Title

From Rags to Enriched: Metagenomic Insights into Ammonia-oxidizing Archaea Following Ammonia Enrichment of a Denuded Oligotrophic Soil Ecosystem.

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Running title: Ammonia-oxidizing archaea in oligotrophic semi-arid soils

Keywords: Ammonia-oxidizing archaea, Bioprospecting, MAGs, Oligotrophic soil, Semi-arid soils.

Originality-Significance Statement

The unique edaphic properties present across the Western Australian landscape makes it a valuable environment for bioprospecting, especially for taxa that provide essential ecological processes. Unfortunately, the natural terrestrial ecosystems of WA are also largely unknown.
We have discovered candidate taxa that could potentially provide an effective role in ecological restoration, which is critical as soil ecosystem disturbances are common due to the prevalent mining operations that occur in WA. The lessons learnt may also be applicable to other bioremediation settings.

**Reviewer link to data:**
https://dataview.ncbi.nlm.nih.gov/object/PRJNA775848?reviewer=ljdvtt4aplo4eg5ptsaovi8rk&t&archive=sra

**Summary**

Stored topsoil acts as a microbial inoculant for ecological restoration of land after disturbance, but the altered circumstances frequently create unfavorable conditions for microbial survival. Nitrogen cycling is a critical indicator for ecological success and this study aimed to investigate the cornerstone taxa driving the process. Previous *in-silico* studies investigating stored topsoil discovered persistent archaeal taxa with the potential for re-establishing ecological activity. Ammonia oxidization is the limiting step in nitrification and as such, ammonia oxidizing archaea (AOA) can be considered as the one of the gatekeepers for the re-establishment of the nitrogen cycle in disturbed soils. Semi-arid soil samples were enriched with ammonium sulfate to promote the selective enrichment of ammonia oxidizers for targeted genomic recovery, and to investigate the microbial response of the microcosm to nitrogen input. Ammonia addition produced an increase in AOA population, particularly within the genus *Candidatus Nitrosotalea*, from which metagenome-assembled genomes (MAGs) were successfully recovered. The *Ca. Nitrosotalea* archaeon candidates’ ability to survive in extreme conditions and rapidly respond to ammonia input makes it a potential bioprospecting target for application in ecological restoration of semi-arid soils and the recovered MAGs provide a metabolic blueprint for developing potential strategies towards isolation of these acclimated candidates.
Introduction

Mining operations often involve the clearing of vegetation, and removal and subsequent storage of topsoil, which serves as a critical reserve of soil microbial communities for post-mining rehabilitation and restoration of ecosystem processes (Williams et al., 2019). Long term storage has been shown to change community structure and diversity, which may have negative implications for nutrient cycling and other vital ecological processes (Ezeokoli et al., 2019). As soil microbial populations are essential for plant community development, successful restoration of degraded terrestrial ecosystems requires the input of effective soil inoculum (Wubs et al., 2016). In determining the microbial diversity and functional capacity of stored topsoil and substrate blends (topsoil and mine site waste substrates) in mine site restoration scenarios in Western Australian (WA) conditions, we showed phylogenetic divergence but “potential” functional redundancy for microbe-related functions and reported enrichment of archaeal taxa across different topsoil blends, indicating a persistent group of taxa (Kumaresan et al., 2017). Long-term storage of topsoil can contribute to loss of the seed bank, leading to the decline of the reemergence of native and endemic flora (Golos and Dixon, 2014). In a follow up study, we also showed that biocrusts, the aggregated layer containing crucial microbial community and the majority of biological activity within arid soils, can play an important role in nitrogen cycling within arid landscapes in WA soils (Moreira-Grez et al., 2019b). Soil biodiversity is key for ecosystem multifunctionality (Wagg et al., 2014) and ensuring microbial functional diversity in mine site restoration scenario is critical to enable critical nutrient cycling processes.

Nitrogen is considered a major limiting nutrient for plant growth within terrestrial ecosystems, and microbial processes facilitate availability of nitrogen compounds for plant growth i.e., nitrogen fixation, ammonia oxidation (Moreau et al., 2019). In the nitrification step of nitrogen cycling, ammonia oxidation is the initial and limiting step and is performed by
ammonia-oxidizing archaea (AOA), ammonia-oxidizing bacteria (AOB) and complete ammonia oxidation (comammox) bacteria (Kuypers et al., 2018; Lehtovirta-Morley, 2018). As such, these ammonia-oxidizing taxa are arguably the gatekeepers of nitrogen cycling and understanding their ecological presence is key to determining the sustainability of an ecosystem (Amoo and Babalola, 2017). AOA, in comparison to AOB, dominate soil microbiomes with low ammonium supply. The metabolic and ecological differences between archaeal and bacterial ammonia oxidizers that modulate this niche partitioning have been previously described in dryland soils (Trivedi et al., 2019). For instance, within the weathered lateritic agricultural WA soils, a reverse trend has been observed, where AOB are present in higher numbers compared to AOA. The authors speculate that the low copper levels (a key co-factor ammonia oxidation in AOA) in WA agricultural soils could drive the functions (Jenkins et al., 2016). Western Australian soils could be host to unique microbial communities that are adapted to the lateritic soils, which are often depleted of other nutrients, such as phosphorus, potassium and critically, nitrogen, due to extensive weathering and the binding of nutrients to lateritic compounds that make them unavailable for uptake (O’Brien et al., 2019).

In our previous work, we highlighted the higher relative abundance of novel 16S rRNA gene sequences related to archaeal taxa involved in nitrogen cycling. Unlike agricultural soils in Western Australia (Jenkins et al., 2016), the nitrifiers in natural ecosystems, particularly rich in iron-ore deposits are largely unknown. Shotgun metagenome sequencing has the potential to pinpoint cornerstone taxa that may play critical roles throughout the ecological restoration due to their capacity to link both taxonomic and functional information (Hart et al., 2020). Here, we aim to recover genomic data of ammonia oxidizing taxa from ammonium-enriched stored topsoil from the mining sites in semi-arid WA soils, with particular focus on the persistent archaeal presence observed in our previous work (Kumaresan et al., 2017; Moreira-Grez, et al., 2019a). Enrichment of the microcosm using inorganic ammonium coupled with metagenomic
sequencing led to improved genomic recovery of AOA that provided *in-silico* blueprints of the metabolic capacity of recovered candidates that can aid in developing isolation strategies for potential bioprospecting ventures. The enrichment was also used to compare the response of AOA and AOB groups to ammonium input in oligotrophic conditions. The metagenomic approach utilized in this study provided insight into the ecological drivers of ammonia-oxidizing taxa in the semi-arid soils from natural ecosystems in WA, with a focus towards their nitrogen cycling potential in the restoration of disturbed soils.

**Results**

*Temporal taxonomic profiles of the soil metagenomes*

Semi-arid soils subjected to enrichment with (NH$_4$)$_2$SO$_4$ showed similar proportions in the overall bacterial populations, with only slight variation in relative abundance observed across time points between both the amplicon (Figure 1a) and WMS data (Figure 1b). The most substantial increase in relative abundance after three-weeks of N enrichment were sequences related to archaeal phyla Thaumarchaeota, Aigarchaeota, Crenarchaeota and Korarchaeota (TACK) superphylum (Guy and Ettema, 2011) which contains the canonical AOA. This is most notable in the Archaea; Thaumarchaeota phylum from the WMS data, with an increase in relative abundance from 0.155% (baseline) to 5.629% (three weeks), a roughly 34 fold increase in metagenomic reads classified as Thaumarchaeota. The disparity between the phylum naming for the amplicon sequences and WMS data is likely due to the differences in the respective databases utilized by each method. The publicly available versions of the Greengenes 16S rRNA gene database uses older taxonomic terms, likely from when Thaumarchaeota was still classified as a ‘mesophilic Crenarchaeota’ (Pester et al., 2011). As the WMS classification is based on the latest NBCI non-redundant protein database, it provides a more up to date classification consistent with current literature. Another point of note is the replacement of Cyanobacteria from amplicon sequences to Gemmatimonadetes in the WMS data, most likely
due to misidentification of chloroplast sequences among the 16S amplicons (Hanshew et al., 2013). The diversity and evenness of the soil samples was shown to decrease over time post enrichment, likely due to the substantial change in relative abundance observed within the archaeal phyla, however this change was determined to be statistically not significant according to one-way ANOVA (Figure 2a). The clustering of the soil sample data in the non-metric multidimensional scaling (NMDS) analysis showed a clear successional trend observed in the microbial communities as time progressed post (NH₄)₂SO₄ enrichment (Figure 2b).

**Figure 1.** Taxonomic distribution at the phylum level of the soil metagenome from amplicon and whole metagenome shotgun sequencing data. Comparisons between baseline soil and time points post enrichment with ammonium sulfate. Amplicon sequencing data was used solely to determine relevant time points for the subsequent whole metagenome shotgun sequencing.
Figure 2. Measurements of microbial community structure at the genus level from Kaiju outputs using whole metagenome shotgun data. Baseline and treatment populations were investigated for temporal changes in A) Alpha-diversity though Shannon diversity index with differences between samples determined to be not significant via one-way ANOVA and B) Beta-diversity via non-metric multidimensional scaling of microbial community dissimilarity.

Effects of enrichment on ammonia oxidizing prokaryote populations

Sequences affiliated to AOA showed greater increase in relative abundance compared to AOB at the genus level across successive time points post enrichment with (NH₄)₂SO₄ (Figure 3a). In the baseline soils, AOA abundances were either at similar levels or lower than those of AOB with respect to relative proportion within the overall population. Over the course of the experimental period, the relative abundances of all detected AOA increased, whereas the AOB population remained similar or showed only slight increases on a subset of taxa. At the end of the three-week period, AOA from the genus Candidatus Nitrosotalea and Nitrososphaera showed higher relative abundances across all samples, with Ca. Nitrosotalea showing at least a 10-fold increase in relative abundance within two replicate samples compared to the highest reported AOB genus. Nitrospira showed a marginal increase at week three, however, as no MAGs were recovered from this genus, we were unable to determine if the reported populations
were composed of comammox or nitrite oxidizers. Correlations of ammonia-oxidizing taxa showed strong associations between the pairs of AOA and AOB (Figure 3b). In the baseline soils, multiple bacterial and archaeal taxa had a strong negative correlation, indicating competitive interactions between both groups. After three weeks, aside from *Nitrosospira*, these associations were no longer observed between archaeal and bacterial groups. Coupled with the substantial increase seen in AOA but not AOB numbers in the relative abundance data, suggests that within this subset of ammonia-oxidizing prokaryotes, archaea may have out-competed most bacteria groups after three weeks post enrichment.
Figure 3. Taxonomic profiling of a subset of ammonia-oxidizing taxa at the genus level from soil metagenome. A) Relative counts of archaea (circles) and bacteria (triangles) from baseline and three-week post ammonia sulfate enrichment. B) Correlation plot of AOA vs AOB (including Nitrospira). Area within the rectangle seen in the correlation plot highlights all archaea-bacteria associations.

Differential abundances of the metabolic profile of the metagenome post enrichment

At the level 3 subsystem category, the overall diversity and evenness of metabolic genes displayed a marginal increase across all soil samples as time progressed post enrichment,
however the one-way ANOVA testing determined that the changes observed were determined to be not significant (Figure 4a). In comparison, certain individual metabolic genes within the level 3 subcategory were significantly enriched or suppressed in the three-week post enrichment treatment sample compared to the baseline soil (Figure 4b, Supplementary Table 1). The genes of interest that were significantly enriched were those associated with archaeal biosynthesis and transcription. This pattern of diversity is also reflected in the taxonomic abundance data, in which the overall population showed little to marginal change, but substantial increases were seen in sequences associated with archaeal taxa.

**Figure 4.** Changes in metabolic gene distributions at SEED subsystem level 3 observed in the baseline and post ammonium sulfate enriched soils. A) Box plot of Shannon diversity and evenness index of metabolic genes showing overall changes from all time points. One-way ANOVA determined differences between samples were not significant. B) Volcano Plot produced from ALDEx2 analysis of metabolic genes from baseline samples vs three-weeks post enrichment. Data points in red indicate that the metabolic gene in question was either significantly enriched or suppressed in comparison to others according to Welch’s t and Wilcoxon rank test ($\alpha < 0.05$).
Archaeal genomes recovered from soil metagenome

A total of 5 archaeal MAGs were recovered from across all soil metagenomes that passed the completeness and contamination protocols. As GTDB-tk was used to classify the MAGs, the GTDB naming convention will be used to categorize the taxonomic labels for all resultant data involving the MAGs recovered in the study. Four MAGs were classified from canonical AOA genera, three from genus *Nitrosotalea* (CheckM reported completeness - nc_117: 64.56%, t2_109: 58.90% and t3_48: 92.23%) and one from genus *Nitrososphaera* (completeness - t2_80: 78.80%). The remaining MAG was an archaeon from order Woesearchaeales. Additional information on all recovered MAGs such as quality and assembly statistics, assigned classification, classification method, predicted tRNAs and rRNA can be found in Supplementary Table 2.

Effects of enrichment on diversity capture and sequence utilization in metagenomic assemblies

The redundancy estimation by Nonpareil showed increased rates of coverage for the diversity observed within the post enrichment soil metagenome compared to that of the baseline soils, given similar sequencing efforts (Figure 5). The improved coverage likely reflects the decreased diversity observed in the post-enrichment metagenomes (Figure 2a), which is also supported by the values reported by the Nonpareil diversity index (Baseline: 23.81, Two-weeks: 23.24, Three-weeks: 22.64). Read mapping of the baseline, two-week and three-week metagenomic reads to their respective assembled sequences reported an overall alignment rate of 46.69%, 53.51% and 57.71% respectively, which closely mirrors the values presented by the estimated average coverage seen on the Nonpareil curves to their respective datasets. Improvements in overall alignment rates from the post-enrichment metagenomes were also seen downstream in the recovered AOA candidate MAGs, with a clear pattern observed in the *Nitrosotalea* candidates, nc_117 (baseline): 0.02%, t2_109 (two-weeks): 0.07% and t3_48
(three-weeks): 0.17%, along with the sole *Nitrososphaera* candidate MAG t2_80 (two-weeks): 0.29%. The increased alignment rates in the post-enrichment metagenomes and MAGs demonstrate that more reads are being utilized in assembled sequences, likely due to the reduction in diversity resulting in better coverage. Increased read maps in the *Nitrosotalea* candidate MAGs from successional metagenomes also suggest that the more abundant *Nitrosotalea* subpopulations (Figure 3a) observed in the post-enrichment microbiomes have improved the genomic capture rate of this taxa.

**Figure 5.** Estimation of the coverage of diversity between the baseline and post enrichment soil metagenomes using Nonpareil. The empty circles on the curves show the actual sequencing volume of the respective datasets. The dashed and solid red horizontal lines show the 95% and 100% estimated coverage threshold respectively. The arrows are a visual indicator of the Nonpareil diversity index of each dataset, with arrows further right indicating more diverse communities.
Phylogenomic comparison and distributions of nitrogen metabolism pathways of interest

The four archaeal MAGs classified as AOA candidates appeared to be closely related to the known ammonia-oxidizers *Nitrososphaera* and *Nitrosotalea*, which is congruent with the genus classification assigned by GTDB-tk (Figure 6). All the MAGs identified as genus *Nitrosotalea* (nc_117, t2_109 and t3_48) were found to contain *amoA* and the t3_48 MAG which had the highest completion, showed a nitrogen metabolic pathway pattern consistent with several *Nitrosotalea* and other AOA genomes. Although the genus *Nitrososphaera* MAG (t2_80) was phylogenetically assigned within the *Nitrososphaera* clade, it is missing the AmoCAB node or results from Pfam/TIGRFAM searches required to definitively confirm it as an AOA. Given the phylogenetic placement, this is most likely due to missing data resulting from the MAG being incomplete, rather than an actual absence of ammonia monooxygenase.
Figure 6. Maximum likelihood tree and nitrogen pathways of archaeal genomes. Ammonia-oxidizing archaea (AOA) genomes from RefSeq were phylogenetically compared to the potential AOA MAGs recovered from this study (indicated by bold labels). Completeness of MAGs were reported by CheckM. Nitrogen pathways displayed were predicted by MinPath using the KEGG metabolic pathway models and are represented by shaded boxes indicating the predicted presence of a node within the respective nitrogen metabolic pathway. Shaded stars are nodes that were not present in the MinPath prediction but were detected via HMM search using the Pfam A database. Bootstrap values of \( \geq 95\% \) are shown by solid dots. Coloured backgrounds indicate AOA groups: Marine Group I (MG-I); South African Gold Mine Crenarchaeotic Group 1 (SAGMCG-1); Soil Crenarchaeotic Group (SCG); Hot Water Crenarchaeotic Group III (HWCGIII) and Thermophilic AOA (ThAOA).

Discussion

Metagenomics provides a primary approach into the observations of the \textit{in-situ} metabolic potential of environmental microorganisms and their response to ecological changes. Furthermore, the recovery of MAGs allows us to understand the metabolic capacity of a microbiome within the context of the individual microorganism present. The prediction of metabolic potential, however, does not automatically guarantee function. A pertinent example is the denitrification process where varying ecological factors determine whether transcription occurs or even if functional enzymes are produced, despite the presence of denitrification genes within the genome (Lycus et al., 2017). Another issue is that due to the limitations of short-read sequencing, the MAGs often remain drafts, with the missing data obscuring valuable ecological and functional information. Nonetheless, extensive \textit{in-silico} approaches such as this study, provide the groundwork to explicitly select candidate taxa for potential bioprospecting.
ventures and the ability to further explore other biosynthetic pathways present, within the context of the ecological question at hand.

From the denuded environment present in the stored topsoil, archaea appeared to have displayed the greatest relative response to the \((\text{NH}_4\text{)}_2\text{SO}_4\) enrichment. This is reflected in the significant increases seen within Thaumarchaeota at the phylum level, AOA at the genus level, as well as in archaeal-associated metabolic gene groups among the metagenomic reads. Studies into archaeal involvement in plant-microbe interactions have shown that not only do archaea participate in nutrient cycling, but also promote growth, improve disease resistance, and mitigate stress factors in plants (Jung et al., 2020). This suggests that stored topsoil treated with \((\text{NH}_4\text{)}_2\text{SO}_4\) to stimulate the growth of AOA populations, can be used to improve soils for post land-use rehabilitation.

In this pilot study, we investigated the potential of using inorganic nitrogen addition within the context of soil restoration and how it can modulate nitrifier activity/diversity. However, the experiments on the basal soils did not account for the growth of AOAs without enrichment. Further AOA propagation studies within stored topsoil should consider baseline community succession i.e., without nutrient input, to account for other potential ecological interactions within the microcosm. Future field-scale trials will certainly account for other contributing factors such as nitrogen fixation, plant-microbe interactions, among other variables, that can modulate nitrogen availability. Here, our focus was to identify keystone taxa and recover the genomes of indigenous archaeal nitrifiers.

Variations in the observed AOA and AOB population numbers between baseline and enriched soils provided valuable insight into the potential effects of N addition within natural ecosystems in WA. Short-term addition of inorganic ammonia in the form of \((\text{NH}_4\text{)}_2\text{SO}_4\) appeared to promote the rapid growth of AOA in the semi-arid topsoil. The large increase in AOA population numbers compared to AOB post \((\text{NH}_4\text{)}_2\text{SO}_4\) enrichment may be due to the
inhibitory effect of high ammonia concentrations, which has been observed to affect the growth of AOB but not AOA (Liang et al., 2020). This niche differentiation of AOA and AOB could be utilized when developing fertilization strategies for the promotion of AOA growth for nitrification within the semi-arid soils present in WA.

Further studies exploring ammonia-oxidizing taxa is crucial for understanding the ecological processes that drive nitrifying microorganisms within the WA natural soil ecosystems. Ecological studies of ammonia-oxidizers have been carried out in managed soils in WA, particularly within agricultural soils (O’Sullivan et al., 2013; Banning et al., 2015; Fisk et al., 2015; Jenkins et al., 2016), but very little work has been done in native terrestrial environments. A thorough understanding of the edaphic properties of soil in natural ecosystems and how they affect ammonia-oxidizing taxa are essential for the restoration of disturbed soils to their state prior to land use. This process is vital for screening ideal candidate taxa acclimated to the ecological conditions present within the unique soil ecosystems present within the natural WA landscape.

Within the dominant post-enrichment AOA population, the genus *Ca. Nitrosotalea* displayed the greatest relative abundance in archaeal reads and provided the highest number of recovered archaeal MAGs. The *Ca. Nitrosotalea* archaeon thrive in acidic environments (pH 4.5 – 5.5) through high-affinity substrate acquisition and pH homeostasis systems which confer the ability to oxidize ammonia at low pH, conditions that have been found to inhibit AOB activities (Lehtovirta-Morley et al., 2016a). High rates of horizontal gene transfer discovered in other *Ca. Nitrosotalea* genomes likely contributed to the lineage’s ability to be successful in extreme environments, by facilitating the procurement of physiological systems allowing for survival in harsh environments (Herbold et al., 2017). The properties associated with *Ca. Nitrosotalea* likely contributed to the relative success in propagation observed in the candidate
taxa found within this study, as the stored topsoil was derived from acidic soils (Kumaresan et al., 2017).

AOA that can persist in extreme conditions are desirable bioprospecting targets for isolation as they can provide effective soil remediation in a broader range of ecological conditions. Candidatus Nitrosocosmicus franklandus, an AOA species isolated from arable soil, has been observed to thrive in high ammonia concentrations potentially providing contributions towards the nitrification of fertilized soils due to its physiological diversity (Lehtovirta-Morley et al., 2016b). The Ca. Nitrosotalea candidates sequenced in this study comparatively, have also been shown to respond well to high ammonium input but are able to survive in denuded oligotrophic conditions commonly seen in stored topsoil. Combined with the known Ca. Nitrosotalea archaeon ability to survive in acidic environments, makes the Ca. Nitrosotalea candidates discovered in this study prime targets for isolation and utilization as indigenous microorganism for the restoration of semi-arid soil ecosystems present in WA. Novel isolation or in-situ enrichment techniques that facilitate the inoculation of Ca. Nitrosotalea can potentially speed up the re-establishment of the N-cycle upon disturbed soils.

In tandem, AOA produce less N₂O emissions compared to AOB under both high and low ammonium supply, making AOA ideal candidates for the reduction in N₂O production (Hink et al., 2018). Within terrestrial environments, excess use of nitrogen-based fertilizer has led to an increase in the production of nitrous oxide (N₂O), a major greenhouse gas, from nitrifying and denitrifying microbes (Thompson et al., 2019). There has been increased interest in investigating the physiology and niche differentiation in AOA, AOB and comammox bacteria to develop strategies that optimize microbial processes within the nitrogen cycle, whilst mitigating the effects of greenhouse gas production (Prosser et al., 2019). Microbial bioprospecting of natural soil ecosystems with low nitrogen content, such as oligotrophic soil
ecosystems could produce candidate AOA that can potentially operate under conditions with minimal nitrogen input and may further reduce N₂O production.

Conclusion

Stored topsoil is vital for restoring post land-use ecosystems and understanding the microbial communities that persist within the less-than-ideal conditions during storage is crucial for successful rehabilitation. Metagenomic approaches coupled with the enrichment of the topsoil biome has provided insight on taxa involved with critical nitrogen cycling capacity that can persist in the denuded oligotrophic conditions, as well as those that can respond rapidly to nitrogen input. Through MAGs, we were able to further explore metabolic capacity within the context of taxa allowing us to explicitly shortlist candidate taxa with diverse nitrogen metabolism. These metagenomic insights can be applied to develop strategies in bioprospecting for acclimated taxa that can perform essential ecological functions needed to revitalize degraded ecosystems. In addition, further insight into the nitrogen metabolism in nitrifying and denitrifying taxa in different ecological niches may aid in the understanding of microbial N₂O production in a bid to reduce climate footprints by finding suitable candidates for use in anthropogenic activities.

Experimental procedures

Sample collection

Soil samples were collected from topsoil storage from an active mine site 200 kilometers east of Geraldton, Western Australia (−29.164658°, 116.786696°). Soil collection protocols are detailed under the ‘Materials and Methods’ section in the study by Kumaresan et al., (2017), with details of soil chemistry composition in Supplementary Table 3.
Experimental setup

Samples were incubated under 150 mg Kg\(^{-1}\) of ammonium sulfate \((\text{NH}_4)_2\text{SO}_4\). The concentration of inorganic nitrogen was selected as it reflects the levels of nitrogen addition commonly used in the WA agricultural sector for soils as demonstrated in a prior study (Moreira-Grez, et al., 2019a). Briefly, 5 g of topsoil and 625 µL of sterilized water were added to a 120 ml serum bottle, aseptically. No other nutrient sources were added to avoid the enrichment of possible copiotroph communities that might hinder the growth of the targeted oligotrophic communities present in the study. Bottles were sealed with a rubber stopper and clamped to prevent air exchange throughout the experiment. Twelve replicates were prepared with destructive sampling over a 3-weeks period (3 replicates per timepoint). Bottles were incubated at 25°C in dark conditions. Soil from the topsoil storage facility was sampled and kept stored at -20°C until downstream analyses in order to assess baseline community composition (herein, timepoint zero).

DNA extraction and 16S rRNA gene amplicon PCR

DNA was extracted in triplicate from 0.3 g of soil samples using a Powersoil-htp 96 Soil DNA isolation kit, following manufacturer’s guidelines with minor modifications (freeze-thaw cycle (x3) after the addition of solution C1; 50 µL of solution C6 passed twice through the silica filter). Extracted DNA was quantified using a Qubit 2.0 fluorometer (Life Technologies, United States). For the 16S rRNA analysis, 2 ng of DNA was used as template for subsequent PCR amplification using the 515F/806R primer set, targeting the 16S rRNA V4 region for both bacteria and archaea domains (Liu et al., 2007). For the in-depth protocols, refer to Kumaresan et al. (2017) for PCR reagent and thermal conditions.

16S rRNA gene amplicon analysis
To identify relevant time points for shotgun metagenomic analysis, amplicon sequencing was used as a first step. PCR amplicons were sequenced using the Ion Torrent PGM platform (Thermo Fisher Scientific, Australia). Analysis was performed within QIIME wrapper (v. 1.9; Caporaso et al., 2010) using parameters described in the paper by Moreira-Grez et al. (2019b). Briefly, low quality sequences were rejected (phred score <20, size between 130 and 350). No primer mismatch and/or barcode error were allowed. Chimeric sequences were filtered using USEARCH (v. 6.1; Edgar, 2010). Quality-passed reads were then clustered using UCLUST4 at 97% sequence similarity cut-off for the identification of representative OTUs. Taxonomic assignment of representative sequences was done using RDP classifier based on the Greengenes database (v. 13.8; McDonald et al., 2012). Resulting taxonomic frequency table was used for determining if notable community differences were observed between time points for further shotgun metagenome sequencing.

After considering the microbial community composition based on the 16S rRNA gene analysis, DNA extracted from the baseline community (time zero), two- and three-weeks post enrichment replicates were selected for whole metagenome shotgun (WMS) sequencing, due to minimal differences in the relative abundance of the microbial communities between the baseline and one-week enrichment samples. Extracted DNA samples for WMS were sent to Australian Genome Research Facility (https://www.agrf.org.au/) for sequencing. Library preparation was performed using the Takara ThruPLEX protocol (version: QAM-108-003) and sequenced via the HiSeq 4000 (Illumina) sequencing platform utilizing the 150bp paired end technology.

**Functional and taxonomic profiling of soil metagenomes.**

Raw reads sequenced from the soil metagenomes were processed with Trimmmomatic (v. 0.39; Bolger et al., 2014) to remove low quality bases and Illumina adapter sequences with
the parameters ‘PE --phred33 ILLUMINACLIP:TruSeq3-PE.fa:2:30:10 LEADING:3 TRAILING:3 SLIDINGWINDOW:4:20 MINLEN:36’. Reads in which both pairs passed the trimming process were kept for further analysis in the workflow. An estimation of coverage and diversity capture between the temporal samples was performed with Nonpareil (v. 3.3.4; Rodriguez-R et al., 2018), using the alignment algorithm option via the ‘-T alignment’ flag. Nonpareil was used to estimate required sequencing efforts needed to cover the calculated diversity between the baseline soil and post-enrichment samples. The associated Nonpareil R package was used to plot the redundancy curves and report the estimated diversity index between the temporal samples.

Taxonomic profiling of microbial abundance was done at the read level using Kaiju (v. 1.7.4; Menzel et al., 2016), with default parameters using the ‘nr_euk’ database (https://kaiju.binf.ku.dk/database/kaiju_db_nr_euk_2021-02-24.tgz) obtained from the Kaiju website. Outputs were combined from samples with the same time point (baseline, two- and three weeks) for determining the taxonomic abundance at the phylum level via the ‘kaiju2table’ module using the ‘-r phylum’ flag (Supplementary Table 4). Entries under ‘cannot be assigned to a (non-viral) phylum’ and ‘unclassified’ were combined under the label ‘Unassigned/unclassified’. Relative abundance computed at the phylum level with 16S rRNA amplicon sequences (QIIME) and metagenomic reads (Kaiju) were used to assess ammonium-dependent enrichment over the three-week period. The eight most abundant phyla were selected for plotting on both datasets, although ‘Unassigned/unclassified’ bin was also included for the metagenomic reads due to its large proportion throughout the dataset. Remaining taxa (other bacteria and archaea, which within the metagenomic subset also includes eukaryotes and viruses) were classified as “other”. Stacked bar charts were computed using triplicates means and were visualized through the ggplot2 R package (v. 3.3.5; Wickham and Sievert, 2016).
For a more in-depth analysis, the ‘kaiju2table’ module was also used to explore the effect of the enrichment on the differential abundance for all replicates across all samples (3 each for baseline, two- and three-weeks post enrichment) at the genus level using the ‘-r genus’ flag (Supplementary Table 5). The genus level output from Kaiju was imported into R v. 4.1.0 and features were rarefied at 39292701 reads per sample, to allow for meaningful comparison when needed. Alpha- and beta-diversity, between the soil community taxa spanning different time points was performed on via the Shannon diversity index and non-metric multidimensional scaling analysis (NMDS) based on the Bray-Curtis dissimilarity coefficient, respectively; implemented in the Vegan R package (v. 2.5-7; Oksanen et al., 2020). Statistical difference between Alpha diversity values was tested using one-way ANOVA with Tukey as a post hoc test as implemented on the multcomp R package (v. 1.4-17; Hothorn et al., 2008). Resulting figures for the alpha- and beta-diversity plots of microbial community structures was visualized by ggplot2.

Prokaryotes known to oxidize ammonia - AOA Nitrososphaera, Nitrosocosmicus, Nitrosocaldus, Nitrosotalea, Nitrosopumilus Nitrosoarchaeum, Nitrosotenuis and Nitrosopelagicus; AOB Nitrosomonas, Nitrosospira and Nitrosococcus; and including comammon Ox Nitrospira, were plotted to determine the effects of (NH4)2SO4 enrichment on the relative abundance of this subset of taxa. Corrplot R package (v. 0.90; Wei and Simko, 2021) was used to create a correlation matrix to analyze the strength of association between the relative abundance of AOA and AOB within the baseline soil and soil three-week post enrichment. The subsequent scatter and correlation plots for representing relative abundance and the correlation matrix, respectively, between ammonia-oxidizing prokaryotes were created using ggplot2.

Functional profiling of the metagenome was performed at the read level using SUPER-FOCUS (v. 0.34; Silva et al., 2016), which groups metabolic genes into functionally similar
groups based on a reduced SEED subsystem (Overbeek et al., 2005). SUPER-FOCUS was run using default parameters, with DIAMOND (v. 0.9.14; Buchfink et al., 2015) as the aligner of choice among all individual samples. The Shannon diversity index was used to determine significant differences between metabolic genes at the Level 3 SEED subcategory comparing data from replicates from the baseline and three-week post enrichment samples through the Vegan R package with the results visualized by ggplot2. ANOVA-like differential expression analysis was used to compare metabolic gene counts between the baseline and three-week post enrichment samples via the ALDEx2 v. 1.24.0 R package using the ‘t’ option for Welch’s t and Wilcoxon rank test (Fernandes et al., 2013; Fernandes et al., 2014; Gloor et al., 2016). The conversion of metabolic count data through centered log scale transformation in ALDEx2 into a scale invariant form removed the need for prior rarefication of read counts and the transformed data underwent 128 Dirichlet Monte-Carlo Instances and 10000 iterations with Benjamini-Hochberg correction. Results from ALDEx2 were then visualized via a volcano plot using the EnhancedVolcano package (v. 1.12.0; Blighe et al., 2021). Genes with Log2 fold change lower than -2.5 and higher than 2.5 while having a significance level below alpha level ($\alpha < 0.05$) were identified as significant microbial enrichments/suppressions (Supplementary Table 1).

Genome recovery from soil metagenomes

De novo assembly of the metagenomes were performed using metaSPAdes (v. 3.15.1; Nurk et al., 2017) using default parameters and the ‘--only-assembler’ option enabled. MetaSPAdes was selected as the assembler of choice as it has been recommended for studies that aim to reconstruct representative genomes from the environment, which was one of the main focuses of this paper (Vollmers et al., 2017). Experimental replicates within each temporal sample (baseline, two weeks, and three weeks enrichment) were respectively co-assembled to improve
the rate of genomic recovery. Read mapping of the assembled contigs and MAGs were performed using Bowtie 2 (v. 2.3.4.1; Langmead and Salzberg, 2012). Post processing of read alignment files such as sorting and indexing of read maps for downstream analyses was carried out using Samtools (v. 1.10; Li et al., 2009). Binning was done using metaBAT2 (v. 2.12.1; Kang et al., 2019) using default parameters, with the quality and assembly statistics of the binned MAGs reported by CheckM (v. 1.1.2; Parks et al., 2015). Archaeal MAGs with reported completion >50% and <5% contamination was selected for further downstream analysis.

Characterization of metagenome-assembled genomes

MAGs were taxonomically classified via GTDB-tk (v. 1.5.0; Chaumeil et al., 2020) using the Genome Taxonomy Database release 06-RS202 data (Parks et al., 2020). Functional annotation of the MAGs was done using MetaErg (v. 1.2.3; Dong and Strous, 2019), with the associated database (released on Jan. 8, 2021, from: http://ebg.ucalgary.ca/metaerg/db.tar.gz). MetaErg is an annotation pipeline that utilizes various HMM and sequence search tools, as well as metabolic pathway prediction tools to automate the functional profiling of metagenomes and MAGs. Within MetaErg, output from Aragorn (v.1.2.41.c; Laslett and Canback, 2004) was used to predict tRNA and rRNAfinder (v. 1.1.0; Dong and Strous, 2019) was used for the 5S, 16S and 23S rRNA sequences for the purposes of MIMAG reporting (Bowers et al., 2017). MetaErg functional profile output results from HMMER (v. 3.1; Eddy, 2011) utilizing the Pfam-A (Finn et al., 2014) and TIGRFAM (Haft et al., 2013) databases as well as the metabolic pathway prediction by MinPath (v. 1.5; Ye and Doak, 2009) for KEGG pathways (Kanehisa and Goto, 2000) were investigated to detect proteins related to nitrogen metabolism.

Phylogenomic comparisons of MAGs and nitrogen pathways
To prepare data for genome-wide phylogenetic comparison, proteins were predicted for each individual MAG via Prodigal (v. 2.6.3; Hyatt et al., 2010). Only archaeal MAGs classified by GTDB from canonical AOA groups were selected for comparison in phylogenetic and pathway distributions. These were then processed via PhyloPhlAn (v. 3.0; Asnicar et al., 2020), using the ‘supermatrix_aa.cfg’ configuration file generated by ‘phylophlan_write_default_configs.sh’, which uses DIAMOND to map proteins to a marker database and MAFFT (v. 7.487; Katoh and Standley, 2013) for multiple sequence alignment. The database used was the PhyloPhlAn database via the ‘-d phylophlan’ option, containing 400 universal markers that are detailed in the paper by Segata et al., (2013) with the ‘--accurate’ option enabled. PhyloPhlAn was first run using default parameters to determine the minimum number of markers present in the genomes used in the study via the standard output using the ‘--verbose’ option. As one MAG returned only 99 markers, the minimum number of marker option was set using the flag ‘--min_num_markers 99’. Archaeal MAGs recovered in the study were compared to reference genomes and as the highest level of taxonomy used to distinguish genomes was at the family level, the diversity setting was set to ‘--diversity medium’. Genomes used as comparative references for AOA candidate MAGs were from canonical AOA members and obtained from NCBI’s RefSeq database (Supplementary Table 6).

To create a bootstrapped tree, the concatenated alignment outputs for bacterial and archaeal marker genes produced by the PhyloPhlAn pipeline were run separately through RAxML-HPC (v. 8.2.12; Stamatakis, 2014) with the parameters ‘-p 1989 -x 12345 -N 100 -m PROTCATLG -f a’. The bootstrapped trees produced by RAxML-HPC were visualized via iTOL (v. 6.3; Letunic and Bork, 2021). Visualization also included the presence of the nodes within selected KEGG pathways modules for nitrogen metabolism: Denitrification (M00529) and Complete nitrification (M00804) (Supplementary Table 7). In cases where the AmoCAB node was absent from Minpath predicted KEGG pathway, the functional profile created by
MetaErg utilizing HMMER and the Pfam A database was used to confirm the presence of ammonia monooxygenase proteins.

*Availability of data and materials*

All sequencing data and genomes have been deposited at the National Center for Biotechnology Information (NCBI) under the accession number PRJNA775848.

*Acknowledgements*

We gratefully acknowledge the resources provided by University of Western Australia and additional computational resources and support from the Pawsey Supercomputing Centre with funding from the Australian Government and the Government of Western Australia.

*Conflict of Interest*

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

*Author contribution*

D.K. and B.M. designed and performed the experiments. P.V. and B.M. performed the analyses. P.V. wrote the manuscript with contributions from D.K., B.M., M.J.W., A.S.W. and P.K. All authors read the manuscript and approved the content.

*Supporting information*

**Supplementary Table 1:** ALDEEx2 output of significantly a) suppressed and b) enriched metabolic genes under the SEED subsystem level 3 subcategory from the baseline and three-week post NH4 enrichment samples.
Supplementary Table 2: GTDB-tk taxonomic classification, CheckM assembly statistics, predicted tRNA and rRNA of recovered archaeal MAGs.

Supplementary Table 3: Chemical characteristics of stored topsoil.

Supplementary Table 4: Taxonomic distribution of the metagenome at the read level from Kaiju. Relative abundance of phyla/taxa across combined treatment samples at each time point.

Supplementary Table 5: Taxonomic distribution of the metagenome at the read level from Kaiju. Read counts of taxa at the genus level across individual treatment samples at each time point.

Supplementary Table 6: RefSeq ammonia-oxidizing archaea genomes used in study (Accessed 27/08/2021)

Supplementary Table 7: Nitrification and denitrification pathways predicted from a) recovered archaeal MAGs and b) RefSeq genomes, including CheckM completeness/contamination statistics.

References

Amoo, A., and Babalola, O. (2017) Ammonia-oxidizing microorganisms: Key players in the promotion of plant growth. Journal of soil science and plant nutrition 17: 935-947.

Asnicar, F., Thomas, A.M., Beghini, F., Mengoni, C., Manara, S., Manghi, P. et al. (2020) Precise phylogenetic analysis of microbial isolates and genomes from metagenomes using PhyloPhlAn 3.0. Nature Communications 11: 2500.
Banning, N.C., Maccarone, L.D., Fisk, L.M., and Murphy, D.V. (2015) Ammonia-oxidising bacteria not archaea dominate nitrification activity in semi-arid agricultural soil. *Scientific reports* 5: 11146-11146.

Blighe, K., Rana, S., and Lewis, M. (2021) EnhancedVolcano: Publication-ready volcano plots with enhanced colouring and labeling.

Bolger, A.M., Lohse, M., and Usadel, B. (2014) Trimmomatic: a flexible trimmer for Illumina sequence data. *Bioinformatics (Oxford, England)* 30: 2114-2120.

Bowers, R.M., Kyrpides, N.C., Stepanauskas, R., Harmon-Smith, M., Doud, D., Reddy, T.B.K. et al. (2017) Minimum information about a single amplified genome (MISAG) and a metagenome-assembled genome (MIMAG) of bacteria and archaea. *Nature Biotechnology* 35: 725-731.

Buchfink, B., Xie, C., and Huson, D.H. (2015) Fast and sensitive protein alignment using DIAMOND. *Nature Methods* 12: 59-60.

Caporaso, J.G., Kuczynski, J., Stombaugh, J., Bittinger, K., Bushman, F.D., Costello, E.K. et al. (2010) QIIME allows analysis of high-throughput community sequencing data. *Nature Methods* 7: 335-336.

Chaumeil, P.-A., Mussig, A.J., Hugenholtz, P., and Parks, D.H. (2020) GTDB-Tk: a toolkit to classify genomes with the Genome Taxonomy Database. *Bioinformatics* 36: 1925-1927.

Dong, X., and Strous, M. (2019) An Integrated Pipeline for Annotation and Visualization of Metagenomic Contigs. *Frontiers in Genetics* 10.

Eddy, S.R. (2011) Accelerated Profile HMM Searches. *PLOS Computational Biology* 7: e1002195.
Edgar, R.C. (2010) Search and clustering orders of magnitude faster than BLAST. *Bioinformatics* **26**: 2460-2461.

Ezeokoli, O.T., Mashigo, S.K., Paterson, D.G., Bezuidenhout, C.C., and Adeleke, R.A. (2019) Microbial community structure and relationship with physicochemical properties of soil stockpiles in selected South African opencast coal mines. *Soil science and plant nutrition (Tokyo)* **65**: 332-341.

Fernandes, A.D., Macklaim, J.M., Linn, T.G., Reid, G., and Gloor, G.B. (2013) ANOVA-Like Differential Expression (ALDE) Analysis for Mixed Population RNA-Seq. *PLOS ONE* **8**: e67019.

Fernandes, A.D., Reid, J.N.S., Macklaim, J.M., McMurrough, T.A., Edgell, D.R., and Gloor, G.B. (2014) Unifying the analysis of high-throughput sequencing datasets: Characterizing RNA-seq, 16S rRNA gene sequencing and selective growth experiments by compositional data analysis. *Microbiome* **2**: 15-15.

Finn, R.D., Bateman, A., Clements, J., Coggill, P., Eberhardt, R.Y., Eddy, S.R. et al. (2014) Pfam: the protein families database. *Nucleic Acids Research* **42**: D222-D230.

Fisk, L.M., Maccarone, L.D., Barton, L., and Murphy, D.V. (2015) Nitrapyrin decreased nitrification of nitrogen released from soil organic matter but not amoA gene abundance at high soil temperature. *Soil Biology and Biochemistry* **88**: 214-223.

Gloor, G.B., Macklaim, J.M., and Fernandes, A.D. (2016) Displaying Variation in Large Datasets: Plotting a Visual Summary of Effect Sizes. *Journal of computational and graphical statistics* **25**: 971-979.
Golos, P.J., and Dixon, K.W. (2014) Waterproofing Topsoil Stockpiles Minimizes Viability Decline in the Soil Seed Bank in an Arid Environment. *Restoration ecology* **22**: 495-501.

Guy, L., and Ettema, T.J. (2011) The archaeal 'TACK' superphylum and the origin of eukaryotes. *Trends Microbiol* **19**: 580-587.

Haff, D.H., Selengut, J.D., Richter, R.A., Harkins, D., Basu, M.K., and Beck, E. (2013) TIGRFAMs and Genome Properties in 2013. *Nucleic Acids Research* **41**: D387-D395.

Hanshew, A.S., Mason, C.J., Raffa, K.F., and Currie, C.R. (2013) Minimization of chloroplast contamination in 16S rRNA gene pyrosequencing of insect herbivore bacterial communities. *Journal of microbiological methods* **95**: 149-155.

Hart, M.M., Cross, A.T., D’Agui, H.M., Dixon, K.W., Van der Heyde, M., Mickan, B. et al. (2020) Examining assumptions of soil microbial ecology in the monitoring of ecological restoration. *Ecological Solutions and Evidence* **1**: e12031.

Herbold, C.W., Lehtovirta-Morley, L.E., Jung, M.Y., Jehmlich, N., Hausmann, B., Han, P. et al. (2017) Ammonia-oxidising archaea living at low pH: Insights from comparative genomics. *Environmental microbiology* **19**: 4939-4952.

Hink, L., Gubry-Rangin, C., Nicol, G.W., and Prosser, J.I. (2018) The consequences of niche and physiological differentiation of archaean and bacterial ammonia oxidisers for nitrous oxide emissions. *The ISME Journal* **12**: 1084-1093.

Hothorn, T., Bretz, F., and Westfall, P. (2008) Simultaneous inference in general parametric models. *Biom J* **50**: 346-363
Hyatt, D., Chen, G.-L., LoCascio, P.F., Land, M.L., Larimer, F.W., and Hauser, L.J. (2010) Prodigal: prokaryotic gene recognition and translation initiation site identification. *BMC Bioinformatics* **11**: 119.

Jenkins, S.N., Murphy, D.V., Waite, I.S., Rushton, S.P., and O’Donnell, A.G. (2016) Ancient landscapes and the relationship with microbial nitrification. *Scientific Reports* **6**: 30733.

Jung, J., Kim, J.-S., Taffner, J., Berg, G., and Ryu, C.-M. (2020) Archaea, tiny helpers of land plants. *Computational and Structural Biotechnology Journal* **18**: 2494-2500.

Kanehisa, M., and Goto, S. (2000) KEGG: Kyoto Encyclopedia of Genes and Genomes. *Nucleic Acids Research* **28**: 27-30.

Kang, D.D., Li, F., Kirton, E., Thomas, A., Egan, R., An, H., and Wang, Z. (2019) MetaBAT 2: an adaptive binning algorithm for robust and efficient genome reconstruction from metagenome assemblies. *PeerJ* **7**: e7359-e7359.

Katoh, K., and Standley, D.M. (2013) MAFFT Multiple Sequence Alignment Software Version 7: Improvements in Performance and Usability. *Molecular Biology and Evolution* **30**: 772-780.

Kumaresan, D., Cross, A.T., Moreira-Grez, B., Kariman, K., Nevill, P., Stevens, J. et al. (2017) Microbial Functional Capacity Is Preserved Within Engineered Soil Formulations Used In Mine Site Restoration. *Scientific Reports* **7**: 564.

Kuypers, M.M.M., Marchant, H.K., and Kartal, B. (2018) The microbial nitrogen-cycling network. *Nature Reviews Microbiology* **16**: 263-276.

Langmead, B., and Salzberg, S.L. (2012) Fast gapped-read alignment with Bowtie 2. *Nature methods* **9**: 357-359.
Laslett, D., and Canback, B. (2004) ARAGORN, a program to detect tRNA genes and tmRNA genes in nucleotide sequences. *Nucleic Acids Research* **32**: 11-16.

Lehtovirta-Morley, L.E. (2018) Ammonia oxidation: Ecology, physiology, biochemistry and why they must all come together. *FEMS Microbiology Letters* **365**.

Lehtovirta-Morley, L.E., Sayavea-Soto, L., Gallois, N., Schouten, S., Stein, L.Y., Prosser, J.I., and Nicol, G.W. (2016a) Identifying potential mechanisms enabling acidophilia in the ammonia-oxidising archaeon 'Candidatus Nitrosotalea devanaterra'. *Applied and environmental microbiology* **82**: 2608-2619.

Lehtovirta-Morley, L.E., Ross, J., Hink, L., Weber, E.B., Gubry-Rangin, C., Thion, C. et al. (2016b) Isolation of 'Candidatus Nitrosocosmicus franklandus', a novel ureolytic soil archaeal ammonia oxidiser with tolerance to high ammonia concentration. *FEMS microbiology ecology* **92**: 1-10.

Letunic, I., and Bork, P. (2021) Interactive Tree Of Life (iTOL) v5: an online tool for phylogenetic tree display and annotation. *Nucleic Acids Research* **49**: W293-W296.

Li, H., Handsaker, B., Wysoker, A., Fennell, T., Ruan, J., Homer, N. et al. (2009) The Sequence Alignment/Map format and SAMtools. *Bioinformatics (Oxford, England)* **25**: 2078-2079.

Liang, D., Ouyang, Y., Tiemann, L., and Robertson, G.P. (2020) Niche Differentiation of Bacterial Versus Archaeal Soil Nitrifiers Induced by Ammonium Inhibition Along a Management Gradient. *Frontiers in Microbiology* **11**.
Liu, Z., Lozupone, C., Hamady, M., Bushman, F.D., and Knight, R. (2007) Short pyrosequencing reads suffice for accurate microbial community analysis. *Nucleic Acids Research* **35**: e120-e120.

Lycus, P., Bøthun, K.L., Bergaust, L., Shapleigh, J.P., Bakken, L.R., and Frostegård, Å. (2017) Phenotypic and genotypic richness of denitrifiers revealed by a novel isolation strategy. *The ISME Journal* **11**: 2219-2232.

McDonald, D., Price, M.N., Goodrich, J., Nawrocki, E.P., DeSantis, T.Z., Probst, A. et al. (2012) An improved Greengenes taxonomy with explicit ranks for ecological and evolutionary analyses of bacteria and archaea. *The ISME Journal* **6**: 610-618.

Menzel, P., Ng, K.L., and Krogh, A. (2016) Fast and sensitive taxonomic classification for metagenomics with Kaiju. *Nature Communications* **7**: 11257.

Moreau, D., Bardgett, R.D., Finlay, R.D., Jones, D.L., and Philippot, L. (2019) A plant perspective on nitrogen cycling in the rhizosphere. *Functional Ecology* **33**: 540-552.

Moreira-Grez, B., Muñoz-Rojas, M., Kariman, K., Storer, P., O’Donnell, A.G., Kumaresan, D., and Whiteley, A.S. (2019a) Reconditioning Degraded Mine Site Soils With Exogenous Soil Microbes: Plant Fitness and Soil Microbiome Outcomes. *Frontiers in Microbiology* **10**.

Moreira-Grez, B., Tam, K., Cross, A.T., Yong, J.W.H., Kumaresan, D., Nevill, P. et al. (2019b) The Bacterial Microbiome Associated With Arid Biocrusts and the Biogeochemical Influence of Biocrusts Upon the Underlying Soil. *Frontiers in Microbiology* **10**.

Nurk, S., Meleshko, D., Korobeynikov, A., and Pevzner, P.A. (2017) metaSPAdes: a new versatile metagenomic assembler. *Genome research* **27**: 824-834.
O'Sullivan, C.A., Wakelin, S.A., Fillery, I.R.P., and Roper, M.M. (2013) Factors affecting ammonia-oxidising microorganisms and potential nitrification rates in southern Australian agricultural soils. *Soil research (Collingwood, Vic)* **51**: 240-252.

O’Brien, F.J.M., Almaraz, M., Foster, M.A., Hill, A.F., Huber, D.P., King, E.K. et al. (2019) Soil Salinity and pH Drive Soil Bacterial Community Composition and Diversity Along a Lateritic Slope in the Avon River Critical Zone Observatory, Western Australia. *Frontiers in Microbiology* **10**.

Oksanen, J., Blanchet, F.G., Friendly, M., Kindt, R., Legendre, P., McGlinn, D. et al. (2020) vegan: Community Ecology Package.

Overbeek, R., Begley, T., Butler, R.M., Choudhuri, J.V., Chuang, H.-Y., Cohoon, M. et al. (2005) The subsystems approach to genome annotation and its use in the project to annotate 1000 genomes. *Nucleic acids research* **33**: 5691-5702.

Parks, D.H., Imelfort, M., Skennerton, C.T., Hugenholtz, P., and Tyson, G.W. (2015) CheckM: assessing the quality of microbial genomes recovered from isolates, single cells, and metagenomes. *Genome research* **25**: 1043-1055.

Parks, D.H., Chuvchenko, M., Chauveil, P.A., Rinke, C., Mussig, A.J., and Hugenholtz, P. (2020) A complete domain-to-species taxonomy for Bacteria and Archaea. *Nat Biotechnol* **38**: 1079-1086.

Pester, M., Schleper, C., and Wagner, M. (2011) The Thaumarchaeota: an emerging view of their phylogeny and ecophysiology. *Current opinion in microbiology* **14**: 300-306.
Prosser, J.I., Hink, L., Gubry-Rangin, C., and Nicol, G.W. (2020) Nitrous oxide production by ammonia oxidizers: Physiological diversity, niche differentiation and potential mitigation strategies. *Global Change Biology* **26**: 103-118.

Rodriguez-R, L.M., Gunturu, S., Tiedje, J.M., Cole, J.R., Konstantinidis, K.T., and Fodor, A. (2018) Nonpareil 3: Fast Estimation of Metagenomic Coverage and Sequence Diversity. *mSystems* **3**: e00039-00018.

Segata, N., Börnigen, D., Morgan, X.C., and Huttenhower, C. (2013) PhyloPhlAn is a new method for improved phylogenetic and taxonomic placement of microbes. *Nature Communications* **4**: 2304.

Silva, G.G.Z., Green, K.T., Dutilh, B.E., and Edwards, R.A. (2016) SUPER-FOCUS: a tool for agile functional analysis of shotgun metagenomic data. *Bioinformatics (Oxford, England)* **32**: 354-361.

Stamatakis, A. (2014) RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. *Bioinformatics* **30**: 1312-1313.

Thompson, R.L., Lassaletta, L., Patra, P.K., Wilson, C., Wells, K.C., Gressent, A. et al. (2019) Acceleration of global N2O emissions seen from two decades of atmospheric inversion. *Nature Climate Change* **9**: 993-998.

Trivedi, C., Reich, P.B., Maestre, F.T., Hu, H.-W., Singh, B.K., and Delgado-Baquerizo, M. (2019) Plant-driven niche differentiation of ammonia-oxidizing bacteria and archaea in global drylands. *The ISME Journal* **13**: 2727-2736.
Vollmers, J., Wiegand, S., and Kaster, A.-K. (2017) Comparing and Evaluating Metagenome Assembly Tools from a Microbiologist’s Perspective - Not Only Size Matters! *PLOS ONE* **12**: e0169662.

Wagg, C., Bender, S.F., Widmer, F., and van der Heijden, M.G.A. (2014) Soil biodiversity and soil community composition determine ecosystem multifunctionality. *Proceedings of the National Academy of Sciences* **111**: 5266.

Wei, T., and Simko, V. (2021) R package 'corrplot': Visualization of a Correlation Matrix.

Wickham, H., and Sievert, C. (2016) *ggplot2: Elegant Graphics for Data Analysis*. Cham: Springer International Publishing AG, z.Hd. Alexander Grossmann.

Williams, W., Chilton, A., Schneemilch, M., Williams, S., Neilan, B., and Driscoll, C. (2019) Microbial biobanking - cyanobacteria-rich topsoil facilitates mine rehabilitation. *Biogeosciences* **16**: 2189-2204.

Wubs, E.R.J., van der Putten, W.H., Bosch, M., and Bezemer, T.M. (2016) Soil inoculation steers restoration of terrestrial ecosystems. *Nature Plants* **2**: 16107.

Ye, Y., and Doak, T.G. (2009) A Parsimony Approach to Biological Pathway Reconstruction/Inference for Genomes and Metagenomes. *PLOS Computational Biology* **5**: e1000465.