Profiles and drivers of antibiotic resistance genes distribution in one-stage and two-stage sludge anaerobic digestion based on microwave-H$_2$O$_2$ pretreatment

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HIGHLIGHTS
- Pretreatment with subsequent anaerobic digestion enhanced ARGs abundance reduction.
- Two-stage system showed some advantages over the ARGs abundance reduction.
- Anaerobic digestion was more effective on pathogens control than pretreatment.
- Changes of bacterial community was significantly correlated with ARGs profiles.
- Anaerobic digestion was a potential technology for ARGs control.

GRAPHICAL ABSTRACT

ABSTRACT
Three anaerobic digestion (AD) processes of waste activated sludge (WAS) were established including the control (mono-WAS), one-stage AD and two-stage AD along with microwave-H$_2$O$_2$ pre-treatment (MW-H$_2$O$_2$) to investigate the profiles and drivers of antibiotic resistance genes (ARGs) distribution concerning co-selection from heavy metals, intI1 and microbial community through qPCR and high-throughput sequencing method. Results showed that MW-H$_2$O$_2$ could reduce the absolute gene copies of all ARGs while increased the relative abundance of most ARGs. After subsequent AD, both total ARGs quantities and relative abundance were enriched while two-stage AD showed some advantages over ARGs abundance reduction. Besides, AD was more effective on the potential pathogens reduction than MW-H$_2$O$_2$. AD could reduce the role of intI1 on the spread of ARGs, while mantel test and procrustes analysis indicated that the variation of ARGs abundance was closely associated with the discrepancy of bacterial community.

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Abbreviations: ARGs, antibiotic resistance genes; ARB, antibiotic resistance bacteria; AD, anaerobic digestion; WAS, waste activated sludge; MW, microwave pretreatment; MW-WAS, microwave pretreatment of waste activated sludge; MW-H$_2$O$_2$, microwave pretreatment along with H$_2$O$_2$ addition; MW-Acid, the acidification phase of the two-stage anaerobic digestion; MW-CH$_4$, the methane-producing phase of the two-stage anaerobic digestion; WWTPs, wastewater treatment plants.

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1. Introduction

The rapid spread of antibiotic resistance bacteria (ARB) indicated that we were close to a point where we may not be able to prevent or treat everyday infections or diseases (United Kingdom Department of Health & DEFRA, 2013). Antibiotic resistance genes (ARGs), as the culprit of bacterial resistance to antibiotics, were gradually considered to be an emerging pollutant in the environment, and municipal wastewater treatment plants (WWTPs) become one of the most important reservoirs of ARGs (Mao et al., 2015), and antibiotic resistance factors from WWTPs have been demonstrated to be associated with clinical pathogens (Kumaraswamy et al., 2014; Su et al., 2015). Waste activated sludge (WAS), as the by-product of the WWTPs, contained high amounts of ARB and ARGs, and 6.25 million tons of dry WAS were produced in 2013 in China, and total sludge production grew by 13% annually from 2007 to 2013 (Yang et al., 2015). The spread of ARGs and ARB due to the release of the huge amounts of WAS should be emphasized (Munir et al., 2011; Singer et al., 2016).

Although antibiotic resistance could not be eradicated due to its ancient origins, it could be managed to limit the threat and minimize the impacts (United Kingdom Department of Health & DEFRA, 2013), and WAS treatment technology could be the key point (Burch et al., 2014). Anaerobic digestion (AD) was widely used for the WAS treatment since it could economically and efficiently realize the resource utilization, volume reduction and harmless of WAS (Zhang et al., 2015). However, the rate-limiting step for the AD efficiency the slow hydrolysis, and thus sludge pretreatment to enhance the hydrolysis of WAS was obviously necessary. Microwave pretreatment (MW), especially with H2O2, was widely demonstrated to be able to enhance the cell destruction of WAS, which further availed the hydrolysis of WAS in AD, and volatile solids removal, methane production and AD stabilization were simultaneously improved (Liu et al., 2015a; b; Wang et al., 2009; Kuglarz et al., 2013).

A few studies have investigated the fate of ARGs during AD of WAS, and they suggested that AD could be used to reduce ARGs abundance (Zhang et al., 2016b; Ma et al., 2011), while our previous studies further confirmed that MW-WAS further facilitated ARGs reduction (Zhang et al., 2016b; Tong et al., 2016). However, it was indicated that the concentration of ARGs in AD effluents may not be directly related to the influent concentrations, because the presence of ARGs was influenced greatly by AD operating conditions and the composition of the microbial community (Miller et al., 2013; Ma et al., 2011; Youngquist et al., 2016), that is, the operating conditions of AD may be the primary factor that governed the composition of the bacterial community and the subsequent prevalence of ARGs (Youngquist et al., 2016). Additional research would be of value to determine the optimum AD conditions for removal of ARB or ARGs and to increase the understanding of the fate of ARGs during AD. Two-stage AD with hydrolytic acidification in the first stage and with methanogenesis in the second stage could enhance the methane production compared to one-stage systems (Liu et al., 2016), but its effectiveness on the ARGs control needs to be investigated. Although Wu et al., (2016) investigated the influence of the two-phase AD of WAS on the fate of ARGs, they focused on the temperature in the different phase, while WAS pretreatment and the comparison between one-stage and two-stage systems concerning the ARGs control were unavailable.

Besides, it was also demonstrated that the co-selection from heavy metals caused by the cross- or co-resistance phenomena and horizontal gene transfer (HGT) through mobile genetic elements (MGEs) played an important role on the spread of ARGs in various environments (Pal et al., 2015; Riber et al., 2014; Xu et al., 2017). Considering the co-occurrence of significant amounts of heavy metals and antibiotics in WAS, the co-selection from heavy metals on the distribution of ARGs could not be negligible (Bondarczuk et al., 2016), while the immense biomass and intense interactions between the microorganisms provided the hotbed for the HGT of ARGs in WAS (Zhang et al., 2011). Thus, the investigation of the effects of co-selection from heavy metals and HGT on the ARGs profiles during AD would be of great importance.

In this study, three different AD processes were established including the control (mono-WAS), one-stage AD (mono-MW) and two-stage AD (MW-Acid and MW-CH4) to (1) find out the different profiles of ARGs between one-stage and two-stage sludge AD comparing with the control (mono-WAS); (2) determine the relationship of the fate of ARGs and its potential drivers including bacterial community evolution, HGT through MGEs and co-selection from heavy metals; (3) provide insight thoughts on the ARGs control during sludge AD.

2. Material and methods

2.1. Sludge anaerobic digestion process

The feed sludge was the dewatered WAS (total solids, TS, ca. 20%) from the Qinghe WWTP, and then the dewatered WAS were diluted with deionized water to the TS concentration of ca. 8%. Then, the sludge was stored at 4 °C and screened through 18-mesh sieve before use and MW-H2O2 pretreatment to remove the sands and other small inorganic matters existed in the WAS (Liu et al., 2015b; Liu et al., 2016b). The TS concentration of WAS after sieving was ca.7.6%. The detailed MW-H2O2 pre-treatment process was described elsewhere (Liu et al., 2015a). Briefly, NaOH solution at 5 M was firstly added to adjust pH of the raw sludge to 10.0, and then the sludge was heated to 80 °C by microwave irradiation for inhibiting activity of catalase which is present in aerobic living cells to avoid the decomposition of H2O2 by the catalase that exists in the sludge (Liu et al., 2016a; Wang et al., 2009). After that, H2O2 (30%, w/w) was added at a dosage of 0.2 g H2O2/g TS as suggested previously (Wang et al., 2009). Finally, the sludge was continuously heated to 100 °C with microwave irradiation operating at 600 W and ambient pressure. The pretreatment process was carried out in 1 L beakers with plastic cap but was not sealed. 300 ml of waste activated sludge in the beaker was heated at the rate of 20 °C/min to reach the target temperature, not holding at 80 or 100 °C.

Three mesophilic sludge AD reactors were established in 2-L glass bottles with 1.8 L of effective working volume and equipped with a motor and stirring paddle each. All of the reactors operated semi-continuously, and the activated sludge or pretreated sludge with TS of approximately 8% was fed into the reactors once per day as previously reported (Liu et al., 2016a). Briefly, one was fed with WAS without pretreatment as the control (mono-WAS). Another was fed with a mixture of MW-H2O2 pretreated sludge and raw sludge (Mass_pretreated_sludge/Mass_raw_sludge = 1/1). Additionally, a two-stage reactor was set up with a 0.65 L bottle (the first stage reactor) in series with a 2 L bottle (the second stage reactor). The first stage reactor (MW-Acid) was fed with the mixture of MW-H2O2 pretreated sludge and raw sludge, and the discharged sludge was fed into the second stage reactor (MW-CH4). MW-H2O2 was adopted due to its most effective on the enhancement of the release of soluble substances in waste activated sludge compared with other MW combined process (Wang et al., 2009), while two-stage sludge AD was used to reduce the inhibitory effects of H2O2 on the biogas production as suggested previously (Liu et al., 2015a), and the first stage reactor was used for the pre-acidification of the feed sludge. The three digesters were
manipulated in the same SRT and an organic loading rate (OLR) of approximately 2.92 g Volatile Solids (VS)/(L·d), and the detail anaerobic digesters operational parameters were listed in Table 1. The produced biogas was automatically recorded by an AMP11 II instrument (Bioprocess Control Company, Sweden). The ARGs and microbial community profiles were investigated during all the digesters were operated at optimal steady-state performance in terms of biogas production.

2.2. Sampling and genomic DNA extraction

Samples collected in this study included the feed sludge, pretreated sludge by MW-H2O2 and the digested sludge of the three digesters, as well as the discharged sludge from the reactor of the MW-Acid, when the reactors were in stable operation for three solid retention times (SRT), about 90 days after startup. Sampling was continuously conducted for three days, and the three samples were then mixed to get a representative sample. There were six samples in total, including feed sludge as influent (IS), pretreated sludge by MW-H2O2 (MW), the effluent of mono-WAS, mono-MW, MW-Acid and MW-CH4. 1 mL sludge sample was centrifuged at 10,000 rpm for 10 min, and the pellet was used for DNA extraction using a FastDNA SPIN Kit for Soil (MP Biomedicals, USA) according to the manufacturer’s instructions. DNA extraction of each sample was conducted in triplicate, and the triplicate DNA extracts were then merged together for further analysis. Extracted genomic DNA was detected and quantified using 1% agarose gel electrophoresis and NanoDrop 2000 (Thermo Scientific, USA), respectively, and then stored at \(-20^\circ\text{C}\) before use.

2.3. Quantitative PCR (qPCR)

Eleven frequently detected ARGs encoding resistance to \(\beta\)-lactam (bla\textsubscript{OXA-1} and bla\textsubscript{TEM}), macrolides (erm\textsubscript{A}, erm\textsubscript{B}, ermF and mef\textsubscript{A}/E); sulfonamides (sul\textsubscript{I} and sul\textsubscript{II}) and tetracyclines (tet\textsubscript{C}, tet\textsubscript{M} and tet\textsubscript{X}) were quantified. These ARGs were selected according to types of antibiotics and main resistance mechanisms (antibiotic deactivation or degradation, efflux pump and target protection) (Selvam et al., 2012). The class I integrase gene (int\textsubscript{I}) was monitored as an indicator of horizontal gene transfer (HGT) and multiple antibiotic resistance (Amos et al., 2015; Gillings et al., 2014). While the three MRGs (pco\textsubscript{A} and cop\textsubscript{A} encoding resistance to copper, c\textsubscript{C}A encoding resistance to cadmium, cobalt and zinc) represent the changes of heavy metals selective pressure during the process. Zinc and copper were selected due to its dominant presence in WAS in China (Zhang et al., 2016c) and its widely demonstrated co-selection for antibiotic resistance in various environments (Di Cesare et al., 2016; Garner et al., 2016; Xu et al., 2017). The changes of 16S rRNA gene copies were followed to represent the evolution of microbial biomass. The detail information of qPCR cycle conditions was described in Supporting information. The primers, annealing temperature and amplification efficiencies in this primer were summarized in Table S1 and S2.

2.4. High-throughput sequencing and bioinformatics analysis

In order to gain insights into the potential mechanisms driving ARGs profiles, bacterial community was studied in detail through high-throughput sequencing method. The bacterial community was investigated using the PCR primers 515F and 806R targeting the 16S rRNA V4 region, which has been demonstrated to better reveal the bacterial community composition compared to other primers (Caporaso et al., 2010; Peiffer et al., 2013). The amplicons sent for small-fragment library construction and pair-end sequencing were prepared as previously described (Zhang et al., 2016a), and the high-throughput sequencing was done by Majorbio Co., Ltd., in Shanghai using the Illumina MiSeq sequencing system (Illumina, USA).

The detail bioinformatics analysis was conducted as suggested previously (Zhang et al., 2016a). Briefly, sequencing reads were assigned to each sample according to the unique barcode, and pair-end reads were merged using FLASH (Magoc and Salzberg, 2011), and then were filtered using QIIME quality filters \((r = 3, p = 0.75, q = 3, n = 0); (r)\) max\_bad\_run\_length: maximum number of consecutive low-quality base calls allowed before truncating a read; \((p)\) min\_per\_read\_length: minimum number of consecutive high-quality base calls to retain a read (as a percentage of total read length); \((q)\) phred\_quality\_score: last quality score considered low quality; \((n)\) sequence\_max\_n: maximum number of ambiguous \((N)\) characters allowed in a sequence. PCR chimeras were filtered out using UCHIME (Edgar et al., 2011). To fairly compare the samples at the same sequencing depth, the sequence number was normalized by extracting the first 29,269 from each sample for further analysis, and then the normalized sequences were uploaded to MG-RAST (http://metagenomics.anl.gov/linkin.cgi?project=17628). The sequences were also submitted to the NCBI Sequence Read Archive (SRA) under the project number of PRJNA387350.

The taxonomic classification of the sequences was carried out using the RDP Classifier selecting the gene bacterial 16S in the RDP pipeline, and the confidence cutoff was set as 50% as suggested by the RDP (Wang et al., 2007). Potential pathogens were identified according to the virulence factor database (VFDB) at the taxonomy level of genera (Chen et al., 2016). In addition, diversity and richness indices were calculated using the Rarefaction, Shannon Index and Chao1 estimator in the pipeline.

2.5. Data analysis

The absolute copy numbers indicated the relevant gene copies per unit of dry weight (DW), while the relative abundance was calculated through the normalized gene copies by 16S rRNA. The significance of the correlation between ARGs, MRGs and int\textsubscript{I} was checked by SPSS 21.0 (IBM, USA), and a p value <0.05 was considered statistically significant. Principal component analysis (PCA), redundancy analysis (RDA) and procrustes analysis were conducted by Canoco 5.0 (Microcomputer Power, USA), and mantel test was done through PAST 3.07 (Hammer et al., 2001). The heat maps in this study were built by Heml 1.0 (Deng et al., 2014).

| Process         | SRT (d) | Feed Description | T (°C) | Working Volume | TS (g) | VS (g) | OLR (g VS/(L·d)) |
|-----------------|---------|------------------|--------|----------------|--------|--------|-----------------|
| Mono-WAS       | 20      | Untreated sludge  | 37      | 1.8 L          | 6.00 ± 0.04% | 3.73 ± 0.05% | 2.89            |
| Mono-MW        | 20      | Mixture\textsuperscript{c} | 37     | 1.8 L          | 5.54 ± 0.01% | 3.16 ± 0.03% | 2.87            |
| MW-Acid        | 2       | Mixture\textsuperscript{c} | 37     | 0.18 L         | 6.70 ± 0.18% | 4.90 ± 0.17% | 28.65\textsuperscript{c} |
| MW-CH\textsubscript{4} | 18      | Discharge of MW-Acid | 37    | 1.62 L         | 5.50 ± 0.03% | 3.12 ± 0.04% | 2.72\textsuperscript{c} |

\textsuperscript{a} Mass\textsubscript{pretreated sludge}/Mass\textsubscript{raw sludge} = 1/1.

\textsuperscript{b} The VS loss during the first stage was ca. 3.6%.

\textsuperscript{c} The OLR of the whole two-stage system was 2.87 g VS/(L·d).
3. Results and discussion

3.1. Effects of MW-H2O2 pretreatment on reduction of ARGs quantities and abundance

MW-H2O2 could reduce all the ARGs quantities especially for ermB, ermF, blaOXA-1, and tetX as shown in Fig. 1. The total ARGs quantities decreased by 0.70 logs, and also the biomass reflected by the 16S rRNA gene copies was reduced by 1.22 logs. Interestingly, the change of the abundance of ARGs varied significantly (Fig. 2). MW-H2O2 increased the abundance of most ARGs except ermB and tetX, and the mefA/E increased the most in abundance by 16.4 times. The total ARGs relative abundance was increased by 3.41 times.

The decrease of ARGs gene copies after MW-H2O2 (Fig. 1) was associated with the death of live microbes and destruction of their DNA due to the heating effects reflected by the reduction of microbial biomass, while the increase of ARGs relative abundance (Fig. 2) indicated that ARGs may resist the degradation caused by the MW-H2O2, and it seemed like that host bacteria of ARGs were more prone to surviving from MW-H2O2. The dominant ARG was sulI (8.84 logs) in IS, and blaOXA-1, the least (3.98 logs), while MW-H2O2 did not change the composition of ARGs significantly.

The gene copies and abundance of intI1 both increased a lot after MW-H2O2, and intI1 was the only target gene which not only increased in quantities by 0.18 logs but also increased the most in abundance by 25.4 times. This indicated the higher survival of class 1 integrons, which could explain its dominance in various environments (Gaze et al., 2011). As for MRGs, czcA was the dominant MRG (8.04 logs) followed by copA (6.88 logs) and pcoA (6.57 logs), and MW-H2O2 could reduce MRGs by 0.64–0.89 logs. However, the abundance of MRGs all increased significantly after MW-H2O2 by 2.19–3.85 times with czcA increasing the most.

3.2. Changes of absolute gene copies of ARGs during subsequent AD

The microbial biomass increased significantly in all the subsequent AD, along with MW-Acid increasing the most followed by MW-CH4, mono-WAS and mono-MW (Fig. 1). Total ARGs quantities showed similar changes to microbial biomass, and there was significantly positive correlation between total ARGs and microbial biomass (p < 0.05). This indicated that the proliferation of microbes during AD inevitably led to the increase of ARGs quantities as suggested previously in various environments (Zhang et al., 2016a, b; Burch et al., 2013). Concerning each ARG quantities, there was some difference between the different digesters. The blaOXA-1 decreased to ND in both MW-CH4 and mono-MW, while it increased to 4.62 logs in mono-WAS similar to MW-Acid. This may indicate that the host bacteria of blaOXA-1 mainly belonged to the acid-forming bacteria. ARGs quantities were enriched by 1.05–4.95 logs after subsequent AD (Fig. 3), and two-stage sludge AD showed the least enrichment of blaTEM, sulI, sulII, tetM and tetX; one-stage AD, ereA, ermF and tetG; mono-WAS, ermB and mefA/E (Fig. 1). The ermB (4.23–4.95 logs) followed by ermA (3.08–3.83 logs) were enriched the most in each digester, while sulI (1.04–1.37 logs) and tetG (1.10–1.40 logs) increased the least. The dominant ARGs also changed significantly, sulI dominated in the feed, and its dominance did not change after MW. However, ermB (8.68–9.40 logs) and ermA (9.89–10.63 logs) dominated after AD (Fig. 1). This indicated that physical effects could not change the composition of ARGs much, and it may be the changes of microbial community caused by the environmental conditions that at last changed the composition.

The intI1 and MRGs gene copies also increased significantly after AD. The intI1 increased by 2.21–2.55 logs with two-stage sludge anaerobic digestion enriching the least followed by mono-MW and mono-WAS (Fig. 1). The czcA was still the dominant MRG, and it was enriched the most in all digesters after AD.
Two-stage sludge AD enriched the least (0.25–0.99 logs) MRGs (Fig. 1). Together, MW-H2O2 subsequent AD increased all the target genes due to the increase of microbial biomass as reflected by 16S rRNA gene copies. Two-stage sludge AD showed slight advantage on ARGs quantities control due to the less enrichment of intI1, MRGs and more kinds of ARGs compared with mono-MW.

Fig. 2. Changes of the relative abundance of ARGs after MW-H2O2 pretreatment and subsequent anaerobic digestion. IS: Influent sludge; MW: Sludge after microwave pretreatment; MW-Acid: The acidification phase of the two-stage AD; MW-CH4: The methane-producing phase of the two-stage AD; mono-WAS: AD of WAS without pretreatment; mono-MW: AD of WAS along with MW-H2O2 pretreatment.

Fig. 3. Changes of absolute gene copies and abundance of total ARGs in different operational parameters of sludge anaerobic digestion. IS: Influent sludge; MW: Sludge after microwave pretreatment; MW-Acid: The acidification phase of the two-stage AD; MW-CH4: The methane-producing phase of the two-stage AD; mono-WAS: AD of WAS without pretreatment; mono-MW: AD of WAS along with MW-H2O2 pretreatment.
3.3. Reduction of the abundance of ARGs during subsequent AD

The Normalizing ARGs gene copies to 16S rRNA genes copies, as indicated by the relative abundance, could not only correct for the variations in extraction efficiencies and analytical efficiencies but also provide an indicator of the proportion of bacteria carrying ARGs (Knapp et al., 2010; McKinney et al., 2010), and it was crucial to determine the changes of relative abundance of ARGs during AD of WAS. Total ARGs abundance was further enriched slightly (1.64 and 1.69 times for two-stage AD and one-stage AD, respectively) along with mono-WAS increased the most (3.12 times). This indicated that MW-H2O2 pre-treatment availed the ARGs control in WAS. Total ARGs abundance was further enriched slightly (1.64 times) for two-stage AD and one-stage AD, respectively. This may indicate that two-stage could reduce some risks from ARGs, and mono-WAS without MW-H2O2 showed the highest risks concerning HGT. Each MRG abundance was decreased a lot, and this was comparable with previous study, which indicated that MRGs could be reduced due to the passivation of heavy metals in AD (Zhang et al., 2016b).

Above all, the relative abundance of ARGs was reduced after AD in all the digesters. Some ARGs relative abundance including bltTEM, sull, sulII and tetG in all the digesters, while ereA, tetM and tetX only in two-stage AD. Two-stage AD could reduce more of ARGs abundance. The ermB, ermF and mefa/E were enriched in all the digesters, and ermB was enriched the most especially in two-stage AD (439.13 times) followed by ermF (33.46 times). The significant increase of ermB, ermF and mefa/E during sludge anaerobic digestion has been elucidated previously (Zhang et al., 2016b). The reason may be that host bacteria of these ARGs functioned necessarily during AD, which indicated that the enrichment of these ARGs just elucidated the special proliferation of the host bacteria. Previous study suggested that antibiotics in WAS should be not enough to select for ARB and pose the selective pressure for the balance of the fitness cost of ARBs (Youngquist et al., 2016). Sludge AD also supported that ARBs could be reduced effectively, while ARGs increased significantly after AD (Tong et al., 2016). We speculated that host bacteria may not function as ARB at all, instead, its primary function was for biogas production, that is, although, ARGs like ermB, ermF and mefa/E were enriched, the risks of these ARGs may be reduced after AD. As for the highest ARGs concentrations in MW-Acid, it was attributed that the fast proliferation for the acid-production microbes compared to the methogens, and this has been proved by the highest 16 s rRNA gene copies in the MW-Acid. The highest abundance of ARGs in the MW-Acid could be because of that the host of ARGs could also function as the important acid-production microbes, and the accumulation of ARGs, especially in the extracellular DNA, could not be overlooked.

Mobile genetic elements like Class I integrons (intI1), were often used to represent the HGT and multiple antibiotic resistance (Dang et al., 2017). Although they cannot mobilize and transfer themselves between microbes, they are often associated with mobile genetic elements which can, such as conjugative plasmids and insertion sequences (Berglund, 2015). Thus, intI1 was suggested as the proxy for anthropogenic pollution of ARGs (Gillings et al., 2014). Two-stage AD could reduce the abundance of intI1, while mono-MW and mono-WAS enriched it by 2.06 and 55.60 times, respectively. This may indicate that two-stage could reduce some risks from ARGs, and mono-WAS without MW-H2O2 showed the highest risks concerning HGT. Each ARGs abundance was decreased a lot, and this was comparable with previous study, which indicated that MRGs could be reduced due to the passivation of heavy metals in AD (Zhang et al., 2016b).

3.4. Bacterial community and potential pathogens

According to the diversity and richness analysis, the diversity of MW-Acid was the lowest, and MW-ChA < mono-MW < mono-WAS. Two-stage AD showed more specificity compared with mono-WAS and mono-MW. The changes of bacterial community at the phylum level were shown in Fig. 4A. It showed that phylum Proteobacteria dominated in the feed, while Bacteroidetes dominated after AD, and this corresponded to the sequential dominate of sulf and ermF. MW-H2O2 changed the bacterial community little, the dominant phylum was Proteobacteria (59.2%) in the feed sludge followed by Bacteroidetes (14.8%). After MW-H2O2, the dominant phylum was the same, and the abundance of Proteobacteria and Bacteroidetes were 57.2% and 14.3%, respectively. The little change in microbial composition could explain the small changes of prevalence ARGs in sludge, while Firmicutes increased from 4.6% to 9.7% and Actinobacteria from 4.9% to 7.8%.

The dominant phylum changed significantly after AD as shown in Fig. 4A. Bacteroidetes become dominant in the three AD digesters accounting for 78.1%, 51.1% and 39.9% in two-stage AD, one-stage AD and mono-WAS, respectively. Meanwhile, Proteobacteria...
decreased to 7.3%, 7.9% and 13.8%, respectively. The significant increase of Bacteroidetes abundance might explain the increase of \textit{ermB} and \textit{ermF} abundance. It was speculated that the host bacteria of \textit{ermB} and \textit{ermF} which increased the most concerning the abundance mainly existed in Bacteroidetes (Liu and Pop, 2009), while the host bacteria of \textit{sulI}, \textit{sulII}, \textit{bla}_{\text{TEM}} and \textit{tetG} mainly existed in Proteobacteria, and the reduction of these ARGs abundance might be associated with the changes of bacterial community. Firmicutes changed little after AD, except in mono-MW (23.3%). Also Tenericutes (26.5%) was only dominant in MW-Acid.

In order to elucidate the changes of bacterial community in detail, the first top ten genera, the genera abundance above 1% and potential pathogens were selected in each sample (Fig. 4B). Acinetobacteria belonging to Proteobacteria was the dominant genera in both IS (13.4%) and MW (8.4%), and it has been demonstrated to be the host bacteria of over 41 kinds of ARGs including \textit{sulI}, \textit{bla}_{\text{TEM}}, \textit{tetG} and \textit{tetM} (Liu and Pop, 2009). It was speculated that the dominance of \textit{sulI} may be associated with Acinetobacteria. While Acinetobacteria decreased to below 0.01% in all the digesters after AD, corresponding the reduction of the abundance of \textit{sulI}, \textit{bla}_{\text{TEM}}, \textit{tetG} and \textit{tetM}. Concerning the dominant genera after AD, unfortunately, 69.4%–83.1% sequences could not be classified to the genus due to the limited dataset and technology compared with IS (23.8%) and MW (35.4%), and the unclassified sequences mainly belonged to the phylum of Bacteroidetes. Petrimonas (3.5%), Cloacamonas (10.3%) and Syntrophomonas (4.1%) were the identified dominant genus in MW-CH4, mono-WAS and mono-MW, respectively. As shown in Fig. 5, MW-\textit{H}_{2}\text{O}_2 changed the bacterial community little as demonstrated by PcoA, while the difference of the bacterial community after AD between different digesters was distinct (Fig. 5).

According to the VFDB, 12 genera were classified to the potential pathogens in the feed sludge (Fig. 4B). MW-\textit{H}_{2}\text{O}_2 could reduce them to some extent, while AD could reduce them more effectively. For instance, the dominant potential pathogens in IS were \textit{Acinetobacter} and \textit{Pseudomonas}, and \textit{MW-\textit{H}}\textit{_2}\text{O}_2 decreased the abundance from 13.4% to 8.4% and 8.2% to 1.1%, respectively, while AD could reduce them to below 0.01%. However, some potential pathogens rebound after AD, like \textit{Clostridium} and \textit{Mycobacterium}, and \textit{MW-\textit{H}}\textit{_2}\text{O}_2 could reduce them from 0.51% and 0.26% to 0.24% and 0.05%, respectively, while they both rebound to 0.74%–0.86% and 0.21%–0.3%, respectively. This indicated that these genera should be paid more attention after AD.

3.5. Relationship among ARGs, bacterial community, \textit{intI1} and MRGs

In order to figure out the relationship between evolution of ARGs and changes of bacterial community, procrustes analysis was conducted as suggested previously (Zhang et al., 2016a,b). The results indicated there existed significantly positive correlation between them (Fig. 5). Changes of bacterial community accounted for the 86.6% of the variation of ARGs abundance, and the correlation between the two axes was significantly positive (R = 0.9750 and 0.9436, respectively). A Mantel test based on Bray–Curtis distance confirmed the significantly positive correlation between them (\textit{R} = 0.7436, \textit{p} = 0.0001, permutation \textit{N} = 9999). This indicated that bacterial community contributed significantly to the evolution of ARGs during these treatments. This kind of relationship between ARGs and bacterial community has been established in various environments (Li et al., 2015; Zhang et al., 2016b).

The class I integron plays an important role in gene transfer among bacteria. There is no significantly positive correlation between \textit{intI1} and ARGs increased except \textit{ereA} (\textit{p} = 0.042) (Table S3). This may indicate that the role of mobile genetic elements on the occurrence of ARGs during AD was limited. MRGs could reflect the real response of bacteria to the selective pressures of heavy metals better compared with the concentration of heavy metals. There was significantly positive correlation between \textit{sulI}, \textit{sulII}, \textit{tetG} and \textit{bla}_{\text{TEM}} and MRGs which decreased significantly after AD. This may indicate that the reduction of heavy metals selective pressure facilitate these ARGs reduction during AD, which was favoured previously in AD (Zhang et al., 2016b).

RDA analysis was conducted to investigate the relationships between bacterial community, \textit{intI1} and MRGs and ARGs (Fig. 6), and the results showed that the selected variables accounted for 94.9% of the total variation of the evolution of ARGs quantified in this study. The primary contributors for the different ARGs profiles in different stages varied significantly (Fig. 6). Firmicutes and \textit{intI1} mainly accounted for the evolution of the ARGs including \textit{tetX}, \textit{mefA/E}, \textit{tetM} and \textit{ereA}, while \textit{Bacteroidetes} mainly accounted for the evolution of \textit{ermB} and \textit{ermF}, and MRGs were associated with the abundance reduction of \textit{sulI}, \textit{sulII}, \textit{tetG} and \textit{bla}_{\text{TEM}}. Besides, there existed significantly positive correlations between potential pathogens including \textit{Acinetobacter}, \textit{Aeromonas}, \textit{Enterococcus} and \textit{Pseudomonas} and ARGs which decreased including \textit{sulI}, \textit{sulII}, \textit{tetG} and \textit{bla}_{\text{TEM}}. This indicated that the pathogens reduction due to AD contributed to the ARGs reduction. However, the potential pathogens including \textit{Clostridium} and \textit{Mycobacterium} which increased after AD should be paid more attention, because they have been demonstrated to host bacteria of the 23 and 8 resistance gene types, respectively, including \textit{ermB}, \textit{ermF} and \textit{mefA/E} which increased significantly after AD in this study.

It looks like that ARGs after AD was only associated with the operation parameters not the feed sludge. The results suggested that bacterial community composition of the sludge digestion process drives the profiles of ARGs, rather than the influent ARG composition. The role of \textit{intI1} may be also limited during AD especially...
in two-stage AD, and the evolution of ARGs was mainly associated with the changes of bacterial community. Concerning the risks of ARGs, we should not only consider the amount of ARGs but also the role of MGEs and co-selection from heavy metals, and from the respective of this, AD was a potential technology for the ARGs control during sludge treatment. Besides, it was demonstrated that thermophilic system outperform than mesophilic system for ARGs removal, and the role of temperature on the ARGs profiles and drivers of the one-stage and two-stage systems deserve further investigation.

4. Conclusions

The potential of sludge AD for ARGs control through MW-H2O2 was investigated in different operational parameters. MW-H2O2 along with subsequent AD could realize some control on ARGs abundance. Two-stage AD showed some advantages on the ARGs abundance reduction and enriched the least ARGs quantities. MW-H2O2 could reduce gene copies of all ARGs while increased the abundance of most ARGs. AD enriched both total ARGs gene copies and abundance while ARGs associated with MGEs were reduced. AD could reduce the role of intI1 on the spread of ARGs, and the ARGs proliferation was closely related with the evolution of microbial community.

Conflict of interest

The authors declare no conflict of interest.

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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.2017.05.157.

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