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1 Effects of the residual ammonium concentration on NOB repression

2 during partial nitritation with granular sludge

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8

9 Abstract

10 Partial nitritation was stably achieved in a bench-scale airlift reactor (1.5L) containing granular sludge. Continuous operation at 20°C treating low-strength synthetic wastewater (50 11 12 mg N-NH₄⁺/L and no COD) achieved nitrogen loading rates of 0.8 g N-NH₄⁺/(L·d) during 13 partial nitritation. The switch between nitrite-oxidizing bacteria (NOB) repression and NOB proliferation was observed when ammonium concentrations in the reactor were below 2-5 mg 14 15 N-NH₄⁺/L for DO concentrations lower than 4 mg O_2/L at 20°C. *Nitrospira* spp. were detected to be the dominant NOB population during the entire reactor operation, whereas 16 Nitrobacter spp. were found to be increasing in numbers over time. Stratification of the 17 18 granule structure, with ammonia-oxidizing bacteria (AOB) occupying the outer shell, was found to be highly important in the repression of NOB in the long term. The pH gradient in 19 20 the granule, containing a pH difference of ca. 0.4 between the granule surface and the granule centre, creates a decreasing gradient of ammonia towards the centre of the granule. Higher 21 residual ammonium concentration enhances the ammonium oxidation rate of those cells 22 23 located further away from the granule surface, where the competition for oxygen between

AOB and NOB is more important, and it contributes to the stratification of both populationsin the biofilm.

26 Keywords: Stratification; pH gradient; *Nitrobacter*; *Nitrospira*; mainstream conditions.

27 **1. Introduction**

48

28 Partial nitritation-Anammox processes are currently under development for the treatment of 29 pretreated sewage (Wett, 2007; Lotti et al., 2014a; Gilbert et al., 2014; Wang et al., 2016; Reino et al., 2016). Advantages of these systems compared to the conventional nitrification-30 31 denitrification treatment are found in economic and environmental aspects. OPEX and 32 CAPEX for nitritation-Anammox can be reduced because of less aeration and COD 33 requirement, and less sludge production. From the environmental point of view, N₂O and CO₂ 34 emissions can be reduced since these greenhouse gasses are not produced in the Anammox process, whereas they are produced during heterotrophic denitrification (Fux and Siegrist, 35 36 2003; Kartal et al., 2010). However, autotrophic nitrogen removal processes in mainstream 37 conditions still cope with some challenges. One of the main problems concerns the process stability in the long term (Winkler et al., 2011; De Clippeleir et al., 2013; Han et al., 2016). 38 Nitrite oxidizing bacteria (NOB) tend to proliferate in long-term partial nitritation operations, 39 40 affecting the process by oxidising nitrite into nitrate and therefore making the effluent 41 unsuitable for further treatment by autotrophic denitrification by Anammox. 42 Process control is needed to repress NOB activity and maintain aerobic oxidation of 43 ammonium into nitrite by ammonium oxidising bacteria (AOB). Proposed NOB repression strategies utilize the control of dissolved oxygen (DO) (Blackburne et al., 2008; Lotti et al., 44 2014b; Ma et al., 2015) or even the DO/ammonium concentrations ratio in the bulk liquid (45 46 Bougard et al., 2006; Bartrolí et al., 2010). These strategies are based on the general reported 47 higher oxygen affinity of AOB compared to NOB (Guisasola et al., 2005; Blackburne et al.,

2008; Pérez et al., 2009). The lower oxygen affinity of NOB together with the oxygen

49 limitation imposed in biofilm systems leads to NOB repression (Garrido et al., 1997; 50 Picioreanu et al., 1997; Sliekers et al., 2005; Peng and Zhu, 2006; Pérez et al., 2009; Brockmann and Morgenroth, 2010, among many others). However, Isanta et al. (2015) 51 52 reported that besides a system operating under oxygen limiting conditions and a higher oxygen affinity for AOB than NOB, a residual ammonium concentration should be 53 maintained in order to keep the growth rate of AOB higher than that of NOB, see Eq. 1. 54 55 Control of the bulk ammonium concentration influences the ammonium oxidation rate. If Eq. 56 1 is used to describe the AOB growth rate, then the residual ammonium concentration affects the ammonium saturation term (or Monod term) and therefore controls the growth rate of 57 58 AOB. Pérez et al. (2014) reported a modelling study in which this concept is used for control of NOB repression. However, until now the influence of the residual ammonium 59 60 concentration on NOB repression was tested mainly in the long term, to obtain stable partial 61 nitritation in mainstream conditions (Isanta et al., 2015; Reino et al., 2016). No further 62 explanations for the success of the strategy and the repression of NOB have been reported.

$$\mu_{AOB} = \mu_{AOB}^{max} \left(\frac{C_{NH_4^+}}{K_{NH_4^+} + C_{NH_4^+}} \right) \left(\frac{C_{O_2}}{K_{O_2} + C_{O_2}} \right)$$
(1)

In this study, a better understanding of the role of the residual ammonium concentration has 63 64 been pursued. Therefore, instead of aiming to demonstrate the long-term stability of the NOB repression (as done recently at low temperatures in Isanta et al., 2015 and Reino et al., 2016), 65 66 assessment of the short term effects of the residual ammonium concentration was specifically targeted. Several techniques were used during the research. Batch test experiments, 67 measurements of the hydroxylamine concentration (an intermediate in nitritation), off-gas 68 69 measurements to monitor NO and N₂O emissions, pH profiles in the granule and FISH on 70 granules slices obtained through cryosectioning were used to investigate the effect of the residual ammonium concentration. Here, we present findings showing the mechanisms that 71

explain the positive effects of the residual ammonium concentration on NOB repression. 72 73 These mechanisms are novel and provide explanation to several reported observations for this type of reactors that were poorly understood. The conclusions of the study provide crucial 74 75 insight in the stability of nitritation and they are very valuable for the next steps in the implementation of anammox in the main water line, to achieve sustainable sewage treatment. 76 77 78 79 2. Materials and Methods 80 2.1 Reactor set-up and inoculum An air-lift reactor with a working volume of 1.5 L was used (Fig. S1). The air flowrate was 81 82 regulated with a mass flow controller (2 L/min capacity, BROOKS, The Netherlands). DO and pH were measured but not controlled. 83 84 The granular sludge was originally obtained from the sidestream reactor in WWTP Olburgen, 85 The Netherlands(Abma et al., 2010). The reactor is performing one-stage nitrogen removal through partial nitritation/anammox process. However, a period of acclimation (ca. two 86 months) of the sludge to mainstream conditions was carried out in the pilot plant of the LIFE 87 project CENIRELTA (Cost Effective NItrogen REmoval by Low-Temperature Anammox) in 88 the WWTP Dokhaven (The Netherlands). The pilot plant treats wastewater obtained from a 89 90 large part of Rotterdam (south, east, centre) after COD removal in a highly loaded aerobic 91 COD removal reactor or A-stage (see a description in Lotti et al., 2014a). When the inoculum was obtained, the effluent concentrations in the CENIRELTA pilot plant were 21 ± 2 mg N-92 93 NH_4^+/L , 0.6 ± 0.3 mg N-NO₂ $^-/L$, 7 ± 1 mg N-NO₃ $^-/L$ and ca. 45 mg COD/L at 23 ± 1 °C. 94 The reactor inoculum was 1 L, containing 4 gVSS/L. Initial maximum activity tests yielded $29 \pm 3 \text{ mg N-NO}_2^{-7}$ (gVSS·d) for AOB, $56 \pm 7 \text{ mg N-NO}_3^{-7}$ (gVSS·d) for NOB and 21 ± 0.6 95

- 96 mg N-NH₄⁺/(gVSS·d) for AMX. At the day of inoculation, the average granule diameter was 97 ca. 0.9 mm.
- 98

99 2.2 Wastewater

- 100 Synthetic wastewater was used containing (per litre of tap water) 0.73 g K₂HPO₄, 0.104 g
- 101 KH₂PO₄, 1.26 g NaHCO₃, 0.236 g (NH₄)₂SO₄, 0.25 mL Fe²⁺-solution and 0.12 mL trace
- elements solution. The Fe^{2+} -solution consisted of (per litre demineralised water) 6.37 g EDTA
- and 9.14 g FeSO₄·7H₂O, and the pH was adjusted to 2.5. The trace elements solution
- 104 contained (per litre Milli-Q water) 19.11 g EDTA, 0.43 g $ZnSO_4 \cdot 7H_2O$, 0.24 g $CoCl_2 \cdot 6H_2O$,
- $105 \qquad 1.0 \ g \ MnCl_2 \cdot 4H_2O, \ 0.25 \ g \ CuSO_4 \cdot 5H_2O, \ 0.22 \ g \ (NH_4)_6Mo_7O_{24} \cdot 4H_2O \ (=1.25 \ mM \ Mo), \ 0.20 \ g \ Mo) \ (=1.25 \ mM \ Mo), \ 0.20 \ g \ Mo)$
- 106 NiCl₂· $6H_2O$, 0.09 g HNaSeO₃, 0.014 g H₃BO₃ and 0.054 g Na₂WO₄· $2H_2O$. The pH was
- adjusted to 6 with solid NaOH.

108

109 **2.3 Reactor operation**

110 The reactor was operated in continuous mode at atmospheric pressure and temperature was 111 controlled at 20°C. At this temperature the advantage of AOB compared to NOB in terms of 112 the maximum specific growth rate is assumed to be rather small (Hunik et al., 1994; Hellinga et al., 1998). The inflow rate was controlled manually (in the range 8-20 L/d) to explore the 113 114 role of the residual ammonium concentration in both the short and long term. During the continuous operation the reactor pH was rather constant at 7.7 ± 0.1 . 115 116 The reactor operation has been divided into 5 phases (Fig. 1). For details of the pseudo-steady states achieved see Table 1. 117

118

119 Calculation of specific ammonium oxidation and nitrate production rates

- 120 To calculate specific rates, the biomass concentration was linearly interpolated and the
- accumulation term was also taken into account, to have a better estimation during transient
- 122 states. For the accumulation term, the first derivative of the (ammonium or nitrate)

123 concentration in time was approached by the incremental ratio: $\frac{dC}{dt} \cong \frac{\Delta C}{\Delta t}$.

124 Diameter distribution

125 The diameter distribution of the granules was measured with the aid of image analysis

126 following the method described in Tijhuis and van Loosdrecht (1994). Surface-based average

127 diameter of the granules was obtained and number of granules and size distribution

128 histograms are detailed in the supplementary information for each one of the measurements.

129 Batch tests

The batch tests were performed in the same (airlift) reactor used for the continuous operation.
Continuous operation was stopped and an ammonium pulse was added. During the batch test
the DO and pH were not controlled. For the Anammox batch test the reactor was switched
from sparging air to supplying nitrogen gas to obtain anaerobic conditions. When the DO was
0%, the medium flowrate was stopped and samples were withdrawn from the top section of
the reactor.

136

137 2.4 Analytical procedures

Ammonium, nitrite and nitrate concentrations were measured offline with Hach Lange cuvette
kits. Dry weight (TSS), ash content and volatile suspended solids (VSS, dry weight minus ash
content) were determined according to standard methods (APHA, 2012). Hydroxylamine
concentrations were measured using a colorimetric method (Frear and Burrell, 1955),

following an *ad hoc* procedure for sample preparation described in Soler-Jofra et al. (2016).
N₂O and NO off-gas concentrations were periodically measured online with a Servomex 4900
infrared gas analyser.

145

146 **2.5 Fluorescence In Situ Hybridization (FISH)**

147 For analysis of the microbial population, the granules were pottered, washed, fixed and loaded onto with gelatine pre-coated Teflon slides according to the procedure described in (Third et 148 149 al., 2001). For cryosectioning of the granules, the granules were washed (3h) in 1x PBS 150 before being fixed (1h). Teflon slides were coated with 0.01% poly-L lysine solution. 151 Granules were put in freeze-medium and cut with a freeze-microtone (Leica CM 1990) at -152 25°C. The obtained slices (10-15 µm thick) were placed on the pre-coated slides and washed 153 with 50% ethanol solution for 5 minutes, to remove the freeze-medium and regain 154 hydrophobicity. Probe hybridization to both pottered samples and cryosectioned slices was 155 again performed as described in (Third et al., 2001). Oligonucleotide probes used are listed in 156 Table S1. Image analysis was done with a Zeiss Axioplan 2 Imaging microscope, together 157 with an AxioCam MRm camera (Zeiss), an ebq100 lamp for fluorescent light and the 158 Axiovision software.

159

160 **2.6 pH profile in the granular sludge**

To determine the pH profile, a granule was fixed in the middle of a flow chamber with a small steel clip (see also the supplementary information, section S1.3). Medium was sparged with air and pumped from the bottom to the top. For the measurements of the pH difference between bulk liquid and granule inside, the pH microelectrode was placed closely above the granule. The pH of the bulk liquid was measured, followed by 1 step of 1000 µm, to measure

the pH inside the granule. The complete experiment was performed at ammonium

167 concentrations of 49 and 11 mg N/L (a different granule was used for each ammonium168 concentration).

169

170 **3. Results**

171 **3.1 Reactor operation**

172 During the entire operation period (223 days) the wastewater inflow rate was used as

173 manipulated variable to control the residual ammonium concentration (Fig. 1A). However,

also the inflow ammonium concentration was lowered from 50 to 40 mg $N-NH_4^+/L$ during

175 phase II (Fig. 1D). The entire performance was divided into 5 phases (Fig. 1), and achieved

pseudo-steady states are summarized in Table 1.

177 Phase I

The start-up period (days 0-11 in phase I, phase I: day 0-67) was used for adaptation of the 178 179 biomass and partial nitritation-Anammox was targeted. Nevertheless, the Anammox activity 180 decreased very fast and eventually was totally lost (see details in section 3.4). As a 181 consequence, nitrite built up in the effluent and the reactor was mainly performing nitritation. 182 From day 50 onwards, the single targeted process was nitritation. The airflow rate was 183 increased step wise to reach a higher DO concentration in the range of $0.7-0.8 \text{ mg O}_2/\text{L}$ (Fig. 184 1C). During days 53 to 67 a pseudo-steady state was reached with reactor and effluent 185 concentrations of $16 \pm 1 \text{ mg N-NH}_4^+/\text{L}$, $24 \pm 2 \text{ mg N-NO}_2^-/\text{L}$, $6 \pm 1 \text{ mg N-NO}_3^-/\text{L}$ and $0.7 \pm 1 \text{ mg N-NO}_3^-/\text{L}$ 186 $0.1 \text{ mg O}_2/L$. This indicates that nitritation was the main process taking place, NOB 187 repression was efficient, although still some residual nitrite oxidation was present. To test the 188 influence of residual ammonium in NOB repression, in a next phase the effluent ammonium 189 concentration was decreased.

190

191 Phase II

192 In phase II (days 68-139) the reactor contained low bulk ammonium concentrations, with an average of ca. 2 mg N-NH₄⁺/L. This was obtained by the decrease in the inflow ammonium 193 194 concentration from 50 to 40 mg N- NH_4^+/L (Fig. 1D). Immediately after the step-down in 195 residual ammonium concentration the nitrate concentration increased (Fig. 1E), although there 196 was not a complete switching towards nitrification, and nitrite was still at high values (ca. 25 197 mg N/L). During days 127 to 137, the residual ammonium concentration decreased and the 198 system switched from oxygen limitation to ammonium limitation resulting in the complete 199 oxidation of ammonium into nitrate (i.e., nitrification). The stoichiometry of the nitrification process makes that 3.43 g O_2/g N-NH₄⁺ are required for the oxidation of ammonium to nitrite 200 and 4.57 g O_2/g N-NH₄⁺ is required for the complete oxidation of ammonium to nitrate. By 201 202 taking into account ammonium and oxygen diffusivities (Picioreanu et al., 1997), the 203 threshold value for the switch from oxygen-limitation to ammonium-limitation could be 204 calculated using Eq. 2 (Harremoes, 1982; Bartrolí et al., 2010).

$$\frac{C_{O_2}}{C_{NH_4^+}} < \frac{\gamma_{O_2/N - NH_4^+} D_{NH_4^+}}{D_{O_2}} = \frac{3.43 \times 1.9 \times 10^{-4}}{2.2 \times 10^{-4}} = 3.0 \frac{gO_2}{gN}$$
(2)

205 During the last part of phase II the values of the DO/ammonium concentrations ratio exceeded 206 $3.0 \text{ g O}_2/\text{g N}$, meaning the switch from oxygen limitation to ammonium limitation (Fig. 1B). 207 Due to ammonium limitation, the ammonium oxidation rate decreased and the DO 208 concentration increased. For days 117-139 a pseudo-steady state was reached with 209 concentrations of $0.8 \pm 0.3 \text{ mg N-NH}_4^+/\text{L}$, $24 \pm 11 \text{ mg N-NO}_2^-/\text{L}$, $14 \pm 11 \text{ mg N-NO}_3^-/\text{L}$ and 210 $1.7 \pm 1.0 \text{ mg O}_2/\text{L}$. When bulk ammonium concentration reaches such low values, NOB 211 repression is not possible, and therefore most ammonium is converted to nitrate. 213 Phase III

214 In the beginning of phase III (phase III: day 140-168) the bulk ammonium concentration was 215 increased to ca. 12 mg N/L. The system switched from ammonium limitation to oxygen 216 limitation (see Fig. 1). During phase III intentional disturbances in the residual ammonium 217 concentration were targeted (see section 3.2 for further explanations about short term effects). 218 Therefore no steady state was achieved. Nitrate built up at higher concentrations when 219 residual ammonium concentration was slightly decreased, indicating a direct and fast effect 220 between high residual ammonium and NOB repression. The fast transitions (within 24 hours) 221 cannot be explained by a community shift. 222 At day 141, due to increasing biomass activity, the inflow rate needed to maintain a certain 223 ammonium effluent concentration had increased to levels that gave practical problems. 224 Therefore, roughly half of the biomass was removed from the reactor to be able to operate at 225 lower inflow rates again (Fig. 1A). After day 151 the airflow rate was increased from 4.2 to 226 6.6 L/h. The DO concentration was increased to enhance the activity of AOB to better 227 develop the AOB layer on the granule surface and completely outcompete NOB from the granule surface. At day 168 the inflow rate was lowered again to ca. 10 L/d to decrease the 228 229 residual ammonium concentration.

230

231 Phase IV

During phase IV (day 169-186) an average bulk ammonium concentration of ca. 2 mg N/L
was reached (Table 1). NOB activity increased rapidly and effluent nitrate concentration
increased to ca. 36 mg N/L (day 186). This was indicating that an ammonium concentration
of ca. 2 mg N/L was not high enough to repress NOB effectively, even under oxygen

limitation. During days 175 to 182 a pseudo-steady state was reached with concentrations of $1.8 \pm 0.1 \text{ mg N-NH}_4^+/\text{L}$, $27 \pm 6 \text{ mg N-NO}_2^-/\text{L}$, $21 \pm 7 \text{ mg N-NO}_3^-/\text{L}$ and $3.6 \pm 0.2 \text{ mg O}_2/\text{L}$. Also during phase IV, the specific biomass activity had increased by more than the double compared to the specific biomass activity before the removal (Table 1).

At day 173 it was noted that the effluent tube, from which samples were withdrawn,

241 contained biofilm which contributed to the measured concentrations of ammonium, nitrite and nitrate. Comparison of a sample after the effluent tube and a sample directly from the reactor 242 243 provided the insight that during the previous measurements, in general the ammonium and 244 nitrite concentration were underestimated (measured errors of ca. 3 mg $N-NH_4^+/L$ and 3 mg N-NO₂/L) and the nitrate concentrations were overestimated (measured error of ca. 7 mg N-245 246 NO_3^{-}/L), indicating that the NOB repression was effective in the reactor, but not in the tube-247 biofilm. From day 173 onwards samples used for the water quality measurements were 248 withdrawn directly from the top section of the reactor. The measured errors were evaluated once the biofilm grown on the tube developed for more than 4 weeks, providing the 249 250 maximum possible bias. In earlier stages of biofilm development on the inner tube wall, a 251 more reduce impact on the results presented is expected.

252

253 Phase V

At day 187 (start of phase V, phase V: days 187-223) the bulk ammonium concentration was increased from ca. 2 to ca. 25 mg N/L. The change in the residual ammonium concentration resulted in a very fast NOB repression, and effluent nitrate concentration rapidly decreased. After a week of operation the DO was decreased by lowering the airflow rate from 6.6 to 4.2 L/h to repress even more the NOB activity. A slight decrease in the nitrate concentration during this phase was observed. A pseudo-steady state was obtained (days 193-214), 27.2 ± 0.8 mg N-NH₄⁺/L, 17.1 ± 1.6 mg N-NO₂⁻/L, 4.9 ± 1.3 mg N-NO₃⁻/L and 2.7 ± 0.5 mg O₂/L. 261

262 **3.2** Short-term effects of the residual ammonium concentration

Especially during phases II and III of the continuous operation, the residual ammonium 263 264 concentration influenced the nitrate build-up (Fig. 1E). With increasing and decreasing 265 ammonium concentrations, a fast (inverse) response was measured for nitrate concentrations. 266 The corresponding change in the nitrate concentration resulted from the change in the ammonium oxidation rate of AOB (Table S2). Higher specific ammonium oxidation rates 267 268 were observed when residual ammonium concentration was increased, which contributed to a 269 lower DO concentration (Table S2). In parallel with the short term increase on the specific 270 ammonium oxidation rate, a decrease in specific nitrate production rate was measured (Table 271 S2). The change in the bulk ammonium concentrations impacts the nitrate concentration 272 immediately, in a period of hours. This fast response is a clear indication that the residual 273 ammonium concentration can be used as controlled variable for nitritation as pointed out 274 previously (Jemaat et al., 2013).

275 Additionally, to present in a more direct way the short term effects of residual ammonium 276 concentration on NOB repression, all data from day 50 onwards has been plotted in Fig. 2A. 277 There is a clear trend in Fig. 2A, showing how NOB repression is achieved at ammonium 278 concentrations higher than ca. 5 mg N/L, regardless to the DO concentration applied, which 279 overall was in a wide range, from $0.7-3.7 \text{ mg O}_2/\text{L}$. When the time between measurements 280 was less than 1 day, the corresponding data were highlighted in Fig. 2. For those points the 281 sample was withdrawn 2.5 hours after the previous measurement, which is in the order of 282 magnitude of the hydraulic retention time, therefore too short to washout the nitrate accumulated at low residual ammonium even if NOB repression is effective. 283

For comparison, a similar graph was plotted by including the bulk DO/ammonium

concentrations ratio in the bulk liquid in Fig. 2B. In the inset graph in Fig. 2B a zoomed in

version of the graph is also given. The correlation between the bulk DO/ammonium
concentrations ratio and NOB repression is less evident (compared to Fig. 2A), mainly due to
the scale and the effect of the ratio itself, which produces small values at high bulk
ammonium concentrations. For values of the ratio lower than 1, NOB repression is more
effective (Fig. 2B, inset graph).

291

Batch test

A batch-test was performed at day 159 (Fig. 3) to further investigate the residual ammonium concentrations range causing the switch from effective NOB repression to nitrate production. An ammonium pulse was added after the inflow rate was stopped (time zero in Fig. 3). For bulk ammonium concentrations in the range 2-4 mg N/L the nitrate concentration increased at a higher rate (in accordance with the continuous operation results in Fig. 3), indicating the ammonium concentration causing the switch between effective NOB repression and nitrification was occurring.

300 The oxygen consumption rate increased ca. 8% immediately after the ammonium pulse.

301 Interestingly, when at t= 45min the bulk ammonium concentration is back to the initial 10 mg

302 N/L, the DO concentration is still well below the initial value, as indicated in Fig. 3 by ΔDO .

303 This increased oxygen consumption rate at the same bulk ammonium concentration (10 mg

N/L happens despite the pH (which is not controlled) decreased by ca. 0.2.

305

306 Step-up increase in residual ammonium concentration

307 The step-up disturbance in the bulk ammonium concentration at day 187 produced a decrease

in the DO concentration due to the increase in specific ammonium oxidation rate (Fig. 4A).

309 As a result, the nitrate concentration rapidly decreased (Fig. 4A). The stabilization of the

310 ammonium oxidation rate occurred several days after the step-up disturbance, with higher 311 rates measured immediately after the disturbance (Fig, 4A). Interesting to emphasize that the 312 DO concentration decreased only during the transient state (3 days). Hydroxylamine at steady 313 state conditions was not detected throughout the operation period. However, the increase in 314 residual ammonium concentration after the step-up disturbance, resulted in hydroxylamine released into the bulk liquid, achieving a maximum value of 0.056 mg N-NH₂OH/L after 7 315 316 hours (Fig. 4C). However the monitoring of the hydroxylamine was not continued until the 317 next morning, when hydroxylamine was not detected anymore. In addition, an increase in N₂O emission was also measured in the off-gas (Fig. 4B). No significant nitric oxide (NO) 318 319 emission was observed. During the stabilisation of the residual ammonium concentration the N₂O emissions decreased again. 320

321

322 **3.3** Biomass characteristics and sludge retention time

323 The biomass concentration in the reactor was plotted in Fig. 1A. The average diameter of the 324 granules was 0.9, 1.4 and 1.3 mm, at days 0, 47 and 123 respectively (Table S2). Due to the 325 wide size distribution, the full size distribution curve was presented in the supplementary 326 information (Figs. S2-S4). Sludge retention time was 75 days at day 55 of continuous operation. From day 118 onwards stabilized at ca. 210±18 days. An average solids 327 concentrations mass ratio of 0.91 gVSS/gTSS was determined. A clear colour change of the 328 329 biomass over time was noticeable (Figs. S5-S7). At the day of inoculation the granules had a dark (brownish) colour, indicating the presence of heterotrophic bacteria near the granule 330 331 surface and not stratification of an AOB layer. Over time the granules became orange 332 coloured indicating the presence of active AOB bacteria in the outermost layer of the 333 granules.

334

335 **3.4 Fluorescence In Situ Hybridization (FISH)**

336 Cryosectioned samples of the granular sludge were used for FISH analysis. Granules from 337 day 148 and 223 were obtained from periods at high residual ammonium concentration, 338 whereas those at day 187 were from a period at low residual ammonium concentration (see Fig. 1E). The granule structure from the three samples was highly similar (Fig. 5), presenting 339 340 a clear stratification: a shell consisting of AOB colonies and behind it, the majority of the 341 NOB colonies. The size of AOB and NOB microcolonies was difficult to measure on the 342 pictures, because individual colonies were difficult to distinguish in both layers, but in 343 particular in the AOB shell. Comparing the granule structure obtained in this study (Fig. 5) to 344 the original inoculum (Fig. S8), the degree of stratification was enhanced during the operation 345 of the reactor. 346 Regarding the predominant NOB species in the granular sludge, at day 148 only Nitrospira

spp. were detected (Fig. S9) (but not *Nitrobacter* spp.). However, at day 223 both *Nitrospira*spp. and *Nitrobacter* spp. were detected (Fig. S10). *Nitrobacter* spp. were found in lower
amounts than *Nitrospira* spp., indicating the development of this population during the reactor
operation.

Although the quantification of the relative abundances of AOB and NOB in the granular
sludge was not specifically targeted, a healthy NOB population was retained in the granular
sludge during the whole period of operation, since a very fast and significant nitrate
production was noticeable as soon as the imposed conditions did not efficiently repress NOB.

356 **3.5 pH gradient in the granule and apparent ammonium half-saturation coefficient**

357 The gradient of pH in the granule was assessed by measuring the pH difference between the 358 core of the granule and the bulk liquid, for a pH range of 7.0-8.4 (a complete pH profile in a 359 granule is also presented as an example, see Fig. S11). Granules were withdrawn from the 360 reactor during phase V. The ammonium consumption in the measuring chamber was 361 negligible. For the entire investigated range of bulk pH, a lower pH was measured inside the 362 granule (Fig. 6). The pH curves in Fig. 6 show that at a pH in the bulk of 7.7, which is the pH 363 inside the reactor during continuous operation, the pH difference between the bulk liquid and 364 inside the granule was 0.44 for both ammonium concentrations tested (11 and 49 mg N/L). A rough estimation of the AOB apparent half-saturation coefficient for ammonium $(K_{S,NH4+}^{App})$ 365 was obtained by using a ratio of average specific ammonium oxidation rates (Eq. 3). These 366 AOB rates 17.3 ± 0.4 mg N-NH₄⁺/(g VSS·h) from days 158-165 with an average ammonium 367 concentration of 9 mg N-NH₄⁺/L ($r_{AOB}^{9mgN/L}$), and 11.2±0.2 mg N-NH₄⁺/gVSS/h from days 368 175-187 with an average ammonium concentration of 2 mg N/L ($r_{AOB}^{2mgN/L}$)) were obtained 369 370 from periods with different ammonium concentrations, but with similar bulk DO concentrations, in order to simplify for the oxygen Monod term (see Eq. 1). Solving Eq. 3 371 resulted in a $K_{S,NH4+}^{App}$ of 1.7 mg N-NH₄⁺/L. 372

$$\frac{r_{AOB}^{2mgN/L}}{r_{AOB}^{9mgN/L}} \approx \frac{\frac{2}{(K_{S,NH4+}^{App} + 2)}}{\frac{9}{(K_{S,NH4+}^{App} + 9)}}$$
(3)

373

374 3.6 Anammox

375 Within a couple of weeks after inoculation, Anammox activity in the reactor was lost. Until

day 12 during the start-up of the reactor, Anammox activity increased as can be seen from the

377 nitrogen balance (Fig. 1D). From day 12 onwards, the activity decreased.

At day 48 an anoxic batch-test was performed (see results in Fig. S12). During the test no
clear signs of Anammox activity were detected. The decrease in ammonium, nitrite and
nitrate concentrations are possibly linked to salt precipitation (for instance struvite).
Ammonia stripping could also have contributed to decrease the ammonium concentration in
time. FISH results from day 167 of the operation also showed a significant amount of dead
cell material (no hybridization with EUB338), whereas FISH results from the last day of
operation confirmed the decay of Anammox (no hybridization with AMX820) (Fig. S13).

385

386 4. Discussion

387 4.1 Nitritation and NOB repression

The control of the residual ammonium concentration confirmed its effectiveness on NOB repression at 20°C and pH 7.6-7.8. Stable nitritation was maintained above bulk ammonium concentrations of ca. 5 mg N/L and nitrate production was enhanced at a residual ammonium concentration of ca. 2 mg N/L (Fig. 2A).

392 In the conditions tested, rather than the DO/ammonium concentrations ratio (Fig. 2B), the ammonium concentration was the main factor regulating NOB repression (Fig. 2A). The 393 394 DO/ammonium concentrations ratio required for efficient NOB repression was ca. 1 mg 395 O₂/mg N or lower (Fig. 2B). Bartrolí et al., 2010 operating at 30°C found that the required value of the ratio was ca. 0.18 mg O₂/mg N or lower. Reasons for this difference remain until 396 397 now unclear. We hypothesize that the difference in behavior comes from the difference in 398 granule structure. In our study, the inoculum was a granular sludge containing anammox in 399 the granule core (Fig. 5). However, in Bartrolí et al. (2010), or in similar trials using the 400 DO/ammonium concentration ratio as main criterion, the granular sludge did not contain 401 anammox.

The production of nitrate in the biofilm grown on the effluent tube inner wall is probably due to the diffusion of oxygen through the tube wall (that type of silicone tube is permeable to oxygen). The counter-diffusion of oxygen makes oxygen available to NOB and stratification is useless to keep nitritation stable.

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407

408 **4.2 Stratification of AOB and NOB populations**

Stratification of AOB and NOB populations in granular sludge has been sometimes reported 409 410 when removing nitrogen through one stage partial nitritation / anammox (Vlaeminck et al., 2010; Winkler et al., 2011). In such systems, anammox bacteria are located in the core of the 411 412 granule and act as a sink for nitrite, facilitating NOB repression and perhaps stratification. 413 Nevertheless, for nitrifying granules, to the best of our knowledge, only one study reported 414 stratification of AOB and NOB in granular sludge reactor (Tsuneda et al., 2003). Their 415 granular sludge was cultivated in an aerobic upflow fluidized bed treating high strength 416 ammonium wastewater. The reasons why the stratification developed and the significance of 417 their findings were not discussed, not even in subsequent reports when mathematical 418 modelling was used to describe the experimental findings (Matsumoto et al., 2010). In fact both mathematical models used (one and two dimensional biofilm models) failed to describe 419 420 the stratification (Matsumoto et al., 2010). In this study, we found stratification of AOB and 421 NOB for the first time when treating low strength wastewater and operating at 20°C. There 422 are two aspects associated to the stratified structure: (i) the position of the AOB 423 microcolonies is better for oxygen competition because they are much closer to the granule 424 surface, enhancing NOB repression; (ii) the outer dense AOB shell acts as a protective layer for NOB microcolonies against detachment, delaying washout of NOB from the granular 425 426 sludge.

In such stratified granule, the oxygen penetration depth could therefore play a clear role in
NOB repression. When AOB preferentially occupy the external shell of the granule, the
competition for oxygen between AOB and NOB is deeply impacted, as demonstrated through
a 3-dimensional modelling study in which the effect of the presence of cell clusters was
specifically targeted (Picioreanu et al., *submitted*).

432 Secondly, the NOB colonies occupying inner layers are protected against detachment. Their 433 residence time in the reactor is expected to be longer than that of AOB. Moreover, a larger 434 cluster size (compared to that of AOB microcolonies) could be achieved in time. In general, 435 larger NOB colonies behind the AOB layer would be easier to repress due to smaller surface 436 to volume ratios. However, due to the intensity of the signal, it is not possible to estimate a 437 representative average size for AOB and NOB cell clusters, and therefore this hypothesis 438 could not be proven at this stage.

439 In this type of granular sludge, NOB is known to persist for long periods of time (several 440 months), despite nitrate production was measured to be at very low levels (Bartrolí et al., 441 2010; Lotti et al., 2014b; Isanta et al., 2015, among others). In our study, also the same trend 442 is observed. This would indicate an alternative metabolic NOB route to survive in absence of 443 oxygen. The ability of some NOB to reverse their main oxidative reaction (i.e. to reduce 444 nitrate into nitrite) has been reported, when there is absence of oxygen but availability of 445 COD (e.g. formate) (Koch et al., 2015). In this case, where an autotrophic synthetic medium 446 is used, this possibility might be only plausible if NOB could use the organic matter formed 447 from decay products. Additionally, complete ammonium oxidation (comammox) Nitrospira 448 were found at high abundances in an autotrophic culture in anoxic conditions, although their 449 primary metabolic route remained unknown (van Kessel et al., 2016).

Some NOB colonies were located closer to the granule surface, surrounded by AOB colonies
(Fig. 5). These NOB colonies were assumed to be the reason for the residual nitrate
concentration in the reactor.

453 Previous studies reported the presence of Nitrobacter spp. as the dominant NOB species when 454 controlling the residual ammonium to repress NOB and hypothesized that a prerequisite to 455 obtain stable partial nitritation could be to select *Nitrobacter* spp. instead of *Nitrospira* spp. (Isanta et al., 2015). Wang et al. (2016) reported that the strategy of controlling residual 456 ammonium at high concentrations would only be successful in the case of Nitrobacter spp. (r-457 458 strategist) being the dominant NOB population. However, here we found that a high residual ammonium concentration enhanced AOB stratification in the external granule layer, which 459 460 demonstrated to be a successful strategy independently of the initial NOB genus found in the 461 sludge.

462

463 **4.3 Linking the effects of the DO/ammonium concentrations ratio to stratification**

464 Higher residual ammonium concentrations result in higher ammonium oxidation rates (Table 1, Fig. 4A, Table S2) which in turn would allow to apply higher DO concentrations in a 465 reactor without compromising the stability of nitritation (in agreement with Bartrolí et al., 466 2010). Simply because the oxygen penetration depth is shorter at higher ammonium oxidation 467 468 rates. This is therefore the fundamental mechanism explaining the correlation found between 469 the bulk DO/ammonium concentrations ratio and NOB repression in Bartrolí et al. (2010). In 470 that study, at 30°C, NOB repression was achieved at residual ammonium concentration of 40 mg N/L and DO = 7 mg O_2/L (DO/ammonium= 0.18 g O_2/g N) and for 20 mg N/L and DO = 471 472 5 mg O_2/L (DO/ammonium= 0.25 g O_2/g N), but complete nitrification at residual ammonium concentration of 20 mg N/L and DO = 7 mg O_2/L (DO/ammonium= 0.35 g O_2/g N). 473

To effectively repress NOB in wastewater treatment systems containing granular sludge, 474 475 stratification of AOB and NOB inside the granule structure is identified here as a requirement. 476 Without the stratification, NOB colonies can grow closer towards the granule surface where 477 they have better access to oxygen resulting in nitrate production. This is in agreement with the 478 assessment of oxygen competition through 3-D modelling of granules containing cell clusters 479 (Picioreanu et al., *submitted*). A complete and dense AOB layer on the granule surface would 480 result in a limited oxygen penetration depth, and no oxygen available for the inner layers 481 where NOB are located. Stratification of AOB on the granule surface can be created by 482 operating at high residual ammonium concentrations, to enhance high ammonium oxidation 483 rates.

484 By applying high residual ammonium concentrations, AOB consume most of the oxygen 485 resulting in the repression of NOB. When in time the nitrate production becomes low enough, 486 indicating good stratification, the possibility arises to decrease the residual ammonium 487 concentration. However, the residual ammonium concentration has its lower limits for 488 successful NOB repression, as reported in this study. Maintaining a high residual ammonium 489 concentration would not be preferred in all autotrophic nitrogen removal systems. The 490 strategy would be suited for a two stage nitrogen removal process, where in the first stage 491 partial nitritation is desired (so in combination with Anammox in a second stage). In this 492 system the residual ammonium concentration has to be high, due to design requirements, 493 since only 50% of the ammonium has to be oxidised to nitrite in order to supply Anammox 494 with the right distribution in N substrates. However, this strategy would not be suited for 495 single stage autotrophic nitrogen removal, as high residual ammonium concentrations in this 496 system are not desired, since the aim is the removal of nitrogen from the wastewater. Plug-497 flow hydrodynamics or SBR operation could be used instead in one-stage nitrogen removal 498 systems, to enhance the use of high residual ammonium concentrations as previously

highlighted in the literature (Pérez et al., 2014). For full scale applications, diurnal variability
of the wastewater, seasonality and rainy events might be also hampering the control of the
residual ammonium concentration in the partial nitritation reactor (Pérez et al., 2015). The use
of reject water might assist to overcome (some of) these issues, as already assessed by
mathematical modelling (Pérez et al., 2015).

504

505 **4.4 pH gradient in the granule**

506 Because ammonia is reported to be the true substrate for AOB (Suzuki et al., 1974), the lower 507 pH inside the granule leads to a lower ammonia concentration in the inner parts due to the 508 ammonium-ammonia acid-base equilibrium. The pH difference between bulk liquid and 509 granule core ($\Delta pH = 0.44$ see Fig. 6) was in the same range found for similar systems (de 510 Beer et al., 1993; Gieseke et al., 2006; Schreiber et al., 2009; Uemura et al., 2011; Winkler et 511 al., 2011) or calculated through mathematical models (Park et al., 2010). Since an increase in 512 the bulk ammonium concentration results in higher ammonium oxidation rates, the pH 513 towards the centre of the granule would decrease even further due to the increase in proton 514 production by AOB. Therefore, higher residual ammonium concentrations lead to an even higher K_{S,NH4+} value towards the centre of the granule due to the larger decrease in pH in 515 516 these regions, making these inner located cells even less saturated in ammonia. This creates 517 the possibility of further increases in the residual ammonium concentration to obtain higher ammonium oxidation rates, resulting in both enhancement of the stratification and in NOB 518 519 repression. The limitation of the enhancement of the rate is that at pH too distant from the 520 optimal pH range of AOB, the maximum specific ammonium oxidation rate would 521 significantly decrease.

522 Suzuki et al. (1974) measured how the ammonium half-saturation coefficient (K_{S,NH4+},

expressed in units of nitrogen ammonium) changes with pH. The lower pH leads to a higher K_{S,NH4+} value inside the granule. With use of the measured pH gradient, the pH effect on the ammonium half-saturation coefficient for AOB ($K_{S,NH4+}$ (pH)) was assessed (see Eqs. S1-S2 in the supplementary information, section S2.8) (Table 2).

527 The apparent ammonium half-saturation coefficient would increase by a factor of 2.7 times 528 with a decrease in pH of 0.44 (Table 2), indicating that AOB cells exposed to a lower pH 529 (those located further away from the granule surface) could be less saturated in ammonium 530 than those at the granule surface. Therefore, these cells would have an advantage when the 531 bulk ammonium concentration is increased (see the corresponding change in the ammonium 532 Monod term in Table 2).

However, the pH also affects the maximum specific growth rate of AOB (μ_{max}^{AOB}). To assess 533 the overall impact of pH on the ammonium oxidation rate, the influence on both μ_{max}^{AOB} and 534 535 K_{S.NH4+} was taken into account as shown in Table 2. Values were used to assess qualitatively 536 how the pH gradient could explain the increase in oxygen consumption detected in the batch test presented in Fig. 3. Comparing only the ammonium Monod term at the pH of the granule 537 538 core for ammonium concentrations of 10 and 20 mg N/L, there is a clear advantage (16% increase). Nevertheless, the μ_{max}^{AOB} value is also smaller at the lower pH (with a decrease of ca. 539 540 -15%, between pH of the bulk and pH of the granule core, see Table 2), which would decrease 541 the overall contribution to the observed ammonium oxidation rate. Also for the batch test 542 conditions, the bulk DO decreased from 3.1 to 2.6 mg O_2/L , which should also penalize the 543 ammonium oxidation rate through the oxygen Monod term (see Eq. 1), even more for cells in the inner layers, at a lower pH. Additionally, the pH decreased just after the pulse. 544 545 Interestingly, despite the negative effects (decrease in DO and pH), the oxygen consumption rate increased. 546

547 When the microsensor is used into the granule for measuring the pH, it is unlikely that the 548 microcolonies (i.e. the dense cell clusters in which AOB and NOB grow in the biofilm) are 549 perforated, due to the strong adhesion properties of the EPS in the microcolony (Larsen et al., 550 2008). The microsensor tip probably would push away those colonies. The pH profile inside 551 the microcolony is therefore expected to be even steeper than that measured in the biofilm 552 matrix, because of the high density in the cell cluster (ca. 600 gCOD/L, Coskuner et al., 553 2005). Therefore, although the pH gradients are here discussed as being one dimensional 554 along the biofilm depth, they would also develop inside the colonies. This applies not only for pH, but also for oxygen and substrate. 555

556 Overall, a truly quantitative impact of the pH gradient on AOB activity is at this stage not 557 conclusive. It would require of three-dimensional biofilm modelling, including the description 558 of the cell clusters. The model might help to clarify if the pH gradient would explain the 559 higher measured oxygen consumption and the higher ammonium oxidizing rates when 560 residual ammonium concentrations are increased.

561 **4.5 Ammonia gradient in the granule**

562 The ammonia gradient in the granule is influenced by both diffusion and the pH gradient. 563 Through diffusion the ammonium concentration tends to decrease in the inner layers of the 564 granules (i.e. ammonia is consumed by AOB, and overall the total ammoniacal nitrogen is 565 therefore decreasing). However, the expected decrease would be rather low, because oxygen 566 is stoichiometrically limiting. Additionally, the pH decreases in the inner layers of the granule 567 due to the protons produced by AOB. Therefore at a lower pH the fraction of free ammonia is 568 even lower. The effect of the pH dominates the gradient of ammonia. To numerically clarify 569 the contributions, we used as example the following conditions: $DO = 3.5 \text{ mg } O_2/L$ and 570 temperature 20°C. Assuming a concentration of 20 mg N/L and pH 7.7 in bulk liquid, the free 571 ammonia concentration is 0.67 mg N/L (see Table S3). Since the oxygen is limiting and the

stoichiometry of the nitritation makes that 3.43 g O_2/g N-NH₄⁺ are required for the oxidation 572 573 of ammonium to nitrite. Using this factor, with the assumed DO (3.5 mg O2/L), the decrease 574 in ammonium would be ca. 1 mg N/L. Therefore the gradient of ammonia coming from the 575 decrease due to consumption by AOB (i.e. diffusion limitation) would be only 0.02 mg N-576 NH₃/L. Assuming a decrease in the pH from 7.7 to 7.26, the decrease in ammonia would be of 577 0.3 mg N-NH₃/L, being therefore 15 times larger than the gradient due to diffusion limitation. 578 Even considering oxygen saturation, the decrease in ammonium would be from 20 to 17.7, 579 which would mean a decrease in ammonia of 0.05, still three times lower than the effect of pH. In conclusion, the gradient of ammonia is dominated by the pH gradient, rather than due 580 581 to diffusion limitation (due to ammonia consumption by AOB). However, both effects contribute and decrease the ammonia towards the inner layers of the granule. 582

4.6 Implications of hydroxylamine release after a step-up increase in residual ammonium concentration

585 Hydroxylamine has been reported to be able to increase the AOB growth rate, in case of the mixotrophic growth of AOB on ammonia and hydroxylamine under substrate-limited growth 586 conditions (De Bruijn et al., 1995; Harper et al., 2009). Hydroxylamine produced an increase 587 588 in the ammonia uptake rate of AOB in the short term (De Bruijn et al., 1995). In addition, 589 hydroxylamine has been reported to be highly inhibitory for NOB (Yang and Alleman, 1992; 590 Blackburne et al., 2004; Noophan et al., 2004). Both effects of hydroxylamine could in theory 591 support the repression of NOB, when increasing the residual ammonium concentration from 592 low concentrations to a high residual ammonium concentration. The hydroxylamine that is 593 temporarily accumulated (as reported in this study), could enhance the growth rate of AOB 594 and simultaneously inhibit NOB.

595 The strong gradients of oxygen and pH that develop in the dense AOB cell clusters might596 create different niches, in which hydroxylamine released by ammonia saturated cells might be

597 cometabolized by other AOB cells, that are more interior in the AOB layer, or in the cell 598 cluster. This cometabolization would require of cells that have oxygen availability, but still 599 are not suffering ammonium saturation. This is plausible given the pH gradient found, where 600 the ammonium saturation condition depends on the pH, as already discussed. In addition, 601 studies of the kinetics and pH-dependency of ammonia and hydroxylamine oxidation by 602 *Nitrosomonas europaea* revealed that hydroxylamine oxidation is moderately pH-sensitive, 603 whereas ammonia oxidation decreases strongly with decreasing pH (Frijlink et al., 1992). 604 Which would support that, the steep pH gradients produce a pool of ammonia non-saturated 605 cells that use hydroxylamine in aerobic environments without being much affected by the low 606 pH values attained. This hypothesis would therefore provide a new mechanism for the 607 positive effects of applying high residual ammonium concentrations for NOB repression. This 608 could be linked with the transient effects of the increase in residual ammonium concentration 609 as highlighted in the short term effects (Fig. 3 and 4). Particularly interesting is the large 610 increase in the specific ammonium oxidation rate (from 11 to 21mg N/(g VSS·h), Fig. 4) 611 during the first hours after the increase in ammonium (Fig. 4). The specific ammonium 612 oxidation rate was calculated also based on the nitrite and nitrate production (summing up both, Fig. 4A), to rule out any potential absorption process in the granular sludge, since 613 614 ammonium absorption in granular sludge is known to happen (Bassin et al., 2011). However, 615 further research is required to be able to obtain conclusive evidence about the effects of the 616 hydroxylamine release on the ammonium oxidation rate.

Hydroxylamine diffusing to deeper layers (either in the granule or in the AOB cell cluster)
where there is no oxygen availability triggers nitrifier denitrification, since nitrite is also
present, as suggested previously for biofilms in a theoretical model based study (Sabba et al.,
2015). Therefore the simultaneous detection of hydroxylamine and a significant increase in
N₂O emissions, could be associated to the nitrifier denitrification pathway.

622	NOB inhibited by hydroxylamine produced by AOB would not be a very plausible
623	explanation, because the levels detected in this study are very low as to be inhibitory
624	(Blackburne et al., 2004; Noophan et al., 2004). In addition, for long term exposure to the
625	inhibitory compound, acclimation of the bacteria would be expected.
626	
627	5. Conclusions
628	• The control of the residual ammonium concentration has proven to be effective for
629	repression of Nitrospira spp. at 20°C. The switch in NOB repression to NOB
630	proliferation was determined to be located in a bulk ammonium concentration range of
631	2-5 mg N/L for DO concentrations lower than 4 mg O_2/L .
632	• Operating at higher residual ammonium concentration triggers higher ammonium
633	oxidation rates and higher oxygen consumption rates, both in the short and long term.
634	• Stratification of an outer AOB layer in the granule structure was found to be highly
635	important to maintain stable partial nitritation in the long term. The AOB layer is
636	important to achieve oxygen limitation for NOB due to the oxygen penetration depth
637	in combination with bulk ammonium concentrations which are high enough to prevent
638	rate-limiting conditions for AOB.
639	• The pH gradient found provides an explanation for the direct effect of residual
640	ammonium in the ammonium oxidation rate, because cells located further away from
641	the granule surface are less saturated in ammonia due to the decrease in pH. This
642	contributes to NOB repression.

643

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TABLES

Table 1. Characterization of the pseudo-steady states attained during reactor operation: average concentrations of N-compounds and DO, specific ammonium oxidation rate (r_{AOB}) and specific nitrate production rate (r_{NOB}). Phase III did not achieve a pseudo-state state operation; therefore, no details are given in this table but the measurements for phase III are shown in Fig. 1.

Phase	Period	$\mathbf{NH_4}^+$ Eff	NO _{2 Eff}	NO _{3 Eff}	Tot-N _{in}	Tot-N _{out}	DO	r _{AOB}	r _{NOB}
	(d)	(mg N/L)	(mg N/L)	(mg N/L)	(mg N/L)	(mg N/L)	$(mg O_2/L)$	(mg N/(gVSS·h))	(mg N/(g VSS·h))
Ι	53-67	16 ± 0.9	24 ± 2	6.0 ± 1	51 ± 0.8	46 ± 0.9	0.7 ± 0.1	5.7 ± 0.6	1.0 ± 0.2
II	117-139	0.8 ± 0.3	24 ± 11	14 ± 11	43 ±4	40 ± 4	2 ± 1	5 ± 1	2 ± 2
IV	175-182	1.8 ± 0.1	27 ± 6	21 ± 7	50 ± 1	50 ± 0.7	3.6 ± 0.2	11.1 ± 0.2	5 ± 2
V	193-214	27 ± 0.8	17 ± 2	5 ± 1	51 ± 1	49.2 ± 0.4	2.7 ± 0.5	13 ± 0.9	2.7 ± 0.7

Table 2. Ammonium half-saturation coefficients for AOB at the bulk liquid pH and at the pH inside a granule at T=20°C, together with the effect of the K_{S,NH4+} on the ammonium Monod term at different ammonium concentrations ($M_{NH4,i}$, where *i* is the value of ammonium concentration in mg N/L): $M_{NH4,i} = C_{S,i}/(C_{S,i}+K_{S,NH4+})$. Maximum specific growth rate (μ_{max}^{AOB}) was also calculated (at 20°C and for the corresponding pH, as in Jubany et al., 2008) to assess the overall impact in ammonium oxidation rate.

Microelectrode pH		<i>K</i> _{<i>S</i>,<i>NH</i>⁴+}	M _{NH4,10}	M _{NH4,20}	μ_{max}^{AOB}	
position		(mg N/L)	(dimensionless)	(dimensionless)	(1/d)	
Bulk	7.70	1.7	0.85	0.92	0.78	
Granule centre	7.26	4.6	0.68	0.81	0.68	

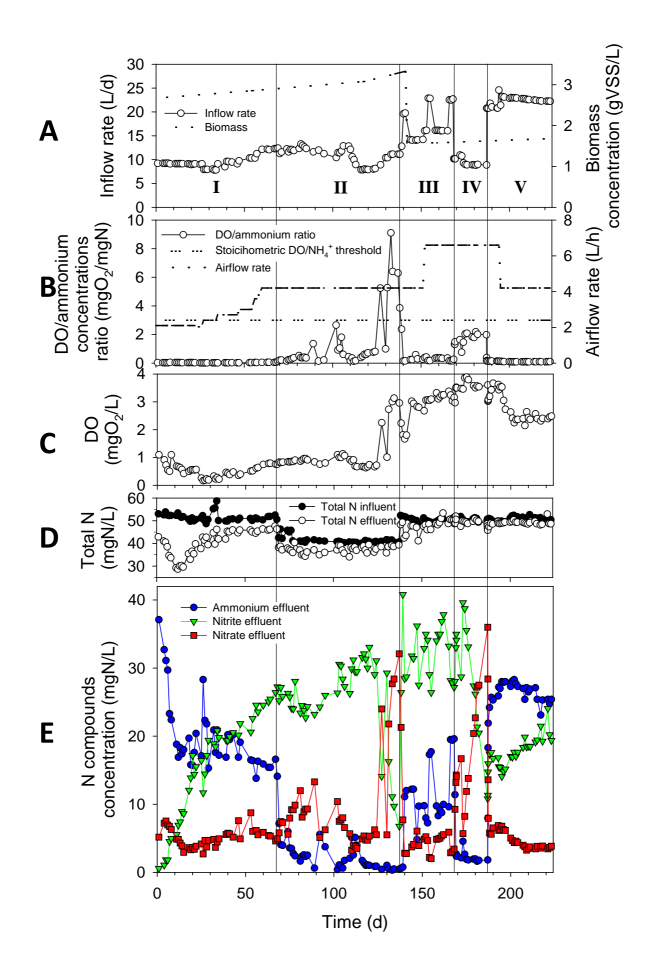


Figure 1. Reactor operation. **A**: Inflow rate and biomass concentration. **B**: DO/ammonium concentrations ratio and threshold value indicating when oxygen is the stoichiometrically limiting compound for AOB (see Eq. 2 for details). Airflow applied in the reactor (the superficial air velocity was in the range 3-9 m/h). **C**: Total nitrogen concentration in the inflow and total nitrogen in the effluent. **D**: Ammonium, nitrite and nitrate and DO concentrations in the reactor.

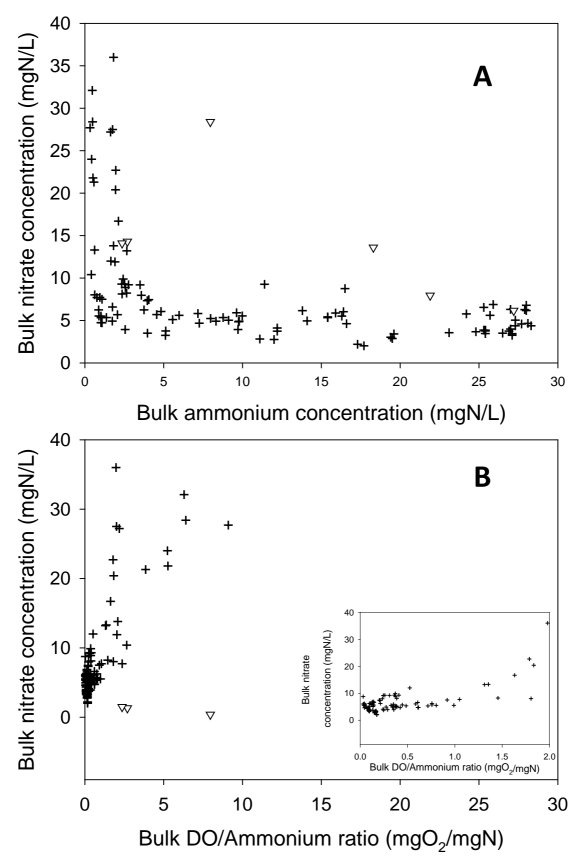


Figure 2. Effluent nitrate concentrations from day 50 onwards correlated with residual ammonium concentration (A) and bulk DO/ammonium concentrations ratio (B). The inset in graph B is a zoomed in version of the main graph, to show variations in the low range of the bulk DO/ammonium concentrations ratio (0-2 g O_2/g N). When the time between

measurements was less than 1 day, the corresponding data were highlighted using triangles, whereas the rest of data points (crosses) were obtained at a slower sampling frequency.

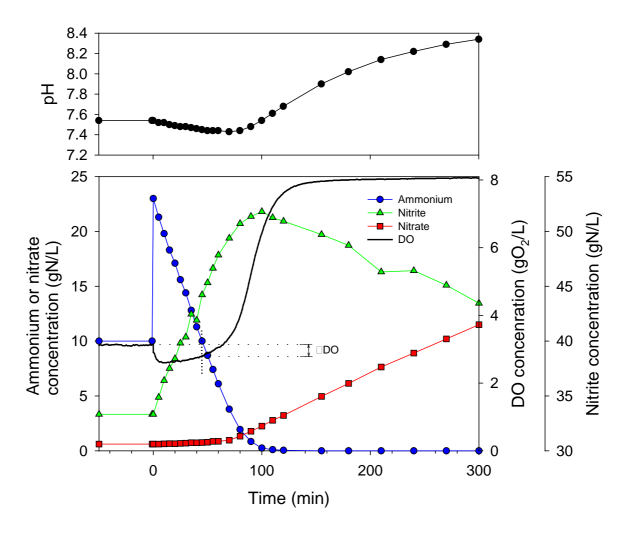


Figure 3. Batch-test performed at day 159 of the operation. An ammonium pulse was added after the inflow rate was stopped (time zero in the graph). The period before time zero indicated the continuous reactor operation before the batch-test. pH was measured but not controlled. The Δ DO highlighted in the figure corresponds to the improved oxygen consumption after the pulse of ammonium, once the bulk ammonium concentration is back to 10 mg N/L (time = 45min).

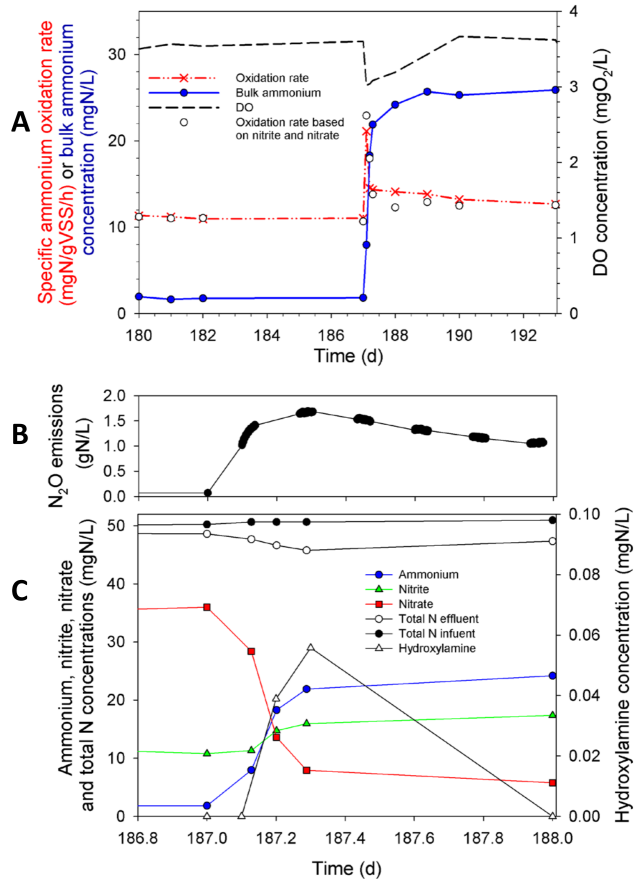


Figure 4. Step-up disturbance in residual ammonium by increasing the flowrate at day 187. **A**: Time course concentrations of ammonium, nitrite and nitrate and hydroxylamine

concentrations. Total nitrogen the influent and effluent has been also included. Specific ammonium oxidation rate has been computed based on the ammonium concentrations (red dashed line) and based on the sum of nitrite and nitrate produced (circles). **B**: N_2O emissions. **C**: Time course bulk ammonium and DO concentration together with the specific ammonium oxidation rate.

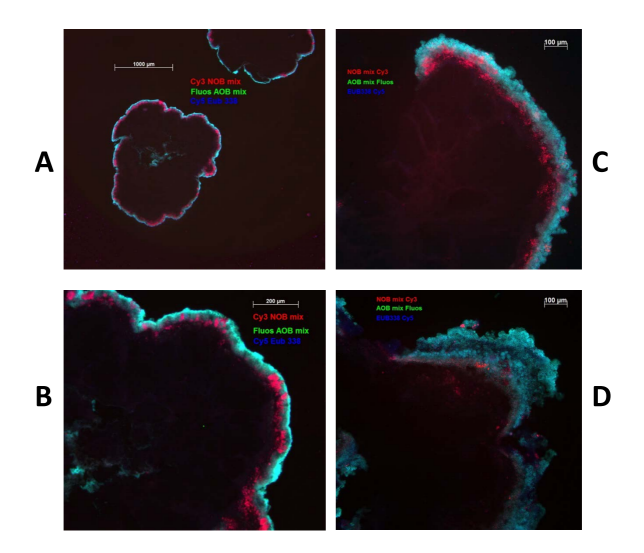


Figure 5. FISH-cryosectioning of granules. Cy3 (red) was used to detect NOB, Fluos (green) to detect AOB and Cy5 (dark blue) to detect most bacteria. Combinations of Cy3 and Cy5 visualised NOB as red/pink and the combination of Fluos and Cy5 visualised AOB as light blue. **A**: Granule slice during period of high residual ammonium concentrations (phase III, day 148)(4x). **B**: Granule slice during period of high residual ammonium concentrations (phase III, day 148, same slice than in Fig. 5A but at 10x magnification).**C**: Granule slice during period of low residual ammonium concentrations (phase V, day 187)(40x). **D**: Granule slice during period of high residual ammonium concentrations (phase V, day 223)(40x).

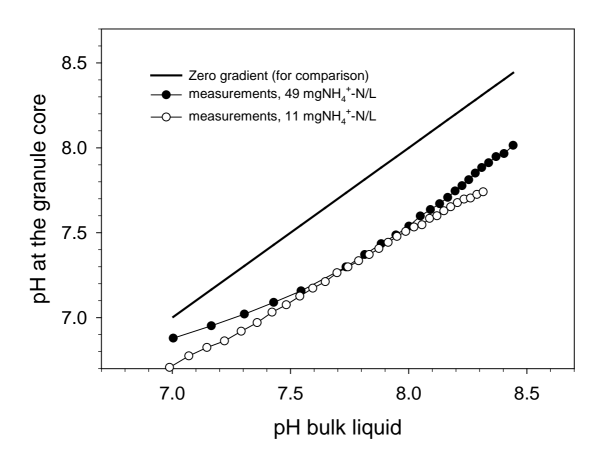


Figure 6. pH difference between the bulk liquid and inside a granule at a bulk pH range of 7.0-8.4. The position for equal pH between the bulk liquid and the granule center has been highlighted by a solid thick line (zero gradient). Points plotted below the grey line indicate a higher pH in the bulk liquid than inside the granule.