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To the Graduate Council:

I am submitting herewith a dissertation written by Guang He entitled "Novel microbial guilds implicated in N2O reduction." I have examined the final electronic copy of this dissertation for form and content and recommend that it be accepted in partial fulfillment of the requirements for the degree of Doctor of Philosophy, with a major in Plant, Soil and Environmental Sciences.

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Novel microbial guilds implicated in N2O reduction

A Dissertation Presented for the

**Doctor of Philosophy** 

Degree

The University of Tennessee, Knoxville

**Guang He** 

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#### ABSTRACT

 $N_2O$  is a long-recognized greenhouse gas (GHG) with potential in global warming and ozone depletion. Terrestrial ecosystems are a major source of N<sub>2</sub>O due to imbalanced N<sub>2</sub>O production and consumption. Soil pH is a chief modulating factor controlling net  $N_2O$  emissions, and  $N_2O$  consumption has been considered negligible under acidic conditions (pH <6). In this dissertation, we obtained solids-free cultures reducing N<sub>2</sub>O at pH 4.5. Furthermore, a co-culture (designated culture EV) comprising two interacting bacterial population was acquired via consecutive transfer in mineral salt medium. Integrated phenotypic, metagenomic and metabolomic analysis dictated that the Serratia population excreted certain amino acid to support *Desulfosporosinus* population growth. Characterization of co-culture EV demonstrated that organisms synthesizing functional NosZ exist in acidic soils. To further close the knowledge gap of low pH  $N_2O$  reduction, we conducted enrichment experiments on two contrasting soils representing natural and agricultural soils, respectively. With varying combination of carbon sources and H<sub>2</sub>, N<sub>2</sub>O reduction activity was observed in a total of six solids-free cultures at pH 4.5. Comparative growth experiments documented that N<sub>2</sub>O was essential for N<sub>2</sub>O-reducing organisms' growth. Reconstruction of the communities suggested the cultures were highly enriched following consecutive transfer efforts. Surprisingly, N<sub>2</sub>O-reducing organisms recovered from the cultures were all identified as clade II lineage or a novel clade lineage. These results together suggested that clade II and the putative novel clade N<sub>2</sub>O-reducing organisms are closely associated with low pH N<sub>2</sub>O reduction. The observation of low pH N<sub>2</sub>O reduction in current research was inconsistent to field observation, where the abiotic-biotic interaction occurs frequently. To simulate potential abiotic-biotic interaction of low pH  $N_2O$  reduction, we tested the impact of geochemistry factors on N<sub>2</sub>O reduction by culture EV. The impeded N<sub>2</sub>O reduction of co-culture EV in the presence of trace amount (as low as 0.01 mM for nitrate and 0.001 mM for nitrite) of nitrogen oxyanions was unexpected, which may explain the weak  $N_2O$  reduction activity under field conditions. These observations indicated N<sub>2</sub>O reduction in acidic soils may be limited by interacting abiotic factors, instead of pH itself.

iii

# **TABLE OF CONTENTS**

Chapter 1 Introduction	1
1.1 Sources and sinks of N2O	1
1.2 Acidic soils are mainly N <sub>2</sub> O source	3
1.3 Research objectives	5
Chapter 2 Sustained bacterial N <sub>2</sub> O reduction at acidic pH	7
Abstract	8
2.1 Introduction	9
2.2 Materials and methods Soil sampling locations and microcosms Enrichment cultures Microbial community analysis Isolation and classfication Quantitative PCR (qPCR) Nutritional interactions in the co-culture Metagenome sequencing Comparative analysis of nos gene clusters Phylogenomic analysis NosZ phylogenetic analysis	10 10 11 12 13 14 14 15 16 16
Metabolome analysis Phenotypic characterization of co-culture EV	
Analytical procedures	
2.3 Results	19
A consortium consisting of two populations reduces N <sub>2</sub> O at pH 4.5	
Isolation efforts	
Identification of auxotrophies pH range of acidophilic N2O reduction by the Desulfosporosinus sp	
Phylogenomic analysis	
Genetic underpinning of N <sub>2</sub> O reduction in 'Ca. Desulfosporosinus nitrousreducens' strain PR Genomic insights for a commensalistic relationship Distribution of the novel N <sub>2</sub> O reducer in soil metagenomes	
2.4 Discussion	29
Appendix	35
Chapter 3 Clade III N <sub>2</sub> O-respiring microbes implicated in N <sub>2</sub> O reduction Abstract	
3.1 Introduction	<b>67</b> iv

3.2 Materials and Methods69
Soil sampling, microcosms and enrichment cultures
Comparative genomic analysis70
Phylogenomic analysis
Computational analysis of nos operon cluster and NosZ structure modeling 71
Comparative growth assays with enrichment cultures harboring clade III N <sub>2</sub> O
reducers
Quantitative PCR72
3.3 Results
Low pH N <sub>2</sub> O reducers enriched from circumneutral and acidic soils
Novel species involved in low pH N <sub>2</sub> O reduction
Identification of clade III nosZ
Structural and functional validation of clade III nosZ in enrichment cultures 78
3.4 Discussion79
Appendix
Chapter 4 Geochemical factors control N2O reduction under low pH conditions . 108
Abstract
4.1 Introduction110
4.2 Materials and Methods112
Microorganisms and cultivation112
Abiotic controls of N <sub>2</sub> O reduction at low pH
Analytical procedures
4.3 Results
Effects of copper and Na <sub>2</sub> S on N <sub>2</sub> O reduction
Effects of nitrogen oxyanion on low pH N <sub>2</sub> O reduction
4.4 Discussion116
Appendix119
Summary and Conclusion 123
References125
Vita

# LIST OF TABLES

Table 2-1. qPCR primers and probe designed for the enumeration of Serratia sp. strain MF and 'Ca	•
Desulfosporosinus nitrousreducens' strain PR 16S rRNA genes	56
Table 2-2. Sequence information submitted to the NCBI database	57
Table 2-3. Organismal and genomic information of microorganisms with reported N2O reduction capacity under different pH.	58
Table 2-4. Genomes used in phylogenomic analysis and classification of 'Ca. Desulfosporosinus nitrousreducens'	59
Table 2-5. Amino acids and concentrations used to augment the growth medium for cultivation of 'Ca. Desulfosporosinus nitrousreducens'. a	51
Table 2-6. Genome features of 'Ca. Desulfosporosinus nitrousreducens' strain PR and Serratia sp.         strain MF, and comparison to closest relatives.	53
Table 2-7. Genome features of 'Ca. Desulfosporosinus nitrousreducens' strain PR and Serratia sp.	64
Table 3-1. Metadata associated with soils used for enrichment procedure	)9
Table 3-2. Substrates provided to the N2O-reducing cultures during the enrichment process10	
Table 3-3. Analyses of high-quality MAGs using the Microbial Genome Atlas (MiGA) a10	)1
Table 3-4. Statistics of the high-quality nosZ-carrying MAGs.	)6
Table 3-5. Growth of clade III N <sub>2</sub> O reducers is N <sub>2</sub> O- and pH- dependent. <sup>a</sup> 10	)7

#### LIST OF FIGURES

- Figure 2-1. Establishment of N<sub>2</sub>O-reducing microcosms and enrichment cultures yielding co-culture EV. (A) Schematic of the workflow leading from a soil sample to a solids-free enrichment, and to a co-culture. (B) Community structure profiling based on 16S rRNA gene sequence analysis documents the enrichment process. Soil microbial community structure was profiled with 16S rRNA genes extracted from shotgun metagenomic reads. Community profiling of 6<sup>th</sup> and 9<sup>th</sup> transfers was based on 16S rRNA gene amplicon sequencing. Sequences with abundances <2%were grouped as "Others". (C) Percent of the metagenomic short read fragments obtained from El Verde soil and the 15<sup>th</sup> transfer culture that recruited to the genomes of *Serratia* sp. or Desulfosporosinus sp. A representative graph showing the high identity (>95%) of reads mapping evenly across Desulfosporosinus sp. genome is presented in Supplementary Figure 2-3. The Serratia sp. or Desulfosporosinus sp. genomes were not detected in the soil metagenome dataset, with less than 0.01% of the metagenomic reads mapping to the two genomes (white bar). (D) Pyruvate fermentation (Phase I, white circles) and N2O consumption (Phase II, black circles) in co-culture EV. Serratia (blue diamonds) and Desulfosporosinus (red diamonds) cell numbers were determined with specific, 16S rRNA gene-targeted qPCR. (E) Amplicon sequencing illustrates the population shifts in co-culture EV following pyruvate consumption (day 7) and following  $N_2O$  consumption (day 18). Relative abundance of *Serratia* (blue bars) and Desulfosporosinus (red bars) in co-culture EV following Phase I (day 7) and Phase II (day 18) in vessels with pyruvate plus  $N_2O$  (left) and pyruvate only (right). Sequencing was performed on single representative cultures. All other data represent the averages of triplicate incubations and error bars represent standard deviations (n=3). Error bars are not shown if
- Figure 2-2. Interspecies cross-feeding supports low pH N<sub>2</sub>O reduction in co-culture EV. (A) Pyruvate fermentation (Phase I, light blue background) in vessels inoculated with axenic Serratia sp. strain MF and N<sub>2</sub>O consumption (Phase II, red background) following inoculation with coculture EV comprising the Serratia sp. and the N<sub>2</sub>O-reducing Desulfosporosinus sp. (indicated by the arrow). The bottom part of Panel A shows the amino acid profile in the supernatant immediately after inoculation with the Serratia sp. (green background), during Phase I, and during Phase II following inoculation with co-culture EV (day 7; 3% inoculum). Samples for untargeted metabolome analysis were collected immediately after inoculation with Serratia sp. (light green), during Phase I (light blue, pyruvate fermentation) and Phase II (light red, N<sub>2</sub>O reduction). The stacked bars show the ratio (%) of areas under the curve (AUC) of the respective amino acids and amino acid derivatives. Metabolites not assigned to structures representing amino acids or its derivatives are not shown. (B) Principal coordinate analysis (PCoA) of amino acid profiles. The enclosing ellipses were estimated using the Khachiyan algorithm with the ggforce package. Permanova analysis was conducted with the vegan Community Ecology package. Black, blue, and red circles represent samples collected at day 0, during Phase I, and during Phase II, respectively. (C) N<sub>2</sub>O consumption in co-culture EV in medium amended with mixtures comprising 5 (blue), 6 (orange), or 15 (black) amino acids (see SI for composition of mixtures), H<sub>2</sub>, and N<sub>2</sub>O. The *Desulfosporosinus* sp. cell numbers (red diamonds) were determined with 16S rRNA gene-targeted qPCR and show growth in medium receiving the 15-amino acid mixture. Various other amino acid mixtures tested resulted in no or negligible N<sub>2</sub>O consumption. All growth assays with amino acid mixture supplementation

- Figure 3-1. Novel microbial guilds implicated in N<sub>2</sub>O reduction. (A) Distribution of putative N<sub>2</sub>Oreducing population in transfer cultures with (+) or without (-) N<sub>2</sub>O amendment. The relative abundance (% of the total metagenome fragments) of MAGs was calculated using CoverM (https://github.com/wwood/CoverM). Taxonomy was assigned to MAGs using GTDB-TK (91). Displayed are MAGs predicted to have at least one nosZ gene based on Prokka annotation (137). The comprehensive microbial community profiled with the 43 high-quality MAGs are summarized in Fig. S4. (B). Phylogenomic analyses of nosZ-carrying MAGs recovered from low pH N<sub>2</sub>O-reducing cultures. The tree topology was obtained from the Maximum Likelihood (ML) tree by using RAxML-NG (92), inferred from a 120 conserved marker genes alignment of 20 complete genomes (selected based on GTDB-TK results) (91) and 7 MAGs. The top scale bar indicates 0.2 amino acid substitution rates per site. Sizes of open circles alongside the branches of each sublineage indicate bootstrap percentages obtained by RAxML-NG. The pairwise genome-wide AAI calculated for each MAGs are shown on the right, with the scale displayed on the bottom. Values for each comparison for a given MAGs are connected by lines, and the comparison to self is indicated as 100%. Pair-wise AAI values of 65-95% (i.e., data points fall between the dashed lines) are indicative of species boundary (179). The color legend corresponds to the bar and line colors in panel (A) and (B)......85
- Figure 3-2. Phylogenetic trees showing the placement of acidic cultures derived NosZ (green) among the reference NosZ (black) and putative NosZ (red). (A) Unrooted phylogenetic tree of NosZ showing three independent clades (i.e., I, II and III). The tree topology was inferred from RAxML-NG. Bootstrap values are displayed as white (> 90%), grey (70 < BP < 90) and black circles (< 70%). (B) Phylogenetic tree showing clade I NosZ with branches belonging to clade II and clade III collapsed. (C) Phylogenetic tree showing clade II NosZ with branches belonging to clade I and clade III collapsed. (D) Phylogenetic tree showing clade III NosZ with branches</p>

- Figure 3-3. Structural and functional validation of clade III NosZ. (A). The NosZ enzyme is a homodimer in head-to-tail orientation with the tetranuclear Cuz active site located in the Nterminal, seven-bladed β-propeller domain and the binuclear CuA site in the C-terminal cupredoxin domain. The Cu<sub>A</sub>-Cu<sub>Z</sub> distance within one monomer is around 42.5 Å (i.e., Cu<sub>A</sub>-A to Cuz-A) and thus too large for efficient electron transfer. Instead, the CuA site of one monomer is at a distance of only 10.5 Å from Cuz in the other monomer (CuA-A to Cuz-B) (B). Conserved residues associated with Cu<sub>A</sub> and Cu<sub>Z</sub> sites. Two histidine and cysteine residues are conserved in Cu<sub>A</sub> site, and seven histidine residues are conserved in Cu<sub>Z</sub> site. The Y163 residue enclosed in red is an unexpected (lacking in clade I and clade II NosZ) tyrosine associated with Cuz site. (C) Comparative growth assays performed with 27th Sabana and (D) 15th WS2 transfers cultures. Green and red lines indicate N2O reduction performance in different pH (4.5 and 7) medium. Green bars represent the cells (pH 4.5) enumerated with specific primers targeting the clade III Desulftobacterium nosZ. Fold changes of nosZ transcripts (orange bars) and assembled NosZ protein (yellow bars) are calculated based on corresponding cultures grown in pH 4.5 medium with (+) and without (-) N<sub>2</sub>O amendment. Of note, growth of clade III N<sub>2</sub>O reducers was not substantial in cultures without N<sub>2</sub>O. Data represent averages of triplicate incubations and error bars represent standard deviations (n=3). Error bars are not shown
- *aromatica* strain RCB (b) in the presence or absence of Na<sub>2</sub>S. N<sub>2</sub>O reduction in culture EV (c) and *D. aromatica* strain RCB (d) received either CuCl<sub>2</sub> or Na<sub>2</sub>S and cultures amended with excess CuCl<sub>2</sub> and Na<sub>2</sub>S. The data represent the averages of triplicate incubations and error bars represent the standard deviations (n=3). Error bars are not shown if smaller than the symbol.

- Supplementary Figure 2-1. Phenotypic characterization of co-culture EV. Cultures were grown in 160 mL glass serum bottles containing 100 mL of medium amended with pyruvate (50 or 250 µmol), H<sub>2</sub> (416 µmol) and N<sub>2</sub>O (416 µmol) (A and B), pyruvate and H<sub>2</sub> (C and D), or pyruvate and N<sub>2</sub>O (E and F). The arrows in panels D and F indicate additional doses of N<sub>2</sub>O or H<sub>2</sub> to corresponding culture vessels lack N<sub>2</sub>O (D) or H<sub>2</sub> (F) to resume N<sub>2</sub>O reduction activity. Pyruvate consumption and associated product formation are documented in panels A, C, and E. N<sub>2</sub>O and H<sub>2</sub> consumption are documented in panels B, D, F. Panel G shows the performance of co-culture EV grown with a lower amount of pyruvate. Panel H depicts pyruvate, H<sub>2</sub> and

- Supplementary Figure 2-3. Fragment recruitment plot of the 'Ca. Desulfosporosinus nitrousreducens' strain PR genome to the metagenome dataset derived from the 15th generation co-culture EV. The recruitment was performed via processing a BLAST search of the metagenome fragments against the 'Ca. Desulfosporosinus nitrousreducens' strain PR genome. The tabular BLAST result was parsed using BLASTab.catsbj.pl, and graphical representation was generated with the BlasTab.recplot2.R embedded in Enveomics collection (88). (A) The bar on the left shows the number of fragments with different identity recruited to each position on the genome. (B) indicates the sequencing depth across the 'Ca. Desulfosporosinus nitrousreducens' strain PR genome on a logarithmic scale. (C) Sequencing depth histogram with peaks from values above 95% identity automatically identified as skewed normal distribution. (D) Metagenome fragments recruited to the 'Ca. Desulfosporosinus nitrousreducens' strain PR genome, placed by location (x-axis) and identity (y-axis). (E) Identity histogram of mapping fragments (light gray) and smoothed spline (black). The backgrounds in panels D and E, and the line colors in panels B and C, correspond to identity matches above (dark blue) and below (light blue) 95 %. A recruitment plot of the 'Ca. Desulfosporosinus nitrousreducens' strain PR genome to the metagenome datasets derived from El Verde soil is not shown as the covered fraction of the 5.6 Mbp genome was below 10%

- Supplementary Figure 2-8. Gene clusters found on the '*Ca*. Desulfosporosinus nitrousreducens' strain PR genome. Displayed are formate dehydrogenase (*fdh* gene cluster), b-type Ni/Fe hydrogenase (*hyp* gene cluster), group 1 Ni/Fe hydrogenase (*hya* gene cluster), FeFe type Nitrogenase (*anf* gene cluster), Mo-Fe Nitrogenase (*nif* gene cluster), Nitric-oxide reductase (*nor* gene cluster), High-affinity amino acid transport system (*ilv* gene cluster). The arrows indicate operon length and orientations. Preliminary annotation of coding genes was done with Prokka (137), MicrobeAnnotator (138), and the RAST server (87), and genes of interests were curated via blasting to the NCBI nr database. Cy-b: b-type cytochrome; ABC: ATP-binding cassette. The black arrow in the *fdh* gene cluster represents a gene encoding a small protein of unknown function.
- Supplementary Figure 2-10. Distribution of 'Ca. Desulfosporosinus nitrousreducens' strain PR and Serratia sp. strain MF in select publicly available soil metagenomes. Metagenome datasets were downloaded from the European Nucleotide Archive (accession number PRJEB44414) with associated metadata available in a Supplemental Information/Source Data file (https://doi.org/10.1038/s41467-022-29161-3). (A) Geographical distribution of the soil metagenomes included in the analysis. Blue dots indicate a soil pH >7, red dots indicate a soil pH <7. (B) The partial importance (%) of different parameters, including the presence of Serratia sp. (based on short reads that align to the Serratia sp. strain MF genome), organic matter (OM content), NO<sub>3</sub> content, NH<sub>4</sub><sup>+</sup> content, C/N ratio, and pH for the relative abundance of 'Ca. Desulfosporosinus nitrousreducens' strain PR based on Random Forest models (139). (C) Correlation between percentages of short read fragments aligned to the Serratia sp. strain MF and the 'Ca. Desulfosporosinus nitrous reducens' strain PR genomes. The significance values for the correlations were calculated with the Pearson test (140) implemented in the ggpubr package (https://github.com/kassambara/ggpubr). .....54 Supplementary Figure 3-1. Low pH N<sub>2</sub>O reduction of transfer cultures derived from farm and tropical soils. Numbers inside parenthesis indicate the transfer generations. For example, panel labeled as WS1 (4th) showed N2O data collected from 4th WS1 transfer culture. Data were

- Supplementary Figure 3-7. AlfphaFold2 predicted structures of proteins located closely to NosZ. Signalp 6.0 was used to identify and remove signal peptides prior to structure prediction. The corresponding membrane orientations of proteins were predicted using MembraneFold. .......98

#### **Chapter 1 Introduction**

### 1.1 Sources and sinks of N<sub>2</sub>O

Nitrous oxide (N<sub>2</sub>O), an odorless gas, is the dominant ozone-depleting substance emitted in the 21<sup>st</sup> century (1). Soils and industrial processes have contributed considerably to the increased N<sub>2</sub>O emissions over the last decades (2, 3). Agricultural and natural soils are the major contributors of the current global annual N<sub>2</sub>O budget as high as 17.0 Tg N yr<sup>-1</sup> (3). However, the N<sub>2</sub>O uptake of soils was as low as 0.01 Tg N yr<sup>-1</sup>, which was much lower than that produced from soils (9.4 Tg N yr<sup>-1</sup> in total). The imbalance may be related to the multiple N<sub>2</sub>O sources (N<sub>2</sub>O-producing process) versus the single N<sub>2</sub>O sink (N<sub>2</sub>Oconsuming process) (4).

Coupled abiotic-biotic processes are the predominant N<sub>2</sub>O sources in terrestrial ecosystems (5). During the Great Oxidation Event (GOE), nitrite (NO<sub>2</sub><sup>-</sup>) and nitrate (NO<sub>3</sub><sup>-</sup>) were produced as a result of oxidization of inorganic nitrogen (6). In the Fe<sup>2+</sup>-rich environments, Fe minerals could have mediated the formation of nitric oxide (NO) and N<sub>2</sub>O via chemical reaction with NO<sub>2</sub><sup>-</sup> (7). In addition, the chemical process is also coupled to biotic process when Fe<sup>3+</sup> and NO<sub>3</sub><sup>-</sup> reduction occur simultaneously. For example, *Anaeromyxobacter dehalogenans* reduced both Fe<sup>3+</sup> and NO<sub>3</sub><sup>-</sup> under anoxic conditions, the resulting Fe<sup>2+</sup> and NO<sub>2</sub><sup>-</sup> subsequently react to form N<sub>2</sub>O chemically (4Fe<sup>2+</sup> + 2NO<sub>2</sub><sup>-</sup> + 5H<sub>2</sub>O  $\rightarrow$  2Fe<sup>3+</sup> + N<sub>2</sub>O + 6H<sup>+</sup>) (8). In this specific chemical reaction, Mn<sup>2+</sup> could substitute Fe<sup>2+</sup> as the oxidant (2Mn<sup>2+</sup> + 2NO<sub>2</sub><sup>-</sup> + 6H<sup>+</sup>  $\rightarrow$  2Mn<sup>4+</sup> + N<sub>2</sub>O + 3H<sub>2</sub>O) (9). These metal minerals could also react with hydroxylamine, an important intermediate product in nitrification, yielding N<sub>2</sub>O as the end product (4Fe<sup>3+</sup>/2Mn<sup>4+</sup> + 2NH<sub>2</sub>OH  $\rightarrow$ 4Fe<sup>2+</sup>/2Mn<sup>2+</sup> + N<sub>2</sub>O + H<sub>2</sub>O + 4H<sup>+</sup>) (10, 11). Thus, abiotic processes could have been the major contributor to N<sub>2</sub>O production when microbial processes yielding N<sub>2</sub>O were not available in the Archaean.

In modern terrestrial ecosystems, microbial transformations of nitrogen, coupled to abiotic processes, have become the major  $N_2O$  source (12). Following the start-up

reaction of nitrogen fixation, the resulting ammonia (NH<sub>4</sub><sup>+</sup>) become available to microorganisms and ready for microbial conversions. Microorganisms convert NH<sub>4</sub><sup>+</sup> to NO<sub>3</sub><sup>-</sup> via nitrification with NH<sub>2</sub>OH and NO<sub>2</sub><sup>-</sup> is formed as an intermediate product, which could further react with metal minerals to formulate N<sub>2</sub>O. In contrast, modular microbial denitrification processes produce N<sub>2</sub>O as the intermediate product from the reverse reaction (NO<sub>3</sub><sup>-</sup>  $\rightarrow$  NO<sub>2</sub><sup>-</sup>  $\rightarrow$  N<sub>2</sub>O  $\rightarrow$  N<sub>2</sub>) (4). Denitrifying organisms such as fungi that lack the gene coding NosZ would produce N<sub>2</sub>O as end product (13).

To date, transformation of N<sub>2</sub>O has been achieved with three enzymes (14), though two of them are not physiology relevant to N<sub>2</sub>O reduction. The nitrogenase was demonstrated to utilize N<sub>2</sub>O as alternative substrate, with the end-product being N<sub>2</sub> and NH<sub>3</sub> (N<sub>2</sub>O  $\rightarrow$ N<sub>2</sub>  $\rightarrow$  NH<sub>3</sub>). However, the conversion of N<sub>2</sub>O by nitrogenase was documented to be biologically irrelevant, as in vivo N<sub>2</sub>O reduction mainly occurred with NosZ (15). Another enzyme that putatively consumed N<sub>2</sub>O is a multicopper oxidase from the archaeon *Pyrobaculum aerophilum* while the end-product remains unknown (16). The most widely studied enzyme catalyzing N<sub>2</sub>O reduction is the NosZ comprising two copper centers (17), constituting the last step of microbial denitrification pathway (12).

 $N_2O$  reduction is associated with energy conservation, disposal of excess electrons, and detoxification in organisms expressing NosZ (18).  $N_2O$  respiring bacteria grow and build up biomass with  $N_2O$  as the sole electron acceptor (i.e., energy conservation) (19). In contrast, obligate aerobic organisms that only reproduce under oxic conditions utilize  $N_2O$  as a temporary substitute for oxygen, allowing them to survive under hypoxia conditions (20). For corrinoid-dependent organisms, consumption of  $N_2O$  serves as a detoxification mechanisms due to its inhibitory effect on corrinoid-dependent processes (21). Therefore, organisms have acquired  $N_2O$  reduction capacity in response to divergent environmental stress.

Study of NosZ has long been focused on denitrifying organisms until the novel NosZ (hereafter referred as clade II NosZ) was identified recently (12). The conventional NosZ (hereafter referred as clade I NosZ) is comprised entirely of *Proteobacteria* and *Archaea* 

and was distinguished with the Tat signal translocation pathway and likelihood of dentification (22). In contrast, a broader organismic distribution was observed for clade II NosZ (e.g., *Chloroflexi, Bacteroidetes* and *Firmicutes*) with a Sec signal peptide conserved in the genes. In addition to the gene diversification, the structures of Nos operon clusters comprising accessory genes of NosZ also differentiate clade I and II NosZ. For example, a membrane-bounding protein harboring at least four transmembrane helixes were exclusively identified immediately following clade II NosZ (12). Despite extensive efforts applied to investigate NosZ, the growing diversity suggests it is underexplored. For example, *nosZ* was conspicuously absent in the genomes of *Acidobacteria*, which suggested a lower frequency of *nosZ* in associated acidic conditions (23). However, a recent study implicated organisms affiliated with *Acidobacteria* were the predominant N<sub>2</sub>O reducers in tundra soil ecosystems (24).

#### **1.2** Acidic soils are the main N<sub>2</sub>O source

Global soil ecosystems are experiencing acidification due to intensive application of nitrogen fertilizer. Biotic reactions of nitrogen are the main cause of soil acidification. A notable example is the nitrification process generating acidity:  $NH_4^+ + 2O_2 \rightarrow 2H^+ +$  $NO_3^- + H_2O$  (25). In the case of a meta-analysis, soil pH has declined by 0.26 on average as a result of N fertilizer application over the past decades (26). Global excessive nitrogen use (27) would exacerbate soil acidification in the future and cause environmental problems (28) such as increasing net N<sub>2</sub>O emissions (29).

Soil pH is a chief modulating factor affecting the magnitude of N<sub>2</sub>O production and consumption, consequently controlling net N<sub>2</sub>O emissions (30). The mechanisms behind this are the organisms/enzymes involved in the N-cycling network have different pH preferences. For instance, acidification led to increased N<sub>2</sub>O production from both ammonia oxidizing microbes via nitrifier-denitrification process (NH<sub>4</sub><sup>+</sup>  $\rightarrow$  NO<sub>2</sub><sup>-</sup>  $\rightarrow$  N<sub>2</sub>O) (31). Furthermore, pH is a control of competing metabolisms within individual organisms: acidic and circumneutral pH favored *Shewanella loihica* strain PV-4 to carry out denitrification and respiratory ammonification (32), with the former conditions

yielding N<sub>2</sub>O as major product. Therefore, the effects of pH on N<sub>2</sub>O production are multidimensional at organismic level.

The physiological pH optima of enzymes are a determining factor controlling N<sub>2</sub>O production of individual organisms, consequently affecting the soil N<sub>2</sub>O emissions. In contrast to the circumneutral pH (~7.3) optima of N<sub>2</sub>O consumption, the preferred pH of N<sub>2</sub>O production was reported to be more acidic (pH 5 to 6) (30). Thus, acidic soils are hypothesized to be the main N<sub>2</sub>O emitters. However, laboratory incubations of acidic solids-containing microcosms arguing the hypothesis showed N<sub>2</sub>O reduction activity (33, 34). The observed N<sub>2</sub>O reduction activity in acidic microcosms may occur in the microsites associated with solids (i.e., soils) where the pH is different from bulk medium (35). For example, a *Cloacibacterium* isolate greatly reduced net N<sub>2</sub>O emission in acidic solids-containing microcosm (pH 5.5) but showed no N<sub>2</sub>O reduction activity in pH 5.5 liquid medium (36). The observed N<sub>2</sub>O sink effect in acidic microcosms could be that *Cloacibacterium* cells were embedded in solid phase, protected against acidic bulk medium pH.

In contrast to the microcosm-based observation, solid-free laboratory incubations and field measurements of N<sub>2</sub>O showed weak or no N<sub>2</sub>O sink capacity under acidic conditions compared to that under circumneutral pH conditions. For example, net N<sub>2</sub>O emissions were negatively correlated to soil pH (37). Furthermore, the N<sub>2</sub>O emission factors were higher in acidic soils than that in circumneutral pH soils, indicating more N fertilizer applied to acidic soils were lost in the form of N<sub>2</sub>O. Phenotypic studies of clade I N<sub>2</sub>O reducers showed that N<sub>2</sub>O reduction was completely blocked at pH values lower than 6 (38, 39), suggesting the weak N<sub>2</sub>O reduction capacity is the main cause of higher N<sub>2</sub>O emissions from acidic soils. Transcriptional analysis and resting cell assays documented that NosZ assembly is impaired under acidic conditions, while the expression of *nosZ* genes was not affected (40, 41). Consistent to these observations, mechanistic study unraveled a conformational change of clade I NosZ under acidic conditions, which explains its incapability in consuming N<sub>2</sub>O (42). However, the impact

of acidic pH on clade II NosZ remains unclear though extensive studies have been conducted to explore clade I NosZ at organismic, genetic, enzymatic level.

### **1.3 Research objectives**

The proposed work will elucidate the microbiology of low  $pH N_2O$  reduction and address the following hypotheses.

**Hypothesis 1:** N<sub>2</sub>O-reducing microbes in soils have adapted to acidic pH conditions (<5) and can be enriched from acidic soils.

**Rationale**: A metagenomic study focused on Luquillo Experimental Forest soils identified diverse *nosZ* gene sequences, suggesting genomic potential for N<sub>2</sub>O reduction in low pH soils (43). A subsequent microcosm study demonstrated N<sub>2</sub>O reduction activity at a pH of 4.5 (44). These microcosms are available for studying the microbiology involved in low pH N<sub>2</sub>O reduction.

**Hypothesis 2:** Acidophilic N<sub>2</sub>O-reducing microbes also exist in acidic microsites in soils with circumneutral pH values.

**Rationale**: The naturally acidic sites on the Earth are known to thrive a variety of acidophilic microorganisms (45), whereas the circumneutral soil are overlooked source of acidophiles. Soils are complex matrixes with high pH heterogeneity, and pH varies by as much as 1.3 pH among samples less than 50 cm apart (46). The microbial metabolism could have led to more contrasting pH at finer scale: the microscale varied by 2 pH in spots localized closely (less than 5 mm) following straw patch amendment (47). Obviously, acidic microsites could occur in circumneutral CFF soils and harbor the acidophilic N<sub>2</sub>O-reducing microorganisms.

Hypothesis 3: Geochemical factors are controlling N<sub>2</sub>O reduction in acidic soils.

**Rationale**: Growth of most acidophilic N<sub>2</sub>O-reducing microbes acquired in this study requires amendment of complex substrates (e.g., amino acids and peptides). In microbial

network, the reliance on external growth factors would render the acidophilic  $N_2O$ -reducing microbes weak competitors for nutrients and energy.

# Chapter 2 Sustained bacterial N<sub>2</sub>O reduction at acidic pH

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#### Abstract

Nitrous oxide (N<sub>2</sub>O) is a climate-active gas and emissions from terrestrial ecosystems are concerning. Microbial reduction of N<sub>2</sub>O to dinitrogen (N<sub>2</sub>) is the only known consumption process and has been studied extensively at circumneutral pH; however, N<sub>2</sub>O reduction under acidic conditions is thought to be limited. Global soil acidification, accelerated by anthropogenic practices, introduces high uncertainty into N<sub>2</sub>O emission budgets. We obtained an enrichment culture from an acidic tropical forest soil that robustly reduces N<sub>2</sub>O to N<sub>2</sub> at pH 4.5 with the addition of pyruvate and hydrogen. Consecutive transfers at pH 4.5 yielded a co-culture and temporal analyses revealed a bimodal growth pattern with a Serratia sp. growing during the initial pyruvate fermentation phase followed by growth of a novel Desulfosporosinus sp. via hydrogenotrophic N<sub>2</sub>O reduction. The *Desulfosporosinus* sp. produced  $(3.1 \pm 0.11) \times 10^8$ cells per mmol of N<sub>2</sub>O consumed, on par with growth yields reported for clade II N<sub>2</sub>O reducers at circumneutral pH. Genome analysis identified a clade II nos gene cluster, but an incomplete pathway for sulfate reduction, a hallmark feature of the genus Desulfosporosinus. Physiological and metabogenomic characterization revealed interspecies nutritional interactions, with the pyruvate fermenting Serratia sp. supplying amino acids as essential growth factors to the *Desulfosporosinus* sp. The co-culture reduced  $N_2O$  between pH 4.5 and 6 but not at or above pH 7, contradicting the paradigm that sustained microbial  $N_2O$  reduction ceases under acidic pH conditions, thus confirming a previously unrecognized N<sub>2</sub>O reduction potential in acidic soils.

#### 2.1 Introduction

Exploitation, mismanagement, and unsustainable practices deteriorate soil health across the globe (48). pH is a key soil health parameter, but soil acidification, a natural process accelerated by the reliance of synthetic nitrogen fertilizer, the growth of legumes, and acidic precipitation/deposition, plagues regions around the world (49, 50). In addition to pH change, N fertilizer input substantially increases the formation of N<sub>2</sub>O, a compound linked to ozone depletion and climate warming (1, 51-54), and interference with biogeochemical processes such as methanogenesis, mercury methylation, and reductive dechlorination (55, 56). The rise in global N<sub>2</sub>O emissions indicates an imbalance between  $N_2O$  formation versus consumption, which has been explained by the contrasting pH optima of processes involved in N<sub>2</sub>O formation (pH 5-6) versus consumption (circumneutral pH) (30). A few studies reported N<sub>2</sub>O consumption in acidic soil laboratory microcosms (57, 58); however, soil heterogeneity and associated (microscale) patchiness of pH conditions make generalized conclusions untenable (35, 36). Attempts with denitrifying enrichment and axenic cultures have thus far failed to demonstrate growth-linked N<sub>2</sub>O reduction and sustainability of such a process under acidic pH conditions (34, 36, 59).

The only known sink for N<sub>2</sub>O are microorganisms expressing N<sub>2</sub>O reductase (NosZ), a periplasmic enzyme that catalyzes the conversion of N<sub>2</sub>O to environmentally benign dinitrogen (N<sub>2</sub>). NosZ expression and proteomics studies with the model denitrifier *Paracoccus denitrificans* demonstrated that acidic pH interferes with NosZ maturation (38, 60), a phenomenon also observed in enrichment cultures harboring diverse N<sub>2</sub>O-reducing bacteria (40). Studies with *Marinobacter hydrocarbonoclasticus* found active NosZ with a Cu<sub>Z</sub> center in the 4Cu<sub>2</sub>S form in cells grown at pH 7.5, but observed a catalytically inactive NosZ with the Cu<sub>Z</sub> center in the form 4Cu<sub>1</sub>S when the bacterium was grown at pH 6.5 (42). These observations support the paradigm that microbial N<sub>2</sub>O reduction is limited to circumneutral pH environments. A metagenome-based analysis of soil microbial communities in the Luquillo Experimental Forest (Puerto Rico) provided evidence that N<sub>2</sub>O-reducing soil microorganisms are not limited to circumneutral pH

soils and exist in acidic tropical forest soils (43). Microcosms established with acidic Luquillo Experimental Forest soil demonstrated the N<sub>2</sub>O reduction activity, and suggested that the microbial communities in pH 4.5 and pH 7 microcosms differed (44). To shed light on the microorganisms performing N<sub>2</sub>O reduction at pH 4.5, a two-population co-culture was derived from N<sub>2</sub>O-reducing soil microcosms established with acidic tropical forest soil collected in the Luquillo Experimental Forest (El Yunque National Forest) in Puerto Rico. Characterization of the co-culture demonstrated the activity of an acidophilic *Desulfosporosinus* sp. that utilizes N<sub>2</sub>O as electron acceptor at pH 4.5.

#### 2.2 Materials and methods

#### Soil sampling locations and microcosms

Soil samples were collected in August 2018 at the El Verde research station in the El Yunque Natural Forest in Puerto Rico (43). The measured soil pH was 4.45 and characteristic for the region. Vertical distance of the El Verde research station to mean sea level is 434 meters. Fresh soil materials from 9 to 18 cm depth were used to establish pH 4.5 laboratory microcosm that were amended with N<sub>2</sub>O and lactate (44).

#### **Enrichment cultures**

Transfer cultures were established in 160-mL glass serum bottles containing 100 mL of anoxic, completely synthetic, defined basal salt medium prepared as described (61) with the following modifications. NaHCO<sub>3</sub> and Na<sub>2</sub>S were omitted and the total KH<sub>2</sub>PO<sub>4</sub> concentration was increased to 7.0 g L<sup>-1</sup>. l-cysteine (0.2 mM) was the sole reductant unless indicated otherwise. The medium pH was 4.27 to 4.35. The bottles with N<sub>2</sub> headspace were sealed with butyl rubber stoppers (Bellco Glass, Vineland, NJ, USA) held in place with aluminum crimp caps and autoclaved. All subsequent amendments to the medium bottles used sterile plastic syringes and needles to augment the medium with aqueous filter-sterilized (0.2  $\mu$ m polyethersulfone membrane filter, Thermo Scientific, Waltham, MA, USA) stock solutions and undiluted gases (62). Ten mL of N<sub>2</sub>O gas (416  $\mu$ mol, 4.16 mM nominal; 99.5%) was added 24 hours prior to inoculation from an El Verde microcosm showing N<sub>2</sub>O reduction activity. The microcosm was manually shaken before 1 mL aliquots were transferred with a 3-mL plastic syringe and a 2-gauge needle. Initial attempts to obtain solids-free enrichment cultures with 5 mM lactate as carbon source and electron donor showed no N<sub>2</sub>O reduction. As such, the following substrates were subsequently tested in the transfer cultures: 5 mM propionate, 20 mM pyruvate, 20 mM pyruvate plus 10 mL (416  $\mu$ mol, 4.16 mM nominal) hydrogen (H<sub>2</sub>), 1 mM formate plus 1 mM acetate and 5 mL (208  $\mu$ mol, 2.08 mM nominal) CO<sub>2</sub>, and 0.1 or 10 g L<sup>-1</sup> yeast extract. Subsequent transfers (3%, v/v) used medium supplemented with 0.5 or 2.5 mM pyruvate and 10 mL H<sub>2</sub>, and occurred when the initial dose of 10 mL N<sub>2</sub>O had been consumed. All culture vessels were incubated in upright position at 30°C in the dark without agitation.

## Microbial community analysis

16S rRNA gene amplicon sequencing was performed with samples collected from  $6^{th}$ generation transfer culture following complete N<sub>2</sub>O consumption, and 9<sup>th</sup>-generation transfer cultures following complete pyruvate consumption (Phase I) and complete N<sub>2</sub>O consumption (Phase II). Biomass from 1 mL of culture suspension samples were collected by centrifugation (10,000 x g, 20 min, 4 °C), and genomic DNA was isolated from the cell pellets using the DNeasy PowerSoil Kit (Qiagen, Hilden, Germany). 16S rRNA gene-based amplicon sequencing was conducted at the University of Tennessee Genomics Core following published procedures (63). Primer sets used in amplicon sequencing of 6<sup>th</sup> and 9<sup>th</sup> transfer cultures are 341F-785R and 515F-805R, respectively (64).

Analysis of amplicon reads was conducted with nf-core/ampliseq v2.3.1 using Nextflow (65). Software used in nf-core/ampliseq was containerized with Singularity v3.8.6 (66). Amplicon read quality was evaluated with FastQC v0.11.9 (67) and primer removal used

Cutadapt v3.4 (68). Quality control including removal of sequences with poor quality, denoising, and chimera removal was performed, and amplicon sequence variants (ASVs) were inferred using DADA2 (69). Barrnap v0.9 was used to discriminate rRNA sequences as potential contamination (70). ASVs were taxonomically classified based on the Silva v138.1 database (71). Relative and absolute abundances of ASVs were calculated using Qiime2 v2021.8.0 (72). Short read fragments of the El Verde soil metagenome representing 16S rRNA genes were identified and extracted using Parallel-Meta Suite v3.7 (73).

#### **Isolation and classfication**

Following 15 consecutive transfers, 100  $\mu$ L cell suspension aliquots were serially diluted and plated on tryptic soy agar (TSA, MilliporeSigma, Maryland, US) medium. Colonies with uniform morphology were observed, and a single colony was transferred to a new TSA plate. This process was repeated three times before a single colony was transferred to liquid basal salt medium (pH 4.5) amended with 2.5 mM pyruvate, 416  $\mu$ mol N<sub>2</sub>O, and 416  $\mu$ mol H<sub>2</sub>. Following growth, DNA was extracted for PCR amplification with general bacterial 16S rRNA gene-targeted primer pair 8F-1541R (74), and Sanger sequencing of both strands yielded a 1,471-bp long 16S rRNA gene fragment.

Efforts to isolate the N<sub>2</sub>O reducer applied the dilution-to-extinction principle (61). Tenfold dilution-to-extinction series used 20 mL glass vials containing 9 mL of basal salt medium and 0.6% (w/v) low melting agarose (MP Biomedicals, LLC., Solon, OH) with a gelling temperature below 30° were established as described (61). Each glass vial received 2.5 mM pyruvate, 1 mL (41.6  $\mu$ mol, 4.16 mM nominal) H<sub>2</sub> and 1 mL (41.6  $\mu$ mol, 4.16 mM nominal) N<sub>2</sub>O following heat sterilization. Parallel 10<sup>-1</sup> to 10<sup>-10</sup> dilutionto-extinction series were established in liquid basal salt medium without low melting agarose, which were used to inoculate the respective soft agar dilution vials. Additional attempts to isolate the N<sub>2</sub>O reducer used solidified (1.5% agar, w/v) basal salt medium. A 1-mL sample of a 15<sup>th</sup>-generation transfer culture that actively reduced N<sub>2</sub>O was 10-fold serially diluted in liquid basal salt medium, and 100  $\mu$ L of cell suspension aliquots were evenly distributed on the agar surface. The plates were incubated under an atmosphere of  $N_2/H_2/N_2O$  (8/1/1, v/v/v), and colony formation was monitored every 2 weeks over a 6-month period.

#### Quantitative PCR (qPCR)

SYBR Green qPCR assays targeting the 16S rRNA gene of the Serratia sp., and TaqMan qPCR assays targeting the 16S rRNA gene of the Desulfosporosinus sp. were designed using Geneious Prime (Table 2-1). Probe and primer specificities were examined by in *silico* analysis using the Primer-BLAST tool (75), and experimentally confirmed using 1,538 bp- and 1,467 bp-long synthesized linear DNA of the respective complete 16S rRNA genes of the Serratia sp. and the Desulfosporosinus sp., respectively (Integrated DNA Technologies, USA). For enumeration of Serratia 16S rRNA genes, 25 µL qPCR tubes received 10 µL 1X Power SYBR Green, 9.88 µL UltraPure nuclease-free water (Invitrogen, Carlsbad, CA, USA), 300 nM of each primer, and 2 µL template DNA. For enumeration of *Desulfosporosinus* 16S rRNA genes, the qPCR tubes received 10 µL TaqMan Universal PCR Master Mix (Life Technologies, Carlsbad, CA, USA), 300 nM of TaqMan probe (5'-6FAM-AAGCTGTGAAGTGGAGCCAATC-MGB-3'), 300 nM of each primer, and 2 µL template DNA (76). All qPCR assays were performed using an Applied Biosystems ViiA 7 system (Applied Biosystems, Waltham, MA, USA) with the following amplification conditions: 2 min at 50°C and 10 min at 95°C, followed by 40 cycles of 15 s at 95°C and 1 min at 60°C. The standard curves were generated using 10fold serial dilutions of the linear DNA fragments carrying a complete sequence of the Serratia sp. (1,538 bp) or the Desulfosporosinus sp. (1,467 bp) 16S rRNA gene, covering the 70- and 72-bp qPCR target regions, respectively.

The qPCR standard curves established with the linear DNA fragments carrying complete *Serratia* sp. or *Desulfosporosinus* sp. 16S rRNA genes had slopes of -3.82 and -3.404, y-intercepts of 37.408 and 34.181,  $R^2$  values of 0.999 and 1, and qPCR amplification efficiencies of 82.7% and 96.7%, respectively. The linear range spanned 1.09 to 1.09 x  $10^8$  gene copies per reaction with a calculated detection limit of 10.9 gene copies per

reaction. The genome analysis indicated that both the *Serratia* sp. and the *Desulfosporosinus* sp. genomes carry a single 16S rRNA gene, indicating that the enumeration of 16S rRNA gene allows estimates of cell abundances. The 16S rRNA gene sequences of the *Serratia* sp. and the *Desulfosporosinus* sp. are available under NCBI accession numbers OR076433 and OR076434, respectively.

### Nutritional interactions in the co-culture

To explore the nutritional requirements of the Desulfosporosinus sp., a time series metabolome analysis of culture supernatant was conducted. Briefly, the axenic Serratia sp. culture was grown in basal salt medium amended with 2.5 mM pyruvate, 4.16 mM (nominal) H<sub>2</sub>, and 4.16 mM (nominal) N<sub>2</sub>O. Following a 7-day incubation period, during which pyruvate was completely consumed, the bottles received 1% (v/v) co-culture EV inoculum. Cell suspension aliquots (1.5 mL) were collected and centrifuged, and the resulting cell-free supernatants were transferred to 2 mL plastic tubes and immediately stored at -80 °C for metabolome analysis. Additional samples assessed the metabolome associated with supernatant of axenic Serratia sp. cultures that received 1 mM DTT instead of l-cysteine as reductant. The results of the metabolome analysis guided additional cultivation experiments with amino acid mixtures replacing pyruvate. The 100fold concentrated, aqueous 15-amino acid stock solution contained (g  $L^{-1}$ ): alanine (0.5); aspartic acid (1); proline (1); tyrosine (0.3); histidine (0.3); tryptophan (0.2); arginine (0.5); isoleucine (0.5); methionine (0.4); glycine (0.3); threonine (0.5); valine (0.9); lysine (1); glutamate (1); serine (0.8). The aqueous stock solution was filter-sterilized and stored in the dark.

#### Metagenome sequencing

DNA was isolated from the axenic *Serratia* sp. culture grown with 2.5 mM pyruvate, and the N<sub>2</sub>O-reducing 15<sup>th</sup> generation co-culture EV grown on H<sub>2</sub>, N<sub>2</sub>O, and the amino acid mixture. Metagenome sequencing was performed at the University of Tennessee Genomics Core using the Illumina NovaSeq 6000 platform. Shotgun sequencing

generated a total of 494 and 387 Gbp of raw sequences from the axenic *Serratia* sp. culture and co-culture EV. Metagenomic short reads were processed using the nfcore/mag pipeline (77). Short read quality was evaluated with FastQC v0.11.9, followed by quality filtering and Illumina adapter removal using fastp v0.20.1 (78). Short reads mapped to the PhiX genome (GCA\_002596845.1, ASM259684v1) with Bowtie2 v2.4.2 were removed (79). Assembly of processed short reads used Megahit2 v1.2.9 (80). Binning of assembled contigs was conducted with MetaBAT2 v2.15 based on the sequencing depth (81), and metagenome-assembled genomes (MAGs) that passed CheckM (82) were selected for further analysis. Protein-coding sequences on MAGs were predicted using MetaGeneMark-2 (83) and functional annotation used Blastp (84) against the Swiss-Prot database (85), KEGG (86) and the RAST server (87). Amino acid biosynthesis completeness was evaluated using KofamKOALA (86).

Metagenomic datasets of El Verde soil and a 15<sup>th</sup> transfer culture were searched against the '*Ca*. Desulfosporosinus nitrousreducens' strain PR MAG using blastn (84). The best hits were extracted using an in-house script embedded in Enveomics Collection tools (88). Graphical representation showing short reads recruited to the '*Ca*. Desulfosporosinus nitrousreducens' strain PR MAG was generated with BlasTab.recplot2.R. The coverage evenness was assessed based on distribution of high nucleotide identity reads across the reference genome sequences. Nonpareil v3.4.1 using the weighted NL2SOL algorithm was used to estimate the average coverage level of the metagenomic datasets (89). Metagenome data of the original El Verde soil was downloaded from NCBI (accession number PRJEB26500). Metagenomic datasets of coculture EV and axenic *Serratia* isolate were deposited at NCBI under accession numbers SRR24709127 and SRR24709126, respectively (Table 2-2).

#### Comparative analysis of nos gene clusters

Available genomes of select N<sub>2</sub>O reducers were downloaded from NCBI (Table 2-3). Functional annotation of the genomes was conducted using the RAST server. Transmembrane topology of the protein encoded by *nosB*, a gene located immediately

adjacent to *nosZ* was confirmed using DeepTMHMM (90). Accessory genes associated with NosZ functionality were selected from a set of seven genomes and used as PSI-BLAST database to query accessory genes on contigs predicted to harbor *nosZ* on the '*Ca*. Desulfosporosinus nitrousreducens' strain PR genome. The *nos* gene clusters were visualized using the gggenes package (<u>https://wilkox.org/gggenes/index.html</u>).

## Phylogenomic analysis

Phylogenomic reconstruction was performed with genomes of the *Desulfitobacteriaceae* family available in the NCBI database (Table 2-4). Conserved marker genes of the 20 genomes were identified and aligned with GTDB-TK (91). Phylogenetic relationships were inferred based on the alignment of 120 concatenated bacterial marker genes using RAxML-NG (92) with 1,000 bootstrap replicates. A best fit evolutionary model was selected based on the result of Modeltest-NG (93). Calculation of average amino acid identity (AAI) and hierarchical clustering of taxa based on AAI values were conducted with EzAAI (94). Tree annotation and visualization were performed with the ggtree package (95).

## NosZ phylogenetic analysis

NosZ reference sequences were downloaded from pre-compiled models of ROCker (96). The NosZ sequence of 'Ca. Desulfosporosinus nitrousreducens' strain PR was aligned to the NosZ reference sequences using MAFFT (97), and a maximum likelihood tree was created with RAxML-NG based on the best model from Modeltest-NG. The inferred tree and amino acid identity between '*Ca*. Desulfosporosinus nitrousreducens' strain PR, *Desulfosporosinus meridiei* and the NosZ reference sequences were visualized using ggtree package.

#### Metabolome analysis

Cell-free samples were prepared as described (98). Briefly, 1.5 mL of 0.1 M formic acid in 4:4:2 (v:v:v) acetonitrile:water:methanol was added to 100 µL aliquots of supernatant

samples. The tubes were shaken at 4°C for 20 minutes and centrifuged at 16,200 x g for 5 minutes. The supernatant was collected and dried under a steady stream of N<sub>2</sub>. The dried extracts were suspended in 300  $\mu$ L of water prior to analysis. For water soluble metabolites, the mass analysis was performed in untargeted mode (99). The chromatographic separations utilized a Synergi 2.6  $\mu$ m Hydro RP column (100 Å, 100 mm x 2.1 mm; Phenomenex, Torrance, CA) with tributylamine as an ion pairing reagent, an UltiMate 3000 binary pump (Thermo Scientific, San Jose, CA), and previously described elution conditions (98). The mass analysis was carried out using an Exactive Plus Orbitrap MS (Thermo Scientific) using negative electrospray ionization and full scan mode. Following the analysis, metabolites were identified using exact masses and retention times, and the areas under the curves (AUC) for each chromatographic peak were integrated using the open-source software package Metabolomic Analysis and Visualization Engine (99, 100). Dynamic changes of metabolites over time were assessed by comparative analysis of AUC values.

### Phenotypic characterization of co-culture EV

To test for autotrophic growth of co-culture EV, pyruvate was replaced by 5 mL (2.08 mM nominal) of CO<sub>2</sub> (99.5% purity). All experiments used triplicate cultures, and vessels without pyruvate, without H<sub>2</sub>, without N<sub>2</sub>O, or without inoculum served as controls. Growth experiments were conducted to determine the responses of the *Serratia* sp. and the *Desulfosporosinus* sp. to pH. Desired medium pH values of 3.5, 4.5, 5, 6, 7 and 8 were achieved by adjusting the mixing ratios of KH<sub>2</sub>PO<sub>4</sub> and K<sub>2</sub>HPO<sub>4</sub>. To achieve pH 3.5, the pH 4.5 medium was adjusted with 5 M hydrochloric acid. The medium with different pH, 3.5, 4.5, 5, 6, 7 and 8, was achieved by adjusting the mix ratios of KH<sub>2</sub>PO<sub>4</sub> and K<sub>2</sub>HPO<sub>4</sub>. Replicate incubation vessels received 4.16 mM (nominal) N<sub>2</sub>O and 4.16 mM H<sub>2</sub> (nominal), and 2.5 mM pyruvate, following an overnight equilibration period, 1% inocula from the axenic *Serratia* sp. culture or the N<sub>2</sub>O-reducing co-culture EV, both pregrown in pH 4.5 medium. The replicate cultures inoculated with the *Serratia* sp. were incubated for 14-days, after which three vessels received an inoculum of co-culture EV (1%) to initiate N<sub>2</sub>O consumption. Three *Serratia* sp. cultures not receiving a co-culture

EV inoculum served as controls. Consumption rates of pyruvate and N<sub>2</sub>O were calculated based on data points representing linear ranges of consumption according to

$$V = \frac{N}{T_1 - T_0}$$

Equation 1

where V represent the consumption rate; N represent the initial amounts of pyruvate or  $N_2O$ .  $T_1$  refers to timepoints when pyruvate or  $N_2O$  were completely consumed.  $T_0$  for pyruvate consumption refers to day zero (i.e., after inoculation with axenic *Serratia* sp.).  $T_0$  for  $N_2O$  consumption refers to day 14 (i.e., after inoculation with co-culture EV), following which a linear  $N_2O$  consumption was observed.

### **Analytical procedures**

N<sub>2</sub>O, CO<sub>2</sub>, and H<sub>2</sub> were analyzed by manually injecting 100  $\mu$ L headspace samples manually injected into an Agilent 3000A Micro-Gas Chromatograph (Palo Alto, CA, USA) equipped with Plot Q and molecular sieve columns coupled with a thermal conductivity detector as described (101). Aqueous concentrations ( $\mu$ M) were calculated from the headspace partial pressures based on reported dimensionless Henry's law constants for N<sub>2</sub>O (2.4×10<sup>-4</sup>), H<sub>2</sub> (7.8×10<sup>-6</sup>) and CO<sub>2</sub> (3.3×10<sup>-4</sup>) (102) according to

$$HRT = \frac{c_g}{c_{aq}}$$
 Equation 2

Where H = Henry's law constants, R = universal gas constant, T = temperature,  $C_g$  = headspace gas-phase concentration and  $C_{aq}$  = liquid-phase (dissolved) concentration. Five-point standard curves for N<sub>2</sub>O, CO<sub>2</sub> and H<sub>2</sub> spanned concentration ranges of 8,333 to 133,333 ppmv. Pyruvate, acetate and formate were measured with an Agilent 1200 Series high-performance liquid chromatography (HPLC) system (Palo Alto, CA, USA) as described (101). pH was measured in 0.4 mL samples of culture supernatant following removal of cells by centrifugation with a pH electrode.

#### 2.3 Results

#### A consortium consisting of two populations reduces N<sub>2</sub>O at pH 4.5

Microcosms established with El Verde tropical soil amended with lactate consumed N<sub>2</sub>O at pH 4.5; however, this activity was lost in transfer cultures amended with the same electron donor. The addition of acetate, formate (1 or 5 mM each), and CO<sub>2</sub> (208  $\mu$ mol, 2.08 mM nominal), propionate (5 mM), or yeast extract (0.10 – 10 g L<sup>-1</sup>) did not stimulate N<sub>2</sub>O reduction in pH 4.5 transfer cultures. Limited N<sub>2</sub>O consumption was observed in transfer cultures amended with 2.5 mM pyruvate, but complete removal of N<sub>2</sub>O required the addition of H<sub>2</sub> or formate. In transfer cultures with H<sub>2</sub> or formate, but lacking pyruvate, N<sub>2</sub>O was not consumed. Subsequent transfers in completely synthetic basal salt medium amended with both pyruvate and H<sub>2</sub> yielded a robust enrichment culture, which consumes N<sub>2</sub>O at pH 4.5 (Figure 2-1 A). Phenotypic characterization illustrated that pyruvate utilization was independent of N<sub>2</sub>O, while N<sub>2</sub>O reduction only commenced following complete consumption of pyruvate.

Cultivation experiments were performed in 160 mL glass serum bottles containing 100 mL of defined basal salt medium. The consortium consumed  $239 \pm 1.16 \mu$ mol pyruvate within 7 days (Phase I) in the presence or absence of N<sub>2</sub>O, and acetate ( $136 \pm 6.05 \mu$ mol), formate ( $58.5 \pm 3.33 \mu$ mol) and CO<sub>2</sub> ( $201 \pm 5.53 \mu$ mol) were produced (Supplementary Figure 2-1 A). In cultures without N<sub>2</sub>O, formate was stable but was readily consumed when cultures were supplemented with N<sub>2</sub>O (Supplementary Figure 2-1 A, C, E). Measurable acetate consumption did not occur in any of the vessels indicating that acetate is not an electron donor for N<sub>2</sub>O reduction (Supplementary Figure 2-1 A, C, E).

In vessels that received pyruvate,  $H_2$ , and  $N_2O$ ,  $H_2$  and  $N_2O$  were simultaneously consumed following the depletion of pyruvate (day 10, Supplementary Figure 2-1 B).  $N_2O$  consumption (Phase II) continued until  $H_2$  became limiting but resumed when additional  $H_2$  was provided (day 33, Supplementary Figure 2-1 B). In cultures lacking  $N_2O$ ,  $H_2$  was stable, but consumption commenced immediately following the addition of

 $N_2O$  (Supplementary Figure 2-1 D). In replicate vessels without  $H_2$ ,  $N_2O$  was consumed after day 10, apparently coupled to the oxidation of formate, a product of pyruvate fermentation (Supplementary Figure 2-1 E, F). Pyruvate consumption (Phase I) and the production of formate, acetate and CO<sub>2</sub> as fermentation products preceded measurable consumption of N<sub>2</sub>O and H<sub>2</sub> (or formate) (Phase II), suggesting that pyruvate was not a direct electron donor for  $N_2O$  reduction (Supplementary Figure 2-1 A-F). Following an 18-day incubation period, triplicate cultures that received pyruvate ( $239 \pm 1.16 \mu mol$ ), H<sub>2</sub>  $(390 \pm 1.66 \,\mu\text{mol})$ , and N<sub>2</sub>O  $(378 \pm 11.7 \,\mu\text{mol})$  had completely consumed N<sub>2</sub>O and nearly half of the exogenously added H<sub>2</sub> (143  $\pm$  8.96  $\mu$ mol) (Supplementary Figure 2-1 B). Reduction of one molecule N<sub>2</sub>O requires two electrons derived from the oxidation of one molecule of H<sub>2</sub> (i.e., H<sub>2</sub> + N<sub>2</sub>O  $\rightarrow$  H<sub>2</sub>O + N<sub>2</sub>). The amount of H<sub>2</sub> oxidized did not reach the expected 1:1 stoichiometry, indicating that products from pyruvate fermentation (e.g., formate) serve as electron donors for  $N_2O$  reduction. Consistently, in cultures that received a low amount of pyruvate (i.e., 50  $\mu$ mol), H<sub>2</sub> oxidation (361.2 ± 5.68  $\mu$ mol) closely matched the amount of N<sub>2</sub>O reduced ( $377 \pm 3.53 \mu$ mol) (Supplementary Figure 2-1 G). Pyruvate, H<sub>2</sub>, and N<sub>2</sub>O consumption were not apparent in vessels without inoculum (Supplementary Figure 2-1 H). No growth occurred in cultures where  $CO_2$ replaced pyruvate, suggesting no autotrophic activity (Supplementary Figure 2-1 I).

The fermentation of pyruvate resulted in pH increases, with the magnitude of the medium pH change proportional to the initial pyruvate concentration. The amount of pyruvate did not exceed 250  $\mu$ mol per 100 mL of medium (i.e., 2.5 mM), which limited the pH increase following complete pyruvate fermentation to 0.53  $\pm$  0.03 pH units (Supplementary Figure 2-2). N<sub>2</sub>O reduction was also observed in cultures that received 5 mM glucose. N<sub>2</sub>O reduction was oxygen sensitive and N<sub>2</sub>O was not consumed in medium without reductant (i.e., cysteine, dithiothreitol).

Microbial profiling of El Verde soil and solids-free transfer cultures documented effective enrichment in defined pH 4.5 medium amended with pyruvate, H<sub>2</sub>, and N<sub>2</sub>O (Figure 2-1 B). Shotgun metagenome sequencing performed on the original El Verde soil recovered 2,718 short read fragments (150 bp) representing 16S rRNA genes, which

could be assigned to 187 bacterial genera (43). 16S rRNA gene fragments representing uncultured microbes, or not assigned to known genera in the El Verde soil sequence pools, accounted for 33.1% and 27.3% of the total 16S rRNA gene fragments, respectively. The remaining 16S rRNA gene fragments were assigned to *Acidothermus* (3.1%), '*Ca.* Solibacter' (2.0%) and '*Ca.* Udaeobacter' (7.4%). Taxa with less than 2% representation in the sequence pools are grouped as 'Others' and accounted for 27% of the 16S rRNA gene fragments. Following six consecutive transfers (3%, v/v) of the El Verde soil microcosm in defined basal salt medium amended with pyruvate, H<sub>2</sub>, and N<sub>2</sub>O, all 16S rRNA amplicon sequences could be assigned to *Serratia* (68.0%), *Desulfosporosinus* (24.3%), *Desufitobacterium* (7.5%), *Caproiciproducens* (0.19%), *Peptoclostridium* (<0.05%) and *Lachnoclostridium* (<0.05%). Following nine consecutive transfers, *Serratia* and *Desulfosporosinus* each contributed about half of the 16S rRNA amplicon sequences (49.7% and 50.2%, respectively), and less than 0.05% of the sequences represented *Planctomycetota*, *Lachnoclostridium*, *Caproiciproducens*.

Following fifteen sequential transfers in the same medium reduced the microbial diversity to two populations, a *Serratia* sp. and *Desulfosporosinus* sp. In 16S rRNA gene clone libraries generated with DNA collected during Phase II from a 15<sup>th</sup> generation transfer culture, Sanger sequencing revealed 16S rRNA genes matching those of the *Serratia* sp. and *Desulfosporosinus* sp., and no other sequences were found. Deep metagenome sequencing (total of 387 Gbp obtained) yielded a 2,865-fold coverage of the *Serratia* sp. genome and 15,103-fold coverage of the *Desulfosporosinus* sp. genome (Supplementary Figure 2-3), and 99.48% of the short read sequences could be mapped to the contigs representing these two genomes. Of note, the *Serratia* genomes constructed from co-culture EV and the axenic *Serratia* culture were nearly identical (ANI 99.9%), suggesting the sequencing depth was sufficient to study the genomic characteristics in the co-culture. The assembly of 16S rRNA gene fragments generated eight partial and two complete 16S rRNA genes, which shared between 91.1 to 99.5% sequence identity to the respective *Serratia* sp. and *Desulfosporosinus* sp. 16S rRNA gene sequences determined by Sanger sequencing (Supplementary Figure 2-4). Both genomes possess a single 16S

rRNA gene, and the observed differences in 16S rRNA sequences were attributed to errors associated with short read assembly and sequencing. Phase contrast microscopy performed with co-culture EV suspension samples collected during Phase I and Phase II revealed the presence of the reported Serratia sp. (103) and the Desulfosporosinus sp. cell morphologies (104). Taken together, these observations support that 15 consecutive transfers yielded a consortium (i.e., a co-culture) comprising two bacterial populations, a Serratia sp. and a Desulfosporosinus sp. Deep shotgun metagenome sequencing performed on a 15<sup>th</sup> transfer culture recovered two draft genomes representing the Serratia sp. and the Desulfosporosinus sp., accounting for more than 95% of the total short read fragments. All 16S rRNA genes associated with assembled contigs could be assigned to Serratia or Desulfosporosinus (Supplementary Figure 2-4), indicating that the enrichment process yielded a consortium consisting of a Serratia sp. and a Desulfosporosinus sp., designated co-culture EV. Efforts to recover the Serratia and Desulfosporosinus genomes from the original soil metagenome data sets via recruiting the soil metagenome fragments to the two genomes (Figure 2-1 C) were not successful, highlighting the effectiveness of the enrichment strategy. Redundancy-based analysis with Nonpareil (89) revealed that the average covered species richness in the metagenome data set obtained from the 15<sup>th</sup> transfer culture was 99.9%, much higher than what was achieved for the El Verde soil (39.5%), suggesting the metagenome analysis of the original soil did not fully capture the resident microbial diversity.

The application of 16S rRNA gene-targeted qPCR assays to DNA extracted from 9<sup>th</sup> transfer N<sub>2</sub>O-reducing cultures revealed a bimodal growth pattern. During pyruvate fermentation (Phase I), the *Serratia* cell numbers increased nearly 1,000-fold from (2.3  $\pm$  0.8) × 10<sup>2</sup> to (1.8  $\pm$  0.2) × 10<sup>5</sup> cells mL<sup>-1</sup>, followed by a 40-fold increase from (3.5  $\pm$  1.5) × 10<sup>4</sup> to (1.2  $\pm$  0.4) × 10<sup>6</sup> cells mL<sup>-1</sup> of *Desulfosporosinus* cells during N<sub>2</sub>O reduction (Phase II) (Figure 2-1 D). In vessels without N<sub>2</sub>O, *Desulfosporosinus* cell numbers did not increase, indicating that growth of this population depended on the presence of N<sub>2</sub>O. Growth yields of (3.1  $\pm$  0.11) × 10<sup>8</sup> cells mmol<sup>-1</sup> of N<sub>2</sub>O and (7.0  $\pm$  0.72) × 10<sup>7</sup> cells mmol<sup>-1</sup> of pyruvate were determined for the *Desulfosporosinus* and the *Serratia* 

populations, respectively. 16S rRNA gene amplicon sequencing performed on representative samples collected at the end of Phase I (day 7) and Phase II (day 18) confirmed a bimodal growth pattern. Sequences representing *Serratia* increased during Phase I and *Desulfosporosinus* sequences increased during Phase II (Figure 2-1 E). Taken together, the physiological characterization, qPCR, genomic, and amplicon sequencing results indicate that co-culture EV performs low pH N<sub>2</sub>O reduction, with a *Serratia* sp. fermenting pyruvate and a *Desulfosporosinus* sp. reducing N<sub>2</sub>O. Isolation efforts yielded an axenic *Serratia* sp., designated strain MF, capable of pyruvate fermentation. Despite extensive efforts, the *Desulfosporosinus* sp. resisted isolation, presumably due to auxotrophic interactions with strain MF (Supplementary Figure 2-5).

## **Isolation efforts**

Numerous attempts were made to isolate the N<sub>2</sub>O-reducing *Desulfosporosinus* sp. from co-culture EV. Serial 10-fold dilution-to-extinction series in pH 4.5 basal salt liquid medium amended with 2.5 mM pyruvate, 4.16 mM (nominal) N<sub>2</sub>O, and 4.16 mM (nominal) H<sub>2</sub> recovered N<sub>2</sub>O reduction activity from  $10^{-6}$  dilution vials; however, the Serratia sp. was also present. The omission of pyruvate prevented growth of the Serratia sp. but without pyruvate, N<sub>2</sub>O consumption did not commence. Plating of serially diluted culture suspension aliquots on solid Tryptic Soy Agar (TSA) medium yielded uniform colonies. Sanger sequencing of PCR-amplified 16S rRNA genes of four isolated colonies from a 10<sup>-6</sup> dilution plate yielded identical sequences with 100% sequence identity to the Serratia sp. Microscopic analysis revealed motile, short rods about 2 µm long representing Serratia cells that grew readily in Tryptic Soy Broth medium. Growth also occurred in defined basal salt pH 4.5 medium amended with pyruvate, but the Serratia sp. did not consume  $H_2$  or  $N_2O$ . Following complete  $N_2O$  consumption in co-cultures grown with  $2.4 \pm 0.01$  mM pyruvate,  $4.06 \pm 1.16$  mM (nominal) of H<sub>2</sub>, and  $3.96 \pm 1.08$ mM (nominal) of N<sub>2</sub>O, *Desulfosporosinus* cells outnumbered Serratia cells approximately 5-fold. Microscopic observations also documented this population shift and *Desulfosporosinus* cells (motile rods about 6 µm in length) dominated the cell suspensions following N<sub>2</sub>O consumption. Attempts to obtain isolated colonies of the

*Desulfosporosinus* sp. in soft agar (0.8% low melting agarose, w/v) shake tubes were not productive because the reduction of N<sub>2</sub>O generated N<sub>2</sub> gas bubbles preventing the recovery of isolated colonies (Supplementary Figure 2-5). Despite extensive efforts, the *Desulfosporosinus* sp. could not be separated from the *Serratia* sp.

#### **Identification of auxotrophies**

To investigate the specific nutritional requirements of the Desulfosporosinus sp. in coculture EV, untargeted metabolome analysis was conducted on supernatant collected from axenic Serratia sp. cultures growing with pyruvate and during N<sub>2</sub>O consumption (Phase II) following inoculation with co-culture EV (Figure 2-2 A). Peaks representing potential metabolites were searched against a custom library and 33 features were assigned to known structures, including seven amino acids (alanine, glutamate, methionine, valine, leucine, aspartate, and tyrosine). Cystine, the oxidized derivative of the amino acid cysteine, was also detected; however, cystine or cysteine were not found in cultures where dithiothreitol (DTT) replaced cysteine as the reductant, suggesting that Serratia did not excrete either compound into the culture supernatant. Time series metabolome analysis of culture supernatant demonstrated dynamic changes to the amino acid profile following inoculation with the Serratia sp. and the Desulfosporosinus sp. (as co-culture EV) (Figure 2-2 A, B). Alanine, valine, leucine, and aspartate increased during pyruvate fermentation (Phase I) and were not consumed by the Serratia sp. (Supplementary Figure 2-6). Consumption of alanine, valine, leucine, and aspartate did occur following the inoculation of the *Desulfosporosinus* sp. (as co-culture EV) (Figure 2-2 A). These findings suggest that the  $N_2O$ -reducing *Desulfosporosinus* sp. is an amino acid auxotroph, and a series of growth experiments explored if amino acid supplementation (Table 2-5) can replace the requirement for pyruvate fermentation by the Serratia sp. for enabling subsequent N<sub>2</sub>O consumption by the Desulfosporosinus sp. The addition of individual amino acids (n=20) was not sufficient to initiate N<sub>2</sub>O reduction in pH 4.5 medium, as was the combination of alanine, valine, leucine, aspartate, and tyrosine. Incomplete N<sub>2</sub>O consumption (<20% of initial dose) was observed in cultures supplemented with the aforementioned amino acid combination plus methionine.  $N_2O$ 

reduction and growth of the *Desulfosporosinus* sp. occurred without delay in cultures supplied with a 15-amino acid mixture (Figure 2-2 C). Omission of single amino acids from the 15-amino acid mixture led to incomplete N<sub>2</sub>O reduction, similar to what was observed with the 6-amino acid combination. Efforts to isolate the *Desulfosporosinus* sp. in medium without pyruvate but amended with amino acids were unsuccessful because of concomitant growth of the *Serratia* sp., as verified with qPCR.

## pH range of acidophilic N<sub>2</sub>O reduction by the Desulfosporosinus sp

Growth assays with co-culture EV were performed to determine the pH range for  $N_2O$ reduction. Co-culture EV reduced N<sub>2</sub>O at pH 4.5, 5.0 and 6.0, but not at pH 3.5, 7.0 and 8.0. Cultures at pH 4.5 exhibited about two times longer (i.e., 10 versus 5 days) lag periods prior to the onset of N<sub>2</sub>O consumption than cultures incubated at pH 5.0 or 6.0 (Supplementary Figure 2-7 A). In medium without amino acid supplementation, pyruvate fermentation is required for the initiation of N<sub>2</sub>O consumption, raising the question if pH impacts pyruvate fermentation by the Serratia sp., N<sub>2</sub>O reduction by the Desulfosporosinus sp., or both processes. Axenic Serratia sp. cultures fermented pyruvate over a pH range of 4.5 to 8.0, with the highest pyruvate consumption rates observed at pH 6.0 and 7.0 (1.47  $\pm$  0.04 mmol L<sup>-1</sup> day<sup>-1</sup>), and the lowest rates measured at pH 4.5  $(0.43 \pm 0.05 \text{ mmol } \text{L}^{-1} \text{ day}^{-1})$  (Supplementary Figure 2-7 B). The N<sub>2</sub>O consumption rates in co-culture EV between pH 4.5 to 6.0 ranged from  $0.24 \pm 0.01$  to  $0.26 \pm 0.01$  mmol L<sup>-1</sup> day<sup>-1</sup> (Supplementary Figure 2-7 C). Since N<sub>2</sub>O reduction in the absence of amino acids only commenced following pyruvate consumption (Figure 2-1 C), pyruvate fermentation by *Serratia* sp., not  $N_2O$  reduction by *Desulfosporosinus* sp., explains the extended lag periods observed at pH 4.5 (Supplementary Figure 2-7 A). Consistently, shorter lag phase for both N<sub>2</sub>O reduction and *Desulfosporosinus* growth were observed in co-culture EV amended with the amino acid mixture (Figure 2-2 C).

# Phylogenomic analysis

Phylogenomic reconstruction based on concatenated alignment of 120 bacterial marker genes corroborated the affiliation of the N<sub>2</sub>O-reducing bacterium with the genus *Desulfosporosinus* (Figure 2-3). The genus *Desulfosporosinus* comprises strictly anaerobic, sulfate-reducing bacteria, and *Desulfosporosinus acididurans* strain SJ4 and *Desulfosporosinus acidiphilus* strain M1 were characterized as acidophilic sulfate reducers. The N<sub>2</sub>O-reducing *Desulfosporosinsus* sp. in co-culture EV possesses the *aprAB* and *dsrAB* genes encoding adenylyl sulfate reductase and dissimilatory sulfate reductase, respectively, but lacks the *sat* gene encoding sulfate adenylyl transferase sulfurylase (Supplementary Figure 2-8).

To provide experimental evidence that the N<sub>2</sub>O-reducing *Desulfosporosinus* sp. in coculture EV lacks the ability to reduce sulfate, a hallmark feature of the genus *Desulfosporosinus*, comparative growth studies were performed. The N<sub>2</sub>O-reducing *Desulfosporosinsus* sp. in co-culture EV did not grow with sulfate as sole electron acceptor, consistent with an incomplete dissimilatory sulfate reduction pathway (Supplementary Figure 2-9 A). *Desulfosporosinus acididurans* strain D (104), a close relative of the N<sub>2</sub>O-reducing *Desulfosporosinus* sp. in co-culture EV, grew with sulfate in pH 5.5 medium, but did not grow with N<sub>2</sub>O under the same incubation conditions (Supplementary Figure 2-9 B). These observations corroborate that the N<sub>2</sub>O-reducing *Desulfosporosinus* sp. lacks the ability to perform dissimilatory sulfate reduction. Based on phylogenetic and physiologic features, the N<sub>2</sub>O-reducer in culture EV represents a novel *Desulfosporosinus* sp., for which the name '*Candidatus* Desulfosporosinus nitrousreducens' strain PR is proposed.

# Genetic underpinning of N<sub>2</sub>O reduction in 'Ca. Desulfosporosinus nitrousreducens' strain PR

The strain PR genome harbors a single *nosZ* gene affiliated with clade II *nosZ* sequences (Figure 2-3). Independent branch placement of the strain PR NosZ on the clade II NosZ tree suggests an ancient divergence; a finding supported by amino acid identity (AI) value comparisons between the NosZ sequences versus their homologs in the genomes of

*Ca.* Desulfosporosinus nitrousreducens' strain PR and *Desulfosporosinus meridiei* (AI: 44% versus 73.83%) (Figure 2-3 and

Figure 2-4 B). The NosZ of '*Ca*. Desulfosporosinus nitrousreducens' strain PR is more similar (AI: 45%) to the taxonomically distant relative *Desulfotomaculum ruminis*.

Comparative analysis of the strain PR *nos* operon with bacterial and archaeal counterparts corroborated characteristic clade II features, including a Sec translocation system, genes encoding cytochromes and an iron-sulfur protein, and a *nosB* gene located immediately downstream of *nosZ* (Figure 2-5). *nosB* encodes a transmembrane protein of unknown function and has been exclusively found on clade II *nos* operons. The *nos* gene clusters found in closely related taxa (e.g., *Desulfosporosinus meridiei*, *Desulfitobacterium dichloroeliminans*, *Desulfitobacterium hafniense*) show similar organization; however, differences were observed in the *nos* gene cluster of 'Ca. Desulfosporosinus nitrousreducens' strain PR. Specifically, the genes encoding an iron-sulfur cluster protein and cytochromes precede *nosZ* in *Desulfosporosinus meridiei*, but are located downstream of two genes encoding proteins of unknown functions in strain PR (Figure 2-5). Of note, among the microbes with *nos* operons and included in the analyses, only '*Ca*. Desulfosporosinus nitrousreducens' and *Nitratiruptor labii* (105), both with a clade II *nos* operon, were experimentally validated to grow with N<sub>2</sub>O below pH 6 (Table 2-3).

#### Genomic insights for a commensalistic relationship

Functional annotation of the *Serratia* sp. and the '*Ca*. Desulfosporosinus nitrousreducens' strain PR genomes was conducted to investigate the interspecies interactions (Figure 2-6). A *ptsT* gene encoding a specific, high-affinity pyruvate/proton symporter (106) and genes implicated in pyruvate fermentation (i.e., *pflAB*, *poxB*) are present on the *Serratia* genome, but are missing on the strain PR genome, consistent with the physiological characterization results. *fdhC* genes encoding a formate transporter are present on both genomes, but only the strain PR genome harbors the *fdh* gene cluster encoding a formate dehydrogenase complex (Supplementary Figure 2-8), consistent with the observation that the *Serratia* sp. excretes formate, which strain PR utilizes as electron donor for N<sub>2</sub>O

reduction (Supplementary Figure 2-1). Gene clusters encoding two different Ni/Fe-type hydrogenases (i.e., *hyp* and hya gene *clusters*) (Supplementary Figure 2-8) and a complete *nos* gene cluster (Figure 2-5) are present on the strain PR genome, but not on the *Serratia* sp. genome.

Based on the KEGG (86) and Uni-Prot databases (85), the *Serratia* genome contains the biosynthesis pathways (100% completeness) for aspartate, lysine, threonine, tryptophan, isoleucine, serine, leucine, valine, glutamate, arginine, proline, methionine, tyrosine, cysteine, and histidine. In contrast, only aspartate and glutamate biosynthesis are predicted to be complete on the strain PR genome, whereas the completeness level for biosynthesis pathways of other amino acids was below 80%. The *Serratia* genome encodes a complete set of TCA cycle enzymes, indicating the potential for forming aspartate and glutamate via transamination of oxaloacetate and  $\alpha$ -ketoglutarate. In contrast, the strain PR genome lacks genes encoding malate dehydrogenase, citrate synthase, and aconitate hydratase, indicative of an incomplete TCA cycle. Therefore, strain PR is deficient of *de novo* formation of precursors for glutamate, aspartate, alanine, and related amino acids (107). A high-affinity amino acid transport system was found on the strain PR genome (Supplementary Figure 2-8), suggesting this bacterium can efficiently acquire extracellular amino acids to meet its nutritional requirements.

#### Distribution of the novel N<sub>2</sub>O reducer in soil metagenomes

For estimating the distribution of the *Serratia* sp. and '*Ca*. Desulfosporosinus nitrousreducens' strain PR in acidic and some circumneutral pH soils, fragments of soil metagenomes (n=195) from a prior study (29) were aligned to the *Serratia* sp. strain MF and the '*Ca*. Desulfosporosinus nitrousreducens' strain PR genomes (Supplementary Figure 2-10 A). The percentage of metagenome fragments that aligned to strain MF and strain PR ranged between 0.11-1.15% and 0.039-0.094%, respectively. RandomForest analysis revealed weak relationships between the presence of '*Ca*. Desulfosporosinus nitrousreducens' strain NH<sub>4</sub><sup>+</sup> content. In contrast, *Serratia* sp. strain MF showed the strongest relationship to '*Ca*. Desulfosporosinus

nitrousreducens' strain PR, explaining 32.6% of the variability of strain PR abundance in the metagenomes included in the analysis (Supplementary Figure 2-10 B). Furthermore, the relative abundances of the two populations showed a positive correlation (Supplementary Figure 2-10 C), suggesting the observed commensalistic relationship may not have coincidentally developed in co-culture EV, and possibly occurs in soil habitats. Soil organic matter (r = -0.35, p < 0.01) and NO<sub>3</sub><sup>-</sup> (r = -0.31, p < 0.01) content strongly correlated with the presence of '*Ca*. Desulfosporosinus nitrousreducens' strain PR.

#### 2.4 Discussion

Biological processes fix about 180 Tg N per year (108) and conventional agriculture introduces more than 100 Tg N of chemically fixed N each year (109). Biogeochemical cycling converts a substantial fraction of fixed N to N<sub>2</sub>O, and rising atmospheric N<sub>2</sub>O concentrations indicate that N<sub>2</sub>O sources outpace microbial N<sub>2</sub>O sinks. Soil acidification is a major concern relevant to N cycling and N<sub>2</sub>O emissions. Microbial processes (e.g., nitrification and denitrification) involved in N<sub>2</sub>O production have been reported to cause greater N<sub>2</sub>O emissions at pH < 6 than at circumneutral pH (31, 41). In contrast, characterized N<sub>2</sub>O reducers from soil habitats consume N<sub>2</sub>O within a narrow pH window ranging from 6 to 8 with generally weak activity approaching pH 6.0 and no activity at lower pH (38, 39, 110, 111). Interestingly, metagenomic studies revealed that acidic Luquillo Experimental Forest (Puerto Rico) tropical soils and temperate circumneutral pH soils harbor a comparable diversity of *nosZ* genes (43), and that pH selects for different N<sub>2</sub>O reducers in Luquillo tropical soil microcosms (44).

A few studies reported limited N<sub>2</sub>O reduction activity in acidic microcosms, but enrichment cultures for detailed experimentation were not obtained (41, 57, 112). Possible explanations for the observed N<sub>2</sub>O consumption in acidic microcosms include residual activity of existing N<sub>2</sub>O-reducing biomass, or the presence of microsites on soil particles where solid phase properties influence local pH, generating pH conditions not captured by bulk aqueous phase pH measurements (113, 114). Soil microcosms providing

microsites with favorable (i.e., higher) pH conditions can give the false impression of low pH N<sub>2</sub>O consumption; however, removal of solids during transfers eliminates this niche and microorganisms experience bulk phase pH conditions, resulting in the cessation of N<sub>2</sub>O reduction activity. Our work with acidic tropical soils alludes to another crucial issue, explicitly the choice of carbon source for the successful transition from microcosms to soil-free enrichment cultures. Lactate sustained N<sub>2</sub>O reduction only when pyruvate substituted lactate. Lactate has a higher p $K_a$  value than pyruvate (3.8 versus 2.45), indicating that a larger fraction of protonated, and potentially toxic, lactic acid exists at pH 4.5 (115). As discussed above, in soil microcosms, particles with ion exchange capacity (i.e., microsites) can suppress inhibitory effects of protonated organic acids, a possible explanation why lactate supported N<sub>2</sub>O reduction in the microcosms but not in the enrichment cultures.

Fifteen repeated transfers with N<sub>2</sub>O, pyruvate, and H<sub>2</sub> yielded a co-culture comprising a *Serratia* sp. and a *Desulfosporosinus* sp. The rapid enrichment of a co-culture may be surprising considering that pyruvate and H<sub>2</sub> are excellent substrates for many soil microbes. N<sub>2</sub>O was the sole electron acceptor provided to the defined basal salt medium, with some CO<sub>2</sub> being formed during pyruvate fermentation (Phase I); however, no evidence was obtained for H<sub>2</sub>-driven CO<sub>2</sub> reduction to acetate or to methane. N<sub>2</sub>O has been recognized as potent inhibitor of corrinoid-dependent enzyme systems (55, 56), and both CO<sub>2</sub>/H<sub>2</sub> reductive acetogenesis and hydrogenotrophic methanogenesis would not be expected to occur in the enrichment cultures, a prediction supported by the analytical measurements.

Available axenic and mixed cultures obtained from circumneutral pH soils reduce  $N_2O$  at circumneutral pH, but not below pH 6.0 (36, 39, 116). The discovery and cultivation of co-culture EV comprising '*Ca*. Desulfosporosinus nitrousreducens' strain PR extends the pH range for microbial growth with  $N_2O$  as electron acceptor to pH 4.5. Metagenomic studies routinely detect *nosZ* in acidic soils providing evidence that bacteria capable of low pH  $N_2O$  reduction exists and are likely more widely distributed in acidic soil habitats

(43, 117, 118). The distribution of bacteria capable of low pH N<sub>2</sub>O reduction may extend to the deep marine biosphere, a hypothesis supported by the recent discovery of the thermophile *Nitratiruptor labii* strain HRV44 with the capacity to reduce N<sub>2</sub>O over a pH range of 5.4 to 6.4 (105).

<sup>•</sup>*Ca.* Desulfosporosinus nitrousreducens' strain PR reduces N<sub>2</sub>O between pH 4.5 and 6.0, with no N<sub>2</sub>O reduction observed at or above pH 7. This finding implies that '*Ca.* Desulfosporosinus nitrousreducens' cannot be enriched with N<sub>2</sub>O as electron acceptor at or above pH 6.5, suggesting the maintenance of acidic pH conditions during enrichment is crucial to cultivate new microorganisms capable of low pH N<sub>2</sub>O reduction. Available N<sub>2</sub>O-reducing soil isolates enriched and isolated at circumneutral pH cannot grow with N<sub>2</sub>O as electron acceptor below pH 6.0. Apparently, pH selects for distinct groups of N<sub>2</sub>O reducers, with prior research mostly focused on soil isolates obtained at circumneutral pH. The discovery of '*Ca.* Desulfosporosinus nitrousreducens' strain PR lends credibility to the hypothesis that the diverse *nosZ* genes found in metagenomes of acidic soils (43) may indeed be functional. Further, the cultivation of strain PR provides a blueprint for unraveling the unknown microbiology of low pH N<sub>2</sub>O reduction and exploring the geochemical parameters that govern this process in acidic soils.

*Ca.* Desulfosporosinus nitrousreducens' strain PR possesses a clade II *nos* operon similar to those found in clade II neutrophilic N<sub>2</sub>O reducers without distinguishing features based on gene content and *nos* operon structure (Figure 2-5). Experimental work with *Paracoccus denitrificans*, a model organism harboring a clade I *nosZ* and used for studying denitrification, has shown that acidic pH impairs N<sub>2</sub>O reductase assembly and maturation (38). NosZ is a periplasmic enzyme with a distinguishing feature of clade I versus clade II NosZ being the mechanism of protein transport across the cytoplasmic membrane. Clade II NosZ follows the general Secretion route known as Sec-pathway, which translocates unfolded proteins. In contrast, clade I NosZ are translocated in a folded state via the Twin-arginine pathway (Tat-pathway) (119). Differences exist between clade I and cade II *nos* operons, and *nosB*, encoding a transmembrane protein of unknown function, has been exclusively found associated with clade II *nos* clusters

(Figure 2-5) (12, 23). To what extent gene content and the secretion pathway influence the pH range of N<sub>2</sub>O reduction is unclear and warrants further genetic/biochemical studies. Other factors that can impact N<sub>2</sub>O reduction at acidic pH include an organism's ability to cope with the potential toxicity of protonated organic acids and maintain pH homeostasis (120, 121). The '*Ca*. Desulfosporosinus nitrousreducens' strain PR genome harbors multiple genes associated with DNA repair and potassium transport, suggesting this bacterium can respond to pH stress.

Soils harbor diverse microbiomes with intricate interaction networks that govern soil biogeochemical processes, including N<sub>2</sub>O turnover (122), and define the functional dynamics of microbiomes (123-125). Interspecies cooperation between bacteria can enhance  $N_2O$  reduction via promoting electron transfer (126), the provision of essential nutrients (as demonstrated in co-culture EV), or limit N<sub>2</sub>O reduction due to competition for electron donor(s) or metal cofactors (i.e., copper) (127, 128). Metabolomic workflows unraveled that *Serratia* sp. strain MF excretes amino acids during growth with pyruvate, which 'Ca. Desulfosporosinus nitrousreducens' strain PR requires to initiate N<sub>2</sub>O reduction, a finding supported by genome functional predictions (i.e., 15 complete AA biosynthesis pathways in strain MF versus only two complete AA biosynthesis pathways in strain PR). An amino acid mixture supported N<sub>2</sub>O reduction by strain PR in the absence of pyruvate; however, slight growth of *Serratia* sp. strain MF was observed, preventing the isolation of the N<sub>2</sub>O reducer Interspecies interactions based on amino acid auxotrophies have been implicated in shaping dynamic anaerobic microbial communities, bolster community resilience, and thus promote functional stability (123). Other microbes can potentially fulfill the nutritional demands of '*Ca*. Desulfosporosinus nitrousreducens', and the observed interrelationship between the Serratia sp. and strain PR might have developed coincidentally during the enrichment process; however, the cooccurrence pattern does imply some level of specificity and that the interactions may occur in situ.

Members of the genus *Desulfosporosinus* have been characterized as strictly anaerobic sulfate reducers with the capacity to grow autotrophically with H<sub>2</sub>, CO<sub>2</sub>, and sulfate, or,

in the absence of sulfate, with pyruvate (129). Most characterized *Desulfosporosinus* spp. show optimum growth at circumneutral pH  $(\sim 7)$  conditions, except for the acidophilic isolates Desulfosporosinus metallidurans, Desulfosporosinus acidiphilus, Desulfosporosinus acididurans, and Desulfosporosinus sp. strain I2, which perform sulfate reduction at pH 4.0, 3.6, 3.8, and 2.6, respectively (104, 115, 130-132). Among the 10 Desulfosporosinus species with sequenced genomes, only Desulfosporosinus meridiei DSM 13257 carries a nos cluster (12). Desulfosporosinus meridiei DSM 13257 grows within a narrow range of pH 6-8 (129), and its ability to reduce  $N_2O$  has not been demonstrated. 'Ca. Desulfosporosinus nitrousreducens' strain PR lacks the hallmark feature of sulfate reduction and is the first acidophilic soil bacterium capable of growth with  $N_2O$  as electron acceptor at pH 4.5, but not at pH 7 or above. Strain PR couples  $N_2O$ reduction and growth at pH 4.5 with the oxidation of H<sub>2</sub> or formate, and our experimental efforts with co-culture EV could not demonstrate the utilization of other electron donors. The four characterized acidophilic representatives of the genus *Desulfosporosinus* show considerable versatility, and various organic acids, alcohols, and sugars, in addition to H<sub>2</sub>, support sulfate reduction (104, 130, 131). The utilization of H<sub>2</sub> as electron donor appears to be a shared feature of *Desulfosporosinus* spp., and two or more gene clusters encoding hydrogenase complexes were found on available *Desulfosporosinus* spp. genomes (132-134).

Escalating usage of N fertilizers to meet societal demands for agricultural products increases N<sub>2</sub>O emissions due to accelerated N cycling and associated N<sub>2</sub>O formation and due to soil acidification, which presumably slows N<sub>2</sub>O consumption. Liming is commonly employed to ameliorate soil acidity, a practice considered beneficial for curbing N<sub>2</sub>O emissions based on the assumption that microbial N<sub>2</sub>O reduction is limited to circumneutral pH soils (39-41, 47). Our findings demonstrate that soils harbors microorganisms (e.g., '*Ca*. Desulfosporosinus nitrousreducens' strain PR) that utilize N<sub>2</sub>O as growth-supporting electron acceptor between pH 4.5 and 6.5. Apparently, N<sub>2</sub>O consumption is not limited to circumneutral pH conditions and acidophilic respiratory N<sub>2</sub>O reducers exist in acidic soils. Thus, increased N<sub>2</sub>O production due to fertilizer

application could potentially be mitigated, at least partially, by microbial N<sub>2</sub>O consumption in acidic soils. Recent efforts have shown substantial N<sub>2</sub>O emission reductions from soils treated with biofertilizers or augmented with mixed cultures harboring clade II N<sub>2</sub>O reducers (135, 136). Based on metagenomic surveys, bacteria capable of low pH N<sub>2</sub>O are not limited to acidic tropical soils, and are more broadly distributed in terrestrial ecosystems (44). The discovery of a naturally occurring soil bacterium that couples N<sub>2</sub>O consumption to growth under acidic pH conditions offers opportunities for the knowledge-based manipulation of agricultural soils to stimulate microbial N<sub>2</sub>O consumption. Curbing undesirable N<sub>2</sub>O emissions at the field scale would allow farmers to further reduce their greenhouse gas emissions footprint and potentially earn carbon credits.

#### Appendix

Figure 2-1. Establishment of N<sub>2</sub>O-reducing microcosms and enrichment cultures vielding co-culture EV. (A) Schematic of the workflow leading from a soil sample to a solids-free enrichment, and to a co-culture. (B) Community structure profiling based on 16S rRNA gene sequence analysis documents the enrichment process. Soil microbial community structure was profiled with 16S rRNA genes extracted from shotgun metagenomic reads. Community profiling of 6<sup>th</sup> and 9<sup>th</sup> transfers was based on 16S rRNA gene amplicon sequencing. Sequences with abundances <2% were grouped as "Others". (C) Percent of the metagenomic short read fragments obtained from El Verde soil and the 15<sup>th</sup> transfer culture that recruited to the genomes of *Serratia* sp. or *Desulfosporosinus* sp. A representative graph showing the high identity (>95%) of reads mapping evenly across Desulfosporosinus sp. genome is presented in Supplementary Figure 2-3. The Serratia sp. or Desulfosporosinus sp. genomes were not detected in the soil metagenome dataset, with less than 0.01% of the metagenomic reads mapping to the two genomes (white bar). (D) Pyruvate fermentation (Phase I, white circles) and N<sub>2</sub>O consumption (Phase II, black circles) in co-culture EV. Serratia (blue diamonds) and Desulfosporosinus (red diamonds) cell numbers were determined with specific, 16S rRNA gene-targeted qPCR. (E) Amplicon sequencing illustrates the population shifts in co-culture EV following pyruvate consumption (day 7) and following  $N_2O$  consumption (day 18). Relative abundance of Serratia (blue bars) and Desulfosporosinus (red bars) in co-culture EV following Phase I (day 7) and Phase II (day 18) in vessels with pyruvate plus N<sub>2</sub>O (left) and pyruvate only (right). Sequencing was performed on single representative cultures. All other data represent the averages of triplicate incubations and error bars represent standard deviations (n=3). Error bars are not shown if smaller than the symbol.

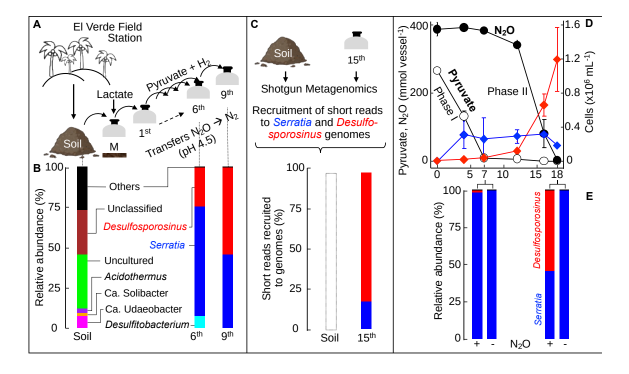
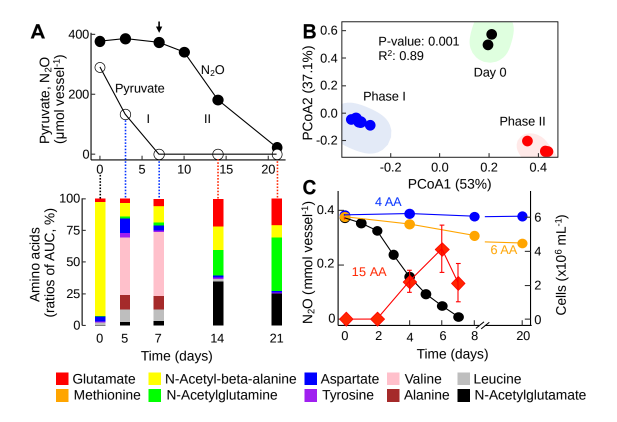


Figure 2-2. Interspecies cross-feeding supports low pH N<sub>2</sub>O reduction in co-culture EV. (A) Pyruvate fermentation (Phase I, light blue background) in vessels inoculated with axenic Serratia sp. strain MF and N<sub>2</sub>O consumption (Phase II, red background) following inoculation with co-culture EV comprising the Serratia sp. and the N<sub>2</sub>Oreducing Desulfosporosinus sp. (indicated by the arrow). The bottom part of Panel A shows the amino acid profile in the supernatant immediately after inoculation with the Serratia sp. (green background), during Phase I, and during Phase II following inoculation with co-culture EV (day 7; 3% inoculum). Samples for untargeted metabolome analysis were collected immediately after inoculation with Serratia sp. (light green), during Phase I (light blue, pyruvate fermentation) and Phase II (light red, N<sub>2</sub>O reduction). The stacked bars show the ratio (%) of areas under the curve (AUC) of the respective amino acids and amino acid derivatives. Metabolites not assigned to structures representing amino acids or its derivatives are not shown. (B) Principal coordinate analysis (PCoA) of amino acid profiles. The enclosing ellipses were estimated using the Khachiyan algorithm with the ggforce package. Permanova analysis was conducted with the vegan Community Ecology package. Black, blue, and red circles represent samples collected at day 0, during Phase I, and during Phase II, respectively. (C) N<sub>2</sub>O consumption in co-culture EV in medium amended with mixtures comprising 5 (blue), 6 (orange), or 15 (black) amino acids (see SI for composition of mixtures), H<sub>2</sub>, and N<sub>2</sub>O. The Desulfosporosinus sp. cell numbers (red diamonds) were determined with 16S rRNA gene-targeted qPCR and show growth in medium receiving the 15-amino acid mixture. Various other amino acid mixtures tested resulted in no or negligible N<sub>2</sub>O consumption. All growth assays with amino acid mixture supplementation were performed in triplicates and repeated in independent experiments.



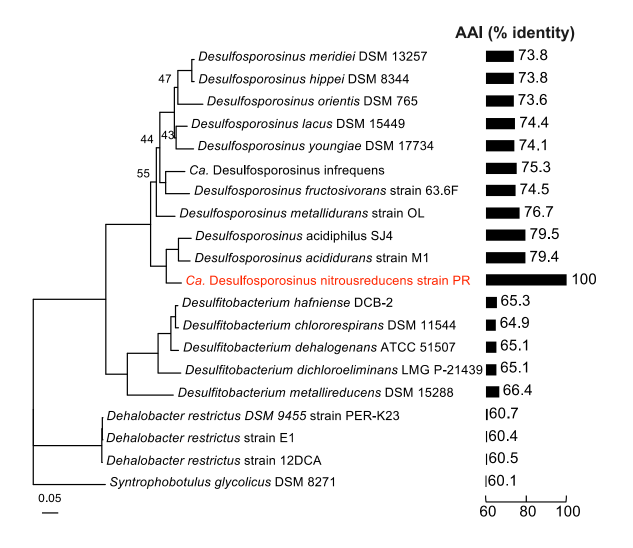
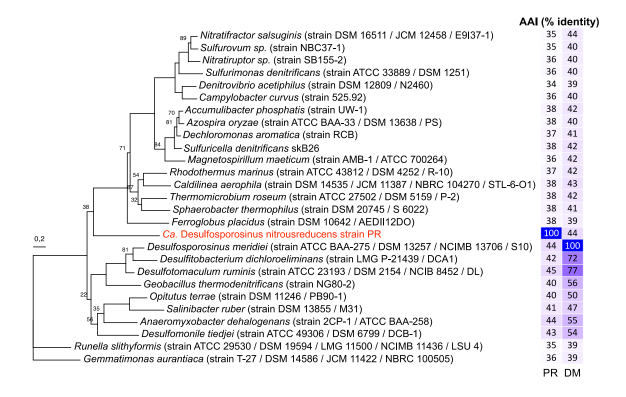
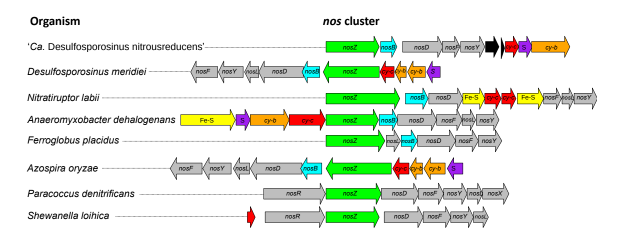


Figure 2-3. Phylogenomic and Average Amino acid Identity (AAI) analyses indicate that the N<sub>2</sub>O-reducing strain PR in co-culture EV represents a new species of the genus *Desulfosporosinus*. Phylogenomic analysis was based on 120 conserved marker genes and included *Peptococcaceae* genomes available from NCBI. Bootstrap values higher than 90 are not displayed. The scale bar indicates 0.05 nucleotide acid substitution per site. Bar plots display the genome-wide AAI (%) between the N<sub>2</sub>O-reducing '*Ca*. Desulfosporosinus nitrousreducens' and related isolates with sequenced genomes.

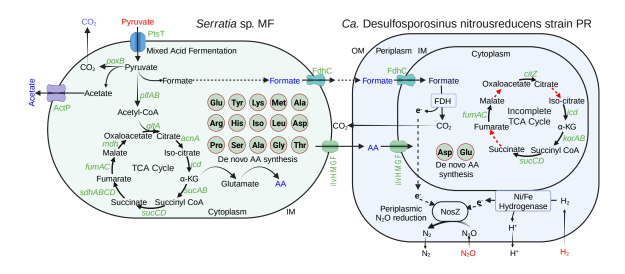


# Figure 2-4. Relatedness and similarity of the clade II NosZ of 'Ca.

**Desulfosporosinus nitrousreducens' strain PR to representative clade II NosZ.** The tree represents a phylogenetic reconstruction of select clade II NosZ protein sequences. The clade II NosZ of *Gemmatimonas aurantiaca* was used to root the tree. The scale bar indicates 0.2 amino acid substitution per site. Numbers at nodes are bootstrap values smaller than 90. The two-column heatmap shows the AAI values between the NosZ of strain PR (PR) and *Desulfosporosinus meridiei* (DM) to other clade II NosZ sequences, with the darker shades of blue indicating higher percent AAI values.

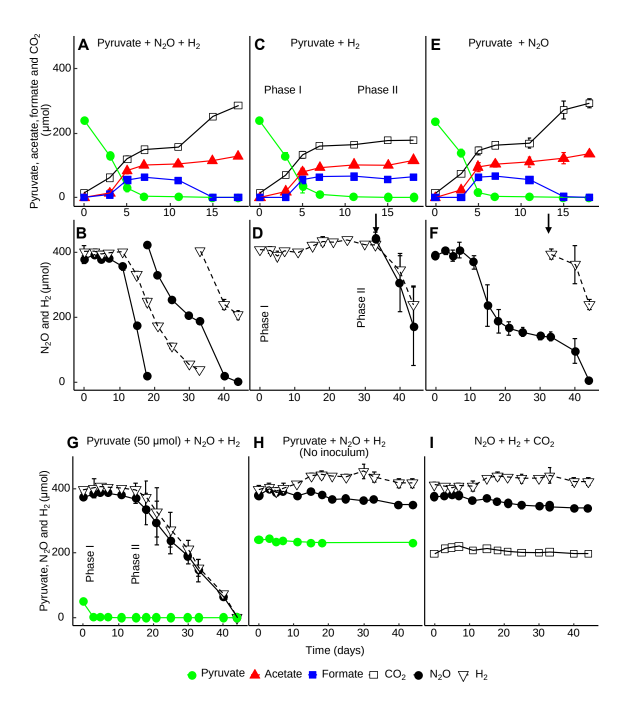


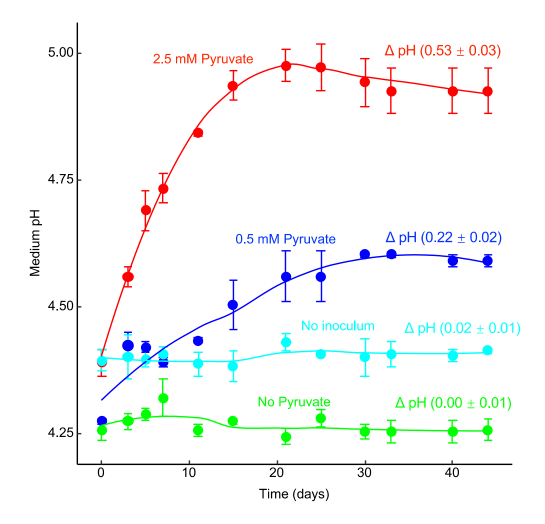
**Figure 2-5.** Comparison of representative *nos* clusters. Included are clade II *nos* clusters encoding select NosZ. and select clade I *nos* clusters of bacteria with confirmed N<sub>2</sub>O reduction activity at circumneutral pH. The colored arrows represent genes with different functions and indicate orientation and approximate length. Green, *nosZ*; gray, *nos* accessory genes (i.e., *nosD*, *nosF*, *nosY*, *nosL*, *nosX* and *nosR*); yellow, genes encoding iron-sulfur (Fe-S) proteins; purple, genes encoding Rieske iron-sulfur proteins (S); orange (cy-b) and red (cy-c), genes encoding b-type and c-type cytochromes, respectively; cyan, *nosB* genes encoding transmembrane proteins characteristic for clade II *nos* operons; black, genes of unknown function. *Desulfosporosinus* and *Desulfitobacterium* spp., *Nitratiruptor* and *Nitratifractor* spp., and *Paracoccus* and *Bradyrhizobium* spp. share similar *nos* cluster architectures, respectively.



**Figure 2-6. Proposed interspecies cross-feeding interactions in co-culture EV.** Genes involved in mixed acid fermentation are only found on the *Serratia* sp. strain MF genome, and genes involved in periplasmic N<sub>2</sub>O reduction are exclusive to '*Ca*. Desulfosporosinus nitrousreducens' strain PR. External substrates (i.e., pyruvate or the 15-amino acid mixture, H<sub>2</sub>, N<sub>2</sub>O) provided to co-culture EV are shown in red font, and metabolites produced by *Serratia* are shown in blue font. Fifteen versus two complete amino acid biosynthesis pathways are present on the *Serratia* sp. strain MF and '*Ca*. Desulfosporosinus nitrousreducens' strain PR genomes, respectively. Strain PR has an incomplete TCA cycle, and the red dashed arrows indicate the absence of the corresponding genes. TCA cycle: tricarboxylic acid cycle; AA: amino acids; FDH: formate dehydrogenase complex; NosZ: nitrous oxide reductase; OM: outer membrane; IM: inner membrane.

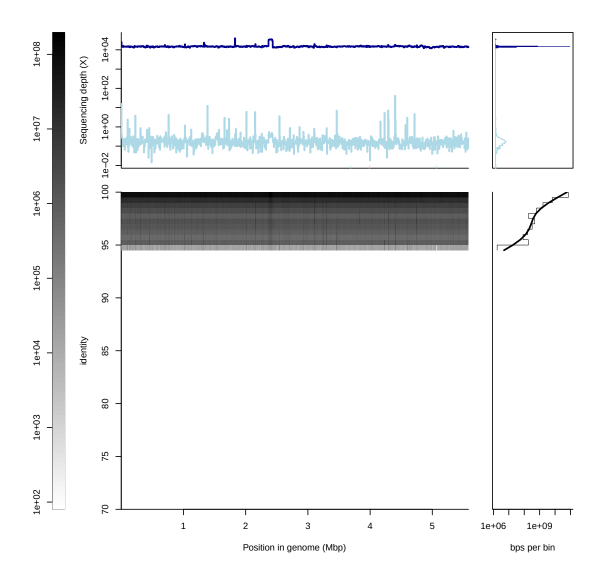
**Supplementary Figure 2-1.** Phenotypic characterization of co-culture EV. Cultures were grown in 160 mL glass serum bottles containing 100 mL of medium amended with pyruvate (50 or 250  $\mu$ mol), H<sub>2</sub> (416  $\mu$ mol) and N<sub>2</sub>O (416  $\mu$ mol) (A and B), pyruvate and H<sub>2</sub> (C and D), or pyruvate and N<sub>2</sub>O (E and F). The arrows in panels D and F indicate additional doses of N<sub>2</sub>O or H<sub>2</sub> to corresponding culture vessels lack N<sub>2</sub>O (D) or H<sub>2</sub> (F) to resume N<sub>2</sub>O reduction activity. Pyruvate consumption and associated product formation are documented in panels A, C, and E. N<sub>2</sub>O and H<sub>2</sub> consumption are documented in panels B, D, F. Panel G shows the performance of co-culture EV grown with a lower amount of pyruvate. Panel H depicts pyruvate, H<sub>2</sub> and N<sub>2</sub>O amounts over time in vessels without inoculum. Autotrophic activity in cultures with CO<sub>2</sub>, but lacking pyruvate, was not observed (I). The light blue shaded areas represent the pyruvate fermentation phase (Phase I) and the light red areas represent the N<sub>2</sub>O reduction phase (Phase I). The data represent the averages of triplicate incubations and error bars represent the standard deviations (n=3). Error bars are not shown when smaller than the symbol.

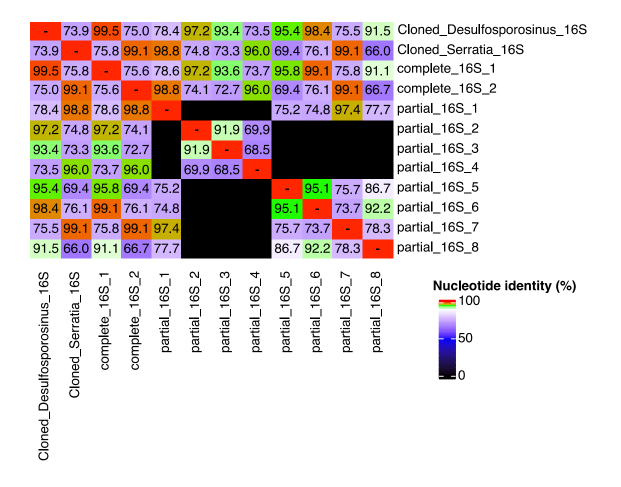




Supplementary Figure 2-2. Medium pH over the course of the incubation. pH profiles were measured in replicate co-culture EV incubation vessels that received 0, 0.5, and 2.5 mM pyruvate. pH changes ( $\Delta$  pH) were calculated by subtracting the pH values measured immediately following inoculation from pH values measured at the end of incubation period. Pyruvate was not consumed in vessels that were not inoculated with co-culture EV. The data shown are the averages of triplicate incubations and error bars represent the standard deviations (n=3). Error bars are not shown when smaller than the symbol.

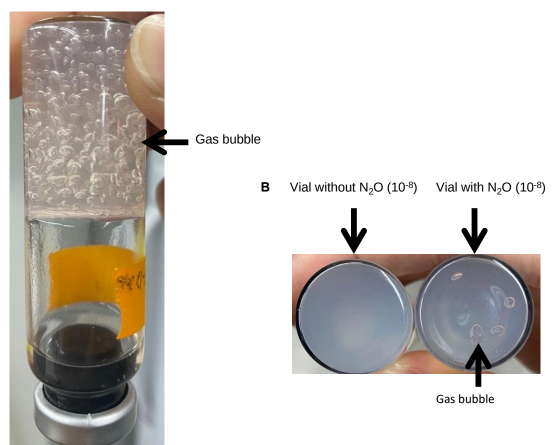
Supplementary Figure 2-3. Fragment recruitment plot of the 'Ca. Desulfosporosinus nitrousreducens' strain PR genome to the metagenome dataset derived from the 15<sup>th</sup> generation co-culture EV. The recruitment was performed via processing a BLAST search of the metagenome fragments against the 'Ca. Desulfosporosinus nitrousreducens' strain PR genome. The tabular BLAST result was parsed using BLASTab.catsbj.pl, and graphical representation was generated with the BlasTab.recplot2.R embedded in Enveomics collection (88). (A) The bar on the left shows the number of fragments with different identity recruited to each position on the genome. (B) indicates the sequencing depth across the 'Ca. Desulfosporosinus nitrousreducens' strain PR genome on a logarithmic scale. (C) Sequencing depth histogram with peaks from values above 95% identity automatically identified as skewed normal distribution. (D) Metagenome fragments recruited to the 'Ca. Desulfosporosinus nitrousreducens' strain PR genome, placed by location (x-axis) and identity (y-axis). (E) Identity histogram of mapping fragments (light gray) and smoothed spline (black). The backgrounds in panels D and E, and the line colors in panels B and C, correspond to identity matches above (dark blue) and below (light blue) 95 %. A recruitment plot of the 'Ca. Desulfosporosinus nitrousreducens' strain PR genome to the metagenome datasets derived from El Verde soil is not shown as the covered fraction of the 5.6 Mbp genome was below 10% (i.e., 0.56 Mbp).



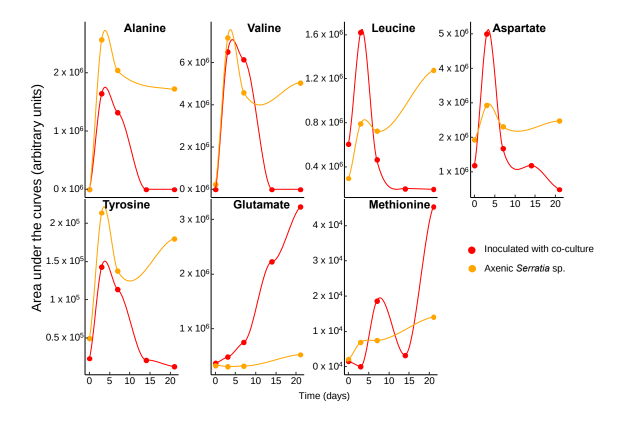


Supplementary Figure 2-4. Comparison of '*Ca*. Desulfosporosinus nitrousreducens' strain PR and *Serratia* sp. strain MF 16S rRNA gene sequences derived from cloned 16S rRNA gene fragments (Sanger sequencing) and co-culture EV (metagenome sequencing). DNA extracted from co-culture EV following N<sub>2</sub>O consumption served as template for the amplification of 16S rRNA gene fragments with general primer pair 8F-1541R. 16S rRNA gene fragments were cloned in *E. coli* and two uniform clone populations were obtained, represented by Cloned\_Desulfosporosinus\_16S and Cloned\_Serratia\_16S.

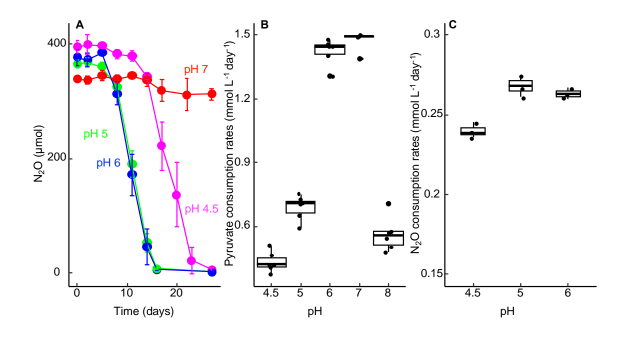
A  $10^{-6}$  diluted vial with N<sub>2</sub>O



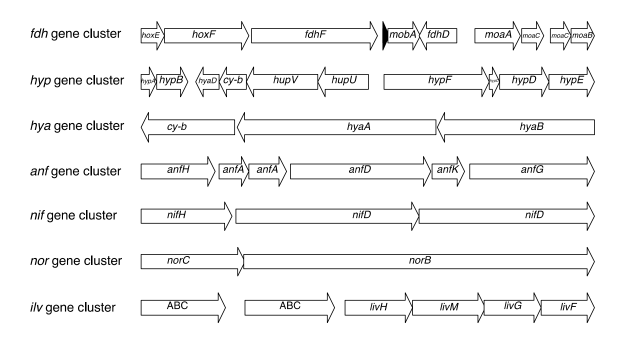
Supplementary Figure 2-5. Formation of gas bubbles in soft agar shake tubes. (A) Visible gas bubbles (presumably N<sub>2</sub>) following N<sub>2</sub>O consumption. The vials were incubated with the stoppers down for 28 days. (B) The formation of gas bubbles was strictly dependent on N<sub>2</sub>O, and no bubbles formed in replicate vials without N<sub>2</sub>O. No gas bubbles formed in control incubations without inoculum.



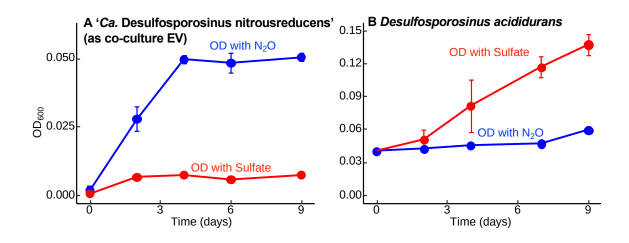
**Supplementary Figure 2-6. Dynamic changes of amino acids in the supernatants of cultures following inoculation with** *Serratia* **sp. and with** *'Ca.* **Desulfosporosinus nitrousreducens' strain PR (as co-culture EV).** Axenic *Serratia* **sp.** cultures amended with pyruvate, N<sub>2</sub>O, and H<sub>2</sub> were inoculated with co-culture EV on day 7. Orange lines represent amino acids in cultures inoculated with axenic *Serratia* **sp.** on day 0. Red lines represent amino acids in cultures inoculated with the axenic *Serratia* **sp.** on day 0 and subsequently inoculated with co-culture EV on day 7. Note that axenic *Serratia* **sp.** cultures without co-culture EV inoculum cannot reduce N<sub>2</sub>O. Shown are representative data obtained from a single culture in medium reduced with 1-cysteine. Cystine was detected in cultures reduced with 1-cysteine; however, in samples from replicates from an independent experiment where DTT served as reductant, cystine was not detected, suggesting *Serratia* **sp.** does not release cystine into the medium.



Supplementary Figure 2-7. Performance of co-culture EV at different medium pH values. (A) N<sub>2</sub>O reduction was measured in triplicate 160 mL serum bottles containing 100 mL of medium (pH 3.5, 4.5, 5, 6, 7, 8) and inoculated with co-culture EV. N<sub>2</sub>O reduction was observed between pH 4.5 and 6, but not at pH 3.5 and at or above pH 7. (B) Pyruvate consumption rates by *Serratia* sp. in cultures adjusted to pH 4.5, 5, 6, 7, and 8. Pyruvate was consumed in all incubation vessels except for those at pH 3.5. (C) N<sub>2</sub>O consumption rates of '*Ca*. Desulfosporosinus nitrousreducens' at pH 4.5, 5, and 6. No N<sub>2</sub>O consumption was observed in pH 3.5, 7.0, and 8.0 cultures. Consumption rates of pyruvate and N<sub>2</sub>O were calculated using data that fell within the linear ranges of consumption. Each black dot in the box plots represent the consumption rates calculated from a single experiment. The data from three and six replicate cultures were used for the calculations of pyruvate and N<sub>2</sub>O consumption rates, respectively.



Supplementary Figure 2-8. Gene clusters found on the '*Ca*. Desulfosporosinus nitrousreducens' strain PR genome. Displayed are formate dehydrogenase (*fdh* gene cluster), b-type Ni/Fe hydrogenase (*hyp* gene cluster), group 1 Ni/Fe hydrogenase (*hya* gene cluster), FeFe type Nitrogenase (*anf* gene cluster), Mo-Fe Nitrogenase (*nif* gene cluster), Nitric-oxide reductase (*nor* gene cluster), High-affinity amino acid transport system (*ilv* gene cluster). The arrows indicate operon length and orientations. Preliminary annotation of coding genes was done with Prokka (137), MicrobeAnnotator (138), and the RAST server (87), and genes of interests were curated via blasting to the NCBI nr database. Cy-b: b-type cytochrome; ABC: ATP-binding cassette. The black arrow in the *fdh* gene cluster represents a gene encoding a small protein of unknown function.

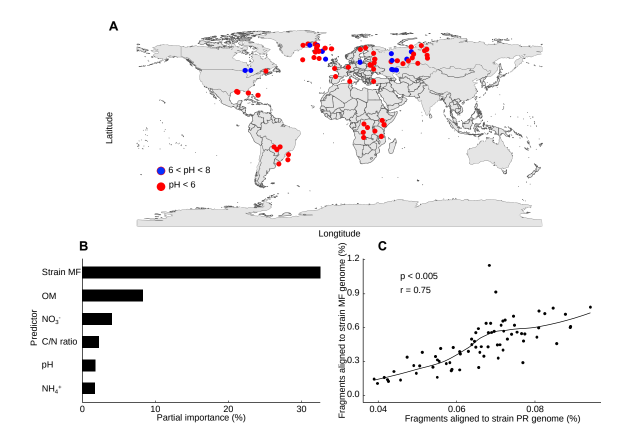


# Supplementary Figure 2-9. Comparative growth studies with 'Ca. Desulfosporosinus nitrousreducens' strain PR (as co-culture EV) and

*Desulfosporosinus acididurans* strain **D**. Panel A shows growth of '*Ca*. Desulfosporosinus nitrousreducens' strain PR with 4.16 mM (nominal) N<sub>2</sub>O (blue) or 5 mM SO<sub>4</sub><sup>2-</sup> (red) as electron acceptor in pH 4.5 medium amended with the 15-amino acid mixture and 4.16 mM (nominal) H<sub>2</sub>. Over the 9-day incubation period, strain PR completely consumed the initial dose of N<sub>2</sub>O, whereas cultures amended with 5 mM SO<sub>4</sub><sup>2-</sup> showed no growth or SO<sub>4</sub><sup>2-</sup> consumption (4.97  $\pm$  0.23 mM sulfate remained). OD values below 0.01 were measured in replicate cultures that did not receive N<sub>2</sub>O as electron acceptor or H<sub>2</sub> as electron donor. Panel B compares growth of *Desulfosporosinus acididurans* strain D with 4.16 mM (nominal) N<sub>2</sub>O (blue) or 5 mM SO<sub>4</sub><sup>2-</sup> (red) as electron acceptors in pH 5.5 medium with 10 mM glycerol as electron donor. Over a 9-day incubation period, *Desulfosporosinus acididurans* strain D reduced 5 mM SO<sub>4</sub><sup>2-</sup>, but N<sub>2</sub>O was not consumed (4.10  $\pm$  0.59 mM N<sub>2</sub>O remained) in replicate cultures, even after an extended 20-day incubation period. Growth was monitored by measuring the optical density at 600 nm (OD<sub>600</sub>).

#### Supplementary Figure 2-10. Distribution of 'Ca. Desulfosporosinus

nitrousreducens' strain PR and *Serratia* sp. strain MF in select publicly available soil metagenomes. Metagenome datasets were downloaded from the European Nucleotide Archive (accession number PRJEB44414) with associated metadata available in a Supplemental Information/Source Data file (https://doi.org/10.1038/s41467-022-29161-3). (A) Geographical distribution of the soil metagenomes included in the analysis. Blue dots indicate a soil pH >7, red dots indicate a soil pH <7. (B) The partial importance (%) of different parameters, including the presence of *Serratia* sp. (based on short reads that align to the *Serratia* sp. strain MF genome), organic matter (OM content), NO<sub>3</sub><sup>-</sup> content, NH<sub>4</sub><sup>+</sup> content, C/N ratio, and pH for the relative abundance of '*Ca*. Desulfosporosinus nitrousreducens' strain PR based on Random Forest models (139). (C) Correlation between percentages of short read fragments aligned to the *Serratia* sp. strain MF and the '*Ca*. Desulfosporosinus nitrousreducens' strain PR genomes. The significance values for the correlations were calculated with the Pearson test (140) implemented in the ggpubr package (https://github.com/kassambara/ggpubr).



**Table 2-1.** qPCR primers and probe designed for the enumeration of *Serratia* sp. strain MF and '*Ca*. Desulfosporosinus nitrousreducens' strain PR 16S rRNA genes.

Target	Primer sequence	Probe
<i>Serratia</i> sp.	TGCTACAATTGGCGTATACAA (Ser_1316qF) GACTACGACATACTTTATGA (Ser_1386qR)	None
Desulfosporosinus sp.	TGCTACAATGGCCGGTACAG (DS_1316qF) CGAACTGAGACCGGCTTTCT (DS_1388qR)	FAM-AAGCTGTGAAGTGGAGCCAATC- MGB <sup>a</sup>

<sup>a</sup> FAM: 6-Carboxyfluorescein attached to the 5' end of the probe. MGB: Minor Groove Binder that attached to the 3' end of the probe.

Dataset	Data type	Accession No.	
Enrichment culture (6 <sup>th</sup> transfer)	Amplicon (16S V3-V4)	SRR24215177	
Enrichment culture (9 <sup>th</sup> transfer)	Amplicon (16S V3-V4)	SRR24083098	
Co-culture EV (15 <sup>th</sup> transfer)	SRA <sup>a</sup>	SRR24709127	
Serratia sp. (isolate)	SRA <sup>a</sup>	SRR24709126	
16S rRNA gene (Desulfosporosinus)	Complete gene	OR076434	
16S rRNA gene (Serratia)	Complete gene	OR076433	
<i>'Ca.</i> Desulfosporosinus nitrousreducens strain PR'	GenBank Assembly <sup>b</sup>	GCA_030954495.1	
<i>Serratia</i> sp. strain MF	GenBank Assembly <sup>b</sup>	GCA_030954505.1	

**Table 2-2.** Sequence information submitted to the NCBI database.

<sup>a</sup> SRA: Sequence Read Archive (raw sequence data)

<sup>b</sup> Genome assembly constructed from SRA data

Organism	Genome NCBI acc no.	Reported pH range for N2O reduction	nosZ clade	Reference
<i>Ca</i> . Desulfosporosinus nitrousreducens' strain PR	GCA_030954495.1	4.5-6	II	This work
Desulfosporosinus meridiei strain DSM 13257	NC_018515.1	ND	II	(12)
Nitratiruptor labii strain HRV44	AP022826.1	5.4-6.4	II	(105)
Anaeromyxobacter dehalogenans strain 2CP-1	NC_011891.1	~7	II	(12)
Ferroglobus placidus	NC_013849.1	ND	II	(12)
Azospira oryzae	GCA_905120365.1	ND	II	(12)
Paracoccus denitrificans strain DSM 413	AL591688.1	7-8	Ι	(38)
Shewanella loihica strain PV-4	NC_009092.1	6-8	Ι	(111)
Alicycliphilus denitrificans strain I51	AP024172.1	6-9	Ι	(110)
Bradyrhizobium diazoefficiens strain 110spc4	NZ_CP032617.1	6-8	Ι	(141)
				58

**Table 2-3.** Organismal and genomic information of microorganisms with reported N<sub>2</sub>O reduction capacity under different pH.

**Table 2-4.** Genomes used in phylogenomic analysis and classification of '*Ca*.Desulfosporosinus nitrousreducens'.

Organism	NCBI accession No. of genomes		
<i>Ca</i> . Desulfosporosinus nitrousreducens' strain PR	GCA_030954495.1		
Desulfosporosinus youngiae DSM 17734	NZ_CM001441.1		
Desulfosporosinus orientis DSM 765	NC_016584.1		
Desulfosporosinus metallidurans strain OL	NZ_MLBF01000001.1		
Desulfosporosinus meridiei DSM 13257	NC_018515.1		
Desulfosporosinus lacus DSM 15449	NZ_FQXJ01000052.1		
Desulfosporosinus hippie DSM 8344	NZ_fncp01000067.1		
Desulfosporosinus fructosivorans strain 63.6F	NZ_SPQQ01000010.1		
Desulfosporosinus acidiphilus SJ4	NC_018068.1		
Desulfosporosinus acididurans strain M1	NZ_LDZY01000001.1		
'Ca. Desulfosporosinus infrequens'	NZ_OMOF01000971.1		
Syntrophobotulus glycolicus DSM 8271	NC_015172.1		
Desulfitobacterium metallireducens DSM 15288	NZ_CP007032.1		
Desulfitobacterium hafniense DCB-2	NC_011830.1		

**Table 2-4 Continued** 

Organism	NCBI accession No. of genomes		
<i>Desulfitobacterium dichloroeliminans</i> LMG P- 21439	NC_019903.1		
Desulfitobacterium dehalogenans ATCC 51507	NC_018017.1		
Desulfitobacterium chlororespirans DSM 11544	NZ_FRDN01000033.1		
Dehalobacter restrictus strain E1	CANE01000102.1		
Dehalobacter restrictus strain 12DCA	NZ_CP046996.1		
Dehalobacter restrictus DSM 9455	NZ_CP007033.1		

Amino acid	Amino acid concentration (µM)	Carbon (µmol 100 mL medium <sup>-1</sup> )		
Alanine	56	16.8		
Valine	77	38.5		
Aspartate	82	33.1		
Isoleucine	38	22.9		
Methionine	27	13.4		
Tyrosine	16	14.9		
Histidine	19	11.6		
Tryptophan	10	10.8		
Glutamate	68	34.0		
Proline	86	43.5		
Arginine	28	17.2		
Glycine	40	8.0		
Threonine	42	16.8		
Lysine	69	41.1		
Serine	76	22.9		

**Table 2-5.** Amino acids and concentrations used to augment the growth medium forcultivation of '*Ca*. Desulfosporosinus nitrousreducens'. <sup>a</sup>

<sup>a</sup> The addition of individual amino acids did not stimulate N<sub>2</sub>O-dependent growth of '*Ca*. Desulfosporosinus nitrousreducens'. Similarly, a 5-amino acid mixture comprising alanine, valine, leucine, aspartate, and tyrosine did not promote N<sub>2</sub>O consumption. Weak and incomplete N<sub>2</sub>O reduction was observed in cultures that received a 6-amino acid mixture comprising alanine, valine, leucine, aspartate, tyrosine, and methionine. The 15-amino acid mixture (i.e., all amino acids listed in Table 5) supported growth of '*Ca*. Desulfosporosinus nitrousreducens' via hydrogenotrophic N<sub>2</sub>O reduction.

Table 2-6. Genome features of 'Ca. Desulfosporosinus nitrousreducens	' strain PR and Serratia sp. strain MF, and comparison to
closest relatives.	

– Feature	Organism <sup>a</sup>				
	(1)	(2)	(3)	(4)	(5)
Genome size (bp)	5,591,411	4,991,181	4,637,866	5,118,938	5,300,955
Completeness (%) <sup>b</sup>	99.5	99.1	92.3	98.0	95.6
Contamination (%) <sup>b</sup>	5.7	0.0	5.14	0.53	4.03
GC content (%)	44	42	41.5	58.8	59.5
5S rRNA genes	8	8	8	8	8

Feature	Organism <sup>a</sup>					
	(1)	(2)	(3)	(4)	(5)	
23S rRNA genes	1	8	11 <sup>d</sup>	1	7	
tRNA genes	105	66	67	81	91	
Coding sequences	5,312	4,554	4,317	4,690	4,941	
Accession number	GCA_030954495.1	GCA_000255115.3	GCA_001029285.1	GCA_030954505.1	GCF_002220655.1	
<sup>a</sup> (1) 'Ca. Desul	fosporosinus nitrousred	ucens' strain PR (this	study)			
(2) Desulfospor	rosinus acidiphilus strai	n SJ4 (130)				
(3) Desulfospor	<i>cosinus acididurans</i> stra	in M1 (104)				

Table 2-7. Genome features of 'Ca. Desulfosporosinus nitrousreducens' strain PR and Serratia sp. strain MF, and close relatives.

(5) Serratia marcescens strain UMH3 (142)

(4) Serratia sp. strain MF (this study)

<sup>b</sup> Genome completeness and contamination were evaluated based on the presence of 120 single copy genes using CheckM.

<sup>c</sup> Only partial 16S and 23S rRNA genes were found on *Desulfosporosinus acididurans* strain M1 genome.

Chapter 3 Clade III N<sub>2</sub>O-respiring microbes implicated in N<sub>2</sub>O reduction.

#### Abstract

Nitrous oxide (N<sub>2</sub>O), a significant contributor to global warming and ozone depletion, has raised concerns over its increasing presence in the atmosphere. N<sub>2</sub>O consumption is limited to microorganisms known to possess clade I and clade II *nosZ* genes, but the N<sub>2</sub>O reduction activity progressively wanes with decreasing pH. Even more troublesome is the widespread phenomenon of soil acidification, a natural process accelerated by agricultural practices. In this study, we established enrichment cultures from acidic tropical forest soils (pH 4.4 to 4.9) and circumneutral regenerative agricultural soils (pH ~ 7), demonstrating sustained, growth-linked N<sub>2</sub>O reduction at an initial pH of 4.5. Characterization of the enrichment cultures suggest previously unknown clade II and a functional clade III N<sub>2</sub>O reducers obligately consume N<sub>2</sub>O at pH < 7. These results provide insight into the biodiversity and physiology of N<sub>2</sub>O-reducing microorganisms, which have pivotal roles in curbing N<sub>2</sub>O emissions, especially in acidic soil systems.

#### **3.1 Introduction**

Nitrous oxide (N<sub>2</sub>O, "laughing gas") is a powerful greenhouse gas with 298 times the warming potential of carbon dioxide (CO<sub>2</sub>) over a lifetime of 100-year (51), and N<sub>2</sub>O is also a strong ozone depleting substance (1). N<sub>2</sub>O also interferes with biogeochemical processes such as methanogenesis, mercury methylation, and reductive dichlorination (55, 56, 143). The Foundation Platform F20 report emphasis of limiting global warming to 1.5°C through curbing CO<sub>2</sub> emission (144); however, this effort could possibly be offset by the growing atmosphere N<sub>2</sub>O concentration (145). Natural and agricultural soils are primary natural and anthropogenic sources of N<sub>2</sub>O, as a consequence of biological nitrogen fixation and increasing load of synthetic nitrogen fertilizer and manure, respectively (146, 147). Compilation of global N<sub>2</sub>O budget suggest that the natural and anthropogenic N<sub>2</sub>O emissions are 9.7 and 7.3 Tg N y<sup>-1</sup>, respectively (3). Without putting N<sub>2</sub>O emissions under tight regulation, atmospheric N<sub>2</sub>O will continue to rise with the increasing demand of the exploding population growth for agricultural product, threatening global climate and ozone layer (148).

The rising N<sub>2</sub>O emissions across the globe indicate an imbalance between N<sub>2</sub>O formation versus consumption, with the rates of former process exceeding the latter. The most ancient origin (i.e., more than 2.7 billion years ago) of N<sub>2</sub>O is the iron mineral catalyzed abiotic process (7), which still occurs in modern tropic soils (5). The early anaerobic microbial life evolved N<sub>2</sub>O reductase, the only known enzyme catalyzing N<sub>2</sub>O reduction to date, to cope with tremendous N<sub>2</sub>O on Archaean earth prior to the great oxygen event (GOE) (7). Denitrification and aerobic nitrification appeared only shortly after the GOE (149), contribute to most N<sub>2</sub>O production on modern earth. Net positive N<sub>2</sub>O fluxes have been reported for peat land (150), forest (151), agricultural (152) and even permafrost soils (153), supporting the generalized paradigm that N<sub>2</sub>O production rates outweigh N<sub>2</sub>O consumption rates; however, net negative N<sub>2</sub>O fluxes, ranging from -0.0014 to -484  $\mu$ g N<sub>2</sub>O-N m<sup>-2</sup> h<sup>-1</sup>, have been reported in pristine and agricultural ecosystems (154), suggesting some soils are a potential N<sub>2</sub>O sink. The net negative N<sub>2</sub>O fluxes measured in field sites could not differentiate N<sub>2</sub>O absorption (physical process) and its

transformation to N<sub>2</sub> (biotic process), and uncertainty exist in the underlying microbiology.

Microorganisms with nosZ gene encoding N<sub>2</sub>O reductase, known to include clade I and II, could reduce N<sub>2</sub>O for the objectives of denitrification, growth-linked respiration, detoxification and electron disposal (18). Soil pH is a key environmental factor influencing microbial N<sub>2</sub>O reduction (30). The pH optima of microbial N<sub>2</sub>O reduction in literature ranges from 7 to 8, and acidic pH was reported to interfere with  $N_2O$  reductase assembly (38, 60), probably via disrupting the Cu cofactor associated histidine residue ligand (5). Most characterized N<sub>2</sub>O reducers with clade I nosZ could not consume N<sub>2</sub>O below pH 6 (38, 39, 110); however, recent advances reported an acidophilic soil bacterium, 'Ca. Desulfosporosinus nitrousreducens' strain PR with a clade II nos cluster, reduces  $N_2O$  between pH 4.5 to 6 in a sustainable circulating manner (i.e., growth linked N<sub>2</sub>O respiration) (Guang, et al., 2023). A coincidental discovery of a clade II thermophile *Nitratiruptor labii*, respiring N<sub>2</sub>O between pH 5.4 to 6.4, extends the low pH N<sub>2</sub>O reduction to deep marine biosphere (105). Physiology and genetic differences exist between clade I and II N<sub>2</sub>O reducers. Clade II N<sub>2</sub>O reducers have higher substrate affinity for N<sub>2</sub>O (101) and a nosB gene, encoding a transmembrane protein of unknown function, has been found exclusively associated with clade II nos clusters (12). As such, a hypothesis was generated that low pH N<sub>2</sub>O reduction is a shared feature among clade II N<sub>2</sub>O reducers that warrants/requires a broader investigation on low pH N<sub>2</sub>O reducers. In this study, we enriched low pH N<sub>2</sub>O-reducing microorganisms from acidic Luquillo Experimental Forest soils (pH 4.5 to 4.9, Puerto Rico) and circumneutral regenerative agricultural soils (pH ~7, Carthage, Tennessee). Characterization of the enrichment cultures revealed that microbial population responsible for low pH N<sub>2</sub>O reduction belong to previously unknown taxa. Clade II N<sub>2</sub>O reducers are enriched under rigorous acidic condition, validating the hypothesis that low pH  $N_2O$  reduction is a shared feature among clade II  $N_2O$  reducers. An unexpected, but surprising, discovery of this study is the identification of a functional novel clade (i.e., clade III) NosZ consuming N<sub>2</sub>O only at acidic pH.

# **3.2 Materials and Methods**

# Soil sampling, microcosms and enrichment cultures

Acidic LEF (i.e., Sabana and Palm Nido sites) microcosms set up and performance were detailed in our prior study (44), and CFF soil (i.e., WS1, WS2, C1 and C1982) containing microcosms were set up in 160 mL glass serum bottles containing 100 mL of anoxic, completely synthetic, acidic (pH 4.5) basal salt medium following established procedure (Guang, et al., under review). A mass of 2.5-gram CFF soils were added to medium containing serum bottles in anaerobic chamber. Lactate and N<sub>2</sub>O served as electron donor and acceptors in all microcosms. N<sub>2</sub>O was monitored via sampling headspace gas during microcosm incubations.

Transfer cultures were established in 20 mL glass serum bottles containing 9 mL of completely synthetic medium as described with the 0.27 g L<sup>-1</sup> Tryptone-Soy Broth was augmented to the medium to meet potential nutrient requirement of auxotrophic microbes (Guang, et al., under review). The bottles with N2 headspace were sealed with butyl rubber stoppers (Bellco Glass, Vineland, NJ, USA) held in place with aluminum crimp caps and autoclaved. Measured medium pH ranges from 4.27 to 4.40 following autoclave. All subsequent amendments (i.e., filter-sterilized [0.2 µm polyethersulfone membrane filter] and undiluted gases) to the medium bottles were augmented with sterile plastic syringes and needles. Carbon source and electron donors used for transfer cultures are summarized in Table S2.  $H_2$  and  $N_2O$ , 1-mL of each, were equilibrated for 24 hours in vessels prior to inoculation from corresponding microcosms showing  $N_2O$  reduction activity. The microcosms were manually shaken for homogenization before 1 mL aliquots were transfer with 1-mL plastic syringes. Attempts of transition from microcosm to solids-free enrichment cultures, with lactate as carbon source and reducing equivalent, were unsuccessful. As such, carbon alternates plus  $H_2$  were tested to resume  $N_2O$ reduction activity in transfer cultures. Of note, 0.27 g L<sup>-1</sup> Tryptone-Soy Broth was augmented to the medium to meet potential nutrient requirement of auxotrophic microbes (Guang, et al., under review). Consecutive transfer in acidic basal salt medium occurred

when the initial dose of 10 mL  $N_2O$  had been consumed, which yielded a total of six solid-free cultures showing  $N_2O$  reduction activity with an initial pH of 4.5. All culture vessels were incubated in upright position at 30°C in the dark without agitation.

#### **Comparative genomic analysis**

Comparative genomic analysis was conducted to identify organisms enriched with  $N_2O$ dosing. Cultures grown in the presence or absence of N<sub>2</sub>O served as positive or negative controls, respectively. Following  $N_2O$  depletion in corresponding cultures, biomass from 5 mL cell suspension samples were collected by centrifugation (10,000 x g, 20 min, 4 °C), and genomic DNA was isolated from the cell pellets using the DNeasy PowerSoil Kit (Qiagen, Hilden, Germany). Short gun metagenomic sequencing was performed with NovaSeq at the University of Tennessee Genomics Core following published procedures (63). Short gun sequencing generated raw sequences ranging from 12.4 to 20.9 Gbp. Nonpareil v3.4.1 using the weighted NL2SOL algorithm was used to estimate the average coverage level of the metagenomic datasets (89). Quality control, assembly and metagenome assembled genome binning of short read were performed with nf-core/mag v2.1.0 (77) with Nextflow (65) using containerized software in singularity (66). Short read quality was evaluated with FastQC v0.11.9 (67), followed by quality filtering and Illumina adapter removal using fastp v0.20.1. (78). PhiX sequences removal was performed using Bowtie2 v2.4.2 (79) referencing genome GCA\_002596845.1, ASM259684v1. Processed short reads from cultures amended with/without N<sub>2</sub>O were combined and co-assembled with Megahit2 v1.2.9 (80). Binning of coassemblys was conducted with MetaBAT2 v2.15 (81) and MaxBin v2.2.7 (155), and the resultant bins were refined using DASTools v1.1.6 (156). Metagenome-assembled genomes (MAGs) that passed quality control (completeness > 80%, contamination < 20%) of CheckM v1.2.1 (82) were dereplicated using dRep v3.4.2 (157) for further microbial community profiling. Protein-coding sequences on MAGs were predicted using Prokka v1.14.6 (137). Genome coverage of metagenomic assembled genomes (MAGs) of each metagenome were calculated with CoverM (https://github.com/wwood/CoverM). The coverage evenness of short reads on MAGs were assessed with in-house script

(https://github.com/rotheconrad/00\_in-situ\_GeneCoverage), based on distribution of high nucleotide identity reads across the reference genome sequences. Taxonomy classification of MAGs were performed using GTDB-TK v2.1.0 with classify workflow (91).

#### **Phylogenomic analysis**

Phylogenomic reconstruction was performed with genomes of the *Desulfosporosinus*, *Desulfitobacterium*, *Calidifontibacillus*, *Paludibacterium* genus available in the NCBI database. Conserved marker genes of the 20 reference genomes and 7 MAGs (constructed in this study) were identified and aligned with GTDB-TK (91). Phylogenomic relationships were inferred based on the alignment of 120 concatenated bacterial marker genes using RAxML-NG (92) with 1,000 bootstrap replicates. Implemented evolutionary model was selected based on the result of Modeltest-NG (93). Calculation of average amino acid identity (AAI) and hierarchical clustering of taxa based on AAI values were conducted with EzAAI (94). Graphical representation of tree topology and associated metadata were generated using the ggtreeExtra package (158).

# Computational analysis of nos operon cluster and NosZ structure modeling

A total of 173 NosZ reference sequences were downloaded from pre-compiled models of ROCker (96), and was further clustered to 80 representative sequences (75% amino acid identity) to reduce tree topology complexity. Eleven NosZ amino acid sequences were identified and translated from *nosZ* genes in seven MAGs, and were aligned with the 80-representative reference NosZ sequences using MAFFT (97). A maximum likelihood tree of NosZ was constructed with RAxML-NG based on the output evolutionary model of Modeltest-NG. Preliminary phylogenetic analysis of NosZ revealed the deep-branching placement of MAG6 and MAG7 NosZ, and subsequent phylogenetic analysis included homolog amino acid sequences of the two NosZ in NCBI database. Amino acid identity of NosZ was calculated using FastAAI package (159). Accessory genes of the NosZ from a set of 9 genomes described in our prior study (12) were used as PSI-BLAST database to

query accessory genes on MAGs with predicted *nosZ* genes (160). Transmembrane helices in the proteins located downstream of NosZ were predicted with DeepTMHMM based on deep neural networks (90). Signal peptides of NosZ proteins were inferred with SignalP 6.0 (161). Visualization and annotation of phylogenetic tree and associated data was performed using ggtreeExtra package implemented in R (158). Signal peptide of Nos operon amino acid sequences were removed prior to structure prediction with the AlphaFold2 implemented in ColabFold (162). Membrane orientation of proteins was predicted using MembraneFold (163). Active sites (i.e., Cu<sub>A</sub> and Cu<sub>Z</sub> sites) of NosZ were predicted using AlphaFill (164). The best-predicted NosZ model was selected and loaded into ChimeraX for visualization (165).

# Comparative growth assays with enrichment cultures harboring clade III N<sub>2</sub>O reducers

To validate the functionality of clade III NosZ, comparative growth assays were performed with cultures Sabana and WS2 in 160 mL vessels containing 100 mL medium described above. Replicate cultures amended with or without N<sub>2</sub>O were compared to verify the growth of clade III N<sub>2</sub>O reducers is N<sub>2</sub>O-dependent. Cultures was also grown in pH 7 medium, where 30 mM bicarbonate replaces 50 mM KH<sub>2</sub>PO<sub>4</sub>, to testify the capacity of N<sub>2</sub>O reduction at circumneutral pH.

#### **Quantitative PCR**

SYBR Green quantitative PCR (qPCR) assays targeting the clade III *nosZ* genes of MAG6 and MAG7 were designed using Geneious Prime. Primer were examined by in silico analysis using the Primer-BLAST tool, and experimentally confirmed using 1,599 bp-long synthesized linear DNA of the respective *nosZ* genes of the MAG6 and MAG7 (Integrated DNA Technologies, USA). The two different nosZ genes could be amplified with the same primer set (395F-527R). For enumeration of clade III *nosZ* genes, 25  $\mu$ L qPCR tubes received 10  $\mu$ L 1X Power SYBR Green, 9.88  $\mu$ L UltraPure nuclease-free water (Invitrogen, Carlsbad, CA, USA), 300 nM of each primer (395F:

GACCGTTTGGACCTCACCAT and 527R: CCGCCGGAAACATCGATTTC), and 2  $\mu$ L template DNA. All qPCR assays were performed using an Applied Biosystems ViiA 7 system (Applied Biosystems, Waltham, MA, USA) with the following amplification conditions: 2 min at 50°C and 10 min at 95°C, followed by 40 cycles of 15 s at 95°C and 1 min at 60°C. The standard curves were generated using 10-fold serial dilutions of the linear DNA fragments carrying a complete sequence of MAG6 (1,599 bp) or MAG7 (1,599 bp) *nosZ* gene, covering the same 133-bp qPCR target region.

The qPCR standard curves established with the linear DNA fragments carrying complete MAG7 *nosZ* gene had slopes of -3.471, y-intercepts of 36.431,  $R^2$  values of 0.998, and qPCR amplification efficiencies of 94.119%, respectively. The linear range spanned 8.54 to 8.54 x 10<sup>8</sup> gene copies per reaction. The functional analysis indicated that both the MAG6 and the MAG7 carry a single *nosZ* gene, indicating that the enumeration of *nosZ* gene allows estimates of cell abundances.

#### 3.3 Results

#### Low pH N<sub>2</sub>O reducers enriched from circumneutral and acidic soils

The in-situ pH of Luquillo Experimental Forest and Caney Fork Farm soils ranged from 4.72 to 4.97 (43) and 6.74 to 6.98, respectively (Table 3-1). As such, the two type soils are representative acidic natural and circumneutral agricultural soils. N<sub>2</sub>O consumption at pH 4.5 was observed in all microcosms with corresponding soils (one microcosm set up for each soil) amended with lactate as a source of carbon and reducing equivalents, and the N<sub>2</sub>O reduction performance differed strikingly between the microcosms with LEF and CFF soils. The initial doses (10 mL, 4.16 mM nominal) of N<sub>2</sub>O in LEF microcosms were consumed following 4-month incubation (44), whereas complete disappearance of same amount of N<sub>2</sub>O was observed in CFF microcosms after two weeks. Transition from microcosm to soils-free enrichment cultures with lactate at pH 4.5 resulted in the loss of N<sub>2</sub>O reduction activity; subsequently alternative carbon sources and electron donors were tested aiming to resume N<sub>2</sub>O reduction activity in transfer cultures. The initiation of N<sub>2</sub>O

reduction in CFF soil- and Sabana (LEF soil) soil-derived transfer cultures occurred with fumarate, whereas Palm Nido (LEF soil) soil-derived cultures required succinate, respectively (Table 3-2). Consecutive transfers performed in closed glass serum bottles (20 mL) containing defined pH 4.5 mineral salt medium (10 mL), tryptone-soy broth (0.1g L<sup>-1</sup>), the respective carbon source (5 mM), and H<sub>2</sub> (10 mL, 4.16 mM nominal) yielded stable enrichment cultures that reduced N<sub>2</sub>O at rates ranging from 0.39 to 1.53 mM day<sup>-1</sup> (Supplementary Figure 3-1). Complete N<sub>2</sub>O consumption was not observed in cultures lacking H<sub>2</sub>, indicating H<sub>2</sub> was the direct electron donor for N<sub>2</sub>O reduction. Over the course of the incubation, an increase in pH by 0.5 to 0.7 units was observed in enrichment cultures with carbon sources added at 5 mM concentration. With lower 0.5 mM concentrations, pH changes were <0.5 pH units, but the lag times for the initiation of N<sub>2</sub>O consumption increased.

Circumneutral (pH 7) soil microcosms were established with the CFF and PR soils, and transition from microcosms to soils-free transfer cultures (pH 7) was easily achieved in defined pH 7 medium with lactate; however, N<sub>2</sub>O reduction activity was abolished following the transitions from pH 7 to pH 4.5 medium. Consecutive transfers and dilution-to-extinction assays were further performed on WS2 and Sabana cultures in pH 7 medium amended with 5 mM lactate plus 10 mL (4.16 mM nominal) N<sub>2</sub>O. The transfer procedure commenced in pH 7 medium rapidly resulted in highly enriched cultures; 16S rRNA based amplicon sequencing showed that *Azospira* population accounting for more than 99.9% of the total sequences in the pH 7 culture derived from Sabana soils.

#### Novel species involved in low pH N<sub>2</sub>O reduction

To identify the microbial taxa involved in low pH  $N_2O$  reduction, shot-gun metagenome sequencing was performed to six different enrichment cultures grown with and without  $N_2O$  (Supplementary Figure 3-2). Metagenomes of transfer cultures and the original soils were compared to evaluate the enrichment effort. Nonpareil analysis (Supplementary Figure 3-3) showed that the microbiota in PR- and CFF-soils and were highly diverse or poorly sequenced with sequence coverages ranging from 26.04% and 40.01% sequence coverages. In contrast, the extracted community DNA of enrichment cultures were completely sequenced; the enrichment cultures had sequence coverages of at least 98.91%.

Population genome binning was performed with contigs from the coassembly (combining metagenomes from the same cultures with/without N<sub>2</sub>O). Following quality assessment (completeness > 80% and contamination < 25%) and dereplication (95% ANI threshold), a total of 43 MAGs were retained for profiling microbial community (Supplementary Figure 3-3). Only small portions (5.53-29.4%) of the short-read sequences could not be recruited to the high-quality MAGs, suggesting the 43 MAGs capture the diversity of each of the six enrichment cultures (Supplementary Figure 3-4). Community structures showed a strong impact of N<sub>2</sub>O on community composition (profiled based on highquality MAGs). Specifically, MAGs representing the class *Desulfitobacteriia* accounted for 14.1% to 81.7% of the total short-read sequences in cultures fed with  $N_2O$  except for C1. In the same cultures without N<sub>2</sub>O, *Desulfitobacteriia* represented no more than 5% of the total sequences. In culture C1, Bacilli population showed strong relationship to N<sub>2</sub>O and contributed 23.2% of the total sequences (versus 1.15% without N<sub>2</sub>O). Recruitment of metagenomes of LEF soils (n=2) and CFF soils (n=18) to the 43 high-quality MAGs was not successful, suggesting they have a low abundance, or are not representative population in the original soils.

To shed light on the N<sub>2</sub>O reducers enriched at pH 4.5, the 43 MAGs were annotated with Prokka and screened for *nosZ* genes. A total of 7 MAGs were predicted to possess at least one *nosZ* gene sequence, and 5 of the 7 MAGs were remarkably more abundant in cultures received N<sub>2</sub>O. Interestingly, MAG7 representing *Desulfitobacterium* 1 population were consistently enriched in culture WS1, WS2, and C1982 though the initial inoculum source (soils) differed (Figure 3-1 A). In culture C1, MAG1 representing *Calidifontibacillus* was more abundant in the presence of N<sub>2</sub>O, suggesting its growth is N<sub>2</sub>O-dependent. MAG3 (*Desulfosporosinus* 1) and MAG2 (*Paludibacterium*) showed contrasting pattern in culture Palm; MAG3 and MAG2 were more abundant in the presence or absence of N<sub>2</sub>O, respectively. Surprisingly, MAG4 (*Desulfosporosinus* 2)

exclusive to culture Palm amended with N<sub>2</sub>O was identical to '*Ca*. Desulfosporosinus nitrousreducens' strain PR (ANI > 99.9%) characterized in a co-culture (Guang, et al., 2023); while the partner bacterium *Serratia* sp. MF is not present in culture Palm. In culture Sabana, MAG6 representing a *Desulfitobacterium* 2 different from that present in CFF cultures (i.e., WS1, WS2, and C1982) was highly abundant (> 80%) in the presence of N<sub>2</sub>O.

An insightful observation is that all *nosZ*-carrying MAGs recovered the low pH  $N_2O$ -reducing cultures represent previously unknown species in the corresponding genera (

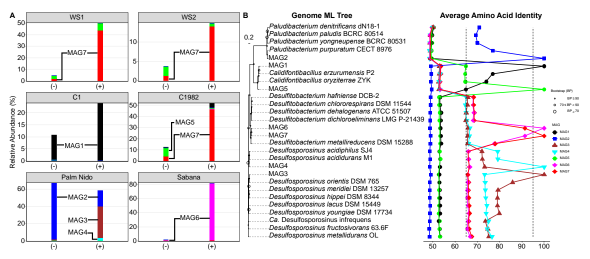


Figure **3-1** B). Albeit assigned to the same genus, the MAG3 and MAG4 are respectively more related to acidophilic *Desulfosporosinus* species (i.e., *D. acididurans* and *acidophilus*) and *Desulfosporosinus orientis*, none of which have the genetic machinery (*nos* operon) for N<sub>2</sub>O reduction. The MAG6 and MAG7 have the highest average amino acid identity (AAI: 91.3%) among the overall pairwise comparison, suggesting they represent different species that closely related to the *Desulfitobacterium metallireducens* lacking the *nos* gene cluster (166). MAG1 and MAG5 are affiliated with *Calidifontibacillus* species, among which two different *nos* gene clusters were found on the genome of *Calidifontibacillus oryziterrae* strain ZYK.

# Identification of clade III nosZ

A total of eleven *nosZ* genes were recovered from the seven high-quality MAGs. To demonstrate the *nosZ* of high-quality MAGs captures the comprehensive *nosZ* diversity in the twelve metagenomes, complete (> 500 bp) NosZ sequences were extracted from the coassembly (six in total) corresponding to each culture. Comparison of the assemblyand high-quality MAGs-derived NosZ (11 vs 11 genes) suggested the latter were sufficient to represent the overall NosZ diversity in the low pH N<sub>2</sub>O-reducing enrichment cultures. MAGs representing Desulfosporosinus (MAG3 and 4), Desulfitobacterium (MAG6 and 7) and *Paludifontibacillus* (MAG2) species were predicted to have a single NosZ sequence, whereas four and two different NosZ sequences were found on MAG1 and MAG5 belonging to *Calidifontibacillus*. Initial phylogenetic analyses placed NosZ sequences derived from acidic enrichment cultures to either clade I or clade II (Supplementary Figure 3-5). However, the deep-branching placement of MAG6- and MAG7-NosZ suggest they do not belong to either clade I or clade II, as the pairwise identity calculated for the two NosZ to other NosZ is no more than 30%. As such, the NosZ sequences of MAG6 and MAG7 were blasted against NCBI nr database, and the top 21 best hit sequences were included in following phylogenetic analyses (Figure 3-2).

Phylogenetic analyses including the reference-, acidic cultures-derived NosZ and the putative NosZ identified three independent NosZ lineages (Figure 3-2A), with the clade I and clade II being described previously (12, 22). MAG2 *nosZ*, associated with a clade I *nos* operon cluster (i.e., *nosZDFYL*, Figure 3-2 B), was clustered with clade I group. The *nosZ* genes of MAG1, 3, 4 and 5, followed immediately by a *nosB*, fell into clade II group. Interestingly, the multiple *nos* operon clusters of the same MAGs (i.e., MAG1 and 5) showed different structures (Figure 3-2 C). For instance, one of the four *nos* operon clusters from MAG1 lacks the *nosB* gene, encoding a transmembrane protein of unknown gene. Moreover, the four *nosZ* recovered from MAG1 shared no more than 72.5% amino acid identity (Supplementary Figure 3-6). These observations suggest *nosZ* diversification occurrs within individual population. The deep-branching *nosZ* genes recovered from MAG6 and 7, and the putative *nosZ*, most of which are recovered from metagenomic based studies, constituted a previously unknown clade and is hereafter

referred as clade III (Figure 3-2 D). Interestingly, clade III harbors MAGs representing *Nitrospira* bacterium, contradicting current recognition that *nosZ* genes are not found in *Nitrospira* (167). Most of the clade III NosZ (20 of 23 sequences) have a N-terminal Sec signal peptide, a feature shared with clade II NosZ. The clade III is coincided with the novel *nosZ* clade previously proposed in a metagenomic study of groundwater (168), yet the N<sub>2</sub>O reduction capacity of the *nosZ* requires experimental verification. The MAG6- and MAG7-*nosZ* and the putative *nosZ* genes all encode a N-terminal seven blade beta-propeller domain (Cu<sub>Z</sub> center) and a C-terminal cupredoxin domain (Cu<sub>A</sub> center), a characteristic found in both clade I and clade II *nosZ* (169). Among the clade I and II type NosZ, the Cuz-binding motifs associated with the first two histidine residues, DxHH, exhibited 100% conservation. Less conserved histidine residues were found in the Cuz-binding motifs of the *putative clade nosZ* (i.e., DxHx).

#### Structural and functional validation of clade III nosZ in enrichment cultures

Structure of the NosZ and closely related proteins (i.e., *nos* gene cluster) recovered from MAG6 were predicted using Alphafold2 with templates in PDB100 database as query structures. High-confidence structures were obtained for the corresponding protein sequences (Supplementary Figure 3-7), as assessed by the sequence coverage and predicted per-residue confidence measure (pLDDT). Among the eight protein sequences, NosZ and two hypothetical proteins were predicted to be periplasmic lipoprotein, whereas other proteins are cytoplasmic membrane-bounding or anchored. Specifically, a homodimer structure (pLDDT > 90) was predicted for NosZ, with two copper containing active sites present (Figure 3-3 A). Two conserved histidine and cysteine residues are associated with the catalytic site (i.e.,  $Cu_A$ ), whereas seven conserved histidine residues are associated with the catalytic site (Figure 3-3 B), but the physiology implication of this tyrosine in  $Cu_Z$  site remains to be demonstrated yet.

Microbial profiling of transfer cultures suggested the same clade III N<sub>2</sub>O reducers, MAG7, was consistently selected in culture WS1, C1982 and WS2

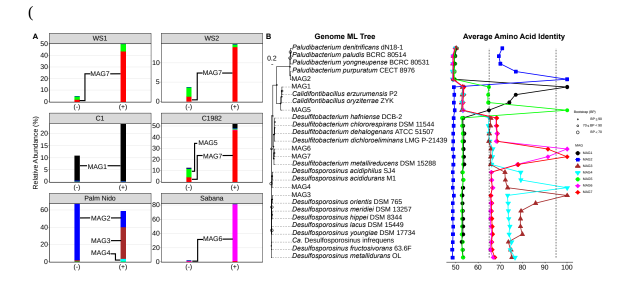


Figure **3-1** A); while MAG6 was only enriched in culture Sabana. As such, the culture WS2 and Sabana was maintained for further study. Following consecutive transfers were performed on culture Sabana and WS2, with which a 27<sup>th</sup> Sabana and 15<sup>th</sup> WS2 transfer cultures harboring the clade III N<sub>2</sub>O reducers were obtained. Subsequently triplicate cultures Sabana and WS2 growing in acidic medium consumed the initial dose of N<sub>2</sub>O (i.e., 416 µmol) over a 3-day incubation (Figure 3-3 C). In contrast, cultures incubated in circumneutral pH medium showed no N<sub>2</sub>O consumption over 10-day incubation. Consistent with N<sub>2</sub>O consumption, cells (measured by qPCR-targeting clade III *nosZ* gene) representing MAG6 in culture Sabana and MAG7 in culture WS2 increased by (25.7 ± 2.9) and (14.5 ± 2.8) folds, respectively (Table 3-5). Growth yields of (8.3 ± 1.1) x 10<sup>8</sup> and (3.2 ± 0.9) x 10<sup>10</sup> cells mmol N<sub>2</sub>O<sup>-1</sup> were determined for MAG6 and MAG7 in culture Sabana and WS2, respectively. In control cultures omitting N<sub>2</sub>O, increase in cells representing MAG6 and MAG7 was not substantial (less than 1-fold increase), suggesting their growth is N<sub>2</sub>O-dependent under the conditions tested. No growth of MAG6 and MAG7 was observed in pH 7 cultures in the presence of N<sub>2</sub>O, demonstrating

the N<sub>2</sub>O reduction activity of the clade III N<sub>2</sub>O reducers (i.e., MAG6 and MAG7) is limited to acidic pH.

Comparative Reverse Transcriptional qPCR (RT-qPCR) analyses were performed on cultures (Sabana and WS2) that were actively consuming N<sub>2</sub>O, with cultures omitting N<sub>2</sub>O served as control. The transcribed *nosZ* in cultures Sabana and WS2 received N<sub>2</sub>O reached (44.7 ± 11.4) and (107 ± 51.8) copies ng RNA<sup>-1</sup>, respectively; whereas the transcribed *nosZ* decreased to (9.65 ± 4.38) and (24.6 ± 20.1) copies ng RNA<sup>-1</sup> in cultures Sabana and WS2 omitting N<sub>2</sub>O, respectively. Significance analyses (p < 0.05) suggest that the *nosZ* transcriptions were enhanced by (4.91 ± 0.9) and (5.13 ± 1.64) folds in cultures Sabana and WS2 in the presence of N<sub>2</sub>O, respectively (Fig. 4, inset graph). The *nosZ* transcripts were lower than (2.9 ± 3.33) copies ng RNA<sup>-1</sup> in all cultures received N<sub>2</sub>O incubated at pH 7, an observation consistent to the ceased N<sub>2</sub>O reduction activity. In line with RT-qPCR results, the amount of assembled clade III NosZ (measured by metaproteomic) increased by (1.4 ± 1.5) and (2.3 ± 0.4) folds in Sabana and WS2 cultures being significant (p < 0.05).

### **3.4 Discussion**

The present study challenged the decade-lasting view that NosZ are divided into clade I and II lineages. The discovery of N<sub>2</sub>O-dependent respiratory growth of clade III N<sub>2</sub>O reducers advances the understanding of *nosZ* diversity involved in N<sub>2</sub>O reduction, and has far-reaching implication in underexplored soil microbial N<sub>2</sub>O reduction potentials. Current approaches (e.g., sequencing and qPCR) estimating soil N<sub>2</sub>O reduction potential only target clade I and clade II NosZ (170), leaving the clade III NosZ out of consideration, tentatively due to the failure in identification and cultivation of members of clade III lineage. A metagenome-based study suggested the putative clade III N<sub>2</sub>O reducers (i.e., *Nitrospinota* and *Nitrospira* bacterium) are present in groundwater and the NosZ was transcriptional active (168); however, the N<sub>2</sub>O reduction capacity of members in clade III was still not resolved. We demonstrated the growth supporting N<sub>2</sub>O reduction

of two population (MAG6 and MAG7) have a clade III *nosZ* gene, which provides experimental evidence the clade III NosZ is of functionality. The function of clade III NosZ was validated with multiple lines of evidence: Comparative growth assays of culture WS2 and Sabana revealed the N<sub>2</sub>O-dependent respiratory growth; AlphaFold prediction of clade III NosZ showed a homo-dimeric structure containing the electron transfer Cu<sub>A</sub> site and catalytic Cu<sub>Z</sub> site; RT-qPCR and proteomics analyses suggest NosZ expression depends on N<sub>2</sub>O amendment. Although the in-situ detection in soil samples (LEF soils) with qPCR assays targeting the clade III *nosZ* (MAG6 and MAG7-NosZ) gene was unsuccessful, enrichment of the clade III N<sub>2</sub>O reducers from distant biomes (CFF and LEF) illustrated they represent rare biosphere but readily responsive to N<sub>2</sub>O. Distribution of the clade III NosZ warrants further investigation of environmental samples, which would possibly leverage the genomic potential of N<sub>2</sub>O reduction in various ecosystems.

In this study, we sampled soils from circumneutral ( $pH \sim 7$ ) sub-tropical agricultural (i.e., CFF) (171) and acidic (4.5 < pH < 5.0) tropical forest sites (i.e., LEF) (43) that represent contrasting soil types. We showed that continuous cultivation and enrichment of microorganisms directly using exogenous N2O under acidic conditions is a successful strategy to obtain acidic N<sub>2</sub>O respiring microorganisms and characterize them. The unusual cultivation strategy (alternate carbon source [fumarate and succinate] plus H<sub>2</sub>) resulted in the enrichment of seven previously unknown nosZ-carrying bacteria, all of which belong to the genus (i.e., Calidifontibacillus, Desulfosporosinus and Desulfitobacterium) have not been implicated in N<sub>2</sub>O reduction yet (i.e., lack experimental validation). In contrast, cultivation and enrichment of the same soils (i.e., Sabana) performed at pH 7 derived well-studied microorganisms (i.e., Azospira) (110, 172), representing most cells (accounting > 99% of the 16S rRNA amplicons) in the cultures following less than fifteen consecutive transfers; however, those pH 7 transfer cultures cannot consume N<sub>2</sub>O in pH 4.5 medium. This finding is consistent to generalized observation: N2O reducers obtained under circumneutral conditions are incapable of low pH N<sub>2</sub>O reduction (36, 38, 39), and maintaining acidic pH is crucial to cultivate acidic

N<sub>2</sub>O reducers (Guang, et al., 2023). Interestingly, the acidophilic '*Ca.* Desulfosporosinus nitrousreducens' strain PR in a consortium consisted of two population, derived from El Verde Field station, was also enriched in the low pH N<sub>2</sub>O-reducing culture derived from Palm Nido soil. The two sites, El Verde field station and Palm Nido, have contrasting soil physicochemical parameters such as elevation (453 m for EV and 634 m for PN) and represent different life zones (43). A coincidental observation is the clade III *nosZ*-carrying MAG7 was consistently selected in cultures enriched from the soils from three separate sites in CFF. The same population could be readily enriched from different soils lend strong experimental credibility to the hypothesis that low pH N<sub>2</sub>O reducing microorganisms are more widely distributed than first thought (Guang, et al., 2023).

pH has been considered the major factor limiting  $N_2O$  reduction activity in acidic biomes (e.g., soils and oceans) (37, 173), and was explained by microbial failure to assemble functional NosZ under acidic conditions (pH < 6) (40). The generalized paradigm was recently challenged by the acidophilic  $N_2O$ -reducing '*Ca*. Desulfosporosinus nitrousreducens' strain PR with a clade II nos operon cluster (Guang, et al., 2023) and the deep-sea isolate, Niratiruptor labii HRV44 (105). The 'Ca. Desulfosporosinus nitrous reducens' strain PR was enriched from the acidic tropical forest (pH < 5), thus raised a question: Does acidophilic N<sub>2</sub>O reducers exist in circumneutral soils? The successful enrichment of N2O-reducing cultures under acidic conditions from circumneutral CFF soils suggest that low pH N<sub>2</sub>O reducers are not limited to acidic soils. The naturally acidic sites on the Earth are known to thrive a variety of acidophilic microorganisms (45), whereas the circumneutral soil are overlooked source of acidophiles. Soils are complex matrixes with high pH heterogeneity, and pH varies by as much as 1.3 pH among samples less than 50 cm apart (46). The microbial metabolism could have led to more contrasting pH at finer scale: the microscale varied by 2 pH in spots localized closely (less than 5 mm) following straw patch amendment (47). Obviously, acidic microsites could occur in circumneutral CFF soils and harbor the acidophilic N<sub>2</sub>O-reducing microorganisms. This finding has far-reaching implications in

predicting N<sub>2</sub>O emissions as the acidophilic N<sub>2</sub>O reducing microorganisms would thrive to cope with increased N<sub>2</sub>O emissions associated with soil acidification.

Prior to this study, N<sub>2</sub>O reducers were known to include clade I and II, with the former mostly possess all genes involved in denitrification of  $NO_3^-$  to  $N_2$  (23). Physiology and genomic features distinguish clade I versus clade II NosZ, suggesting niche differentiation may exist within and between the two clades. The NosZ is a periplasmic enzyme, which must be transported across inner membrane. Clade I NosZ are translocated in a folded state using the Twi-arginine pathway (Tat-pathway) whereas clade II NosZ are translocated as unfolded protein via the general secretion route (Secpathway) (12). The translocation pathway may influence the pH range of clade I and II NosZ. Metagenomic investigation of acidic LEF soils suggest that clade II nosZ dominated the *nosZ* gene pool (> 10% of total *nosZ* reads) (43), and pH selected for different clade II N<sub>2</sub>O reducers in the LEF soils-derived microcosms (44). Coincidently, N<sub>2</sub>O consumption activity in acidic microcosms increased proportionally with the clade II nosZ abundance (5). In line with prior studies, our comparative metagenome analyses with soils-free cultures grown with/without N2O amendment identifies that all active N2O reducing microorganisms possess clade II or clade III (with Sec peptide) nosZ, suggesting low pH N<sub>2</sub>O reduction may be a shared feature in clade II and clade III N<sub>2</sub>O reducers. Growth of the clade I nosZ-carrying microorganism (MAG2, Paludibacterium) in culture Palm was N<sub>2</sub>O-independent at pH 4.5, suggesting its growth depends on fumarate fermentation. This observation is consistent to the failure of N<sub>2</sub>O reduction by Ensifer meliloti (clade I) at pH 6, whereas oxygenic respiration was not influenced at pH 6 (39). A possible explanation is the oxygen respiration and fumarate fermentation occur in cytoplasm, where the pH was maintained relatively stable via homeostasis; however,  $N_2O$ reduction occurs in periplasm, where the pH is more impacted by external environments.

Genomic and Phenotypic differences have been documented in clade I and clade II  $N_2O$  reducers, implicating niche differentiation existing between the two evolutionary distinct lineages (12). For example, the co-occurrence rate of *nosZ* and *nirK/nirS* was much higher in clade I (83%) than clade II  $N_2O$  reducers (49%) (23), suggesting clade II  $N_2O$ 

reducers are more likely to be obligate  $N_2O$  sink. Furthermore, kinetic study demonstrated that clade II and clade I N<sub>2</sub>O reducers are more likely to acclimate oligotrophic and eutrophic conditions, respectively (101). Therefore, oligotrophic conditions rich with  $N_2O$  are proposed to favor growth of clade II over clade I  $N_2O$ reducers. Indeed, laboratory enrichment documented that clade II outweigh clade I N2O reducers when  $N_2O$  was the sole electron acceptor, while clade I  $N_2O$  reducers were more abundant with  $NO_3^-$  as electron acceptor (174). In contrast, most N<sub>2</sub>O reducers isolated on eutrophic medium (i.e., Tryptone-Soy Agar) were close relatives to clade I associated taxa and produced  $N_2O$  via denitrification (116). In current research, clade II  $N_2O$ reducers acclimated cultures growing in oligotrophic medium (containing 1% v/v TSB) in response to N<sub>2</sub>O additives. The enriched clade II N<sub>2</sub>O-reducing organisms are of concern in bioaugmentation perspective as they are predicted to be non-denitrifiers (i.e., lack nirS/nirK genes) and are potential N<sub>2</sub>O scavengers in soils. Soils become net N<sub>2</sub>O sources when  $N_2O$ -producing rates exceed  $N_2O$ -reducing rates (154), while soils turn to be net N<sub>2</sub>O sink when N<sub>2</sub>O-consuming organisms outcompete N<sub>2</sub>O-producing organisms (175). Acidic soils have long been considered major N<sub>2</sub>O emitters as acidic conditions favor N<sub>2</sub>O formation over its consumption in denitrifying organisms (38, 40). As such, the non-denitrifying organisms reducing N<sub>2</sub>O under acidic conditions offer a golden opportunity to shift acidic soils into net  $N_2O$  sinks if their growths are favored over denitrifying organisms.

The N<sub>2</sub>O reduction was hypothesized an ancient trait that appeared before or after the 'great oxygen event (GOE)', 2.8-2.5 Gyr before present (23). Phylogenetic analysis indicated *nosZ* has evolved predominantly via vertical inheritance and some instances of horizontal gene transfer (HGT) (176, 177). Following the definition of clade II lineage, phylogenetic diversity of *nosZ* and microbial taxa have been greatly expanded (12). In contrast to the narrow distribution of clade I *nosZ* among *Proteobacteria* phyla, the clade II *nosZ* appeared to be shared by evolutionary distant phyla (e.g., *Firmicutes* and *Chloroflexi*) (22). Furthermore, the increasing clade II *nosZ* diversity indicated its potential in mitigating N<sub>2</sub>O emissions from diverse soil ecosystems. For example, *nosZ* 

has not been detected from the genomes of *Actinobacteria* or *Acidobacteria* that are associated with low resource availability and low pH conditions in terrestrial ecosystems during the past decades (23). However, in-depth characterization of microbial communities in tundra soils identified *nosZ*-carrying organisms affiliated with *Acidobacteria*, associated with acidic conditions (24), implying its role in N<sub>2</sub>O consumption was underexplored.

The clade II and III nosZ genes acquired in current research expanded the nosZ diversity associated with low pH N<sub>2</sub>O reduction. Organisms carry clade II nosZ genes are classified as Desulfosporosinus, Calidifontibacillus and clade III nosZ-carrying organisms are affiliated with *Desulfitobacterium*. In consistent to previous studies (22, 178), the nosZ phylogeny of Calidifontibacillus and Bacillales showed congruence to the hierarchical taxonomic classification, suggesting the *nosZ* genes were attained via vertical heritance events within these taxa. Incongruence between nosZ and taxonomic phylogeny was observed for *Desulfitobacterium* and *Desulfosporosinus* population, suggesting higher diversification within the taxa. The deep-branching subclade comprising Desulfitobacterium nosZ was unexpected, as the nosZ of Desulfitobacterium are defined as clade II. In supporting the evolutionary difference, the multiple sequence alignment analysis identified partially conserved histidine ligands of the Cuz center within the clade. In addition, the NosD protein (encoded by nosD), delivering Cu<sup>2+</sup> to Cu<sub>z</sub> center in Pseudomonas stutzeri (clade I lineage) (169), was absent in the deep-branching Desulfitobacterium nosZ clade. However, the assembly machinery may differ between clade I, II and III N<sub>2</sub>OR as the folding processes occur in different cell compartments (cytoplasm vs. periplasm) (23). Therefore, the role of nosD in the deep-branching nosZ cluster remains questionable. Taken together, the deep-branching nosZ clade was proposed a putative novel nosZ lineage specialized in low pH N<sub>2</sub>O reduction.

# Appendix

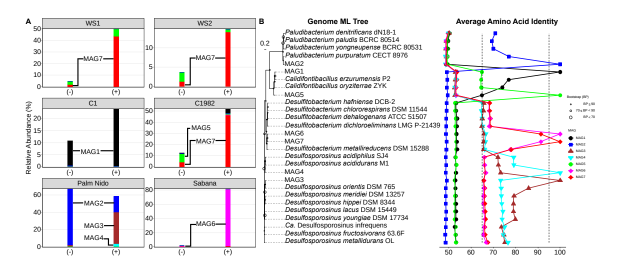


Figure 3-1. Novel microbial guilds implicated in N2O reduction. (A) Distribution of putative N<sub>2</sub>O-reducing population in transfer cultures with (+) or without (-) N<sub>2</sub>O amendment. The relative abundance (% of the total metagenome fragments) of MAGs was calculated using CoverM (https://github.com/wwood/CoverM). Taxonomy was assigned to MAGs using GTDB-TK (91). Displayed are MAGs predicted to have at least one nosZ gene based on Prokka annotation (137). The comprehensive microbial community profiled with the 43 high-quality MAGs are summarized in Fig. S4. (B). Phylogenomic analyses of nosZ-carrying MAGs recovered from low pH N<sub>2</sub>O-reducing cultures. The tree topology was obtained from the Maximum Likelihood (ML) tree by using RAxML-NG (92), inferred from a 120 conserved marker genes alignment of 20 complete genomes (selected based on GTDB-TK results) (91) and 7 MAGs. The top scale bar indicates 0.2 amino acid substitution rates per site. Sizes of open circles alongside the branches of each sublineage indicate bootstrap percentages obtained by RAxML-NG. The pairwise genome-wide AAI calculated for each MAGs are shown on the right, with the scale displayed on the bottom. Values for each comparison for a given MAGs are connected by lines, and the comparison to self is indicated as 100%. Pair-wise AAI values of 65-95% (i.e., data points fall between the dashed lines) are indicative of

species boundary (179). The color legend corresponds to the bar and line colors in panel (A) and (B).

# Figure 3-2. Phylogenetic trees showing the placement of acidic cultures derived NosZ (green) among the reference NosZ (black) and putative NosZ (red). (A)

Unrooted phylogenetic tree of NosZ showing three independent clades (i.e., I, II and III). The tree topology was inferred from RAxML-NG. Bootstrap values are displayed as white (> 90%), grey (70 < BP < 90) and black circles (< 70%). (B) Phylogenetic tree showing clade I NosZ with branches belonging to clade II and clade III collapsed. (C) Phylogenetic tree showing clade II NosZ with branches belonging to clade I and clade III collapsed. (D) Phylogenetic tree showing clade III NosZ with branches belonging to clade I and clade III collapsed. Of note, the MAG4 *nos* gene cluster is identical to '*Ca*. Desulfosporosinus nitrousreducens' in a co-culture reported in a prior study (Guang, et al., 2023). Accessory genes associated with *nosZ* are displayed as oriented arrows and gene designation is represented by color legend in panel D.

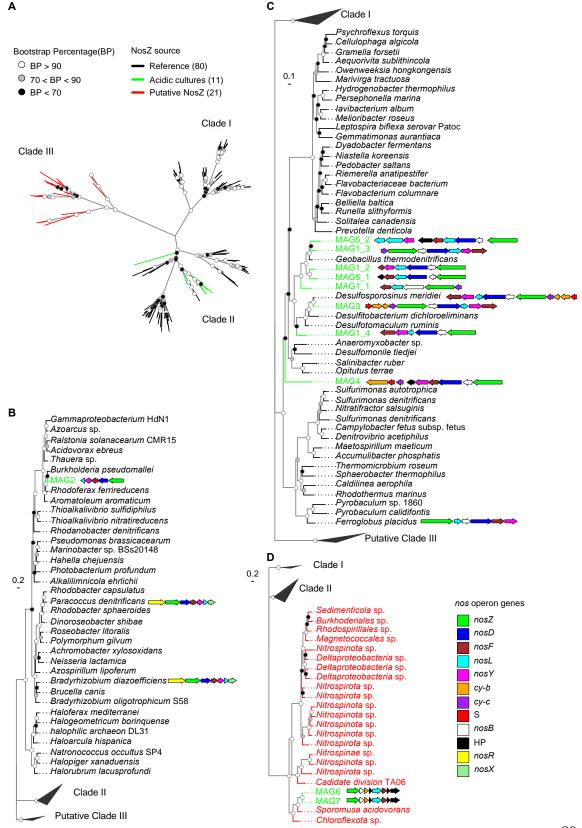
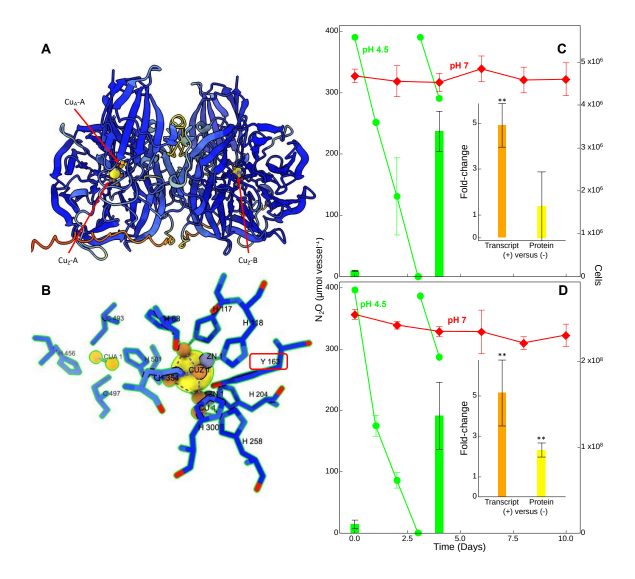
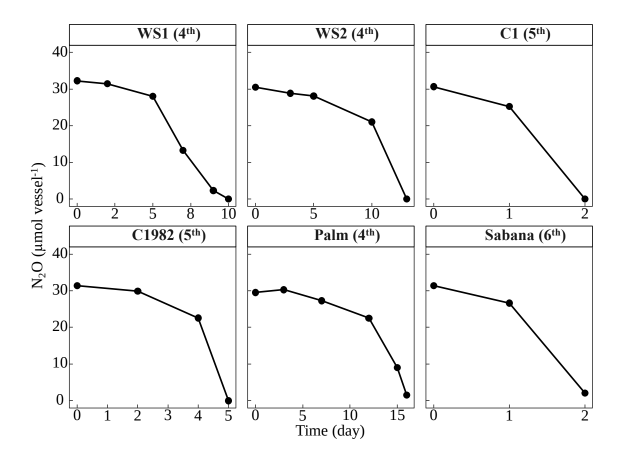
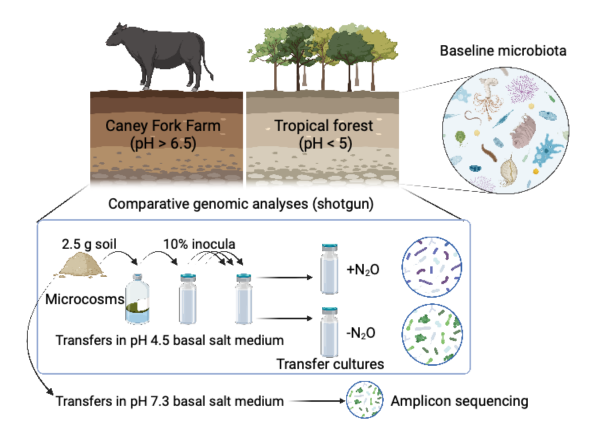


Figure 3-3. Structural and functional validation of clade III NosZ. (A). The NosZ enzyme is a homodimer in head-to-tail orientation with the tetranuclear Cuz active site located in the N-terminal, seven-bladed β-propeller domain and the binuclear Cu<sub>A</sub> site in the C-terminal cupredoxin domain. The Cu<sub>A</sub>-Cu<sub>Z</sub> distance within one monomer is around 42.5 Å (i.e., Cu<sub>A</sub>-A to Cu<sub>Z</sub>-A) and thus too large for efficient electron transfer. Instead, the Cu<sub>A</sub> site of one monomer is at a distance of only 10.5 Å from Cu<sub>Z</sub> in the other monomer (Cu<sub>A</sub>-A to Cu<sub>Z</sub>-B) (B). Conserved residues associated with Cu<sub>A</sub> and Cu<sub>Z</sub> sites. Two histidine and cysteine residues are conserved in Cu<sub>A</sub> site, and seven histidine residues are conserved in Cuz site. The Y163 residue enclosed in red is an unexpected (lacking in clade I and clade II NosZ) tyrosine associated with Cuz site. (C) Comparative growth assays performed with 27<sup>th</sup> Sabana and (D) 15<sup>th</sup> WS2 transfers cultures. Green and red lines indicate  $N_2O$  reduction performance in different pH (4.5 and 7) medium. Green bars represent the cells (pH 4.5) enumerated with specific primers targeting the clade III Desulftobacterium nosZ. Fold changes of nosZ transcripts (orange bars) and assembled NosZ protein (vellow bars) are calculated based on corresponding cultures grown in pH 4.5 medium with (+) and without (-) N<sub>2</sub>O amendment. Of note, growth of clade III N<sub>2</sub>O reducers was not substantial in cultures without N<sub>2</sub>O. Data represent averages of triplicate incubations and error bars represent standard deviations (n=3). Error bars are not shown when smaller than the symbol.





**Supplementary Figure 3-1. Low pH N<sub>2</sub>O reduction of transfer cultures derived from farm and tropical soils.** Numbers inside parenthesis indicate the transfer generations. For example, panel labeled as WS1 (4<sup>th</sup>) showed N<sub>2</sub>O data collected from 4<sup>th</sup> WS1 transfer culture. Data were collected from single experiment.

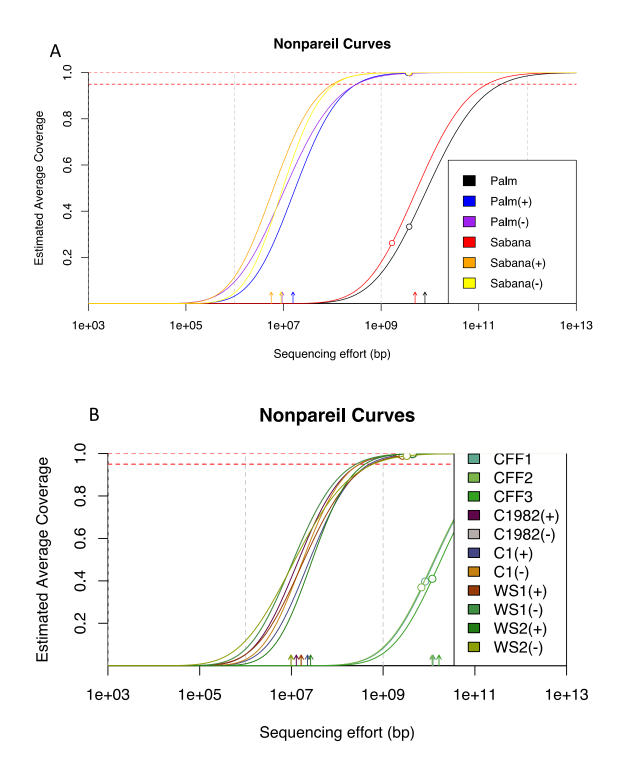


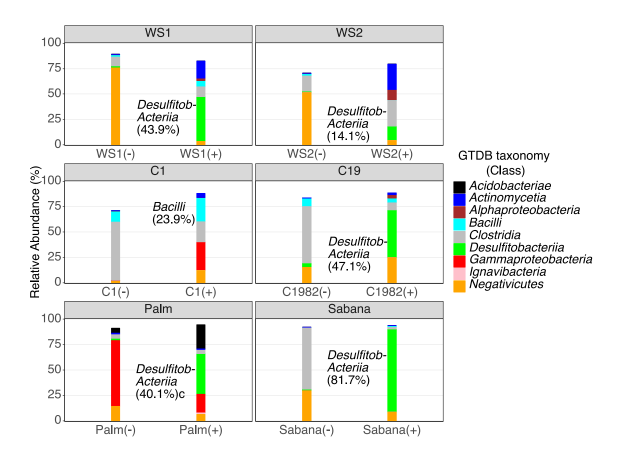
# Supplementary Figure 3-2. Schematic of establishing low pH N<sub>2</sub>O reducing cultures.

Shotgun metagenomic sequencing was applied to original Caney Fork Farm and Puerto Rico Soils to study the baseline microbiota. Soil samples from Caney Fork Farm and Puerto Rico Forest were subsequently used to establish pH 4.5 and 7 soil microcosms. Consecutive transfers were performed in pH 4.5 and 7 basal salt medium and derived solids-free cultures. Comparative shotgun metagenomic analyses were conducted with corresponding pH 4.5 cultures with or without N<sub>2</sub>O substrate. Amplicon sequencing targeting 16S V3-V4 regions was performed on cultures transferred in pH 7 basal salt medium.

# Supplementary Figure 3-3. Nonpareil analysis of soil- and enrichment cultures'

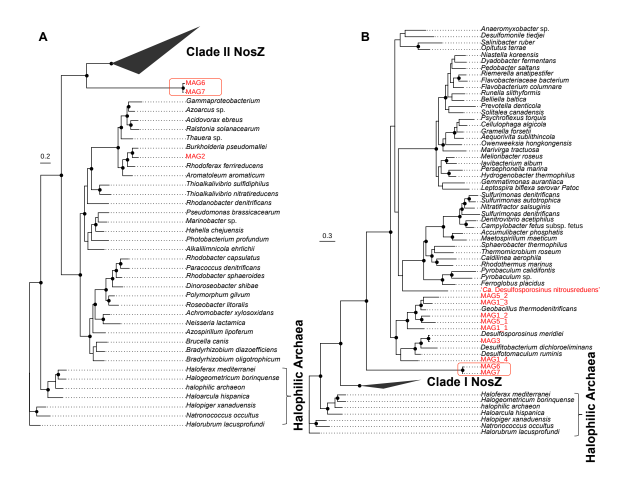
**metagenomes.** (A) Nonpareil curves of PR soil- and derived enrichment cultures. (B) Nonpareil curves of CFF soil- and derived enrichment cultures. Open circles represent the estimated average coverage of the datasets obtained and projections based on model fitting to reach 95% and 99% coverage are indicated (horizontal dashed lines). The arrows at the bottom represent sequencing effort required to achieve 50% coverage.





# Supplementary Figure 3-4. Microbial community constructed with high-quality

**MAGs.** Relative abundance of MAGs in each metagenome was calculated with CoverM. Taxonomy was assigned to MAGs using GTDB-TK. Microbial community was displayed at class level with ggplot package. T axa highlighted with text are population selectively growing in the presence of N<sub>2</sub>O.



**Supplementary Figure 3-5. Preliminary phylogenetic analyses allude to the clade III** *nosZ* genes recovered from acidic N<sub>2</sub>O-reducing cultures. (A) clade I and (B) clade II *nosZ* are shown separately for readers' convenience. The branch labels in black indicate the 80 referecen *nosZ* genes, whereas branch labels in red indicate the *nosZ* genes recovered from acidic N<sub>2</sub>O-reducing cultures. Bifurcation with bootstrap values over 75% are indicated with closed circle symbols. Tree scales indicate amino acid substitution rates. The *nosZ* genes (MAG6 and 7) not cluster with either clade I or clade II are enclosed in red.

NA	64.5	64.2	58.0	31.8	55.7	42.2	60.9	59.9	25.0	24.7	MAG1_1
64.5	NA	72.3	54.6	31.4	52.8	41.7	75.0	66.9	23.6	24.1	MAG1_2
64.2	72.3	NA	56.0	31.6	54.2	42.2	69.5	72.0	23.9	24.2	MAG1_3
58.0	54.6	56.0	NA	32.1	61.1	46.0	54.8	55.9	25.9	25.6	MAG1_4
31.8	31.4	31.6	32.1	NA	32.2	27.5	29.9	29.7	23.1	23.1	MAG2
55.7	52.8	54.2	61.1	32.2	NA	42.9	52.3	53.8	24.8	24.7	MAG3
42.2	41.7	42.2	46.0	27.5	42.9	NA	44.7	41.0	21.4	21.5	MAG4
60.9	75.0	69.5	54.8	29.9	52.3	44.7	NA	67.8	23.5	23.2	MAG5_1
59.9	66.9	72.0	55.9	29.7	53.8	41.0	67.8	NA	23.5	23.5	MAG5_2
25.0	23.6	23.9	25.9	23.1	24.8	21.4	23.5	23.5	NA	97.0	MAG6
24.7	24.1	24.2	25.6	23.1	24.7	21.5	23.2	23.5	97.0	NA	MAG7
MAG1_1	MAG1_2	MAG1_3	MAG1_4	MAG2	MAG3	MAG4	MAG5_1	MAG5_2	MAG6	MAG7	

Supplementary Figure 3-6. Heatmap showing amino acid identity between the NosZ recovered from high-quality MAGs. Numbers inside cells represent the pairwise sequence identity (%) between two NosZ. Four and two different NosZ are found on MAG1 and MAG5, respectively. The MAG6 and MAG7 NosZ share no more than 30% to reference NosZ shown as black labels in Fig. S5.

# AlphaFold models

	NosZ	NosB	Cyt-B	H122	NosL	NosF	H125	H126
	N <sub>2</sub> O reductase	Part of ABC transporter?	Cytochrome	Met-rich β-barrel lipoprotein	Met-rich ABC transporter	Part of ABC transporter	Unknown	β propeller
	periplasmic lipoprotein	cytoplasmic membrane	cytoplasmic membrane	periplasmic lipoprotein	cytoplasmic membrane	cytoplasm	cytoplasmic membrane- anchored	periplasmic lipoprotein
	dimer	monomer	monomer	monomer	monomer?	dimer	monomer	monomer
			Section of the sectio				and the second s	E JUI
monomer		88.9	77.4	88.4	83.9		80.8	84.6
dimer	91.8 (0.92, 0.91)	83.2 (0.8, 0.78)			72.3 (0.52, 0.443)	88.5 (0.77, 0.64)		
								DLDDT DLDDT (pTM, ipTM)

**Supplementary Figure 3-7. AlfphaFold2 predicted structures of proteins located closely to NosZ.** Signalp 6.0 was used to identify and remove signal peptides prior to structure prediction. The corresponding membrane orientations of proteins were predicted using MembraneFold.

Soil Samples (Location <sup>a</sup> )	Longitude	Latitude	pH <sup>b</sup>	Tree <sup>c</sup>
Sabana (LEF)	-65.7296575	18.317758	4.72	Dacryodes excelsa
Palm Nido (LEF)	-65.798597	18.295042	4.97	Sierra Palm
WS1 (CFF)	-85.910731	36.243253	6.74	Winter-East-AA-005
WS2 (CFF)	-85.910401	36.243113	6.87	Winter-East-AA-013
C1 (CFF)	-85.909630	36.241153	6.96	Winter-East-AA-061
C1982 (CFF)	-85.909575	36.241508	6.98	Winter-East-AA-053

 Table 3-1. Metadata associated with soils used for enrichment procedure.

<sup>a</sup> Location: LEF refers to Luquillo Experimental Forest located in Puerto Rico; CFF refers to Caney Fork Farm located in Carthage, Tennessee. <sup>b</sup> pH of LEF soils were reported in a prior study (43) and CFF soil pH was measured by mixing 2.5 gram fresh soils with 10 mL deionized water (1:4) in this study. <sup>c</sup> The CFF soils are associated with labelled trees aiming for long-term monitoring; WS1 and WS2 are associated with trees that inoculated with Symbionts to Advance Food and Energy (SAFE) fertilizer whereas C1 and C1982 are soils close to trees not receiving any fertilizer. Sabana and Palm Nido sites is dominated by Dacryodes excelsa and Sierra Palm, respectively.

Culture <sup>a</sup>	Carbon source <sup>b</sup>	Electron acceptor	Electron donor
wigh (cth)		4.16 mM N <sub>2</sub> O	4.16 mM H <sub>2</sub>
WS1 (6 <sup>th</sup> )	5 mM Fumarate	(nominal)	(nominal)
WGO (cth)	5 mM Essente	4.16 mM N <sub>2</sub> O	4.16 mM H <sub>2</sub>
WS2 (6 <sup>th</sup> )	5 mM Fumarate	(nominal)	(nominal)
C1 (7 <sup>th</sup> )	5 mM From outo	4.16 mM N <sub>2</sub> O	4.16 mM H <sub>2</sub>
C1 (7 <sup>th</sup> )	5 mM Fumarate	(nominal)	(nominal)
C1082 (7 <sup>th</sup> )	5 mM From outo	4.16 mM N <sub>2</sub> O	4.16 mM H <sub>2</sub>
C1982 (7 <sup>th</sup> )	5 mM Fumarate	(nominal)	(nominal)
<b>C</b> -hang (0 <sup>th</sup> )	5 mM Essente	4.16 mM N <sub>2</sub> O	4.16 mM H <sub>2</sub>
Sabana (9 <sup>th</sup> )	5 mM Fumarate	(nominal)	(nominal)
Datus Mida (Oth)	5 m M Smark	4.16 mM N <sub>2</sub> O	4.16 mM H <sub>2</sub>
Palm Nido (9 <sup>th</sup> )	5 mM Succinate	(nominal)	(nominal)

**Table 3-2.** Substrates provided to the N<sub>2</sub>O-reducing cultures during the enrichment process.

<sup>a</sup> Numbers inside parenthesis indicate the corresponding cultures were subjected to shotgun sequencing efforts. <sup>b</sup> In addition to fumarate/succinate, 1% (v/v) Tryptone-Soy Broth was augmented as growth nutrient.

MAG ID <sup>b</sup>	Completeness	Contamination	Closest relative <sup>c</sup>	AAI d
C1982- MAG1 <sup>e</sup>	92.2	5.7	Calidifontibacillus erzurumensis	76.9
Palm- MAG2 <sup>e</sup>	94.4	0.9	Paludibacterium purpuratum	77.0
Palm- MAG3 <sup>e</sup>	97.0	1.6	Desulfosporosinus orientis	85.8
Palm- MAG4 <sup>e</sup>	99.5	6.0	Desulfosporosinus acididurans	79.2
C1982- MAG5 <sup>e</sup>	95.6	2.4	Calidifontibacillus erzurumensis	64.6
Sabana- MAG6 <sup>e</sup>	98.9	2.6	Desulfitobacterium_metallireducens	77.7
WS2- MAG7 <sup>e</sup>	98.9	2.0	Desulfitobacterium_metallireducens	78.0
Palm- MAG8	95.0	0.6	Clostridium celerecrescens	92.0
Sabana- MAG9	99.4	0.6	Clostridium sphenoides	97.4
Sabana- MAG10	95.4	3.7	Variimorphobacter saccharofermentans	64.9

 Table 3-3. Analyses of high-quality MAGs using the Microbial Genome Atlas (MiGA) <sup>a</sup>.

Table 3-3. Continued

MAG ID <sup>b</sup>	Completeness	Contamination	Closest relative <sup>c</sup>	AAI <sup>d</sup>
Palm- MAG11	100	9.0	Pelosinus fermentans	75.4
C1-MAG12	97.9	1.4	Sinanaerobacter chloroacetimidivorans	68.0
Sabana- MAG13	85.2	5.8	Pseudoclostridium thermosuccinogenes	55.0
Sabana- MAG14	98.6	5.0	Ectobacillus_panaciterrae	65.0
C1-MAG15	97.2	3.1	Clostridium homopropionicum	84.2
C1-MAG16	84.8	2.8	Clostridium magnum	83.7
Palm- MAG17	99.9	9.08	Clostridium aciditolerans	89.3
WS2- MAG18	85.1	2.3	Clostridium thailandense	92.2
Sabana- MAG19	89.7	8.9	Clostridium rhizosphaerae	62.9
WS2- MAG20	97.4	1.1	Clostridium swellfunianum	63.3

Table 3-3. Continued

MAG ID <sup>b</sup>	Completeness Contamination		Closest relative <sup>c</sup>	AAI <sup>d</sup>
C1982- MAG21	97.2	6.6	Clostridium rhizosphaerae	93.2
WS2- MAG22	89.7	22.6	Clostridium swellfunianum	63.3
WS2- MAG23	96.6	2.8	Clostridium rhizosphaerae	63.7
C1-MAG24	99.3	19.5	Clostridium rhizosphaerae	64.3
WS2- MAG25	90.8	6.8	Clostridium swellfunianum	60.1
C1-MAG26	98.0	0.7	Clostridium homopropionicum	84.2
Palm- MAG27	98.7	1.0	Caproiciproducens galactitolivorans	67.3
WS2- MAG28	98.0	4.7	Dendrosporobacter quercicolus	58.3
Palm- MAG29	97.8	0	Paracidobacterium acidisoli	50.8
WS2- MAG30	99.4	1.4	Dendrosporobacter quercicolus	58.7

Table 3-3. Continued

MAG ID <sup>b</sup>	Completeness Contamination		Closest relative <sup>c</sup>	AAI <sup>d</sup>
C1-MAG31	97.2	0.5	Propionicimonas paludicola	66.3
Palm- MAG32	97.2	1.8	Propionicimonas paludicola	63.9
WS1- MAG33	99.5	1.2	Microbacterium algeriense	83.9
WS1- MAG34	98.8	0	Microbacterium aurum	95.8
WS2- MAG35	99.2	0.3	Cellulomonas fimi	63.2
WS1- MAG36	99.1	0.8	Sphingomonas echinoides	95.0
WS2- MAG37	98.1	5.0	Sphingomonas echinoides	97.3
C1-MAG38	100	1.4	Pseudomonas nicosulfuronedens	97.2
Palm- MAG39	84.6	10.7	Oscillibacter ruminantium	71.0
Palm- MAG40	85.9	1.36	Muriventricola aceti	66.1

Table 3-3. Continued

MAG ID <sup>b</sup>	Completeness	Contamination	Closest relative <sup>c</sup>	AAI <sup>d</sup>
C1-MAG41	98.0	3.02	Ruminiclostridium hungatei	66.8
Palm- MAG42	90.2	7.0	Sporolituus thermophilus	57.1
Palm- MAG43	98.0	1.1	Melioribacter roseus	50.7

<sup>a</sup> MAG completeness and contamination were calculated using CheckM package. <sup>b</sup> Prefix (e.g., Sabana and Palm) of MAG ID indicates the metagenomes from where a MAG was constructed. <sup>c</sup> Closest relatives and associated <sup>d</sup> Average Amino Acid Identity (AAI) calculated using MiGA. <sup>e</sup> MAGs carry at least one *nosZ* gene.

MAG id	Contigs	Size (Mbp)	GC content (%)	CDS <sup>b</sup>	tRNA <sup>c</sup>	rRNA d
MAG1	69	4.2	38.0	4163	53	15
MAG2	85	3.7	62.1	3461	59	4
MAG3	106	5.3	46.4	4926	70	12
MAG4 <sup>a</sup>	94 (84)	5.6 (5.6)	44.7 (44.8)	5239 (5231)	41 (40)	7 (8)
MAG5	31	4.14	38.3	4052	72	5
MAG6	105	4.5	41.3	4394	40	5
MAG7	45	4.14	41.0	3976	43	5

Table 3-4. Statistics of the high-quality *nosZ*-carrying MAGs.

<sup>a</sup> MAG4 is identical to the genome of '*Ca*. Desulfosporosinus nitrousreducens' strain PR and numbers inside parenthesis are the genome characteristics reported in prior study (Guang, et al., 2023). <sup>b</sup> CDS: coding sequences. <sup>c</sup> tRNA: transfer RNA. <sup>d</sup> rRNA: ribosomal RNA.

Population	N <sub>2</sub> O supply	Medium pH	Cells (mL <sup>-1</sup> ) <sup>b</sup>	Cells (mL <sup>-1</sup> ) <sup>c</sup>
	Yes	4.5	1.3 (0.1) x 10 <sup>5</sup>	3.4 (0.5) x 10 <sup>6</sup>
MAG6 (Sabana)	No	4.5	3.0 (0.6) x 10 <sup>4</sup>	4.7 (0.8) x 10 <sup>4</sup>
	Yes	7	7.5 (2.7) x 10 <sup>3</sup>	1.5 (0.6) x 10 <sup>4</sup>
	Yes	4.5	1.0 (0.5) x 10 <sup>7</sup>	1.4 (0.4) x 10 <sup>8</sup>
MAG7 (WS2)	No	4.5	1.5 (0.3) x 10 <sup>6</sup>	2.8 (1.5) x 10 <sup>6</sup>
_	Yes	7	1.6 (0.2) x 10 <sup>6</sup>	1.7 (0.2) x 10 <sup>6</sup>

Table 3-5. Growth of clade III N<sub>2</sub>O reducers is N<sub>2</sub>O- and pH- dependent. <sup>a</sup>

<sup>a</sup> Growth of clade III N<sub>2</sub>O reducers (i.e., MAG6 and MAG7) were measured in mixed cultures (culture Sabana and WS2, respectively) with *nosZ*-targeting primers. <sup>b</sup> Cells measured immediately following vessels received inoculum. <sup>c</sup> Cells in pH-4.5 and -7 cultures were measured at 4- and 10-day, respectively.

Chapter 4 Geochemical factors control N<sub>2</sub>O reduction under low pH conditions

#### Abstract

Nitrous oxide (N<sub>2</sub>O) is a greenhouse gas exacerbating ozone depletion and global warming. Microorganisms synthesizing  $N_2O$  reductase are the only sink of  $N_2O$ . However, the N<sub>2</sub>O reduction process is regulated by environmental factors (e.g., oxygen content, pH and copper ions). Acidic soils had long been considered  $N_2O$  emitters because laboratory efforts studying denitrification showed a strong preference for circumneutral pH conditions for complete reduction to environmentally benign dinitrogen gas. Field observations and the recent discovery of bacteria with a preference of low pH N<sub>2</sub>O reduction suggest the microbiology is not limiting N<sub>2</sub>O consumption under acidic conditions. Laboratory cultivation experiments explored the geochemical controls of acidophilic and neutrophilic N<sub>2</sub>O reducing co-culture EV and *Dechloromonas aromatica* strain RCB, respectively. The results revealed geochemical factors (i.e., nitrogen oxyanions, Cu<sup>2+</sup> and Na<sub>2</sub>S), instead of pH, were limiting low pH N<sub>2</sub>O reduction by coculture EV. Temporal nitrogen oxyanions (i.e.,  $NO_3^-$  and  $NO_2^-$ ) dosing experiments demonstrated different mechanisms behind NO3<sup>-</sup> and NO2<sup>-</sup> inhibition on low pH N2O reduction. Cu<sup>2+</sup> and Na<sub>2</sub>S dosing experiments revealed multidimensional effect of sulfide on N<sub>2</sub>O reduction. The start-up concentration of copper toxicity (0.5 mM) on N<sub>2</sub>Odependent growth was much lower than that reported for NO<sub>3</sub><sup>-</sup>-dependent growth. These observations together suggest that geochemical factors other than pH are limiting  $N_2O$ reduction in acidic soils.

### 4.1 Introduction

Nitrous oxide (N<sub>2</sub>O) is a long-lived greenhouse gas with ozone destruction potential and is of environmental significance for global warming. Increases in atmospheric  $N_2O$  are predicted to cause a 2-4% decline of the total ozone column by the end of 21<sup>st</sup> century (180). Biotic and coupled biotic-abiotic processes in terrestrial and aquatic ecosystems are the primary N<sub>2</sub>O sources (181, 182). The N<sub>2</sub>O-forming and -consuming processes are regulated by multiple environmental factors such as temperature, oxygen partial pressure, pH, copper availability, and electron donor availability (40, 183-186). The only known biological N<sub>2</sub>O sink is the microbial transformation of N<sub>2</sub>O to N<sub>2</sub> catalyzed by N<sub>2</sub>O reductase  $(N_2OR)$  (17), which was previously considered a weak process in acidic soils (40, 41). Current paradigm of low pH N<sub>2</sub>O reduction is that the assembly of N<sub>2</sub>O reductase is impaired under acidic pH conditions (38, 40, 41). However, a deep-sea isolate, *Nitratiruptor* sp., respires N<sub>2</sub>O between pH 5.4 to 6.4, contradicting the recognition that N<sub>2</sub>O reductase cannot be assembled under acidic conditions (105). We have enriched clade II nosZ-carrying organisms capable of respiring N<sub>2</sub>O under strong acidic conditions (Guang, et al., unpublished work) from tropical forest soils and managed agricultural soils, implicating microbiology is not limiting N<sub>2</sub>O reduction activities in acidic soils. Thus, a better understanding on the limiting factors of N<sub>2</sub>O reduction activities is needed to manage N<sub>2</sub>O emissions from acidic soils.

N<sub>2</sub>OR, encoded by the *nosZ* gene, is a periplasmic multicopper protein consisting of two distinct copper centers, designated Cu<sub>A</sub> and Cu<sub>Z</sub> (17) and strictly requires copper as a metal cofactor (187). The binuclear Cu<sub>A</sub> center channels electrons to the tetranuclear Cu<sub>Z</sub> center where N<sub>2</sub>O reduction to N<sub>2</sub> occurs (188). Copper-limiting conditions impacts the assembly of functional N<sub>2</sub>OR. For example, methanobactin produced by some methanotrophs, and the formation of CuS with a solubility product of 8.5 x 10<sup>-45</sup> (189), effectively sequester copper and inhibits N<sub>2</sub>O reduction (127, 190, 191). Under circumneutral pH conditions, the addition of copper ions (Cu<sup>2+</sup>) recovered N<sub>2</sub>O reduction activity (191, 192). However, due to limited cultivable organisms with capability of N<sub>2</sub>O

reduction at pH < 5, the impact of copper sequestration by sulfide on N<sub>2</sub>O reduction under acidic conditions remains unclear.

 $NO_3^-$  and  $NO_2^-$  are key precursors of N<sub>2</sub>O availabilities and have been implicated as key environmental factors controlling N<sub>2</sub>O formation (32, 193). [need ref that shows that  $NO_3^-$  and  $NO_2^-$  are N2O precursors. Then need refs that show that  $NO_3^-$  and  $NO_2^$ concentrations impacts N2O formation (e.g., Sukhwan Yoon's paper with Shewanella loihica). Net N<sub>2</sub>O emission was result from N<sub>2</sub>O production outpacing N<sub>2</sub>O consumption, presumably due to geochemical favoring N<sub>2</sub>O production or inhibiting N<sub>2</sub>O consumption (38, 128, 185). For example, intracellular competition for electrons directed towards NO<sub>3</sub><sup>-</sup> versus N<sub>2</sub>O reduction in denitrifying organisms (converting NO<sub>3</sub><sup>-</sup> to N<sub>2</sub>) can explain the impact of NO<sub>3</sub><sup>-</sup> on N<sub>2</sub>O reduction activity (194). NO<sub>2</sub><sup>-</sup> also impacts N<sub>2</sub>O reduction and stalled N<sub>2</sub>OR activity was observed in laboratory incubations (195, 196). Previous paradigm that denitrifying organisms are the key  $N_2O$  consumers had been argued by the discovery of clade II  $N_2O$  reducers, most of which lack the *nirK* and *nirS* genes to denitrify (12). Therefore, denitrification is considered a modular microbial process, consisting of step-by-step reductions of oxidized nitrogen species (NO<sub>3</sub><sup>-</sup>  $\rightarrow$  NO<sub>2</sub><sup>-</sup>  $\rightarrow$  N<sub>2</sub>O  $\rightarrow$  N<sub>2</sub>). The impact of nitrogen oxyanions on [new section. now you are bringing up the modularity issue. This needs to be explicitly explained  $N_2O$ -reducing bacteria that cannot denitrify may differ from that on denitrifying N<sub>2</sub>O reducers. The case studies investigating the effect of nitrogen oxyanions on N<sub>2</sub>O reduction was limited to neutral pH conditions (197). Our former work documented that organisms performing low pH N<sub>2</sub>O reduction are incapable of denitrification with *nirK* and *nirS* genes absent (Guang, et al., unpublished work). Therefore, those cultures offer an opportunity to study the impact of nitrogen oxyanion on low pH N<sub>2</sub>O reduction.

While some progress has been made to understand the complex organisms-environment interactions that govern  $N_2O$  reduction at circumneutral pH, our understanding of factors controlling  $N_2OR$  activity at low pH are unclear. This research was conducted to test the hypothesis: geochemical factors, instead of pH, is limiting microbiology of low pH  $N_2O$  reduction. Specifically, copper, sulfide and nitrogen oxyanions limitations on low pH

N<sub>2</sub>O reduction by culture EV were evaluated. The *Desulfosprosinus nitrousreducens* habituating culture EV reduces N<sub>2</sub>O with H<sub>2</sub> as electron donor. To compare the impact of geochemical factors on acidic N<sub>2</sub>O reduction to that on neutral N<sub>2</sub>O reduction, hydrogenotrophic N<sub>2</sub>O-reducing *Dechloromonas aromatica* strain RCB was cultivated in parallel.

### 4.2 Materials and Methods

#### Microorganisms and cultivation.

Co-Culture EV, consisting of pyruvate-fermenting Serratia marcescens strain MF and N<sub>2</sub>O-respiring *Desulfosporosinus nitrousreducens*, was enriched from acidic tropical forest soil (Guang, et al, unpublished work). After the establishment of the stable coculture, it was maintained in the laboratory via repeated transfers in defined mineral salt medium with  $N_2O$  as the sole electron acceptor and  $H_2$  as the electron donor. Co-Culture EV was cultivated in 160-mL serum bottles containing 100 mL phosphate-buffered (50 mM, pH 4.5) anoxic mineral salt medium reduced with L-cysteine (0.2 mM) as described with the following modifications (Guang, et al, unpublished work). A volume of 1 mL stock solution of mixed amino acid was added to the vessels prior to inoculation served as growth nutrient. The stock solution of amino acid contains (g  $L^{-1}$ ): alanine (0.5); aspartic acid (1); proline (1); tyrosine (0.3); histidine (0.3); tryptophan (0.2); arginine (0.5); isoleucine (0.5); methionine (0.4); glycine (0.3); threonine (0.5); valine (0.9); lysine (1); glutamate (1); serine (0.8). Dechloromonas aromatica strain RCB was maintained in 160-mL serum bottles containing 100 mL bicarbonate-buffered (30 mM, pH 7.3) anoxic mineral salt medium reduced with L-cysteine (0.2 mM) as described (101). Ten mL (416 µmol) of H<sub>2</sub> and N<sub>2</sub>O were added using plastic syringes and equilibrated over-night prior to the inoculation of co-culture EV or D. aromatica strain RCB.

#### Abiotic controls of N<sub>2</sub>O reduction at low pH.

N<sub>2</sub>O reduction by the co-culture EV was examined in the presence of copper chloride (CuCl<sub>2</sub>) at the concentration ranging from 0 to 0.5 mM. To precipitate the residual copper carried from the parent culture, sodium sulfide (Na<sub>2</sub>S) was added at a final concentration of 0.2 mM in cultures that did not receive CuCl<sub>2</sub>. N<sub>2</sub>O reduction of co-culture EV and *D. aromatica* strain RCB was evaluated in the presence of Na<sub>2</sub>S (0.2 or 1 mM) with excessive CuCl<sub>2</sub> (0.22 or 1.02 mM). Cultures omitting CuCl<sub>2</sub> severed as negative controls. Additional time-serial dosing experiments were conducted to evaluate the toxicity of Na<sub>2</sub>S to co-culture EV and *D. aromatica* strain RCB. Briefly, cultures amended with Na<sub>2</sub>S (0.2 or 1 mM) were incubated for the first week followed by the addition of excessive CuCl<sub>2</sub> (0.22 or 1.02 mM) to resuscitate N<sub>2</sub>O respiration. To investigate the effect of nitrate and nitrite on N<sub>2</sub>O reduction by co-culture EV, growth experiments were conducted in the presence of nitrate or nitrite (0.1, 0.5 or 1 mM). Na<sub>2</sub>S, nitrate and nitrite were added to the cultures when half of the initially added N<sub>2</sub>O was consumed to assess their impact on the activity of pre-synthesized N<sub>2</sub>O reductase.

### Analytical procedures.

N<sub>2</sub>O were analyzed via manually injecting 100  $\mu$ L headspace samples into an Agilent 3000A Micro-Gas Chromatograph (Palo Alto, CA, USA) equipped with Plot Q and molecular sieve columns coupled with thermal conductivity detection as described (101). Aqueous gas concentrations were calculated from the headspace concentrations based on the Henry's law using published Henry's constant for N<sub>2</sub>O (2.4×10<sup>-4</sup>) (102).

$$C_{aq} = \frac{C_g}{H} \times \frac{1}{24.86}$$
 Equation 1 Henry's law

 $C_{aq}$ ,  $C_g$  and H are the aqueous gas concentration, the headspace gas concentration, and Henry's constant (dimensionless), respectively. Five-point standard curves for N<sub>2</sub>O spanned a concentration rage of 8,333 to 133,333 ppmv. The total amount of N<sub>2</sub>O in vessels was calculated by combining the gaseous and aqueous N<sub>2</sub>O. Average N<sub>2</sub>O reduction rates at different pH were calculated as below:

$$v(N_2 O) = \frac{c(N_2 O_m - N_2 O_n)}{m - n}$$
 Equation 2

 $v(N_2O)$  is the average N<sub>2</sub>O reduction rate; c represents N<sub>2</sub>O concentration (unit: µmole vessel<sup>-1</sup>); n and m represent timepoint prior to measurable N<sub>2</sub>O consumption observed and at N<sub>2</sub>O depletion. NO<sub>3</sub><sup>-</sup> and NO<sub>2</sub><sup>-</sup> were measured using the Dionex IC 2100 system (Sunnyvale, CA, USA) as described (198).

## 4.3 Results

### N<sub>2</sub>O reduction by culture EV and D. aromatica strain RCB require copper

Culture EV enriched from acidic forest soils harbors a non-denitrifying *Serratia* marcescens population and an obligate N<sub>2</sub>O-reducing *Desulfosporosinus nitrousreducens* which only reduces N<sub>2</sub>O to N<sub>2</sub> between pH 4.5 to 6 (Figure 4-1 A). *Dechloromonas* aromatica strain RCB carries all genes required for canonical denitrification and grows with NO<sub>3</sub><sup>-</sup> or N<sub>2</sub>O as sole electron acceptor (Figure 4-1 B). A genomic feature shared by *Desulfosporosinus nitrousreducens* and *Dechloromonas aromatica* strain RCB is the hyptype NiFe hydrogenase, which is consistent to their capability of utilizing hydrogen gas as the sole electron donor.

Culture EV cultivated at pH 4.5 and *Dechloromonas aromatica* strain RCB grown at pH 7 reduced N<sub>2</sub>O at similar rates with Cu<sup>2+</sup> concentrations ranging from 0.0005 to 0.05 mM over a 6-day incubation (Figure 4-1 C, D). N<sub>2</sub>O reduction was not observed in either culture EV or *Dechloromonas aromatica* strain RCB received 0.2 mM sulfide. Sulfide precipitates Cu<sup>2+</sup> to form CuS with a solubility product constant of  $8.5 \times 10^{-45}$  indicating the dissolved, bioavailable Cu<sup>2+</sup> concentration is no more than  $4.25 \times 10^{-38}$  nM. N<sub>2</sub>O reduction was completely inhibited when the Cu<sup>2+</sup> concentration exceeded 0.5 mM, presumably due to toxic effects.

### Effects of copper and Na<sub>2</sub>S on N<sub>2</sub>O reduction

In both culture EV (grown at pH 4.5) and the axenic culture of D. aromatica strain RCB (grown at pH 7) receiving copper at the concentration of 0.02 mM, N<sub>2</sub>O was depleted within 6 days. N<sub>2</sub>O reduction was prohibited in all cultures of EV and D. aromatica strain RCB amended with 0.2 mM Na<sub>2</sub>S or 1 mM Na<sub>2</sub>S alone (Figure 4-2). Average N<sub>2</sub>O reduction rates in culture EV received both Na<sub>2</sub>S and CuCl<sub>2</sub> ( $35.5 \pm 1.22$  and  $34.5 \pm 0.26$  $\mu$ mol N<sub>2</sub>O day<sup>-1</sup>) were about half of that received 0.02 mM CuCl<sub>2</sub> only (66.1 ± 0.46  $\mu$ mol N<sub>2</sub>O day<sup>-1</sup>, Figure 4-2 A). Culture EV amended with 0.2 mM Na<sub>2</sub>S plus 0.22 mM CuCl<sub>2</sub> consumed the initially added N<sub>2</sub>O following an incubation of 11 days. A lag phase of 10day was observed for Culture EV amended with 1 mM Na<sub>2</sub>S plus 1.02 mM CuCl<sub>2</sub> to initiate N<sub>2</sub>O reduction (Figure 4-2 C). Similar average N<sub>2</sub>O reduction rates of D. aromatica strain RCB were observed in cultures with both Na<sub>2</sub>S and CuCl<sub>2</sub> (70.9  $\pm$  0.84 and 70.0  $\pm$  2.32 µmol N<sub>2</sub>O day<sup>-1</sup>), and in cultures with CuCl<sub>2</sub> only (70.1  $\pm$  0.84 µmol N<sub>2</sub>O day<sup>-1</sup>, Figure 4-2 B). No lag phase was observed for *D. aromatica* strain RCB received excessive copper (0.2 mM Na<sub>2</sub>S plus 0.22 mM CuCl<sub>2</sub> or 1 mM Na<sub>2</sub>S plus 1.02 mM  $CuCl_2$ ) to initiate N<sub>2</sub>O reduction (Figure 4-2 D). The different N<sub>2</sub>O reduction phenotype of culture EV responding to low and high Na<sub>2</sub>S concentration raised a question: does Na<sub>2</sub>S exert toxicity to N<sub>2</sub>O-reducing organisms at high concentrations under acidic conditions? Therefore, excessive CuCl<sub>2</sub> (0.22 or 1.02 mM) was added to cultures of El and D. aromatica strain RCB pre-exposed to low and high Na<sub>2</sub>S. CuCl<sub>2</sub> dosing initiated N<sub>2</sub>O reduction in cultures of both EV and D. aromatica strain RCB exposed to low Na<sub>2</sub>S concentration (0.2 mM) but not to high Na<sub>2</sub>S concentration (1 mM) (Figure 4-3 A, B). In contrast, N<sub>2</sub>O reduction was stalled in culture EV and *Dechloromonas aromatica* strain RCB immediately following Na<sub>2</sub>S dosing (Figure 4-3 C, D), suggesting the of Na<sub>2</sub>S over copper against pre-synthesized N<sub>2</sub>O reductase.

### Effects of nitrogen oxyanion on low pH N<sub>2</sub>O reduction.

 $N_2O$  reduction by culture EV was prohibited in the presence of 0.1 mM to 1 mM  $NO_3^$ and 0.01 mM to 1 mM  $NO_2^-$  (Figure 4-4 A, B). Culture EV received a concentration of 0.01 mM NO<sub>3</sub><sup>-</sup> or 0.001 mM NO<sub>2</sub><sup>-</sup> exhibited comparable N<sub>2</sub>O reduction activities as cultures without NO<sub>3</sub><sup>-</sup> or NO<sub>2</sub><sup>-</sup>, suggesting the threshold concentrations of NO<sub>3</sub><sup>-</sup> and NO<sub>2</sub><sup>-</sup> inhibition on N<sub>2</sub>O reduction by culture EV are between 0.01 to 0.1 mM and 0.001 to 0.01 mM, respectively. Following a 3-day incubation, culture EV reduced approximately half of the initial amount of N<sub>2</sub>O, subsequent NO<sub>2</sub><sup>-</sup> (1 mM) but not NO<sub>3</sub><sup>-</sup> dosing (1 mM) stalled N<sub>2</sub>O reduction immediately (Figure 4-4 C). In contrast, NO<sub>2</sub><sup>-</sup> dosing in *Dechloromonas aromatica* strain RCB transiently inhibit N<sub>2</sub>O reduction, suggesting NO<sub>2</sub><sup>-</sup> exerted a monopoly on electron against N<sub>2</sub>O reduction (Figure 4-4 D).

#### 4.4 Discussion

Geochemical factors are important regulators controlling N<sub>2</sub>O flux in soils (199) and their impact on N<sub>2</sub>O reduction have been extensively studied under circumneutral pH conditions (185, 191, 196). However, the regulation of geochemistry over low pH N<sub>2</sub>O reduction remains elusive due to limited cultivable organisms. The isolation of organism that reduces N<sub>2</sub>O under acidic conditions (pH <5) (Guang, et al., unpublished work) offers an opportunity to study how geochemical factors impact N<sub>2</sub>O reduction under acidic conditions. In this study, we demonstrated that N<sub>2</sub>O-dependent growth of culture EV was abolished in the presence of either NO<sub>3</sub><sup>-</sup> or NO<sub>2</sub><sup>-</sup>, implicating that nitrogen oxyanions, instead of pH itself, may be the limiting factor of N<sub>2</sub>O reduction in acidic soils.

Acidic soils are considered as N<sub>2</sub>O emitters because of the inability of N<sub>2</sub>O reduction under acidic conditions. A previous study focused on denitrifying organisms attributed the abolished N<sub>2</sub>O reduction to pH interference on N<sub>2</sub>O reductase assembly (40). However, we enriched a co-culture harboring *Desulfosporosinus* bacteria capable of sustaining N<sub>2</sub>O reduction under acidic conditions (pH ~ 4.5) (Guang, et al., unpublished work). Here, we demonstrate that NO<sub>2</sub><sup>-</sup> has a lethal (inhibitory?) effect on both N<sub>2</sub>Odependent growth and the activity of pre-synthesized N<sub>2</sub>O reductase of culture EV under acidic conditions. Meanwhile NO<sub>2</sub><sup>-</sup> only exert a transient inhibitory effect on N<sub>2</sub>O

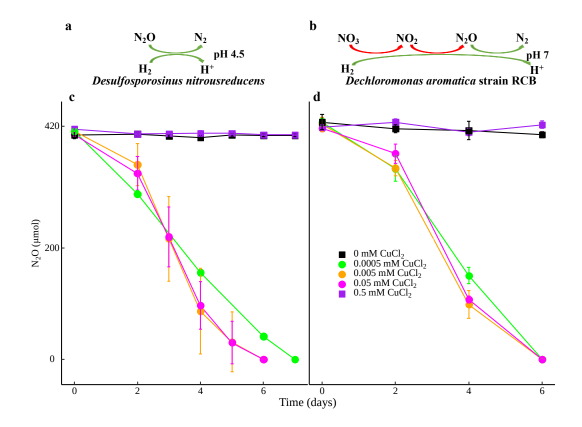
consistent to previous studies focused on neutralophilic denitrifying organisms (110, 196). In addition,  $NO_3^-$  prevents N<sub>2</sub>O-dependent growth while exerts no effect on presynthesized N<sub>2</sub>O reductase activity of co-culture EV. The adverse impact of  $NO_3^-$  on N<sub>2</sub>O-dependent growth of *Desulfosporosinus nitrousreducens* was unexpected. Though high soil  $NO_3^-$  content (10 mM, equivalent to 44.9 mg N kg<sup>-1</sup> soil) was reported to inhibit N<sub>2</sub>O reduction in soil incubations (200), there is no example of  $NO_3^-$  regulation over N<sub>2</sub>O reduction in cultures. The theory of electron competition between  $NO_3^-$  and N<sub>2</sub>O derived from denitrifying *Bradyrhizobia* genera (194, 201) is not sufficient to explain the observation of non-denitrifying culture EV. An alternative explanation is that the presence of  $NO_3^-$  only interferes the assembly of N<sub>2</sub>O reductase while the already assembled enzymes are still functioning. These observations together suggested that genotype varying N<sub>2</sub>O reducers respond differently to nitrogen oxyanions.

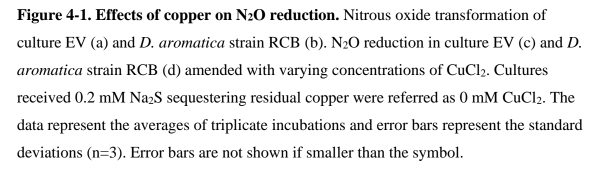
Cu is a co-factor involved in the assembly of functional N<sub>2</sub>O reductase (17). In this research, cultivation experiments with culture EV and *Dechloromonas aromatica* strain RCB demonstrated that Cu at nM level is sufficient for the assembly of N<sub>2</sub>O reductase under both acidic and circumneutral pH conditions. A previous study proposed that H<sub>2</sub>S yielded from cysteine during anaerobic incubation may sequester copper and impede N<sub>2</sub>O reduction (202). However, in our cysteine-amended medium, N<sub>2</sub>O reduction of both culture EV and *Dechloromonas aromatica* strain RCB was not impacted until exogenous Na<sub>2</sub>S was added. Furthermore, the onset concentration of Cu (0.5 mM) toxicity on N<sub>2</sub>O-dependent growth was much lower than that previously reported (20 mM) on NO<sub>3</sub><sup>-</sup>-dependent growth (202). These observations are informative to soil environments where N<sub>2</sub>O is the sole energy source for microbial growth.

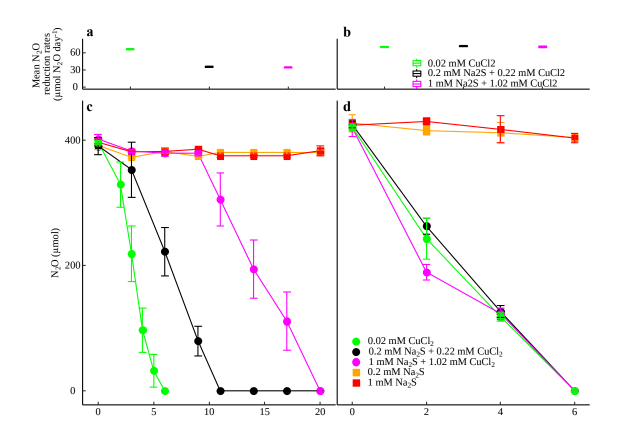
Sulfide has been long considered as a potent inhibitor of  $N_2O$  reduction and previous studies derived two possible mechanisms: (I) ionic S<sup>2-</sup> and HS<sup>-</sup> formulated H<sub>2</sub>S exerts toxicity on organisms (203); (II) sequestration of copper by S<sup>2-</sup> (191). Here we demonstrate that the effect of sulfide on N<sub>2</sub>O reduction is multidimensional by parallelly cultivating two N<sub>2</sub>O-respiring organisms under either acidic or circumneutral pH conditions. Copper dosing resuscitated N<sub>2</sub>O reduction inhibited low concentration of

Na<sub>2</sub>S (0.2 mM) in culture EV grown at pH 4.5 and *Dechloromonas aromatica* strain RCB (grown at pH 7), suggested copper sequestration to be the major cause of stalled N<sub>2</sub>O reduction in the scenario. This observation is consistent with previous incubation experiments with sludge and an axenic culture where divalent metal dosing restored sulfide-inhibited N<sub>2</sub>O reduction (191, 192). However, if organisms could recover N<sub>2</sub>O reduction activity following the exposure to high concentrations of Na<sub>2</sub>S were not considered in those studies. In this study, high concentration of Na<sub>2</sub>S (1 mM) exerted irreversible inhibition on N<sub>2</sub>O reduction in culture EV and *Dechloromonas aromatica* strain RCB, suggesting the onset concentration of sulfide toxicity is pH independent. Furthermore, the toxicity of copper and sulfide could be counteracted via precipitation, implicating interactions between geochemical factors indirectly regulate N<sub>2</sub>O reduction activity.

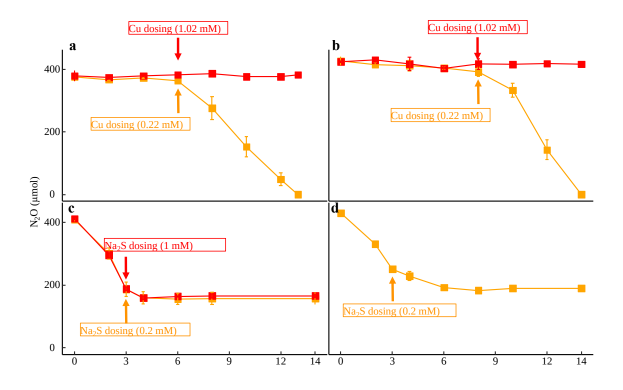
# Appendix







**Figure 4-2. Effect of Na<sub>2</sub>S on N<sub>2</sub>O reduction.** Mean N<sub>2</sub>O reduction rates of culture EV (a) and *D. aromatica* strain RCB (b) in the presence or absence of Na<sub>2</sub>S. N<sub>2</sub>O reduction in culture EV (c) and *D. aromatica* strain RCB (d) received either CuCl<sub>2</sub> or Na<sub>2</sub>S and cultures amended with excess CuCl<sub>2</sub> and Na<sub>2</sub>S. The data represent the averages of triplicate incubations and error bars represent the standard deviations (n=3). Error bars are not shown if smaller than the symbol.



**Figure 4-3. Reversible and irreversible inhibition of Na<sub>2</sub>S on N<sub>2</sub>O reduction.** N<sub>2</sub>O reduction of culture EV following CuCl<sub>2</sub> (a) or Na<sub>2</sub>S dosing (c). N<sub>2</sub>O reduction by *D. aromatica* strain RCB following CuCl<sub>2</sub> (c) or Na<sub>2</sub>S dosing (d). Arrows indicate CuCl<sub>2</sub> or Na<sub>2</sub>S dosing. The data represent the averages of triplicate incubations and error bars represent the standard deviations (n=3). Error bars are not shown if smaller than the symbol.

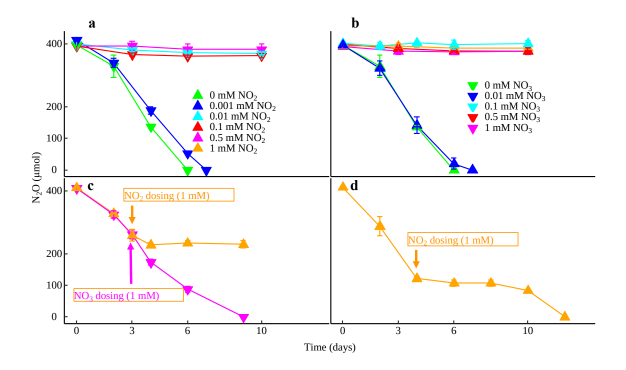


Figure 4-4. Inhibition of nitrogen oxyanions on N<sub>2</sub>O reduction by culture EV. N<sub>2</sub>O reduction in culture EV received varying concentrations of NO<sub>2</sub><sup>-</sup> (a) or NO<sub>3</sub><sup>-</sup> (b). N<sub>2</sub>O reduction in culture EV (c) and *Dechlromonas aromatica* strain RCB (d) following nitrogen oxyanion dosing. The data represent the averages of triplicate incubations and error bars represent the standard deviations (n=3). Error bars are not shown if smaller than the symbol.

## **Summary and Conclusion**

Soil health is being degraded across the globe due to exploitation, mismanagement, and anthropogenic practices. One important parameter of soil health is pH, but regions around the world are experiencing soil acidification due to natural processes accelerated by acidic precipitation/deposition and agricultural practices that rely on nitrogen fertilizer and the growth of legumes. Nitrogen fertilizer input also leads to the formation of N<sub>2</sub>O, a substance linked to climate warming and interference with biogeochemical processes such as methanogenesis, mercury turnover, and reductive dechlorination.

The imbalance between N<sub>2</sub>O formation and consumption is a key issue, with the rise in global N<sub>2</sub>O emission indicating that N<sub>2</sub>O formation is outpacing consumption. This is partly due to the contrasting pH optima of processes involved in N<sub>2</sub>O formation (pH 5-6) versus consumption (pH $\sim$ 7). While some studies have reported N<sub>2</sub>O consumption in acidic soil laboratory microcosms and ecosystems, soil heterogeneity and associated microscale patchiness of pH conditions make generalized conclusions difficult.

Attempts to demonstrate low pH N<sub>2</sub>O reduction in denitrifying enrichment and axenic cultures have so far failed to demonstrate growth-linked N<sub>2</sub>O reduction and the sustainability of such a process. The only known sink for N<sub>2</sub>O are microorganisms expressing N<sub>2</sub>O reductase, a periplasmic enzyme that catalyzes the conversion of N<sub>2</sub>O to environmentally benign dinitrogen. Studies have shown that pH interferes with NosZ maturation, limiting microbial N<sub>2</sub>O reduction to circumneutral pH environments.

However, chapter 2 has shown that organisms capable of low pH N<sub>2</sub>O reduction exist, at least, in acidic soils. We demonstrated an intricate organismal interaction between two populations derived from El Verde soil, revealing a commensal relationship between a pyruvate fermenting *Serratia* population and an N<sub>2</sub>O-reducing *Desulfosporosinus* population. Growth experiments illustrated that the *Serratia* population fermented pyruvate to acetate, formate, and CO2, while the *Desulfosporosinus* population only grew with N<sub>2</sub>O. The *Desulfosporosinus* population reduced N<sub>2</sub>O at a narrow range of pH (4.5

to 6), which contradicts previous observations that model denitrifier *Paracoccus denitrificans* only reduces N<sub>2</sub>O above pH 6. The difference in N<sub>2</sub>O reduction behavior may be due to differences in enzyme assembly strategies.

In chapter 3, Enrichment cultures were established with acidic tropical forest soil and circumneutral soil retrieved from an organic farm practicing regenerative agriculture. Comparative metagenome analysis revealed N<sub>2</sub>O reduction under acidic conditions was associated with microbial growth. Further analysis suggested that low pH N<sub>2</sub>O reducers cannot perform canonical denitrification and that phylogenetic analysis of NosZ genes illustrated a preference of acidic pH over clade II N<sub>2</sub>O reducers. Two novel subclades were also identified with the *nosZ* sequences from the enrichment cultures.

Although the existence of soil microorganisms capable of low pH N<sub>2</sub>O reduction has been demonstrated, the in-situ N<sub>2</sub>O emissions from acidic soils are generally higher than those from circumneutral pH soils. Chapter 4 shows that N<sub>2</sub>O reduction of the El Verde co-culture is sensitive to geochemical factors such as copper and nitrogen oxyanions. These factors could play a role in limiting N<sub>2</sub>O reduction in acidic soils, explaining why N<sub>2</sub>O reducers stay inactive even when they reside in such environments.

In conclusion, the study of low pH N<sub>2</sub>O reduction in soil microorganisms provides valuable insights into the role of microbial communities in mitigating the impacts of anthropogenic activities on soil health and global climate change. The findings suggest that the reduction of N<sub>2</sub>O in acidic soils is not only possible but also linked to the presence of specific microbial populations with unique functional and physiological characteristics. However, the complex interactions between soil biogeochemistry and microbial communities make it difficult to generalize the findings and predict their implications at a larger scale. Further research is needed to better understand the factors that regulate the activity of N<sub>2</sub>O reducers in different soil types and under different environmental conditions to develop effective strategies for sustainable soil management and climate change mitigation.

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Guang He was born in Guangzhou City, Guangdong Province, China in 1995. He spent his childhood (up to 12 year-old) in Guangzhou City and teenager in Leivang City, where his parents were born. As such, Guang He does well in both Mandarin and Cantonese. Guang He moved to Changsha City pursuing the bachelor degree in Agricultural Resource and Environment at Hunan Agriculture University. The undergraduate research was more related to large-scale field study (e.g., on-site camping and sampling). He got a good internship opportunity in Chinese Agricultural Academy Science, where he enrolled in a fundamental research project. These experiences inspired Guang He to pursue a master education in the China Agriculture University. He then joined the Junling Zhang's lab researching the tri-party interaction between host-plant, arbuscular mycorrhizal fungi and N<sub>2</sub>O-cycling bacteria. During his master education, he self-learned versatile molecular techniques (e.g., qPCR and clone sequencing). In addition, Guang He developed great interests in computational analysis (e.g., omics analysis and R visualization). The skills Guang He obtained in master program stand him ready to the Ph.D study in Department of Plant, Soil and Environmental Science at the University of Tennessee, Knoxville, supervised by prestigious Dr. Frank E. Löffler and Dr. Mark Radosevich. In the past five years, Guang He has been researching the microbial reduction of N<sub>2</sub>O under acidic conditions, which has been a decade-lasting retard topic. The most impressing intellectual merit achieved by Guang He is that he obtained different enrichment cultures from either agricultural or tropical soils, all of which have the low pH N<sub>2</sub>O reduction capacity. More strikingly, he has made it available of an acidophilic N<sub>2</sub>O reducing bacterium in pure culture. Guang He is also involved in other collaborative projects including N-PFAS supporting microbial growth, where he contributes to the bioinformatic analyses and data interpretation. After graduation from the University of Tennessee, still driven by the passion, Guang He intends to continue his research career and contribute to resolving the global climate changes induced by N<sub>2</sub>O emissions.