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## **OPEN** Modeling of Nitrous Oxide **Production from Nitritation Reactors Treating Real Anaerobic Digestion Liquor**

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In this work, a mathematical model including both ammonium oxidizing bacteria (AOB) and heterotrophic bacteria (HB) is constructed to predict N<sub>2</sub>O production from the nitritation systems receiving the real anaerobic digestion liquor. This is for the first time that N<sub>2</sub>O production from such systems was modeled considering both AOB and HB. The model was calibrated and validated using experimental data from both lab- and pilot-scale nitritation reactors. The model predictions matched the dynamic N<sub>2</sub>O, ammonium, nitrite and chemical oxygen demand data well, supporting the capability of the model. Modeling results indicated that HB are the dominant contributor to N<sub>2</sub>O production in the above systems with the dissolved oxygen (DO) concentration of  $0.5-1.0 \text{ mg O}_2/\text{L}$ , accounting for approximately 75% of N<sub>2</sub>O production. The modeling results also suggested that the contribution of HB to N<sub>2</sub>O production decreased with the increasing DO concentrations, from 75% at DO =  $0.5 \text{ mg O}_2/L$ to 25% at DO = 7.0 mg O<sub>2</sub>/L, with a corresponding increase of the AOB contribution (from 25% to 75%). Similar to HB, the total  $N_2O$  production rate also decreased dramatically from 0.65 to 0.25 mg N/L/h when DO concentration increased from 0.5 to 7.0 mg O<sub>2</sub>/L.

Nitrous oxide  $(N_2O)$  is not only a potent greenhouse gas, with a global warming potential of approximately 265 times stronger than carbon dioxide<sup>1</sup>, but also a compound destructing the stratospheric ozone layer<sup>2</sup>. N<sub>2</sub>O can be produced in the wastewater treatment systems through both nitrification and denitrification<sup>3-7</sup>. During nitrification, ammonium  $(NH_4^+)$  is first being oxidized to nitrite  $(NO_2^-)$  by ammonium-oxidizing bacteria (AOB). Afterwards, nitrite is further oxidized to nitrate  $(NO_3^-)$  by nitrite-oxidizing bacteria (NOB).  $N_2O$  is not an obligatory intermediate of nitrification, but it can be generated by AOB via two primary pathways: i) N<sub>2</sub>O as the end product of AOB denitrification, called the nitrifier denitrification pathway, and ii) N<sub>2</sub>O as the by-product of incomplete oxidation of hydroxylamine (NH<sub>2</sub>OH), known as the NH<sub>2</sub>OH pathway<sup>7-9</sup>. By contrast, N<sub>2</sub>O is an obligatory intermediate of heterotrophic denitrification, which is composed of reductive reactions transforming  $NO_3^-$  to  $NO_2^-$ , nitric oxide (NO),  $N_2O$  and finally to nitrogen gas ( $N_2$ ). These reactions are carried out by heterotrophic bacteria (HB). N<sub>2</sub>O can accumulate when the N<sub>2</sub>O production is faster than the N<sub>2</sub>O reduction<sup>6-8,10</sup>.

It has become a common practice to remove nitrogen from the anaerobic digestion liquor in a side-stream process in wastewater treatment plants (WWTPs)<sup>11,12</sup>. The anaerobic digestion liquor has a high ammonium concentration of 300-1500 mg N/L and a low ratio of chemical oxygen demand to nitrogen (COD/N) for the conventional nitrification and denitrification process. One treatment option is partial nitritation ( $NH_4^+ \rightarrow NO_2^-$ ) followed by the anammox process<sup>12,13</sup>. The partial nitritation process oxidizes approximately 50% of the ammonium to nitrite. This generates a mixture of nitrite and ammonium with a molar ratio of approximately 1:1, which is suitable for the subsequent anammox process. The other treatment option is nitritation followed by denitritation with the additional addition of an external carbon source such as methanol<sup>14</sup>.

 $N_2O$  emissions from the nitritation systems receiving the anaerobic digestion liquor have been extensively reported. However, the results show a huge variation. For instance, the N<sub>2</sub>O emission factors (mg N<sub>2</sub>O-N emitted

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per mg NH<sub>4</sub><sup>+</sup>-N oxidized) were determined to be from 0.7 to 19.3% of the NH<sub>4</sub><sup>+</sup>-N oxidized<sup>3,12,14-19</sup>. Mathematical modeling is an appropriate method for estimating site-specific N<sub>2</sub>O emissions and is of great importance towards a full understanding of N<sub>2</sub>O production mechanisms from the nitritation reactors treating anaerobic digestion liquor. However, little effort has been dedicated to modeling N<sub>2</sub>O production from such systems, which are characterized by high nitrite accumulation and are significantly different from the main-stream wastewater treatment systems. Although Ni *et al.*<sup>9</sup> modeled N<sub>2</sub>O production from such systems using an electron carrier (EC)-based mathematical model, the modeled nitritation system received organics-free synthetic digestion liquor. Therefore, the model did not include N<sub>2</sub>O production by heterotrophic bacteria (HB), which were recently experimentally demonstrated to play an important role in N<sub>2</sub>O production by HB in a methanol and nitrate-fed denitrifying culture. This model has been proven to be able to better predict N<sub>2</sub>O production than the commonly used four-step denitrification model<sup>21,22</sup>. However, this model has never been applied to model N<sub>2</sub>O production from the nitritation reactors treating anaerobic digestion liquor.

In this work, we aim to develop and calibrate a mathematical model to predict  $N_2O$  production from the nitritation reactors receiving the real anaerobic digestion liquor. This model integrated the EC-based  $N_2O$  production model for AOB with that for HB for the first time. The model was calibrated and validated by comparing the simulation results with the measured data from two different nitritation reactors (4 L and 500 L, respectively). The contributions of AOB and HB to  $N_2O$  production as well as  $N_2O$  production rates of both AOB and HB at various dissolved oxygen (DO) levels from such systems were also evaluated.

#### Results

**Model calibration.** A two-step procedure was applied to calibrate the model. In the first step, the kinetic parameters related to AOB were tested using the ammonium and nitrite data. Then, the maximum COD oxidation rate ( $r_{COD,max}$ ), was further calibrated using the volumetric N<sub>2</sub>O emission rate and effluent COD data in the second step. The calibration procedure is shown in Fig. S1. In this work, the default kinetic parameters related to AOB could describe the nitrogen conversion profiles well (Fig. 1A). Therefore, there is no need to calibrate the kinetic parameters associated with AOB. We then calibrated r<sub>COD,max</sub>. The calibration of the r<sub>COD,max</sub> involved optimizing the value of this parameter by fitting the simulation results to the experiment data from Nitritation reactor I at DO = 0.5 mg O<sub>2</sub>/L. The measured and simulated N<sub>2</sub>O dynamics, along with the effluent COD data are illustrated in Figs 1 and 2. The model captured all these trends well. The good agreement between these simulated and measured NH<sub>4</sub><sup>+</sup>, NO<sub>2</sub><sup>-</sup>, effluent COD and N<sub>2</sub>O dynamics supported that the model could be used to estimate N<sub>2</sub>O production from the nitritation reactors receiving anaerobic digestion liquor.



**Figure 1.** Model calibration results using the experimental data from Nitritation reactor I at  $DO = 0.5 \text{ mg } O_2/L$  (measured data: symbols; model predictions: lines): (A)  $NH_4^+$  and  $NO_2^-$  data; (B) Volumetric  $N_2O$  emission rate; (C) Liquid phase  $N_2O$  concentration.



Figure 2. Measured and simulated effluent COD data in Nitritation reactors I and II. Case 1: DO = 0.5 mg O<sub>2</sub>/L in Nitritation reactor I; Case 2: DO = 1.0 mg O<sub>2</sub>/L in Nitritation reactor I; Case 3: DO = 3.0 mg O<sub>2</sub>/L in Nitritation reactor I; Case 4; Data on 16 March 2011 in Nitritation reactor II; Case 5: Data on 21 March 2011 in Nitritation reactor II.

The calibrated value of  $r_{COD,max}$ , which gives the optimum model fittings with the experimental data, is listed in Table S1, together with its 95% confidence interval. The calibrated  $r_{COD,max}$  value of  $1.33 \pm 0.02$  mmol COD/ (gVSS'h) (with 95% confidence interval) (VSS: volatile suspended solids) is much smaller than that obtained (8.46 mmol COD/(gVSS\*h)) by Pan *et al.*<sup>20</sup>. This is likely because only 35% of the biomass were HB in this study<sup>3</sup>, whereas the majority (>90%) of the biomass were HB in the study of Pan *et al.*<sup>20</sup>. Also, the anaerobic digestion liquor and the product of biomass decay were the source of COD in this study, which is difficult to be utilized<sup>23</sup>. In contrast, the readily biodegradable methanol was used as COD source in the study of Pan *et al.*<sup>20</sup>. The 95% confidence interval of  $r_{COD,max}$  (0.02 mmol COD/(gVSS\*h)) is approximately 2% of the estimated value (1.33 mmol COD/(gVSS\*h)), indicating that parameter  $r_{COD,max}$  has a high-level of identifiability and the estimated value is reliable.

**Model validation and further evaluation.** Model validation was based on the comparison between the model predictions (using the same value of  $r_{COD,max}$  as obtained in model calibration) and the experimental data from Nitritation reactor I under different DO conditions (i.e. 1.0 and 3.0 mg O<sub>2</sub>/L). The model predictions and the experimental results are shown in Figs 2 and 3. The validation results show that the model predictions match the measured effluent COD,  $NH_4^+$ ,  $NO_2^-$  and  $N_2O$  dynamics well in all these validation experiments, which supports the validity of the N<sub>2</sub>O model.

To further evaluate the N<sub>2</sub>O model, the experimental results of the effluent COD,  $NH_4^+$ ,  $NO_2^-$  and N<sub>2</sub>O dynamics from Nitritation reactor II were also used. It should be noted that Nitritation reactor I was conducting partial nitritation, whereas Nitritation reactor II was performing nitritation-denitritation. They represented two different treatment options of the anaerobic digestion liquor. The experimental results collected on 16 March 2011 were used to recalibrate  $r_{COD,max}$ , and the results collected on 21 March 2011 were used for model validation. The value of  $r_{COD,max}$  was first recalibrated. The obtained value of  $r_{COD,max}$  for Nitritation reactor II is  $4.78 \pm 1.35 \text{ mmol COD/(gVSS*h)}$  (with 95% confidence interval). This value is higher than that  $(1.33 \pm 0.02 \text{ mmol COD/(gVSS*h)})$  obtained for Nitritation reactor I. This indicates that the COD in the anaerobic digestion liquor of Nitritation reactor II is easier to be biodegraded than that of Nitritation reactor I. The validation of the resulting value of  $r_{COD,max}$  was further performed through comparison of the model predictions with the experimental data of  $NH_4^+$ ,  $NO_2^-$ , effluent COD and N<sub>2</sub>O dynamics collected on 21 March 2011. Figure 4 shows the modeling and experimental results of the model calibration (Fig. 4A,B) and model validation (Fig. 4C,D). As can be seen in Fig. 3, the model predictions are consistent with the experimental results and no systematic deviations are observed, further suggesting that the model is appropriate for describing the N<sub>2</sub>O production in the nitritation reactor fed with anaerobic digestion liquor.

### Discussion

In this work, a mathematical model including both AOB and HB is constructed to predict N<sub>2</sub>O production from the nitritation reactors receiving real anaerobic digestion liquor. This is for the first time that N<sub>2</sub>O production from such systems was modeled considering both AOB and HB. The kinetic parameter ( $r_{COD,max}$ ) closely related to N<sub>2</sub>O production by HB was estimated from the experimental data. The value obtained was robust in its ability to predict N<sub>2</sub>O dynamics. The validity of this model was confirmed by the independent NH<sub>4</sub><sup>+</sup>, NO<sub>2</sub><sup>-</sup>, COD and N<sub>2</sub>O data from both the lab-scale and the pilot-scale nitritation reactors receiving real anaerobic digestion liquor. The successful application of the model in this work indicates that it is applicable to describe N<sub>2</sub>O production in the nitritation reactors receiving anaerobic digestion liquor.

We also performed additional simulation studies using only the AOB  $N_2O$  model<sup>9</sup> to evaluate the measured  $N_2O$  data from the two nitritation reactors used in this work. The model parameters for the AOB  $N_2O$  model were the same as those in Nitritation reactors I and II. The results showed that this model could not reproduce the measured  $N_2O$  data (see Fig. S2). This is due to the fact that both AOB and HB play a role in  $N_2O$  production



**Figure 3.** Model validation results using the experimental data from Nitritation reactor I at  $DO = 1.0 \text{ mg } O_2/L$  (A–C) and 3.0 mg  $O_2/L$  (D–F) (measured data: symbols; model predictions: lines). (A,D)  $NH_4^+$  and  $NO_2^-$  data; (B,E) Volumetric  $N_2O$  emission rate; (C,F) Liquid phase  $N_2O$  concentration.

in the nitritation systems receiving real anaerobic digestion liquor. This is in contrast to the conclusion of other studies<sup>8,12,14</sup>, in which the contribution of HB to  $N_2O$  production was assumed to be negligible.

Figure 5A shows the model predicted contributions of AOB and HB to the N<sub>2</sub>O production from Nitritation reactors I and II. HB are the dominant contributors to the N<sub>2</sub>O production, accounting for approximately 75% in both Nitritation reactors I ( $DO = 0.5 \text{ mg O}_2/L$ ) and II ( $DO = 0.5-1.0 \text{ mg O}_2/L$ ). In contrast, only 25% of N<sub>2</sub>O production can be attributed to AOB. This for the first time quantifies the contribution of HB and AOB to the N<sub>2</sub>O production in the nitritation systems receiving real anaerobic digestion liquor.

The contributions of AOB and HB to the aerobic  $N_2O$  production at different DO concentrations are also evaluated using Nitritation reactor I as an example. Figure 5B shows that the contribution of HB to the aerobic  $N_2O$  production decreases from 75% at  $DO = 0.5 \text{ mg } O_2/L$  to 25% at  $DO = 7.0 \text{ mg } O_2/L$ , with a corresponding increase of AOB contribution (from 25% to 75%). These results suggest that AOB are the dominant contributor to aerobic  $N_2O$  production only when DO in the nitritation systems is high (e.g.  $> 3.0 \text{ mg } O_2/L$ ), whereas HB would be responsible for the majority of the  $N_2O$  production at low DO levels. Wang *et al.*<sup>3</sup> indicated that both AOB and HB contributed to the  $N_2O$  production in Nitritation reactor I and the contribution of HB to the  $N_2O$  production decreased with increasing DO. Our modeling results support the observations made in Wang *et al.*<sup>3</sup>. Figure 5B also shows the average aerobic  $N_2O$  production rates of both AOB and HB at various DO levels. In general, aerobic  $N_2O$  production rate of AOB increases with increasing DO, which is in agreement with results reported by Law *et al.*<sup>24</sup>. In contrast, aerobic  $N_2O$  production rate of HB decreases with increasing DO, from 0.49 mg N/L/h



Figure 4. Model calibration and validation results using the experimental data from Nitritation reactor II (measured data: symbols; model predictions: lines). (A,B)  $NH_4^+$ ,  $NO_2^-$  and volumetric  $N_2O$  emission rate results for calibration; (C,D)  $NH_4^+$ ,  $NO_2^-$  and volumetric  $N_2O$  emission rate results for validation.



**Figure 5.** Model predictions of (**A**) the contributions of AOB and HB to  $N_2O$  production in Nitritation reactors I and II, and (**B**) average aerobic  $N_2O$  production rates in Nitritation reactors I at various DO levels.

at  $DO = 0.5 \text{ mg } O_2/L$  to 0.07 mg N/L/h at  $DO = 7.0 \text{ mg } O_2/L$ . This is due to the fact that higher DO inhibits heterotrophic nitrite reduction, thereby decreasing N<sub>2</sub>O production by HB. Figure 5B also shows that the total N<sub>2</sub>O production rate (i.e. N<sub>2</sub>O production rate of both AOB and HB) decreased dramatically from 0.65 to 0.28 mg N/L/h when DO concentration increased from 0.5 to 3.0 mg O<sub>2</sub>/L. Afterwards, the total N<sub>2</sub>O production rate remained relatively stable at approximately 0.25 mg N/L/h even if DO concentration increased from 3.0 to 7.0 mg O<sub>2</sub>/L. In contrast, the ammonium oxidizing rate kept increasing when DO concentration increased from 0.5 to 3.0 mg O<sub>2</sub>/L, but remained stable when DO concentration increased from 3.0 to 7.0 mg O<sub>2</sub>/L. (see Fig. S3). This indicates that DO should be maintained at around 3.0 mg O<sub>2</sub>/L from the perspective of minimizing N<sub>2</sub>O production rate and maximizing ammonium oxidizing rate, energy consumption would increase accordingly. This will increase operating costs. Therefore, a trade-off has to be made in practice between minimizing N<sub>2</sub>O production rate and minimizing costs.

It should be noted that the potential existence of NOB was not considered in the current model. This is acceptable due to the fact that only a tiny amount of nitrate was produced (less than 10% of ammonium oxidized) in the studied systems. Also, NOB are known not to contribute to  $N_2O$  production<sup>7</sup>. However, the processes related to NOB could be easily incorporated into the model based on the study of Moussa *et al.*<sup>25</sup> if necessary in future applications. It should also be pointed out that the biomass growth of AOB and HB was negligible in a short batch test of a few hours. Therefore, the simulation results regarding their concentrations were not shown in the study. Also, only  $r_{COD,max}$  regarding N<sub>2</sub>O production by HB was estimated from the experimental data. The parameters with regard to AOB (e.g.  $r_{NH3}$ , ox,  $K_{O2,NH3}$ ) were not calibrated in this work because the adopted values from literature were able to describe the NH<sub>4</sub><sup>+</sup> and NO<sub>2</sub><sup>-</sup> data well. These parameter values could be system-specific, and may need calibration when the model is applied to other systems. It should be highlighted that a free nitrous acid (FNA i.e. HNO<sub>2</sub>)-related Haldane-type kinetics was added to the model structure (see Section of "Mathematical model for N<sub>2</sub>O production) to describe the heterotrophic N<sub>2</sub>O reduction. This is necessary for modeling heterotrophic N<sub>2</sub>O reduction in the nitritation systems receiving anaerobic digestion liquor. FNA has been reported to have an inhibitory effect on heterotrophic N<sub>2</sub>O reduction<sup>26</sup>. For instance, Zhou *et al.*<sup>26</sup> demonstrated that FNA inhibition on N<sub>2</sub>O reduction initiated at an FNA concentration of 0.0002 mg HNO<sub>2</sub>-N/L. When the FNA concentrations in the Nitritation reactors I and II could reach 0.32 and 0.009 mg HNO<sub>2</sub>-N/L, respectively (see Fig. S4 for the FNA concentrations in Nitritation reactors I and II). Therefore, FNA inhibition would occur.

In conclusion, a mathematical model including both ammonium-oxidizing bacteria (AOB) and heterotrophic bacteria (HB) is constructed to predict  $N_2O$  production from the nitritation reactors receiving real anaerobic digestion liquor for the first time. Model calibration and validation show good agreement between the simulation results and the experimental results obtained from both lab- and pilot-scale nitritation reactors receiving real digestion liquor. HB are the dominant contributors to the  $N_2O$  production in both reactors. The contribution of HB to aerobic  $N_2O$  production decreases with increasing DO levels, with a corresponding increase of AOB contribution. Also, the  $N_2O$  production rate of HB decreased with the increasing DO levels, whereas the  $N_2O$  production rate of AOB increased when DO concentration increased. The model is expected to enhance our ability to predict  $N_2O$  production from such systems.

#### Materials and methods

Experimental data for model evaluation. Nitritation reactor I. Experimental data from a culture performing partial nitritation previously reported in Wang et al.<sup>3</sup> were used for the model calibration and validation. The culture was developed in a 4-L lab-scale sequencing batch reactor (SBR) fed with anaerobic digestion liquor collected from an Australian wastewater treatment plant. The anaerobic digestion liquor contained approximately  $860 \pm 13 \text{ mg NH}_4^+$ -N/L and  $345 \pm 15 \text{ mg COD/L}$  with the other composition described in Wang *et al.*<sup>3</sup>. The SBR was operated with a cycle time of 6 h, consisting of 5 min aerobic feeding I, 120 min aerobic reaction I, 35 min anoxic reaction I, 5 min aerobic feeding II, 120 min aerobic reaction II, 35 min anoxic reaction II, 2 min aerobic sludge wasting, 25 min settling, 8 min decanting and 5 min anoxic mixing. 0.5 L of anaerobic digestion liquor was fed in each feeding period giving a hydraulic retention time (HRT) of 24 h. The sludge retention time (SRT) was kept at 11 days. The temperature was controlled at  $33 \pm 1$  °C using a water jacket. During the feeding, aerobic reaction and wasting phases, aeration was supplied to maintain a DO concentration of around  $0.5 \text{ mg O}_2/\text{L}$  using an on/off controller. The pH in the SBR varied between 6.4 and 7.1 during a typical cycle with NaHCO<sub>3</sub> solution (1 M) being added (0.1–3.0 ml per cycle) automatically via a programmable logic controller (PLC) when pH was below 6.4. This culture converted approximately  $50 \pm 5\%$  of the NH<sub>4</sub><sup>+</sup>-N to NO<sub>2</sub><sup>-</sup>-N, resulting in both effluent ammonium and nitrite concentrations of  $430 \pm 40$  mg N/L. Effluent nitrate was below 10 mg N/L at all times in the SBR. The effluent COD was  $245 \pm 16$  mg/L. Microbial community analysis revealed that AOB accounted for 65% of the entire microbial community, with the other 35% being HB. In contrast, NOB were not detected (<1%). Mixed liquor suspended solids (MLSS) and mixed liquor volatile suspended solids (MLVSS) concentrations were  $750 \pm 30$  mg/L and  $610 \pm 30$  mg/L, respectively. Both the gas and liquid phase N<sub>2</sub>O were monitored every 3–4 days using an on-line gas analyzer and a liquid microsensor, respectively. More details of the reactor operation and performance can be found in Wang et al.<sup>3</sup>.

In addition, two tests were conducted to evaluate the effect of DO concentrations on aerobic N<sub>2</sub>O production. In these tests, DO concentrations in the aerobic phases of the SBR were controlled at around 1.0 and 3.0 mg O<sub>2</sub>/L, respectively, by adjusting air flow rate (2 L/min) and using an on/off controller. As the bacterial activity increases with the increasing DO concentrations, length of the aerobic phase was shortened to make sure that the effluent ammonium, and nitrite concentrations were comparable with those achieved at the DO level of 0.5 mg O<sub>2</sub>/L. Other operating conditions were the same as those at DO = 0.5 mg O<sub>2</sub>/L. More details can be found in Wang *et al.*<sup>3</sup>.

The gas phase N<sub>2</sub>O concentration was analyzed with an infrared analyzer (URAS 14 Advance Optima, ABB). The data were logged every 3 s. The liquid phase N<sub>2</sub>O was measured online using a N<sub>2</sub>O microsensor (N<sub>2</sub>O-100, Unisense A/S. Aarhus, Denmark). The N<sub>2</sub>O emission rate (mg N<sub>2</sub>O-N/h) was calculated by multiplying the gas phase N<sub>2</sub>O concentration by the known gas flow rate. The volumetric N<sub>2</sub>O emission rate (mg N<sub>2</sub>O-N/L/h) was calculated by dividing the N<sub>2</sub>O emission rate by the volume of the mixed liquor in the SBR. Mixed liquor samples were taken periodically using a syringe and immediately filtered through disposable Millipore filters (0.22  $\mu$ m pore size) for NH<sub>4</sub><sup>+</sup>, NO<sub>2</sub><sup>-</sup> and NO<sub>3</sub><sup>-</sup> analyses using a Lachat QuikChem8000 Flow Injection Analyzer. COD was analyzed according to Standard Methods<sup>27</sup>.

*Nitritation reactor II.* Experimental data from a culture performing nitritation-denitritation previously reported in Lemaire *et al.*<sup>28</sup> were used to further evaluate the model. The culture was developed in a 500-L pilot-scale SBR receiving anaerobic digestion liquor from a French WWTP. The anaerobic digestion liquor contained approximately  $475 \pm 40 \text{ mg NH}_4^+$ -N/L and  $180 \pm 50 \text{ mg COD/L}$  with the other composition described in Lemaire *et al.*<sup>28</sup>. The SBR cycle was divided into several sub-cycles with each composed of aerobic feeding, aerobic reaction and anoxic reaction. Ethanol was added as a carbon source at the beginning of each anoxic reaction to provide enough

carbon sources for nitrogen removal. After reaching the working volume of the SBR, the sub-cycle was completed. This was followed by a short sludge wastage phase to keep the SRT at around 15–20 days, a 1 h settling phase and a decanting phase to discard the treated effluent. The SBR was operated at 20–25 °C. During the aerobic feeding and aerobic reaction phases, DO was between 0.5 and 1.0 mg  $O_2/L$ . pH varied between 6.7 and 7.1 during the cycle. Ammonium, nitrite and nitrate were measured by on-line sensors. This culture achieved total nitrogen removal of 85–90% and the effluent COD was around 120 mg/L. The gas phase N<sub>2</sub>O was measured continuously using an infrared analyzer (VA-3000, Horiba, Japan). More details of the reactor operation and performance can be found in Lemaire *et al.*<sup>28</sup>. The calculation of the N<sub>2</sub>O emission rate was similar to that for Nitritation reactor I.

**Mathematical model for N<sub>2</sub>O production.** The previously proposed EC-based N<sub>2</sub>O models by Ni *et al.*<sup>9</sup> and Pan et al.<sup>20</sup> were integrated for the first time to describe the N<sub>2</sub>O data. In the N<sub>2</sub>O model developed by Ni et al.<sup>9</sup>, the NH<sub>2</sub>OH oxidation and AOB denitrification pathways were integrated to describe N<sub>2</sub>O production by AOB. In this model, the oxidation and reduction processes were modeled separately, with intracellular electron carriers introduced to link the two types of processes (see Table S1–S4). In the N<sub>2</sub>O model developed by Pan et al.<sup>20</sup>, the electron competition by different denitrification steps was described for the first time, through modeling the carbon oxidation and nitrogen reduction processes separately (see Table S1-S4). Electron carriers were introduced to link carbon oxidation and nitrogen oxides reduction. In addition, the biomass decay process was added to describe the generation of slowly biodegradable COD after cell death. The generated slowly biodegradable COD can then be hydrolyzed to readily biodegradable COD, which can be further utilized to produce the reduced form of the electron carrier during COD oxidation by HB. As free nitrous acid (FNA i.e. HNO2) has an inhibitory effect on N<sub>2</sub>O reduction<sup>26</sup> (see Fig. S4 for FNA concentrations in Nitritation reactors I and II), a Haldane-type kinetics  $(S_{HNO2}/(K_{HNO2} + S_{HNO2} + (S_{HNO2})^2/K_{I,HNO2}))$  was applied to describe the N<sub>2</sub>O reduction (R14 in Table S3). Because the nitrite concentration in Nitritation reactor I was between 390 and 500 mg N/L and a high level of nitrite (>50 mg N/L) has an inhibitory effect on N<sub>2</sub>O production<sup>29</sup>, a Haldane-type kinetics (S<sub>NO2</sub><sup>-/</sup>  $(K_{NQ2}^{-} + S_{NQ2}^{-} + (S_{NQ2}^{-})^2/K_{LNQ2}^{-}))$  was added to describe the NO<sub>2</sub><sup>-</sup> reduction for N<sub>2</sub>O production in Nitritation reactor I (R6 in Table S3). Since ethanol is much more easily biodegradable than the other biodegradable COD that exists in the anaerobic digestion liquor and which is produced from biomass decay, the anoxic ethanol oxidation process  $\left(r_{ethanol, \max} XHB\left(\frac{Sethanol}{Ks + Sethanol}\right)\left(\frac{SMox, HB}{KMox, HB + SMox, HB}\right)\left(\frac{KO2, HB}{KO2, HB + SO2}\right)\right)$  (see Tables S2 and S4) was introduced to model anoxic ethanol oxidation in Nitritation reactor II. The components, kinetic rate expressions, stoichiometric matrix, and kinetic and stoichiometric parameters of the N<sub>2</sub>O model are summarized in Table S1-S4.

The biomass concentrations of AOB and HB for model input were calculated based on microbial community analysis results and MLVSS concentration. The measured MLVSS concentration was apportioned to the bacterial populations based on the determined fractions of various bacterial populations including AOB and HB. This is the commonly used method of determining the concentrations of bacterial populations<sup>9,30</sup>. The biomass concentrations of AOB and HB were then determined as 210 mg/L and 400 mg/L, respectively, in Nitritation reactor I. In Nitritation reactor II, the biomass concentrations of AOB and HB were approximately 280 mg/L and 1600 mg/L, respectively.

**Model calibration and validation.** The  $N_2O$  model includes 45 stoichiometric and kinetic parameters, as summarized in Table S1. Most of these parameters were well established in previous studies. Therefore, literature values were directly adopted for these parameters (Table S1). As the  $N_2O$  production pathway by HB is for the first time included in the model for estimating  $N_2O$  production from nitritation systems receiving anaerobic digestion liquor and sensitivity analysis reveals that the maximum COD oxidation rate ( $r_{COD,max}$ ) is the key parameter governing  $N_2O$  production by HB,  $r_{COD,max}$  was estimated using the experimentally obtained volumetric  $N_2O$  emission rate and COD data.

Experimental data from Nitritation reactor I at DO = 0.5 mg O<sub>2</sub>/L were used to calibrate the model. The value of  $r_{COD,max}$  was estimated by minimizing the sum of squares of the deviations between the measured data and the model predictions using a modified version of AQUASIM 2.1d<sup>31</sup>, with 95% confidence interval for parameter uncertainty analysis. The 95% confidence interval of  $r_{COD,max}$  was calculated from the mean square fitting error. Model validation was then carried out with the calibrated value of  $r_{COD,max}$  using the other two sets of experimental data under different DO conditions (i.e. 1.0 and 3.0 mg O<sub>2</sub>/L).

To further verify the validity and applicability of the  $N_2O$  model, we also applied the model to evaluate the  $N_2O$  data from Nitritation reactor II. The  $r_{COD,max}$  was recalibrated using one cycle data monitored on 16 March 2010 from Nitritation reactor II. The validation step was then carried out with the recalibrated  $r_{COD,max}$  using one cycle monitoring data on 21 March 2010 from Nitritation reactor II with dynamic influent conditions which has not been used to estimate the parameter.

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#### **Author Contributions**

All authors contributed intellectually via scientific discussions during the work. Q.W. developed the methodology, performed modeling work, conducted data analysis, prepared all figures and drafted the manuscript. B.-J.N., R.L., X.H. and Z.Y. contributed to the methodology development. All authors reviewed the manuscript.

### **Additional Information**

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