

Identification of inorganic and organic species of phosphorus and its bio-availability in nitrifying aerobic granular sludge

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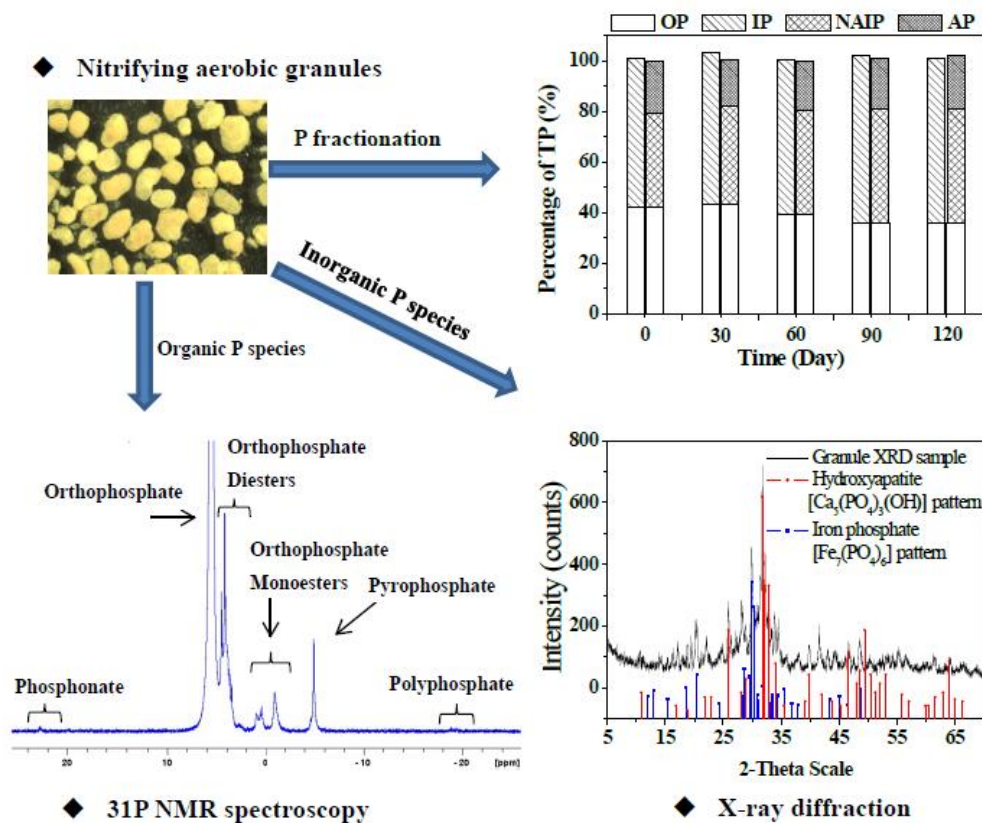
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10 **GRAPHICAL ABSTRACT**



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13 **Abstract**

14 Phosphorus (P) recovery from sewage sludge is necessary for a sustainable
15 development of the environment and thus the society due to gradual depletion of
16 non-renewable P resources. Aerobic granular sludge is a promising biotechnology for
17 wastewater treatment, which could achieve P-rich granules during simultaneous
18 nitrification and denitrification processes. This study aims to disclose the changes in
19 inorganic and organic P species and their correlation with P mobility and
20 bio-availability in aerobic granules. Two identical square reactors were used to
21 cultivate aerobic granules, which were operated for 120 days with influent ammonia
22 nitrogen ($\text{NH}_4\text{-N}$) of 100 mg/L before day 60 and then increased to 200 mg/L during
23 the subsequent 60 days (chemical oxygen demand (COD) was kept constant at 600
24 mg/L). The aerobic granules exhibited excellent COD removal and nitrification
25 efficiency. Results showed that inorganic P (IP) was about 61.4 - 67.7% of total P (TP)
26 and non-apatite inorganic P (NAIP) occupied 61.9 - 70.2% of IP in the granules. The
27 enrichment amount of NAIP and apatite P (AP) in the granules had strongly positive
28 relationship with the contents of metal ions, i.e. Fe and Ca, respectively accumulated
29 in the granules. X-ray diffraction (XRD) analysis and solution index calculation
30 demonstrated that hydroxyapatite ($\text{Ca}_5(\text{PO}_4)_3(\text{OH})$) and iron phosphate ($\text{Fe}_7(\text{PO}_4)_6$)
31 were the major P minerals in the granules. Organic P (OP) content maintained around
32 7.5 mg per gram of biomass in the aerobic granules during the 120 days' operation.
33 Monoester phosphate (21.8 % of TP in extract), diester phosphate (1.8%) and
34 pyrophosphate (0.1%) were identified as OP species by Phosphorus-31 nuclear

35 magnetic resonance (^{31}P NMR). The proportion of NAIP+OP to TP was about 80% in
36 the granules, implying high potentially mobile and bio-available P was stored in the
37 nitrifying aerobic granules. The present results provide a new insight into the
38 characteristics of P species in aerobic granules, which could be helpful for developing
39 P removal and recovery techniques through biological wastewater treatment.

40

41 **Keywords:**

42 Aerobic granules; Nitrification; P fractionation; Phosphorus-31 nuclear magnetic
43 resonance (^{31}P NMR)

44

45 **1. Introduction**

46 Phosphorous (P) is an essential and limiting element for living organisms and human
47 beings, especially for the growth of plants. P deficiency will greatly affect and restrict
48 crop yields worldwide. However, phosphate rock, a non-renewable resource and raw
49 material used for P fertilizers, is reported to be depleted in 50-100 years (Cordell et al.,
50 2009). Thus P resource protection and P recovery is prerequisite for a sustainable
51 agriculture and society on a global scale. On the other hand, sewage sludge generated
52 from wastewater treatment is regarded as a potential phosphorus reservoir due to its
53 high production amount and high P content (Xu et al., 2012). Land application of
54 sewage sludge as P fertilizer has become one of the main sludge disposal practices
55 (Singh and Agrawal, 2008; Xie et al., 2011a).

56 Aerobic granular sludge is a promising biotechnology for nitrogen and

57 phosphorus removal from wastewater. The aerobic granules possess advantages like
58 excellent settleability, high biomass and ability to withstand high loading rate
59 compared to conventional activated sludge processes (Qin et al., 2004; Yilmaz et al.,
60 2008; Adav et al., 2008a). Previous studies indicate that P-rich aerobic granules could
61 be achieved during simultaneous nitrification and denitrification processes (Lin et al.,
62 2012; Li et al., 2014), implying that aerobic granular sludge can be more
63 prospectively used for P fertilizer after being properly treated.

64 As well known that not all the forms of P exhibit similar mobility and
65 bio-availability in the sludge, detailed information about P fraction is necessary for
66 both activated sludge and aerobic granules, especially when land application of sludge
67 is taken into consideration. Moreover, identification of P species in aerobic granular
68 sludge is helpful to understand the characteristics and function of P in aerobic
69 granules and thus the mechanisms of P removal through this new biotechnology. For
70 this purpose, some researchers declared that sewage sludge contained higher
71 mobilizable forms of P, i.e. non-apatite inorganic P than sediment (Medeiros et al.,
72 2005; Xie et al., 2011a and 2011b), which is much meaningful for land application of
73 sewage sludge. To date, as for aerobic granules, lab-scale experiments have been
74 conducted on the mineral forms of P, demonstrating that the major inorganic P species
75 in aerobic granules were greatly dependent on influent characteristics and operation
76 conditions (Angela et al., 2011; Lin et al., 2012; Li et al., 2014). More specifically, in
77 enhanced biological phosphorus removal (EBPR) process, biologically induced
78 precipitation of phosphorus as hydroxyapatite was discovered in the core of granules

79 (Angela et al., 2011). On the other hand, Ca-Mg phosphate and whitlockite
80 ($\text{Ca}_3(\text{PO}_4)_2$) were found to be the major minerals in aerobic granular sludge used for
81 simultaneous phosphorus accumulation by chemical precipitation and biological
82 nitrogen removal via nitrite (Li et al., 2014). In addition, previous works show that
83 orthophosphate, pyrophosphate and polyphosphate are the main inorganic species
84 of phosphorus in the EPS of activated sludge and aerobic granular sludge, accounting
85 for 5-10.5% of total P in the EBPR systems (Zhang et al., 2013a, 2013b). However,
86 the organic P species in aerobic granules, the potential available P resource closely
87 related with the activity of phosphate accumulating organisms (PAOs) (Uhlman et al.,
88 1990), is poorly understood most probably attributable to its complex nature and the
89 limitation of analytical methods.

90 Recent research indicates that phosphorus-31 nuclear magnetic resonance
91 spectroscopy (^{31}P NMR) can be used for analyzing inorganic and organic P species
92 (orthophosphate monoesters, orthophosphate diester and phosphonates) in sediments,
93 soil and activated sludge since it is able to distinguish multiple P compounds among
94 complex substances (Uhlmann et al., 1990; Ahlgren et al., 2011; Li et al., 2013).
95 Therefore it is speculated that this technology could provide more insight into the P
96 species in aerobic granules, especially the organic P fraction.

97 This study aims to reveal the fractionation and distribution of P in aerobic
98 granules and to evaluate the mobility and bio-availability of P in aerobic granules. In
99 addition, organic and inorganic P species in the granules were also determined and
100 characterized by ^{31}P NMR and X-ray diffraction (XRD), respectively. It is expected

101 that this work would not only be useful for P utilization and recovery from aerobic
102 granules but also provide insight into the characteristics of P in aerobic granules,
103 which will help to develop applicable technologies for P removal and recovery from
104 wastewater.

105

106 **2. Materials and methods**

107 **2.1. Experimental set-up and operation**

108 Aerobic granules were cultivated in two identical sequencing batch reactors (SBRs)
109 made of acrylic plastic with height of 60 cm and square cross section of 6 cm × 6 cm.
110 Their effective working volume was 1.4 L. Aeration was provided by an air pump
111 (AK-30, KOSHIN, Japan) through air bubble diffusers at the bottom of each reactor
112 with an air flow rate of 2.0 cm/s and the dissolved oxygen (DO) maintained at 7-9
113 mg/L during aeration. Synthetic wastewater was used in this study, and its
114 composition was as follows: chemical oxygen demand (COD) 600 mg/L (50% of
115 which was contributed by glucose and sodium acetate, respectively); 10 mg PO₄-P/L
116 (KH₂PO₄); 100 mg NH₄-N/L (NH₄Cl) during the first 60 days' operation and then
117 increased to 200 mg NH₄-N/L till the end of experiment; 10 mg Ca²⁺/L (CaCl₂); 5 mg
118 Mg²⁺/L (MgSO₄•7H₂O); 5 mg Fe²⁺/L (FeSO₄•7H₂O); and 1ml/L of trace element
119 solution. The trace element solution contained (in mg/L) H₃BO₃ (50), ZnCl₂ (50),
120 CuCl₂ (30), MnSO₄•H₂O (50), (NH₄)₆Mo₇O₂₄•4H₂O (50), AlCl₃ (50), CoCl₂•6H₂O
121 (50), and NiCl₂ (50) (Adav et al., 2008b). The pH in the reactors was adjusted with
122 sodium bicarbonate to be within 7.5-8.0.

123 Each reactor was inoculated with 0.5 L of seed sludge sampled from a
124 sedimentation tank of the Shimodate Sewage Treatment Plant, Ibaraki Prefecture,
125 Japan. The initial mixed liquor suspended solids (MLSS) concentration was 4.6 g/L
126 with sludge volume index (SVI) of 81.4 ml/g and MLVSS/MLSS of 0.8 in the two
127 reactors. After aerobic granules appeared, the mixed liquor was withdrawn daily from
128 the reactors in order to keep their solids retention time (SRT) around 40 days. The
129 reactors were operated sequentially in a 4-h cycle at room temperature ($25 \pm 2^\circ\text{C}$): 2
130 min of influent filling, 28 min of non-aeration period, 186-206 min of aeration, 2-20
131 min of settling, and 4 min of effluent discharge. The settling time was gradually
132 decreased from 20 min to 2 min due to the increase in settleability of the sludge. The
133 volumetric exchange ratio was kept at 54%, resulting in a hydraulic retention time of
134 7.4 h.

135 **2.2. Chemical and physical analysis**

136 Mixed liquor (volatile) suspended solids (ML(V)SS), sludge volume index (SVI),
137 COD, ammonia nitrogen ($\text{NH}_4\text{-N}$), nitrite nitrogen ($\text{NO}_2\text{-N}$), nitrate nitrogen ($\text{NO}_3\text{-N}$),
138 and phosphorus ($\text{PO}_4\text{-P}$) were measured in accordance with standard methods
139 (APHA, 1998). DO concentration in the bulk liquid was measured with a DO meter
140 (HQ40d, HACH, USA). pH was determined by using a pH meter (Mettler Toledo
141 FE20, Switzerland).

142 Metal ions in sludge particles were quantified after the sludge samples being
143 digested and filtered through 0.22 μm cellulose nitrate membrane filters (Nalgene).
144 0.1 g of dried sludge was digested in a mixture of 3 ml hydrochloric acid (37%,

145 Wako), 1 ml nitric acid (70%, Wako), and 1 ml perchloric acid (60%, Wako) on an
146 electric heating plate for 10 min. The concentration of each metal was measured by
147 inductively coupled plasma mass spectrometry (ICP-MS, ELAN DRC-e, Perkin
148 Elmer, USA). Granular surface elemental analysis was also performed by using an
149 energy dispersive X-ray - spectroscopy (EDX) equipped with a JSM 7000F field
150 emission scanning electron microscope (FE-SEM) operated at an acceleration voltage
151 of 20 kV.

152 The mean granular size was measured by a stereo microscope (STZ-40TBa,
153 SHIMADZU, Japan) with a program Motic Images Plus 2.3S (Version 2.3.0).
154 Morphology characteristics of the granules were observed using a scanning electron
155 microscope (SEM, JSM6330F, Japan). XRD analysis was performed using a
156 Multiflex diffractometer (Rigaku, Japan) with a cobalt tube scattering from 5 to 75° in
157 2θ. The samples used for XRD analysis were previously dried and calcined in an oven
158 at 500°C for 2 h in order to remove the organic fraction.

159 Average values were taken for all the determinations and used for results and
160 discussion.

161 **2.3. Phosphorus fractionation in granular sludge**

162 In this study, the Standards, Measurements and Testing (SMT) Programme extraction
163 protocol was applied to analyze phosphorus fractions in the aerobic granules, which
164 has been widely used in soil, sediment and sewage sludge samples (Ruban et al., 1999;
165 Medeiros et al., 2005). After sequential extraction based on the SMT method,
166 phosphorus in sludge can be fractionated into the following 5 categories: (1)

167 concentrated HCl-extractable P, namely total P (TP), (2) organic P (OP), (3) inorganic
168 P (IP), (4) non-apatite inorganic P (NAIP, the P fraction associated with oxides and
169 hydroxides of Al, Fe and Mn), and (5) apatite P (AP, the P fraction associated with
170 Ca). In order to avoid the transformation of P species in granules during preparation,
171 the samples taken from the reactors were frozen immediately at -80°C , lyophilized
172 and stored at -20°C until analysis. The phosphorus concentration in the supernatant
173 collected after extraction was determined with molybdenum blue method.

174 **2.4. ^{31}P NMR analysis**

175 Granular sludge taken from each reactor was also frozen immediately at -80°C ,
176 lyophilized, and ground to fine powders. Then P was extracted from 2 g of the
177 prepared sludge powder with 40 ml of a solution consisting of 0.25 M NaOH and 0.05
178 M Na_2EDTA (Turner et al., 2003). The mixture was shaken for 6 h at 100 rpm and
179 ambient temperature and then centrifuged at $6000 \times g$ for 20 min at 4°C . 2 ml of the
180 resultant supernatant was taken for TP, IP and OP analysis. The remaining extracts
181 were freeze-dried again and stored at -20°C till ^{31}P NMR analysis.

182 500 mg of freeze-dried extract was re-dissolved in 0.8 ml of 1M NaOH and 0.2
183 ml D_2O and then transferred to a 5 mm NMR tube. The ^{31}P NMR spectrum was
184 obtained by using a Bruker Avance-600MHz NMR Spectrometer at 242.94 MHz for
185 ^{31}P . 90°C of pulse width, 25°C of regulated temperature, and acquisition time of 0.67
186 s (with relaxation delay of 2s) were applied in the experiments. Chemical shifts of
187 signals were determined relatively to an external standard of 85% H_3PO_4 via signal
188 lock. The peaks were assigned to P species according to the reports in literature with

189 peak areas calculated by integration (Ahlgren et al., 2005; Turner et al., 2003; Turner,
190 2004; Zhang et al., 2013).

191

192 **3. Results and discussion**

193 **3.1. Formation and characterization of aerobic granules**

194 The SBRs were operated for 120 days. Granules appeared in the two reactors on day
195 13 after startup and then grew gradually along with the operation. From day 90 on, the
196 granular size averagely stabilized at 0.76-0.78 mm, although the initial diameter of
197 seed sludge was about 0.17 mm (Table 1). This observation is in agreement with the
198 finding by Verawaty et al. (2013) who reported that granules in the reactor
199 equilibrated towards a common critical size of around 0.6-0.8 mm. MLSS and SVI₃₀
200 were determined to be 12-13 g/L and 22 ml/g after 60 days' cultivation. It is worth
201 noting that the MLVSS/MLSS ratio of granules progressively decreased from 86% to
202 73% at the end of experiment, probably due to the accumulation of mineral substances,
203 which will be further demonstrated in the following sections. As shown in Fig. 1a, the
204 yellowish mature granules had irregular shapes, possibly contributed by the square
205 structure of SBRs in this study. SEM observation on day 110 clearly shows the
206 compact and dense structure of the granules, and most of the bacteria are distributed
207 in the outer layer (Figs. 1b and 1d) with little bacteria in the core of the granules (Figs.
208 1b and 1c), which is in agreement with the results from Adav et al. (2008b).
209 Furthermore, EDX results shown in Fig. S1 indicate that the intensities of metal ions
210 and P are decreasing along with the granular radius, signaling the metal ions and P are

211 mainly accumulated in the core of granules. This observation suggests the existence
212 of some relationship between the accumulated metal ions and phosphorus.

213

214 **3.2. Overall pollutants removal performance**

215 In order to evaluate COD, ammonia and phosphate removal rates, typical batch
216 experiments were carried out on days 55 and 115, namely the influent $\text{NH}_4\text{-N}$
217 concentration was 100 mg/L and 200 mg/L, respectively (Fig. 2). It was found that
218 COD removal efficiency was about 80% during non-aeration stage, then increased to
219 around 92% after aeration for 30 min and kept this COD removal rate in both cycle
220 tests (data not shown). Figs. 2a and 2b show that a small amount of $\text{NH}_4\text{-N}$ was
221 removed during the non-aeration period, probably attributable to heterotrophic
222 assimilation and adsorption by the granules (Bassin et al., 2011). In the subsequent
223 aeration stage, $\text{NH}_4\text{-N}$ was observed to first convert to $\text{NO}_2\text{-N}$ and then rapidly to
224 $\text{NO}_3\text{-N}$ by nitrifying bacteria. Both $\text{NH}_4\text{-N}$ and $\text{NO}_2\text{-N}$ were not detectable (99.9% of
225 nitrification) after influent feeding for about 2 h and 3 h under influent $\text{NH}_4\text{-N}$
226 conditions of 100 mg/L and 200 mg/L, respectively (Figs. 2a and 2b). The results
227 indicate that the granules possess excellent nitrification capacity, about 9 mg
228 $\text{NH}_4\text{-N/g-VSS h}$ with the influent COD of 600 mg/L and $\text{NH}_4\text{-N}$ of 200 mg/L.

229 As shown in Fig. 2a, on the other hand, phosphorus was observed to release
230 remarkably during the non-aeration period while P uptake was detected during
231 aeration period under 100 mg/L of influent $\text{NH}_4\text{-N}$ concentration, suggesting the
232 presence and activity of polyphosphate accumulating organisms (PAOs). This

233 phenomenon, however, didn't occur when the influent $\text{NH}_4\text{-N}$ concentration increased
234 to 200 mg/L (Fig. 2b), signaling the activity of PAOs might be inhibited by free
235 ammonia (FA). Zheng et al. (2013) claimed that 8.88 mg-N/L of FA initiated the
236 inhibition on PAOs and its toxic threshold concentration for P metabolism would be
237 17.76 mg-N/L. In this study, the pH of bulk liquor ranged between 7.5 and 8.0, thus
238 FA was maintained at relatively high levels (3.6 - 11.1 mg-N/L). In addition, only
239 about 1 mg-P/L of P removal (10% of influent P) was achieved through microbial
240 assimilation during the two cycle tests, which was mostly associated with a long SRT
241 (40 days) applied in this study. The above results suggest that the aerobic granules
242 cultivated in this study possess high nitrification capability while low P removal
243 efficiency.

244

245 **3.3. P fractionation in aerobic granules by SMT protocol**

246 Table 2 lists the analytical results of P fractions in seed sludge and aerobic granules.
247 Seen from Table 2, it is clearly that the content of each P fraction in seed sludge is
248 much higher than that in aerobic granules. This observation may be resulted from the
249 different influent composition and operation strategy between the seed sludge
250 sampled from the wastewater treatment plant and the aerobic granules cultivated in
251 this study. Table 2 also shows that the OP content maintained at around 7.5
252 mg/g-MLSS in aerobic granules during the whole process, suggesting that OP is
253 relatively stable in quantity under the designed operation conditions, which may play
254 an important role in the P recovery from aerobic granules. Although the OP content of

255 seed sludge (15.6 mg/g) is much higher than that of granules (Table 2), the percentage
256 of OP to TP for both sludges is quite similar (42.3% in the seed sludge and 39.3-43.3%
257 in the granules sampled between day 30 and day 60, Fig. 3a). Specifically, it is worth
258 mentioning that the OP content and percentage in this study are higher than the
259 findings of previous studies (1.4-5.8 mg/g and 9.9-22.3%)(Xie et al., 2011b; Medeiros
260 et al., 2005), which might be brought about by the different drying and storage
261 methods used for sludge preparation in these works.

262 On the other hand, IP is the major P fraction in the seed sludge (57.5% of TP)
263 and aerobic granules (61.4%-67.7% of TP), in which NAIP amounts to 65.1% and
264 61.9-70.2% of IP, respectively (Fig.3a). In addition, NAIP is about 37.4% and
265 38.0-47.5% of TP in the seed sludge and aerobic granules. Compared with OP and
266 NAIP, AP content is relatively lower and much stable, about 20.9% and 18.1-20.1% of
267 TP respectively in the seed sludge and aerobic granules, respectively.

268 When the changes in biomass concentration in the reactors are taken into
269 consideration, P mass stored in the sludge could be used for the assessment of P
270 accumulation capability of the microorganisms (PAOs). Obviously from Fig. 3b it can
271 be discerned that before granular maturation (day 30), little accumulation was
272 detected in all the P fractions, although the biomass concentration doubled in the
273 reactors (Table 1), possibly due to the absence of PAOs and the operation strategy
274 used in this study. Under the same operation conditions including the same influent
275 COD (600 mg/L) and NH₄-N (100 mg/L) concentrations, however, P amount stored in
276 the biomass increased remarkably on day 60 by 80%, 53% and 70% with respect to

277 TP, OP and IP, respectively, although P removal from wastewater was low (Fig.2).
278 Nevertheless, this increase trend seemed to retard to some extent when the influent
279 $\text{NH}_4\text{-N}$ concentration was increased to 200 mg/L, especially for OP. This observation
280 partly coincides with the cycle tests (Fig. 2). Along with the operation from day 60 to
281 day 120, although P mass in inorganic forms (NAIP and AP) continuously increased,
282 the organic P mass showed much less increment (Fig. 3b), partially signaling the
283 activity of PAOs was inhibited. In addition, seen from Fig. 3b the accumulation rates
284 of P species during days 30-60 were about 2 times those of P species during days
285 90-120, also suggesting PAOs in the granules might be inhibited during the later
286 period (under 200 mg/L of influent $\text{NH}_4\text{-N}$ conditions).

287 Interestingly, the increase of IP, NAIP and AP in the sludge was found to have
288 close relationship with the amount of metal ions accumulated in the granules,
289 especially Ca and Fe in this work (Table 3). The linear correlation coefficient (R^2) was
290 0.995 between accumulated Fe and NAIP, which was about 0.971 between
291 accumulated Ca and AP content in the sludge. According to the SMT protocol, NAIP
292 is mainly associated with Al, Fe and Mn, while AP is directly related with Ca (Ruban
293 et al., 1999; Pardo et al., 2003). Due to very less amount of Mn and Al in the sludge, it
294 can be inferred that mineral phosphorus was mainly associated with Fe and Ca in
295 NAIP and AP fractions, respectively in this study. In addition, the accumulation of Fe
296 in granules has a positive effect on the bio-available P because both NAIP and OP are
297 regarded as potentially releasable and bio-available P (Ruban et al., 1999; Xie et al.,
298 2011a). As AP is considered to be non-bioavailable P, Ca accumulation or

299 precipitation will bring about an increase in quantity of AP and then a decrease in the
300 proportion of bio-available P in the granules. Furthermore, Ca precipitation is
301 supposed to have a negative effect on the accumulation of polyphosphate and OP in
302 the sludge, leading to the decreases in bioavailability of phosphorus, PAO activity and
303 MLVSS/MLSS ratios of granules (Barat et al., 2008; Ren et al., 2008). Restated, in
304 this study the IP accumulated in biomass was mainly contributed by the increase in
305 NAIP content (Fig. 3b), implying an increased bio-availability of P in the nitrifying
306 granules during the granulation process.

307 In this study, the proportion of NAIP+OP was about 80% for both seed sludge and
308 aerobic granular sludge (Fig. 3a). Similar results were reported by Xie et al. (2011b)
309 who found that NAIP+OP was greater than 85% and 75% of TP in activated sludge
310 samples fed with domestic and industrial wastewaters, respectively. Moreover, a
311 positive correlation ($R^2= 0.995$) was observed between TP and NAIP+OP in aerobic
312 granules during the whole operation. Therefore, it could be concluded that high
313 potentially mobile and bio-available phosphorus were stored in the nitrifying aerobic
314 granules.

315

316 **3.4. Species of inorganic P in aerobic granules by XRD**

317 XRD, an efficient tool for distinguishing crystalline minerals from those of
318 amorphous structure, was used to identify the species of P minerals in the granule
319 samples. As illustrated in Fig. 4, a number of distinct peaks in the diffractogram
320 reflect the presence of crystalline forms. By comparison with portable document
321 format standard card (2004) using Jade 6.0, most of the stronger peaks coincide with

322 those of hydroxyapatite ($\text{Ca}_5(\text{PO}_4)_3(\text{OH})$) pattern and iron phosphate($\text{Fe}_7(\text{PO}_4)_6$)
323 pattern.

324 Different P minerals have been reported in aerobic granules, such as
325 hydroxyapatite ($\text{Ca}_5(\text{PO}_4)_3(\text{OH})$), whitlockite ($\text{Ca}_3(\text{PO}_4)_2$, $\text{Ca}_{18}\text{Mg}_2\text{H}_2(\text{PO}_4)_{14}$) and
326 struvite ($\text{NH}_4\text{Mg PO}_4 \cdot 6\text{H}_2\text{O}$) in previous studies (Angela et al., 2011; Li et al., 2014;
327 Lin et al., 2012). The result from this study agrees with Angela et al (2011) who
328 reported that hydroxyapatite was a major phosphate mineral in aerobic granules. In
329 the calcium phosphate family, hydroxyapatite is considered as the most stable and
330 insoluble one. Moreover, the dense structure of granules encourages the accumulation
331 of P minerals in the core of granules and represses the solubilization of the crystals.
332 However, no whitlockite ($\text{Ca}_3(\text{PO}_4)_2$) or struvite ($\text{NH}_4\text{Mg PO}_4 \cdot 6\text{H}_2\text{O}$) was detected in
333 the granules. As pH and influent composition play an important role on phosphate
334 precipitation due to their influence on the saturation index (SI) of different minerals
335 (Montastruc et al., 2003; Barat et al., 2011; Juang et al., 2010), in this study SI was
336 calculated using the Visual MINTEQ ver. 3.1 database (PHREEQC software) (Table
337 4). The results show that SI of struvite was negative under the designed conditions in
338 this study, indicating magnesium concentration might be too low to initiate struvite
339 precipitation. The SI of whitlockite ($\text{Ca}_3(\text{PO}_4)_2$) was close to zero, implying
340 whitlockite was poorly or very temporarily formed. On the contrary, high level of SI
341 (between 4.78 and 7.66) was obtained for hydroxyapatite, i.e. the most stable phase
342 among the calcium phosphates signaling its oversaturation state in the tested
343 conditions. Besides the naturally chemical precipitation of hydroxyapatite, biological

344 especially PAOs induced hydroxyapatite precipitation and accumulation should also
345 be considered (Angela et al., 2011). As mentioned above, the accumulation rate of AP
346 during days 30-60 was much higher than that during days 90-120. To some extent, the
347 increased AP content might be partially contributed by PAOs induced hydroxyapatite
348 precipitation, due to that hydroxyapatite is the major component of AP.

349 In addition, high level of SI for vivianite ($\text{Fe}_3(\text{PO}_4)_2 \cdot 8\text{H}_2\text{O}$) was also obtained
350 (Table 4). According to the portable document format standard card, the observed
351 XRD peaks do not agree with those of vivianite ($\text{Fe}_3(\text{PO}_4)_2 \cdot 8\text{H}_2\text{O}$) pattern. However,
352 two major peaks (20.5 and 29.7) of the granular sample could be assigned to another
353 iron phosphate, ($\text{Fe}_7(\text{PO}_4)_6$) pattern. Under the high DO levels (7-9 mg/L) applied in
354 the reactors in this study, part of Fe^{2+} could be inevitably and easily oxidized to Fe^{3+}
355 and then formed the stable crystal of iron phosphate ($\text{Fe}_7(\text{PO}_4)_6$). Moreover, FePO_4
356 crystal can be gradually transformed into iron phosphate ($\text{Fe}_7(\text{PO}_4)_6$) by redox
357 reactions (Gadgil et al., 1994; Wang et al., 2013) created in the granules. Therefore, it
358 can be confirmed that hydroxyapatite (HAP) and iron phosphate ($\text{Fe}_7(\text{PO}_4)_6$) are the
359 major inorganic P in the nitrifying aerobic granules in this study. Still, whether other
360 intermediates present or not remain unknown due to the limitation of XRD for
361 micro-amount minerals.

362

363 **3.5. Identification of P species in granules by ^{31}P NMR analysis**

364 All NMR-spectra show peaks in the areas for orthophosphate (Ortho-P, 5–7 ppm),
365 orthophosphate monoesters (Monoester-P, 3–5 ppm), orthophosphate diester

366 (Diester-P, 2 to -3 ppm), pyrophosphate (Pyro-P, -4 to -6 ppm), polyphosphate
367 (Poly-P, -18 to -21 ppm), and phosphonates (22-24 ppm) (Fig. 5). Monoester-P,
368 Diester-P and phosphonates belong to OP, while Ortho-P, Pyro-P and Poly-P are IP.

369 Although it's difficult to make an exact integration of the peak areas in NMR, the
370 quantification of these compounds with NMR is a suitable method to estimate the
371 relative proportions of P groups (Ahlgren et al., 2005; Zhang et al., 2013). Table 5
372 shows the contents of these P fractions extracted by NaOH+ Na₂EDTA method and
373 their relative proportions (%TP) identified by ³¹P NMR. The average TP content in the
374 NaOH+ Na₂EDTA extracts was 17.3 mg-P/g SS with an average extraction rate of
375 approximately 77.6% of TP, which is similar with the result (about 73%) obtained by
376 Turner et al.(2004).

377 Orthophosphate is the dominant P species in the granule extracts, accounting for
378 74.3% of TP. Ortho-P mainly exists in phosphate form like Fe- and Ca-bound
379 inorganic P and is also the main nutrient for living organisms. It is deduced that
380 accumulation of Fe and Ca could increase the concentration of orthophosphate in the
381 granules. Although a low level of 1.9% (of TP) was detected in the granule extracts,
382 Pyro-P content in the granules is much higher than those (0.2-0.8%) of lake sediment
383 samples (Zhang et al., 2013). Besides, the presence of Pyro-P signals high microbial
384 activity involved in the biological P cycling in samples (Condrón et al., 1985; Ahlgren
385 et al., 2005). On the other hand, only a very small amount of Poly-P was extracted,
386 comprising approximately 0.1% of the extractable TP from granules, which is much
387 lower than those (50% of TP) from activated sludge samples with

388 phosphate-accumulating bacteria (Uhlmann, et al., 1990). The less amount of Poly-P
389 in the granules sampled on day 120 was probably brought about by the inhibition of
390 FA on the activity of PAOs. Extracellular polymeric substances (EPS) was noticed to
391 play an important role on P species in EBPR sludge (Zhang et al., 2013a, 2013b), in
392 which Poly-P was the major P species followed by Ortho-P and Pyro-P. In this study,
393 Poly-P was much lower, about 0.1% of the extractable TP, compared to other IP
394 species like Ortho-P and Pyro-P determined. Thus much less contribution of Poly-P
395 was associated with the change in P bioavailability of the nitrifying granules, possibly
396 due to the inhibition of FA to PAOs, which needs further investigation. Furthermore,
397 EPS, consisting of polysaccharides, proteins, glycoproteins, nucleic acids,
398 phospholipids, and humic acids (Adav et al., 2008b; McSwain et al., 2005), can form
399 a matrix for microbial cells and influence the distribution of poly-P and OP species in
400 the granules.

401 On the other hand, Monoester-P is the major form of OP extracted with NaOH-
402 Na₂EDTA from the aerobic granules, accounting for about 21.8% of TP. This is in
403 agreement with the previous results from soil and sediment samples. Monoester-P can
404 be directly correlated with microbial, for instance, the glycerol-6-phosphate
405 (nucleotides) found in cell membrane belongs to Monoester-P (Ahlgren et al., 2011).
406 In addition, George et al. (2006) reported that the rhizosphere of plant could utilize
407 Monoester-P from oxisols after phosphatase hydrolysis. Moreover, algae can
408 assimilate more than 40% of P from Monoester-P in a temperate mesotrophic lake
409 (Hernandez et al., 1997). Although only 1.8% of extracted TP in granules, Diester-P

410 plays a crucial role in the aerobic granules due to its close relation to
411 deoxyribonucleic acid (DNA-P), lipid (lipids-P) and teichoic acid (Teichoic-P). Thus
412 the quantity of Diester-P can be used to indicate the activity and concentration of
413 microorganisms in aerobic granules. As Whitton et al. (1991) pointed out that
414 Diester-P could be easily absorbed and utilized by plants and blue green algae under
415 certain conditions, the OP fractions including Monoester-P and Diester-P are believed
416 to play an important role in the potential mobility and bio-availability of P resource
417 from aerobic granular sludge.

418

419 **4. Conclusion**

420 This study presents the preliminary results of P species and its bio-availability in
421 nitrifying aerobic granular sludge. The following major conclusions could be arrived
422 at:

423 (1) IP is the primary P fraction in the granules, in which NAIP amounts to
424 62-70%. A positive strong correlation ($R^2=0.971-0.995$) has been found between the
425 content of IP (NAIP and AP) and that of metal ions (especially Fe and Ca) in the
426 granules. OP content was very stable in quantity during the operation. About 80% of
427 TP in aerobic granular sludge possessed high potential mobility and bio-availability.

428 (2) XRD analysis and SI calculation reveal that hydroxyapatite ($\text{Ca}_5(\text{PO}_4)_3(\text{OH})$)
429 and iron phosphate($\text{Fe}_7(\text{PO}_4)_6$) patterns are the main IP species in the nitrifying
430 aerobic granules.

431 (3) Three organic P compounds (Monoester-P, Diester-P, phosphonates) and three

432 inorganic P compounds (Ortho-P, Pyro-P and Poly-P) have been identified by using
433 ³¹P NMR, respectively. Monoester-P was the dominant P in OP, and less amount of
434 Poly-P was detected in the granules, probably attributable to the inhibition of FA on
435 the activity of PAOs.

436 In this study P removal rate was low due to the inhibition of relatively high FA
437 levels, which could be alleviated with enhanced P removal by adjusting operation
438 strategy (data not shown). The results from current work imply that aerobic granules
439 could be further potentially developed as P resource materials with high amount of
440 bioavailable P.

441

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559 **Tables**

560

Operation duration (day)	MLS S (g/L)	MLVSS/ MLSS (%)	S VI ₃₀ (m l/g)	Average diameter (mm)
0	3.6	80	81	0.17
30	7.8	86	28	0.38
60	11.5	82	25	0.62
90	12.8	77	22	0.76
120	13.1	73	22	0.78

561

Operation duration (day)	TP	OP	IP	NA IP	A P	NAI P+AP
Seed sludge	36.9 ±0.9	15.6 ±0.5	21.2 ±0.7	13. 8±0.5	7. 7±0.3	21.5 ±0.8
30	17.1 ±0.4	7.4 ±0.5	10.5 ±0.3	6.5 ±0.4	3. 1±0.2	9.6± 0.7
60	19.6 ±0.7	7.7 ±0.3	12.1 ±0.6	8.0 ±0.6	3. 7±0.2	11.7 ±0.7
90	20.4 ±0.6	7.3 ±0.6	13.5 ±0.5	9.2 ±0.4	4. 1±0.4	13.3 ±0.5
120	22.3 ±0.6	7.6 ±0.4	15.1 ±0.2	10. 6±0.5	4. 4±0.3	15.0 ±0.6

The data are expressed as mean value ± standard deviation. MLSS, mixed liquor suspended solids; TP, total phosphorus; OP, organic phosphorus; IP, inorganic phosphorus; NAIP, non-apatite inorganic phosphorus; AP, apatite phosphorus.

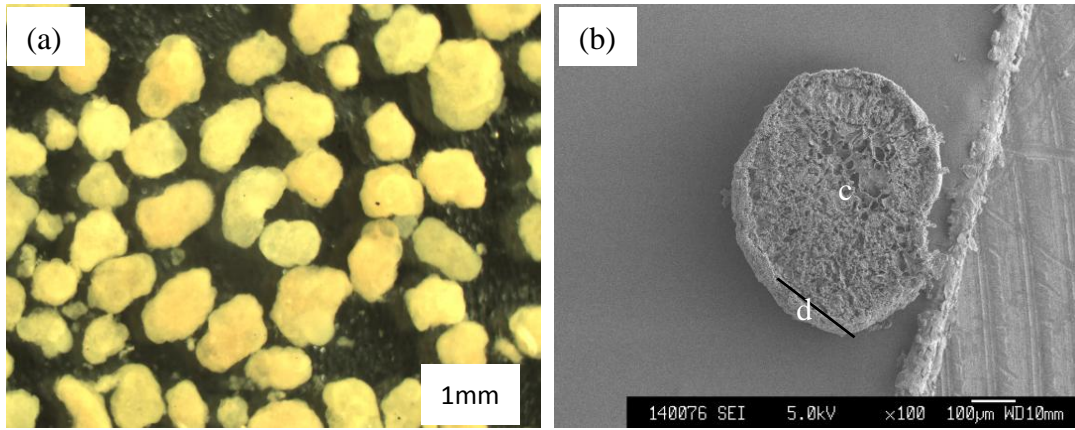
Operation duration (day)	N	K	C	M	F	M	A
30	3	1	4	2	6	0	0
	.71	.46	.60	.67	.02	.21	.15
60	4	2	6	3	8	0	0
	.05	.06	.77	.74	.07	.26	.18
90	4	2	8	3	1	0	0
	.27	.33	.49	.51	0.46	.31	.25
120	5	3	1	3	1	0	0
	.50	.07	0.79	.96	2.48	.45	.25

Species	100 mg NH ₄ -N /L		200 mg NH ₄ -N /L	
	pH 7.5	pH 8.0	pH 7.5	pH 8.0
	Ca ₃ (PO ₄) ₂ (beta)	-0.63	0.47	-1.08
Calcite(CaCO ₃)	-0.42	0.08	-0.25	0.25
Dolomite (CaMg(CO ₃) ₂)	-0.76	0.24	-0.41	0.59
Hydroxyapatite(Ca ₅ (PO ₄) ₃ (OH))	5.53	7.66	4.78	6.89
Struvite(MgNH ₄ PO ₄ •6H ₂ O)	-1.81	-1.26	-1.69	-1.15
Vivianite (Fe ₃ (PO ₄) ₂ •8H ₂ O)	6.26	7.35	6.24	7.33

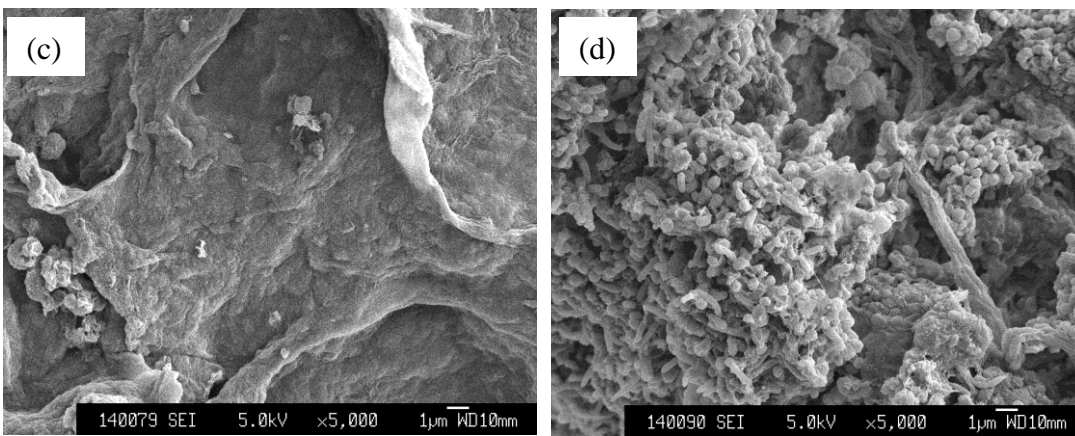
TP (mg/ g-SS)	IP			OP		
	Ort ho-P (%)	Pyr o-P (%)	Pol y-P (%)	Monoester-P (%)	Diester-P (%)	Phosphonate (%)
17.3	74.3	1.9	0.1	21.8	1.8	0.1

SS, suspended solids; Ortho-P, orthophosphate; Monoester-P, orthophosphate monoesters; Diester-P, orthophosphate diester; Pyro-P, pyrophosphate; Poly-P, polyphosphate.

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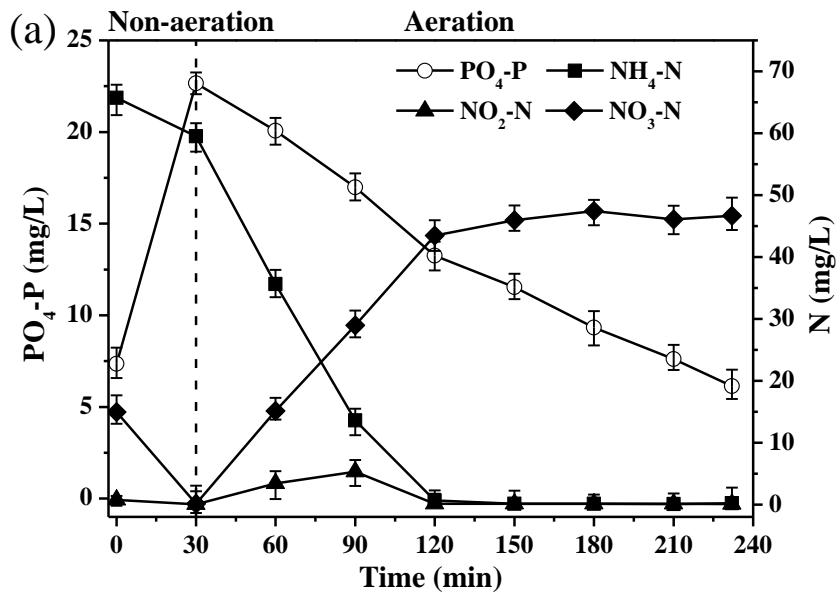
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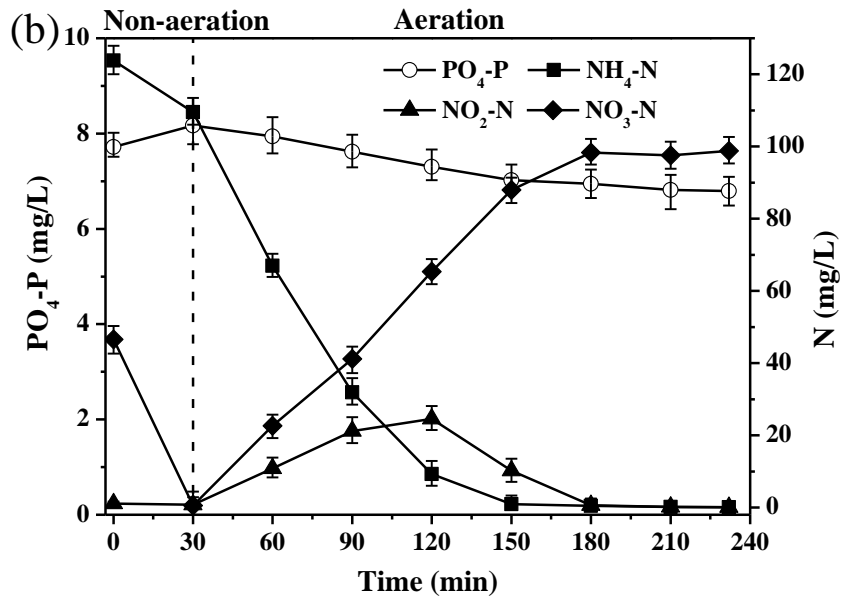
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574 Fig. 1 – Images of the granular sludge on day 110 (Influent $\text{NH}_4\text{-N}$ = 200 mg/L).
575 Digital image of granules (a), and SEM observation of granules from cross section (b),
576 core (c) and edge (d) on day 110, respectively.



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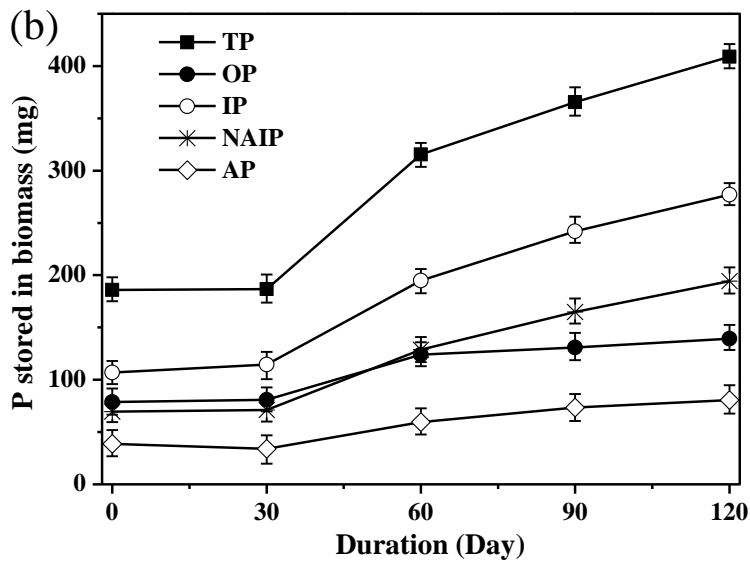
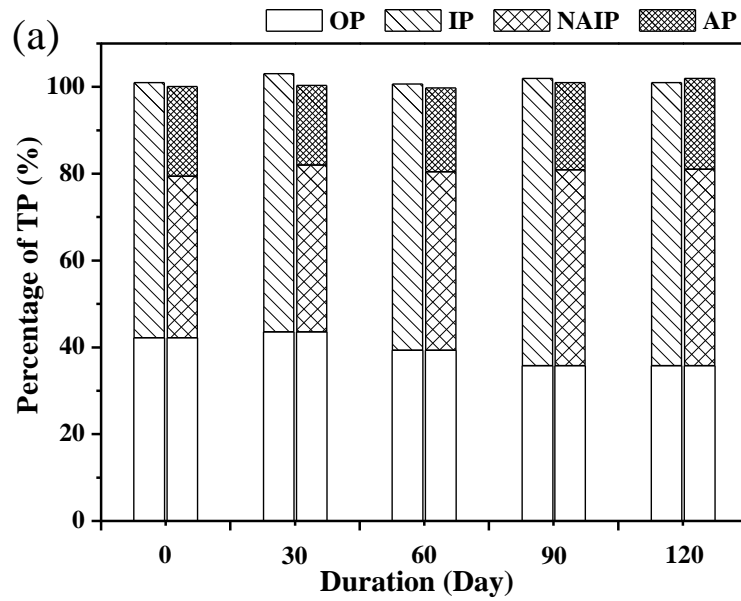


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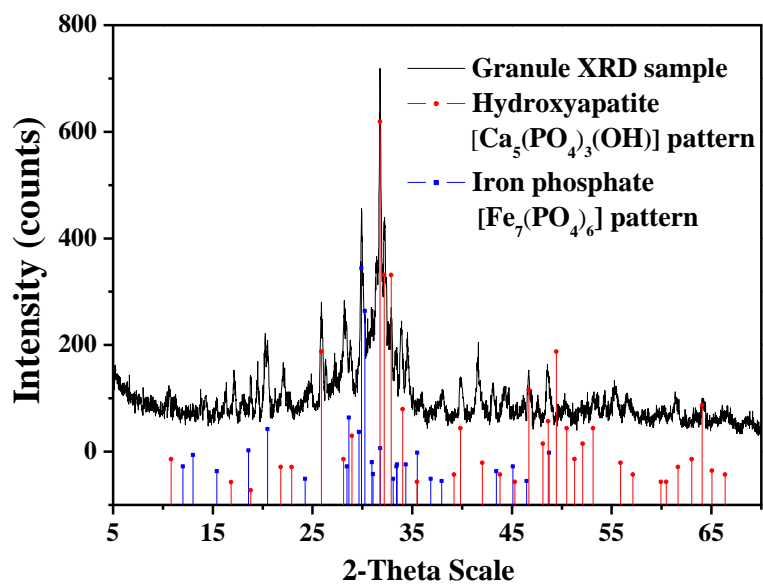
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581 Fig. 2 – Variation of $\text{NH}_4\text{-N}$, $\text{NO}_2\text{-N}$, $\text{NO}_3\text{-N}$ and $\text{PO}_4\text{-P}$ in the bulk liquor during
 582 cycle tests under 100 mg/L (a) and 200 mg/L (b) of influent $\text{NH}_4\text{-N}$ concentration,
 583 respectively.



588 Fig. 3 – Average proportions of phosphorus fractions (%TP) in sludge (a) and changes
 589 in P mass stored in biomass in the reactors (b) during 120 days' operation. TP, total
 590 phosphorus; OP, organic phosphorus; IP, inorganic phosphorus; NAIP, non-apatite
 591 inorganic phosphorus; AP, apatite phosphorus. P stored in biomass is estimated
 592 according to P fraction content and biomass (MLSS) concentration in the reactors.

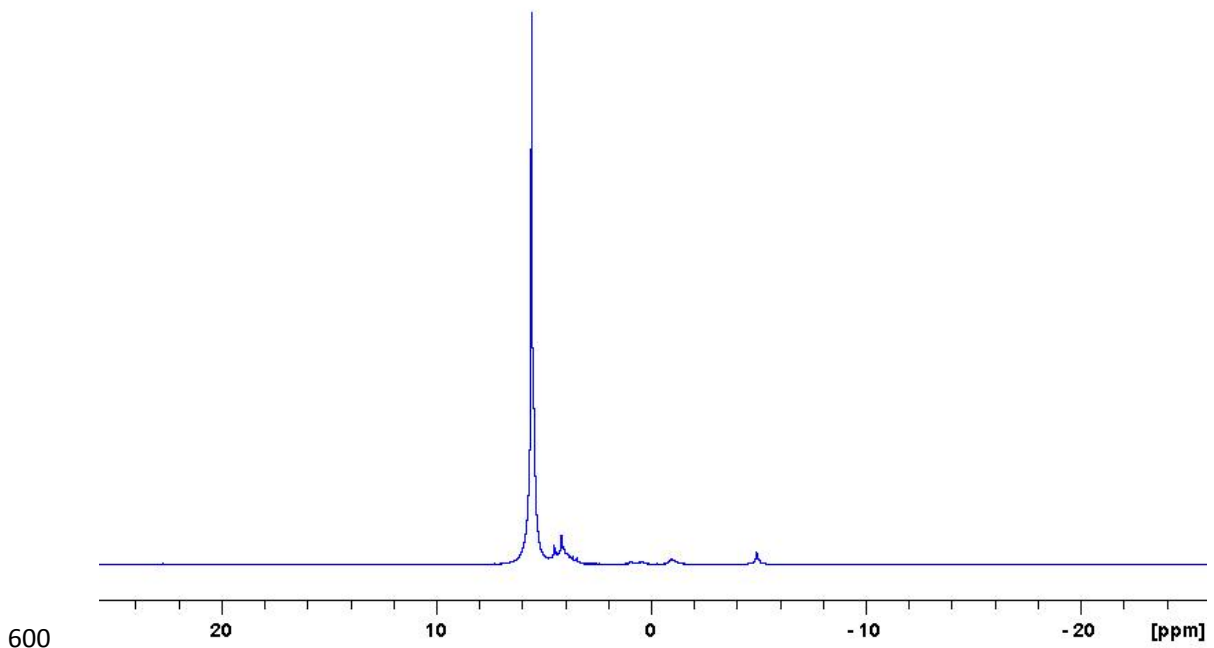
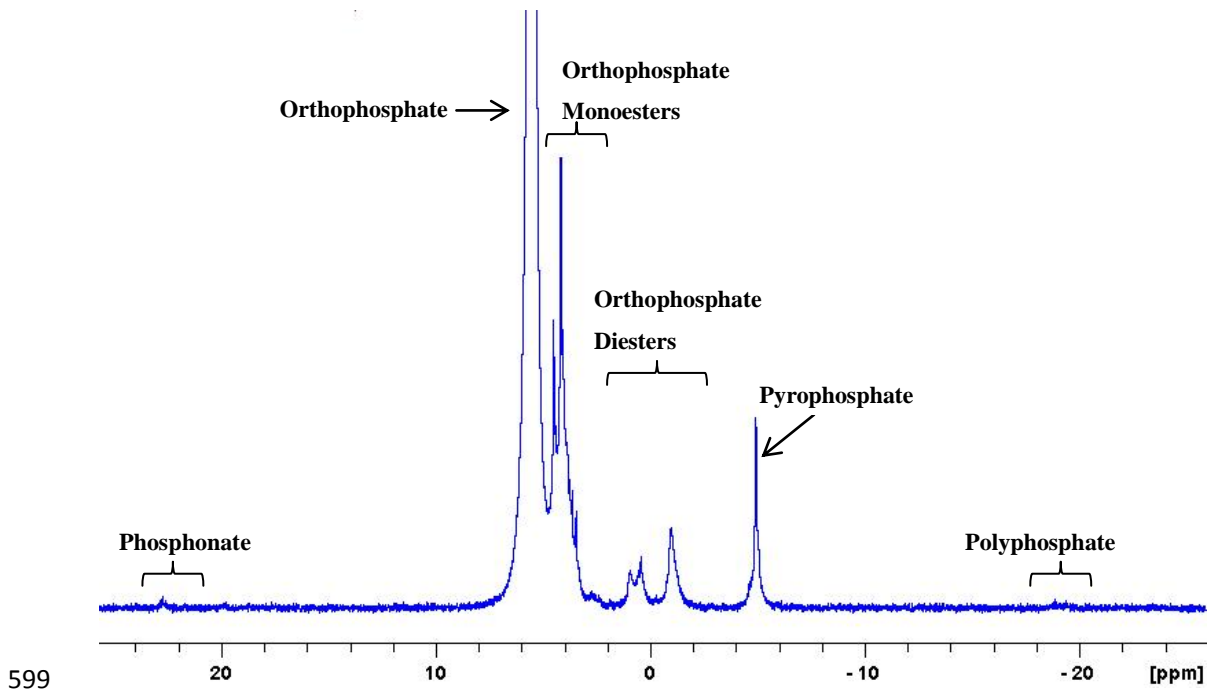


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595 Fig. 4 – XRD diffractogram of granular sludge compared to standard hydroxyapatite
596 and iron phosphate patterns

597
598



601 Fig. 5 – Typical ^{31}P -NMR spectra of NaOH+Na₂EDTA extracts from the aerobic
602 granules sampled on day 120