% from 2007 to 2013, and 6.25 million tons dry solids were produced in 2013 in China, and almost 80% of it has not been properly stabilized (Yang et al., 2015). The treatment of sewage sludge has become a big challenge due to strict legal regulations, land shortages, rising costs, and public concern along with fast urbanization and economic development (Wei et al., 2003). Anaerobic digestion (AD) of SS due to the production of renewable energy and sludge reduction has been carried out widely, and ca. 50% of the total of 2500 waste water treatment plants (WWTPs) were designed with AD up to 2010 in China (Duan et al., 2012). However, its efficiency is largely limited due to the relatively slow hydrolysis process and low C/N ratio (6–9), and hydrolysis has been reported as the rate-limiting step (Toreci et al., 2009).

About 6.0 × 107 tons of food waste (FW) were produced according to China Statistical Yearbook 2011, and the increasing rate of FW production was higher than 10% every year due to population growth and rising living standards. AD is preferred as an efficient
pathway for treatment of FW due to its high organic contents and excellent biodegradability (Zhang et al., 2007). Nevertheless, the accumulation of volatile fatty acids (VFAs) occurred frequently due to the labile organic fraction which was hydrolyzed too fast, and this may lead to process failure due to the acidification of AD (Kawai et al., 2014; Yong et al., 2015). Thus, the anaerobic co-digestion of FW and SS became increasingly popular, with the advantage of adjusting the C/N ratio, increasing the methane yields, diluting harmful substances, and also mediating the hydrolysis of FW and SS (Kawai et al., 2014; Lee et al., 2009; Yong et al., 2015).

Cell wall destruction or disruption is considered as an effective way of enhancing AD efficiency of SS which is mainly composed of microbial cells within an extra-cellular polymeric floc matrix (Baier and Schmidheiny, 1997), because hydrolysis was enhanced through SS pretreatment by releasing the intracellular organics. Microwave pretreatment (MW) of SS (MW-SS) was widely investigated to enhance the AD efficiency of SS, and volatile solids (VS) removal, methane production and AD stabilization were improved (Coelho et al., 2011). Meanwhile, the effects of microwave pretreatment of FW (MW-FW) on AD were investigated, while contrary results existed. Marin et al. (2010) indicated that anaerobic biodegradability could be improved by 9% after MW of model FW. In contrary, Shahriari et al. (2013) recommended that digester staging without MW be employed to maximize methane production in comparison with MW-FW. However, high temperature (175 °C or 145 °C) was adopted in these studies during MW pretreatment, there is little information on the effects of MW-FW under low temperature (<100 °C) on the AD. Besides, although the effects of MW-SS or MW-FW on AD efficiency have been investigated, there is still little information concerning the underlying microbial mechanisms.

In addition, although several studies focusing on the bacterial or archaeal community of anaerobic co-digestion of FW and SS have been conducted previously under hyper-thermophilic and thermophilic conditions using PCR-DGGE (Lee et al., 2009) or dry AD by high-throughput sequencing method (Cho et al., 2013), the information on bacterial and archaeal community of anaerobic co-digestion of FW and SS based on MW is still unclear, and anaerobic co-digestion of SS and FW based on MW has never been investigated till now to our knowledge. Therefore, the aims of this study were as follows: (1) to determine whether MW could enhance anaerobic co-digestion of FW and SS; (2) to find out the optimized ratio of FW and SS based on MW for anaerobic co-digestion; (3) to clarify the evolution of bacterial and archaeal community under optimized ratio using high-throughput sequencing method.

2. Methods

2.1. Pretreatment of substrates and sludge inoculum

Food waste (FW) was collected daily from the dining hall of Research Center for Eco-environmental Sciences, Chinese Academy of Sciences, Beijing, China, which mainly contains leftovers of cooked foods like meats, fishes, rice, breads, noodles and vegetables, and the hard matters such as bones, toothpicks, chopsticks and napkins were discarded. Grease in FW was full of oils and fats, and led to acidification of AD easily (Zhu et al., 2011), thus, grease was trapped through washing the food waste for 3–4 times, and then FW was homogenized and crushed to particle size of ca. 2 mm and stored at 4 °C before use. Dewatered sewage sludge used for co-digestion with FW was collected from Beijing Qinghe WWTP. Due to the low moisture content of dewatered SS, it is hard to mix thoroughly with FW, thus, moisture content was mediated to ca. 90% by adding water. A laboratory industrial microwave reactor operating at 2,450 MHz and ambient pressure was used for MW pretreatment of FW and SS with a maximum power and temperature of 1000 W, 100 °C, respectively. 300–500 ml raw FW or SS was heated to 100 °C by the MW reactor operated at 600 W without additional retention. MW-FW and MW-SS were then cooled down to room temperature before use. Seeding sludge was collected from a mesophilic AD reactor treating SS in Beijing Xiaohongmen WWTP, and sludge after setting down for ca. 2–3 days was used for inoculum. The characteristics of FW, SS and seeding sludge were shown in Table 1.

2.2. BMP (biochemical methane potential) tests

The BMP test, which is considered as the most suitable method for a relatively easy evaluation of the anaerobic digestibility (Nges and Liu, 2009), was used as a tool for evaluating the methane production and biodegradability of the mixtures of FW and SS at different ratios based on MW. In this study, BMP tests were carried out by an Automatic Methane Potential Test System II (Bioprocess Control, Sweden) in which digesters were a series of serum bottles (working volume: 0.4 L or 1.8 L) equipped with plastic caps including agitators and rubber stoppers. The substrates used for BMP tests were mixed thoroughly at a ratio according to total solid (TS) shown in Table 2. 0.4 L or 1.8 L of the mixture of substances and inoculums at a ratio of 5:1 (TS) were transferred to each bottle, and the final TS was adjusted to about 3.5% or 7% as shown in Table 2. After sealing, the headspaces were flushed with nitrogen gas for 3–5 min to remove traces of oxygen. Then the bottles were incubated in a water bath to control temperature at 37 ± 0.5 °C. Before biogas production yields were automatically measured, CO2 was removed by a gas-washing bottle containing 3 M NaOH solution. The termination of BMP tests was judged by no obvious accumulation biogas production from bottles (about 35 days). 19 BMP tests were conducted in total in this study, and there was no sampling for thirteen BMP tests conducted in 0.4 L working volume bottles which was used to find out the optimized ratio of FW and SS based on MW for anaerobic co-digestion. Three 1.8 L working volume bottles in duplicate were mainly used to find out the mechanisms for the optimized ratio of FW and SS based on MW pretreatment, thus, sampling was conducted on days 1, 5, 12, 19, 26 and 33 for further analysis.

Table 1

| Characteristics                | Food waste | Dewatered sewage sludge | Inoculated sludge | Microwaved food waste | Microwaved sewage sludge |
|-------------------------------|------------|--------------------------|-------------------|-----------------------|--------------------------|
| Moisture content (%)          | 88.12 ± 0.34 | 89.35 ± 0.07             | 97.59 ± 0.05      | 87.03 ± 0.05          | 88.10 ± 0.05             |
| Total solids (g/L)            | 136.83 ± 11.65 | 125.87 ± 7.56           | 24.19 ± 1.20      | 139.67 ± 0.19         | 135.77 ± 1.44            |
| Volatile solids (g/L)         | 130.06 ± 11.51 | 84.98 ± 10.88           | 12.92 ± 0.58      | 132.83 ± 0.40         | 101.49 ± 5.45            |
| SCOD (g/L)                    | 59.35 ± 20.65 | 10.88 ± 1.68            | 0.59 ± 0.02       | 44.20 ± 10.56         | 24.58 ± 8.70             |
| NH4-N (mg/L)                  | 111.05 ± 0.90 | 794.13 ± 135.88         | 335.2 ± 14.80     | 201.6 ± 15.70         | 985.0 ± 45.70            |
| Proteins (g/L)                | 1.46 ± 0.10  | 2.33 ± 0.55             | –                 | 2.12 ± 0.30           | 4.83 ± 0.70              |
| Polysaccharides (g/L)         | 40.16 ± 22.05 | 1.10 ± 0.04             | –                 | 19.41 ± 13.70         | 2.90 ± 0.30              |
The detail information of mesophilic BMP tests for anaerobic co-digestion of FW and SS based on MW pretreatment.

| Pretreatment | FW:SS | SCOD removal (%) | Proteins removal (%) | NH₄-N removal (%) | VS | SCOD (g/L) | Proteins (g/L) | NH₄-N (g/L) |
|--------------|-------|------------------|---------------------|------------------|----|-----------|----------------|------------|
| FW           | 32.98 ± 0.51 | 4.84 5.64 ± 0.23 | 44.75 ± 10.7 9.79 ± 1.71 | 2.42 ± 0.51 0.35 ± 0.21 0.25 ± 0.02 | 16.92 | 4 |
| SS           | 35.03 ± 1.31 | 6.2 6.73 ± 0.12 | 36.15 ± 11.2 2.15 ± 0.50 0.2 ± 0.12 0.31 ± 0.18 0.29 ± 0.03 | 36.98 41.05 12.24 18.62 0.88 ± 0.37 7.62 25.22 142.70 |
| FW:SS        | 1:1     | 34.66 ± 0.42     | 0.73 5.95 ± 0.10 | 40.18 ± 13.0 5.86 ± 0.72 | 0.75 ± 0.34 | 53.91 | 82.00 86.58 |
| MW-FW:SS     | 2:1     | 35.40 ± 3.23     | 0.69 6.02 ± 0.05 | 39.12 ± 14.7 7.33 ± 0.81 | 0.55 ± 0.23 | 56.98 | 85.34 94.97 |
| MW-FW:SS     | 3:2     | 35.24 ± 0.22     | 0.69 6.09 ± 0.04 | 38.62 ± 12.7 6.97 ± 0.73 | 1.14 ± 0.34 | 59.27 | 81.35 162.4 |
| MW-FW:SS     | 2:3     | 35.14 ± 0.35     | 0.74 6.35 ± 0.11 | 39.60 ± 8.7 5.18 ± 0.45 | 0.83 ± 0.24 | 52.74 | 69.89 296.98 |
| FW:MW-SS     | 2:1     | 34.39 ± 1.46     | 0.70 5.99 ± 0.07 | 39.71 ± 12.7 8.31 ± 1.01 | 0.43 ± 0.18 | 62.35 | 83.99 80.09 |
| FW:MW-SS     | 1:1     | 34.91 ± 0.82     | 0.73 6.11 ± 0.13 | 38.49 ± 13.7 7.98 ± 0.95 | 0.61 ± 0.18 | 55.45 | 85.30 73.57 |
| MW-FW:SS     | 3:2     | 71.17 ± 3.41     | 0.69 7.14 ± 0.17 | 55.35 ± 13.7 19.2 ± 1.57 | 0.94 ± 0.37 | 39.70 | 54.07 68.58 |
| FW:MW-SS     | 3:2     | 72.31 ± 2.47     | 0.68 7.37 ± 0.03 | 62.75 ± 12.7 18.5 ± 1.73 | 1.08 ± 0.39 | 54.86 | 76.90 66.46 |

2.3. Chemical analysis

TS, TCOD, SCOD, pH, NH₄-N and VS were measured according to standard methods (APHA et al., 2005). Samples were centrifuged at 4000 rpm for 10 min and filtered through 0.45 μm cellulose membrane. The filtrate was analyzed for pH, NH₄-N, SCOD, proteins, polysaccharides, and VFAs. The pH was determined using a pH-meter (pHs-3C, Leici Co. Ltd., Shanghai). Ammonia (NH₄-N) was determined by the colorimetric method. SCOD was measured using Hach 8000 methods with a DR 2800 spectrometer (Hach, USA). VFAs were analyzed by a gas chromatograph (Shimadzu GC-14A Plus, Japan) equipped with a flame ionization detector (GC-FID) and stabilwax-DA column (30 m × 0.32 mm × 0.25 μm). The temperatures of the injection port and the detector were set at 240 °C and 260 °C, respectively, and helium was the carrier gas at the initial pressure of 16.3 kPa. The six VFAs standards including acetic acid, propionic acid, butyric acid, iso-butyric acid, valeric acid and iso-valeric acid were purchased from Sigma (Sigma, USA), and the total VFAs concentration in this study was defined as the sum of them. The polysaccharides and proteins content was measured as previously suggested (Zhang et al., 2015).

2.4. DNA extraction

Five points of each BMP test for the optimized ratio of FW and SS based on MW on days 1, 5, 12, 19 and 33 were chosen for bacterial and archaeal community analysis. 4 mL of each sludge sample was centrifuged at 10,000 rpm for 10 min, and the pellet was used for DNA extraction using FastDNA Spin Kit for Soil (MP Biomedicals, USA), in triplicate according to manufacturer’s instructions, and the resulting extracts were composited to average out bias in sampling and extraction. Quality and concentration of the extracted DNA were determined through 1% agarose gel electrophoresis and NanoDrop ND-1000 (NanoDrop, USA), respectively.

2.5. Bacterial and archaeal community analysis

PCR primers 515F and 806R targeting both bacteria and archaea 16S rRNA V4 region were selected for bacterial community analysis using Illumina high-throughput sequencing method (Caporaso et al., 2010). While the abundance of archaeal community was much lower in comparison with bacterial community, archaeal community was specially analyzed using nested PCR. In the nested PCR approach, the specific archaeal community was firstly amplified using the primers Arch340F (5’-CCCTAYGGGGGYGCASCAG-3’) and Arch1000R (5’-GAGARWGRTGATCATGCCG-3’) as described by Ganttner et al. (2011), and then the PCR product was used as a template in the second PCR using the primers Arch349F (5’-GYGGCAGKCGGGMGA-3’) and Arch806R (5’-GACACTACVGGGTATC-3’) (Takai and Horikoshi, 2000). Barcodes unique to each sample were incorporated before the forward primers, which allowed the identification of each sample in a mixture for an Illumina sequencing run. DNA was amplified in triplicate for each sample, and then PCR amplicons were further purified with a DNA purification kit (Gel Purification kit, Sangon, China), and the concentrations were determined using a spectrometry (Qubit 2.0, Invitrogen, USA). Amplicons from different samples were then mixed to achieve equal mass concentrations in the final mixture, which were sent out to Sangon Co., Ltd., in Shanghai for small-scale fragment library construction and pair-end sequencing using the Illumina MiSeq sequencing system (Illumina, USA).

Sequencing reads were assigned to each sample according to the unique barcode of each sample. Pairs of reads from the original DNA fragments were firstly merged using FLASH (Magoe and Salzberg, 2011), and then PRINSEQ was used for the quality control.
of these merged reads (Schmieder and Edwards, 2011). The barcode and primers then were removed. All the reads were further uploaded to MG-RAST (http://metagenomics.anl.gov/linkin.cgi?project=14442), PCR chimeras were filtered out using UCHIME (Edgar et al., 2011). After the above filtration, the average length of all of the clean reads was 250 bp and 379 bp, and the average sequencing depth was ca. 36,000 and 30,000 clean reads for bacterial and archaeal community analysis, respectively. The taxonomic classification of the sequences was carried out using the Ribosomal Database Project (RDP) Classifier at the bootstrap cutoff of 80% suggested by the RDP. Furthermore rarefaction curves and richness indices were calculated using the relevant RDP pipeline modules as described elsewhere (Zhang et al., 2015).

2.6. Data analysis

The results of chemical parameters were visualized through Origin 9.0 (OriginLab, USA). A heat map of the top 10 genus in each sample based on reads log2 transformed was built using the HemI (http://hemi.biocuckoo.org/). Circos graph showing the evolution of bacterial community of each sample in different treatments at phylum level was produced by the Circos software (http://circos.ca/). According to the relative content of each genus from the classification, a principal component analysis (PCA) was performed using Canoco 5.0 (Microcomputer Power, USA).

3. Results and discussion

3.1. Effects of MW on anaerobic co-digestion of FW and SS

As shown in Fig. 1A, all the biogas production in anaerobic co-digestion of FW and SS with and without MW could be enhanced significantly in comparison with mono-FW and mono-SS. After 35 days of BMP tests, the cumulative methane production was calculated to be 70.72, 142.70, 296.75, 311.26 and 317.92 mLCH4/gVSadded for mono-FW, mono-SS, FW:SS (1:1), MW-FW:SS (1:1) and FW:MW-SS (1:1), respectively. FW mainly consisted of polysaccharides, while SS was mainly constituted by proteins (Table 1). The anaerobic degradability of poly-saccharides was much higher than proteins (Tommaso et al., 2003). This could be confirmed by the maximum daily methane production of mono-FW on day 1, while mono-SS on days 1–8. The least biogas production of mono-FW maybe due to the acidification as previously suggested (Kawai et al., 2014), while the cumulative production of mono-FW maybe due to the acidification as previouly suggested by the RDP. Furthermore rarefaction curves and richness indices were calculated using the relevant RDP pipeline modules as described elsewhere (Zhang et al., 2015).

3.2. Optimized ratio of FW and SS for anaerobic co-digestion based on MW

In order to determine the optimized ratio of FW and SS based on MW, BMP tests of five ratio of FW and SS (2:1, 3:2, 1:1, 2:3 and 1:2) (TS) with MW-FW or MW-SS were further conducted. The optimized ratio of FW and SS was always 3:2 (TS) for both MW-FW and MW-SS as revealed by Fig. 1B and C, and the detail methane production was listed in Table 2. The maximum methane production was 316.24, 338.44 mLCH4/gVSadded for MW-FW, MW-SS at the optimized ratio of 3:2, respectively, and the methane production of MW-SS was 6.94% higher than that of MW-FW at the same ratio. These results further confirmed that the enhancement was higher for co-digestion of FW and SS through MW-SS instead of MW-FW.

The daily biogas production showed a similar profile at different treatments. It peaked on day 1, and then decreased significantly, while increased on day 4. The active phase for methane production ranged from day 5 to 16. Polysaccharides removal rate was generally higher for the MW-FW compared to that of MW-SS, while proteins removal rate was much higher for the MW-SS in comparison with the MW-FW (Table 2). The final NH4+-N was generally lower for MW-SS at the same ratio, which corresponded to higher proteins removal. TS and VS removals decreased along with the increasing addition of SS for both MW-FW and MW-SS (Table 2). This could be explained by the relative low TS and VS removal in comparison with FW. Initial and final NH4+-N also showed the same pattern. Although TCOD removal decreased along with the increasing addition of SS for MW-FW and MW-SS (Table 2). This is easy to understand the effects of MW pretreatment to enhance anaerobic digestion of SS has been widely investigated (Eskicioglu et al., 2007a,b). The main fraction of SS consists of a polymeric network formed by extracellular polymeric substances (EPS) and microbial cells that are resistant to direct anaerobic degradation since cell walls and EPS present physical and chemical barriers, while MW pretreatment can achieve higher SS floc, cell destruction and release of EPS and intracellular materials into the soluble phase, and this increased biogas production. The increased SCOD after MW pretreatment could elucidate this assumption (Table 1).

Maillard compounds start to form at temperatures above 100 °C depending on the retention time, and the formation of more complex compounds, such as acrylamides and other vinylogous compounds, increases at higher temperatures like 180 °C (Stadler et al., 2004), thus, low temperature MW pretreatment of FW was adopted in this study. MW-FW could also further increase the biogas production for the co-digestion of FW and SS. The enhancement of co-digestion of FW to anaerobic digestion has also been confirmed previously, and the solubilization of FW could be increased (Table 1). However, the enhancement of co-digestion was slightly lower for the MW-FW, and this could be explained by higher enhancement of release of SCOD of MW-SS (ca. 2.0 times) in comparison with MW-FW (ca. 1.2 times). However, slightly lower TS and VS removal (Table 2) compared with the co-digestion of FW and SS without MW indicated the formation of complex compounds which were hard to be biodegraded after MW pretreatment.

It is important to highlight that both MW-FW and MW-SS could further enhance the methane production and TCOD removal of SS on the anaerobic co-digestion of FW and SS. The MW pretreatment to enhance anaerobic digestion of SS has been widely investigated (Eskicioglu et al., 2007a,b). The main fraction of SS consists of a polymeric network formed by extracellular polymeric substances (EPS) and microbial cells that are resistant to direct anaerobic degradation since cell walls and EPS present physical and chemical barriers, while MW pretreatment can achieve higher SS floc, cell destruction and release of EPS and intracellular materials into the soluble phase, and this increased biogas production. The increased SCOD after MW pretreatment could elucidate this assumption (Table 1).

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removal at the optimized ratio. The different mechanisms behind MW-FW and MW-SS needed to be further elucidated.

3.3. Mechanisms of anaerobic co-digestion underlying the optimized ratio based on MW

Further studies concerning the underlying mechanisms of anaerobic co-digestion of FW and SS at the optimized ratio based on MW were conducted to find out the different effects on anaerobic co-digestion between MW-FW and MW-SS. Previous studies have shown that MW pretreatment is more effective for sludge anaerobic digestion with high solids concentration (Eskicioglu et al., 2007a,b), thus, the TS was doubled to enhance the anaerobic co-digestion of FW and SS. BMP tests of mono-SS, MW-FW:SS (3:2) and FW:MW-SS (3:2) were further re-conducted in duplicate in a bigger scale to monitor the evolution of relative parameters. Mono-FW anaerobic digestion was not conducted due to its end of methane production untimely as suggested previously (Fig. 1A).

3.3.1. Evolution of chemical parameters and VFAs

Higher solids concentration indeed increased methane production (Table 2), and the methane production was 191.90, 367.62 and 389.64 mLCH₄/gVSadded for mono-SS, MW-FW:SS (3:2) and FW:MW-SS (3:2), respectively. Compared to mono-SS, co-digestion of FW and SS based on MW pretreatment increased by 1.92 and 2.03 times, respectively. Furthermore, MW-SS showed its advantage of enhancement of anaerobic co-digestion in comparison with MW-FW (Fig. 1D). The final methane production of MW-SS was 5.99% higher than that of MW-FW. Concerning the daily methane production, the similar pattern to previous BMP tests was developed, i.e., peak on day 1, another peak on day 10 and active methane production on days 5–16. All the peaks were generally higher for MW-SS than that of MW-FW.

MW-SS showed lower pH and NH₄⁺-N concentrations throughout the tests, and there was a peak value of SCOD on day 5 for all the treatments. However, the peak value for mono-SS (18, 720 mg/L) and SS (31, 810 mg/L) was much higher than that of mono-SS (18, 720 mg/L). The increase of SCOD for mono-SS could be due to the release of refractory substances, and this was further confirmed by the increase of polysaccharides and proteins concentrations. MW-SS showed higher TCOD, SCOD, TS, VS and proteins removal but lower polysaccharides removal in comparison with MW-FW, and the significant reduction of polysaccharides could explain the peak daily methane production of co-digestion.

Changes of VFAs concentration and composition could be used to indicate the performance of anaerobic digestion (Wang et al., 2014). The accumulation of VFAs appeared on day 5 for all the treatments, and then it decreased gradually (Fig. 2). The average peak value of VFAs concentration was 9, 354.34, 23, 022.8 and 24, 537.9 mg/L for mono-SS, MW-FW and MW-SS, respectively. The highest release of VFAs for MW-SS determined its highest methane production. VFAs are subsequently decreased as a result
of their uptake by the anaerobic microorganisms. The composition of VFAs changed significantly along with anaerobic digestion. Acetic acid was predominant on day 0–12, and also peaked on day 5. As for MW-SS and mono-SS, acetic acid decreased along with anaerobic digestion, and little acetic acid left at the final (241.48 and 149.80 mg/L, respectively). Although propionic acid was difficult to be utilized, there was no accumulation for MW-SS and mono-SS. Propionic acid, valeric acid, iso-valeric acid, butyric acid and iso-butyric acid all decreased to ND (not detected). However, it is quite different for MW-FW, propionic acid accumulated to 6262.96 mg/L on day 26, while acetic acid decreased on day 12–19, it rebounded to 2103.57 mg/L at the final. This indicated that higher propionic acid accumulation has led to inhibition of methanogenesis which further contributed to the accumulation of acetic acid. It was assumed that accumulation of propionic acid was the reason for the lower methane production of MW-FW in comparison with MW-SS.

3.3.2. Evolution of bacterial community

Detailed evolution of bacterial community was investigated to further find out underlying mechanisms of the difference between the effects of MW-FW and MW-SS on anaerobic co-digestion of FW and SS. After chimer analysis, a total of 499, 307 high quality reads for the 15 samples ranging from 32, 422 to 34, 461 reads were generated for further analysis. The evolution of bacterial alpha indexes was shown in Supporting information. These high quality reads were assigned to different taxa levels using the RDP classifier. A total of 33 phylum were identified, and Firmicutes, Proteobacteria, Bacteroidetes and Actinobacteria were the predominant phylum in the bacterial community as shown in Fig. 3A. Firmicutes was enriched by 5.05, 6.45 and 4.61 times for mono-SS, MW-SS and MW-FW, respectively. The abundance of Firmicutes all increased significantly on day 5. After that, Firmicutes changed little for mono-SS and MW-SS while the abundance increased much further on day 12 for MW-FW and changed little to the end. The evolution of Proteobacteria was quite different for different treatments especially for mono-SS. The abundance of Proteobacteria decreased significantly for MW-SS and MW-FW in comparison with mono-SS, and it was the maximum contributor to the difference of mono-SS and co-digestion. The phylum Actinobacteria showed similar evolution to Proteobacteria. As for Bacteroidetes, the abundance decreased a little for mono-SS while increased for MW-SS and MW-FW. There was a common phenomenon for these four dominant phylum that the maximum or minimum abundance, that is, greatest changes, all appeared on day 5, which indicated that recombination of bacterial community occurred at acidification phase corresponding to VFAs accumulation.

The top 10 abundant genus in each sample (a total of 61 genus) were selected for a detailed understanding of the evolution of microbial community structure as suggested previously (Zhang et al., 2015). Heat map shown in Fig. 3B indicated the evolution of each genus selected in different treatments. PCA analysis based on these genus showed the changes of microbial community, which indicated the significant difference between mono-SS and
co-digestion of FW and SS based on MW (Fig. 4). Changes of bacterial community could be divided into two phases based on PCA analysis: days 0–5, acidification phase corresponding to accumulation of VFAs and days 5–33, active methane production phase corresponding to consumption of VFAs (Fig. 2). The greatest changes happened at acidification phase for all the treatments, and so the difference between mono-SS and co-digestion based on MW-FW and MW-SS did. Proteiniborus and Parabacteroides were the dominant genus for the VFAs production in all the tests, while Petrimonas was only dominant at the acidification phase for mono-SS. Correspondingly, Bacteroides which changed little for mono-SS increased significantly and dominated on day 5 for co-digestion. After acidification phase, most of the dominant genus decreased along with anaerobic digestion.

Proteiniborus belonging to phylum Bacteroidetes is a kind of anaerobic, mesophilic and protein-specific utilizing bacteria, and the fermentation products mainly include ethanol, acetic acid, hydrogen, carbon dioxide and a trace amount of propionic acid (Niu et al., 2008). It is easy to understand its dominance for anaerobic digestion of SS which was mainly composed of proteins, and it is the first abundant genus for mono-SS (18.81%) and MW-SS (23.64%) anaerobic digestion. Meanwhile, its abundance was the lowest (5.52%) for MW-FW, and this was because MW-SS could help release proteins in bacteria. Parabacteroides also belonging to Bacteroidetes is saccharolytic and produces acetate and succinate as its primary fermentation end products (Tan et al., 2012). That is to say that members of Parabacteroides are responsible for the degradation of polysaccharides in all the treatments. Thus, its abundance was higher in co-digestion due to the addition of FW whose main components were polysaccharides. Petrimonas is described as a mesophilic, strictly anaerobic and fermentative bacterium, and acetic acid was the major end products (Grabowski et al., 2005). It contributed more to VFAs accumulation of mono-SS than that of MW-SS and MW-FW, because its abundance increased significantly on day 12 for MW-SS (14.59%) and MW-FW (9.26%), 7 days later than mono-SS. It is interesting to notice that Bacteroides only dominated for the co-digestion. Members of genus Bacteroides were isolated from various environments, and most of them produce acetic acid, propionate acid, formate and succinic acid as the major end-products of fermentation (Hatamoto et al., 2014). Especially for MW-FW, its abundance accounted for 32.66%, which indicated that it contributed most to the VFAs accumulation of MW-FW.

Besides, other microbes only increased during anaerobic digestion of mono-SS were revealed by PCA analysis (Fig. 4) such as Janibacter, Comamonas, Luteimonas. These genus only increased in mono-SS, while decreased or changed little for the co-digestion based on MW pretreatment. But their specific function for anaerobic digestion is still unclear.
3.3.3. Evolution of archaeal community

Although primers of 16S rRNA V4 targeting both bacterial and archaeal community were used, lower abundance of archaeal community (ranging 0.29–1.01% shown in bacterial community analysis at phylum) made it impossible to analyze the composition of archaea in detail. Thus, a total of 479,911 high quality reads ranging from 30,124 to 34,910 were acquired for detailed archaeal community analysis. According to RDP classifier, most of these reads were assigned to methanogenic archaea. Compared to bacterial community, the diversity of archaeal community was very simple, while significant changes happened along with anaerobic digestion as shown in Fig. 5. The acetoclastic Methanosaeta dominated on day 0 accounting for 81.23%, 52.9% and 70.53% for mono-SS, MW-SS and MW-FW, respectively, while its abundance decreased significantly along with anaerobic digestion and the final abundance was 6.98%, 8.14%, and 5.7%, respectively. Meanwhile, another acetoclastic methanogen, Methanosarcina, increased significantly at the active methane production phase (day 12) for all the treatments. The maximum abundance at this phase was 30.09%, 49.92% and 39.78%, that is, increased by 28.66, 97.88 and 44.64%, while its initial abundance was ND, 0.57% and 0.49% for mono-SS, MW-SS and MW-FW, respectively.

The most interesting phenomenon appeared during anaerobic co-digestion of MW-FW, that is, genus Methanosphaera dominated on day 5 (from 0.84% to 40.76%), and its evolution was quite special for MW-FW and subsequently decreased gradually along with anaerobic digestion to the end (3.83%) (Fig. 5). Members of genus Methanosphaera have one of the most restricted energy metabolisms of all methanogens because it can neither oxidize methanol to CO₂ nor reduce CO₂ to methane (Fricke et al., 2006). Instead, it is dependent on acetate as a main carbon source for growth and uses H₂ to reduce methanol to produce methane (Fricke et al., 2006). Previous study have demonstrated that high levels of methanol and acetate would provide an ideal environment for Methanosphaera to thrive (Facey et al., 2012), and this indicated that MW pretreatment of FW may produce amounts of methanol.

4. Conclusions

Anaerobic co-digestion of FW and SS was generally considered effective for methane production. The effects of MW for the co-digestion have never been investigated. Results showed that the optimized ratio of FW and SS based on MW pretreatment was 3:2 (TS), no matter FW or SS was MW pretreatment, and MW-FW was less efficient for enhancing the methane production than MW-SS due to the accumulation of propionic acid. Co-digestion of FW and SS based on MW changed the composition of bacterial and archaeal community significantly, and the
evolution of them was studied in detail through high-throughput sequencing method.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.biortech.2015.10.037.

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