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Nitrous oxide respiring bacteria in biogas digestates for reduced agricultural emissions

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Inoculating agricultural soils with nitrous oxide respiring bacteria (NRB) can reduce N₂O-emission, but would be impractical as a standalone operation. Here we demonstrate that digestates obtained after biogas production are suitable substrates and vectors for NRB. We show that indigenous NRB in digestates grew to high abundance during anaerobic enrichment under N₂O. Gas-kinetics and meta-omic analyses showed that these NRB's, recovered as metagenome-assembled genomes (MAGs), grew by harvesting fermentation intermediates of the methanogenic consortium. Three NRB's were isolated, one of which matched the recovered MAG of a *Dechloromonas*, deemed by proteomics to be the dominant producer of N₂O-reductase in the enrichment. While the isolates harbored genes required for a full denitrification pathway and could thus both produce and sequester N₂O, their regulatory traits predicted that they act as N₂O sinks in soil, which was confirmed experimentally. The isolates were grown by aerobic respiration in digestates, and fertilization with these NRB-enriched digestates reduced N₂O emissions from soil. Our use of digestates for low-cost and large-scale inoculation with NRB in soil can be taken as a blueprint for future applications of this powerful instrument to engineer the soil microbiome, be it for enhancing plant growth, bioremediation, or any other desirable function.

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INTRODUCTION

Nitrous oxide is an intermediate in the nitrogen cycle and a powerful greenhouse gas emitted in large volumes from agricultural soils, accounting for ~1/3 of total anthropogenic N₂O emissions [1]. Reduced emissions can be achieved by minimizing the consumption of fertilizer nitrogen through improved agronomic practice and reduction of meat consumption [2, 3], but such measures are unlikely to do more than stabilize the global consumption of fertilizer-N [4]. This calls for more inventive approaches to reduce N₂O emissions, targeting the microbiomes of soil [5], in particular the physiology and regulatory biology of the organisms involved in production and consumption of N₂O in soil [6].

N₂O turnover in soil involves several metabolic pathways, controlled by a plethora of fluctuating physical and chemical variables [7–9]. Heterotrophic denitrification is the dominant N₂O source in most soils, while autotrophic ammonia oxidation may dominate in well drained calcareous soils and references therein [10]. Heterotrophic denitrifying organisms are both sources and sinks for N₂O because N₂O is a free intermediate in their stepwise reduction of nitrate to dinitrogen (NO₃⁻ → NO₂⁻ → NO → N₂O → N₂). Denitrification involves four enzymes collectively referred to as denitrification reductases: nitrate reductase (Nar/Nap), nitrite reductase (Nir), nitric oxide reductase (Nor) and nitrous oxide reductase (Nos), encoded by the genes *nar/nap*, *nir*, *nor* and *nosZ*, respectively. Oxygen is a strong repressor of denitrification, both at the transcriptional and the metabolic level [11, 12]. Many

organisms have truncated denitrification pathways, lacking from one to three of the four reductase genes [13, 14], and truncated denitrifiers can thus act as either N₂O producers (organisms without *nosZ*) or N₂O reducers (organisms with *nosZ* only). The organisms with *nosZ* only, coined non-denitrifying N₂O-reducers [15], have attracted much interest as N₂O sinks in the environment [16]. Of note, organisms with a full-fledged denitrification pathway may also be strong N₂O sinks depending on the relative activities and regulation of the various enzymes in the denitrification pathway [17, 18]. Despite their promise, feasible ways to utilize N₂O-reducing organisms to reduce N₂O emissions have not yet emerged.

A soil with a strong N₂O-reducing capacity will emit less N₂O than one dominated by net N₂O producing organisms, as experimentally verified by Domeignoz-Horta et al. [19], who showed that soils emitted less N₂O if inoculated with large numbers (10⁷–10⁸ cells g⁻¹ soil) of organisms expressing Nos as their sole denitrification reductase. As a standalone operation, the large-scale production and distribution of N₂O-respiring bacteria would be prohibitively expensive and impractical. However, the use of N₂O-respiring bacteria could become feasible if adapted to an existing fertilization pipeline, such as fertilization with the nitrogen- and phosphate-rich organic waste (digestate) generated by biogas production in anaerobic digesters. Anaerobic digestion (AD) is already a core technology for treating urban organic wastes, and is expected to treat an increasing proportion of the much larger volumes of waste produced by the agricultural sector

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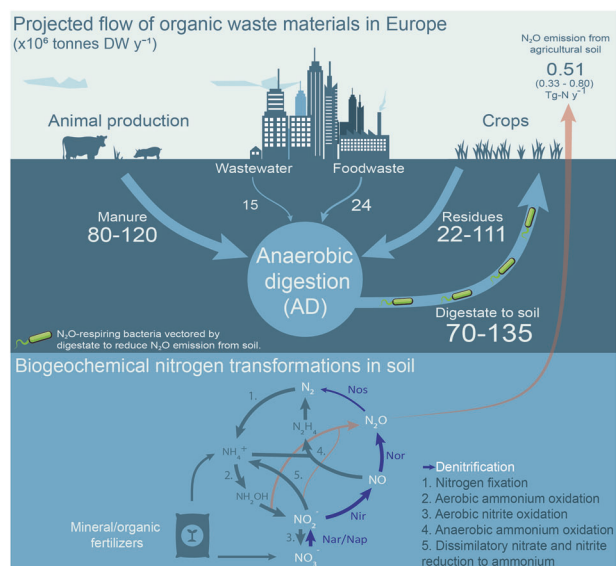


Fig. 1 Possible biomass streams in a future circular economy with a central role for anaerobic digestion. Solid arrows (top section) show streams of biomass available for anaerobic digestion (AD). Numbers indicate known estimates of currently used or potentially available amounts in Europe, in million tonnes dry-weight (DW) per year [50–53]. The arrow from anaerobic digestion to agricultural soil, indicates a credible pathway for digestate enriched with N_2O -reducing bacteria (assuming enrichment after AD); fertilization with such enriched digestates strengthens the N_2O sink capacity of the soil, hence reducing N_2O emissions. N_2O emissions from agricultural soil in Europe are estimated at 0.51 tG per year (min 0.33 – max 0.80), representing some 48% of total European N_2O emissions [1], which account for ~3.5% of the global warming effect from European greenhouse gas emissions and 35% of the global warming effect from European agriculture [54]. The lower half of the picture shows the microbial nitrogen transformations underlying these N_2O emissions, which are fed by fertilizers. Today, AD is primarily used for treating urban organic wastes, which comprise only ~10% of the biomass potentially available for AD. The amount of biomass treated by AD is expected to increase by an order of magnitude when adopted on a large scale in the agricultural sector. This would generate 70–135 Mt DW of digestate annually (assuming 50% degradation by AD), which is equivalent to 400–780 kg DW $ha^{-1} y^{-1}$ if spread evenly on the total farmland of Europe (173 million ha).

(Fig. 1), as an element of the roadmap towards a low-carbon circular economy [20]. This means that digestates from AD are likely to become a major organic fertilizer for agricultural soils, with a huge potential for reducing N_2O emissions if enriched with N_2O -respiring bacteria prior to application.

Here we provide the first proof of this promising concept. Firstly, we demonstrate selective enrichment and isolation of fast-growing digestate-adapted N_2O -respiring bacteria using a digestate from a wastewater treatment plant. Secondly, we demonstrate that the use of digestates enriched with such organisms as a soil amendment reduces the proportion of N leaving soil as N_2O , confirming the suitability of such digestates for this purpose. Analysis of the enrichment process with multi-omics and in-depth monitoring of gas kinetics provides valuable insights into *Nos*-synthesis by the various enriched taxa, and the metabolic pathways of the anaerobic consortium providing substrates for these enriched N_2O -respiring organisms.

MATERIALS AND METHODS

The digestates were taken from two anaerobic digesters, one mesophilic (37°C) and one thermophilic (52°C), which were running in parallel,

producing biogas from sludge produced by a wastewater treatment plant. The sludge was a poly-aluminum chloride (PAX-XL61™, Kemira) and ferric chloride (PIX318™, Kemira) precipitated municipal wastewater sludge, with an organic matter content of 5.6% (w/w). Both digestors reduced the organic matter by approximately 60%, producing digestates containing ~2.1% organic matter, 1.8–1.9 g $NH_4^+-N L^{-1}$, ~16 and 32 Meq VFA L^{-1} , pH = 7.6–7.8 and 8.2; mesophilic and thermophilic, respectively (see Supplementary Methods 1 for further details). The digestates were transported to the laboratory in 1 L insulated steel-vessels and used for incubation experiments 3–6 h after sampling.

The robotized incubation system developed by Molstad et al. [21, 22] was used in all experiments where gas kinetics was monitored. The system hosts 30 parallel stirred batches in 120 mL serum vials, crimp sealed with gas tight butyl rubber septa, which are monitored for headspace concentration of O_2 , N_2 , N_2O , NO, CO_2 and CH_4 by frequent sampling. After each sampling, the system returns an equal volume of He, and elaborated routines are used to account for the gas loss by sampling to calculate the production/consumption-rate of each gas for each time interval between two samplings. More details are given in Supplementary Methods 2.

Enrichment culturing of N_2O -respiring bacteria (NRB) in digestate was done as stirred (300 rpm) batches of 50 mL digestate per vial. Prior to incubation, the headspace air was replaced with Helium by repeated evacuation and He-filling [21] and supplemented with N_2O . The N_2O in the headspace was sustained by repeated injections in response to depletion. Liquid samples (1 mL) were taken by syringe, for metagenomic and metaproteomic analyses, and for quantification of volatile fatty acids (VFA) and 16S rRNA gene abundance. The samples were stored $-80^\circ C$ before analyzed. The growth of NRB in the enrichments was modeled based on the N_2O reduction kinetics. The modeling and the analytic methods (quantification of VFA and 16S rRNA gene abundance) are described in detail in Supplementary Methods 3.

Metagenomics and metaproteomics

Sequencing of DNA (Illumina HiSeq4000), and the methods for Metagenome-Assembled Genome (MAG) binning, and the phylogenetic placement of the MAGs is described in detail in Supplementary Methods 4. Proteins were extracted and digested to peptides, which were analyzed by nanoLC-MS/MS, and the acquired spectra were inspected, using the metagenome-assembled genomes (149 MAGs) as a scaffold (Supplementary Methods 5).

Isolation of N_2O -respiring bacteria (NRB) (Supplementary Methods 6). NRB present in the enrichment cultures were isolated by spreading diluted samples on agar plates with different media composition, then incubated in an anaerobic atmosphere with N_2O . Visible colonies were re-streaked and subsequently cultured under aerobic conditions, and 16S-sequenced. Three isolates, AS (*Azospira* sp), AN (*Azonexus* sp) and SP (*Pseudomonas* sp) (names based on their 16S sequence), were selected for genome sequencing, characterization of their denitrification phenotypes, and for testing their effect as N_2O sinks in soil.

Genome sequencing and phenotyping of isolates

Three isolates were genome sequenced and compared with MAG's of the enrichment culture (Supplementary Methods 7). The isolates' ability to utilize various organic C substrates was tested on BiOLOG Phenotype MicroArray microtiter plates, and their characteristic regulation of denitrification was tested through a range of incubation experiments as in previous investigations [17, 18, 23, 24], by monitoring the kinetics of O_2 , N_2 , N_2O , NO and NO_2^- throughout the cultures' depletion of O_2 and transition from aerobic to anaerobic respiration in stirred batch cultures with He + O_2 ($\pm N_2O$) in the headspace (Supplementary Methods 8). The kinetics of electron flow throughout the oxic and anoxic phase in these experiments were used to assess if the organisms were *bet hedging*, as demonstrated for *Paracoccus denitrificans* [17], i.e. that only a minority of cells express nitrate- and/or nitrite-reductase, while all express *Nos*, in response to oxygen depletion. Putative *bet hedging* was corroborated by measuring the abundance of nitrate-, nitrite- and nitrous oxide reductase (Supplementary Methods 9).

N_2O mitigation experiments (Supplementary Methods 10). To assess the capacity of the isolates to reduce the N_2O emission from soil, they were grown aerobically in sterilized digestate, which was then added to soil in microcosms, for measuring the NO^- , N_2O - and N_2 - kinetics of denitrification in the soil. For comparison, the experiments included soils amended with sterilized digestate, live digestate (no pretreatment), and digestate in

which N_2O -reducing bacteria had been enriched by anaerobic incubation with N_2O (as for the initial enrichment culturing).

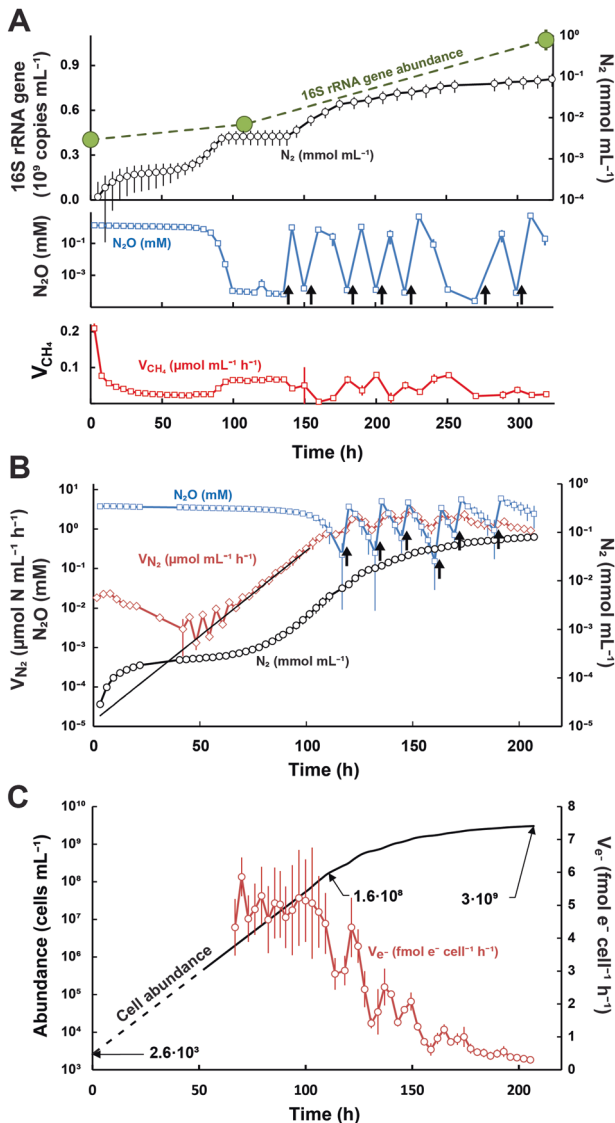
RESULTS AND DISCUSSION

Enrichment of indigenous nitrous oxide respiring bacteria (NRB) in digestates

We hypothesized that suitable organisms could be found in anaerobic digesters fed with sewage sludge, since such sludge contains a diverse community of denitrifying bacteria stemming from prior nitrification/denitrification steps [25]. We further hypothesized that these bacteria could be selectively enriched in digestates by anaerobic incubation with N_2O . We decided to enrich at 20 °C, rather than at the temperatures of the anaerobic digesters (37 and 52 °C), to avoid selecting for organisms unable to grow within the normal temperature range of soils.

The digestates were incubated anaerobically as stirred batch cultures with N_2O in the headspace (He atmosphere), and the activity and apparent growth of N_2O reducers was assessed by monitoring the N_2O -reduction to N_2 . Figure 2A shows the results for the first experiment, where liquid samples were taken at three time points (0, 115 and 325 h) for metagenomics, metaproteomics, and quantification of 16S rRNA gene copy abundance and volatile fatty acids (VFAs). N_2O was periodically depleted (100–140 h) in

Fig. 2 Gas kinetics in anaerobic enrichment cultures with digestate. Panel A shows results for the enrichment culture (triplicate culture vials) sampled for metagenomics, metaproteomics, quantification of volatile fatty acids (VFAs) and 16S rRNA gene abundance (sampling times = 0, 115 and 325 h). The top panel shows the amounts of N_2 produced ($\text{mmol } N_2 \text{ L}^{-1}$ digestate, log scale) and 16S rRNA gene copy numbers. The mid panel shows the concentration of N_2O in the digestate (log scale), which was replenished by repeated injections (20 mL N_2O , resulting in 5 mM N_2O in the liquid) from $t=140$ h and onwards (indicated by black arrows). The bottom panel shows the rate of methane production. Standard deviations ($n=3$) are shown as vertical lines in all panels. Panels B and C show the results of a repeated enrichment experiment where N_2O -depletion (as seen at $t=100$ –140 h in panel A) was avoided, to allow more precise assessment and modeling of growth kinetics. Panel B: N_2O concentration in the digestate (mM N_2O), rate of N_2 -production (V_{N_2}) and N_2 produced ($\text{mmol } N_2 \text{ mL}^{-1}$ digestate), all log scaled. The curved black line shows the modeled V_{N_2} assuming two populations, one growing exponentially ($\mu=0.1 \text{ h}^{-1}$), and one whose activity was dying out gradually (rate = -0.03 h^{-1}). The dotted black line is the activity of the exponentially growing population extrapolated to time = 0. Panel C shows the modeled density (cells mL^{-1}) of cells growing by N_2O respiration, extrapolated back to $t=0$ h (dashed line), and the respiratory electron flow rate per cell (V_{e^-} , $\text{fmol electrons cell}^{-1} \text{ h}^{-1}$), which declined gradually after 110 h. Standard deviations ($n=3$) are shown as vertical lines. Supplementary Fig. S1 provides additional data for the experiment depicted in Panel A, as well as a detailed description of the modeling procedures and their results.



this experiment, precluding detailed analysis of the growth kinetics throughout. This was avoided in the second enrichment, for which complete gas data are shown (Fig. 2B, C). Apart from the deviations caused by the temporary depletion of N_2O in the first experiment, both experiments showed very similar N_2 production rates (Fig. 2B and Supplementary Fig. S1B). The gas kinetics of the second enrichment are discussed in detail below.

The rates of N_2 -production (V_{N_2}) declined during the first 50 h, followed by exponential increase (Fig. 2B). This was modeled as the activity of two groups of NRB, one growing exponentially from low initial abundance, and one which was more abundant initially, but whose activity declined gradually (further explained in Supplementary Fig. S1). The modeling indicated that the cell density of the growing NRB increased exponentially (specific growth rate, $\mu = 0.1 \text{ h}^{-1}$) from a very low initial density ($\sim 2.5 \cdot 10^3$ cells mL^{-1}) to $1.6 \cdot 10^8$ cells mL^{-1} after 110 h, and continued to increase at a gradually declining rate to reach $\sim 3 \cdot 10^9$ cells mL^{-1} at the end of the incubation period (215 h). The modeled electron flow rate per cell for the growing NRB (cell specific e-flow, V_{e^-} , Fig. 2C) was sustained at around 5 $\text{fmol } e^- \text{ cell}^{-1} \text{ h}^{-1}$ during the exponential growth, and declined gradually thereafter (note that the model assumes constant cell yield per mole electrons), as the number of cells continued to increase, while the overall rate of N_2O -respiration remained more or less constant (V_{N_2} , Fig. 2B). Enrichment culturing was repeated several times, demonstrating that the characteristic N_2 production kinetics was highly reproducible (Supplementary Figs. S2 and S14).

The provision of substrate for the N_2O -respiring bacteria can be understood by considering the enrichment culture as a continuation of the metabolism of the anaerobic digester (AD), albeit slowed down by the lower temperature (20 °C versus 37 °C in the digester). In AD, organic polymers are degraded and converted to CO_2 and CH_4 through several steps, conducted by separate guilds of the methanogenic microbial consortium: 1) hydrolysis of polysaccharides to monomers by organisms with carbohydrate-active enzymes, 2) primary fermentation of the resulting monomers to volatile fatty acids (VFAs), 3) secondary fermentation of VFAs to acetate, H_2 and CO_2 , and 4) methane production from acetate, CO_2 , H_2 , and methylated compounds. By providing N_2O to

this (anaerobic) system, organisms that respire N_2O can tap into the existing flow of carbon, competing with the methanogenic consortium for intermediates, such as monomeric sugars, VFAs (such as acetate) and hydrogen [26]. Thus, the respiration and growth of the N_2O -respiring bacteria is plausibly sustained by a flow of carbon for which the primary source is the depolymerization of organic polymers. It is possible that the retardation of growth after ~ 100 h of enrichment was due to carbon becoming limiting. Thus, at this point, the population of N_2O -respiring organisms may have reached high enough cell densities to reap most of the intermediates produced by the consortium.

Parallel incubations of digestates without N_2O confirmed the presence of an active methanogenic consortium, sustaining a methane production rate of $\sim 0.2 \mu\text{mol CH}_4 \text{ mL}^{-1} \text{ h}^{-1}$ throughout (Supplementary Fig. S3). Methane production was inhibited by N_2O , and partly restored in periods when N_2O was depleted (Fig. 2A, Supplementary Figs. S3 and S4). We also conducted parallel incubations with O_2 and NO_3^- as electron acceptors. These incubations showed that methanogenesis was completely inhibited by NO_3^- , and partly inhibited by O_2 (concentration in the liquid ranged from 20 to 90 $\mu\text{M O}_2$) (Supplementary Fig. S3). The rates of O_2 and NO_3^- reduction indicated that the digestate contained a much higher number of cells able to respire O_2 and NO_3^- than cells able to respire N_2O (Supplementary Fig. S5A–C). During the enrichment culturing with NO_3^- almost all reduced NO_3^- appeared in the form of N_2O -N during the first 50 h (Supplementary Fig. S5E), another piece of evidence that in the digestate, prior to enrichment culturing, the organisms reducing NO_3^- to N_2O outnumbered those able to reduce N_2O to N_2 . The measured production of CH_4 and electron flows to electron acceptors deduced from measured gases (N_2 , O_2 and CO_2) were used to assess the effect of the three electron acceptors (N_2O , NO_3^- and O_2) on C-mineralization. While oxygen appeared to have a marginal effect, NO_3^- and N_2O caused severe retardation of C-mineralization during the first 50 and 100 h, respectively (Supplementary Fig. S5A–D). This retarded mineralization is plausibly due to the inhibition of methanogenesis, causing a transient accumulation of H_2 and VFAs until the N_2O -reducing bacteria reach a cell density that allowed them to effectively reap these compounds. This was corroborated by measurements of H_2 and VFAs (Supplementary Fig. S13).

To track the origin of the enriched N_2O -respiring bacteria in the digestate, we considered the possibility that these are indigenous wastewater-sludge bacteria that survive the passage through the anaerobic digester, which had a retention time of 20–24 days. We assessed survival of N_2O -respiring bacteria by comparing the N_2O reduction potential of wastewater sludge and the digestate. The results indicated that $\leq 1/3$ of the N_2O -respiring bacteria in the sludge survived the passage (Supplementary Fig. S6). We also did enrichment culturing with a digestate from a thermophilic digester (52 °C) operated in parallel with the mesophilic digester (the same feed), and found that it contained N_2O reducers that could be enriched, although the estimated initial numbers were orders of magnitude lower than in the mesophilic digestate (Supplementary Fig. S7).

MAG-centric metaproteomic analysis of the enrichment cultures

We analyzed the metagenome and metaproteome at three timepoints (0, 115 and 325 h) during the first enrichment culturing (gas kinetics and sampling times shown in Fig. 2A), to explore the effect of the anaerobic incubation with N_2O on the entire microbial consortium, and to identify the organisms growing by N_2O reduction. Metagenomic sequences were assembled and resultant contigs assigned to 278 metagenome-assembled genomes (MAGs), of which 149 were deemed to be of sufficient quality (completeness > 50% and contamination < 20%, Supplementary Data S1) for downstream analysis. The phylogenetic

relationship and the relative abundance of the MAGs throughout the enrichment are summarized in Fig. 3, which also shows selected features revealed by the combined metagenomic and metaproteomic analyses, including information about genes and detected proteins involved in N_2O reduction, other denitrification steps, methanogenesis, syntrophic acetate oxidation and methane oxidation.

Closer inspections of the abundance of individual MAGs, based on their coverage in the metagenome and metaproteome, showed that the majority of the MAGs had a near constant population density throughout the incubation, while two MAGs (260 and 268) increased substantially (Fig. 4; further analyses in Supplementary Section B, Supplementary Figs. S8–S11). The stable abundance of the majority indicates that the methanogenic consortium remained intact despite the downshift in temperature (20 °C versus 37 °C) and the inhibition of methanogenesis by N_2O . Only 9 MAGs showed a consistent decline in abundance throughout the enrichment (Supplementary Table S1). These MAGs could theoretically correspond to microbes whose metabolism is dependent on efficient H_2 scavenging by methanogens [27], but we found no genomic evidence for this, and surmise that organisms circumscribed by the declining MAGs were unable to adapt to the temperature downshift from 37 °C to 20 °C.

Six MAGs, including the two that were clearly growing (MAG260 & MAG268) contained the *nosZ* gene and thus had the genetic potential to produce N_2O -reductase (Nos) (Fig. 4). Nos proteins originating from five of these MAGs were detected in the metaproteome. Importantly, while all but one of these MAGs contained genes encoding the other denitrification reductases, none of these were detected in the metaproteome, suggesting that the organisms can regulate the expression of their denitrification machinery to suit available electron acceptors, in this case N_2O . Three of the MAGs with detectable Nos in the proteome (MAG004, MAG059, MAG248) appeared to be non-growing during the enrichment. The detected levels of their Nos proteins remained more or less constant, and their estimated abundance in the metagenome and -proteome did not increase (Fig. 4B). It is conceivable that these three MAGs belong to the initial population of N_2O reducers whose N_2O -reduction activity was present initially but gradually decreased during the early phase of the enrichment (Fig. 2A). The two growing MAGs (MAG260 and MAG268) showed increasing Nos levels and increasing abundance both in terms of coverage and metaproteomic detection (Fig. 4B), in proportion with the N_2 produced (Supplementary Fig. S11). MAG260 reached the highest abundance of the two and accounted for 92% of the total detectable Nos pool at the final time point. MAG260 is taxonomically most closely affiliated with the genus *Dechloromonas* (GTDB, 97.9% amino acid similarity). Interestingly, Nap rather than Nar takes the role of nitrate reductase in MAG260 (Fig. 4), which makes it a promising organism for N_2O mitigation since organisms with Nap only (lacking Nar) preferentially channel electrons to N_2O rather than to NO_3^- [18]. MAG260, MAG004 and MAG088 contain a clade II *nosZ*, characterized by a *sec*-dependent signal peptide, in contrast to the more common *tat*-dependent clade I *nosZ* [16]. The physiological implications of clade I versus clade II *nosZ* remains unclear. Organisms with *nosZ* Clade II have high growth yield and high affinity (low k_s) for N_2O , compared to those with *nosZ* Clade I [28], suggesting a key role of *nosZ* Clade II organisms for N_2O reduction in soil, but this was contested by Conthe et al. [29], who found that Clade I organisms had higher “catalytic efficiency” (V_{max}/k_s) than those with Clade II.

The apparent inhibition of methanogenesis by N_2O seen in the present study has been observed frequently [30] and is probably due to inhibition of coenzyme M methyltransferase [31], which is a membrane bound enzyme essential for methanogenesis and common to all methanogenic archaea [32]. The gas kinetics demonstrate that the inhibition was reversible, being partly

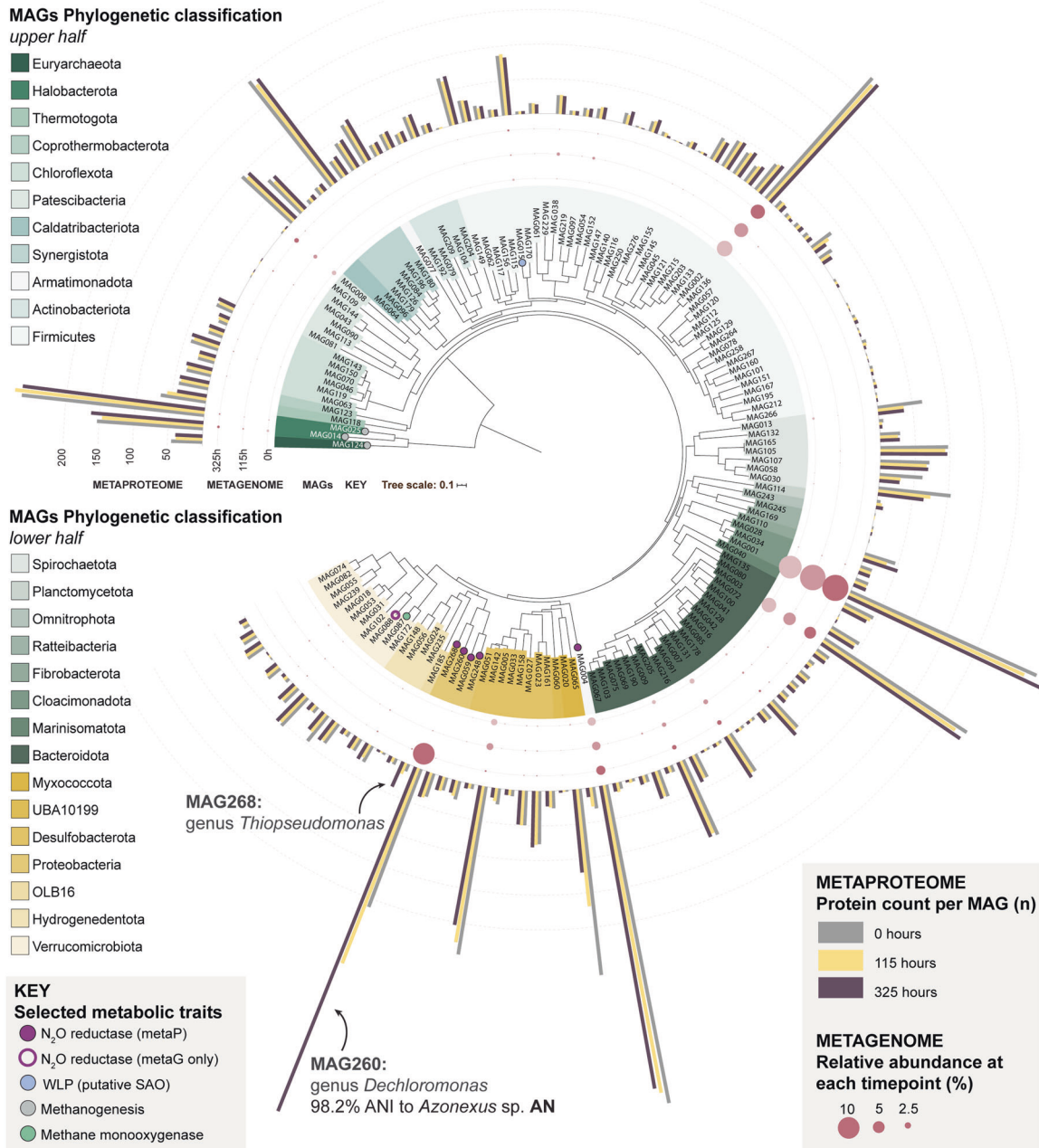


Fig. 3 Metagenome Assembled Genomes (MAGs) from the anaerobic enrichment culture with the mesophilic digestate. Maximum likelihood tree indicating the phylogenetic placement of MAGs from the anaerobic enrichment. The tree was constructed from a concatenated set of protein sequences of single-copy genes. Taxonomic classification of the MAGs was inferred using the Genome Taxonomy Database (GTDB) and is displayed at the phylum level by label and branch coloring. Branch label decorations indicate the presence of genes involved in selected metabolic traits in the MAGs. The relative abundance of the MAG in the community as calculated from sequence coverage is indicated by bubbles at branch tips and bar charts indicate the number of detected proteins affiliated with each MAG at the three time points during incubation. Four of the 149 MAGs that met the completeness and contamination threshold for construction of the metaproteome database were lacking the universal single-copy marker genes and were omitted from the tree. Total protein counts per MAG were calculated by aggregating both secretome and cell-associated proteomes.

restored whenever N₂O was depleted (Fig. 2). In the enrichment culture where metagenomics and metaproteomics was monitored, several such incidents of N₂O depletion occurred (Fig. 2A) and during these periods CH₄ accumulated to levels amounting to 10% of levels in control vials without N₂O (Supplementary Fig. S4B). These observations suggest that methanogens would be able to grow, albeit sporadically, during the enrichment, which is corroborated by the sustained detection of the complete methanogenesis pathway, including the crucial coenzyme M methyl-transferase, of *Methanotherix* (MAG025), *Methanoregulaceae*

(MAG014) and *Methanobacterium* (MAG124) at high levels in the metaproteome. In fact, both MAG coverage data and 16S rRNA gene copy numbers assessed by ddPCR suggested that the majority of the original methanogenic consortium continued to grow (Supplementary Section B). A tentative map of the metabolic flow of the methanogenic consortium, including the reaping of intermediates (monosaccharides, fatty acids, acetate and H₂) by N₂O-respiring bacteria is shown in Supplementary Fig. S12. Since methane production was inhibited from the very beginning of the incubation, while it took ~100 h for the N₂O-respiring bacteria to

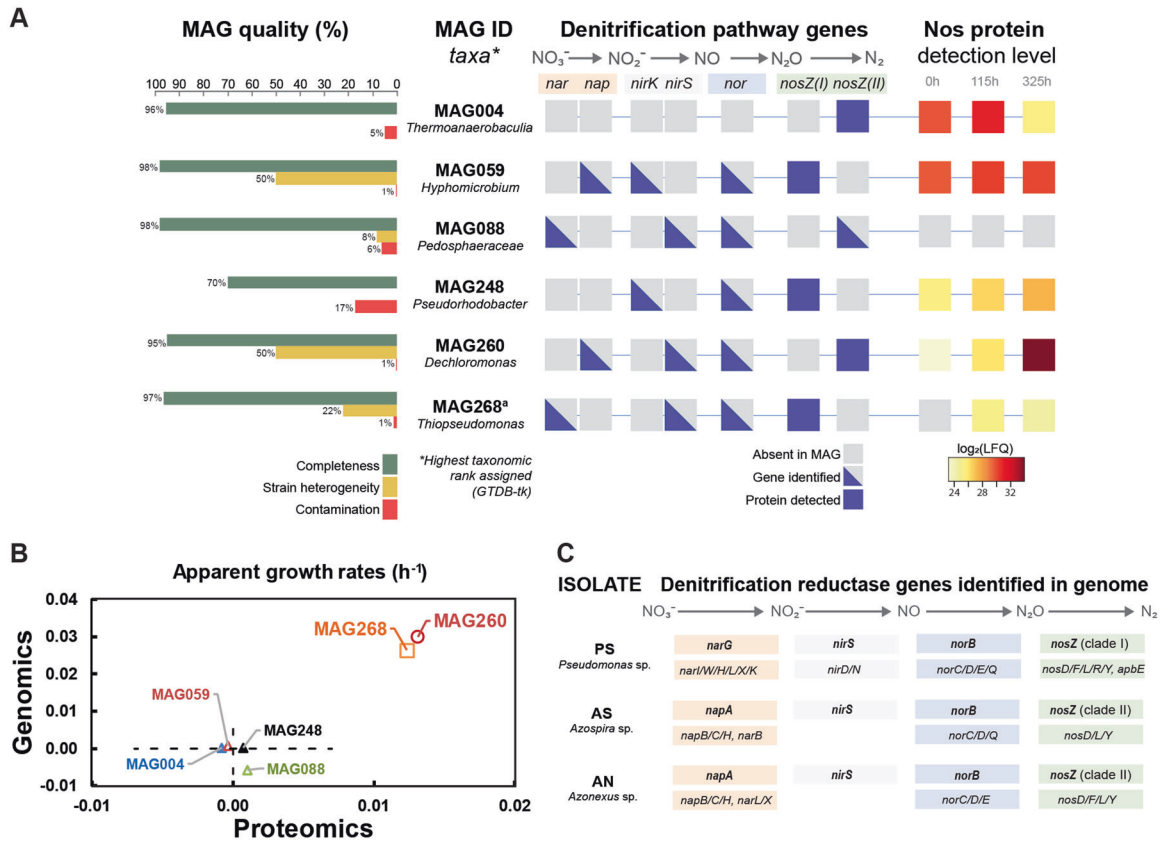


Fig. 4 Overview of MAGs with *nosZ* and denitrification genes in isolated organisms. Panel **A** shows the quality (completeness, strain heterogeneity and contamination), taxonomic classification based on GTDB and NCBI, presence of denitrifying genes and proteins, and the detected levels of Nos (N_2O reductase, encoded by *nosZ*) throughout the enrichment culturing for the six MAGs containing the *nosZ* gene (Supplementary Data S1 and S2). Nos was detected in the proteome of five MAGs, but the detection level increased significantly throughout for only MAG260 and 248, respectively. None of the MAGs produced detectable amounts of the other denitrification reductases. (a) Lable Free Quantification (LFQ) values for one of the two detected predicted Nos proteins for MAG268 is shown. Panel **B** shows the apparent growth rates of the MAGs, based on their coverage in the metagenome and metaproteome (regression of $\ln(N)$ against time; see Supplementary Fig. S11 for more details). Panel **C** shows the taxonomic classification (16S rDNA), working names (abbreviations) and denitrification genotypes of three isolates from the enrichment culturing. The genes coding catalytic subunits of denitrification reductases are shown in bold, above the accessory genes [55] that were also identified. More information about accessory genes is presented in Supplementary Fig. S15. The isolate AN has 98.2% ANI to MAG260.

reach high enough numbers to become a significant sink for intermediates (Fig. 2), one would expect transient accumulation of volatile fatty acids and H_2 , which was corroborated by measurements of these metabolites (Supplementary Fig. S13).

Of note, we detected methane monooxygenase and methanol dehydrogenase proteins from MAG087 and MAG059, respectively, in the metaproteome. This opens up the tantalizing hypothesis of N_2O -driven methane oxidation, a process only recently suggested to occur [33, 34]. However, a close inspection of the N_2O - and CH_4 -kinetics indicated that N_2O -driven methane oxidation played a minor role (Supplementary Fig. S4CD).

In a follow-up experiment, 7 parallel enrichment cultures were analyzed by 16S rRNA gene amplicon sequencing, demonstrating reproducibility of the selective enrichment of organisms circumscribed by MAG 260 (Fig. S14).

Isolation of N_2O -respiring bacteria and their geno- and phenotyping

Whilst this enrichment culture could be used directly as a soil amendment, this approach is likely to have several disadvantages. First, it would require the use of large volumes of N_2O for enrichment, a process which would be costly and require significant infrastructure. An alternative approach would be to introduce an axenic or mixed culture of digestate-derived, and

likely digestate-adapted, N_2O -respiring bacteria to sterilized/sanitized digestates. This approach has multiple benefits: (1) it would remove the need for N_2O enrichment on site as isolates could be grown aerobically in the digestate material, (2) one could chose organisms with favorable denitrification genotypes and regulatory phenotypes, (3) the sanitation would eliminate the methanogenic consortium hence reducing the risk of methane emissions from anoxic micro-niches in the amended soil, and (4) sanitation of digestates aligns with current practices that require such a pretreatment prior to use for fertilization. For these reasons an isolation effort was undertaken to obtain suitable digestate-adapted N_2O -respiring microorganisms from the N_2O -enrichment cultures (Supplementary Material and Methods, Chapter 6). These efforts resulted in the recovery of three axenic N_2O -respiring bacterial cultures, which were subjected to subsequent genomic and phenotypic characterization.

The isolates were phylogenetically assigned to *Pseudomonas* sp. (PS), *Azospira* sp. (AS) and *Azonexus* sp. (AN) (working names in bold) based on full length 16S rRNA genes obtained from the sequenced genomes (accessions ERR4842639 – 40, Supplementary Table S2, phylogenetic trees shown in Supplementary Fig. S15). All were equipped with genes for a complete denitrification pathway (Fig. 4C). AN and AS carried *napAB*, encoding the periplasmic nitrate reductase (Nap) and *nosZ* clade

II, whilst PS carried genes for the membrane bound nitrate reductase (Nar), encoded by *narG*, and *nosZ* clade I. All had *nirS* and *norBC*, coding for nitrite reductase (NirS) and nitric oxide reductase (Nor), respectively. Pairwise comparison of average nucleotide identities (ANI) with MAGs from the enrichment metagenomes showed that the isolate AN matched the *Dechloromonas*-affiliated MAG260 with 98.2% ANI, suggesting the isolate is circumscribed by MAG260 [35]. Given the GTDB phylogeny of AN and MAG260 and the 16S rRNA gene homology of AN (95.2% sequence identity to *Azonexus hydrophilus* DSM23864, Supplementary Fig. S15C), we conclude that AN likely represents a novel species within the *Azonexus* lineage. Unfortunately, the 16S rRNA gene was not recovered in MAG260, preventing direct comparison with related populations. No significant ANI matches in our MAG inventory were identified for the genomes of PS and AS indicating they were not captured via our metagenomic approaches, which highlights the complementarity of applying culture-dependent methods in parallel.

The carbon catabolism profiles of the isolates were assayed using Biolog PM1 and PM2 microplates, to screen the range of carbon sources utilized (Supplementary Section E: Supplementary Table S3). PS utilized a wide spectrum of carbon sources (amino acids, nucleic acids, volatile fatty acids (VFA), alcohols, sugar alcohols, monosaccharides and amino sugars), but only one polymer (laminarin). AN and AS could only utilize small VFAs (e.g. acetate, butyrate), intermediates in the TCA cycle and/or the β -oxidation/methyl malonyl-CoA pathways of fatty acid degradation (e.g. malate, fumarate, succinate), and a single amino acid (glutamate). Thus, all three would be able to grow in a live digestate by reaping the VFA's produced by the methanogenic consortium. While the utilization of VFAs as C-substrates is one of several options for PS, AN and AS appear to depend on the provision of VFAs. This was confirmed by attempts to grow the three isolates in an autoclaved digestate: while PS grew well and reached high cell densities without any provision of extra carbon sources, AN and AS showed early retardation of growth unless provided with an extra dose of suitable carbon source (glutamate, acetate, pyruvate or ethanol) (Supplementary Figs. S26 and S27). A high degree of specialization and metabolic streamlining may thus explain the observed dominance of AN (MAG260) during enrichment culturing.

To evaluate the potentials of these isolates to act as sinks for N_2O , we characterized their denitrification phenotypes, by monitoring kinetics of oxygen depletion, subsequent denitrification and transient accumulation of denitrification intermediates (NO_2^- , NO, N_2O). The experiments were designed to assess properties associated with strong N_2O reduction such as 1) *bet hedging*, i.e. that all cells express N_2O reductase while only a fraction of the cells express nitrite- and/or nitrate-reductase, as demonstrated for *Paracoccus denitrificans* [17]; 2) strong metabolic preference for N_2O -reduction over NO_3^- -reduction, as demonstrated for organisms with periplasmic nitrate reductase [18]. Supplementary section F (Supplementary Figs. S16–S25) provides the results of all the experiments and a synopsis of the findings. In short: *Azonexus* sp. (AN) had a clear preference for N_2O over NO_3^- reduction, but not over NO_2^- reduction, ascribed to *bet hedging* with respect to the expression of nitrate reductase (a few cells express Nap, while all cells express Nos), which was corroborated by proteomics: the Nos/Nap abundance ratio was ~ 25 during the initial phase of denitrification (Supplementary Fig. S18). *Azospira* sp. (AS) had a similar preference for N_2O over NO_3^- reduction, albeit less pronounced than in AN, and no preference for N_2O over NO_2^- . *Pseudomonas* sp. (PS) showed a phenotype resembling that of *Paracoccus denitrificans* [17], with denitrification kinetics indicating that Nir is expressed in a minority of cells in response to O_2 depletion, while all cells appeared to express N_2O reductase. This regulation makes PS a more robust sink for N_2O than the two

other isolates, since it kept N_2O extremely low even when provided with NO_2^- .

In summary, PS appeared to be the most robust candidate as a sink for N_2O in soil for two reasons; 1) it can utilize a wide range of carbon substrates, and 2) its N_2O sink strength is independent of the type of nitrogen oxyanion present (NO_2^- or NO_3^-). In contrast, AN and AS appear to be streamlined for harvesting intermediates produced by anaerobic consortia, hence their metabolic activity in soil could be limited. In addition, they could be sources rather than sinks for N_2O if provided with NO_2^- , which is likely to happen in soils, at least in soils of neutral pH, during hypoxic/anoxic spells [36].

Effects on N_2O emissions

To assess if fertilization with digestates containing N_2O -reducing bacteria could reduce N_2O emissions from denitrification in soil, we conducted a series of incubation experiments with soils fertilized with digestates with and without N_2O -reducing bacteria. The fertilized soils were incubated in closed culture vials containing He + 0.5 vol % O_2 , and O_2 , NO, N_2O and N_2 were monitored during oxygen depletion and subsequent denitrification. The experiments included soils amended with digestates in which indigenous N_2O -reducing bacteria had been enriched by anaerobic incubation with N_2O (Fig. 2), as well as autoclaved digestates in which the isolates from the current study had been grown by aerobic cultivation (see Supplementary Figs. S26 and S27 for cultivation details). The experiments included three types of control digestates: digestate (directly from the digester), digestate heated to 70 °C for 2 h (to eliminate most of the indigenous consortium), and autoclaved digestate in which the strain PS had been grown aerobically and then heated to 70 °C for 2 h, to kill PS. In all cases, 3 mL of digestate was added to 10 g of soil. Since soil acidity has a pervasive effect on the synthesis of functional N_2O reductase [24], we tested the digestates with two soils from a liming experiment [37] with different pH ($pH_{CaCl_2} = 5.5$ and 6.6).

The transient N_2O accumulation during denitrification was generally higher in the acid than in the near-neutral soil (Fig. 5), which was expected since the synthesis of functional Nos is hampered by low pH [23, 24]. Based on the kinetics of both N_2 and N_2O (see Supplementary Figs. S28 and S29), we calculated the N_2O -index (I_{N_2O}) which is a measure of the molar amounts of N_2O relative to $N_2 + N_2O$ in the headspace for a specific period (0–T), see equation at top of Fig. 5). Low values of I_{N_2O} indicate efficient N_2O -reduction. In this case, we calculated I_{N_2O} for the incubation period until 40% of the available NO_3^- had been recovered as $N_2 + N_2O$ ($=I_{N_2O_{40}}$) and for the incubation period until 100% was recovered ($I_{N_2O_{100}}$). Statistical analyses showed significant effects of digestate treatments and soil pH, and the interaction between the two ($p < 0.001$).

Extremely low I_{N_2O} values were recorded for the treatments with digestate in which N_2O -reducing bacteria were enriched by anaerobic incubation with N_2O , even in the acid soil. This is in line with the current understanding of how pH affects N_2O -reduction: low pH slows down the synthesis of functional Nos, but once synthesized, it remains functional even at low pH [23]. Functional Nos had already been expressed during the enrichment and was evidently active after amendment to the soils.

The presence of the isolates in the digestates had clear but variable effects on I_{N_2O} . The most relevant control-treatment for evaluating the effect of the isolates would be PS_70 °C, because this digestate had been treated exactly the same way as that with metabolically active isolates present (autoclaved, aerated and aerobic cultivation). These contrasts (I_{N_2O} for PS_70 °C versus PS, AS and AN) were all statistically significant (confidence intervals did not overlap), thus all strains reduced I_{N_2O} compared to PS_70 °C. AN and AS resulted in much higher I_{N_2O} in the acid than in the neutral, suggesting that the expression of functional N_2O reductase in these strains was hampered by low pH. In contrast

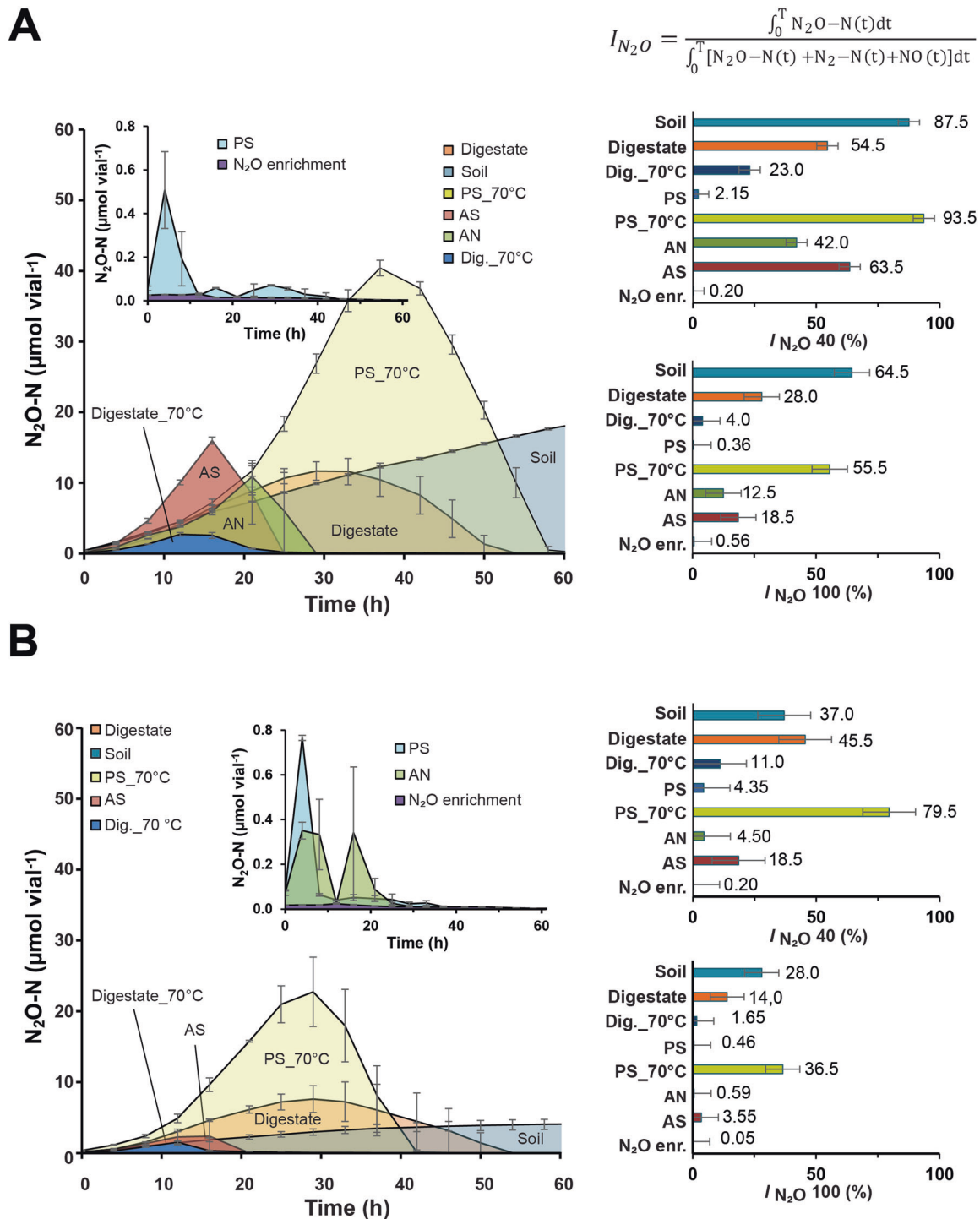


Fig. 5 Soil incubations, effects on N_2O emission. N_2O kinetics during incubation of soils amended with seven different digestates and a control (soil only); panel **A**: pH 5.5 soil, panel **B**: pH 6.6 soil. The digestates treatments are: Soil = soil without any amendment; Digestate = digestate directly from the anaerobic digester; Dig_70°C = digestate heat treated to 70 °C for 2 h; PS, AN and AS = autoclaved digestate in which isolates PS, AN and AS (respectively) had been grown aerobically (see Supplementary Figs. S26 and S27); PS_70°C = digestate in which PS had been grown (as for the PS treatment), subsequently heated to 70 °C for 2 h, N_2O enr. = digestate in which indigenous N_2O -respiring bacteria had been enriched (as shown in Fig. 2). The left panels show measured N_2O throughout soil incubations; the insets with altered scaling show N_2O levels for treatments that accumulated very little N_2O . The bar graphs to the right show the N_2O indexes (I_{N_2O}) expressed as % (equation shown in the panel), with confidential intervals (ANOVA + Tukey's range test). I_{N_2O} is a proxy for potential N_2O emission from denitrification in soil [56]. Two I_{N_2O} values are shown: one for the timespan until 40% of the NO_3^- -N was recovered as $N_2 + N_2O + NO$ (I_{N_2O} 40), and one for 100% recovery (I_{N_2O} 100). More details (including N_2 and NO kinetics) are shown in Supplementary Figs. S28 and S29.

to AN and AS, PS resulted in very low I_{N_2O} values in both soils, suggesting that this organism has an exceptional capacity to synthesize functional Nos at low pH.

These results show that the emission of N_2O from soil fertilized with digestates can be manipulated by tailoring the digestate microbiome. Interestingly, measurements of methane in these soil incubations showed that the methanogenic consortia in digestates that had not been heat treated (i.e. the live digestate and the N_2O enrichment) remained metabolically intact in the soil, and started producing methane as soon as N_2O and nitrogen oxyanions had been depleted, while no methane was produced in the soils amended with autoclaved digestate, and that heated to 70 °C (Supplementary Fig. S30).

In an effort to determine the survival of the N_2O -scavenging capacity of a digestate enriched with N_2O reducers, we also tested its effect on soil N_2O emissions after a 70-h aerobic storage period (in soil or as enrichment culture, at 20 °C). These experiments demonstrated a sustained beneficial effect on I_{N_2O} after 70 h of aerobic storage (Supplementary Fig. S31). This result indicates that the enrichment strategies discussed here are robust, although long-lasting storage experiments as well as field trials are needed.

CONCLUDING REMARKS

This feasibility study identifies an avenue for large scale cultivation of N_2O reducers for soil application, which could be low cost if implemented as an add-on to biogas production systems. Further efforts should be directed towards selecting organisms that are both strong sinks for N_2O and able to survive and compete in soil, to secure long-lasting effects on N_2O emissions. A tantalizing added value would be provided by selecting organisms (or consortia of organisms) that are not only strong N_2O -sinks, but also promote plant growth and disease resistance [38, 39].

Gas kinetics, metagenomics and metaproteomics revealed that the methanogenic consortium of the digestate remains active during anaerobic incubation with N_2O , and that bacteria with an anaerobic respiratory metabolism grew by harvesting fermentation intermediates. The inhibition of methanogenesis by N_2O implies that the respiring organisms would have immediate access to the electron donors that would otherwise be used by the methanogens, i.e. acetate and H_2 , while they would have to compete with fermentative organisms for the “earlier” intermediates such as alcohols and VFA. The importance of fermentation intermediates as a carbon source for the N_2O -respiring bacteria would predict a selective advantage for organisms with a streamlined (narrow) catabolic capacity, i.e. limited to short fatty acids, and our results lend some support to this: the catabolic capacity of the organism that became dominant (MAG260, isolate AN) was indeed limited, as was also the case for isolate AS. Such organisms are probably not ideal N_2O -sinks in soil because their ability to survive in this environment would be limited. Organisms with a wider catabolic capacity, such as the isolated *Pseudomonas* sp. (PS), are stronger candidates for long term survival and N_2O -reducing activity in soil. The ideal organisms are probably yet to be found, however, and refinements of the enrichment culturing process are clearly needed.

The digestate used in this study contained N_2O -respiring bacteria, most likely survivors from the raw sludge, which however, were clearly outnumbered by bacteria that are net producers of N_2O . We surmise that the relative amounts of N_2O -producers and N_2O -reducers in digestates may vary, depending on the feeding material and configuration for the anaerobic digestion. This could explain the observed variable effects of digestates on N_2O emission from soils [40, 41]. The high abundance of both NO_3^- - and O_2 -respiring organisms in digestates has practical implications for the attempts to grow isolated strains in digestates: they could be outnumbered by the indigenous NO_3^- - and O_2 -respiring organisms (Supplementary

Fig. S5). Hence, we foresee that future implementation of this strategy will require a brief heat treatment or other sanitizing procedure. A bonus of such sanitation is that it eliminates methane production by the digestate in soil.

We failed to enrich organisms lacking all other denitrification genes than *nosZ*; the only reconstructed genome with *nosZ* only (MAG004) did not grow at all. Failure to selectively enrich such organisms by anaerobic incubation with N_2O was also experienced by Conthe et al. [29]. The organisms that did grow by respiring N_2O in our enrichment, were all equipped with genes for the full denitrification pathway, although the only denitrification enzyme expressed/detected during the enrichment was Nos. This agrees with the current understanding of the gene regulatory network of denitrification; *nosZ* is the only gene whose transcription does not depend on the presence of NO_3^- , NO_2^- or NO [42], which were all absent during the enrichment.

Two of the reconstructed MAGs had genes encoding periplasmic nitrate reductase (*nap*), as was the case for two of the three isolates (AN and AS). This in itself would predict preference for N_2O - over NO_3^- reduction at a metabolic level [43], but otherwise their potential for being N_2O sinks cannot be predicted by their genomes. The phenotyping of the isolates revealed conspicuous patterns of *bet hedging* as demonstrated for *Paracoccus denitrificans* [17]. The *bet hedging* in *P. denitrificans* is characterized by expression of Nir (and Nor) in a minority of the cells, while Nos is expressed in all cells, in response to oxygen depletion, hence the population as a whole is a strong sink for N_2O . The isolated *Pseudomonas* sp. (PS) displayed denitrification kinetics that closely resembles that of *P. denitrificans*. The two other isolates (AN and AS) showed indications of *bet hedging* as well, but of another sort: Nap appears to be expressed in a minority of the cells. This different regulatory phenotype had clear implications for the ability of organisms to function as N_2O -sinks: while all isolates were strong N_2O sinks when provided with NO_3^- only, AN and AS accumulated large amounts of N_2O if provided with NO_2^- .

The N_2O sink capacity of the organisms was tested by fertilizing soils with digestates with and without the organisms, and monitoring the gas kinetics in response to oxygen depletion, thus imitating the hot spots/hot moments of hypoxia/anoxia [44]. Since the isolates were raised by aerobic growth in autoclaved digestates, they would have to synthesize all denitrification enzymes in the soil, hence the synthesis of functional Nos was expected to be hampered by low pH [24]. The results for isolates AS and AN lend support to this (high I_{N_2O} in the soil with pH = 5.5). AN was also dominating in the digestate enrichment culture, and in this case the organism had a strong and pH-independent effect on N_2O emission, plausibly due synthesis of Nos prior to incorporation into the soils.

In summary, we have demonstrated that a digestate from biogas production can be transformed into an effective agent for mitigating N_2O emission from soil, simply by allowing the right bacteria to grow to high cell densities in the digestate prior to fertilization. The technique is attractive because it can be integrated in existing biogas production systems, and hence is scalable. If we manage to treat a major part of waste materials in agroecosystems by AD, the resulting digestates would suffice to treat a large share of total farmland, as illustrated by Fig. 1. Estimation of the potential N_2O -mitigation effect is premature, but the documented feasibility and the scalability of the approach warrant further refinement as well as rigorous testing under field condition. Our approach suggests one avenue for a much needed valorization of organic wastes [45] via anaerobic digestion. Future developments of this approach could extend beyond the scope of climate change mitigation and include the enrichment of microbes for pesticide- and other organic pollutant degradation [46], plant growth promotion [47] and inoculation of other plant symbiotic bacteria [48].

DATA AVAILABILITY

The sequencing data for this study have been deposited in the European Nucleotide Archive (ENA) at EMBL-EBI under accession number PRJEB41283 (isolates AN, AS and PS) and PRJEB41816 (metagenome) (<https://www.ebi.ac.uk/ena/browser/view/PRJEBxxxx>). Functionally annotated MAGs and metagenomic assembly are available in FigShare (<https://doi.org/10.6084/m9.figshare.13102451> and <https://doi.org/10.6084/m9.figshare.13102493>). The proteomics data has been deposited to the ProteomeXchange Consortium (<http://proteomecentral.proteomexchange.org>) via the PRIDE partner repository [49] with the dataset identifier PXD022030* and PXD023233** for the metaproteome and proteome of *Azonexus* sp. AN, respectively.

* Reviewer access: Username: reviewer_pxd022030@ebi.ac.uk. Password: GdTR3biE

** Reviewers access: Username: reviewer_pxd023233@ebi.ac.uk Password: nMz62S8O

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AUTHOR CONTRIBUTIONS

KRJ conducted the research, analyzed and interpreted the data, and wrote the manuscript, LHH conducted genomic and proteomic analyses of the enrichments, SHWV conducted the research and wrote the manuscript, MA contributed to analysis of the genomics and proteomics, VE and ÅF contributed to the writing of the manuscript, PL did proteomics on the isolate, LM was responsible for gas measurements and kinetics analyses, PBP contributed to the analysis of genomics and proteomics and to the writing of the manuscript, LRB designed the experiment, analyzed the data and wrote the manuscript.

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COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

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