Feasibility of short-term fermentation for short-chain fatty acids production from waste activated sludge at initial pH10: Role and significance of rhamnolipid

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Short-term fermentation for SCFAs production from WAS had been proposed. RL pretreatment enhanced WAS hydrolysis/acidification under alkaline conditions. Role and significance of EPSs had been investigated. The cumulative SCFAs were better fitted with pseudo-first-order kinetic model. RL + initial pH 10 caused positive synergies on anaerobic fermentation process.

1. Introduction

The enhanced biological nutrients removal system has been widely employed to treat wastewater all over the world. However, the available biodegradable chemical oxygen demand (COD) contained in influent, cannot meet the carbon source (CS) demand for both denitrifying bacteria and phosphorous accumulating organisms [1,2]. It is unambiguous that the chemically synthesized CS (e.g., acetate, propionate and glucose) serving as the external CS is neither cost-effective nor sustainable in full scale wastewater treatment plant (WWTP) [3,4]. Recently, some literatures have reported the feasibility of fermentation liquid (enriched in short-chain fatty acids (SCFAs)) from waste activated sludge (WAS) as the external CS for nutrients removal, and satisfactory results were achieved [3–6].

ABSTRACT

Short-chain fatty acids (SCFAs), preferred carbon source for enhanced biological nutrients removal system, can be produced from anaerobic fermentation of waste activated sludge (WAS). Hydrolysis is known as the rate-limiting step for SCFAs production. This study presents a novel technology using rhamnolipid (RL) pretreatment connected alkaline anaerobic fermentation to enhance SCFAs production. Experiment results showed that the integration treatments performed significant enhancement of SCFAs production, and the fermentation time was remarkably shortened. The maximum SCFAs production, 378 mg COD/g VSS (volatile suspended solid), reached at 72-h fermentation time under 0.2 g RL/g TSS (total suspended solid) and initial pH 10 treatment, which was 4.31, 1.32 and 1.24 times higher than that of control, initial pH 10 and 0.2 g RL/g TSS treatment, respectively. The mechanism study showed that integration treatments could enhance the release of constituents from the cells and/or extracellular polymeric substances to suspension, and hydrolysis rate constant was improved greatly in integration treatment, 8.68, 2.14 and 1.55 times higher than that of control, sole RL and initial pH 10 treatments, respectively. The results suggested that positive synergy led to improvement of WAS solubilization and SCFAs production under integration conditions. The pseudo-first-order model was successfully built to present the SCFAs accumulation. Therefore, the integration method in this work was a promising technology for SCFAs production enhancement from WAS.
As the by-product of WWTPs, 6.25 million tons dry sludge has been produced in China in 2013 and is still showing a rapid increasing rate [7]. The WAS disposal has been considered as a main issue for the sustainable development of WWTPs, because the cost of efficient WAS disposal is very high, accounting for approximately 40–60% operation fee of WWTPs [7–9]. As well known, the main component of WAS is organic matter (e.g., proteins, polysaccharides and lipid), which makes it be a potential substrate for SCFAs production [10,11]. Currently, many studies have proved the feasibility of SCFAs production from WAS with anaerobic digestion process [11–18]. Generally, anaerobic digestion process of particulate organic matters (POMs) usually includes three stages: hydrolysis, acidification and methanogenesis [19,20]. Hydrolysis is known as the rate-limiting step, only little of total COD (TCOD) of WAS can be biodegraded unless the POMs are significantly solubilized [21–24]. Aiming to strengthen the performance of anaerobic digestion of WAS, some efforts have been developed, such as chemical, mechanical, biological, thermal co-fermentation treatments [7,19,20,24–26]. Alkaline, one of chemical pretreatment, has been selected as a potential way for improving SCFAs production [7,17,24,27]. On one hand, this method can not only enhance the performance of WAS hydrolysis but also inhibit the activities of methanogens, which results in positive effects on SCFAs production [17,27]. On the other hand, it is cost–effective and operation–convenient, which is of great strategic significance. However, though the SCFAs production can be improved at pH 10, more than 60% of volatile suspended solids (VSS) could not be effectively degraded by microorganisms [27,28]. Thus, it is necessary to couple alkaline with other methods to further harvest improvement of SCFAs production from WAS.

Recently, some researchers have reported that bio-surfactant can accelerate the rate of sludge hydrolysis, and benefit to SCFAs production by improving POMs dissolution and inhibiting methanogenesis [12,13,21]. Though the chemically synthesized surfactants (e.g., sodium dodecyl sulfate and sodium dodecyl benzenesulfonate) can improve the efficiency of anaerobic fermentation of WAS, their addition and residual present a potential risk to the ecosystem because of biotoxicity in an accumulated concentration [12,21]. Rhamnolipid (RL), as a kind of bio-surfactants, is eco-friendly, this makes it a desirable replacement of traditionally chemical surfactants. It has been proved that RL can be produced in situ by some microorganisms, such as Pseudomonas, during WAS anaerobic fermentation [12,21]. And an external addition of RL can trigger this self-synthesis process. In our previous study, we have found that RL pretreatment effectively enhanced SCFAs production, meanwhile, RL concentration in liquid phase at 96-h fermentation time was 1.49-fold higher than that at the beginning fermentation time, which proved that RL generated in situ, and further contributed to WAS hydrolysis and acidification [21]. Thus, RL pretreatment method is a promising technology for WAS treatment because it is eco-friendly, renewable and cost–effective, and may be coupled with alkaline method successfully for further enhancing WAS dissolution and acidification.

The main objective of this study is to evaluate the feasibility of RL pretreatment integrated into alkaline anaerobic fermentation for enhancing SCFAs production from WAS. The mechanisms of proposed method for enhancing SCFAs production from WAS were explored by analyzing extracellular polymeric substances (EPSs) and kinetic models of WAS hydrolysis. Meanwhile, the mathematical models were also built to better understand the kinetics of SCFAs accumulation. The outcome of this work may have potential application to solve the CS shortage of a WWTP by a cost-effective and renewable way.

2. Materials and methods

2.1. Source of WAS and bio-surfactant RL

The WAS was taken from the secondary sedimentation tank of Taiping Municipal Wastewater Treatment Plant, running with anaerobic–anoxic–aerobic (A2/O) system, Harbin, China. The WAS firstly was thickened by gravitational sedimentation for 24 h at 4 °C, then screened with a 1 mm sieve to remove impurities, finally stored at 4 °C prior for later use and test. And the main characteristics of WAS were: pH 6.51 ± 0.02, total suspended solid (TSS) 23.577 ± 0.487 g/L, VSS 14.425 ± 0.428 g/L, total COD (TCOD) 22.711 ± 500 mg/L, soluble COD (SCOD) 231 ± 34 mg/L, soluble proteins 114 ± 15 mg COD/L, soluble polysaccharides 41 ± 8 mg COD/L, and SCFAs 86 ± 22 mg COD/L.

The bio-surfactant RL was purchased from Victex Company, China, and the purity was 80%, which was a blend of RhaC10G10 (C26H48O9, m/z 503) and RhaRhaG10 (C32H58O13, m/z 649), shown in Fig. S1.

2.2. Batch experiments for SCFAs production

To investigate the effects of the sole RL and RL and initial pH = 10 on SCFAs production from WAS, two groups of experiments were carried out (Table 1). Group 1 was to investigate the effects of sole RL levels on SCFAs production. And group 2 was employed to explore the effects of RL pretreatment integrated into alkaline anaerobic fermentation on SCFAs production. The demanded pH achieved by adding 4 M NaOH and 1 M HCl. Batch laboratory-scale anaerobic fermentation experiments were conducted in 500 mL serum bottles filled with 300 mL raw WAS each. Nitrogen gas was introduced to the reactors for 10 min to remove oxygen, then the reactors were capped, sealed, and stirred in an air-bath shaker (100 rpm) at 35 ± 1 °C for 5 days. All the fermentation experiments were carried out in triplicate.

2.3. EPSs extraction

Microbial EPSs are the main components of the WAS floc matrix [29,30], and their presence are of great importance for maintaining the functional integrity and strength of sludge as well as biodegradability [31], so the changes of their characteristics are of great concern to understand the effects of pretreatment methods on WAS hydrolysis and further SCFAs production. The EPSs had been divided into two parts, including loosely bound EPSs (LB-EPSs) and tightly bound EPSs (TB-EPSs) [29,32]. The extraction methods of LB-EPSs and TB-EPSs referenced from the previous literatures [29,32], and were modified appropriately. The specific method was as follows: firstly, 10 mL samples were centrifuged at 4000 g for 10 min, the supernatant was filtered with 0.45 μm cellulose nitrate membrane filters, and the filtrate was considered

| Group | No. | RL additions (g/g TSS) | pH |
|-------|-----|------------------------|----|
| 1. Effects of RL addition levels | 1 | 0 | 6.51 ± 0.02 |
| | 2 | 0.005 | 6.51 ± 0.02 |
| | 3 | 0.05 | 6.51 ± 0.02 |
| | 4 | 0.1 | 6.51 ± 0.02 |
| | 5 | 0.2 | 6.51 ± 0.02 |
| 2. Effects of RL and initial pH = 10 | 6 | 0 | 10.08 ± 0.04 |
| | 7 | 0.005 | 10.03 ± 0.04 |
| | 8 | 0.05 | 10.02 ± 0.01 |
| | 9 | 0.1 | 10.03 ± 0.02 |
| | 10 | 0.2 | 10.03 ± 0.01 |

Table 1 Batch experiments for SCFAs production from WAS.
as the dissolved organic matters (DOMs). Then, the residue was treated by the EPSs extraction method for LB-EPSs [29], and the filtrate was served as the LB-EPSs. Finally, after LB-EPSs extraction, the residue was treated according to the EPSs extraction method for TB-EPSs [29], and the filtrate was regarded as the TB-EPSs.

2.4. Analytical methods

Sludge samples collected from reactors were centrifuged at 10,000 rpm for 10 min, then supernatant samples were filtered by 0.45 μm cellulose nitrate membrane filters and finally filtrated samples were stored at 4 °C prior to analysis. The filtrate was immediately used to analyze SCFAs, polysaccharides, and proteins. The measurements of SCFAs, SCOD, TCOD, polysaccharides, proteins, TSS, VSS and pH were the same as the methods mentioned in our previous publications [11,14,21,33]. The SCFAs were regarded as the sum of acetic (HAc), propionic (HPr), n-butyric (n-HBu), iso-butyric (iso-HBu), n-valeric (n-HVa), and iso-valeric acids (iso-HVa) [11,33,34].

Three-dimensional fluorescence excitation-emission matrix (EEM) spectroscopy (FP-6500, Jasco, Tokyo, Japan) was applied to characterize the DOMs, LB-EPSs, and TB-EPSs extracted from raw sludge, and WAS after sole RL = 0.1 g/g TSS, initial pH = 10 and initial pH = 10 and 0.1 g RL/g TSS treatments. Details of spectra scan, elimination of inner filtering effect and Raman scattering, and parallel factor analysis (PARAFAC) were used to model EEM fluorescence data, could be found in our previous publication [11,14,33].

2.5. Kinetic modeling

Two kinetic models (pseudo-first-order (PFO), pseudo-second-order (PSO)) were applied to understand both the effects of fermentation time and kinetic behavior on WAS hydrolysis and SCFAs production. The PFO model assumed that the rate of WAS hydrolysis and SCFAs accumulation with fermentation time was directly proportional to the difference between the relative equilibrium capacity of SCOD/SCFAs and the amount of SCOD/SCFAs at any time t, respectively. The PFO model could be determined by the following equation [35]:

\[
\ln (q_t - q_{eq}) = \ln q_{eq} - kt
\]

where \( q_t \) (mg COD/L) and \( q_{eq} \) (mg COD/L) were the amount of SCOD/SCFAs at time t (h), \( q_{eq} \) (mg COD/L) were the relative equilibrium capacity of SCOD/SCFAs, and \( K \) (h\(^{-1}\)) was the rate constant of PFO model. As the accumulation of SCOD/SCFAs depended on their rates of production and consumption, here, the specific equilibrium capacity of SCOD/SCFAs were defined as the increased SCOD/SCFAs, caused by the hydrolysis/acidification of particulate organic matters, were approximately equal to the decreased SCOD/SCFAs, caused by biodegradation such as gasification [36].

The PSO model assumed that the SCFAs accumulation rate was proportional to the square of the driving force [37]. The PSO model could be expressed by the following equation [35]:

\[
\frac{1}{q_t} - \frac{1}{q_{eq}} = \frac{t}{k_s q_{eq}^2} + \frac{1}{q_{eq}}
\]

where \( k_s \) was the rate constant of PSO model, L/h (mg COD h). The normalized standard deviation (NSD, \( \Delta q \) (%)) and the average relative error (ARE (%)), calculated by the following Eqs. (3) and (4), were employed to evaluate the error of both PFO and PSO models [38,39]:

\[
\text{ARE} (%) = 100 \frac{1}{N-1} \sum_{i=1}^{N} \left( \frac{q_{i,\text{exp}} - q_{i,\text{cal}}}{q_{i,\text{exp}}} \right)^2
\]

where \( q_{i,\text{exp}} \) and \( q_{i,\text{cal}} \) (mg COD/L) were the experimental and calculated values, respectively, at time 't' and N was the number of measurements made.

3. Results and discussion

3.1. Performance of SCFAs production from WAS

3.1.1. Effect of different RL levels on SCFAs production

SCFAs accumulations under different RL dosages are shown in Fig. S2, it was clear that the control test achieved the minimum SCFAs production at equilibrium time, 1263 mg COD/L (i.e., 88 mg COD/g VSS). By contrast, the corresponding equilibrium SCFAs production increased with the RL dosage increasing (from 0.025 g/g TSS to 0.2 g/g TSS). When the RL addition was 0.2 g/g TSS, the maximum SCFAs production reached, 4392 mg COD/L (i.e., 304 mg COD/g VSS), which was 3.47 times higher than that of the control test. However, the SCFAs production at dosage 0.2 g/g TSS had no significant increase (p > 0.05) compared to that (4102 mg COD/L, i.e., 285 mg COD/g VSS) obtained at dosage 0.1 g/g TSS (as shown in Fig. S2), indicating that a reduced RL addition to 0.1 g/g TSS could be applied to pretreat WAS for SCFAs production. Furthermore, all the RL tests achieved the maximal SCFAs production at 96-h (i.e., 4-day) fermentation time, which was much less than that reported by previous studies, such as 8 days in the research of Yuan et al. [27] and 6 days in the research of Jiang et al. [13], suggested that short-term fermentation for SCFAs production from WAS was possible under RL pretreatment, which was beneficial to both improve the efficiency and reduce the cost of sludge treatment.

3.1.2. Enhancement of SCFAs production under combination of RL and alkaline treatments

Previous studies have reported that SCFAs production from WAS could be significantly improved by controlling the anaerobic fermentation pH at constant 10.0 [27]. The SCFAs production from WAS anaerobic fermentation had been investigated at both initial and constant 10.0 in this work. The equilibrium SCFAs production at initial pH 10 was 4130 mg COD/L (i.e., 287 mg COD/g VSS), which was 3.27 times higher than that obtained in the control test, indicating the potential application of initial pH 10 for SCFAs production from WAS. The maximum SCFAs production from WAS under a constant pH of 10 was 4758 mg COD/L (i.e., 330 mg COD/g VSS) at 10-day fermentation time, which was 1.15 times higher than that under initial pH10. Though a slight improvement of SCFAs production achieved under a constant pH of 10 compared to that of initial pH10, the fermentation time of the maximum SCFAs production reached was 3.33 times longer, which went against the high efficiency demand of WAS disposal. Moreover, a system continuously kept at pH 10 was neither economic nor convenient because of the continuous acid/alkaline adjustment. Thus, compared with the constant pH 10, initial pH 10 is relatively more efficient, economical and sustainable for SCFAs production from WAS. It was obvious that SCFAs productions were substantially promoted by RL addition, alkaline, and RL and alkaline pretreatment compared with control test (Fig. 1). Meanwhile, it was significant, while using RL to enhance the SCFAs production under initial pH 10 anaerobic fermentation. The maximum SCFAs production achieved under 0.2 g RL/g TSS integrated with initial pH 10 treatment, 5445 mg COD/L (i.e., 378 mg COD/g VSS), which was 4.31, 1.32 and 1.24 times higher than that of control, initial pH 10 and
0.2 g RL/g TSS tests, respectively. Also, the fermentation time of equilibrium SCFAs accumulations reached (72 h, i.e., 3 days) in integration tests relatively less than that using sole RL treatment, demonstrating that when the integration methods were employed, the fermentation time could be further shortened. The above results showed that the production of SCFAs could be significantly improved at initial pH 10 and further enhanced under the integration treatment, suggesting that RL pretreatment integrated into alkaline anaerobic fermentation of WAS could result in positive synergies on SCFAs production.

### 3.1.3. Composition of SCFAs under different conditions at different fermentation time

The composition of SCFAs would influence the nutrient removal efficiency, while the fermentation liquid of WAS was used as the external CS for enhanced biological nutrients removal system [11,6,40]. It was observed that with fermentation time going on, the composition was significantly affected by the dosages of RL and pH (Fig. 2). For all the tests, the two highest individual SCFAs were HAc and HPr, with a total percentage of 47.7–66.9%, in accordance with the previous studies [11,21,33]. Ordering the average percentage of individual SCFAs was HAc > HPr > iso-HVa > n-HBu > n-HVa > iso-HBu. The possible reasons was that HAc, n-HBu and HPr were formed directly from the fermentation of polysaccharides and proteins, yet the higher molecular weight SCFA such as n-HVa, were largely relevant to the fermentation of proteins [11]. And n-HBu, HPr and n-HVa were easily biodegraded to form HAc in the anaerobic fermentation system [11,27].

Moreover, RL levels had a significant influence on the ratio of HAc to HPr (Fig. 2). For sole RL treatments, under lower RL dosage (≤0.05 g/g TSS), the ratio of HAc to HPr decreased quickly with fermentation time going on. However, the ratio of HAc to HPr was kept stable in combined treatment of RL and alkaline throughout the fermentation process. It was indicated that an alkaline condition (initial pH 10) would maintain a stable composition of SCFAs, which preferred to the enhanced biological nutrients removal system while SCFAs were used as the internal CS.

### 3.2. Role and significance of EPSs under different pretreatment conditions

#### 3.2.1. Organic fractions in EPSs

As the main components of EPSs were polysaccharides, proteins, and humic acid, etc., especially polysaccharides and proteins, accounted for 50–72% of EPSs [29], changes of SCOD, soluble polysaccharides, and proteins were used to evaluate the effects of sole RL and integration pretreatments on WAS solubilization. Fig. 3 shows the changes of SCOD, soluble proteins, and polysaccharides in DOMs, LB-EPSs, and TB-EPSs under different conditions at 12-h fermentation time. It was obvious all the pretreatment methods benefitted to SCOD increase (Fig. 3(a)). Integration treatments performed better than either sole RL or initial pH = 10 treatments. In the control test, SCOD of DOMs only increased by approximately 540 mg/L. By contrast, SCOD of DOMs rose by approximately 3877 mg/L under RL dosage 0.2 g/g TSS and 3313 mg/L under initial pH = 10 treatments, which were 7.18 and 6.14 times higher than that of control test, respectively. A further increase of SCOD of DOMs was obtained in the case of combined treatment, increasing by 6199 mg/L (11.48 times higher than control test). The more release of SCOD implied that more microbial cells and/or EPSs became soluble substrates from POMs under different pretreatment conditions [11,14,33]. A similar increasing trend was also observed in the case of soluble proteins and polysaccharides (Fig. 3(b) and (c)). This corroborated the SCOD results, demonstrated more intracellular and/or extracellular constituents were released from the cells and/or EPSs [19,31].

As shown in Fig. 3(a), with RL addition increasing from 0 to 0.2 g/g TSS, the SCOD of LB-EPSs was increased from 131 to 918 mg/L in sole RL treatment, and from 623 to 978 mg/L in integration treatments. The SCODs of TB-EPSs all decreased, either in sole RL or integration treatment. A similar trends was also observed in the case of soluble proteins and polysaccharides both in LB-EPSs and TB-EPSs (Fig. 3(b) and (c)). This also corroborated the SCOD results.

The intracellular released capacity (IRC) was developed to evaluate the organics release from the cells under different pretreatment conditions. It can be expressed by the following equation:
IRC = \( \frac{(\text{DOM}_{\text{exp}}^x - \text{DOM}_{\text{raw}}^x)}{(\text{EPS}_{\text{exp}}^x - \text{EPS}_{\text{raw}}^x)} \) (5)

where \( \text{EPS}_{\text{exp}}^x \) and \( \text{EPS}_{\text{raw}}^x \) (mg/L) were the sums of SCOD in LB-EPSs and TB-EPSs under different pretreatment conditions at 12-h fermentation time and raw sludge, respectively. \( \text{DOM}_{\text{exp}}^x \) and \( \text{DOM}_{\text{raw}}^x \) (mg/L) were the corresponding SCOD in suspension, respectively. And \( x \) was the given experiment No., shown in Table 1.

As shown in Fig. 3, with RL dosage increasing, IRC increased in both sole RL and integration treatments. For SCOD, the highest IRC obtained at 0.2 g RL/g TSS under sole RL treatments, 4130 mg/L (11.76 times higher than control). In the integration treatments, the highest IRC obtained under 0.2 g RL/g TSS and initial pH = 10 treatment, 5822 mg/L (16.75 and 1.41 times respectively higher than that of control and sole RL dosage 0.2 g/g TSS).

A similar trend was also observed in the case of soluble proteins and polysaccharides (Fig. 3(b) and (c)), whose changes contributed to SCOD changes. These above results further demonstrated the positive effects of RL addition on the release of intracellular constituents, and RL integrated into alkaline anaerobic fermentation of WAS could result in positive synergies on organics release from the cells. As SCFAs mainly formed from the fermentation of polysaccharides and proteins [11,27], both sole RL and integration treatments could provide substrates efficiently for WAS anaerobic fermentation to produce SCFAs, but the integration treatment was more efficient.

3.2.2. EEM fluorescence spectra analysis of EPSs

Previous studies had reported that three-dimensional EEM could be applied to characterize complete information of some fluorescent substances (such as proteins, metabolites (e.g., amino, fulvic and hemic acids), enzymes and pigments) by changing excitation and emission wavelength simultaneously [41,42]. We used EEM spectra to comprehensively reveal the component changes of WAS under different conditions in terms of DOMs, LB-EPSs, and TB-EPSs [11,14,43,44]. And the PARAFAC, a chemometric method with three-way decomposition, was employed to separate the spectra of the main components from overlapped EEM spectra [41–44].

Two fluorescence peaks could be identified from the EEM spectra by the PARAFAC (Fig. 4 and Fig. S3). The two peaks were associated with tryptophan (Ex/Em 275/340, named component 1 (Com.1)) and tyrosine (Ex/Em 275/305, named component 2 (Com.2)) protein-like substances, respectively [11,12,14,42]. As shown in Fig. 4 and Fig. S3, all the samples were similar in the peak locations with main fluorophore of protein-like substances, but have different fluorescence intensities (FIs) for different pretreatment methods. In DOMs, compared with raw sludge, after 12-h pretreatment time, the FI all increased significantly, especially in experimental tests. The increased fluorescence signals could be attributed to the increase of protein-like substances in WAS mixed liquor [14,42,43]. This was consistent with the results of proteins.
under different conditions (Section 3.2.1). The maximum FI of both Com.1 and Com.2 were all obtained under the integration treatment, 351 and 450, respectively, which were respectively 1.73, 2.53 and 15.87 times higher for Com.2. The above results proved that positive synergies had been formed on WAS hydrolysis under integration treatment [11,14,33]. And all FI of Com.2 were higher than that of Com.1, indicating that tyrosine protein-like substances presented the main abundant compounds of released matters, the same results could be found in previous literature [14]. As the protein-like substances in DOMs must be released from either EPSs or intracellular materials, and were relatively easy to be biodegraded [42,43], the pretreatment methods applied, especially the integration method, were efficient for WAS hydrolysis, which closely related to WAS acidification.

The FI of LB-EPSs had the similar trend with DOMs (Fig. S3). This was also consistent with both the results provided in Section 3.2.1 and previous studies, in which, the changes of protein-like substances in DOMs must be released from either EPSs and/or cells of WAS were effectively ameliorated by pretreatment methods. Results above demonstrated that the release of protein-like substances in EPSs and/or cells of WAS were effectively ameliorated under integration treatment, which was beneficial to the followed acidification process.

3.3. Kinetic modeling of WAS hydrolysis and acidification

3.3.1. Effects of pre-treatment on hydrolysis of WAS

The level of soluble proteins and polysaccharides could be taken as an index to evaluate the efficiency of WAS hydrolysis [11], and the changes of SCOD mainly attributed to the contents of soluble proteins and polysaccharides [11,19,47]. Thus, released SCOD was used to evaluate the WAS hydrolysis under different conditions.

Fig. 5(a), (b) shows the hydrolysis kinetics of WAS under both sole RL and integration treatments. It was obvious that good agreement achieved between fitted and observed cumulative SCOD (%) with the strong correlation coefficients ($R^2$: 0.9503–0.9967), small $\Delta q$ (%) (1.95–9.37 < 10) and ARE (%) (0.038–0.86 < 1) (Table 2), indicating that WAS hydrolysis obeyed the PFO kinetics. A similar result had been reported by Feng et al. [48], hydrolysis of particulate COD in WAS was successfully presented by first-order kinetic model. With the RL levels increasing, the hydrolysis rate constant increased linearly in both sole RL and integration treatments (Fig. S4a). The highest hydrolysis rate constant was obtained in the 0.2 g RL/g TSS test, 4.06 times higher than control test. And it was further enhanced under combined treatment, which was 2.14 times higher than sole RL treatment.

It was clear that the equilibrium SCFAs production had a strongly linear relationship with hydrolysis rate constant (Fig. S4b), demonstrated that improving WAS hydrolysis could enhance SCFAs production significantly.

3.3.2. Kinetic modeling of SCFAs accumulation

The kinetic studies were of great importance for gaining insight on SCFAs accumulation during the anaerobic fermentation. In this work, in order to evaluate the kinetics of SCFAs accumulations, both PFO and PSO models were fitted to interpret the experimental data. The fitted plots showed that PFO was much better than PSO model (Fig. 5c–f) and fitted cumulative SCFAs production from Eq. (1) (Fig. S5c, d) were more similar with those observed experimentally than that from Eq. (2) (Fig. S5e, f), indicating that SCFAs accumulations under all conditions were better fitted PFO than PSO kinetic model. Results above were supported by the statistical indices (Table 2). The correlation coefficient ($R^2$), $\Delta q$ (%) and ARE (%) were used for model comparison and goodness-of-fit evaluation for a given pretreatment method. It was obvious that $R^2$'s of the PFO model were more than that of the PSO model for all the tests. By contrast, $\Delta q$ (%)s and ARE (%)s of PFO model were all less than that of PSO model, suggested that PFO model fitted to the experimental data better. This was similar with previous literature, in which, the accumulation of SCFAs obeyed first-order kinetics.
Fig. 4. EEM fluorescence spectra of DOMs, LB-EPSs, and TB-EPSs at 12-h fermentation time under different conditions.
Thus, PFO model was successfully built to present the SCFAs accumulation production.

### 3.4. RL pretreatment integrated into alkaline anaerobic fermentation for exploiting internal CS to enhance nutrients removal in a WWTP

Our previous study had reported that RL pretreatment prior to anaerobic digestion is an economically attractive and environmentally favorable method for enhancing SCFAs production from WAS [21], this study revealed that further enhancement of SCFAs production could be achieved by RL pretreatment integrated into alkaline anaerobic fermentation of WAS. Compared with our previous work [21], some new findings have been discussed and concluded. Firstly, it was proved that more carbon recovery was increased under the new integrated condition using RL and alkaline (shown in Section 3.1.2). The more internal CS produced in situ in a WWTP,

Fig. 5. Kinetic modeling of WAS hydrolysis and SCFAs accumulation under different conditions. (a), (b) Kinetic modeling of WAS hydrolysis, (c), (d) PFO model of SCFAs accumulation, and (e), (f) PSO model of SCFAs accumulation.

| Table 2 | Kinetic parameters for WAS hydrolysis and SCFAs accumulations under different conditions. |
|---------|------------------------------------------------------------------------------------------|
|         | WAS hydrolysis                                                                          |
|         | $k_1$ (h$^{-1}$)                                                                         |
|         | $q_e,cal$ (mg/L)                                                                         |
|         | $R^2$                                                                                   |
|         | $\Delta q$ (%)                                                                           |
|         | $ARE$ (%)                                                                                |
|         | $q_e,exp$ (mg/L)                                                                         |
|         | Pseudo-first-order of SCFAs accumulation                                                  |
|         | $k_1$ (h$^{-1}$)                                                                         |
|         | $q_e,cal$ (mg COD/L)                                                                     |
|         | $R^2$                                                                                   |
|         | $\Delta q$ (%)                                                                           |
|         | $ARE$ (%)                                                                                |
|         | Pseudo-second-order of SCFAs accumulation                                                |
|         | $k_2$ (L/(mg COD h))                                                                     |
|         | $q_e,cal$ (mg COD/L)                                                                     |
|         | $R^2$                                                                                   |
|         | $\Delta q$ (%)                                                                           |
|         | $ARE$ (%)                                                                                |
|         | $q_e,exp$ (mg COD/L)                                                                     |

[48]. Thus, PFO model was successfully built to present the SCFAs accumulation production.

Our previous study had reported that RL pretreatment prior to anaerobic digestion is an economically attractive and environmentally favorable method for enhancing SCFAs production from WAS [21], this study revealed that further enhancement of SCFAs production could be achieved by RL pretreatment integrated into alkaline anaerobic fermentation of WAS. Compared with our previous work [21], some new findings have been discussed and concluded. Firstly, it was proved that more carbon recovery was increased under the new integrated condition using RL and alkaline (shown in Section 3.1.2). The more internal CS produced in situ in a WWTP,
the less external CS for the operation of a WWTP needed [5,6,18,26]. Secondly, the efficient treatment led to a shortened fermentation time when using the new method. The maximum SCFAs accumulations achieved at 96-h (i.e., 4-day) or more fermentation time in the previous researches [6,12,13,21,27], but it was reduced to 72-h (i.e., 3-day) in this study (shown in Section 3.1.1), which was attractive for both improving the efficiency of WAS disposal and preparing SCFAs from WAS to meet the CS demand of a WWTP [6,12,18,27]. Thirdly, the mechanisms of the enhanced SCFAs accumulations under either sole RL treatments or integrated treatments have been identified by investigating the role of EPSs on WAS solubilization (shown in Section 3.2) and kinetic modeling of WAS hydrolysis and acidification (shown in Section 3.3), which were not disclosed well in previous studies [12,21,27].

Currently, some methods had been applied to improve SCFAs production from WAS with anaerobic fermentation process, and Table S2 exhibits some typical methods reported in the previous publications. It was clear that, among these available methods, pH = 10 had been known as the most common way to enhance SCFAs production [27,28]. More interesting, SCFAs production could be further enhanced by adding RL into alkaline anaerobic fermentation process with a short-term fermentation time in this study. As well known, while using WAS as the substrate to produce internal CS in a WWTP, the SCFAs production should be maximized, in contrast, the input should be minimized. Unlike other pretreatment methods (e.g., ultrasonic), RL was a renewable chemical that could be in situ generated during the SCFAs production, proposed by our previous study [21], suggested that RL pretreatment integrated into alkaline anaerobic fermentation for SCFAs production from WAS was a potential and practical technology.

Therefore, a sustainable and economical technology was urgently needed to apply to enhance nutrient removal in WWTPs (Fig. 6). And Tong et al. had reported that WAS alkaline fermentation liquid as the additional CS for municipal wastewater biological nitrogen and phosphorus removal, could obtain good performance, but the released nitrogen and phosphorus should be firstly separated from fermentation liquid [5]. Thus, the proposed concept in Fig. 6 may be potential and practical to reduce both operation cost and sludge production simultaneously of WWTPs.

4. Conclusions

The feasibility of short-term fermentation for enhancing SCFAs production from WAS with a novel proposed method, RL pretreatment integrated into alkaline (initial pH10) anaerobic fermentation, has been successfully investigated in this work. The main conclusions are: (1) positive synergies on anaerobic fermentation process had been formed under integration conditions. The maximum SCFAs production was increased to 378 mg COD/g VSS at 72-h fermentation time, when WAS was pretreated by initial pH 10 and RL dosage 0.2 g/g TSS. (2) Compared with control, initial pH 10 and sole RL methods, the integration method performed best to release intracellular and/or extracellular constituents from the cells and/or EPSs to suspension, and hydrolysis rate was 2.14 times higher than that of sole RL treatment. (3) The PFO model was successfully built to present the SCFAs accumulation production. (4) RL pretreatment integrated into alkaline anaerobic fermentation was an attractive technology for enhancing sustainable CS (SCFAs) production from WAS in situ in a WWTP.

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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/jcej.2016.01.033.

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