Batch enrichment of anammox bacteria and study of the underlying microbial community dynamics

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ABSTRACT

The anaerobic ammonium oxidation (anammox) consists on the biological conversion of ammonium (NH₄⁺) into dinitrogen gas under absence of oxygen. Nitrite (NO₂⁻) is a substrate of the anammox reaction, but also an inhibitor at high concentrations. This study investigates the effect of nitrite on the microbial community during the batch enrichment of anammox sludge. Six inoculums collected from different environments were enriched after a conditioning pretreatment and under controlled conditions during 4 months. Concerning the mineral medium used, two different nitrite supply strategies were applied; i.e., (i) initially low concentration at 25 mg NO₂⁻-N/L and progressive increase to 150 mg NO₂⁻-N/L, and (ii) constant high concentration at 150 mg NO₂⁻-N/L. All tested inoculums developed anammox activity but only when the enrichment was started at low nitrite concentration. In such case, the specific ammonium conversion rates finally obtained ranged from 21 ±1 to 118 ±1 mg NH₄⁺-N/g VS/d (VS, volatile solids). Abundance of the functional gene encoding
for the enzyme hydrazine oxidoreductase (hzo) was assessed using the real-time quantitative polymerase chain reaction (q-PCR) showing positive correlation with the anammox activity finally reported. In addition, high-throughput DNA sequencing helped to elucidate the underlying microbial community dynamics. The raw inoculum source, the conditioning pretreatment, and the cultivation conditions applied were jointly determinants of the final microbial community structure of the enrichments despite a clear convergence at the end of the experimental period. On the other hand, the cultivation conditions alone determined the selection of anammox species belonging to the genus Candidatus Brocadia.

KEYWORDS

Anaerobic ammonium oxidation; Biomass enrichment; Nitrite; Real-time quantitative polymerase chain reaction; Next-generation sequencing.

1. INTRODUCTION

The anaerobic ammonium oxidation (anammox) consists on the biological conversion of ammonium (NH$_4^+$) into dinitrogen gas (N$_2$) under absence of oxygen. This is a chemolithoautotrophic microbial process where nitrite (NO$_2^-$) acts as the electron acceptor. The anammox reaction also involves the production of a minor fraction of nitrate (NO$_3^-$). According to Strous et al. [1], the corresponding molar ratios for NH$_4^+$ consumption, NO$_2^-$ consumption, N$_2$ production, and NO$_3^-$ production are 1.00:1.32:1.02:0.26, respectively. Metagenomic studies also suggest that nitric oxide (NO) and hydrazine (N$_2$H$_4$) are intermediates in the anammox reaction [2]. The anammox process was discovered in the early 1990’s in a denitrifying fluidized bed reactor [3] and the interest in this bioprocess has ever since been rising in fundamental and applied research fields, such as marine ecology and environmental biotechnology [4,5].

So far, six “Candidatus” anammox bacterial genera have been enriched [6,7] from wastewater treatment facilities and freshwater environments (Brocadia, Kuenenia, Jettenia, Anammoxoglobus and
Anammoxomicrobium), as well as from marine environments (Scalindua). All these genera make a monophyletic branch within the phylum Planctomycetes. In physiological terms, they feature a specific cytoplasmatic membrane-bound organelle known as anammoxosome, which is the locus of the anammox catabolism. They are also characterized by a low growth rate, with doubling times of 2.1-11 days (at ~30°C) equivalent to a maximum specific growth rate of 0.065-0.334 d⁻¹ [1,8]. Because of this slow biomass development, and the specialized metabolism, anammox bacteria may be difficult to culture. Yet, phylotypes related to the anammox genera have increasingly been observed by molecular means in diverse environments, such as activated sludge from wastewater treatment plants (WWTP) [9,10], marine sediments [11,12], freshwater environments [13], and terrestrial ecosystems [14].

Anammox bacteria have not been isolated in pure culture yet, thus pointing to the fact that they may coexist with other microbial species, even in bioreactors fed exclusively with mineral substrates [8,15,16]. The most frequent enrichment strategies have been based on different types of continuously operated bioreactors [17]; e.g., sequencing batch reactor (SBR), rotating biological contactor, up-flow biofilm reactor, or membrane bioreactor [1, 18-20]. Alternatively, enrichments have also been developed in batch cultures [9, 21-23].

Many studies have shown that successful cultivation of anammox bacteria from conventional sludge takes long time; i.e., generally from 4 months to 1 year [24]. Such time will be influenced by factors like (i) the ecological characteristics of the seeding sludge including initial concentration and relative abundance of anammox bacteria [24], (ii) effective biomass retention inside the reactor [25], and (iii) the environmental conditions applied: temperature, pH, and concentration of NH₄⁺, NO₂⁻, dissolved oxygen (DO), organic carbon, sulphide and other inhibitors like metals and antibiotics [26-28].

Monitoring the anammox activity usually involves the chemical analysis of relevant nitrogen (N) compounds (i.e., NH₄⁺, NO₂⁻, and NO₃⁻) in the liquid phase. However, this strategy might be unsuccessful during the initial stages of the enrichment process, when the number of anammox cells is too low and their activity can still not be detected macroscopically. The use of culture-independent molecular methods has been proposed as a suitable method in such cases and various protocols for the DNA amplification by polymerase chain reaction (PCR) of target genes have been described in the literature, as summarized by Li et al. [12]. Anammox specific primers have been developed for amplifying ribosomal genes (16S rRNA) or functional
genes such as that encoding for the enzyme hydrazine oxidoreductase (hzO) that dehydrogenates hydrazine to N₂. These primers have been used to assess the abundance of anammox bacteria genes within environmental samples by real-time quantitative polymerase chain reaction (q-PCR), and to analyse the microbial community diversity by molecular typing and sequencing methods [9, 29-32]. Emerging next-generation sequencing (NGS) have also been applied for providing an in-depth characterization of the microbial biodiversity in anammox systems [33-36]. Yet, not so much information is available in the literature concerning the microbial community structure and dynamics during enrichment of the anammox biomass as determined by quantitative and qualitative culture-independent molecular methods.

Autotrophic nitrogen removal (ANR) applications based on anammox are promising for N-removal from municipal side-/mainstreams, industrial and agricultural wastewaters [37-40], particularly after anaerobic digestion once biodegradable organic carbon is depleted. However, anammox enriched sludge is not always available, and biomass enrichment can become the critical point for the start-up of the process. An appropriate selection of the environmental conditions applied is decisive for a successful enrichment. In this regard, NO₂⁻ is a substrate of the anammox reaction but may also become an inhibitor at high concentrations. Such inhibition has been reported as highly case-specific; i.e., concentrations as low as 5 and 30 mg NO₂⁻-N/L were found as inhibitory in some studies [41,42] whereas much higher inhibitory boundaries of 210-274 mg NO₂⁻-N/L were determined in other cases [43-45]. Concerning this variability, Kimura et al. [44] suggested that differences in NO₂⁻ concentration tolerance may be caused by the cultivation conditions used.

The aim of this study is to investigate the presence of anammox populations in different inoculum sources and to assess the feasibility of enrichment in batch under two different strategies concerning NO₂⁻ supply. Thus, final concentrations of 150 mg NO₂⁻-N/L were targeted in the mineral medium used as feeding solution but testing two different supply strategies (i.e., initially low vs. high concentration) in order to evaluate the effect of the NO₂⁻ concentration when starting anammox batch enrichments. Use of molecular techniques will help to detect anammox bacteria and to establish correlations between macroscopically observed process parameters and the underlying microbial community dynamics. Microbial monitoring will be conducted using q-PCR and 16S rRNA gene targeted NGS.
2. MATERIALS AND METHODS

2.1. Inoculum sources

Six different biomass sources collected in conventional N-removal facilities were considered as inoculum (I) for batch enrichment; i.e., (I1) activated sludge collected in a municipal WWTP that combine the use of a Modified Ludzack-Ettinger (MLE) bioreactor unit and a membrane filtration loop (Betton, Brittany, France), (I2) mixture of activated and settled sludge collected in a pig slurry treatment plant with intermittent aeration and gravity settling (Meslin, Brittany, France), (I3) activated sludge collected in a pig slurry treatment plant with MLE configuration (Calldetenes, Catalonia, Spain), (I4) settled sludge -sediments- collected in a receiving lagoon treating municipal wastewater (Amanlis, Brittany, France), (I5) settled sludge -sediments- collected in polishing lagoons treating municipal wastewater (Amanlis, Brittany, France), and (I6) settled sludge collected in an intermittently aerated lagoon treating pig slurry (Almacelles, Catalonia, Spain). The volatile solids (VS) content of the samples was 0.44%, 1.82%, 0.62%, 0.82%, 0.48%, and 1.92% of the wet weight, respectively; whereas, the corresponding VS/TS ratio (TS, total solids) was 0.52, 0.64, 0.54, 0.09, 0.09, and 0.59, respectively. In order to favour biodegradation of available organic carbon before incubation for anammox biomass enrichment, a conditioning pretreatment based on promoting denitrification was carried out at room temperature during the first days after sampling by adding a NO$_3^-$ source such as KNO$_3$ in pulses of 722 mg/L (100 mg N/L) and controlling the pH within the range 7.0-8.0 (HCl 2M). Batch enrichment was started once denitrification declined (after 2-4 weeks).

2.2. Mineral medium

The synthetic nutritive solution was prepared using tap water according to a modification of the mineral medium described by Magrí et al. [45]; i.e., NH$_4$Cl (variable: 95-573 mg/L), NaNO$_2$ (variable: 123-739 mg/L), KNO$_3$ (361 mg/L), KHCO$_3$ (1000 mg/L), FeSO$_4$·7H$_2$O (9 mg/L), EDTA (5 mg/L), MgSO$_4$·7H$_2$O (240 mg/L), CaCl$_2$·2H$_2$O (143 mg/L), and trace element solution 0.3 mL/L. The trace element solution contained ZnSO$_4$·7H$_2$O (1247 mg/L), MnSO$_4$·H$_2$O (1119 mg/L), CuSO$_4$·5H$_2$O (44 mg/L), Al$_2$(SO$_4$)$_3$·14H$_2$O (201.5 mg/L), Na$_2$MoO$_4$·2H$_2$O (129 mg/L), CoCl$_2$·6H$_2$O (30 mg/L), KCl (100 mg/L), and EDTA (975
mg/L). Once dissolved the mineral salts, the DO was purged by bubbling with N₂ (< 0.2 mg/L) and the pH was adjusted to 7.0 (HCl 2M). The chemical KNO₃ was added aiming to strengthen the anoxic conditions and to prevent potential sulphate reduction to sulphide (which could inhibit the anammox reaction) in case of total NO₂⁻ consumption during the enrichment. Unfortunately, any phosphorus source was added to the nutritive solution throughout the enrichment due to a mistake, discovered later, in the labelling of the corresponding chemical (K₂O (27 mg/L) was supplied instead of KH₂PO₄). Thus, it seems that the phosphorus released by the decaying biomass was enough to avoid limitation in the availability of this nutrient.

2.3. Enrichment procedure

Twelve glass bottles (total volume: 575 mL; working volume: 500 mL) containing inoculum and mineral medium were flushed with N₂, sealed with a rubber stopper plus an aluminium cap, and placed in an incubator shaker (KS4000i control, IKA, Germany) at 150 rpm, 35ºC, and in dark conditions. Initial VS content within the bottles was adjusted to 3 g/L. Biomass settling was allowed once per week (for 1h). Each bottle was then opened and the supernatant was manually withdrawn to avoid accumulation of inhibitory compounds while keeping the settled biomass inside of the bottles. Subsequently, the bottles were refilled with new nutritive mineral medium, closed, and flushed with N₂. Two different strategies concerning NO₂⁻ supply were applied (i.e., initially low vs. high concentration). Thus, mineral medium was prepared with low amount (25 mg NO₂⁻-N/L) or high amount (150 mg NO₂⁻-N/L) of NO₂⁻ whereas NH₄⁺ was added at a constant rate of 25 mg NH₄⁺-N/L. In those bottles running at low NO₂⁻ concentration, once anammox activity was detected, a second weekly addition of NO₂⁻ and NH₄⁺ (NO₂⁻-N/NH₄⁺-N = 1.2) was carried out. Following this procedure, NO₂⁻ content in the mineral medium was progressively increased from 25 to 150 mg N/L at increments of 12.5 mg N/L. For the six bottles running at high NO₂⁻ concentration, the NO₂⁻ content was kept constant at 150 mg N/L with only one feeding event per week throughout the experimental period.

Concerning NH₄⁺ content, it was proportionally increased at a ratio of 1.2 g NO₂⁻-N per gram of NH₄⁺-N for those first six bottles initially running at low NO₂⁻ concentration but it was kept constant at 25 mg NH₄⁺-N/L for the others. The pH within the bottles was controlled in the range from 7.0 to 8.0 using HCl 2M. N₂ flushing was used to displace air in the bottles headspace every time they were opened. The liquid volume
exchanged when renewing the mineral medium was variable depending on the settling capability of the biomass, but usually between 60-80%. This is a higher value than those generally considered in anammox SBRs [46-48]. Enrichment lasted 4 months. Liquid samples were taken before and after each new feeding event and filtered using 0.45 µm polypropylene membrane filters prior to storage in the refrigerator. Biological samples were taken once per month, centrifuged at 10,000g for 4 min and supernatants were discarded. Pellets were stored at -20°C.

2.4. Final anammox activity test

Final anammox activity was assessed using batch tests at the end of the enrichment period. According to the aforementioned conditions, after renewing the mineral medium, liquid samples were collected using syringes at regular time intervals for N-compounds analysis. VS contents were also measured. The batch experiments were done in duplicate. Linear regression analyses were used to describe N-conversion.

2.5. Chemical analyses

NH$_4^+$, NO$_2^-$ and NO$_3^-$ were measured by ion chromatography (850 Professional IC, Metrohm, Switzerland). TS were measured after sample drying to constant weight at 105°C and VS were measured after further ignition in a muffle furnace at 550°C. The pH and DO were measured using portable meters pH 197i and Oxi 197 (WTW, Germany), respectively.

2.6. Molecular analyses

2.6.1. DNA extraction

Total DNA was extracted from approximately 0.25 g of pellet with the PowerSoil™ DNA Isolation Kit (MoBio Laboratories Inc., USA), according to the instructions of the manufacturer. The concentration and purity of the extracted DNA were checked spectrophotometrically (ND-1000, NanoDrop Technologies, USA) and in TBE 1X - 0.7% agarose gel. The extracted DNA was stored at -20°C until further analysis.
2.6.2. Real-time q-PCR

All real-time PCR amplifications were performed using the iQ™ SYBR® Green supermix (2x) (Bio-Rad Laboratories, USA) and the CFX96 real-time system equipped with the CFX Manager™ software v3.1 (Bio-Rad Laboratories), according to the instructions of the manufacturer. Quantification of total bacteria used universal eubacterial forward 1055F (5’-ATGGCTGTCGTCCAGCT-3’) and reverse 1392R (5’-ACGGGCGGTGTGTAC-3’) primers to amplify the hypervariable V3-V5 region from the 16S rRNA gene, as previously reported by Ferris et al. [49]. Concerning anammox bacteria, the gene encoding specific enzyme HZO (hzo gene) was used as functional biomarker. The primer set used to selectively amplify anammox bacterial fragment was hzoel1F1 (5’-TGYAAGACYTGYCAYTGG-3’) and hzoel1R2 (5’-ACTCCAGATRTGCTGACC-3’) as described by Schmid et al. [50]. The PCR reactions were run in a 25 µL volume containing 12.5 µL of iQ™ SYBR® Green supermix (2x), 1.5 or 0.625 µL of each primer (eubacteria-16S rRNA or anammox-hzo, respectively; 10 µM), 2 µL of diluted DNA template, and 7.5 or 9.25 µL of sterile water (eubacteria or anammox, respectively). To check against potential inhibition of the PCR amplification, reactions were done on DNA templates diluted 10- and 100-folds and each reaction was carried out in triplicates. The amplification program applied for eubacteria was 95ºC for 10 min, followed by 45 cycles of denaturation at 95ºC for 30 s, annealing at 60ºC for 50 s, and extension at 72ºC for 30 s. In case of anammox bacteria, the program applied was 95ºC for 2 min, followed by 40 cycles of denaturation at 95ºC for 30 s, annealing at 51ºC for 60 s, and extension at 72ºC for 60 s. The fluorescence intensity of the amplified DNA was measured after each extension step and a melting curve analysis was performed after completion of PCR. The microbial quantification was based on a mean slope value derived from standard curves obtained by q-PCR amplification of 10-fold successive dilutions of DNA fragments of the targeted genes [51], and which were purified using the Wizard® Plus SV Minipreps DNA Purification System (Promega, USA) according to the instructions of the manufacturer. The absolute copy number of DNA fragments per µL of standard solution was calculated from the concentration measured spectrophotometrically (ND-1000) and the molecular mass of the DNA fragment. The standard curves covered a range from 10² to 10¹⁰ gene copies per µL of reaction. These series of standard dilutions were
amplified along with the unknown samples during each q-PCR run, which allowed building the standard curve ($R^2 > 0.98$) corresponding to each run. Results were expressed as gene copy number per mL of mixed liquor within the enrichment bottle.

2.6.3. High-throughput DNA sequencing

16S rRNA genes high-throughput DNA sequencing was performed at the NGS facility of the BIOMIC Team of IRSTEA (Antony, France) using Ion Torrent™ (Life Technologies, USA) technology and methods, according to the procedures described by Poirier et al. [52]. Briefly, the bacterial and archaeal hypervariable region V4-V5 of the 16S rRNA genes was amplified using the “universal” fusion-primer 515F (5’-GTGYCAGCMGGCCGCGTAA-3’) and 928R (5’-CCCGGYCAATTCTTTRAGTTT-3’) [53] modified to allow the tagging and sequencing of the amplified products. Amplification was performed in a 50 µL reaction mixture containing 5 µL of Pfx buffer (10X), 1.5 µL of dNTP mix (10 mM each), 1 µL of MgSO$_4$ (50 mM), 1.5 µL of each primer (10 µM), 0.4 µL of Platinum® Pfx DNA polymerase, 38.1 µL of water and 1 µL of extracted DNA (10 to 200 pg) (Pfx SuperMix protocol from Life Technologies). The mixture was held at 94°C for 5 min, followed by 30 cycles at 94°C for 15 s, 50°C for 30 s and 68°C for 1 min, and a final extension step at 68°C for 5 min. PCR products were cleaned using the Agencourt® AMPure® XP magnetic beads purification system (Beckman Coulter, USA) and quantified with a capillary electrophoresis bioanalyzer (2100 Electrophoresis Bioanalyzer, Agilent Technologies, USA). Purified libraries were diluted (in a first step at 500 pM and later at 100 pM). Equal volumes of amplicons were combined in equimolar concentrations (100 pM) for sequencing using the Ion OneTouch™ 2 Instrument with the Ion PGM™ Template OT2 400 Kit and using a Ion Personal Genome Machine (PGM™) System with the Ion 316™ Chip Kit v2 and the Ion PGM™ Sequencing 400 Kit, according to the instructions of the manufacturer. Low quality and polyclonal sequence reads were filtered out by the PGM™ System software, and resulting data was exported as a FastQ file. 16S rRNA genes high-throughput DNA sequencing was performed directly on total DNA extracts to analyze global microbial community structure, and when anammox bacteria were not detected on PCR products obtained by specific amplification of the Planctomycetes phylum using the primers Pla46F and 1392R, as described in Bae et al. [9]. Sequences were analyzed using the Quantitative Insights into Microbial Ecology (QIIME v1.8.0) pipeline [54]. Sequences shorter than 200 bp, containing
chimeras, and found as singletons were removed. Operational taxonomic units (OTUs) were subsequently defined using UPARSE implemented in USEARCH (v8.0.1623) [55] at a 97% similarity level. MOTHUR (v1.25.0) [56] and SILVA (v119) [57] were used as the classifier tool and database for taxonomic association (with a minimum similarity threshold of 80%), respectively.

2.6.4. Statistical analyses

Statistical analysis to evaluate microbial community structure evolution was carried out through the non-metric multidimensional scaling (NMDS) method using the open-source software R (v3.2.3) [58] including functions from the vegan package (v2.3-2) [59]. Shannon-Weaver, Simpson, and Inverse Simpson diversity indices, as well as species richness and Pielou’s evenness were calculated according to the procedures described in Oksanen et al. [59].

3. RESULTS AND DISCUSSION

3.1. Performance of the anammox enrichment

All tested inoculums developed positive anammox activity throughout the 4-month experimental period when the enrichment was started at low NO$_2^-$ concentration (Fig. 1A). However, biomass still maintained overall appearance of activated sludge and brownish colour at the end of the enrichment. Three main enrichment phases were observed: (P1) endogenous heterotrophic denitrification was the dominant process and NH$_4^+$ may even slightly increase during incubation due to the hydrolysis of the remaining organic matter, (P2) occurrence of NH$_4^+$ consumption and subsequent speed up at increasing NO$_2^-$ concentration, and (P3) consolidation of NH$_4^+$ consumption at high NO$_2^-$ concentration with evidence of NO$_3^-$ production (while the NO$_2^-$-N/NH$_4^+$-N reaction ratio became closer to the expected value of 1.32 [1]). Time for detecting NH$_4^+$ consumption under anaerobic conditions (beginning of P2) was variable (Table 1), ranging from 0 days in bottle seeded with I3 to 92 days in bottle seeded with I2 (time for anammox activity appearance was longer in those bottles seeded with the inoculums containing higher VS). According to this fact, and the progressive
increase in the NO$_2^-$ supplied, concentrations of 150 mg NO$_2^-$-N/L at the beginning of a new batch were applied at the end of the experimental period in all cases except in bottle with I2, where the maximum concentration used was 50 mg NO$_2^-$-N/L. On the other hand, no anammox activity was developed by any of the inoculums when enrichment was started at high NO$_2^-$ concentration (Fig. 1B), which evidences the importance of the feeding strategy adopted when targeting anammox bacteria enrichment and although NO$_2^-$ levels of 150 mg N/L were not previously evaluated as inhibitory elsewhere [43-45].

Results of the final activity test did not evidence link between time of anammox activity appearance and measured NH$_4^+$ conversion rate (Table 1). Good linearity in the evolution of the N-forms was observed during the activity test, which was indicative of no substrate inhibition ($R^2 > 0.98$). Thus, specific activity was assessed within the range from 21 to 118 mg NH$_4^+$-N/g VS/d which is equivalent to total N-conversion rates of 59-297 mg N/g VS/d. These values are within the range of specific N-removal activities of 60-1600 mg N/g VSS/d (n=14; VSS, volatile suspended solids) obtained for a variety of anammox sludges of diverse origins as reported by Van Hulle et al. [39].

The procedure followed in this study for the enrichment of anammox biomass consisted on the use of bottles as bioreactors (where substrates were supplied in pulses to the biomass). Eventual singularities of this method with respect to the use of the SBR technology are: long cycle (7 d), long hydraulic residence time (> 7 d), high volume exchange ratio (60-80%), and exposure of the biomass to a wide range of substrate concentrations (0-150 mg NO$_2^-$-N/L). These factors could influence on the evolution of the microbial community since they may have implications on the biomass retention within the reactor, the existence of famine periods during the enrichment, and the tolerance of high substrate concentrations by the biomass.

3.2. Monitoring the enrichment of anammox bacteria by real-time q-PCR

The evolution of total and anammox bacteria was monitored by real-time q-PCR throughout the 4 months that lasted the experimental period. For a given batch culture, the bacterial 16S rRNA gene was the most abundant and the least fluctuant (Fig. 2). Average values in the enrichments ranged from $2.5 \pm 0.2 \cdot 10^6$ to $1.0 \pm 0.2 \cdot 10^9$ copies/mL depending on the inoculum source and the NO$_2^-$ supply strategy applied. Concerning the
anammox hzo gene, it was quantified from the beginning of the experiment in all inoculums (Table 1), and a significant increase throughout the enrichment was evidenced in those cases where NO$_2^-$ was initially supplied at low concentration (Fig. 2). Conversely, that was not the case when NO$_2^-$ was initially supplied at high concentration. Thus, at day 0, the hzo gene copy number in the enrichments started at low NO$_2^-$ concentration ranged from 3.3 ±0.3·10$^4$ to 1.2 ±0.3·10$^5$ copies/mL, and after 4 months reached values from 1.1 ±0.1·10$^6$ to 1.5 ±0.1·10$^7$ copies/mL. At that time, it was also evidenced a positive correlation between the hzo gene copy number and the anammox activity (Fig. 3), similarly as previously reported by Tsushima et al. [29] using primers targeting the 16S rRNA gene of the anammox bacteria, and although there was no evident link between the hzo gene copy number in the inoculum at day 0 and the time of anammox activity appearance (Table 1). Lack of linkage between the initial hzo gene copy number and the time of anammox activity appearance suggests that other factors besides the initial anammox biomass concentration (measured as hzo gene copy number) determined the time when such activity became evident macroscopically. Among these factors, we might include competition for substrate, coexistence of the anammox bacteria with other microbial populations, and change of the growing anammox species (see section 3.3). In this regard, mineralization of organic-N forms and denitrification could have masked an earlier detection of anammox activity. Furthermore, Tao et al. [24] pointed out dependence of the lag phase length not only on the initial biomass concentration but also on other ecological characteristics of the inoculum used and concluded that an evenly distributed community benefits the start-up of the anammox process with shorter times and higher activities. Yet, differences exist between these findings and our study; i.e., we dealt with inoculums with much lower initial relative abundances of anammox bacteria and also we observed a change of the growing anammox species during the enrichment process (see section 3.3). On the other hand, final hzo gene copy numbers were below 2.0 ±0.2·10$^5$ copies/mL in all cultures started at high NO$_2^-$ concentration.

3.3. Microbial community structure and dynamics during the enrichment

The 16S rRNA high-throughput DNA sequencing was performed for all inoculums (I1-I6) at initial time (t0 = 0 months) and final time (t4 = 4 months) when the enrichment was started at low NO$_2^-$ (LN) and high NO$_2^-$ (HN) concentrations, and for I3 at all times (t0-t4) and NO$_2^-$ supply conditions (LN, HN). Reads obtained yielded between 1933 and 22475 high-quality sequences per sample that made up to 633 OTUs. A
systematic random depletion was applied on the different datasets to equalize the number of sequences per sample; i.e., dataset for all inoculums including samples at initial and final time (18 samples) was adjusted to 11058 sequences per sample whereas dataset for I3 including samples at all times (9 samples) was adjusted to 1933 sequences per sample. The representativeness and identification of these OTUs was analysed to better understand the process of anammox enrichment.

3.3.1. Evolution of the microbial communities

The diversity indices reported in Table 2; i.e., Shannon-Weaver, Simpson, and Inverse Simpson, globally show a systematic decrease of the microbial diversity throughout the enrichment (average index reductions for the cultures started at low NO$_2^-$ concentration of 25.7 ±4.5%, 8.8 ±3.4%, and 67.9 ±14.3%, respectively). In addition, both the species richness and Pielou’s evenness decreased in this period (average reductions for the cultures started at low NO$_2^-$ concentration of 30.2 ±7.6% and 20.5 ±4.7%, respectively). These data may imply concomitant disappearance of some species and larger segregation between the low and highly represented taxons [60-61]. The impact seems to be more important when the enrichment was started at low NO$_2^-$ concentration (Table 2). No relation was found between the initial microbial diversity and the effectiveness of the anammox enrichment in terms of both time required and final activity achieved.

Statistical analysis of the microbial community structure evolution was also performed through the NMDS method. For a given inoculum, the evolution of the microbial community was different depending on the NO$_2^-$ supply strategy applied during the enrichment. Relative differences in the composition of both microbiomes (enrichments started at low vs. high NO$_2^-$ concentration) over time were evidenced as it is shown for the bottle seeded with I3 in Fig. 4. Thus, although both microbiomes evolved similarly with a diminution in the diversity throughout the enrichment, divergence in the microbial community structure progressively increased and was maximal after 4 months of enrichment. Such divergence included the enrichment of anammox bacteria only under conditions of initially low NO$_2^-$ concentration. In addition, the NMDS analysis also revealed that, despite the significant relative differences in the composition of the microbial community concerning the inoculums used in this study, the enrichment conditions applied forced the convergence of such communities (Fig. 5). The most similar microbial community structures were
observed for those inoculums coming from systems treating the same kind of wastewater; i.e., municipal (I1, I4, and I5) vs. pig slurry (I2, I3, and I6), at both initial and final time. The clear trend to converge independently of the inoculum source and the feeding strategy applied indicates the high selective pressure exerted by the experimental conditions here implemented.

3.3.2. Description of the microbial community structures

As aforementioned, the microbial community structure in the batch cultures evolved significantly throughout the enrichment. However, main phyla detected in inoculums such as *Proteobacteria, Bacteroidetes, Firmicutes, Chloroflexi, Chlorobi, Acidobacteria,* and *Planctomycetes* were also present at the end of the enrichment period (Fig. 6A-6B; Table 3 and 4). The phylum *Proteobacteria,* which is commonly found in wastewater treatment bioreactors, anaerobic digesters, and soils [62-64] was one of the most predominant at both initial time (relative abundances of 10.3-46.7%) and final time (relative abundances of 23.9-55.5%). For those enrichments started at low NO$_3^-$ concentration, the evolution of this phylum in terms of abundance was dependent on the inoculum source. Thus, the proportion of sequences belonging to this phylum followed an increase of 93.1 ±60.4% (in average) for those inoculums coming from pig slurry treatment plants (I2, I3 and I6) whereas followed a decrease of 19.6 ±16.0% (in average) for those inoculums coming from municipal WWTPs (I1, I4, and I5). On the other hand, for those enrichments started at high NO$_3^-$ concentration, the abundance of sequences belonging to this phylum increased at an average rate of 55.9 ±37.9% regardless the inoculum source. For I3 (Fig. 6B), the increased proportion of *Proteobacteria* is essentially attributable to the evolution of genera belonging to the class *Betaproteobacteria* and family *Rhodocyclaceae* (OTUs 2 and 11) that contain several denitrifying species; e.g. *Azoarcus. Bacteroidetes* was another of the most represented phyla at initial time with proportions ranging from 6.1% to 30.9% in I2 and I6, respectively. However, despite this initial high abundance, a systematic reduction was observed regardless the inoculum source and feeding strategy applied, with final relative abundances lower than 2.7%. This is not surprising since the phylum *Bacteroidetes* contains both aerobic and anaerobic bacteria such as *Sphingobacterales* and *Bacteroides,* respectively. In the case of I3 (Fig. 6B), the reduced proportion of *Bacteroidetes* is clearly linked to the disappearance of OTU 4 identified as a *Sphingobacterales Saprospiraceae.* Conditions applied during the anoxic enrichment including the progressive abatement of residual organic compounds may be the
cause of the depletion of this widespread phylum [62-64]. Concerning the phylum *Firmicutes*, relative abundance of sequences belonging to this phylum (0.7-18.6%) was higher for those inoculums coming from pig slurry treatment facilities than for those others coming from municipal WWTPs; i.e., $12.7 \pm 6.2\%$ vs. $1.0 \pm 0.2\%$ in average, respectively. The corresponding quantitative evolution was slightly different depending on the NO$_3^-$ supply strategy. The coexistence of bacteria belonging to the phylum *Chloroflexi* is usually reported in anammox reactors being suggested that they can use decaying anammox bacterial cell materials [65].

Here, the evolution of this phylum (initial relative abundances of 2.1-12.9%) was highly variable ranging from 67.7\% decrease to 302.4\% increase for I2 enriched at initially low NO$_3^-$ concentration and I6 enriched at high NO$_3^-$ concentration, respectively. Sequences corresponding to the phylum *Chlorobi* (initial relative abundances of 0.1-9.3\%) represented from 1.8\% to 14.1\% of total sequences at final time for I6 and I1, respectively. Concerning the phylum *Acidobacteria*, final relative abundance was lower than 4.0\% in all cases. *Planctomycetes*, which contains all known anammox genera, did not become dominant in any of the enrichments that developed anammox activity (final relative abundances ranged from 0.6\% to 7.7\% for I4 and I1, respectively), which is in accordance with other microbial characterizations performed in anammox dedicated reactors [34-35]. Within this group, the final relative abundance of sequences belonging to anammox species was between 31.7\% and 75.3\% for I2 and I1, respectively. For I3 (Fig. 6B), a significant enrichment of OTU 1 belonging to phylum *Armatimonadetes* (formally called the candidate phylum OP10) was observed (similarly for other inoculums). Species belonging to the phylum *Armatimonadetes* have been detected in different natural environments such as the Obsidian Pool in Yellowstone National Park and freshwater lakes and rivers [66]. However, the reason of their enrichment in this study is not clear. OTU 36 was identified as Candidatus_Brocadia. In addition, 6 OTUs listed in Table 4 (from a total of 15) were identified as identical to sequences retrieved from anammox systems and 2 other OTUs were related to sequences retrieved from N-removal systems. This fact suggests that the conditions applied to perform the enrichment also favoured selection of other specific microbial groups that could either behave as partners or competitors of the anammox bacteria.

### 3.3.3. Anammox genera

Concerning the presence of anammox genera in the inoculums at initial time, Ca. *Brocadia*, Ca. *Kuenenia*,
Ca. *Anammoximicrobium* and Ca. *Jettenia* were all detected (Fig. 7), although with an important disparity in terms of distribution and relative abundance. Thus, for instance, these four genera were detected together in I3 but not in the other inoculums. Otherwise, any of these genera were detected in I1. Ca. *Jettenia* was only detected when DNA samples were first amplified by means of selective PCR using *Planctomycetes* primers, which allowed lowering the detection threshold but preclude their relative quantification (Fig. 7). Before enrichment, higher relative abundances were measured for Ca. *Anammoximicrobium* in I4 and I5 (lagoon WWTP), with 0.27% and 0.37%, respectively, of the total sequences. In fact, this anammox genus (which was recently described) may grow optimally at temperatures around 20°C [7]. On the other hand, the genus Ca. *Brocadia* (maximum initial relative abundance of 0.011% for I3) became dominant at the end of the enrichment period regardless of the inoculum considered, with relative abundances ranging from 1.0 to 6.1% of the total sequences. Thus, similarly to Costa et al. [34], the obtained results indicated that while the type of inoculum and the culture conditions are both key determinants of the global microbial composition of the enriched biomass, the operational conditions alone determined the selection of the anammox species. The niche differentiation between Ca. *Brocadia* and other anammox genera often dominant in enrichments performed at lab-scale such as Ca. *Kuenenia* is still not clear. According to several studies [36, 67-69], the genus Ca. *Brocadia* would presumably be an r-strategist (i.e., relatively high growth rate and low substrate affinity), while the genus Ca. *Kuenenia* could be a K-strategist (i.e., relatively low growth rate and high substrate affinity). In this study, the feeding of the enrichments was based on the intermittent supply of substrates which would favour the proliferation of an r-strategist population.

Overall, our findings showed that anammox bacteria were ubiquitous in all the inoculums collected from different waste/water treatment environments which made feasible their eventual enrichment in batch under controlled conditions. However, such enrichment was only achieved when the process was started at low NO₂⁻ concentration (≤ 25 mg NO₂⁻-N/L) evidencing the importance of carefully controlling the conditions applied (i.e., substrate concentration). The procedure used prompted the enrichment of the anammox genus Ca. *Brocadia* regardless of the inoculum source. A higher initial anammox biomass concentration (measured as *hzo* gene copy number) did not necessarily imply a faster process start-up. Thus, other physicochemical and ecological characteristics of the inoculum affected the evolution of the enrichment since they determined issues such as the competition for substrate, coexisting microbial groups, dominant anammox species, and
associated relative abundances at the beginning of the process.

4. CONCLUSIONS

The effect of NO$_2^-$ supply during the batch enrichment (4 months) of anammox sludge was investigated using six different biomass sources. Concerning the mineral medium used as feeding solution, two different NO$_2^-$ supply strategies were applied; i.e., (i) initially low concentration at 25 mg NO$_2^-$-N/L and progressive increase to 150 mg NO$_2^-$-N/L, and (ii) constant high concentration at 150 mg NO$_2^-$-N/L.

- All tested inoculums developed anammox activity only when the enrichment was started at low NO$_2^-$ concentration. In such case, the final specific NH$_4^+$ conversion rate was measured within the range from 21 to 118 mg NH$_4^+$-N/g VS/d.
- Abundance of the hzo functional gene showed positive correlation with the anammox activity finally reported.
- The biomass source, a conditioning pretreatment, and the cultivation conditions applied were determinant factors of the final microbial composition of the enrichments despite a clear convergence at the end of the experimental period. However, the cultivation conditions alone determined the selection of anammox species belonging to the genus Ca. Brocadia.

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Table 1. Results of the anammox activity test at the end of the 4-month enrichment period when process was started at low nitrite concentration and comparison with the day of activity appearance and initial $hzo$ gene copy number.

<table>
<thead>
<tr>
<th>Inoculum (source)</th>
<th>Activity appearance (days)</th>
<th>ACR $^a$ (mg NH$_4^+$-N/L/d)</th>
<th>sACR (mg NH$_4^+$-N/g VS/d)</th>
<th>TNCR (mg N/L/d)</th>
<th>NO$_2^-$-N/NO$_3^-$-N</th>
<th>Initial $hzo$ gene copy number (copies/mL) $^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>I1 (WWTP)</td>
<td>43</td>
<td>222 ±2</td>
<td>118 ±1</td>
<td>560 ±11</td>
<td>1.53 ±0.03</td>
<td>6.3 ±1.4 · 10$^7$</td>
</tr>
<tr>
<td>I2 (PSTP)</td>
<td>92</td>
<td>60 ±9</td>
<td>42 ±6</td>
<td>160 ±25</td>
<td>1.67 ±0.01</td>
<td>1.2 ±0.3 · 10$^4$</td>
</tr>
<tr>
<td>I3 (PSTP)</td>
<td>0</td>
<td>164 ±6</td>
<td>62 ±2</td>
<td>373 ±17</td>
<td>1.28 ± 0.02</td>
<td>5.2 ±0.8 · 10$^3$</td>
</tr>
<tr>
<td>I4 (LWWTP)</td>
<td>57</td>
<td>78 ±1</td>
<td>26 ±0</td>
<td>208 ±8</td>
<td>1.64 ±0.08</td>
<td>3.3 ±0.3 · 10$^6$</td>
</tr>
<tr>
<td>I5 (LWWTP)</td>
<td>43</td>
<td>84 ±4</td>
<td>21 ±1</td>
<td>238 ±14</td>
<td>1.83 ±0.02</td>
<td>5.4 ±0.9 · 10$^5$</td>
</tr>
<tr>
<td>I6 (LPSTP)</td>
<td>77</td>
<td>162 ±12</td>
<td>70 ±5</td>
<td>370 ±37</td>
<td>1.28 ±0.05</td>
<td>1.1 ±0.2 · 10$^3$</td>
</tr>
</tbody>
</table>


$^b$ Quantification of those samples labelled with * was carried out at the detection limit.
Table 2. Diversity indices for the six tested inoculums (I1-I6) at initial time (t0 = 0 months) and final time (t4 = 4 months) when the enrichment was started at low nitrite (LN) and high nitrite (HN) concentrations.

<table>
<thead>
<tr>
<th>Indices</th>
<th>I1</th>
<th>I2</th>
<th>I3</th>
<th>I4</th>
<th>I5</th>
<th>I6</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>t0</td>
<td>t4-LN</td>
<td>t4-HN</td>
<td>t0</td>
<td>t4-LN</td>
<td>t4-HN</td>
</tr>
<tr>
<td>Shannon-Weaver</td>
<td>4.02</td>
<td>2.99</td>
<td>3.15</td>
<td>4.22</td>
<td>2.92</td>
<td>3.44</td>
</tr>
<tr>
<td>Simpson</td>
<td>0.96</td>
<td>0.89</td>
<td>0.87</td>
<td>0.97</td>
<td>0.88</td>
<td>0.94</td>
</tr>
<tr>
<td>Inverse Simpson</td>
<td>28.0</td>
<td>8.77</td>
<td>7.55</td>
<td>36.3</td>
<td>8.19</td>
<td>15.7</td>
</tr>
<tr>
<td>Species richness</td>
<td>245</td>
<td>165</td>
<td>187</td>
<td>255</td>
<td>163</td>
<td>209</td>
</tr>
<tr>
<td>Pielou’s evenness</td>
<td>0.73</td>
<td>0.59</td>
<td>0.60</td>
<td>0.76</td>
<td>0.57</td>
<td>0.64</td>
</tr>
</tbody>
</table>
Table 3. Relative abundances (% of total sequences) at the phylum level in the six tested inoculums (I1-I6) at initial time (t0 = 0 months) and final time (t4 = 4 months) when the enrichment was started at low nitrite (LN) and high nitrite (HN) concentrations. “Others” includes all groups with relative abundance <1%.

<table>
<thead>
<tr>
<th></th>
<th>I1</th>
<th>I2</th>
<th>I3</th>
<th>I4</th>
<th>I5</th>
<th>I6</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>t0</td>
<td>t4-LN</td>
<td>t4-HN</td>
<td>t0</td>
<td>t4-LN</td>
<td>t4-HN</td>
</tr>
<tr>
<td>Euryarchaeota</td>
<td>0.09</td>
<td>0.04</td>
<td>0.07</td>
<td>0.18</td>
<td>0.20</td>
<td>0.25</td>
</tr>
<tr>
<td>Acidobacteria</td>
<td>3.74</td>
<td>0.43</td>
<td>2.18</td>
<td>1.43</td>
<td>1.10</td>
<td>1.26</td>
</tr>
<tr>
<td>Actinobacteria</td>
<td>5.29</td>
<td>1.01</td>
<td>4.00</td>
<td>5.47</td>
<td>0.32</td>
<td>3.04</td>
</tr>
<tr>
<td>Armatimonadetes</td>
<td>6.33</td>
<td>24.43</td>
<td>15.59</td>
<td>4.72</td>
<td>26.41</td>
<td>9.98</td>
</tr>
<tr>
<td>Bacteroidetes</td>
<td>13.07</td>
<td>0.95</td>
<td>0.48</td>
<td>6.13</td>
<td>0.99</td>
<td>0.51</td>
</tr>
<tr>
<td>Candidate_division_JS1</td>
<td>*</td>
<td>*</td>
<td>0.01</td>
<td>0.70</td>
<td>0.10</td>
<td>1.02</td>
</tr>
<tr>
<td>Chlorobi</td>
<td>2.93</td>
<td>14.12</td>
<td>3.46</td>
<td>3.50</td>
<td>4.41</td>
<td>8.73</td>
</tr>
<tr>
<td>Deinococcus-Thermus</td>
<td>*</td>
<td>0.03</td>
<td>2.31</td>
<td>8.75</td>
<td>0.88</td>
<td>4.83</td>
</tr>
<tr>
<td>Firmicutes</td>
<td>1.22</td>
<td>1.57</td>
<td>0.80</td>
<td>13.21</td>
<td>13.47</td>
<td>13.39</td>
</tr>
<tr>
<td>Gemmatimonadetes</td>
<td>1.24</td>
<td>0.96</td>
<td>3.11</td>
<td>1.44</td>
<td>0.58</td>
<td>2.50</td>
</tr>
<tr>
<td>Nitrospirae</td>
<td>*</td>
<td>*</td>
<td>0.00</td>
<td>*</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Planctomycetes</td>
<td>4.54</td>
<td>7.68</td>
<td>3.74</td>
<td>4.19</td>
<td>2.95</td>
<td>1.73</td>
</tr>
<tr>
<td>Proteobacteria</td>
<td>32.47</td>
<td>31.68</td>
<td>49.15</td>
<td>22.04</td>
<td>39.40</td>
<td>36.43</td>
</tr>
<tr>
<td>SHA-109</td>
<td>*</td>
<td>0.27</td>
<td>*</td>
<td>0.01</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Spirochaetae</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Tenericutes</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Others</td>
<td>16.18</td>
<td>5.69</td>
<td>7.78</td>
<td>15.45</td>
<td>5.05</td>
<td>6.96</td>
</tr>
</tbody>
</table>

* Relative abundance <0.01%.
Table 4. Identification of the main operational taxonomic units (OTUs) presented in Fig. 6 and source of their closest relative sequences.

<table>
<thead>
<tr>
<th>OTU</th>
<th>Identification (RDP-II)</th>
<th>Closest relative sequence (BLAST)</th>
<th>Accession number</th>
<th>Similarity (%)</th>
<th>Source of the closest relative sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Bacteria; Armatimonadetes; Armatimonadetes_grp5</td>
<td>clone AMX001BXPT7</td>
<td>LC094877</td>
<td>99</td>
<td>Sludge from an anammox UASB reactor</td>
</tr>
<tr>
<td>2</td>
<td>Bacteria; Proteobacteria; Betaproteobacteria; Rhodocyclales; Rhodocyclaceae; Azoarcus</td>
<td>Azoarcus sp. PA01 16S rRNA</td>
<td>KT784536</td>
<td>99</td>
<td>Unknown</td>
</tr>
<tr>
<td>3</td>
<td>Bacteria; Proteobacteria; Betaproteobacteria; Burkholderiales; Burkholderiaceae</td>
<td>clone AMX001BA4AW</td>
<td>LC094801</td>
<td>99</td>
<td>Sludge from an anammox UASB reactor</td>
</tr>
<tr>
<td>4</td>
<td>Bacteria; Bacteroidetes; Sphingobacteria; Sphingobacteriales; Saprospiraceae</td>
<td>clone E7-201bp</td>
<td>KJ993903</td>
<td>99</td>
<td>Earthworm gut</td>
</tr>
<tr>
<td>5</td>
<td>Bacteria; Firmicutes; Clostridia; Clostridiales; Clostridium sensustricto_1</td>
<td>DGGE gel band RB2-79</td>
<td>KT835589</td>
<td>99</td>
<td>SBRs performing N-removal via nitrite treating swine wastewater</td>
</tr>
<tr>
<td>6</td>
<td>Bacteria; Gemmatimonadetes; Gemmatimonadetes; Gemmatimonadaceae</td>
<td>clone SEAB1AA061</td>
<td>KC432372</td>
<td>99</td>
<td>Wetland</td>
</tr>
<tr>
<td>7</td>
<td>Bacteria; Proteobacteria; Betaproteobacteria; Rhodocyclales; Rhodocyclaceae</td>
<td>clone L1A.6H12</td>
<td>AY989011</td>
<td>97</td>
<td>Soil</td>
</tr>
<tr>
<td>8</td>
<td>Bacteria; Chlorobii; Ignavibacteria; Ignavibacteriales</td>
<td>clone AMX001CIOAL</td>
<td>LC094916</td>
<td>99</td>
<td>Sludge from an anammox UASB reactor</td>
</tr>
<tr>
<td>10</td>
<td>Bacteria; Planctomycetes; Planctomycetacia; Brocadiales; Brocadaceae; Candidatus Brocadia sinica clone MBR_day_30</td>
<td>Candidatus Brocadia sinica clone MBR_day_30</td>
<td>KT023580</td>
<td>99</td>
<td>Anammox biomass in a membrane bioreactor</td>
</tr>
<tr>
<td>11</td>
<td>Bacteria; Proteobacteria; Gammaproteobacteria; unclassified</td>
<td>clone MISEQ01_89</td>
<td>KP356069</td>
<td>99</td>
<td>Biofilm in a bioelectrochemical system</td>
</tr>
<tr>
<td>12</td>
<td>Bacteria; Bacteroidetes; unclassified</td>
<td>cloneAMX001B2NAV</td>
<td>LC094861</td>
<td>99</td>
<td>Sludge from an anammox UASB reactor</td>
</tr>
<tr>
<td>13</td>
<td>Bacteria; Chlorof flexi; Anaerolineae; Anaerolineaceae</td>
<td>clone HAD103</td>
<td>HG380607</td>
<td>99</td>
<td>Simultaneous autotrophic and heterotrophic denitrification process</td>
</tr>
<tr>
<td>14</td>
<td>Bacteria; Acidobacteria; unclassified</td>
<td>DGGE band ANAMMOX11</td>
<td>AM900571</td>
<td>99</td>
<td>Anammox batch culture</td>
</tr>
</tbody>
</table>
FIGURE CAPTIONS

Fig. 1. Example of evolution of the N-forms throughout the 4-month experimental period in the bottle seeded with I1 when process was started at (A) low nitrite concentration and (B) high nitrite concentration.

Fig. 2. Average copy number of bacterial 16S rRNA and anammox hzo genes throughout the 4-month experimental period in the bottle seeded with I1 when process was started at (A) low nitrite concentration and (B) high nitrite concentration. Error bars represent ± standard deviation.

Fig. 3. Correlation between the ammonium conversion rate (ACR) and the hzo gene copy number at the end of the enrichment period (4 months) in the 6 batch cultures positive in anammox activity.

Fig. 4. Non-metric multidimensional scaling (NMDS) plot showing the evolution of the microbial community structure in both bottles seeded with I3 throughout the 4-month experimental period. Circles correspond to the bottle started at low nitrite concentration whereas squares correspond to the bottle started at high nitrite concentration. Numbers inside circles and squares indicate the experimental time elapsed (in months).

Fig. 5. Non-metric multidimensional scaling (NMDS) plot showing the evolution of the microbial community structure for all the inoculums tested (1-6) throughout the 4-month experimental period. Triangles correspond to initial time, circles correspond to final time for cultures started at low nitrite concentration and squares correspond to final time for cultures started at high nitrite concentration.

Fig. 6. Evolution of the microbial community structure level in both bottles seeded with I3 (LN: process started at low nitrite concentration, HN: process started at high nitrite concentration) throughout the 4-month experimental period (t0-t4). Analysis was performed (A) at the phylum level and also (B) at the genus level.

Abbreviations: Arc, Euryarchaeota; B<1%, sum of all operational taxonomic units (OTUs) with relative abundance <1%; A/A, Actinobacteria/Acidobacteria; Bc, Bacteroidetes; Cb, Chlorobi; Cx, Chloroflexi; D, Deinococcus-Thermus; F, Firmicutes; G, Gemmatimonadetes; A, Armatimonadetes; Pl, Planctomycetes;
Proteobacteria (α, Alphaproteobacteria; β, Betaproteobacteria; δ, Deltaproteobacteria; γ, Gammaproteobacteria); u, unknown. * is for Candidatus_Brocadia. Main OTUs are numbered and identified in Table 4.

Fig. 7. Percentage of sequences related to anammox genera in the enriched inoculums (I1-I6) at initial time (t0 = 0 months) and final time (t4 = 4 months). Only cultures started at low nitrite concentration are included. *, not detected when high-throughput sequencing was performed on DNA extracts using “universal” 16S rRNA primers and neither when sequencing was performed using PCR products obtained using Planctomycetes primers. **, only detected when sequencing was performed using PCR products obtained using Planctomycetes primers, so relative abundance cannot be quantified (<0.009%).
FIGURES

Fig. 1
Fig. 2
Fig. 3
Fig. 4
Fig. 5
Fig. 6
Fig. 7
HIGHLIGHTS

- Effect of nitrite supply during batch enrichment of anammox sludge was investigated.
- Anammox activity was only developed when starting at low nitrite concentration.
- Final abundance of the $hzo$ gene and anammox activity were positively correlated.
- Inoculum and enrichment conditions determined final microbial community structure.
- Enrichment conditions prompted selection of anammox species belonging to Ca. *Brocadia*. 