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Abstract: Archaeal communities in arable soils are dominated by Nitrososphaeria, a class within Thaumarchaeota comprising all known ammonia-oxidizing archaea (AOA). AOA are key players in the nitrogen cycle and defining their niche specialization can help predicting effects of environmental change on these communities. However, hierarchical effects of environmental filters on AOA and the delineation of niche preferences of nitrososphaerial lineages remain poorly understood. We used phylogenetic information at fine scale and machine learning approaches to identify climatic, edaphic and geomorphological drivers of Nitrososphaeria and other archaea along a 3000 km European gradient. Only limited insights into the ecology of the low-abundant archaeal classes could be inferred, but our analyses underlined the multifactorial nature of niche differentiation within Nitrososphaeria. Mean annual temperature, C:N ratio and pH were the best predictors of their diversity, evenness and distribution. Thresholds in the predictions could be defined for C:N ratio and cation exchange capacity. Furthermore, multiple, independent and recent specializations to soil pH were detected in the Nitrososphaeria phylogeny. The coexistence of widespread ecophysiological differences between closely related soil Nitrososphaeria highlights that their ecology is best studied at fine phylogenetic scale.

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Diversity of archaea and niche preferences among putative ammonia-oxidizing Nitrososphaeria dominating across European arable soils

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Summary

Archaeal communities in arable soils are dominated by Nitrososphaeria, a class within Thaumarchaeota comprising all known ammonia-oxidizing archaea (AOA). AOA are key players in the nitrogen cycle and defining their niche specialization can help predicting effects of environmental change on these communities. However, hierarchical effects of environmental filters on AOA and the delineation of niche preferences of nitrososphaerial lineages remain poorly understood. We used phylogenetic information at fine scale and machine learning approaches to identify climatic, edaphic and geomorphological drivers of Nitrososphaeria and other archaea along a 3000 km European gradient. Only limited insights into the ecology of the low-abundant archaeal classes could be inferred, but our analyses underlined the multifactorial nature of niche differentiation within Nitrososphaeria. Mean annual temperature, C:N ratio and pH were the best predictors of their diversity, evenness and distribution. Thresholds in the predictions could be defined for C:N ratio and cation exchange capacity. Furthermore, multiple, independent and recent specializations to soil pH were detected in the Nitrososphaeria phylogeny. The coexistence of widespread ecophysiological differences between closely related soil Nitrososphaeria highlights that their ecology is best studied at fine phylogenetic scale.

Introduction

Archaea are pivotal for the functioning of all major biomes, as they play a critical role in both carbon (C) and nitrogen (N) cycles (Falkowski et al., 2008; Offre et al., 2013). In terrestrial ecosystems, archaeal communities tend to be phylogenetically clustered and are commonly dominated by Thaumarchaeota (Auguet et al., 2010; Bates et al., 2011; Tripathi et al., 2015). This phylum harbours the globally important ammonia-oxidizing archaea (AOA), restricted to the class Nitrososphaeria (Alves et al., 2018). All AOA characterized so far use the
ammonia monooxygenase enzyme (encoded by amoABC genes) to catalyze the first step of nitrification, the oxidation of ammonia to hydroxylamine (Vajrala et al., 2013). Globally, nitrification contributes to the circulation of N (Kuypers et al., 2018), but locally this process causes N losses, directly through nitrate leaching and production of the greenhouse gas nitrous oxide and indirectly by fuelling denitrification leading to gaseous N losses as dinitrogen gas or nitrous oxide. Altogether, this corresponds to an average loss of 50% of the N added to arable soils (Lassaletta et al., 2014). Thus, nitrification affects N use efficiency in cropping systems, causes eutrophication of watersheds and contributes to global warming. In arable soils, AOA are typically abundant and important contributors to nitri-

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of 97%, respectively. Most of the diversity was found within the order Nitrososphaerales (322 ASVs or approximately 91% of putative AO ASVs and approximately 98% of the reads), whereas Nitrosopumilales and Nitrosotaleales represented a minor fraction of the nitrososphaerial communities (approximately 1% and 8% of ASVs; < 1% and < 2% of the reads, respectively). Thermoplasma and Woesearchaeia were diverse, respectively representing approximately 27% and 25% of the ASVs, but also relatively rare (< 3% of the reads).

The combination of PCA and PERMANOVA showed that the overall archaeal communities were structured following a spatial gradient of pH and temperature along PC1, with a shift from alkaline pH and warm temperatures to acidic soils and colder climate (Fig. 2A). This is reflected by the increasing dissimilarity between communities with increasing spatial distance between sampling sites (Fig. 2B). Besides mean annual temperature (MAT) and pH ($R^2 = 0.11$ and 0.10, respectively), soil C:N ratio ($R^2 = 0.07$) and soil moisture ($R^2 = 0.06$) were also important contributors to $\beta$-diversity ($p < 0.001$; Table S1). Climatic, edaphic and spatial factors collectively explained 36% of the variation in the archaeal community composition and all three groups of factors were significant ($p < 0.01$; Fig. 2C). When partitioning the variation, spatial distance (i.e. dispersal limitation) explained nearly as much variation in community composition as the edaphic factors (i.e. environmental filtering; approximately 10%), whereas the climatic component defined solely by MAT explained only 1%.

**Environmental predictors of diversity of the individual archaeal classes and abundance of AOA**

Random forest-based variable selection analyses revealed that different environmental variables (Table 1) contributed to the PD and evenness of the taxa-specific archaeal communities, although MAT was important for diversity of nearly all taxonomic groups (Figs. 3A and B, S1 and S2). Across the archaeal domain, edaphic factors were more important for PD than for evenness. For the methanogens, PD and evenness of Methanobacteria and Methanomicrobia displayed the same relationship to calcium, elevation, soil organic carbon (SOC) and, to a lesser degree, MAT. All three categories of environmental factors significantly influenced the PD of Woesearchaeia, while soil texture and MAT were major predictors of the diversity and evenness of Thermoplasma.

For the Nitrososphaeria, elevation, cation exchange capacity (CEC), C:N ratio and moisture had strong effects on the diversity, whereas evenness was driven by multiple climatic, edaphic and geomorphological (i.e. elevation) variables. Here, diversity and pH exhibited a u-shaped relationship, with an increase associated with pH below 7 and above 7.5. More acidic soils were associated with an increase in both PD and evenness in Group 1.1c, i.e. a defined lineage within the Thaumarchaeota that likely does not oxidize ammonia (Weber et al., 2015). The abundance of AOA, measured as the copy number of the archaeal amoA gene (Fig. S3A), was positively influenced by elevation, total N and total P, but tended to decrease with increasing silt content and bulk density (Figs. 3C and S3B).

**Ecological preferences of nitrososphaerial taxa**

Spatial distance explained a larger proportion of the variation in the structure of the nitrososphaerial communities than edaphic factors (14.6% and 11.2%, respectively), whereas MAT explained 1.4% (Fig. S4). The abundance of archaeal amoA genes, being a biotic factor predicting functionality, explained only 0.8% to the variation.

The sequential effect of several environmental filters was determined using multivariate regression trees
(MRT), which delineate clusters of samples where the variation in environmental conditions is minimized. Each of the eight clusters (labelled A–H) contained 10–31 soil samples (Fig. 4), with the exception of the four samples from northern Sweden that formed a separate cluster (cluster A in Fig. 4). Eight variables, among edaphic (bulk density, calcium, CEC, C:N ratio, pH, SOC and total C) and climatic (MAT) factors, were selected in the regression analyses, with MAT also reflecting spatial distance. The MRT identified two to four levels of environmental filtering, with MAT being the most important driver. The importance of geographic distance was evidenced by the origin of the samples present in each cluster, with a clear North–South gradient. Several variables contributed equally to the split between clusters A and B (CEC, C:N ratio, MAT, SOC and total C) and clusters G and H (calcium and SOC), indicating that soil C content also plays a role in defining ecological preferences. It should be noted that this eight-cluster partition does not imply within-cluster homogeneity for the rest of the measured variables (Table S2).

Balances, which depict relative changes in ASV abundance between two neighbouring clades relative to each other, were calculated for all inner nodes of the phylogenetic tree in each MRT cluster. They revealed extensive niche differentiation throughout the phylogeny (Fig. 4). Within the less abundant orders Nitrosopumilales and Nitrosotaleales, balances at both deep (depicting ancient evolutionary events conserved in the phylogeny) and shallow (depicting more recent adaptations) nodes in the

Fig. 2. Factors driving the variation in archaeal community composition and structure across the European gradient.
A. Principal component analysis (PCA) showing differences in archaeal communities between all samples and the associated changes in pH and MAT, identified as the two best explanatory variables (PERMANOVA, \( p < 0.001 \)).
B. Distance-decay relationship between geographic distance and community similarity. The red line indicates the ordinary least squares linear regression (\( p < 0.001 \)).
C. Variation partitioning analysis (VPA) between climatic, edaphic and spatial components. All fractions were significant (\( p < 0.01 \)) and the variation explained is indicated (%).
Both PCA and VPA were performed on the philr-transformed data. [Color figure can be viewed at wileyonlinelibrary.com]
Table 1. Environmental variables used in this study.

| Category          | Variable         | Range        |
|-------------------|------------------|--------------|
| Climatic          | Mean annual temperature (MAT; °C) | 3.7–19.5     |
| Geomorphological  | Elevation (m)    | 7.8–1022.0   |
| Edaphic           | Bulk density (g cm⁻³) | 0.7–1.7     |
|                   | C:N ratio        | 7.8–63.9     |
|                   | C:P ratio        | 7.3–1042.9   |
|                   | Calcium (cmol kg⁻¹) | 0.2–36.1    |
|                   | Cation exchange capacity (CEC; cmol kg⁻¹) | 5.6–49.6 |
|                   | Clay (%)          | 8.6–54.9     |
|                   | Magnesium (cmol kg⁻¹) | 0.3–5.6   |
|                   | Moisture (%)      | 0.03–0.35    |
|                   | N:P ratio         | 0.1–19.1     |
|                   | pH               | 5.4–8.3      |
|                   | Silt (%)          | 12.4–63.1    |
|                   | SOC (g kg⁻¹)      | 3.2–36.1     |
|                   | Total C (g kg⁻¹)  | 6.6–90.2     |
|                   | Total N (g kg⁻¹)  | 0.5–4.8      |
|                   | Total P (g kg⁻¹)  | 0.1–1.7      |

The congruence between the phylogeny of Nitrososphaeria and their ecological preferences was relatively limited, suggesting the existence of a considerable ecotypic intra-diversity within this class (Alves et al., 2018) due to high diversification rates (Gubry-Rangin et al., 2015). These observations corroborate work conducted on marine systems, where genomic and physiological data indicate that closely related AOA isolates could differ in a range of traits, including motility, pH preferences and substrate utilization (Qin et al., 2014; Bayer et al., 2016). Extensive niche specialization due to differences in affinity for ammonia (Lehtovirta-Morley et al., 2016; Hink et al., 2017; Jung et al., 2021) and for different organic C compounds in mixotrophic/heterotrophic growth have been discussed (Prosser and Nicol, 2012). However, organic acids have been shown to be used to detoxify the hydrogen peroxide produced during ammonia oxidation in several AOA isolates rather than used as an energy source (Kim et al., 2016) and the question of whether there are AOA able to grow heterotrophically remains open. Observed differences in the balances between lineages at fine phylogenetic scale also reflect a low level of intra-genomic heterogeneity in the 16S rRNA gene (Sun et al., 2013), and highlight the relevance of using ASVs to study the environmental determinants of soil Nitrososphaeria (i.e. more than 10 times as many nitrososphaerial ASVs than OTUs were detected).

By using MRT, we could evaluate the hierarchical effects of environmental filters in the delineation of ecological preferences of nitrososphaerical lineages, whereas the random forest modelling allowed us to go beyond linear correlations when exploring drivers of archaeal diversity. Across our broad geographical and environmental gradients, MAT was the most prominent variable in the MRT analysis, as indicated by the major grouping between clusters A–B, C–E and F–H, and ranked among the best predictors of evenness. Temperature affects composition and nitrification activity of AOA communities (Tourna et al., 2008; Alves et al., 2013; Gubry-Rangin et al., 2017), but since MAT and latitude correlated (Spearman’s r = 0.9, p < 0.001), MAT likely also reflects the importance of spatial distance. Results of the variation partitioning analysis, where spatial distance explained approximately 15% of the variation in the structure of the nitrososphaerical communities and MAT only approximately 1%, further support this interpretation. Soil pH was another significant predictor of phylogenetic diversity and evenness of Nitrososphaeria across the European gradient. This agrees with the dominant idea that soil pH drives diversification of terrestrial Thaumarchaeota at broad phylogenetic scales (Nicol et al., 2008; Gubry-Rangin et al., 2011, 2015; Oton et al., 2016). However, we detected multiple balance shifts in the shallow nodes in the phylogenies corresponding to low-pH MRT clusters (i.e. A, B and E), which suggest that closely related nitrososphaerial taxa display distinct preferences with regards to soil pH. This would imply multiple independent and recent specializations to acidic pH in Nitrososphaeria, which aligns with the conclusions of a study based on analyses of the amoA gene (Alves et al., 2018). The limited genomic information available on Nitrososphaericales prevents us from using the congruence between 16S rRNA and amoA phylogenies (Oton et al., 2016; Alves et al., 2018) to match the clades observed in this study for identifying amoA-based low-pH AOA lineages (within clades NS-α, -β, -γ and -ζ in Alves et al., 2018). Nevertheless, since pH controls the equilibrium between ammonia and ammonium in soils, these adaptations could reflect differences in substrate affinity (Lehtovirta-Morley et al., 2016; Hink et al., 2017), possibly through distinct molecular adaptations of the ammonia monooxygenase (Macqueen and Gubry-Rangin, 2016). Different N-related variables...
such as CEC and C:N ratio were important predictors of the diversity and/or evenness of Nitrososphaeria; and for the abundance of archaeal amoA genes, it was total N. MAT (or distance) and elevation were thus the only variables linking $\alpha$-diversity and predicted function (archaeal amoA genes). The abundance of archaeal...
amoA genes was also a poor predictor of nitrososphaerical β-diversity, suggesting a weak link between diversity of this class and the genetic potential for ammonia oxidation at broad geographic scale. Our results also illustrate the expected selective role played by soil N, and are consistent with demonstrated effects of soil C:N ratio (Bates et al., 2011; Jiang et al., 2014) and ammonium supply (Verhamme et al., 2011; Hink et al., 2018) on the abundance and structure of soil AOA communities. Random forest modelling revealed that nearly all of the above-mentioned relationships were nonlinear. Furthermore, some accumulated local effect curves, showing how the predictions change on average over the range of each individual environmental variable, suggested the existence of thresholds (TH). The increase of nitrososphaerical PD in the low range of the C:N ratio (TH ~ 10) could be due to preferences for mineralized N from organic matter (Levínčik-Höfferle et al., 2012). This aligns with the observed sharp increase of archaeaial nitritification rates at C:N ratios below 20 (Lu et al., 2015), indicating a coupling between nitrososphaerical diversity levels and nitritification activity. However, the high number of significant shallow nodes in the phylogeny of both high and low C:N ratio clusters in Southern Europe shows that niche specialization in Nitrososphaeria extends past preferences in soil C:N. For example, total N was a better predictor for AOA abundances than C:N, showing a strong positive effect across the measured range, although it appeared weaker between 2 and 3.5 g N kg⁻¹ soil. The soils’ CEC was also important, likely by influencing the retention of ammonium. We identified a threshold above which CEC had a strong negative effect on nitrososphaerical PD (TH ~ 15 cmol kg⁻¹). Soil CEC is linked to soil texture and clay content has previously been reported to negatively affect the abundance of AOA at the local scale (Wessén et al., 2011). We did not observe any effect of neither clay content or CEC on AOA abundance, which suggests that soil texture and its implications on CEC has a stronger effect on diversity than on abundance of AOA at the continental scale.

For the overall soil archaeal communities, MAT was an important factor and the main driver of β-diversity. As for the Nitrososphaeria, the importance of MAT also reflects the effect of distance on the changes in community composition, as shown by the higher explanatory power of spatial distance compared to MAT in the variation partitioning analysis and the high fitness value of the distance-decay relationship ($R^2_{adj} = 0.22$) along the geographical gradient. This latter observation contrasts with a recent survey where substantially weaker decays of community similarity were found in archaeal communities in maize and rice fields across Eastern China (Jiao et al., 2019b). At large spatial scales, distance-decay relationships are typically influenced by dispersal limitation (stochastic process) and species sorting (deterministic process), i.e. the combined effect of environmental filtering and biotic interactions (Hanson et al., 2012). In the present study, dispersal limitation and environmental filtering had comparable effects on the overall β-diversity of archaea, whereas the few other studies of archaeal communities across broad spatial scales in arable soils have reported larger sorting: dispersal effect ratios (Liu et al., 2019; Jiao et al., 2019b). The relative importance of dispersal limitation in the present study could have been overestimated, since some of the variation explained by spatial factors alone likely encompass unmeasured environmental variables and we could have missed environmental variables that are relevant for archaea. Nevertheless, the use of ASVs instead of OTUs and the removal of the rare taxa combined with broader environmental and geographic gradients, as we did, should lead to a more accurate assessment of the relative importance of environmental filtering versus dispersal limitation for the assembly of archaeal communities when compared to other studies (Liu et al., 2019; Jiao et al., 2019b). Our results thus indicate that the structure of archaeal communities associated to fields under cereal cultivation could be less influenced by changes in environmental conditions than previously suggested; an important finding in the context of ongoing climate change. Moreover, balanced effects of stochastic versus deterministic processes were recently found to promote diverse, yet uneven ecosystem functions in a study comparing three types of agroecosystems (Liu et al., 2021). This interpretation of effects of balanced stochastic and deterministic processes fits with the observed low evenness of the archaeal community and the presence of a high diversity of low abundant groups harbouring potentially diverse functional capabilities at the continental scale.

Fig. 3. Relationship between environmental variables and β-diversity indices or the abundance of AOA, based on random forest (RF) analyses. Variables selected by VSURF (x-axis; see Table 1 for units and range) were used to generate accumulated local effects plots, which show how the prediction of the response variables (phylogenetic diversity, evenness or abundance of AOA) changes along the range of each environmental variable in each of the RF models (γ-axis; range indicated in brackets). Model parameters and fit are indicated in Table S3. Full plots are available as Figs. S1, S2 and S3B. A. Phylogenetic diversity of archaeal communities. B. Evenness of archaeal communities. C. Abundance of AOA, measured as the copy number of the archaeal amoA gene (g⁻¹ dw soil). [Color figure can be viewed at wileyonlinelibrary.com]
Fig. 4. Legend on next page.
Archaenal classes present at low abundance exhibited different responses to the environmental factors evaluated. Soil pH displayed a negative relationship with both PD and evenness in Group 1.1c thaumarchaeota. Accordingly, these microorganisms have mainly been detected in acidic environments (Lehtovirta et al., 2009; Tripathi et al., 2013). The few known representatives of this group do not have amoA homologues and do not produce nitrite or nitrate in culture. It is therefore hypothesized that Group 1.1c is a non-ammonia oxidizing lineage within the Thaumarchaeota (Lin et al., 2015; Weber et al., 2015), which is supported by the lack of significant effects of N-related variables observed in this study. Woesearchaeia and the two classes of methanogens shared only a few environmental preferences (i.e. elevation and clay content) with regards to α-diversity, despite recent findings that they that are potential metabolic partners (Liu et al., 2018). This partnership or their importance in arable soils thus remain elusive. Finally, diversity and evenness of the Thermoplasmata, which was the second most abundant class in terms of both ASVs and relative abundance, appeared to be predominantly driven by MAT and soil texture along the gradient. Their ecology in agricultural soils is largely unknown as Thermoplasmata have mainly been studied in acid mine drainage (Baker and Banfield, 2003) and hot environments (Massello et al., 2020). Members of this class however account for approximately 5% of archaeal sequences in global soil samples (Auguet et al., 2010) and have been detected at high abundances in deeper soil layers in both boreal (Kemnitz et al., 2007) and temperate, acidic deciduous forests (Isoda et al., 2017). There are few cultivated representatives, but more than 400 genomes available that suggest a versatile metabolic potential for this class, including methanogenesis, sulfur (S) cycling and even dinitrogen fixation (Baker et al., 2020). Sulfur cycling was also detected in a genome obtained from peat soil (Lin et al., 2015), while methanogenic Thermoplasmata appear to be widespread in wetlands (Söllinger et al., 2016). Several low-abundant archaenal lineages could thus be involved in C and S cycling and play a more important role than previously thought in arable soils.

This study provides novel insights into the ecology of archaea, particularly putative ammonia-oxidizing Nitrososphaeria, in arable soils across Europe. Our results suggest extensive ecophysiological differences between closely related Nitrososphaeria and underline the multifactorial nature of niche differentiation in this class. Both MAT (i.e. distance) and soil C:N ratio were unexpectedly better predictors of their diversity and distribution than soil pH, for which multiple, independent adaptations were inferred. Thresholds for soil C:N and CEC were also identified. Overall, we show that future studies aiming at deciphering the ecology of Nitrososphaeria should be performed at fine phylogenetic scale, using methods accounting for non-linear relationships between environmental drivers and the diversity of these functionally important archaea in arable soils.

**Experimental procedures**

**Sampling and measurement of edaphic parameters**

Soil samples were collected across a north–south gradient in Europe (Sweden, Germany, Switzerland, France and Spain) in a total of 151 agricultural fields (Garland et al., 2021) (Fig. S5). To homogenize variation in plant development stages and associated farming practices, the sampling was performed around flowering time (i.e. anthesis) between May and August 2017 depending on country and location. Only fields under cereal cultivation (barley, oat or wheat) and conventional tillage were surveyed. In each field, eight soil cores (Ø5 × 20 cm) were taken within a 10 m radius. Five cores were sieved (2 mm) and homogenized into a composite sample, which was air-dried before measuring soil parameters (Table 1) using the Swiss standard protocols (Fal et al., 1996). A fresh subsample of the composite soil was taken before drying for DNA extraction and stored at −20°C until DNA extraction (one extraction per sample). The three remaining cores were kept intact and used to measure bulk density of the fine soil (< 8 mm). Mean annual atmospheric temperature (MAT) data (1987–2017) were obtained for each sampling site using their GPS coordinates and the closest weather station in the NOAA database (https://www.noaa.gov/), through the R
package ‘moaa’ (v. 0.8.4; Chamberlain, 2019). The same GPS coordinates were also used to gather elevation data using the R package ‘elevatr’ (v. 0.2.0; Hollister and Shah, 2017).

DNA extraction, amplification and sequencing

DNA was extracted on the homogenized soil samples using the DNeasy PowerSoil-htp 96 well DNA isolation kit (Qiagen, Hilden, Germany), according to the manufacturer’s instructions. Archaeal 16S rRNA gene fragments, encompassing the V3–V4 hypervariable regions, were amplified using the primer pair S-D-Arch-0349-a-S-17 (Takai and Horikoshi, 2000) and S-D-Bact-0785-a-A-21 (Herlemann et al., 2011) to capture both Thaumarchaeota and low-abundant and under-studied groups typically present in soils (e.g. different classes of methanogens, Thermoplasmata; George et al., 2019; Jiao et al., 2019a; Liu et al., 2019). The PCRs were run in duplicate 15 μl reactions under the following conditions: 3 min at 98°C, followed by 30 cycles of 98°C for 30 s, 65°C for 30 s and 72°C for 30 s and a final extension step of 10 min at 72°C. The PCR products were then pooled and inspected by gel electrophoresis. For the second (indexing) PCR, a single 30 μl reaction was performed using 0.2 μM of primers with Nextera adaptors and index sequences, and 3 μl of the pooled PCR product from the first PCR as template. Conditions were the same as in the first PCR, except an annealing temperature of 55°C, an extension time of 45 s, and 8 cycles. The final PCR products were purified using AMPure XP PCR purification beads (Beckman Coulter, Indianapolis, IN, USA) following the manufacturer’s protocol. The amplicons were checked by gel electrophoresis and using a 2100 BioAnalyzer (Agilent, Santa Clara, CA, USA) to verify the correct size and that there were no additional amplicons. All samples passed the control. After quantification using a Qubit™ fluorometer (Thermo Fisher Scientific, Waltham, MA, USA), a single library was created by pooling equal amounts of purified amplicons from all the 151 samples. Sequencing was performed by SciLifeLab (Uppsala, Sweden) on the Illumina MiSeq (2 × 250 bp) platform.

Quantitative PCR analysis

The abundance of AOA was determined by quantitative real-time PCR (qPCR) based on SYBR green detection and the archaeal amoA gene (encoding the ammonia monoxygenase subunit A). The qPCR reactions were carried out in duplicate runs on a ViiA7 (Life Technologies, Carlsbad, CA, USA) and a 15 μl reaction volume containing 7.5 μl of Takyon Master Mix (Eurogentec, Liège, Belgium), 1 μM of each primer (CrenamoA23f and CrenamoA616r; Tourna et al., 2008), 250 ng of T4 gene DNA (QBiogene, Carlsbad, CA, USA) and 1 ng of DNA). Cycling conditions were 15 min at 95°C, 35 cycles of 15 s at 95°C, 30 s at 55°C, 30 s at 72°C and a plate read of 15 s at 80°C (efficiency: 88%). Standard curves were obtained by serial dilutions of linearized plasmids with cloned fragments of the specific gene. The amplifications were validated by melting curve analyses. Potential inhibition of PCR reactions was checked by amplifying a known amount of the pGEM-T plasmid (Promega, Madison, WI, USA) with the plasmid specific T7 and SP6 primers when added to the DNA extracts or non-template controls. No inhibition was detected with the amount of DNA used.

Sequence processing and phylogenetic reconstruction

All sequence analyses were performed using the R software (v. 3.6.4, R Core Team, 2019). The archaeal 16S rRNA gene amplicons were processed with the ‘dada2’ package (v. 1.6.0; Callahan et al., 2016) to infer amplicon sequence variants (ASVs), which allow detection of ecological preferences at the finest phylogenetic scale (Hunt et al., 2008; Larkin and Martiny, 2017). Briefly, primer sequences were removed and the reads merged using default parameters. Chimeras were discarded using a denovo approach with the removeBimeraDenovo function (‘consensus’ method). The resulting ASVs were aligned to the SILVA reference database (SSU132 Ref NR) using the SINA aligner (v. 1.6.0; Pruesse et al., 2012) and classified using SINA’s least common ancestor algorithm. After elimination of the bacterial ASVs, 2042 archaeal ASVs remained based on a total of 63 167 075 high-quality 16S rRNA gene amplicons (corresponding to approximately 70% of the original dataset). The reads were also clustered into OTUs at a similarity cut-off of 97% (Supplementary Methods) to compare the diversity of nitrosoarchaeal taxa based on ASVs and OTUs.

Rarefaction curves of species richness were generated from the raw ASV table using the rarecurve function in ‘vegan’ (v. 2.5.5; Oksanen et al., 2018) (Fig. S6) and a rarefied table (n = 1079 ASVs) was obtained by averaging the ASV counts over 1000 computations using the rarefy function in ‘vegan’. A phylogeny was built with the rarefied ASVs and a broad taxonomic selection of reference sequences extracted from SILVA. Sequences were aligned using the SINA aligner and the phylogeny generated using FastTreeMP (v. 2.1.10; Price et al., 2010) with the GTR + CAT model of nucleotide evolution.

Partitioning and transformation of sequence data

The ASV abundance distributions were examined to further partition the rarefied table between frequent and rare
community members. An index of dispersion corresponding to the ratio of the variance to the mean abundance multiplied by the occurrence was calculated (Hubbell, 2001) to split the dataset according to the frequency of occurrence of each ASV (Magurran, 2007). This index was then used to model whether ASVs followed a stochastic (Poisson) distribution and those falling below the 2.5% confidence limit of the $\chi^2$ distribution were discarded (Krebs, 1999) (Fig. S7). By focusing on the frequent community members, we minimized the risk of sampling artefacts that would bias the distribution of the ASVs and thus increased the likelihood to detect relevant ecological patterns. Such partitioning remains rarely used in microbial ecology (but see Fillol et al., 2016; Jeanbille et al., 2016; Liu et al., 2018), despite being more statistically robust than traditional approaches using arbitrary cut-offs of local and regional abundances (e.g. Mo et al., 2018). The resulting community included 718 ASVs representing 99.6% of the reads in the rarefied dataset. Zero count ASVs were replaced by an imputed value using the Bayesian-multiplicatives replacement method available in the ‘zCompositions’ package (v. 1.2.0; Martín-Fernández et al., 2015). An isometric log-ratio transformation (Egozcue et al., 2003), as implemented in the ‘phlr’ package (default parameters, v. 1.10.0; Silverman et al., 2017), was then applied to the zero replaced dataset using the phylogenetic tree of the ASVs as the sequential binary partition. The output of this transformation consists of a matrix of sites x nodes containing the balances calculated on the inner nodes of the phylogeny. Balances were computed for each node and depict relative changes in ASV abundance between two neighbouring clades relative to each other, while ignoring abundances in the remainder of the tree. This approach accounts for the compositional nature of amplicon datasets (Gloor et al., 2017) by inferring changes between phylogeny-based subcommunities (i.e. the two set of branches stemming from any given node) rather than changes of individual taxa (i.e. branch tips) (Morton et al., 2017; Silverman et al., 2017).

Statistical analyses

Alpha-diversity indices and taxonomic composition were calculated using the full rarefied table. The alpha diversity metric Faith’s phylogenetic diversity (PD), which measures the shared phylogenetic history among taxa occurring in a sample (Faith, 1992), was calculated at both domain and class levels using the phylogenetic tree of the ASVs and the ph function in the ‘picante’ package (v. 1.8; Kembel et al., 2010). Piérou’s evenness (Piérou, 1966) was computed with the diversity function in ‘vegan’.

Random forest (RF) based variable selection was performed on the entire set of variables (Table 1) using the ‘VSURF’ package (v. 1.1.0; Genuer et al., 2015) to identify the best predictors for PD and evenness of archaeal groups and the absolute abundance of AOA (response variables). Random forests represent an ensemble machine learning algorithm that is well suited to model non-linear relationships between predictors and response variables and can deal with non-normality and high collinearity among predictors (Breiman, 2001). Briefly, variables were first ranked according to a variable importance score, averaged across 50 RFs. The set of variables leading to the model with the smallest out-of-bag error, averaged across a nested collection of 25 RFs starting from one with only the most important variable, was selected. To account for the random nature of RFs, the algorithm was run with default parameters 100 times and only the variables selected in the second step in >95% of the runs were retained. Random forest analyses, as implemented in the ‘randomForest’ package (v. 4.6–14; Liaw and Wiener, 2002), were then used to study the relationship between the selected environmental variables and the response variables. A grid search was first conducted to find the optimal combination of tuning parameters (with ntree = 500): the number of variables to randomly sample as candidates at each split (mtry; range 1–10, step = 1), the minimal number of samples within the terminal nodes (node_size; range 2–10, step = 1) and the fraction of samples to train the model on (samp_size; 55%, 63.25% (default), 70% and 80%). The search was run 100 times and the combination of parameters corresponding to the best model fit (or lowest out-of-bag root-mean-square error) was selected (Table S3). The relationship between each environmental variable and PD, evenness and the absolute abundance of AOA was visualized using accumulated local effects plots (grid.size = 30) implemented in the ‘iml’ package (v. 0.9.0; Molnar et al., 2018). These plots show how the prediction of a response variable in a given RF model (PD, evenness or abundance of AOA) changes on average over the range of each individual environmental variable, while accounting for potential correlations among explanatory variables (Apley and Zhu, 2020).

All statistical analyses on $\beta$-diversity were conducted on the phlR-transformed (Silverman et al., 2017) ASV table. Differences in community composition and structure were visualized with a principal component analysis (PCA) using the rda function in ‘vegan’. Distance-decay curves were calculated as the linear regression relationship between geographical distance and Euclidean distance-based community similarity. The relative influence of climatic, edaphic and spatial factors on the patterns of $\beta$-diversity was estimated by variation partitioning analysis (VPA; varpart function in ‘vegan’). To this end,
cartesian coordinates of each sampling site were obtained from the GPS data (geoXY function, package ‘SoDA’ v. 1.0.6) and used to construct a matrix of distance-based Moran's eigenvectors maps (dbhMEM). The edaphic factors were selected following a procedure similar to Power et al. (2018). First, individual permutational multivariate analyses of variance (PERMANOVA) and Mantel tests were conducted between overall β-diversity and each variable using the adonis and mantel functions implemented in ‘vegan’ (number of permutations = 9999), respectively (Table S1). Collinearity among edaphic factors was assessed by pair-wise Spearman correlations; only the variable with the highest mantel statistic was retained in each collinear group (|r| > 0.7). Thereby, the selected edaphic factors were calcium, clay, C:N and C:P ratios, magnesium, moisture, pH, silt, SOC and total C. Finally, a forward selection step (p < 0.05) was applied to select the final set of variables before running the VPA (forward.sel function, package ‘adespatial’ v. 0.3.4). The significance of each component of the VPA was estimated by a permutation test (adonis function). Elevation, although significant, only improved the explanatory power by 0.4% and was not included in the final analysis. The VPA was also applied to the nitrososphaerai community data and a subset of the full phlir-transformed ASV table was generated by extracting the nodes corresponding to Nitrososphaeria. The abundance of the archaeal amoA gene was fitted onto the nitrososphaerai ordination using the envfit function in ‘vegan’ (permutations = 9999). Since the correlation was significant (p < 0.001), the abundance of amoA genes was included in the VPA as a biotic component (functional predictor), along edaphic, climatic and spatial factors.

Since Nitrososphaeria represent putative AOA (Alves et al., 2018), their ecological preferences were predicted at a finer taxonomic scale by recursive partitioning of the corresponding phlir-transformed ASV table with the entire set of metadata (Table 1). Multivariate regression trees (MRT) were computed using the ‘mvpart’ package (v. 1.6.2; De'ath, 2002), to predict the relationships between the set of variables (Table 1) and the community structure of Nitrososphaeria. This approach has the advantage to allow for the examination of the sequential effect of several environmental filters (contrary to Gubry-Rangin et al., 2011, 2015; Oton et al., 2016; Alves et al., 2018 who assessed effects of each variable individually) and delineate clusters of samples in which the variation in environmental conditions is minimized (Table S2). The selected tree represented the most parsimonious solution within one standard error above the minimal cross-validated relative error (Fig. S8; n = 10,000 trees), following (Breiman et al., 1984), and explained 42% of the variation. Indicator nodes were searched for in each partition (i.e. clusters of samples) and corresponded to balances that significantly differed from the mean across all partitions according to Tukey’s HSD (p < 0.01). The balances within each MRT cluster were plotted on the nitrososphaerai phylogeny with ggtree (v. 1.16.0; Yu et al., 2017), with blue and red branches indicating an increase and a decrease, respectively. Sequences of Group 1.1c thaumarchaeota were used to root the phylogeny.

Availability of data and materials

Sequencing data has been deposited at the European Nucleotide Archive (ENA) under the accession number PRJEB35080. The datasets and code generated during the current study are available in the Zenodo repository (http://doi.org/10.5281/zenodo.4095504). They include metadata, R code, amoA gene abundances and the phylogeny in newick format.

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Author contribution

A.S., S.H., M.G.A.v.d.H., F.T.M., L.P. and M.C.R. initiated the study and planned the field work. A.S., S.B., F.D., A.E., P.G.P., G.G., C.H., D.S.P., and S.R. contributed to data collection. A.S. performed the analyses and drafted the manuscript together with S.H. All authors commented on and approved the final manuscript.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

**Table S1.** Results of Mantel tests and PERMANOVAs between the archaeal communities and the set of environmental variables (**p < 0.001, *p < 0.01; *p < 0.05).
Table S2. Range of environmental variables in the MRT clusters presented in Fig. 4 (see Table 1 for units).
Table S3. Model parameters and fit (residual mean squared error—RMSE) of the random forest analyses.
Fig. S1. Relationship between environmental variables and the phylogenetic diversity of archaeal communities, based on random forest analyses. Predictor variables selected by VSURF were used to generate accumulated local effects plots, which show the relationship between the predictor (x-axis) and response variables in the model (y-axis) while accounting for potential correlations among predictor values. N.S. Non-significant variable. See Table 1 for units. Model parameters and fit are indicated in Table S3.
Fig. S2. Relationship between environmental variables and the evenness of archaeal communities, based on random forest analyses. Predictor variables selected by VSURF were used to generate accumulated local effects plots, which show the relationship between the predictor (x-axis) and response variables in the model (y-axis) while accounting for potential correlations among predictor values. N.S. Non-significant variable. See Table 1 for units. Model parameters and fit are indicated in Table S3.
Fig. S3. Relationship between environmental variables and the abundance of AOA (number of gene copies g$^{-1}$ dw soil), based on random forest analyses. (A) Abundances of AOA in the 151 samples. (B) Predictor variables selected by VSURF were used to generate accumulated local effects plots, which show the relationship between the predictor (x-axis) and response variables in the model (y-axis) while accounting for potential correlations among predictor values. N.S. Non-significant variable. See Table 1 for units. Model parameters and fit are indicated in Table S3.
Fig. S4. Factors driving the variation in nitrososphaerial community composition and structure across the European gradient according to variation partitioning analysis (VPA). The biotic component corresponds to the abundance of the archaeal amoA gene, which correlated to the nitrososphaerial community structure ($p < 0.001$). All fractions were significant ($p < 0.001$) and the variation explained is indicated (%). Values < 0 are not shown. The VPA was performed on the philr-transformed data corresponding to the Nitrososphaeria.
Fig. S5. Sampling sites across Europe.
Fig. S6. Rarefaction curves of archaeal communities.
Fig. S7. Occurrence of each ASV plotted against its dispersion index. The dotted line represents the 2.5% confidence limit of the $\chi^2$ distribution, with frequent ASVs depicted in orange and rare ASVs in blue.
Fig. S8. Relationship between the size of trees and the relative error (green) and the cross-validated relative error (blue) computed in the multivariate regression tree analysis. The best predictive tree is indicated by a red dot. The orange dot corresponds to the smallest tree one standard error above the best tree. The green vertical bars indicate the number of times that a given solution was selected as the best one during the cross-validation iterations.