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Cassava stillage and its anaerobic fermentation liquid as external carbon sources in biological nutrient removal^{*}

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Abstract: The aim of this study was to investigate the effects of one kind of food industry effluent, cassava stillage and its anaerobic fermentation liquid, on biological nutrient removal (BNR) from municipal wastewater in anaerobic anoxic-aerobic sequencing batch reactors (SBRs). Experiments were carried out with cassava stillage supernatant and its anaerobic fermentation liquid, and one pure compound (sodium acetate) served as an external carbon source. Cyclic studies indicated that the cassava by-products not only affected the transformation of nitrogen, phosphorus, poly-β-hydroxyalkanoates (PHAs), and glycogen in the BNR process, but also resulted in higher removal efficiencies for phosphorus and nitrogen compared with sodium acetate. Furthermore, assays for phosphorus accumulating organisms (PAOs) and denitrifying phosphorus accumulating organisms (DPAOs) demonstrated that the proportion of DPAOs to PAOs reached 62.6% (Day 86) and 61.8% (Day 65) when using cassava stillage and its anaerobic fermentation liquid, respectively, as the external carbon source. In addition, the nitrate utilization rates (NURs) of the cassava by-products were in the range of 5.49–5.99 g N/(kg MLVSS·h) (MLVSS is mixed liquor volatile suspended solids) and 6.63–6.81 g N/(kg MLVSS·h), respectively. The improvement in BNR performance and the reduction in the amount of cassava stillage to be treated *in-situ* make cassava stillage and its anaerobic fermentation liquid attractive alternatives to sodium acetate as external carbon sources for BNR processes.

Key words: Biological nutrient removal (BNR), Denitrification, Enhanced biological phosphorus removal (EBPR), External carbon source

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1 Introduction

Due to increasingly stringent limits for nitrogen and phosphorus removal, the amount of organic compounds in influent wastewater often is not sufficient for efficient denitrification and enhanced biological phosphorus removal (EBPR) in biological nutrient removal (BNR) systems. In such cases, specific available commercial compounds ("conventional" external carbon sources), such as methanol, ethanol, acetic acid, and sodium acetate, are usually added to improve the overall BNR performance and obtain low effluent levels for nitrogen and phosphorus. However, these commercial external carbon sources are expensive and usually require acclimation periods. Such attributes do not recommend their application in full-scale plants. On the other hand, in many cases, the problem of high costs of commercial organic compounds can be overcome if industrial wastewater, such as brewery wastewater, is readily accessible (Henze *et al.*, 1995). Various industrial by-products or waste materials have recently received

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more attention as "alternative", cost-effective external carbon sources for denitrification (Makinia *et al.*, 2012). Among these materials, food industry effluents stand out as being potential alternatives for "conventional" external carbon sources due to their high C/N ratios and high content of readily biodegradable organic compounds (Lim *et al.*, 2000; Cappai *et al.*, 2004; Quan *et al.*, 2005; Sage *et al.*, 2006; Swinarski *et al.*, 2009a). However, the use of food industry effluents may result in some operating problems, including variation in quality and quantity resulting from their seasonal production cycles.

Different food industry effluents as external carbon sources have been shown to enhance denitrification in BNR systems. Swinarski et al. (2009b) investigated three different types of industrial wastewater (effluents from a distillery, brewery, and fish-pickling factory, respectively) on the denitrification capability of process biomass originating from a full-scale BNR plant in northern Poland. These three food industry effluents were characterized by a high content of easily biodegradable chemical oxygen demand (COD) (33.7%-88.5% of the total COD) and low nutrient concentrations (COD/total nitrogen (TN) ratios in the range of 15.1-80.6). The observed single nitrate utilization rates (NURs) for three industrial wastewaters ranged from 4.8 to 5.1 mg N/(g VSS \cdot h), 2.4 to 6.0 mg N/(g VSS \cdot h), and 4.0 to 6.0 mg N/ (g VSS h) (VSS is volatile suspended solids), respectively, compared with the rates associated with the use of readily biodegradable COD in settled wastewater. Quan et al. (2005) recommended hydrolyzed molasses as an economical and effective external carbon source for nitrogen removal. They found that with hydrolyzed molasses the denitrification rate was 2.9–3.6 mg N/(g VSS \cdot h) and that a higher TN removal efficiency ($(91.6\pm1.6)\%$) was obtained than when using methanol ((85.3±2.0)%). Rodríguez et al. (2007b) studied six different agro-food wastewaters from cheese production, milk bottling, potato processing, beet sugar processing, tomato processing, and a winery, respectively. Although the measured denitrification rates, in the range of $2.0-4.1 \text{ mg N/(g VSS \cdot h)}$, were lower than those obtained for acetate (4.7 mg N/(g VSS·h)) and municipal wastewater (4.3 mg N/ (g VSS·h)), the addition of tomato processing wastewater as the external carbon source raised the denitrification capacity significantly, with 99% of nitrate nitrogen removed in a full-scale wastewater treatment plant (WWTP). In addition, examples of noticeably higher denitrification rates resulting from the use of industrial effluents can be found in the literature. Cappai *et al.* (2004) tested wastewaters from the ice-cream industry and a beet-sugar factory and found that the denitrification rates were 6.7 and 5.5 mg N/(g VSS·h), respectively. Similar values were obtained with dairy effluent, glycerin and meat processing wastewater. Some other studies, for instance, with glycerol, fusel oil, brewery effluent, and wine distillery effluent revealed values exceeding 13 mg N/(g VSS·h) (Monteith *et al.*, 1980; Bernet *et al.*, 1996; Hinojosa *et al.*, 2008; Kampas *et al.*, 2009).

The effect of industrial effluents as external carbon sources in enhancing denitrification in BNR systems has been investigated and reported extensively, but very little is known about their effects as external carbon sources for EBPR. It is well known that in EBPR metabolism, the composition of poly-β-hydroxyalkanoates (PHAs), including poly-βhydroxybutyrate (PHB), poly-β-hydroxyvalerate (PHV) and poly-3-hydroxy-2-methylvalerate (PH2MV), produced by different types of phosphorus accumulating organisms (PAOs) and denitrifying phosphorus accumulating organisms (DPAOs) during the anaerobic period, varies based on the carbon source. However, PAOs and DPAOs are unable to metabolize these different types of PHA at the same rate in the subsequent anoxic phase (Oehmen et al., 2005a). Therefore, the use of different carbon sources may lead to different properties of phosphorus metabolism and removal. Rodríguez et al. (2007a) evaluated the performance of the use of winery wastewater to enhance the BNR process. A dose of winery wastewater significantly enhanced nitrogen removal (from 58% to 75%). However, the phosphorus removal remained constant. Similar findings were reported by Rodríguez et al. (2007b), who found that the optimum dosage of tomato processing wastewater applied as an external carbon source enhanced nitrate removal, but the COD and phosphorus removal were not significantly affected. However, other studies showed that both nitrogen and phosphorus removals were enhanced by adding external carbon sources. Lee et al. (2003) investigated the feasibility and applicability of anaerobically fermented leachate of food waste (AFLFW) as an external carbon source in a lab-scale BNR process treating domestic sewage with a low C/N ratio of 4.4. As AFLFW was supplied, not only did the average removal efficiency of TN increased from 53% to 74%, but also the total phosphorus (TP) removal efficiency increased from 21% to 68%. Higher values were reported by Chae *et al.* (2004) who found that when AFLFW was supplied, the average removal efficiencies of TN and TP increased from 53% to 97% and from 55% to 93%, respectively. They attributed the denitrification efficiency and EBPR activity to the concentration of volatile fatty acids (VFAs) in the external carbon source.

In this study, the effects of one kind of food industry effluent, cassava stillage and its anaerobic fermentation liquid, used as external carbon sources, on denitrification and EBPR were investigated. The behavior of intracellular materials including PHA and glycogen affecting the EBPR mechanism was also examined.

2 Materials and methods

2.1 Sequencing batch reactor (SBR) set-up and operation

We compared the performance, in terms of nitrogen and phosphorus removal, of three bench-scale anaerobic/anoxic/aerobic SBRs, each with a working volume of 3.0 L. The carbon sources for the three SBRs (SA-SBR, CS-SBR and CSF-SBR) were sodium acetate (SA), cassava stillage supernatant (CS) and its anaerobic fermentation liquid (CSF), respectively. The SBRs were completely automated with time controllers regulating all peristaltic pumps, stirrers, air pumps, and phase lengths. Sludge originated from a sewage treatment plant (Quyang Sewage Treatment Plant, Shanghai, China) that employed a BNR process. The three SBRs were operated on three 8-h cycles per day, with each cycle consisting of 10 min feeding, 90 min anaerobic, 210 min anoxic, 40 min aerobic, followed by 60 min settling, 10 min decanting and 70 min idle periods. During the first 10 min feeding period in each reactor, 1.5 L of synthetic wastewater was pumped into the reactor, while 50 ml of KNO₃ stock solution was pulse-added into the reactors at the beginning of the anoxic period, giving an initial NO_3 -N concentration of 35 mg/L. During the aerobic phase, the dissolved oxygen (DO) concentration in each SBR was around 3 mg/L, and

50 ml of sludge was removed daily at the end of the aerobic period, giving a solids retention time (SRT) of about 20 d. Mixed liquor suspended solids (MLSS) concentrations of the three SBRs were around 3000 mg/L, and the hydraulic retention time (HRT) was about 16 h. The SA-SBR and CS-SBR were operated from August to November 2012, while the CSF-SBR was operated from November 2012 to January 2013.

2.2 Characteristics of the food industry effluents

The cassava stillage used as a carbon source in this work originated from a full-scale cassava ethanol plant (Jiangsu, China). Before experiments, the stillage was concentrated by settling at 4 °C for 24 h and the decanted supernatant (CS) was subsequently used for 5-d anaerobic alkaline fermentation at pH 9.0 and 35 °C. After fermentation, the liquid phase (CSF) was centrifuged at 1000 r/min for 10 min and stored at 4 °C before use. Sample analyses of CS and CSF revealed high COD concentrations and low total nutrient concentrations (COD/TN ratios of 84±5.4 and 53±4.2 for CS and CSF, respectively) (Table 1). The VFAs produced in CSF contained 35.8% acetate, 20.9% propionate, 42.8% butyrate, and 0.5% iso-butyric acid. The VFAs were the major COD component with an average VFA/COD ratio of (73 ± 0.9) %, which was comparable to the obtained (79%-80%) when using AFLFW as the external carbon source (Lee et al., 2003). The residual carbohydrate in the fermentation liquid contributed to

 Table 1
 Characteristics of cassava stillage supernant

 (CS) and its anaerobic fermentation liquid (CSF)

()	-	()
Constituent	CS	CSF
pH	4.57±0.02	8.10±0.12
COD (mg/L)	43720±320	31580 ± 260
NH ₃ -N (mg/L)	$12.0{\pm}1.0$	20.0±1.6
TN (mg/L)	520±21	600±27
TP (mg/L)	1072±25	1116±42
Soluble carbohydrate (mg/L)	13290±43	2858±292
Soluble protein (mg/L)	3514±257	983±57
Ethanol (mg/L)	299.8±13.2	0 ± 0
Acetic acid (mg/L)	249.3±10.6	8216±476
Propionic acid (mg/L)	0 ± 0	4801±33
Butyric acid (mg/L)	0±0	9820±201
Iso-butyric acid (mg/L)	0 ± 0	126±6
Valeric acid (mg/L)	0±0	0 ± 0
Iso-valeric acid (mg/L)	0 ± 0	0±0
COD/TN	84.0±5.4	53.0±4.2

The values are expressed as mean±SD (n=3)

the remaining COD in the CSF. Compared with the CSF, the main compounds besides acetate contributing to the COD in the CS were carbohydrate and protein. More detailed characteristics of the tested carbon sources are presented in Table 1.

2.3 Synthetic wastewater

The synthetic wastewater used in this study contained (per liter): 13.16 mg of KH₂PO₄ (3.0 mg of P), 16.84 mg of K₂HPO₄ (3.0 mg of P), 57.4 mg of NH₄Cl (15.0 mg of N), 85 mg of MgSO₄·7H₂O, 10 mg of CaCl₂, and 110 mg of NaHCO₃. The three SBRs were supplemented with SA, CS, and CSF, respectively, as the sole carbon source to achieve a feed COD of about 300 mg/L. Additionally, 0.5 ml of mineral salt solution was added per liter of synthetic wastewater. Each liter of mineral salt solution was composed of 5 g FeCl₃·6H₂O, 0.15 g H₃BO₃, 0.03 g CuSO₄·5H₂O, 0.18 g KI, 0.12 g MnCl₂·4H₂O, 0.06 g Na₂MoO₄·2H₂O, 0.12 g ZnSO₄·7H₂O, 0.15 g CoCl₂·6H2O, and 10 g ethylene diamine tetraacetic acid (EDTA). The wastewater was fed into the three SBRs at the beginning of each cycle, giving the feed of each SBR about 300 mg/L COD, 6 mg/L soluble ortho-phosphorus (SOP), and 15 mg/L NH_4^+ -N.

2.4 Batch experiments

2.4.1 Conventional NURs

Fresh activated sludge from the three steadystate SBRs at the end of the decanting period and the three carbon sources were mixed, respectively, to obtain 1.5 L of mixed liquor with volatile suspended solids (MLVSS) concentration of about 2000 mg/L. Measurements were recorded for 6 h after injecting KNO₃ stock solution to raise the initial NO₃⁻-N concentration to 30 mg/L. Samples were withdrawn at a frequency of 5–30 min and filtered through Millipore filter units (0.45 µm pore size) for analyses of COD, NO₂⁻-N, and NO₃⁻-N.

2.4.2 Anaerobic-anoxic/aerobic tests

Mixed liquor (1.5 L) containing active biomass from the three steady-state SBRs during the aerobic period was incubated in the SBRs and maintained in anaerobic conditions in the presence of one of the three carbon sources to reach a COD of 300 mg/L. The sludge was homogenized in the anaerobic phase to promote phosphate release and COD uptake for 4 h. Subsequently, mixed liquor in each batch reactor was divided into two sets to perform the anoxic/aerobic phosphate uptake experiments: one set was exposed to anoxic conditions for 3 h where KNO₃ stock solution was added at the beginning of anoxic to produce an NO₃⁻N concentration of 30 mg/L; the other set was exposed to aerobic conditions for 3 h where air was injected into the mixture to maintain the DO concentrations at around 3 mg/L. Samples were withdrawn at a frequency of 20-30 min and filtered through Millipore filter units (0.45 µm pore size) for analyses of COD and PO_4^{3-} -P. The phosphate uptake rates (PURs) under anoxic and aerobic conditions were estimated from the linear regression of phosphate concentrations. The ratio of the anoxic PUR to the aerobic PUR was used as an index (denoted K) reflecting the fraction of DPAOs (Wachtmeister et al., 1997).

2.5 Analytical methods

The liquid samples from the three SBRs were immediately filtered through Millipore filter units (0.45 μ m pore size) for analyses of COD, NH₄⁺-N, NO₂⁻-N, NO₃⁻-N, TN, PO₄³⁻-P, and TP. COD, TN, PO₄³⁻-P, TP, MLSS, and MLVSS were measured according to standard methods (APHA, 1998). NH₄⁺-N, NO₂⁻-N, and NO₃⁻-N were analyzed by ion chromatography using two conductivity detectors and two sets of columns (Dionex ICS-3000). The soluble carbohydrate concentration was measured by the phenol sulfuric acid method, with glucose as standard (Dubois et al., 1956). Soluble protein was determined by the Lowry-Folin method with bovine serum albumin as standard (Lowry et al., 1951). Samples for the analyses of ethanol and VFAs (acetic, propionic, butyric, iso-butyric, valeric, iso-valeric acids) were diluted with a 3% (0.03 g/ml) H₃PO₄ solution (sample/ acid 1:1, v/v) and were analyzed by gas chromatography (GC; Agilent, 6890 N) using a flame ionization detector (FID) and analytical column CPWAX52CB $(30 \text{ m} \times 0.25 \text{ mm} \times 0.25 \text{ } \mu\text{m})$. The temperatures of the injector and FID were 200 and 220 °C, respectively. Nitrogen gas was used as the carrier gas with a flow rate of 25 ml/min. The temperature of GC-oven was programmed to begin at 110 °C and remain there for 2 min, then increase at a rate of 10 °C/min to 220 °C, and hold at 220 °C for an additional 2 min. The sample injection volume was 1.0 µl.

The analyses of PHB, PHV, and PH2MV were conducted according to the method described by

Oehmen et al. (2005b). Lyophilized sludge samples were digested, methylated, and extracted with chloroform. The extracted methyl esters were then identified using an HP 4890 GC equipped with an FID detector and a 30 m×0.53 mm×0.88 µm HP-5 column. Helium was the carrier gas with a flow rate of 30 ml/min. The GC-oven temperature was programmed to begin at 120 °C and remain there for 2 min, then increase at a rate of 8 °C/min to 200 °C and hold at 200 °C for an additional 2 min, and lastly increase at a rate of 15 °C/min to 270 °C and hold at 270 °C for an additional 5 min. The standards of PHB/PHV (88%/12%) and poly-2-hydroxycaproic acid were purchased from the Sigma-Aldrich Chemical Company (USA) and used for the establishment of the calibration curve. Note that no standard sample was available for the direct measurement of PH2MV. As poly-2-hydroxycaproic acid and PH2MV are isomers, poly-2-hydroxycaproic acid was used for the standard curve of PH2MV. The total PHA in the samples was calculated as the sum of the measured PHB, PHV, and PH2MV. Glycogen was determined using the method described by Fang et al. (2002).

3 Results

3.1 Comparison of COD transformations with different carbon sources

Cyclic studies were performed in the three SBRs after stable nutrient and phosphorus conversions were achieved. Fig. 1 shows the COD concentration profiles in the three SBRs. During the anaerobic stage, carbon sources were rapidly consumed in both the SA-SBR and CSF-SBR, even if a little slowly in CSF-SBR, with the COD concentrations at 64 and 82 mg/L at the end of the anaerobic period in the SA-SBR and CSF-SBR, respectively. In the CS-SBR, the COD concentration was higher at 189 mg/L, due to the high amount of compounds of large molecular size, such as carbohydrate and protein, in CS. Small molecules like VFAs, can be easily used by the storage of PHA and the release of phosphorus in the anaerobic period. In the subsequent anoxic period, the COD concentration decreased continuously in the CS-SBR. The COD concentrations at the end of anoxic period were 29, 42, and 82 mg/L in the SA-SBR, CSF-SBR, and

CS-SBR, respectively. The higher effluent COD concentration in the CS-SBR indicated that pre-treatment of CS, such as anaerobic alkaline fermentation, was necessary to facilitate the bio-degradation of large molecules.



Fig. 1 Variation in COD in three SBRs fed with different carbon sources

The values are expressed as mean \pm SD (n=3)

3.2 Comparison of nitrogen transformations with different carbon sources

The variations in NH₄⁺-N, NO₂⁻-N, and NO₃⁻-N in the three SBRs are shown in Fig. 2. Ammonium was rapidly oxidized and decreased in the aerobic stages, accompanied by increases in NO2-N and NO₃⁻N. The three SBRs all achieved high ammonia oxidation rates in the aerobic stage, and the ammonia was eventually removed. In the anoxic period, once KNO₃ solution was added, the denitrification process started. The NO₃-N concentration became depleted and then remained constant in the SA-SBR after 60 min, while in the CS-SBR and CSF-SBR it continued to go down. During the anoxic stage, the specific denitrification rates in the CS-SBR and CSF-SBR were 2.71 and 3.19 mg NO_{r} -N/ (g MLVSS·h), respectively, while the SA-SBR demonstrated a higher rate of 4.45 mg NO_x -N/ (g MLVSS h). At the end of the anoxic stage, the NO_x⁻-N concentrations in the CSF-SBR, CS-SBR and SA-SBR were (6.63±0.71), (12.83±0.77), and (15.50± 0.52) mg/L, respectively, resulting in a higher TN removal efficiency in the CSF-SBR ((73.1±1.3)%) and CS-SBR ((61.6±0.9)%) than in the SA-SBR $((55.2\pm2.0)\%)$ at the end of the aerobic period. Note that significant NO₂⁻-N accumulation occurred during denitrification in the three reactors. This can be

attributed to the slower rate of reduction of nitrite compared with when there is limited availability of biodegradable organic carbon (Oh and Silverstein, 1999). The timing of the peak NO₂⁻-N concentration in the SA-SBR (2.25 h) was earlier than that of the CS-SBR (2.5 h) and CSF-SBR (3 h) (Fig. 2c). However, the peak NO₂⁻-N concentrations in the CSF-SBR and CS-SBR were (6.26 ± 0.31) and (16.06 ± 0.75) mg/L, respectively, much lower than that in the SA-SBR ((24.01 ± 1.02) mg/L) during the anoxic stage. These observations suggest that NO₂⁻-N accumulation is linked to the carbon source and denitrification rate (Zhou *et al.*, 2008; Wang *et al.*, 2011).

3.3 Comparison of PO₄³⁻-P, PHA, and glycogen transformations with different carbon sources

3.3.1 Effects of carbon sources on the transformation of $PO_4^{3^-}$ -P

As mentioned above, the carbon sources were rapidly consumed during the anaerobic stage, and this process was accompanied by the release of phosphorus, the formation of PHA, and the degradation of glycogen. The variation in PO_4^{3-} -P during cyclic studies in the three SBRs is illustrated in Fig. 3. Significant anaerobic phosphate release and aerobic phosphate uptake occurred in the three SBRs. Note that in the anoxic stage, significant anoxic P uptake occurred concurrently with denitrification in both the CS-SBR and CSF-SBR, but not in the SA-SBR. This indicates that EBPR may occur in the CS-SBR and CSF-SBR.

The specific anaerobic phosphorus release in the SA-SBR ((4.45 ± 0.28) mg PO₄³⁻-P/g MLVSS) was higher than that in the CS-SBR ((2.13±0.15) mg $PO_4^{3-}-P/g$ MLVSS) and the CSF-SBR ((3.27± 0.26) mg PO_4^{3-} -P/g MLVSS), as shown in Table 2. However, the largest difference between P uptake and release occurred in the CSF-SBR ((2.46±0.17) mg PO₄³⁻-P/g MLVSS), followed by the SA-SBR $((2.36\pm0.18) \text{ mg PO}_4^{3-}-P/g \text{ MLVSS})$ and the CS-SBR $((1.80\pm0.17) \text{ mg PO}_4^{3-}-P/g \text{ MLVSS})$. Better phosphorus removal efficiencies were achieved in the CSF-SBR ((67.1±2.5)%) and the CS-SBR ((78.6± 3.7)%) than in the SA-SBR ((56.5±3.1)%). Better phosphorus removal was also obtained using VFAs derived from the fermenter as the carbon source than by dosing an equivalent amount of acetic acid (Thomas et al., 2003).



Fig. 2 Variations in NH_4^+ -N (a), NO_3^- -N (b), and NO_2^- -N (c) in three SBRs fed with different carbon sources The values are expressed as mean \pm SD (n=3)



Time (min) Fig. 3 Variation in PO₄³⁻-P in three SBRs fed with different carbon sources

The values are expressed as mean \pm SD (*n*=3)

Table 2 Transformation of PO₄³⁻-P, PHA, and glycogen in three SBRs with different carbon sources

Parameter	SA	CS	CSF
Initial P concentration (mg/L)	6.06±0.37	5.99±0.40	6.05±0.42
Specific anaerobic P release (mg PO ₄ ³⁻ -P/g MLVSS)	4.45±0.28	2.13±0.15	3.27±0.26
Anoxic P uptake (mg PO ₄ ³⁻ -P/g MLVSS)	0.47 ± 0.10	2.19±0.23	3.59±0.29
Aerobic P uptake (mg PO_4^{3-} -P/g MLVSS)	6.34±0.37	1.74 ± 0.14	2.14±0.20
Difference between P uptake and release (mg PO_4^{3-} -P/g MLVSS)	2.36±0.18	1.80 ± 0.17	2.46±0.17
Aerobic end P concentration (mg/L)	2.64 ± 0.25	1.28 ± 0.20	1.99±0.26
P removal efficiency (%)	56.5±3.1	78.6±3.7	67.1±2.5
PHA synthesis (mmol C/g MLVSS)	3.27±0.14	2.62±0.11	2.07±0.10
Glycogen degradation (mmol C/g MLVSS)	2.14±0.11	1.23 ± 0.05	0.98 ± 0.06
PHA degradation (mmol C/g MLVSS)	2.78±0.09	2.45±0.06	1.57±0.06
Glycogen synthesis (mmol C/g MLVSS)	2.29 ± 0.06	1.31±0.04	1.18±0.06

The values are expressed as mean±SD (n=3)

3.3.2 Effects of carbon sources on the transformation of PHA and glycogen

The structure of PHA formed in cells depends on the characteristics of substrates and microorganisms in the system (Chae et al., 2004; Li et al., 2008; Pijuan et al., 2009). The transformation of PHA in the three SBRs is shown in Fig. 4. When SA was used as a substrate, PHB was the major compound, with the average compositions of PHB, PHV, and PH2MV being 97.6%, 2.0%, and 0.4%, respectively. When CS and CSF were used as the substrates, a decrease in PHB and an increase in PHV and PH2MV of PHA occurred, with the average compositions of PHB, PHV, and PH2MV being 78.6%, 18.0%, and 3.4%, respectively, in CS-SBR and 49.3%, 41.1%, and 9.6%, respectively, in CSF-SBR. These findings correspond well with the results of other studies showing that the ratio of PHB synthesis/PHA synthesis decreases when the carbon source has a higher ratio of propionate/acetate (Lemos et al., 2003; Wang et al., 2011). In addition, negligible or very little amounts of PH2MV were anaerobically synthesized in the three SBRs. These findings correspond with those of other studies (Pijuan et al., 2009; Wang et al., 2011).

The reducing power necessary for anaerobic PHA accumulation is primarily derived from the degradation of stored glycogen (Fig. 4c). The amount of anaerobic glycogen degradation decreased when the carbon source was changed from SA ((2.14 ± 0.11) mmol C/g MLVSS) to CS ((1.23 ± 0.05) mmol C/g MLVSS) or CSF ((0.98 ± 0.06) mmol C/g MLVSS), which was responsible for a trend similar to the amount of PHA synthesis in each reactor (Fig. 5, Table 2). On the other hand, glycogen transformation showed a trend similar to PHA degradation. The



Fig. 4 Variation in PHA, PH2MV (a), PHB, PHV (b), and glycogen (c) in three SBRs fed with different carbon sources

The values are expressed as mean \pm SD (*n*=3)

glycogen replenishments in the CS-SBR ((1.31 ± 0.04) mmol C/g MLVSS) and CSF-SBR ((1.18 ± 0.06) mmol C/g MLVSS) were lower than that in the SA-SBR ((2.29 ± 0.06) mmol C/g MLVSS). This suggests that more PHA would be saved for taking up more phosphate, although the PHA accumulation was lower in the CS-SBR and the CSF-SBR. Thus, higher TP removal efficiencies were observed in the CS-SBR and CSF-SBR than in the SA-SBR.



Fig. 5 Effects of carbon sources on PHA synthesis (a), PHA degradation (b), and glycogen synthesis/ degradation (c) in three SBRs

The values are expressed as mean \pm SD (*n*=3)

3.4 Batch tests

3.4.1 Conventional NURs

The specific conventional NUR values measured with SA, CS, and CSF are listed in Table 3, and the experimental results are illustrated in Fig. 6. In the experiments with the CS, the NURs associated with use of the readily biodegradable COD (RBCOD) (NUR1) and the slowly biodegradable fraction (NUR2) varied in the ranges of 5.49-5.99 and 0.80-1.06 g N/(kg MLVSS h), respectively. These results suggested that the RBCOD was used first, but many of the other CODs (e.g. particulate COD or macromolecular compounds) were used subsequently and more slowly. Moreover, the explicit double rates observed in this study resulted from the content of various COD fractions in CS, revealing different rates of biodegradability, similar to other industry wastewaters used as external carbon sources (Rodríguez et al., 2007a; 2007b; Swinarski et al., 2009a). In the case of CSF, a single NUR value was observed. This could be explained by taking into account that the fermentation substrates were the most abundant COD fraction in the CSF studied. The NUR of CSF remained in the range of 6.63-6.81 g N/ (kg MLVSS h), which was slightly higher than the range associated with the use of the RBCOD in the CS (5.49-5.99 g N/(kg MLVSS · h)). The measured NURs of the CS and CSF were similar to or higher than those of other industry wastewaters as external carbon sources (Table 3), indicating that CS and CSF could be economically and environmentally sound external COD sources for the denitrification process.

The biodegradable organic compounds were less effective during the experiments with the industrial wastewater, which was reflected by much higher $\Delta COD/\Delta N$ ratios (ΔCOD is the amount of removed COD and ΔN is the amount of nitrogen removed) with CS (11.11–11.56 g COD/g N) and CSF (10.07–10.38 g COD/g N) than with SA (9.20–9.84 g COD/g N). This can be explained as follows: it is most likely that the 0.45 µm filters used to determine soluble COD (SCOD), allowed colloidal matter to be included in the SCOD determination. This colloidal matter may be removed instantaneously by sorption and may not be associated with the SCOD removal determined during denitrification (Swinarski *et al.*, 2009a).

~ .	Process temperature	$\Delta COD/\Delta N$ ratio	NUR	Literature	
Carbon source	(°C)	(g COD/g N)	(g N/(kg MLVSS·h))		
AFLFW	20		8.2	Lee et al., 2003	
Ice cream production industry wastewater	20		3.28	Cappai et al., 2004	
Beet-sugar industry wastewater	20		2.72	Cappai et al., 2004	
Hydrolized molasses			2.9-3.6	Quan et al., 2005	
Dairy effluent			3.4-8.0	Sage et al., 2006	
Beet-sugar processing wastewater	20		2.5 (NUR 1) 1.7 (NUR 2)	Rodríguez et al., 2007b	
Potato processing wastewater	20		4.1 (NUR 1) 1.7 (NUR 2)	Rodríguez <i>et al.</i> , 2007a; 2007b	
Milk bottling wastewater	20		3.8 (NUR 1) 1.3 (NUR 2)	Rodríguez <i>et al.</i> , 2007a; 2007b	
Cheese industry wastewater	20		2.8 (NUR 1) 1.1 (NUR 2)	Rodríguez <i>et al.</i> , 2007a; 2007b	
Tomato processing wastewater	20		2.7 (NUR 1) 1.9 (NUR 2)	Rodríguez <i>et al.</i> , 2007a; 2007b	
Winery wastewater	20		2.0 (NUR 1) 1.8 (NUR 2)	Rodríguez <i>et al.</i> , 2007a; 2007b	
Brewery effluent	20		15.6	Kampas et al., 2009	
Distillery wastewater	23.1-24.0	11.1-12.7	4.8-5.1	Swinarski et al., 2009b	
Bewery wastewater	19.7-22.7	6.5-20.8	2.4-6.0	Swinarski et al., 2009b	
Fish-picking wastewater	21.4-23.0	11.7-13.1	4.0-6.0	Swinarski et al., 2009b	
SA	18.9-23.1	9.20-9.84	16.69–13.62	This study	
CS	19.6–21.9	11.11–11.56	5.49–5.99 (NUR 1) 0.80–1.06 (NUR 2)	This study	
CSF	12.8-13.1	10.07-10.38	6.63–6.81	This study	

1 able 5 Comparison of NUKS when using different carbon source	Table 3	Comparison	of NURs	when using	g different	carbon sources
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AFLFW: anaerobically fermented leachate of food waste; NUR1: the NURs associated with use of the readily biodegradable COD (RBCOD); NUR2: the NURs associated with use of the slowly biodegradable fraction

3.4.2 Anaerobic-anoxic/aerobic tests

Cyclic studies demonstrated that the SBRs fed with CS and CSF as the sole carbon source displayed EBPR. Therefore, anaerobic-anoxic/aerobic tests were carried out to quantify the PAO and DPAO activity in the systems. Anoxic P uptake indeed occurred in both the CS-SBR and CSF-SBR, but not in the SA-SBR (Fig. 7). Thomas et al. (2003) proposed that glycogen accumulating organisms (GAOs) have a competitive advantage by using acetate and PAOs have a competitive advantage by using propionate, butyrate, or some other SCOD compounds in the fermenter effluent. This may be why there was no EBPR in the SA-SBR, in which acetate facilitated the GAO growth. However, further research is needed to clarify this issue. It is important to observe that P release in anaerobic period and P uptake in the aerobic period increased during the experimental period (Table 4), similar to the results of Monclús et al. (2010). Since the energy production efficiency of DPAOs growing with nitrate instead of oxygen is about 40% lower than that of PAOs (Kuba et al., 1996a), the anoxic PUR value of DPAOs is generally lower than the aerobic PUR value of PAOs (Table 4). This means that the P uptake activity by PAOs was higher than by that DPAOs (Peng et al., 2006; Oehmen et al., 2007). As the lower energy production efficiency of DPAOs growing under anoxic conditions results in a 20%-30% lower cell yield (Kuba et al., 1996a), it is expected that PAOs would outcompete DPAOs and that the DPAO/PAO ratio would decrease with time (Monclús et al., 2010). On Day 86, 62.6% of the population in PAOs was in the form of DPAOs in the CS-SBR, while this proportion decreased to 49.1% on Day 100. Nevertheless, simultaneous nitrogen and phosphorus removal by DPAOs requires less demand for oxygen and carbon sources, which leads to reduced plant operational costs (Kuba et al., 1996b). Therefore, the use of DPAOs for BNR processes is very economical and attractive.



Fig. 6 NO_x⁻-N and COD consumption in NUR measurements with SA (a), CS (b), or CSF (c) as the sole carbon source The values are expressed as mean \pm SD (n=3)



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Carbon source	Time	P _{rea} rate (mg PO ₄ ³⁻ -P/(g MLVSS·h))	Anoxic P _{upt} rate (mg PO ₄ ³⁻ -P/(g MLVSS·h))	Aerobic P _{upt} rate (mg PO ₄ ³⁻ -P/(g MLVSS · h))	DPAO (%)
SA	Day 86	2.93	0	4.13	0
	Day 100	3.36	0	4.81	0
CS	Day 86	3.61	2.26	3.61	62.6
	Day 100	4.73	2.32	4.72	49.1
CSF	Day 65	2.15	2.43	3.93	61.8
	Day 79	2.51	2.29	4.26	53.8

 P_{rea} : P release in anaerobic period; P_{upt} : P uptake in the anoxic or aerobic period

3.5 SBR performance

Table 5 depicts the performance of the three SBRs at steady-state conditions in terms of removal efficiencies of COD, NH4⁺-N, TN, and TP. It is evident that the effluent COD concentrations in the CS-SBR were relatively high compared with those in the SA-SBR and CSF-SBR. This suggests that the addition of CS as carbon source may have led to the increase in effluent COD concentration. Lee et al. (2003) stated that compounds with a molecular size of more than 30 kDa are not easily used by microorganisms. NH₄⁺-N removal was not adversely affected in the case of the addition of CS and CSF as the sole carbon source in comparison with SA. The TN removal efficiency was higher than 60% in all three SBRs, and highest in the CSF-SBR ((73.2±2.6)%). In addition, comparatively higher TP removal efficiency was observed in the CS-SBR ((74.1±10.8)%) and CSF-SBR ((69.6±7.3)%). The improvement in both nitrogen and phosphorus removal and the advantage of reducing the amount of cassava stillage to be treated in-situ, suggest that the use of cassava stillage or its anaerobic fermentation liquid as an external carbon source for BNR processes has considerable potential.

Table 5Treatment performance of three SBRs asdifferent carbon sources during the whole experimental period

menum periou			
Parameter	SA	CS	CSF
Effluent COD	25.5±4.5	36.2±3.5	27.1±6.3
(mg/L)			
COD removal	91.5±1.5	88.7±1.2	91.4±3.7
efficiency (%)			
Effluent NH4 ⁺ -N	0.6 ± 0.6	$0.4{\pm}0.4$	0.5 ± 0.6
(mg/L)			
NH4 ⁺ -N removal	96.0±4.0	97.3±2.5	96.5±3.0
efficiency (%)			
Effluent TN	18.7±1.7	13.8 ± 2.1	12.9±2.5
(mg/L)			
TN removal	62.6±3.5	72.4±3.2	73.2±2.6
efficiency (%)			
Effluent TP	2.50 ± 0.49	1.42 ± 0.64	1.82 ± 0.42
(mg/L)			
TP removal	58.3±8.1	74.1±10.8	69.6±7.3
efficiency (%)			

The values are expressed as mean \pm SD (*n*=3)

4 Conclusions

In this study, we investigated the performance of cassava stillage and its anaerobic fermentation liquid

used as the external carbon source for BNR in anaerobic-anoxic-aerobic SBRs. Based on the results, the following conclusions may be drawn:

1. Cyclic studies indicated that the cassava stillage and its anaerobic fermentation liquid affected transformations of COD, nitrogen, phosphorus, PHA, and glycogen of the BNR process. During the whole experimental period, the use of cassava stillage and its anaerobic fermentation liquid showed better TN removal efficiency $((72.4\pm3.2)\%$ and $(73.2\pm2.6)\%)$, respectively, than that of an equivalent amount of sodium acetate ((62.6 ± 3.5)%). EBPR occurred when the reactors were fed with cassava stillage and its anaerobic fermentation liquid as the carbon sources. Sludge analysis demonstrated that the ratio of DPAO/PAO reached 62.6% (Day 86) and 61.8% (Day 65) using cassava stillage and its anaerobic fermentation liquid as the sole carbon source, respectively.

2. Based on measurements, the NURs of the cassava stillage (5.49-5.99 g N/(kg MLVSS·h)) and its anaerobic fermentation liquid (6.63-6.81 g N/(kg MLVSS·h)) were similar to or higher than those of other industry wastewaters as external carbon sources.

3. The performance of the SBRs indicated that addition of the cassava stillage and its anaerobic fermentation liquid as the external carbon sources enhanced both TN and TP removals, while COD or NH_4^+ -N removal was not significantly affected.

4. The results presented indicate that cassava stillage and its anaerobic fermentation liquid could be used as potential external carbon sources for BNR processes.

Compliance with ethics guidelines

Fan BU, Xiang HU, Li XIE, and Qi ZHOU declare that they have no conflict of interest.

This article does not contain any studies with human or animal subjects performed by any of the authors.

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<u>中文概要</u>

- 题 目:木薯酒糟及其厌氧发酵液作为外加碳源强化城市 生活污水生物营养盐去除研究
- **目** 的:针对目前城市污水处理中普遍存在的进水碳源不 足引起的脱氮除磷效率不高的问题,而大量具有

很高碳氮比(C/N)和可生物降解有机成分的食品工业废水亟需处理的现状,本文考察了木薯酒精厂废水(木薯酒糟及其厌氧发酵液)分别作为 外加碳源资源化利用的可行性。

- **创新点:**对比考察了木薯酒糟及其厌氧发酵液分别作为碳 源对活性污泥系统脱氮除磷的影响,为处理低 C/N比城市污水寻求廉价优质的外加碳源提供理 论依据。
- 方 法:通过运行三组平行的厌氧/缺氧/好氧序批式活性 污泥反应器(SBR),以乙酸钠作为对比,考察 木薯酒糟及其厌氧发酵液作为碳源对活性污泥 系统碳、氮、磷变化规律的影响,并通过硝酸盐 利用速率(NUR)和厌氧-缺氧/好氧批次试验对 污泥特性进行分析。
- 结 **论:** (1) 在整个试验过程中,与乙酸钠作为碳源相 比,木薯酒糟及其厌氧发酵液在相同的运行条件 下取得了更高的总氮(TN)去除率,分别为 (72.4±3.2)%和(73.2±2.6)%,高于乙酸钠的(62.6± 3.5)%。NUR 试验结果表明,木薯酒糟及其厌氧 发酵液污泥的反硝化速率分别为 5.49~5.99 g N/ (kg MLVSS·h)和 6.63~6.81 g N/(kg MLVSS·h), 与其他研究中报道的食品工业废水的反硝化速率 相当或略高。(2)以木薯酒糟及其厌氧发酵液作 为碳源的系统发生了显著的反硝化聚磷现象,两 体系中的反硝化聚磷菌分别占总聚磷菌的 62.6% (86天)和61.8%(65天)。(3)以木薯酒糟 上清液及其厌氧发酵液作为碳源的生物营养盐 去除(SBR)系统均取得了良好稳定的脱氮除磷 效果。在 BNR 工艺中投加木薯酒糟及其厌氧碱 性发酵液作为外加碳源,不仅可以提高系统氮磷 去除效果,还能解决这部分废水的处理问题,是 一种很有潜力的替代碳源。
- 关键词: 生物营养盐去除; 外加碳源; 反硝化; 强化生物 除磷