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1 **Formation of aerobic granular sludge and the influence of the pH on**  
2 **sludge characteristics in a SBR fed with brewery/bottling plant**  
3 **wastewater**

4  
5 Short title: Aerobic granular sludge for the treatment of brewery wastewater

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24  
25 **Abstract**

26 A lab-scale sequencing batch reactor (SBR) was operated for 450 days to assess  
27 aerobic granule formation when treating brewery/bottling plant wastewater by  
28 consistent application of a feast/famine regime. The experiment was divided into  
29 three major periods according to the different operational conditions: (I) no pH  
30 control and strong fluctuations in organic loading rate (OLR)  
31 ( $1.18 \pm 0.25 \text{ kgCOD} \cdot (\text{m}^3 \cdot \text{day})^{-1}$ ), (II) pH control and aeration control strategy to  
32 reduce OLR fluctuations ( $1.45 \pm 0.65 \text{ kgCOD} \cdot (\text{m}^3 \cdot \text{day})^{-1}$ ) and (III) no pH control  
33 and stable OLR ( $1.42 \pm 0.18 \text{ kgCOD} \cdot (\text{m}^3 \cdot \text{day})^{-1}$ ). Aerobic granule formation was

34 successful after 80 days and maintained during the subsequent 380 days. The  
35 aerobic granular sludge was characterized by SVI<sub>5</sub> and SVI<sub>30</sub> values below  
36 60mL/g and dominated by granular, dense structures. An oxygen uptake rate  
37 (OUR) based aeration control strategy insured endogenous respiration at the end  
38 of the aerobic phase, resulting in stable SBR operation when the influent  
39 composition fluctuated. The quantitative polymerase chain reaction (qPCR)  
40 results show no significant enrichment of *Accumulibacter* or *Competibacter*  
41 during the granulation process. The 16S rRNA sequencing results indicate  
42 enrichment of other, possibly important species during aerobic granule formation  
43 while treating brewery wastewaters.

44 Keywords: Industrial wastewater ; phosphate accumulating organisms (PAO);  
45 glycogen accumulating organisms (GAO); Oxygen uptake rate (OUR); pH control

## 46 Introduction

47 The brewing sector is considered a major contributor to the European economy providing  
48 over 2 million jobs and producing approximately 40 million m<sup>3</sup> of beer per year (The  
49 Brewers of Europe 2016). Awareness and interest in the development of sustainable water  
50 and wastewater management, improving environmental performance and reducing water  
51 and energy usage is increasing within the sector. Globally great effort is put into  
52 decreasing water usage for beer production and reducing the carbon footprint by sourcing  
53 renewable energy (Modic et al. 2015). Research into sustainable wastewater treatment  
54 systems contributes to decreasing the environmental impact of the brewing industry and  
55 its waste and wastewater management.

56 Generally, brewery wastewater has a high organic matter content due to the presence of  
57 starch, sugar, volatile fatty acids (VFA), etc. which are typically easily biodegradable  
58 compounds (Driessen & Vereijken 2003). For this type of wastewater both biological  
59 anaerobic and aerobic wastewater treatment plants (WWTP) are used globally. The main  
60 advantages of anaerobic systems are the energy production (biogas) and the low sludge  
61 production and space requirements. To insure a higher degree of effluent quality an  
62 aerobic treatment is required. Both conventional activated sludge systems (CAS) and  
63 sequencing batch reactors (SBR) can be used for aerobic wastewater treatment (Driessen  
64 & Vereijken 2003; Wang 2007). To date, the aerobic granular sludge (AGS) technology  
65 in a SBR is accepted as a sustainable alternative to the CAS system for treatment of  
66 municipal wastewater and is already applied in many full-scale applications (Niermans et  
67 al. 2014; Pronk et al. 2015a). The technology provides strong improvements in  
68 settleability of the activated sludge (SVI<sub>5</sub>~SVI<sub>30</sub> < 100mL.g<sup>-1</sup>) due to the presence of  
69 large, dense granular structures. The excellent settling characteristics of aerobic granules  
70 implicate the possibility for higher biomass concentration resulting in the design of more  
71 compact bioreactors. In addition, full-scale references show more energy-efficient SBR  
72 operation and nutrient removal capacities (Niermans et al. 2014; Pronk et al. 2015a).  
73 More recently, lab-scale studies showed successful aerobic granulation while treating a  
74 variety of industrial wastewaters, some examples being: petrochemical (Caluwé et al.  
75 2017), malting (Schwarzenbeck et al. 2004), potato industry (Dobbeleers et al. 2017) and

76 brewery (Wang et al. 2007; Corsino et al. 2017) suggesting great potential of the AGS  
77 technology for industrial applications. Stable aerobic granule formation is based on the  
78 enrichment of phosphate accumulating (PAO) and glycogen accumulating (GAO)  
79 organisms. The selection for these bacteria is conducted by applying a feast/famine  
80 regime consisting of an anaerobic feeding during which conversion of VFA into  
81 intracellular storage polymers, i.e. poly-hydroxyalkanoates (PHA), occurs. The  
82 subsequent aerobic phase involves consumption of these polymers for microbial growth  
83 and maintenance. GAO obtain energy for anaerobic VFA uptake through intracellular  
84 glycogen degradation (Oehmen et al. 2007). The PAO metabolism typically shows  
85 anaerobic phosphate release as a result of intracellular poly-phosphate degradation,  
86 generating additional energy for VFA uptake. Intracellular phosphate accumulation  
87 (poly-P) takes place during aeration, resulting in a net phosphate removal. Previous  
88 research showed that pH has a direct influence on the PAO/GAO ratio, favouring PAO  
89 under alkaline conditions ( $\text{pH} > 7.2$ ). This is explained by the fact that additional energy  
90 is required for VFA conversion when pH increases. Stored poly-P provides a secondary  
91 energy source favouring PAO growth over GAO at higher pHs (Filipe et al. 2001). The  
92 influent P/COD ratio as well as the temperature and carbon source are considered to be a  
93 less decisive factor regarding the competition between PAO and GAO. To our  
94 knowledge, research concerning the influence of the pH on the microbial composition has  
95 so far only been conducted using synthetic wastewater (Oehmen et al. 2007; Weissbrodt  
96 et al. 2013).  
97 The objective of this study consists out of (1) stable aerobic granule formation and long  
98 term maintenance while insuring good effluent quality and (2) investigation of the  
99 influence of the pH on the selection of GAO and PAO while treating a complex industrial  
100 wastewater, i.e. brewery/bottling plant wastewater.

## 101 **Material and methods**

### 102 *Reactor set-up*

103 A fully automated lab-scale SBR with a working volume of 11L, a diameter of 24 cm  
104 and a H/D ratio of 1.7 was operated at room temperature (18-22°C) for 450 days. A  
105 mechanical stirrer (100rpm) was used to keep sludge in suspension during mixed phases  
106 of the SBR operation. Process operation of the SBR was controlled by a Siemens PLC.  
107 Process settings and visualisation of the online measurements of the dissolved oxygen  
108 concentration (DO), pH, oxidation reduction potential (ORP) and conductivity were  
109 controlled by LabView™ (National Instruments, Austin – Texas, USA).

### 110 *Industrial wastewater and seed sludge*

111 Wastewater from a local brewery/bottling WWTP was regularly collected and instantly  
112 stored at 4°C to minimize COD degradation. To insure sufficient nutrient availability for  
113 microbial growth,  $\text{NH}_4\text{Cl}$  and  $\text{K}_2\text{HPO}_4$  were added to the raw wastewater providing a  
114 COD:N:P ratio of 100:2:0.2. During the first 250 days of the experiment strong  
115 fluctuations in wastewater composition were observed due to the absence of an  
116 equalization tank at the existing WWTP. Periodically, high strength brewery wastewater  
117 with a COD concentration up to 13540  $\text{mg O}_2\cdot\text{L}^{-1}$  is added to the medium strength  
118 wastewater from the bottling plant with a COD concentration of 1200  $\text{mg O}_2\cdot\text{L}^{-1}$ . On

119 day 252, an equalization tank was taken into service at the full scale site resulting in a  
 120 slightly more constant wastewater composition. Table 1 gives an overview of the  
 121 wastewater composition before and after the presence of the equalization tank.

122 Table1: Wastewater composition before and after presence of the equalization tank –  
 123 incl.  $\text{NH}_4\text{Cl}$  and  $\text{K}_2\text{HPO}_4$  dosage

	Equalization tank absent						Equalization tank present					
	n	Min.	Max.	Av.	Stdev	CV(%)	n	Min.	Max.	Av.	Stdev	CV(%)
CODt (mg $\text{O}_2\cdot\text{L}^{-1}$ )	37	2195	13540	6846	2461	35.94	20	1690	6495	4037	1476	36.55
CODs (mg $\text{O}_2\cdot\text{L}^{-1}$ )	37	2025	9940	6053	2005	33.13	20	1540	5010	3606	1294	35.88
TOC (mg $\text{C}\cdot\text{L}^{-1}$ )	26	365	2908	1580	562	35.6	13	282	1444	806	432	53.6
$\text{NH}_4^+$ - N (mg $\text{N}\cdot\text{L}^{-1}$ )	37	5.74	198	103	45.8	44.3	22	4.00	148	76.7	49.2	64.1
$\text{PO}_4^{3-}$ - P (mg $\text{P}\cdot\text{L}^{-1}$ )	38	2.0	100	19	17	89	22	1.7	19	9.2	3.7	41

124 Seed sludge from the same local brewery/bottling WWTP was used to inoculate the lab-  
 125 scale SBR. The full-scale installation consists of a side-stream membrane bioreactor  
 126 (MBR) where high sludge concentrations between  $12\text{-}15\text{g}\cdot\text{L}^{-1}$  are used. Initial MLSS  
 127 and MLVSS concentrations were  $14.0$  and  $12.6\text{g}\cdot\text{L}^{-1}$ , respectively. The morphology  
 128 was characterised by a filamentous/flocculent structure with  $\text{DSVI}_{30}$  of  $285\text{mL}\cdot\text{g}^{-1}$   
 129 indicating very poor settling capacities. The full-scale installation is operated at a  
 130 sludge retention time (SRT) of approximately 60 days and a highly variable food to  
 131 mass ratio (F/M) varying from 0.01 to  $0.30\text{kg COD}\cdot(\text{kg MLSS}\cdot\text{day})^{-1}$ .

### 132 **Reactor operation**

133 The lab-scale SBR was operated for approximately 450 days which were divided into  
 134 three major periods due to significant changes in operational conditions. Table 2 gives  
 135 an overview of the different experimental periods. During period I and III the total cycle  
 136 length of the SBR was fixed at 12 and 8h, respectively, consisting of following phases:  
 137 an aerated iddle phase (60-30min.), followed by a short non-mixed, non-aerated feed  
 138 preparation period (10min) insuring anaerobic conditions. Subsequently, a non-mixed  
 139 anaerobic pulse feed (10min) was imposed during which 1L of brewery/bottling  
 140 wastewater was fed on top of the SBR resulting in a constant volume exchange ratio  
 141 (VER) of 9%. Thereafter, an extended mixed anaerobic phase was prolonged from 90  
 142 up to 120min on day 25 to promote anaerobic COD storage. The duration of the  
 143 subsequent aerobic phase was fixed at 500-545 and 225min during period I and III,  
 144 respectively. Aeration times were only adjusted to insure a constant total cycle length  
 145 due to shortening of the settling phase from 60 down to 15min. To finalize, a  
 146 withdrawal phase (5min) is imposed for effluent discharge. During period II, all phases

147 remained unchanged with the exception of (1) the settling phase which was reduced to  
 148 10min and (2) the aeration phase during which an aeration control strategy was applied  
 149 as described in the following paragraph. At the end of period II, i.e. from day 319 to  
 150 346, the reactor was mixed without feeding. This was in agreement with the period  
 151 during which brewing and bottling activities were ceased and therefore no wastewater  
 152 was produced. Due to the lack of wastewater production, it was of importance to  
 153 determine the resistance of the AGS to non-active periods.

154 Table2: Overview of experimental periods

		Period I	Period II	Period III
Day		1-128	129-325	325-450
<b>Full-scale WWTP</b>	Equalization tank	Absent	Absent/present	Present
<b>Lab-scale SBR</b>	Aeration Strategy	Fixed duration DO: 1-3 mgO <sub>2</sub> .L <sup>-1</sup>	OUR regulation DO: 1-3 mgO <sub>2</sub> .L <sup>-1</sup>	Fixed duration DO: 1-3 mgO <sub>2</sub> .L <sup>-1</sup>
	Total cycle time	12h	5.3-15.9h	8h
	Aeration time	500-545min	120-720min	225min
	Settling time	60 – 15min	10min	10min
	pH control	No pH control	pH control at 7.0±0.2	No pH control

155 During period I and II , SRT was kept constant at 30 days by removing a volume of  
 156 330mL excess sludge every day. During period III no sludge was removed.

### 157 *Aeration control strategy and pH control*

158 Online oxygen uptake rate (OUR) calculations were conducted during the aerobic  
 159 phase. Aeration was provided using an automatic on-off regulation between two DO  
 160 set-points. When the maximum DO setpoint was reached, aeration stopped and oxygen  
 161 levels dropped due to oxygen usage by the microorganisms, resulting in a decreasing  
 162 linear DO versus time curve. From this data, the OUR is automatically calculated as the  
 163 absolute value of the slope from the linear curve between the two DO set-points. Due to  
 164 the combination of strong variations in organic load of the wastewater and a constant  
 165 volume exchange ratio during period I, feeding of low strength wastewater resulted in  
 166 prolonged famine conditions characterised by constant and low specific OUR (sOUR)  
 167 measurements ( $< 2 \text{ mgO}_2 \cdot (\text{gMLVSS} \cdot \text{h})^{-1}$ ). By introducing an aeration control strategy  
 168 during period II, sOUR values varied between  $2\text{-}5 \text{ mgO}_2 \cdot (\text{gMLVSS} \cdot \text{h})^{-1}$  at the end of  
 169 the aerobic phase so prolonged aeration was avoided. Aeration continued until the  
 170 occurrence of an inflection point in the OUR curve indicating the degradation of all  
 171 internal and external carbon. Therefore, a minimum OUR value was set at  
 172  $20 \text{ mgO}_2 \cdot (\text{L} \cdot \text{h})^{-1}$ , corresponding with sOUR values between  $2\text{-}5 \text{ mgO}_2 \cdot (\text{gMLVSS} \cdot \text{h})^{-1}$ .  
 173 When reaching the aimed set-point, the next cycle phase was initiated.

174  
 175 The pH in the full-scale installation treating the brewery/bottling wastewater typically  
 176 varies between 8.0-8.5 which is considered to influence GAO growth negatively and  
 177 therefore is expected to favour PAO growth when applying anaerobic feeding followed  
 178 by an aerobic famine period (Filipe et al. 2001; Oehmen et al. 2007; Weissbrodt et al.  
 179 2013). The composition of the used brewery wastewater requires addition of phosphate  
 180 to insure sufficient nutrient availability for microbial growth. Since PAO are known to

181 accumulate excess phosphate intracellularly but only limited amounts are present in the  
182 wastewater, enrichment of PAO could be influenced negatively and therefor complicate  
183 enrichment of slow growing micro-organisms. The influence of the pH on sludge  
184 characteristics is investigated by controlling pH at  $7.0 \pm 0.2$  during period II. The  
185 correction was applied by the automatic addition of a 1M HCl solution.

### 186 *Analytical methods*

187 Total and soluble chemical oxygen demand concentrations, i.e. COD<sub>t</sub> and COD<sub>s</sub>, were  
188 measured with tests tubes (HI 93754A-25 and HI 93754B-25) from Hanna Instruments  
189 (Temse, Belgium). Also NH<sub>4</sub><sup>+</sup>-N (HI 93715-01), NO<sub>2</sub><sup>-</sup>-N (HI 93707-01), NO<sub>3</sub><sup>-</sup>-N (HI  
190 93766-50) and PO<sub>4</sub><sup>3-</sup>-P (HI 93717-01) concentrations were measured using test kits  
191 from Hanna Instruments. Sulphate concentration of the influent was measured using a  
192 LCK153 test kit from Hach (Mechelen, Belgium). Dissolved Organic Carbon  
193 concentrations (DOC) were measured using a Sievers InnovOx Laboratory Total  
194 Organic Carbon Analyzer (GE Analytical instruments). Before measurement of the  
195 DOC, COD<sub>s</sub>, NH<sub>4</sub><sup>+</sup>-N, NO<sub>2</sub><sup>-</sup>-N, NO<sub>3</sub><sup>-</sup>-N and PO<sub>4</sub><sup>3-</sup>-P concentration, samples were  
196 filtered using 1.2µm glass microfiber filters (VWR International, Belgium). Effluent  
197 suspended solids (SS) concentrations were calculated by the ratio of the difference  
198 between effluent COD<sub>t</sub> and COD<sub>s</sub> concentrations to the conversion factor of 1.42 as  
199 presented by Eckenfelder et al. (2009).

200

201 All sludge samples were taken at the end of the aerobic phase to insure endogenous  
202 conditions for all analyses. The sludge volume (SV) and mixed liquor (volatile)  
203 suspended solids concentration (ML(V)SS) were determined according to the standard  
204 methods described by (AHA/AWWA/WEF 1998). Particle size distribution by volume,  
205 DV, measurements of the sludge mixture were performed using a Malvern Mastersizer  
206 3000 (Malvern, UK). Settings: refractive index: 1.52, absorption index: 1.0, particle  
207 size: non-spherical, dispersant: water, both blue and red measurement duration: 30sec.,  
208 number of measurements: 10, obscuration: 8-12%, stirrer speed: 600rpm. To investigate  
209 evolution of the sludge morphology, microscopic analysis was performed using a  
210 MOTIC BA310 microscope (Opti-service, Belgium).

### 211 *In-situ cycle measurement during SBR operation*

212 In order to evaluate anaerobic DOC uptake and phosphate release, associated with the  
213 presence of GAO and PAO bacteria, in-situ cycle measurements were performed during  
214 multiple SBR cycles. Grab samples were taken every 30min. during the anaerobic phase  
215 and at the end of the SBR cycle in order to obtain DOC and phosphate profiles.

$$216 \quad \text{Anaerobic DOC removal (\%)} = 100 \times \frac{DOC_{t_0} - DOC_{t_1}}{DOC_{t_0} - DOC_{t_e}}$$

217 With  $t_0$  just after anaerobic feed,  $t_1$  at the end of the anaerobic phase and  $t_e$  at the end of  
218 the SBR cycle. Grab samples were filtered using glass microfibre filters before  
219 analytical measurements were performed.

### 220 *DNA extraction*

221 To investigate the evolution in abundance of PAO and GAO and perform 16S rRNA

222 gene sequencing analysis, sludge samples were taken in triplicate on different days  
223 during the experiment and preserved at -80°C. Subsequently, DNA extractions were  
224 performed according to the method described by Mcilroy et al. (2008). A Qubit 3.0  
225 Fluorometer (ThermoFisher Scientific, Massachusetts) was used for quantification of  
226 the resulting DNA concentrations.

### 227 ***Molecular quantification by qPCR***

228 Target genes were quantified as described in (Caluwé et al. 2017) by the use of  
229 PAO541f/PAO846r primers for PAO (16S rRNA *Candidatus Accumulibacter*  
230 *phosphatis*), GAOQ989f/GAM1278r primers for GAO (16S rRNA *Candidatus*  
231 *Competibacter phosphatis*) and Universal1055f/Universal1392r primer for the total  
232 amount of bacteria (16S rRNA Universal bacteria). From the measured DNA  
233 concentrations, the amount of target cells per g MLVSS was calculated.

### 234 ***Microbial community composition by 16S rRNA gene amplicon sequencing***

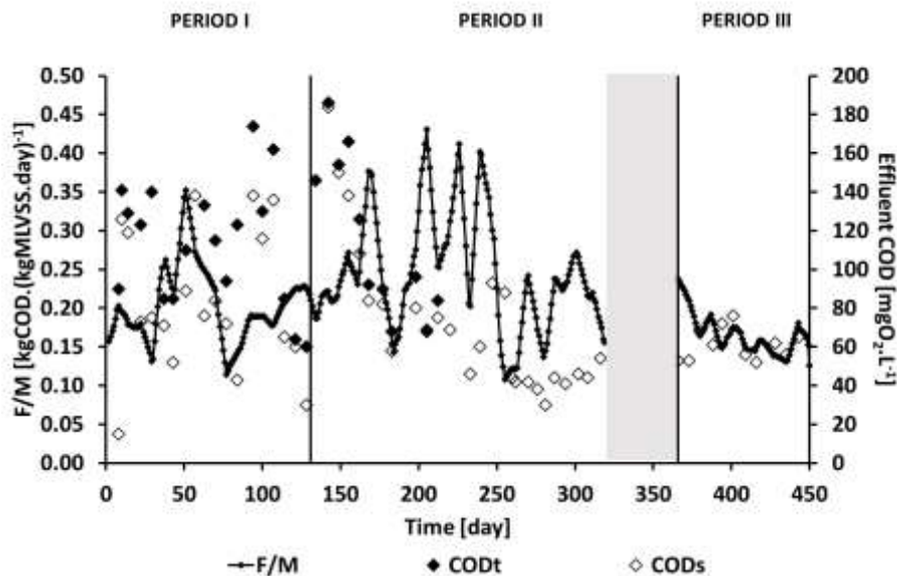
235 The 16S rRNA gene amplicon sequencing analysis was performed on two sludge  
236 samples, i.e. seed sludge and a grab sample taken at the end of operational period II  
237 (day 302). Amplicons targeting the V4 region of the 16S rRNA gene were generated  
238 with barcoded primers (IDT), and Phusion High-Fidelity DNA polymerase (Thermo  
239 Scientific) as described by Kozich et al. (2013). PCR products were purified using the  
240 SequalPrep Normalization plate kit (Invitrogen), and pooled. The resulting library was  
241 further purified by gel extraction using NucleoSpin Gel and PCR Clean-up (Macherey  
242 Nagel), and diluted to obtain a 2nM library. Amplicon sequencing was carried out on a  
243 Illumina Miseq system at the Centre for Medical Genetics (Edegem, Belgium) with the  
244 MiSeq Reagent Kit v2 (Illumina).  
245 The obtained paired-end reads were processed with the UPARSE pipeline (Edgar 2013).  
246 Taxonomy predictions of the OTU sequences was done with MiDAS (version 2.1) as the  
247 reference database: manually curated SILVA 16S rRNA taxonomy (release 1.23 Ref  
248 NR99) that proposes a name for all the abundant genus-level taxa present in activated  
249 sludge, anaerobic digesters and influent wastewater (Mcilroy et al. 2015).

## 250 **Results and discussion**

### 251 ***Reactor performance***

252 A constant applied volume exchange ratio (VER) of 9% and strong variations of the  
253 influent COD concentration resulted in distinct variations in organic loading rate (OLR)  
254 during period I. Low strength wastewater resulted in prolonged famine conditions  
255 causing possible unnecessary aeration. Since aeration is one of the main operational  
256 costs in full-scale installations, excess aeration should be avoided when assessing  
257 industrial applicability of the technology (Niermans et al. 2014). During period I, the  
258 average OLR was  $1.18 \pm 0.25 \text{ kgCOD} \cdot (\text{m}^3 \cdot \text{day})^{-1}$  resulting in strong fluctuation of the  
259 F/M ratio between 0.11-0.36  $\text{kgCOD} \cdot (\text{kgMLVSS} \cdot \text{day})^{-1}$ . During period II, an aeration  
260 control strategy based on the oxygen uptake rate (OUR) was introduced to reduce  
261 fluctuation of the OLR resulting in an average OLR of  $1.45 \pm 0.65 \text{ kgCOD} \cdot (\text{m}^3 \cdot \text{day})^{-1}$ .

262 After the equalization tank was taken into service at the full-scale WWTP (day 252), a  
 263 slightly more constant wastewater composition was observed so again fixed aeration  
 264 times were applied during period III (tables 1 and 2). As a result OLR and F/M ratio  
 265 varied between 1.1-1.9 kgCOD.(m<sup>3</sup>.day)<sup>-1</sup> and 0.13-0.25 kgCOD.(kgMLVSS.day)<sup>-1</sup>,  
 266 respectively. Similar OLR were applied by Corsino et al. (2017) resulting in stable  
 267 granule formation. Figure 1 shows the evolution of the F/M applied during operation  
 268 and the results of the total and soluble effluent COD concentration throughout the  
 269 experiment.



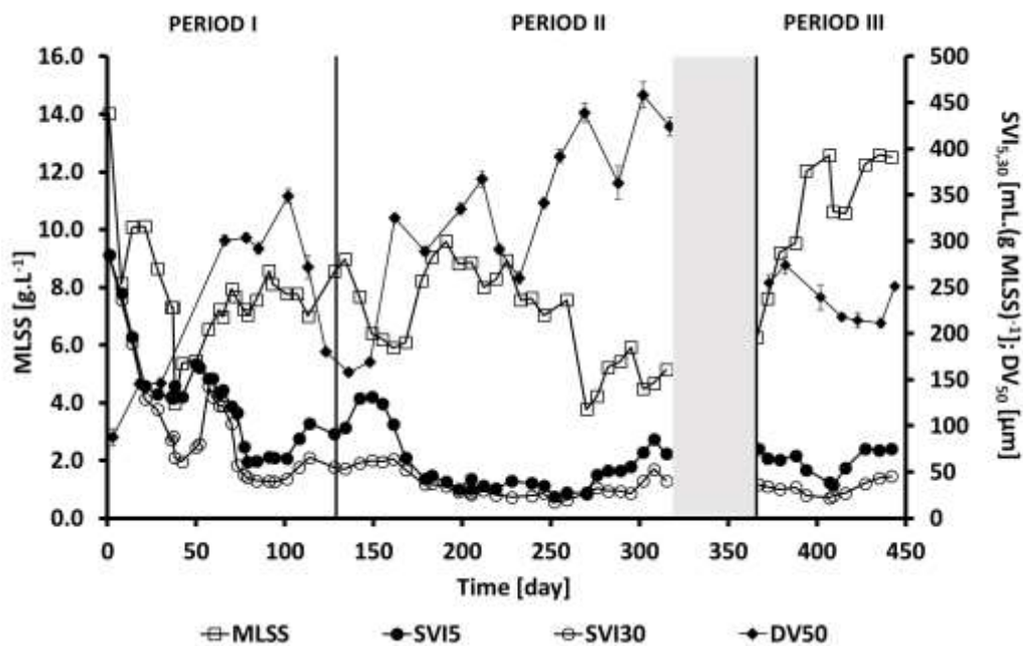
270  
 271 Figure 1 Evolution of the F/M and effluent CODs concentration during the experiment  
 272 (grey zone: 30days mixed without feeding)

273 Adjustment of the seed sludge to the new operation strategy and the distinct variations  
 274 in OLR caused poor effluent quality with CODt concentrations up to 141mgO<sub>2</sub>.L<sup>-1</sup>  
 275 during start-up. Thereafter, CODt and CODs concentrations differed strongly indicating  
 276 wash-out of suspended solids up to 56 mg SS.L<sup>-1</sup> on day 84. However, at the end of  
 277 period I, good effluent quality was already achieved. From day 129, i.e. start of period  
 278 II, an aeration control strategy was introduced to handle strong fluctuations of the  
 279 influent COD concentration and additionally pH control was imposed. During the first  
 280 40 days of period II, the presence of suspended solids in the effluent decreased with an  
 281 average of 8.1±5.7 mg SS.L<sup>-1</sup>, far below the discharge limit of 60 mg SS.L<sup>-1</sup>, so  
 282 thereafter only effluent CODs concentration was measured. After a possible adjustment  
 283 period of 40 days, stable effluent quality well below the Flemish discharge limit (125  
 284 mgO<sub>2</sub>.L<sup>-1</sup>) was established. Low effluent COD concentrations were maintained  
 285 throughout the remainder of the experiment even though strong variations in applied  
 286 F/M still occurred and pH control was switched off again during period III. Due to the  
 287 lack of wastewater production, the resistance of the AGS to non-active periods was  
 288 investigated. Therefore, a 30 day period of mixing without feeding was imposed as a  
 289 strategy to preserve AGS. Results in Figure 1 show high effluent quality and therefore  
 290 high COD removal efficiencies up to 99% immediately after reinitiating normal SBR  
 291 operation at the start of period III. Therefore, mixing without feeding can be assumed as  
 292 an appropriate strategy to preserve AGS during periods where no wastewater is  
 293 produced. Throughout the experiment, the CODs removal efficiency varied between 95-  
 294 99%. Results show improvement in effluent quality as granule formation evolved and

295 large structures started to dominate the sludge mixture (see below). An OUR based  
 296 aeration control strategy is proposed to result in proper famine conditions, i.e. sOUR  
 297 values  $< 5\text{mgO}_2\cdot(\text{gMLVSS}\cdot\text{h})^{-1}$ , at the end of the aeration period. Consequently, stable  
 298 SBR operation and effluent quality were obtained.

### 299 *Sludge characteristics and morphology*

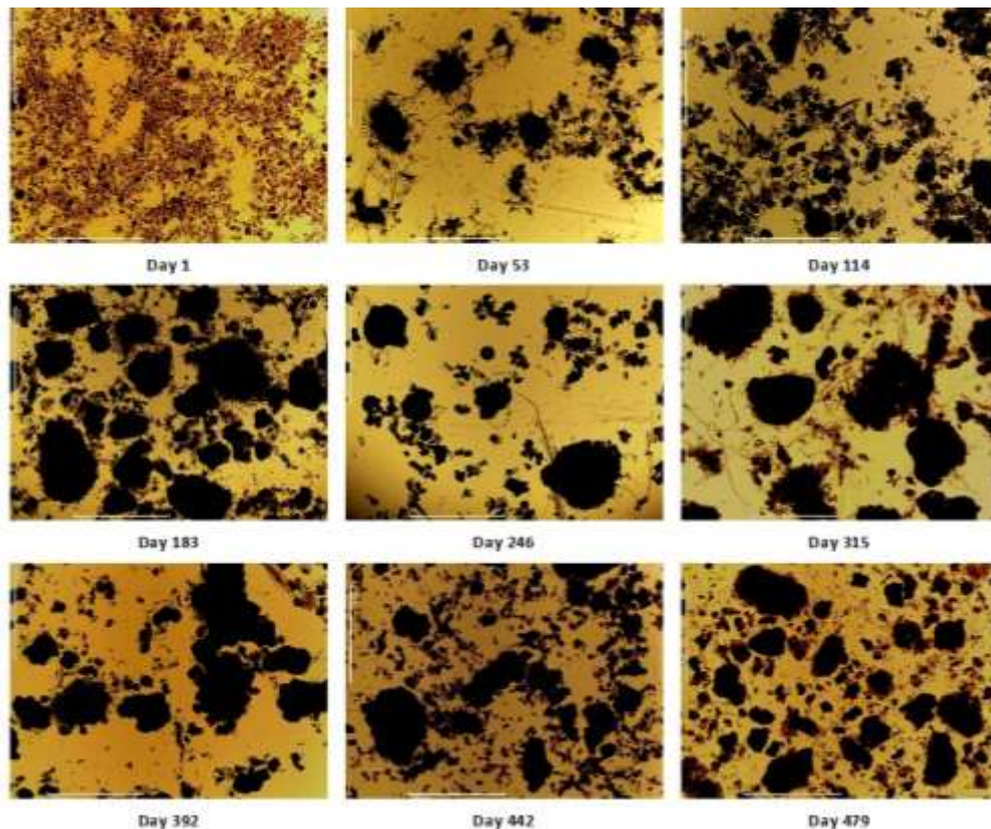
300 By applying a feast/famine regime consisting of an anaerobic pulse feeding followed by  
 301 prolonged anaerobic and aerobic phase, sludge characteristics changed significantly  
 302 during the experiment. Figure 2 shows the evolution of the MLSS, the median particle  
 303 size by volume ( $DV_{50}$ ) and the SVI after 5 and 30 minutes of settling.



304  
 305 Figure 2 Evolution of the MLSS, SVI<sub>5</sub> and SVI<sub>30</sub> and DV<sub>50</sub> during the experiment  
 306 (grey zone: 30days mixed period without feeding)

307 The objective of this study was to stimulate aerobic granule formation while respecting  
 308 the required effluent quality. Therefore, a long initial settling time (60min.) was  
 309 imposed. Due to the extremely poor settling characteristics of the membrane bioreactor  
 310 (MBR) seed sludge, i.e.  $DSVI_{30}=285\text{mL}\cdot\text{g}^{-1}$ , which was dominated by flocculent and  
 311 filamentous sludge structures (Figure 3), biomass washout occurred during start-up  
 312 resulting in a decrease of the MLSS concentration. The second drop of the MLSS  
 313 concentration (day 38) was caused by removing half of the sludge volume for start-up  
 314 of a second experiment. The MLVSS/MLSS ratio remained stable during the entire  
 315 experiment at  $92\pm 3\%$ . A rapid increase of  $DV_{50}$  is observed from  $87.5\pm 9.4\mu\text{m}$  up to  
 316  $349\pm 7.6\mu\text{m}$  during period I. Concurrently, strong improvement of the sludge settling  
 317 characteristics were observed:  $SVI_5$  and  $SVI_{30}$  decreased from  $285\text{mL}\cdot\text{gMLSS}^{-1}$  down to  
 318 66 and  $40\text{mL}\cdot\text{gMLSS}^{-1}$ , respectively. These results correspond to the significant reduced  
 319 presence of filamentous and flocculent sludge structures and the development to a more  
 320 dense granule morphology (Figure 3). From day 102, decreased particle size and a  
 321 temporary reduction of sludge settling capacities was observed. Nevertheless, results  
 322 show successful aerobic granule formation after approximately 80 days of SBR  
 323 operation. Similar results were observed by Wang et al. (2007) where mature granule

324 formation also fed with brewery wastewater was observed after nine weeks of  
325 operation.  
326 After introducing the pH control and the OUR based aeration strategy (period II), the  
327 effluent quality and settling characteristics temporary worsened. The sudden lowering  
328 of the  $pH_{max}$  from 8.1 to 7.2 may have created an excessive change in environmental  
329 conditions. After a 30 days adjustment period, stable SBR operation was recovered.  
330 Thereafter, significant increase of  $DV_{50}$  was observed from  $180 \pm 3.8 \mu m$  up to  
331  $424 \pm 10 \mu m$  on day 317. In addition, large, dense granules dominated the biomass  
332 mixture (Figure 3) while settling capacities improved further to stable  $SVI_5$  and  $SVI_{30}$   
333 values of 35 and  $30 \text{ mL.gMLSS}^{-1}$ , respectively. Subsequently, 30 days of mixing  
334 without feeding was applied. When SBR operation was reinitiated (period III), the  
335 median granule size was smaller which may indicate that this period had a slightly  
336 negative influence on granule stability. However, the median granule size remained  
337 stable between 211 and  $255 \mu m$  during the subsequent 84 days indicating good  
338 resistance of the AGS to periods without feeding. Sludge settling characteristics were  
339 good during this period with a final  $SVI_{30}$  value of  $46 \text{ mL.gMLSS}^{-1}$  while MLSS  
340 concentrations were as high as  $12.5 \text{ g.L}^{-1}$ . At this point biomass consisted of a mixture of  
341 large and small granules.  
342  
343 Considering the overall duration of the experiment, our results indicate the good  
344 resistance to variable operational conditions to which the aerobic granular sludge was  
345 exposed, such as significant fluctuations in F/M ratio, the variations in pH (with and  
346 without control) and the 30 day period of mixing without feeding. These variable  
347 conditions had no negative influence on carbon removal efficiencies, settling  
348 characteristics and the overall sludge morphology. Only the sudden lowering of the pH  
349 at the start of period II seemed to negatively influence both the sludge settling capacities  
350 and effluent quality. Generally, all results indicate that aerobic granule formation was  
351 successful and long term stability was maintained under changing operational  
352 conditions during the treatment of brewery wastewater.



353  
354

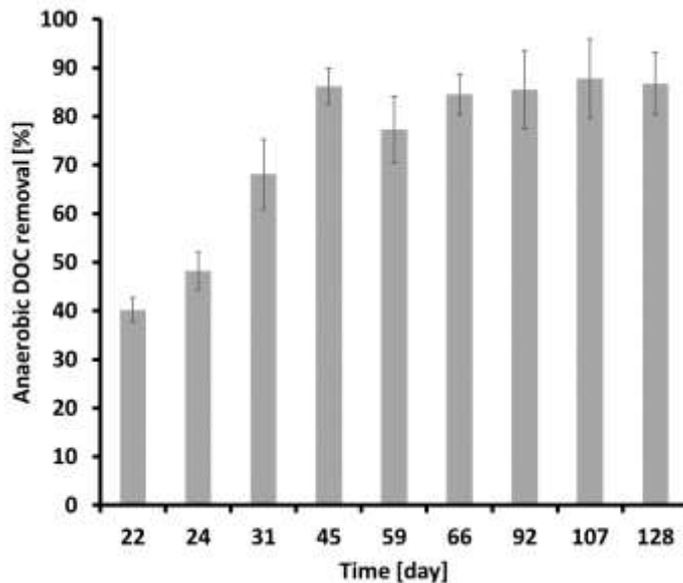
Figure 3 Microscopic analysis of sludge morphology (calibration curve: 1000 $\mu$ m)

355 It should be mentioned that minor filamentous outgrowth was constantly present at the  
 356 granule surface (Figure 3). This corresponds to previous findings on the use of more  
 357 complex wastewaters containing large organic compounds (i.e. starch, polysaccharides,  
 358 etc.) and particulate matter, which are also expected to be present in brewery  
 359 wastewater. Slowly biodegradable compounds are found to adsorb on the granule  
 360 surface where hydrolytic conversion take place. Hydrolytic products diffuse into the  
 361 granule structure and subsequently, anaerobic conversion into storage polymers can take  
 362 place. Depending on the anaerobic hydrolysis rate and duration of the anaerobic phase,  
 363 aerobic hydrolysis may occur at the granule surface which is suggested to favour  
 364 filamentous growth (de Kreuk et al. 2010; Pronk et al. 2015b). In contrast to our  
 365 findings, no filaments and larger granules were observed by Wang et al. (2007) and  
 366 Corsino et al. (2017) using brewery wastewater as substrate for granule formation. In  
 367 both studies, influent was fed at the bottom of the reactor into the sludge bed to insure  
 368 high substrate concentrations to stimulate anaerobic carbon uptake. In our study, it was  
 369 decided to feed influent on top of the SBR since this feeding strategy is believed to be  
 370 more easily applicable. Other possible reasons for these varying results may be found in  
 371 the high initial hydraulic selection pressure applied by Wang et al. (2007) (18 min of  
 372 settling) and Corsino et al. (2017) (4.5min of settling). In our study, the initial hydraulic  
 373 selection was low due to the 1h settling time, so complete initial wash-out of the  
 374 biomass was avoided. The difference in feeding strategy and/or hydraulic selection  
 375 pressure may have had an impact on the overall sludge composition and morphology. It  
 376 is however unclear which factor is more decisive. Since the wastewater composition  
 377 was highly variable, the 2h anaerobic phase may have periodically been insufficient to  
 378 hydrolyse all carbon compounds, resulting in possible aerobic hydrolysis which may  
 379 explain the minor but constant presence of filaments.

380 To summarize, without feeding at the bottom of the reactor and without significant  
381 hydraulic selection pressure, aerobic granule formation and long-term maintenance was  
382 successful. Consistent submission of a fixed anaerobic feast phase, followed by a  
383 dynamically regulated (OUR based) aerated famine phase, seems to be a promising  
384 strategy to maintain a stable aerobic granular sludge system using wastewater with a  
385 highly fluctuating composition.

### 386 *Evolution of the anaerobic DOC removal*

387 Multiple in-situ cycle measurements were performed during the SBR cycle in order to  
388 evaluate the anaerobic DOC removal.



389 Figure 4 Evolution of the anaerobic DOC uptake during period I of SBR operation  
390  
391

392 The gradual increase of anaerobic DOC uptake during period I is shown in Figure 4.  
393 Initially, only  $41 \pm 3\%$  of the DOC was removed from the bulk liquid during the 90min.  
394 anaerobic phase. At that point, sludge mixture was dominated by flocculent and  
395 filamentous structures and settling characteristics were poor ( $SVI_{5,30} > 100 \text{ mL/g}$ ). To  
396 promote anaerobic DOC uptake, the duration of the anaerobic phase was increased up to  
397 120 min. Consequently, anaerobic DOC removal rapidly increased up to  $87 \pm 7\%$   
398 suggesting strong improvement in anaerobic conversion of biodegradable substrate into  
399 storage polymers. Additional analysis of the evolution of the internal PHA content is  
400 necessary to confirm these findings. During period II, anaerobic DOC removal  
401 increased further up to  $94 \pm 7\%$  while sludge structures developed to mature granules.  
402 During period III the percentage of anaerobic substrate removal from the bulk liquid  
403 remained stable with an average of  $97 \pm 2\%$ . The high percentage of DOC removal  
404 during the anaerobic phase indicates that substrate was converted into storage polymers,  
405 resulting in a DOC removal from the bulk liquid, creating stable operational conditions  
406 favouring growth of slow growing micro-organisms associated with aerobic granulation  
407 and additionally avoiding filamentous outgrowth.

## 408 *Influence of pH on selection of GAO and PAO*

409 The pH is suggested to be one of the main factors influencing PAO/GAO competition.  
410 Due to the presence of an additional energy source (poly-P) to convert VFA into  
411 intracellular storage polymers, PAO outcompete GAO when  $\text{pH} > 7.2$  (Filipe et al.  
412 2001). Similar results, showing direct selection of PAO under slightly alkaline  
413 conditions, were obtained by Weissbrodt et al. (2013). Caluwé et al. (2017) showed an  
414 increased PAO activity and abundance treating petrochemical wastewater with a COD/P  
415 ratio of 300 for the formation of aerobic granular sludge. In this study the average pH in  
416 the reactor was 7.8 which is known to stimulate PAO growth. Remarkably, even though  
417 no excess phosphate was present in the wastewater, minor enrichment of PAO was  
418 observed. These results implicate that other factors than COD/P ratio, like pH, may play  
419 a more decisive role in the selection for PAO. When treating brewery wastewater,  
420 maximum pH values ( $\text{pH}_{\text{max}}$ ) are typically obtained at the end of the aeration phase due  
421 to  $\text{CO}_2$  stripping. No pH control was applied during period I, resulting in an average  
422  $\text{pH}_{\text{max}}$  of  $8.1 \pm 0.4$ . With this knowledge, it was suggested that growth of GAO could be  
423 affected due to unfavourable conditions which consequently may have a negative  
424 impact on the granule formation. On day 30, in-situ measurements of DOC and  
425 phosphate concentrations were conducted during the anaerobic phase. The anaerobic  
426 DOC removal was  $68 \pm 7\%$  while no phosphate release was observed. Subsequently on  
427 day 128, anaerobic DOC removal had increased to  $87 \pm 7\%$  while anaerobic phosphate  
428 release was also observed from  $4.8 \text{ mgP.L}^{-1}$  after feed up to  $6.5 \text{ mgP.L}^{-1}$  at the end of  
429 the anaerobic phase. A yield of phosphate release to carbon uptake ( $Y_{\text{p/C,An}}$ ) was  $0.008$   
430  $\text{Pmol}_{\text{PO}_4}.\text{Cmol}_{\text{DOC}}^{-1}$  indicating minor enrichment of PAO since start-up of the  
431 experiment. Since anaerobic DOC removal increased significantly, enrichment of  
432 carbon-storing microorganisms is assumed to be established.

433  
434 Molecular analysis by qPCR was performed to quantify enrichment of PAO  
435 (*Candidatus Accumulibacter phosphatis*) and GAO (*Candidatus Competibacter*  
436 *phosphatis*), commonly associated with aerobic granule formation. A slight increase of  
437 PAO abundance was observed during period I to  $0.87 \pm 0.12\%$  confirming the findings  
438 above regarding a minor increase of the  $Y_{\text{p/C,An}}$  ratio due to some PAO enrichment.  
439 Abundance of GAO showed no increase, even a slight decrease from  $2.91 \pm 0.21\%$  down  
440 to an average abundance of  $2.00 \pm 0.25\%$  during the first experimental period. At this  
441 stage, no enrichment of the common PAO or GAO group was observed while the  
442 granulation process evolved. During period II, pH control was introduced using a 1M  
443 HCl solution to lower the pH resulting in an average  $\text{pH}_{\text{max}}$  of  $7.3 \pm 0.1$  to influence  
444 microbial competition to stimulate GAO growth since anaerobic carbon removal is less  
445 energy consuming under these conditions. In addition it was assumed PAO activity  
446 would not occur. In contrast to our expectations, the abundance of GAO did not  
447 increase, i.e.  $2.46 \pm 0.03\%$  during period I and  $2.44 \pm 0.20\%$  during period II. Introducing  
448 the pH control did not seem to enhance enrichment of this specific group of GAO.  
449 However, granulation remained stable suggesting good selection for granule forming  
450 organisms other than the well-known PAO or GAO. In-situ measurements on day 317  
451 showed that phosphate release occurred to a lesser extent from  $1.6 \text{ mgP.L}^{-1}$  to  $2.4$   
452  $\text{mgP.L}^{-1}$  at the end of the anaerobic phase while  $94 \pm 7\%$  of DOC was anaerobically  
453 removed resulting in a lower  $Y_{\text{p/C,An}}$  of  $0.006 \text{ Pmol}_{\text{PO}_4}.\text{Cmol}_{\text{DOC}}^{-1}$ . The qPCR results  
454 confirm the nearly complete absence of the PAO fraction during this period with an  
455 average PAO abundance of  $0.07 \pm 0.03\%$ . After switching off pH control in period III,  
456 the GAO abundance decreased down to  $1.42 \pm 0.22\%$ . A slight increase in anaerobic

457 phosphate release from 2.2 mgP.L<sup>-1</sup> to 4.4 mgP.L<sup>-1</sup> at the end of the anaerobic phase  
458 was once more observed while 99±7% of DOC was taken up resulting in a Y<sub>p/C,An</sub> of  
459 0.009 Pmol<sub>PO<sub>4</sub></sub>.Cmol<sub>DOC</sub><sup>-1</sup>. The qPCR results show low percentages of PAO during  
460 period III with an average of 0.12±0.02% confirming these findings. Even when no pH  
461 control was applied (pH>7.2), low Y<sub>p/C,An</sub> values were obtained compared to results  
462 reported by Weissbrodt et al. (2013) indicating that other carbon-storage  
463 microorganisms were more dominant compared to PAO. This can be accredited to the  
464 very high COD/P ratio of the wastewater, i.e. 419±100 mgCOD/mgP. The measured  
465 GAO concentrations are all relatively low (<6%) so no decisive conclusion can be made  
466 regarding the effective enrichment of these specific groups of slow growing organisms.  
467 It is however unclear which group of organisms played a key role in the successful  
468 aerobic granule formation.

#### 469 *Selection of slow growing micro-organisms for aerobic granule formation*

470 Aerobic granule formation was successfully maintained for approximately 350 days  
471 while treating brewery wastewater. The strategy used in this study was based on the  
472 selection and enrichment of slow growing micro-organisms by applying a feast/famine  
473 regime. qPCR results indicate that no significant enrichment of *Accumulibacter* nor  
474 *Competibacter* took place during granule formation. Alongside, 16S rRNA gene  
475 amplicon sequencing analysis was performed which showed that read abundances for  
476 *Accumulibacter* and *Competibacter*, were negligible (<0.1%) for both seed sludge and a  
477 sludge sample taken at the end of period II. Additionally, read abundance of a second  
478 GAO genus also associated with aerobic granule formation, i.e. *Defluviicoccus* (Pronk  
479 et al. 2017) decreased towards the end of period II. However, results concerning  
480 anaerobic DOC uptake, settling characteristics and morphology suggest strong  
481 enrichment of slow growing microorganisms who are able to store carbon intracellularly  
482 during the anaerobic phase and show strong granule formation. According to the 16S  
483 rRNA gene amplicon data, it was only *Candidatus Obscuribacter* (PAO) which seemed  
484 to have increased in read abundance from 0.2 to 0.3±0.1%. Molecular analysis do  
485 therefore not confirm the enrichment of GAO in the system. Interestingly, the read  
486 abundance of *Rhodobacter* and *Thiothrix* were amplified by a factor 3.5 and 10 at the  
487 end of period II compared to the seed sludge. Resulting read abundances were 1.4±0.1  
488 and 10.2±0.6%, respectively. The presence of *Rhodobacter* in aerobic granular sludge  
489 has been reported previously, suggesting a possible role in granule formation due to the  
490 fact they have the capacity to produce and secrete EPS (Lv et al. 2014). In addition,  
491 *Rhodobacter* is also known for its sulphate reducing capacities and since sulphur-  
492 containing compounds were present in the brewery wastewater (i.e. 18.4mg SO<sub>4</sub><sup>2-</sup>-S,  
493 1.21 mg SO<sub>3</sub><sup>2-</sup>-S.L<sup>-1</sup>, 0.154 mg S<sup>-2</sup>.L<sup>-1</sup>) this should be taken into account. As a result,  
494 sulphate present in the brewery wastewater may be anaerobically converted into  
495 sulphide, resulting in possibly significant amounts of sulphide in the system. Recent  
496 studies showed a reduced anaerobic activity of *Accumulibacter* when total sulphide  
497 concentrations in wastewater increased from 0 to 189 mg TS-S.L<sup>-1</sup>. In addition, the  
498 aerobic metabolism of PAO, i.e. phosphate uptake, was also affected negatively by the  
499 presence of sulphide at total sulphide concentrations above 42 mg TS-S.L<sup>-1</sup> (Rubio-  
500 Rincón et al. 2017a). One specific filamentous species of the *Thiothrix* genus, i.e.  
501 *Thiothrix Caldifontis* was found to be able to convert VFA into storage polymers, i.e.  
502 PHA, indicating a competition between *T. Caldifontis* and *Accumulibacter* for FVA  
503 when sulphide is present in the wastewater (Rubio-Rincón et al., 2017b). Since  
504 enrichment of *Rhodobacter* and *Thiothrix* was observed, it may be so that these

505 microbial groups may have had an influence on the selection of *Accumulibacter*. These  
506 results suggest that aside from the well-known GAO and PAO species, enrichment of  
507 other groups of bacteria could influence the evolution of the biomass composition and  
508 therefore the aerobic granule formation and stability. When using complex, industrial  
509 effluents, more fundamental research is needed concerning the influence of other  
510 specific compounds. For example, the influence of S-compounds on the biochemical  
511 conversions and competition between microbial groups should be looked at. During this  
512 study, this was not investigated more intensively since the main focus consisted of  
513 stable aerobic granulation and the influence of the pH on the selection for PAO and  
514 GAO. However, it is highly possible that microorganisms with similar metabolic  
515 pathways are selected by the aerobic/anaerobic SBR process who are not yet associated  
516 with granule formation nor the operational strategy. In future research, 16S rRNA gene  
517 amplicon sequencing should be used in research concerning the treatment of industrial  
518 wastewaters to give more insight in enrichment and role of other species during the  
519 granulation process.

## 520 **Conclusion**

521 Aerobic granule formation in an SBR fed with brewery wastewater was successful after  
522 80 days of operation by applying feast/famine regime, while, in our opinion, a more  
523 applicable feeding strategy was applied by feeding on top of the SBR and complete  
524 initial biomass wash-out was avoided by applying long initial settling periods. Stable  
525 sludge settling capacities characterised by  $SVI_5$  and  $SVI_{30}$  values consistently below  
526  $60\text{mL}\cdot\text{g}^{-1}$  and COD removal efficiencies between 95-99% were maintained for  
527 approximately 280 days. With these findings the stability of AGS for the treatment of an  
528 industrial effluent under changing operational conditions was determined showing good  
529 resistance to fluctuations in F/M ratio and changing pH conditions. Results show that  
530 for industrial applications, where wastewater composition is strongly variable, the  
531 presence of a balancing tank or a dynamic system is highly recommended to insure  
532 feast/famine conditions, maintain stable reactor performances and sludge  
533 characteristics. In this study, the integration of an OUR based aeration insured  
534 endogenous respiration at the end of the aerobic phase. It is thereby found to be suitable  
535 as a strategy when strong fluctuations in wastewater composition occur. In addition,  
536 mixing without feeding is found to be a suitable strategy to preserve AGS in periods  
537 when no wastewater is produced due to periods of shutdown of the industrial activities.  
538 The selection of slow growing organisms is found to be one of the main drivers for  
539 successful granulation. The molecular analysis however do not confirm a significant  
540 enrichment of common PAO or GAO groups associated with successful aerobic  
541 granulation. The lack of enrichment of common PAO and GAO groups, associated with  
542 stable granule formation, is highly remarkable in this study. Combined with the fact that  
543 anaerobic DOC removal increased up to 100%, it should be taken into account that,  
544 when using complex, industrial effluents, it is likely that other, possible slow growing  
545 microorganisms with similar metabolic pathways are selected by the aerobic/anaerobic  
546 SBR process who are not yet associated with granule formation nor the operational  
547 strategy. However, this was not investigated more intensively since it was not the main  
548 focus of this study. More fundamental research is needed concerning the influence of  
549 other specific compounds, for example S-compounds, on the biochemical conversions  
550 and competition between microbial groups.  
551

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