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BRIEF COMMUNICATION





Indications for enzymatic denitrification to N₂O at low pH in an ammonia-oxidizing archaeon

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Abstract

Nitrous oxide (N₂O) is a key climate change gas and nitrifying microbes living in terrestrial ecosystems contribute significantly to its formation. Many soils are acidic and global change will cause acidification of aquatic and terrestrial ecosystems, but the effect of decreasing pH on N₂O formation by nitrifiers is poorly understood. Here, we used isotope-ratio mass spectrometry to investigate the effect of acidification on production of N₂O by pure cultures of two ammonia-oxidizing archaea (AOA; *Nitrosocosmicus oleophilus* and *Nitrosotenuis chungbukensis*) and an ammonia-oxidizing bacterium (AOB; *Nitrosomonas europaea*). For all three strains acidification led to increased emission of N₂O. However, changes of ¹⁵N site preference (SP) values within the N₂O molecule (as indicators of pathways for N₂O formation), caused by decreasing pH, were highly different between the tested AOA and AOB. While acidification decreased the SP value in the AOB strain, SP values increased to a maximum value of 29‰ in *N. oleophilus*. In addition, ¹⁵N-nitrite tracer experiments showed that acidification boosted nitrite transformation into N₂O in all strains, but the incorporation rate was different for each ammonia oxidizer. Unexpectedly, for *N. oleophilus* more than 50% of the N₂O produced at pH 5.5 had both nitrogen atoms from nitrite and we demonstrated that under these conditions expression of a putative cytochrome P450 NO reductase is strongly upregulated. Collectively, our results indicate that *N. oleophilus* might be able to enzymatically denitrify nitrite to N₂O at low pH.

Eugene L. Madsen deceased on Aug 9, 2017

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N₂O is an important ozone-depleting substance with a high global warming potential [1–3]. The processes of biological N₂O production include partial dissimilatory nitrate (NO₃⁻) or nitrite (NO₂⁻) reduction (denitrification), nitrifier denitrification [2], ammonia [hydroxylamine (NH₂OH)] oxidation, and NOx detoxification (also known as the "nitrosative stress" pathway) [1]. AOB are recognized as a major source of N₂O production from terrestrial environments [4] and AOA are also considered to be important contributors to N₂O production in various environments, based on their high abundance in many ecosystems and the documented formation of N2O during AOA-mediated ammonia oxidation [5–8]. Despite the presence of a nitrite reductase gene (nirK) in almost all AOA genomes, canonical nitric oxide reductase (NOR) genes have not been detected in these organisms. Instead, isotope labeling experiments suggested hybrid N₂O formation in the AOA Nitrososphaera viennensis from nitrite and an intermediate of ammonia oxidation, and was attributed to either an enzymatically catalyzed (codenitrification) or abiotic N-nitrosation reaction [8]. Under low-oxygen conditions, abiotic formation of N₂O from hydroxylamine or NO was observed in experiments with killed *N. viennensis* biomass in AOA media [9].

2634 M.-Y. Jung et al.

Furthermore, it was recently demonstrated that N_2O is formed abiotically under aerobic conditions from hydroxylamine and nitrite produced by aerobic ammonia-oxidizing microbes and that this hybrid pathway can account for a large proportion of the aerobically produced N_2O [10]. Interestingly, we observed in a previous study for selected AOA strains variable N_2O isotopomer SP (enrichment of ^{15}N at the alpha, or beta site of N_2O) values (ca. 20-30%e) and a very variable contribution of both N atoms in N_2O from nitrite (8.4–53% of the N_2O produced by the AOA strains had both N from nitrite) at high initial nitrite concentrations. These results are consistent with the hypothesis that nitrifier denitrification might contribute to N_2O production by some AOA in the presence of excess nitrite [6].

Terrestrial and ocean environments experience acidification due to natural or human activities [11, 12] and a large part (ca. 30%) of the world's natural and arable soils are acidic (pH < 5.5) [13]. However, most of the studies of N₂O production by AOA and AOB have been performed in circumneutral pH ranges only, although various biological and chemical reactions involved in N₂O production are pHdependent [11]. Thus, responses of N₂O production by AOA and AOB to acidification need to be better understood to estimate the present and future contribution of soil N₂O production to the global N₂O budget. The present study was designed to reveal the impact of acidification upon N₂O production by two selected AOA strains and by a model strain of AOB. We analyzed N₂O production and SP of the N₂O molecule under varying pH conditions. In addition, pH-dependent changes of the source of nitrogen of N₂O were investigated using ¹⁵N-nitrite tracer experiments in order to obtain insights into potential differences of the N₂O production pathways between the analyzed strains.

In order to assess the effect of acidification on N₂O production, axenic AOA (Nitrosotenuis chungbukensis MY2 of thaumarchaeal group I.1a; Nitrosocosmicus oleophilus MY3 of thaumarchaeal group I.1b) and AOB (Nitrosomonas europaea ATCC 19718) strains were incubated in growth media spanning a pH range of 5.5-8.5 (at intervals of 0.5 pH units). More details on cultivation methods, media, and incubation conditions are described in the Supplementary Materials and Methods. The lowest pH used in these assays was selected after screening the three cultures for retention of ~20% of the growth rate found at optimal pH. Thus, pHs of 5.5, 6.0, and 6.5 were used as most acidic incubation conditions for N. oleophilus MY3, N. chungbukensis MY2, and for N. europaea ATCC 19718, respectively. For all tested ammonia oxidizers the N₂O yield (N₂O-N/oxidized NH₄⁺-N) increased as pH decreased (Supplementary Fig. S1). The N2O yield of N. chungbukensis MY2 at low pH was much higher than that of N. oleophilus MY3. The increased N2O production at acidic conditions may be caused by increased production of enzymes involved in N₂O production [14], an acidic pH optimum of N₂O-producing enzymes [15, 16], or acceleration of abiotic hybrid N₂O formation [8, 17]. In coculture experiments of the AOA N. oleophilus MY3 with the nitrite-oxidizing bacterium (NOB), Nitrobacter winogradskyi Nb-255 nitrite accumulation was not observed during nitrification at pH 7.5 and pH 5.5 (Supplementary Fig. S2). This *Nitrobacter* strain has no known enzymatic repertoire to produce or oxidize N2O, but encodes a reversible nitrite oxidoreductase that is able to catalyze the oxidation of nitrite to nitrate [18]. Interestingly, for both AOA and AOB at the lowest pH tested, the presence of the NOB in the cultures caused a significant decrease of N₂O yields (28% and 48%, respectively) (P < 0.001), suggesting that the accumulation of nitrite in the experiments without addition of NOB contributed to the increased N2O production under these conditions (Supplementary Fig. S1). Thus, it is tempting to speculate that the higher N₂O production at lower pH may possibly be connected to an upregulation of nitrite detoxification due to increased formation of the reactive compound nitrous acid (HNO₂) from nitrite (NO_2^- , pKa = 3.39) at lower pH.

Many isotopic studies have documented a positive SP value (around 30%) that is the characteristic for N₂O produced by ammonia oxidizers via the formation of NH₂OH [19], with similar values reported for fungal denitrification [20] or chemical formation of N₂O from hydroxylamine [21]. In contrast, SP values for N₂O produced by bacterial heterotrophic denitrification or nitrifier denitrification are both near or below zero [19]. The SP values of N₂O produced by the strains in the present study (N. oleophilus MY3, N. chungbukensis MY2, and N. europaea ATCC 19718) at pH 7.5 were 26%, 29%, and 28%, respectively, resembling the signatures of N₂O produced mostly via NH₂OH formation (Fig. 1a). The SP values of N₂O produced by the AOB strain ATCC 19718 decreased dramatically from $\sim 30\%$ at pH 8.5 to $\sim 0.5\%$ (P < 0.001) at pH 6.5 (Fig. 1a). This shift is consistent with prior observations that increased nitrifier denitrification activity of AOB at decreasing pH [14] may be associated with removal of HNO₂ the toxic form of NO₂. In contrast, for N. chungbukensis MY2 only a slight decrease of SP values of N2O (from 29 to 27%; P = 0.05) was observed when the pH dropped from pH 7.5 to 6.0 (Fig. 1a). Intriguingly, for N. oleophilus MY3 the SP values of N2O even increased from 26 to 29% (P < 0.05), when the pH decreased from 7.5 to 5.5 suggesting no involvement of canonical nitrifier denitrification. Meanwhile, continuous nitrite removal from the N. oleophilus MY3 culture by cocultivation with a NOB had no significant effect on the SP values of N₂O at pH 5.5 (Fig. 1a) indicating independence of N₂O production mechanisms on the external nitrite concentration at low pH. Our observations of opposing and/or variable trends in SP

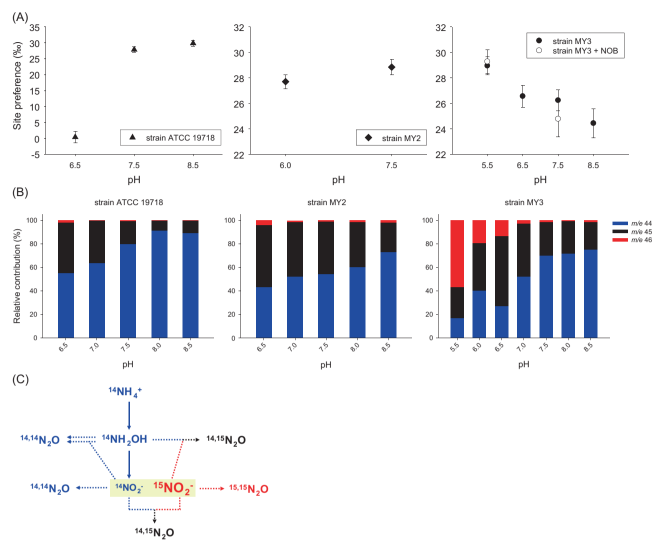


Fig. 1 a ¹⁵N site preference (SP) values of N₂O at various pH conditions for *N. europaea* ATCC 19718, *N. chungbukensis* MY2, and *N. oleophilus* MY3. The SP values of N₂O were measured after ammonia oxidation was completed in the incubation experiments. *Nitrobacter winogradskyi* Nb-255 was the cocultured NOB. The error bars are based on replicate experiments to show the standard deviation and the raw data used in this plot are presented in Supplementary Table S3. **b** The panels depict the composition of labeled N₂O produced during the tracer experiment by *N. europaea* ATCC 19718, *N. chungbukensis* MY2, and *N. oleophilus* MY3. The ammonia oxidizers were incubated in the presence of 0.2 mM unlabeled NH₄⁺ and 0.2 mM ¹⁵N-labeled nitrite at different pH conditions. The values for the different masses of N₂O at each pH condition for each strain are presented as mean values

of triplicate experiments (standard deviations of all values were <5%). c Proposed (bio)chemical processes showing the pathways leading to N_2O production in the tracing experiment. Two different nitrogen sources (from unlabeled-ammonia or ^{15}N -labeled nitrite) permitted three possible mass combinations in the produced N_2O . It should be noted that hydroxylamine is converted to NO by the hydroxylamine dehydrogenase in AOB and then further oxidized to nitrite by an unknown enzyme ([23]; not shown). NO has also been suggested as an important intermediate in the energy metabolism of AOA, but its exact role is still under debate ([9, 23]; not shown). Unlabeled N_2O (m/e 44) can be produced enzymatically by AOB by conversion of N_2OH by cytochrome P460 [29] or chemically in the presence of N_2OH for N_2OH by N_2OH by N

values of N_2O in the different strains as a function of pH indicates that, although N_2O yields increased under acidic conditions in the tested AOA and AOB cultures (see Supplementary Fig. S1), the underlying mechanisms responding to acidification likely to differ.

The SP value of the N₂O produced by the tested AOA does not exclude production of N₂O by a chemical reaction of NO₂⁻ with NH₂OH during ammonia oxidation [10], as

the SP of N_2O abiotically produced from NH_2OH and NO_2 –(or HNO_2) at acidic conditions was reported to be ~34‰ [21]. However, in our abiotic control experiments at pH 5.5 and 7.5, N_2O production was much lower in the presence of 500 μ M of NO_2 –with or without addition of 10 and 50 μ M of NH_2OH compared to corresponding experiments with active AOA and AOB cultures (see Supplementary Table S1 and 2). The NH_2OH concentrations in the abiotic

2636 M.-Y. Jung et al.

controls were selected based on the recently published data that showed maximum concentrations of extracellular NH₂OH of <10 μ M in AOA, AOB, and comammox strains during oxidation of 2 mM ammonia [10]. In this context it should be noted that previously higher N₂O production in abiotic control incubations that we report here has been detected [9, 10]. This difference likely reflects differences in media composition (for example, we used a ~10 × lower trace metal concentration than Kozlowski et al. [9] and Liu et al. [10] for our biotic and abiotic experiments) and/or incubation conditions (e.g., our abiotic experiments were performed fully aerobic in contrast to Kozlowski et al. [9]) between studies.

The increase of the N_2O yield by both tested AOA (and the increase of the SP value of N_2O for *N. oleophilus* MY3) at lower pH cannot be explained by conventional nitrifier denitrification for HNO₂ detoxification as this would be expected to strongly lower the SP value (as observed for *N. europaea*). To further investigate the underlying processes of N_2O production, nitrogen incorporation into N_2O produced by the three nitrifiers was traced by using ^{15}N -labeled nitrite. With this setup, most of m/e 44 ($^{14,14}N_2O$) and all of m/e 46 ($^{15,15}N_2O$) is produced by conversion of unlabeled ammonia and ^{15}N -labeled nitrite, respectively (see Fig. 1b, c). Interestingly, the relative contribution of labeled nitrite to N_2O production (m/e 45 – one labeled N atom + m/e 46) was increased by acidification in all strains (Fig. 1b).

For N. europaea ATCC 19718, the increased N₂O yield and decreased SP at low pH suggests production of N2O via nitrifier denitrification by the combined action of its nitrite reductase and nitric oxide reductase enzymes. Dissolved O2 concentrations in our cultures of AOA and AOB were >95% saturation and those enzymes are also known to be expressed under aerobic conditions [14, 22]. Unexpectedly however, in the ¹⁵N-nitrite labeling experiment doublelabeled N₂O (m/e 46) was not much increased at low pH and instead more hybrid-N₂O formation (m/e 45) was observed (Fig. 1b). It is important to keep in mind that N. europaea ATCC 19718 produces much more NO than AOA at circumneutral pH [9], and the expression of nitrite reductase is further induced by low pH and high nitrite [14], which will lead to increased NO production from nitrite under these conditions. Furthermore, recent biochemical experiments demonstrated that the hydroxylamine dehydrogenase (HAO) of N. europaea produces NO and not nitrite during ammonia oxidation [23]. Consequently, in our labeling experiment at acidic conditions (unlabeled) ¹⁴NO formed from ammonia oxidation by the HAO activity will mix with (labeled) ¹⁵NO formed from nitrite via nitrite reductase. Reduction of this partially labeled NO by NorB (or NorSY [24], or cytochrome c554 [25]; with possibly different contributions of the different NO reductases with varying pH) will result in hybrid N_2O formation (m/e 45).

This hypothesis might also explain the results from recent stable isotope labeling experiments with a natural AOB community in a lake that also indicated increased hybrid N_2O formation with decreasing pH [11].

For *N. chungbukensis* MY2, the nitrite-labeling experiment also suggested an increased contribution of hybrid-N₂O formation (*m/e* 45) at pH 6.5 (Fig. 1b). Lowering the pH might have increased the N₂O yield in *N. chungbukensis* MY2 by increasing chemical hybrid N₂O formation from nitrite and hydroxylamine as previously described [10–12]. However, this is inconsistent with the data from our abiotic control experiments with nitrite and hydroxylamine that showed a much lower N₂O production at pH 6.5 than measured in the corresponding biotic experiment. Thus, our data indicate that in *N. chungbukensis* MY2 N₂O formation at low pH might be catalyzed by an unknown NO-reducing enzyme.

Surprisingly, for N. oleophilus MY3 up to 56.5% of the N₂O produced in our labeling experiment had both N from nitrite (15,15N₂O) implicating a substantial involvement of an unusual nitrifier denitrification process in N2O formation (Fig. 1b). In this context it is interesting to note that the SP values for N₂O produced from fungal denitrification show values of up to 35% [19, 20], similar to those observed by us for N. oleophilus MY3 at low pH. The most characteristic feature of the fungal-denitrifying system is the involvement of cytochrome P450, as NOR (P450nor) [20, 26]. The proposed overall mechanisms for the reduction of NO by the enzyme P450nor is $[2NO + NAD(P)H + H^+ \rightarrow N_2O]$ $+ H_2O + NAD(P)^+$ and the enzyme accept two electrons directly from NAD(P)H, in contrast to other P450 enzymes (non-NOR type) where the electrons are donated one at a time via redox partners involving flavins and iron-sulfur centers [26]. The fungal denitrifying system seems to lack N₂O reductase (NOS) and therefore evolves N₂O as the final product [26]. Interestingly, putative cytochrome P450encoding genes were detected by us in the genomes of N. oleophilus MY3 and some other thaumarchaeal group I.1b members, as well as in some nitrifying bacterial strains (AOB, NOB, and comammox) (Supplementary Fig. S3). The cytochrome P450 of *Nitrosocosmicus* spp., like the fungal P450nor clade (CYP55 NOR), is related to bacterial members of this enzyme superfamily. However, the fungal P450nor clade and the respective enzyme superfamily members of nitrifying bacteria, Nitrososphaera spp. and Nitrosocosmicus spp. are polyphyletic, and without biochemical data no direct proof of their specific enzymatic activity can be obtained. However, at pH 5.5 (compared to pH 7.5), we observed significantly increased transcription of two cytochrome P450-like genes (MY3_00641 and MY3_01637) of strain N. oleophilus MY3 compared to transcription of housekeeping genes (see Supplementary Table S4 for information on qPCR primers) such as those encoding the 16S rRNA, AmoA, and enzymes required for

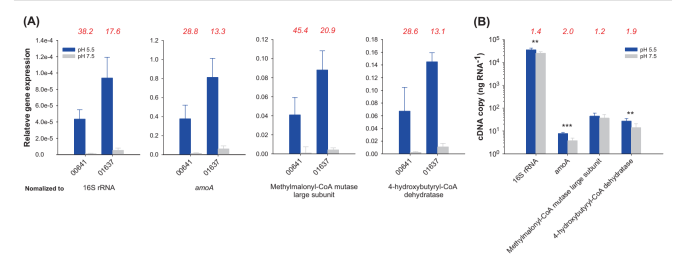


Fig. 2 a *N. oleophilus* MY3 cytochrome P450 cDNA gene expression ratios at two different pH conditions (pH 5.5 and pH 7.5) normalized to other expressed genes. Average of two independent qPCR experiments performed on reverse-transcribed total RNA from cells grown at pH 5.5 and 7.5 and harvested at exponential stage are presented. Relative mRNA expression of two different copies of cytochrome P450 transcripts (locus: MY3_00641 and MY3_01637) to those of representative housekeeping genes [16S rRNA and mRNA of *amoA*, methylmalonyl-CoA mutase large subunit (MY_02370), and 4-hydroxybutyryl-CoA dehydratase (MY3_03315)], were calculated for obtaining normalized data. The ratio of relative expression at pH 5.5–7.5 is indicated above the graphs in red. Normalized expression of both P450 genes is significantly higher at low pH than at high pH in all panels (significance of differences of normalized expression level of

cytochrome P450 gene between pH 5.5 and 7.5 were determined by a t-test (P < 0.05)). Error bars indicate standard deviation of duplicate experiments. The difference between the y-axes of the four panels should be noted. Expression at low pH between the two P450 genes was also found to be significantly different (P < 0.05). **b** Nonnormalized RT-qPCR data cDNA for 16S rRNA, amoA, and methylmalonyl-CoA mutase large subunit (MY_02370) and 4-hydroxybutyryl-CoA dehydratase (MY3_03315) at pH 5.5–7.5. The cDNA transcripts of each gene were quantified per 1 ng of RNA. The ratio of relative expression at pH 5.5–7.5 is indicated above the graphs in red. Error bars indicate standard deviation from duplicate experiments. For each gene, significance of difference in measured cDNA copy number between pH 5.5 and 7.5 was determined by a t-test (*t)-t0.5, **t0.1, and ***t0.05)

CO₂ fixation (Fig. 2 and Supplementary Fig. S4). N₂O yields are significantly increased due to nitrifier denitrification in AOB under low-oxygen conditions [8, 27, 28], which can be higher than those from nitrifier denitrification in *N. oleophilus* MY3 under aerobic conditions at acidic conditions (Supplementary Table S1). Altogether, this suggests that in *N. oleophilus* MY3 cytochrome P450 might be possibly involved in the production of N₂O via nitrifier denitrification acting as NOR under aerobic conditions at a lower pH, a hypothesis that warrants further experimental investigation. If confirmed, cytochrome P450-catalyzed N₂O production in AOA and possibly some other nitrifiers would significantly expand our perception of the metabolic repertoire of these important N-cycle microorganisms and their contribution to global change.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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2638 M.-Y. Jung et al.

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